ORIGINAL ARTICLE

The ciliogenic transcription factor Rfx3 is required for the formation of the thalamocortical tract by regulating the patterning of prethalamus and ventral telencephalon

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Abstract

Primary cilia are complex subcellular structures that play key roles during embryogenesis by controlling the cellular response to several signaling pathways. Defects in the function and/or structure of primary cilia underlie a large number of human syndromes collectively referred to as ciliopathies. Often, ciliopathies are associated with mental retardation (MR) and malformation of the corpus callosum. However, the possibility of defects in other forebrain axon tracts, which could contribute to the cognitive disorders of these patients, has not been explored. Here, we investigate the formation of the corticothalamic/thalamocortical tracts in mice mutant for Rfx3, which regulates the expression of many genes involved in ciliogenesis and cilia function. Using DiI axon tracing and immunohistochemistry experiments, we show that some Rfx3−/− corticothalamic axons abnormally migrate toward the pial surface of the ventral telencephalon (VT). Some thalamocortical axons (TCAs) also fail to leave the diencephalon or abnormally project toward the amygdala. Moreover, the Rfx3−/− VT displays heterotopias containing attractive guidance cues and expressing the guidance molecules Slit1 and Netrin1. Finally, the abnormal projection of TCAs toward the amygdala is also present in mice carrying a mutation in the Inpp5e gene, which is mutated in Joubert Syndrome and which controls cilia signaling and stability. The presence of identical thalamocortical malformations in two independent ciliary mutants indicates a novel role for primary cilia in the formation of the corticothalamic/thalamocortical tracts by establishing the correct cellular environment necessary for its development.
Introduction

Primary cilia are complex subcellular organelles protruding from the cell surface, which mediate several sensory functions and are involved in regulating the cellular response to several signaling pathways and in particular to Sonic hedgehog (Shh) signaling (1). Moreover, primary cilia have important roles in human diseases, and defects in primary cilia formation and/or function underlie several human syndromes commonly referred to as ciliopathies (2,3). These disorders are associated with a large variety of manifestations that often include neurological features such as MR, but much less is known about the specific signaling pathways and the pathogenesis at the cellular and tissue level that ultimately result in neurological disease phenotypes (4,5).

A common feature of ciliopathies is malformations of the corpus callosum (CC) which as the major forebrain commissure provides the interhemispheric exchange of information between the two cortical hemispheres. While the significance of callosal abnormalities for the MR pathogenesis of ciliopathy patients remains unclear, this finding raises the possibility that other prominent axon tracts of the forebrain are affected in ciliopathies. To explore this, we here investigate the formation of the corticothalamic/thalamocortical tracts in two ciliary mouse mutants. The thalamocortical tract connects the dorsal thalamus with the cerebral cortex thereby conveying sensory information from the external environment to the cortex. The corticothalamic tract in turn sends processed sensory information back to the thalamus thereby providing the feedforward and feedback mechanisms essential in this processing unit (6). The formation of the thalamocortical tract requires a complex navigation of thalamocortical axons (TCAs) through the prethalamus and ventral telencephalon (VT), and involves several guidepost cues along the thalamocortical path that control the navigation of thalamic axons. Two populations of pioneer neurons are located in the prethalamus and the VT projecting their axons into the thalamus and providing scaffolds for TCAs growing into the prethalamus and across the diencephalic/telencephalic boundary, respectively (7–10). In addition, cells from the lateral ganglionic eminence (LGE) migrate into the medial ganglionic eminence (MGE) to form a permissive corridor and guide TCAs through the otherwise non-permissive MGE (11). These corridor cells also mediate the sorting of TCAs according to their rostral/caudal origin in the thalamus by expressing the Slit1 and Nestin1 guidance factors (12). Interestingly, the development of the VT and of the prethalamus, which provide these guidepost cues, is under the control of Shh signaling, raising the possibility that the formation of the thalamocortical tract might be affected in primary cilia mutants. However, this has not been explored yet.

RFX transcription factors have been shown to play fundamental roles in ciliogenesis by regulating the expression of genes involved in cilia assembly or function (13–15). Accordingly, Rfx3-deficient mouse mutants exhibit several hallmarks of ciliopathies, in particular left–right asymmetry defects and hydrocephalus (13,14), yet importantly they survive until birth, providing a rare opportunity to study the formation of forebrain axon connections in a cilia mouse mutant. Indeed, the Rfx3 mutation interferes with the formation of the CC (16). Here, we investigate the formation of the corticothalamic/thalamocortical tracts in Rfx3-deficient mice. In these animals, some corticothalamic axons (CTAs) abnormally migrate toward the amygdala. Moreover, only a small proportion of TCAs reach the dorsal telencephalon but many fail to migrate through the prethalamus, whereas others enter the VT but subsequently project ventrally toward the amygdala. These defects correlate with abnormal patterning of the prethalamus. In addition, the rostroventral telencephalon forms neural heterotopias at its pial surface containing a mixture of MGE- and LGE-derived cells expressing the Slit1 and Nestin1 guidance molecules. Finally, the abnormal ventral deflection of TCAs toward the amygdala is also found in a second ciliary mouse mutant, which is deficient for inpp5e, a Joubert syndrome disease gene. Taken together, these analyses identify a novel role for primary cilia in the development of thalamocortical tract.

Results

Rfx3−/− mutant embryos display defects in the development of thalamocortical connections

Recently, we showed that the ciliogenic transcription factor Rfx3 is required for CC development (16). Here, we investigate whether an Rfx3 null mutation also affects the formation of the thalamocortical/corticothalamic tracts. TCAs progress in a multistep fashion over their intermediate territories, the diencephalon and the VT (17). Thalamic neurons send their axons ventrally through the prethalamus toward the diencephalic–telencephalic boundary. After entering the telencephalon at E13.5, thalamic axons migrate through the VT via the internal capsule (ic) to reach the pallial–subpallial boundary (PSPB) and finally enter the cortical intermediate zone at E14.5. CTAs leave the cortex around E14.5 and follow the same trajectory through the VT as TCAs but in the opposite direction. This trajectory is revealed in E18.5 control embryos by placement of DiI crystals into the cortex. This analysis also reveals the path of TCAs by retrograde labeling (Fig.1A–D). Rfx3−/− mutant brains show a similar labeling pattern; however, we noted a number of backlabeled nuclei in the rostroventral telencephalon close to a region of the VT containing a heterotopia. This analysis also revealed an abnormal ventral projection toward the amygdala at intermediate and caudal levels (Fig.1E–H). To more specifically label TCAs, we placed DiI crystals into the thalamus. TCAs are highly fasciculated in the ic, are arranged in roughly parallel bundles in the striatum and reach the cortex (Fig.1I and J). In contrast, in Rfx3−/− mutant brains, thalamic axons appear disorganized in the striatum with some axons projecting toward the PSPB in more ventral regions (Fig.1M and N). At caudal levels, TCAs abnormally project into the VT toward the amygdala similar to our findings on the CTA trajectory.

Immunostaining for the panaxonal marker Neurofilament (NF) confirmed the pathfinding defects revealed by DiI labeling. In control embryos, this analysis showed parallel organized axons in the thalamus, prethalamus, ic, the striatum and in the intermediate zone of the cerebral cortex (Fig. 1K and L). In Rfx3−/− mutants, NF staining revealed several abnormal fiber tracts. Many axons are already disorganized in the thalamus, and the exit and entry zones of CTAs and TCAs into and out of the cortex appear broader (Fig.1O, arrowheads). Many NF+ fibers ectopically project toward the amygdala forming a highly fasciculated axon bundle at more caudal levels (Fig.1P, arrow).

To gain insights into the origin of these axonal defects, we performed NF immunofluorescence analyses and DiI labeling on E14.5 control and Rfx3−/− mutant brains. At this stage, NF labels axons in the developing thalamus, prethalamus and VT of control embryos (Supplementary Material, Fig. S1A and B). The ic contains highly fasciculated NF+ axons, whereas axons are more loosely organized in the striatum (Supplementary Material, Fig. S1C and D). In Rfx3−/− embryos, NF+ axons are disorganized in the VT projecting in several directions within the LGE and MGE.
Figure 1. Axon guidance defects in the E18.5 Rfx3−/− brains. Coronal sections through the brain of control (A–C and I–L) and Rfx3−/− (E–G and M–P) E18.5 embryos. (A–C and E–G) DiI placements in the cortex (asterisks) reveal the anterior commissure (ac) (A and E) and the corticothalamic and thalamocortical tract through anterograde and retrograde labeling, respectively. In Rfx3−/− mutants, both tracts are formed but note the backlabeled neurons close to a heterotopia at the ventral telencephalic surface (inset and white asterisks in E) and the abnormal projections toward the amygdala at intermediate and caudal levels (arrowheads in F and G). (I, J, M and N) Thalamic DiI labeling (asterisks) revealed the TCA trajectory in control embryos (I and J). In Rfx3−/− mutants, TCAs cross the FSP98 in a broader entry zone (arrowheads in M and N). Several axons leave the ic prematurely and form an ectopic axon bundle running ventrally toward the pial surface (arrows in M and N). (K, L, O and P) Immunostainings for NF showed axonal pathfinding defects in the diencephalon (arrowheads in O) and an ectopic axon bundle running ventrally toward the amygdala of Rfx3−/− embryos (arrows in O and P). (D and H) Schematic drawings summarizing the CTA and TCA pathfinding defects in Rfx3−/− mutants. Scale bar: 100 µm.
and a thick axon bundle runs ectopically toward the pial surface (Supplementary Material, Fig. S1F and G, arrow), whereas caudally no axons reached the PSPB (Supplementary Material, Fig. S1H and I). In addition, several NF+ axons mis-project in the prethalamus resulting in the formation of abnormal bundles (Supplementary Material, Fig. S1I). These findings were confirmed by Dil labeling experiments of TCAs. In contrast to control embryos, very few TCAs traverse the VT with no axons reaching the PSPB in Rfx3−/− mutants (Supplementary Material, Fig. S1K, L and Q), but a large fiber bundle abnormally projects ventrally toward the telencephalic pia (Supplementary Material, Fig. S1R). To further dissect the path of thalamic axons, we performed calretinin (CR) immunostaining and DiI labeling experiments, many CR+ axons form abnormal clusters in the thalamus. Finally, placing Dil crystals into the cortex revealed CTAs in control embryos having penetrated into the VT and retrogradely labeled neurons in the thalamus (Supplementary Material, Fig. S1M and N). In Rfx3−/− mutants, backlabeled neurons were not observed, instead a large axon bundle projects from the cortex toward the pia of the VT (Supplementary Material, Fig. S1S and T). Collectively, these analyses indicate severe disruptions of these genes as shown by qRT-PCR (Fig. 2B, C, E, F, H and I). Next, we investigated prethalamic patterning. In E14.5 control embryos, the prethalamic progenitor domain is characterized by Nkx2.2 expression (25) (Fig. 3A and B). This expression domain is expanded in Rfx3−/− mutants (Fig. 3C and D). Olig2, another marker for diencephalic progenitor cells, displays a more complex expression pattern. Immunofluorescence analysis showed a high-level Olig2 expression domain in prethalamic progenitors, and lower-level expression in thalamic progenitors, with a high-center to low-dorsal expression gradient (Fig. 3F, I). In the Rfx3−/− mutant diencephalon, the dorsal thalamic expression of Olig2 and its expression gradient is maintained (Fig. 3G, H, K and L). However, Olig2 expression is expanded in the prethalamus of Rfx3−/− mutants (Fig. 3H and L). Finally, the expression domain of the Pax6 transcription factor normally extends ventrally from the zli encompassing progenitors from the prethalamus and from the eminentia thalami, thereby delineating the boundary between the thalamus and prethalamus (Fig. 3E, F, I and J). In the Rfx3−/− mutants, Pax6 expression is maintained in prethalamic progenitors, but its expression in the diencephalic mantle is reduced (Fig. 3G, H, K and L). Taken together, these results suggest regionalization defects in the prethalamus of the Rfx3−/− mutants.

Abnormal MGE development and formation of heterotopias in the Rfx3−/− VT

Since Rfx3−/− mutants show several corticothalamic/thalamocortical pathfinding defects in the VT, we next investigated its development. Recently, we reported a slight down-regulation of Shh signaling in the MGE of Rfx3−/− mutants using in situ hybridization (16), but qRT-PCR showed no effect on Shh and Gli1 expression, whereas Ptc1 expression was mildly up-regulated (P = 0.0384; n = 6; Supplementary Material, Fig. S4). Correct levels of Shh signaling are crucial for establishing the PSPB separating the dorsal telencephalon and VT. However, expression analyses for Dlx2, Gsh2, Pax6 and Dbx1 did not reveal abnormalities at this boundary (Fig. 4A and F, and Supplementary Material, Fig. S5). We next analyzed the subdivision of the VT into the MGE and LGE. Dlx2 is expressed throughout the proliferative zone of the VT, whereas Nkx2.1 expression is restricted to the MGE (Fig. 4A and B). Moreover, Nkx6.2 expression is confined to progenitors on either side of the interganglionic sulcus (Fig. 4C). In Rfx3−/− embryos, these expression patterns are maintained except for a slight Nkx6.2 down-regulation (Fig. 4E–G). We also noted an ectopic accumulation of Dlx2 and Nkx2.1 expressing cells outside the VT at its ventral surface (Fig. 4E–G) reminiscent of the heterotopia we observed at E18.5 (Fig. 1E). These heterotopias were detected on both sides of the rostral telencephalon in all Rfx3 mutants but not in more caudal regions. Also, they varied in size with some extending toward the midline, whereas others were confined to the surface of the ventrolateral telencephalon. In addition, the heterotopias contain a mixture of progenitor cells and neurons. Expression of the MGE-specific genes Lhx6 and Lhx7, which regulate Shh expression in the MGE and which are in turn Shh target genes (26), was detected in the heterotopias (Fig. 4H and M) as well as that of Tuj1 characteristic of differentiating neurons (Fig. 4N). Tuj1 immunofluorescence also showed a size reduction of the MGE mantle. Finally, we investigated whether disruption of the basal lamina, which normally surrounds the telencephalon, could account for heterotopia formation. Laminin immunofluorescence showed that the basal lamina is disrupted in several positions at the ventral pial surface of E12.5 and E14.5 Rfx3−/− mutants (Fig. 4O and P). Taken together, these analyses show a size reduction of the MGE mantle and a disruption of the basal lamina.
lamina concomitant with the formation of neural heterotopias in Rfx3−/− embryos.

Cellular and molecular guidance cues are affected in the VT of Rfx3−/− embryos

Given these defects in the formation of the VT, we next analyzed whether ventral telencephalic guidance cues essential for TCA projection are also affected. Pioneer neurons located in the rostral MGE extend their axons into the thalamus and serve as scaffolds to guide TCAs across the diencephalic–telencephalic boundary (7–9). Since no markers are available to selectively label these MGE pioneer neurons, we investigated their formation by DiI placement into the E12.5 thalamus. In control embryos, these placements retrogradely labeled the pioneer axons and their cell bodies located in the rostral MGE (Fig. 5A and B).

While this analysis also revealed pioneer neurons cell bodies and axons in the MGE mantle of Rfx3−/− mutants, their distribution appeared less dense (Fig. 5C and D). In addition, some DiI-labeled cell bodies were located in the heterotopic tissue outside the MGE mantle.

In addition to the MGE pioneer neurons, neurons derived from the LGE migrate into the MGE and form a corridor along which TCAs project through the otherwise non-permissive environment of the MGE. In control E12.5 and E14.5 brains, these guidepost cells are marked by both Ebf1 and Islet1/2 expression from their origin in the LGE and during their migration into the MGE, where they form the ic (Fig. 6A–C). In E12.5 and E14.5 Rfx3−/− embryos, Ebf1-expressing cells originate normally in the LGE and migrate toward the MGE forming the corridor at caudal levels (Fig. 6F). However, at rostral levels, Ebf1-expressing cells migrate toward the heterotopias thereby abnormally connecting the

Figure 2. Shh signaling in Rfx3−/− mutant diencephalon. Coronal sections through the diencephalon of E12.5 control (A–C) and Rfx3−/− mutant embryos (D–F) hybridized with the indicated probes. (A and D) No major differences are observed in Shh expression in the Rfx3−/− zli compared with control embryos. Note the enlargement of the third ventricle in mutant embryos (asterisks in D, E and F). (B and C) Ptc1 and Gli1 are expressed in the diencephalon of control brains, but not in the zli. (E and F) Expanded expression of both Ptc1 and Gli1 is found in the prethalamus of Rfx3−/− mutants (arrowheads in E and F). (G–I) Quantification of Shh (G), Ptc1 (H) and Gli1 (I) expression in the diencephalon using qRT-PCR revealed no significant changes in the expression levels of these markers (n = 6). Scale bar: 100 µm.

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surface of the MGE mantle with the LGE (Fig. 6D and E). These abnormalities were confirmed by immunostainings for Isl1/2. As in control embryos, Isl1/2+ cells normally populate the corridor (Fig. 6G, H, J and K), but were also found ectopically in the heterotopic tissue in Rfx3−/− embryos (Fig. 6J and K). Finally, Nkx2.1 expression labels the MGE ventricular zone and the globus pallidus of E14.5 control embryos, whereas the ic remains Nkx2.1 negative (Fig. 6I). The Nkx2.1 expression pattern in the Rfx3−/− MGE mantle appears unaltered in Rfx3 mutants but as with the corridor markers, Nkx2.1-expressing cells were also detected in the ectopic tissue (Fig. 6I). These findings suggest that the MGE corridor forms at caudal levels, but that corridor cells abnormally migrate toward the heterotopias at rostral levels.

Next, we investigated whether the defects in forebrain patterning and in the formation of cellular guidance cues correlate with altered expression patterns of axon guidance molecules. Interestingly, Sema6a−/− mice show a very similar thalamocortical pathfinding defect as Rfx3−/− mutants with TCAs being deflected toward the amygdala after entering the VT (27). Sema6a has a highly complex expression pattern in the developing forebrain. Rostrally, Sema6a expression was mainly confined to the ventricular zone of the VT, but weak expression was also found in the heterotopias of Rfx3−/− embryos (Fig. 7A and E). Caudally, Sema6a transcripts were detected in thalamic neurons and in groups of cells along the path of TCAs in the prethalamus and dorsally and ventrally to the permissive corridor. This expression pattern is maintained in Rfx3−/− embryos with no Sema6a expression in the heterotopias (Fig. 7I, J, M and N).

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Figure 3. Patterning defects in the prethalamus of Rfx3−/− mutant brains. (A–L) Coronal sections through the diencephalon of E14.5 control (A, B, E, F, I and J) and mutant (C, D, G, H, K and L) brains were stained with the indicated antibodies. B, D, F, H, J and L are higher magnifications of A, E, G, I and K, respectively. (A and B) Nkx2.2 immunostaining labels the prethalamic progenitor domain. (C and D) Nkx2.2 expression was expanded in the diencephalon of Rfx3−/− mutants (arrowhead in D). (E, F, I and J) Pax6 is highly expressed in the diencephalon of control brains at the boundary between prethalamus and dorsal thalamus. (G, H, K and L) In the Rfx3−/− diencephalic ventricular zone, the Pax6 expression domain in the diencephalic mantle is reduced (arrows in H and L). Scale bars: 100 µm.
expressed in ventral telencephalic progenitor cells, and Netrin1 transcripts were also found in the MGE mantle (Fig. 7K and L). These expression patterns remain unaltered in Rfx3−/− embryos, but both genes are ectopically expressed in cells located in the heterotopias (Fig. 7O and P). Taken together, these expression analyses did not reveal changes in the overall Sema5b, Sema6a...
and PlexinA4 expression patterns, but show differences in the organization of TCAs with respect to the expression domains of these guidance molecules and also indicate ectopic expression of axon guidance factors in the heterotopias.

The Inpp5e ciliary mouse mutant phenocopies the TCA pathfinding defects of Rfx3−/− mutants

The Rfx3 transcription factor has a prominent role in ciliogenesis by regulating the expression of many genes involved in cilia assembly and function, suggesting that the thalamocortical abnormalities of Rfx3−/− embryos are likely caused by structural and/or functional ciliary defects. However, scanning electron microscopy revealed that cilia are present in the diencephalon and telencephalon of E10.5 Rfx3−/− embryos without obvious morphological abnormalities (Supplementary Material, Fig. S6 and data not shown). To gain support for the involvement of cilia in the pathogenesis of these Rfx3−/− phenotypes, we characterized thalamocortical development in an additional ciliary mouse mutant that carries a mutation in the Inpp5e gene (29). Inpp5e encodes inositol polyphosphate-5-phosphatase E that hydrolyzes the 5-phosphate of the second messengers PI(4,5)P2 and PI(3,4,5)P3. The Inpp5e protein is localized at the axoneme of the primary cilium and has a role in cilia-mediated signaling and in regulating cilia stability (29,30). Moreover, its human homolog INPP5E is mutated in MORM and in Joubert syndrome (29,30). Inpp5e mouse mutants show defects in limb and kidney development typical for cilia dysfunction (29), but thalamocortical development has not been investigated yet. Similar to Rfx3−/− embryos, labeling of the thalamocortical tract in E18.5 Inpp5e−/− embryos by inserting DiI crystals into the thalamus revealed an abnormal broad trajectory of TCAs into the rostral MGE and project their axons caudally toward the diencephalon. Scale bar: 200 µm.

Figure 5. Abnormal formation of MGE pioneer neurons in Rfx3−/− mutants. (A) Dil placements into the E12.5 thalamus of control embryos retrogradely labeled cell bodies located in the MGE and their axons. (C) Dil placed in the E12.5 Rfx3−/− thalamus revealed pioneer neuron cell bodies and axons in the MGE mantle and in ectopic locations in the heterotopias (arrows in C label the cell bodies and the asterisk the heterotopia). (B and D) Schematic representation of a series of coronal sections showing the location and axonal projections of MGE pioneer neurons. Note that these neurons reside in the rostral MGE and project their axons caudally toward the diencephalon. Scale bar: 200 µm.
Tbr1 and Ctip2, characteristic of corticofugal projection neurons, also appeared normal in Inpp5e−/− embryos (Supplementary Material, Fig. S9). In contrast, the MGE corridor was broader and the groups of Sema6a- and PlexinA4-expressing cells delineating the corridor were not discernible in mutant embryos (Supplementary Material, Figs S10 and S11), suggesting that defective ventral telencephalic development underlies the TCA pathfinding defects. Moreover, the similar ventral deflection of TCAs in Rfx3−/− and Inpp5e−/− embryos strongly implicates cilia dysfunction as the major cause of this phenotype in Rfx3−/− mutants.
Figure 7. The Rfx3−/− heterotopias express thalamocortical guidance molecules. Coronal sections through the brain of control (A–D, I–L) and Rfx3−/− mutants (E–H, M–P) hybridized with the indicated probes. (A, B, E and F) At rostral levels, Sema6a shows expression in the ventral telencephalic ventricular zone and weak expression in the heterotopia of Rfx3−/− embryos (arrow). Caudally, Sema6a is expressed along the path of TCAs in the prethalamus and VT, and its expression is not affected in Rfx3−/− mutants. (C) PlexinA4 is expressed in the thalamus and delineates the corridor ventrally and dorsally (arrows). (D) PlexinA4-expressing cells also reside in the caudal VT where TCAs normally do not project. (G and H) In Rfx3−/− embryos, PlexinA4 expression is largely maintained in the VT and thalamus; however, in the caudal VT, it is expressed adjacent to an ectopic axon bundle projecting toward the amygdala (arrow). (I, J, M and N) Sema5b expression in telencephalic progenitors and in neurons of the piriform cortex (pc) is maintained in Rfx3−/− embryos. (K, L, O and P) Netrin1 and Slit1 are expressed in ventral telencephalic progenitor cells, and Netrin1 transcripts were also found in the MGE mantle in both control and Rfx3−/− embryos, but both genes are ectopically expressed in the heterotopias (arrows in O and P). Scale bar: 100 µm.
The Gli3 repressor does not rescue TCA pathfinding and heterotopia formation in Rfx3−/− mutants

Finally, we addressed the cilia-regulated signaling pathway(ies) which could underlie the TCA and heterotopia phenotype in Rfx3 mutants. During forebrain development, primary cilia are essential for the processing of the Gli3 protein thereby controlling the balance between Gli3 repressor (Gli3R) and activator forms (Gli3A). Recently, we showed that this balance is disturbed in the Rfx3−/− forebrain (16). Rfx3 mutants also phenocopy the thalamocortical abnormalities of the Gli3 hypomorphic mutant Gli35tm (10). In addition, the olfactory bulb phenotype of the Ftm cilia mutant is rescued in Ftm−/−;Gli35.699/+ embryos (31) in which the Gli35.699 allele exclusively produces a short Gli3 isoform that resembles Gli3R (32) in a cilia-independent manner. These findings raised the possibility that the TCA pathfinding defects in Rfx3−/− embryos are caused by disrupting the Gli3R/Gli3A ratio, and we therefore analyzed thalamocortical tract and heterotopia formation in Rfx3−/−;Gli35.699/+ mutants. However, Rfx3−/−;Gli35.699/+ mutants still showed an abnormal projection of TCAs toward the amygdala and formed heterotopias in the ventrostral telencephalon (Supplementary Material, Fig. S12). Therefore, re-introduction of Gli3R is not sufficient to rescue these defects unlike the previously described rescue of the Ftm olfactory bulb phenotype (31). This finding could be explained by a requirement for Gli activator and not Gli repressor function in VT patterning and indeed, Gli3 western blots showed that the levels of Gli3R are not altered in the VT of Rfx3−/− embryos (Supplementary Material, Fig. S13).

Discussion

Ciliopathies, syndromes caused by dysfunction of the primary cilium, are often associated with MR; however, the underlying pathogenesis remains largely unknown. Here, we investigated the development of the forebrain in mice mutant for the ciliogenic transcription factor Rfx3. We show that Rfx3 mutants display the formation of heterotopias in the VT and patterning defects in the prethalamus. Moreover, the development of guidepost neurons, which are provided by the VT and which guide TCAs from the thalamus into the cortex, is affected. In fact, many TCAs are unable to leave the thalamus or mis-project toward the amygdala after entry into the VT. As a similar ventral deflection of TCAs into the VT is also found in Inpp5e mutants, these findings provide a novel role for primary cilia in the development of the thalamocortical tract.

Rfx3 is required for patterning the VT and the diencephalon

Recent analyses have shown a prominent role of primary cilia in forebrain development, but only few cilia mouse mutants with forebrain phenotypes have been described yet. These analyses were also limited by severe patterning defects (33,34) or did not analyze diencephalic development (31). Therefore, roles of primary cilia in the VT and in the diencephalon remain largely unknown or have not been analyzed at all. Rfx3 mutant mice provide a useful tool for studying such functions. In contrast to previously analyzed cilia mutants, these animals show a relatively mild alteration in the Gli3 activator/Gli3 repressor ratio in the forebrain (16), an important determinant of Shh controlled patterning processes.

Interestingly, Rfx3−/− embryos show multiple and complex abnormalities in the VT. First, the size of the Rfx3−/− MGE mantle is reduced. Secondly, Rfx3−/− embryos form heterotopias in which the basal lamina is disrupted and telencephalic cells are located outside the mantle zone. These heterotopias are a complex mixture of different cell types including progenitor cells (Dlx2+ and Dlx2+).
Nkx2.1+ and MGE- (Lhx6+ and Lhx7+) and LGE- (Ebf1+ and Isl1/2+) derived neurons. The heterotopias also express Shh (16) and several genes encoding axon guidance molecules (Fig. 7). The molecular mechanisms underlying their formation remain unknown, but could relate to altered Shh signaling which could lead to an early overproduction of MGE neurons. While qRT-PCR analysis using the whole VT revealed a mild up-regulation for Ptc1 but not for Gli1 expression, in situ hybridization indicated reduced expression of the Shh target genes Ptc1, Gli1 and Nkx6.2 in the ventricular zone of the VT. This discrepant finding could be attributed to the Shh expression in the heterotopias (16) and a concomitant up-regulation of Shh signaling in this ectopic tissue. Alternatively, the heterotopias could be formed by an over-migration of ventral telencephalic neurons (35), which could also contribute to the size reduction of the MGE mantle region, with neurons settling outside the MGE mantle.

The Rfx3 mutation also affects the patterning of the prethalamus as indicated by an expansion of Nkx2.2 and Olig2 expression and a concomitant reduction of Pax6 expression. These alterations in marker gene expression are consistent with known roles of Shh signaling. Prethalamic Nkx2.2 expression is reduced in the diencephalon of mouse embryos in which Shh signaling is inhibited (25), whereas Nkx2.2 and Olig2 expression is expanded after increasing Shh signaling in the diencephalon (25,36). In addition, tissue transplant studies in chick embryos show that ectopic expression of Shh negatively controls diencephalic Pax6 expression (37). Similarly, the expression of the Shh target genes Gli1 and Ptc1 is expanded in the prethalamus; however, our qRT-PCR analysis could not detect significant changes in the expression levels of these markers in the diencephalon. Due to the comparatively small size of the prethalamic progenitor domain within the diencephalon, it is possible that changes in Gli1 and Ptc1 expression in the prethalamus are too small to be detected within the whole diencephalic tissue. Alternatively, alterations in other signaling pathways could be responsible for the prethalamic defects. Wnt/β-catenin plays well-characterized roles in establishing the zli (38) and in specifying progenitor identity in the thalamus (39), whereas Fgf signaling controls the development of GABAergic neurons in the prethalamus and rostral thalamus (40). However, roles for either pathway in prethalamic patterning have to be identified yet. In contrast, Rfx3−/− embryos showed no obvious defect in thalamic development. The specification of the thalamic nuclei appears to be delayed compared with the differentiation of prethalamic neurons and is concomitant with a down-regulation of Shh signaling, suggesting that molecules others than Shh play a more prominent role in specifying thalamic nuclei (41). Regardless of the exact mechanism, our data clearly indicate novel roles for primary cilia in the development of the VT and in the patterning of the prethalamus.

Corticothalamic/thalamocortical tract defects in Rfx3−/− mutants

Our analyses also revealed pathfinding defects of thalamicortical and CTAs in Rfx3−/− mutants occurring subsequently to pathfinding defects. Some thalamic axons reach the cortex whereas others are either not able to exit the dorsal thalamus, or after having entered the VT mis-project toward the amygdala. Since thalamic patterning, production of post-mitotic neurons and initial growth of CTAs are not obviously affected, cell-autonomous defects in thalamic neurons are unlikely to underlie these pathfinding defects, but this requires further testing in transplantation experiments. In contrast, development of the prethalamus and the VT is compromised in Rfx3−/− mutants. Both structures give rise to multiple neuronal cell types playing important roles in the guidance of thalamocortical/CTAs (17). Indeed, we show here an abnormal migration of corridor cells and MGE pioneer neurons into the ventral telencephalic heterotopias of Rfx3−/− mutants. Similarly, mis-patterning of the prethalamus might have affected the development of prethalamic pioneer neurons, which normally project their axons into the thalamus thereby providing a scaffold for CTAs. Given the importance of pioneer neurons in the development of forebrain connections (10,42-46), defects in the prethalamic pioneer neurons might be responsible for the clustering of CTAs in the diencephalon; however, we currently lack the molecular markers to test this hypothesis.

Moreover, our analysis provides first insights into the cellular and molecular mechanisms by which Rfx3 controls thalamocortical tract formation. The Rfx transcription factors regulate the transcription of genes important for cilogenesis and function, suggesting that ciliary defects may underlie the abnormal TCA pathfinding in Rfx3−/− embryos. While cilia are present without obvious morphological abnormalities in the Rfx3 mutant forebrain, their function might be affected as in J688 hypomorphic mutants which have severe telencephalic pathfinding defects (34). Taken together with our finding that inactivating two genes with essential but different roles in cilia, namely Rfx3 or Inpp5e, which encode a transcription factor controlling cilia assembly or an enzyme regulating cilia signaling and stability, respectively, result in an highly similar TCA phenotype suggests that cilia dysfunction underlies this TCA pathfinding defect. The ectopic projection of CTAs toward the amygdala in Rfx3−/− and Inpp5e−/− embryos also phenocopies the TCA pathfinding errors in the Gli3 hypomorphic mutant Gli3Pdn/Pdn (10), suggesting altered Gli3 processing as the key cilia controlled pathway responsible for this phenotype. However, while the Gli3R/Gli3FL ratio is altered in the whole forebrain (16), Gli3 western blotting did not reveal significant changes in this ratio in the VT of Rfx3−/− embryos, similar to our previous finding in Gli3Pdn/Pdn−/− mutants (10). Moreover, re-introducing a Gli3R allele in an Rfx3−/− mutant background does not ameliorate the TCA abnormalities, unlike the rescue of olfactory bulb formation in Ftm−/− embryos (31) or of the callosal defects in Rfx3−/− embryos (B. Durand, manuscript submitted). These findings rule out the involvement of altered Gli3 to Gli3R processing. Alternatively, these data suggest that there could be different requirements for Gli3 repressor and activator functions in the dorsal telencephalon and VT. It will therefore be interesting to aim at rescuing the Rfx3 phenotype using a Gli3 mutant that predominantly forms Gli3A (47).

Finally, the formation of neural heteropias might also affect CTA pathfinding in the VT. The ventral TCA defect and heterotopia formation occur at different rostral/caudal levels. In contrast, CTAs are deflected at rostral levels where heterotopias are present, and this defect is absent in Inpp5e−/− and Gli3Pdn/Pdn−/− embryos which lack heterotopias. Moreover, the heterotopias contain Ebf1+ and Isl1/2+ neurons. These cells normally migrate from the LGE in the MGE to form a corridor permissive for CTAs and CTAs, but are abnormally scattered throughout the MGE mantle and even form a stream of cells connecting the LGE with the heterotopias. These also contain cells expressing Nettin1 and Slit1, which individually act as chemotactant or repellent of CTAs, respectively, but their combined action is required for the rostral/caudal sorting of CTAs (12). Therefore, the heterotopias could provide alternative migration routes for CTAs toward the thalamus (27).
Conclusion

Our study highlights essential roles of Rfx3 for the correct development of the VT and the prethalamus and emphasizes how subtle defects in these tissues can lead to severe pathfinding defects in the thalamocortical tract. These findings are highly relevant to human ciliopathies, which are often associated with MR. The thalamocortical tract as a major forebrain axon tract conveys most of the sensory information from the environment to the cerebral cortex and its disruption is predicted to interfere with the normal functioning of the cortex. Indeed, development of the thalamus and cortex is tightly linked and abnormalities in the thalamocortical pathways correlate with sensory and motor deficits in preterm born children (48,49). In addition, Bardet-Biedl syndrome patients have recently been reported with a reduced thalamic size (50,51). Moreover, we show here that an identical TCA pathfinding defect is present in mice mutant for the Inpp5e gene, whose human homolog is mutated in Joubert Syndrome (29,30). It will therefore be interesting to analyze thalamocortical tract formation in cilopathy patients which could lead to a new understanding of the pathogenesis of MR in these patients.

Materials and Methods

Mice

All animal research has been conducted according to relevant national and international guidelines. Rfx3−/− deficient embryos were generated and genotyped as previously described (14). Gli3+/− and Inpp5e null mutant mice used in this work have previously been described (29,32). For each marker and each stage, 3–6 mutant embryos were analyzed and compared with 3–6 controls. All reported phenotypes were fully penetrant.

In situ hybridization and immunohistochemistry

Antisense RNA probes for Dlx1 (52); Dlx2 (53); Ebf1 (11); Gbx2 (54); Gli1 (55); Ngn2 (56); Nkx2.1 (57); Lhx2 (EST2101448); Lhx6 and Lhx7 (58); Nkx6.2 (59); Ptcr (60) and Shh (61) were labeled with digoxigenin. In situ hybridization on 10 µm serial paraffin sections of mouse brains were performed as described (62).

Immunohistochemical analysis was performed as described previously (62) using antibodies against the following molecules: β-III-tubulin (TuJ1 antibody; 1 : 1000, Sigma); CR (1 : 1000, CHEMICON); Gsh2 (1 : 2500, a gift from K. Campbell, Cincinnati Children’s Hospital Medical Center, OH); Isl1/2 (1 : 100; DSHB); laminin (1 : 100, Sigma); NF (1 : 5; DSHB); Nkx2.2 (1 : 50, DSHB); Olig2 (1 : 1000, Millipore) and Pax6 (1 : 200; DSHB). Primary antibodies for immunohistochemistry were detected with Alexa- or Cy2/3-conjugated fluorescent secondary antibodies, and sections were incubated the nuclear counterstain TOPRO3 (0.2 µM) overnight at 4°C.

Scanning electron microscopy

E10.5 embryos were dissected in PBS at 400 mOsm. Brains were removed, cut into small fragments and fixed overnight at 4°C in PBS (200 mOsm)/2% glutaraldehyde. Brain samples were then washed several times in PBS (400 mOsm) and postfixed for 15 min in PBS (400 mOsm)/1% OsO4 (Euromedex, France). Fixed brain samples were washed extensively with distilled water and dehydrated in a graded series of ethanol solutions and finally in acetone. Brain samples were then prepared for scanning electron microscopy by the critical point freeze–dry procedure (Balzers–Union, CPD020). Samples were surface-coated using a gold–palladium spattering device (Hummer 2, Technics) under optimal conditions for 1 min 30 s, and observed with a scanning electron microscope (S800, Philips) at 15 keV. Observations were performed at the Centre for Microstructure Analysis of the University of Lyon.

Quantitative RT-PCR

Total RNA was extracted from the dorsal telencephalon, the VT and the diencephalon of six wild-type and six Rfx3−/− E12.5 embryos using the Nucleospin RNA XS kit (Macherey Nagel). cDNAs were synthesized using 0.8 µg of total RNA, 200 ng of random primers (Promega) and 200 units of RevertAid H Minus M-MuLV Reverse Transcriptase (Fermentas) according to the manufacturer’s instructions in a final volume of 20 µl. Real-time PCR was performed as previously described (63) on 2 µl of cDNA diluted one-fifth using the SYBR Green fluorescent mix (Roche) in a LC480 LightCycler (Roche). Primer sequences are available upon request. According to melting point analysis, only one PCR product was amplified. RNA extracted from heterozygous samples was used to generate a standard quantification curve for each gene, allowing the calculation of relative amounts of transcripts in the samples. The expression of each gene was normalized using the housekeeping gene Tbp. Statistical analysis was performed with the non-parametric Mann–Whitney test using the GraphPad Prism software.

Western blotting

Protein was extracted from the VT of E13.5 wild-type and Rfx3−/− embryos as described previously (10). About 10 µg protein were subjected to gel electrophoresis on a 3–8% gradient Tris-acetate gel (Invitrogen), and protein was transferred to a nitrocellulose membrane, which was incubated with rabbit polyclonal anti-Gli3 antibody (1 : 500; Abcam). After incubating with a horseradish peroxidase-conjugated anti-rabbit IgG secondary antibody (1 : 2000; Dako), signal was detected using an ECL Plus detection kit (Amer sham GE healthcare). Band intensity was determined using the ImageJ software. The levels of Gli3R, Gli3A and the Gli3R/Gli3A ratio were compared between wild-type and mutant tissue using the Mann–Whitney test.

Supplementary Material

Supplementary Material is available at HMG online.
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