Precise Somatotopic Thalamocortical Axon Guidance Depends on LPA-Mediated PRG-2/Radixin Signaling

Highlights

- PRG-2 at the growth cone mediates axonal LPA sensitivity of thalamocortical fibers
- LPA/PRG-2/RDX signaling in the growth cone is a novel axon guidance mechanism
- PRG-2 deficiency disrupts precision of thalamocortical somatotopy
- PRG-2\textsuperscript{−/−} mice have altered thalamocortical targeting and specific sensory defects

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In Brief

Cheng et al. describe a new axon guidance mechanism depending on PRG-2 at the growth cone of thalamocortical axons and extracellular LPA synthesized below the cortical plate. Molecular analysis revealed LPA/PRG-2/RDX signaling at the growth cone important for thalamocortical targeting.
Precise Somatotopic Thalamocortical Axon Guidance Depends on LPA-Mediated PRG-2/Radixin Signaling

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SUMMARY

Precise connection of thalamic barreloids with their corresponding cortical barrels is critical for processing of vibrissal sensory information. Here, we show that PRG-2, a phospholipid-interacting molecule, is important for thalamocortical axon guidance. Developing thalamocortical fibers both in PRG-2 full knockout (KO) and in thalamus-specific KO mice prematurely entered the cortical plate, eventually innervating non-corresponding barrels. This misrouting relied on lost axonal sensitivity toward lysophosphatidic acid (LPA), which failed to repel PRG-2-deficient thalamocortical fibers. PRG-2 electroproomation in the PRG-2−/− thalamus restored the aberrant cortical innervation. We identified radixin as a PRG-2 interaction partner and showed that radixin accumulation in growth cones and its LPA-dependent phosphorylation depend on its binding to specific regions within the C-terminal region of PRG-2. In vivo recordings and whisker-specific behavioral tests demonstrated sensory discrimination deficits in PRG-2−/− animals. Our data show that bioactive phospholipids and PRG-2 are critical for guiding thalamic axons to their proper cortical targets.

INTRODUCTION

The thalamocortical projection is essential for sensory processing. In rodents, vibrissae provide input to specific thalamic nuclei and these nuclei have well-defined connections with the somatosensory cortical region where sensory information is processed (Woolsey and Van der Loos, 1970). These connections have a somatotopic organization, and sensory information from each vibrissa reaches a specific barreloid in the thalamic ventrobasal (VB) complex and is relayed to a specific cortical barrel field in the primary somatosensory cortex. During their developmental outgrowth, thalamocortical axons are guided by a variety of cues (Molnár et al., 2012). After crossing the subpallial-pallial border, thalamocortical axons advance within the intermediate zone (IZ) and, approaching the cortex, they accumulate below the cortical plate (CP) at the subplate (Ghosh et al., 1990). The interaction of thalamocortical axons with the subplate is one of the most enigmatic processes in the development of thalamocortical fibers. Even in Reeler or p35 knockout (KO) mice, where the subplate is aberrantly located, thalamocortical axons cross the CP toward the misplaced subplate before connecting to their final targets (Hoerder-Suabedissen and Molnár, 2015). However, the molecular mechanisms that control thalamocortical axon-subplate interaction and ultimately the correct targeting of thalamic projections are largely unknown.

In the developing brain, bioactive phospholipids like lysophosphatidic acid (LPA) play important roles in cortical migration (Fukushima et al., 2000) and neuronal apoptosis (Kingsbury et al., 2003). These effects are mediated by LPA receptors, namely LPA1-R and LPA2-R, which are expressed in the developing brain (Kingsbury et al., 2003). LPA is a well-described repellent factor for axons, eventually leading to growth cone (GC) collapse (Campbell and Holt, 2003). However, while in vitro experiments suggested an involvement of LPA1-R in LPA-mediated axonal retraction, deletion of specific LPA receptors did not lead to significant alterations of fiber tracts in the brain (Contos et al., 2002) and did not affect inhibitory LPA effects on retinal GCs (Birgbauer and Chun, 2010).
Together, this suggests additional LPA-dependent signaling pathways independent of LPA receptors.

LPA is enzymatically inactivated by dephosphorylation via lipid phosphate phosphatases (LPPs), which are cell-surface lipid enzymes and constitute a large family of phosphatidic acid phosphatases (PAP2). Recently, it was shown that LPA signaling at the synapse is controlled by PRG-1/Lppr4 (Trimbach et al., 2009), which belongs to a new class of LPA-interacting molecules (plasticity-related genes, PRGs) sharing transmembrane features of the LPPs (Brauer et al., 2003) but which seem to have a different mode of action (McDermott et al., 2004). Among these, PRG-2/Lppr3 is expressed in developing thalamocortical axons and has a high expression level in GCs. Using both PRG-2−/− animals and a thalamus-specific PRG-2E12E12/Gbx2CreER/R26tdTomato mouse line, we set out to analyze the role of this new LPA regulatory mechanism in outgrowing axons.

RESULTS

PRG-2−/− Mice Have an Aberrant Thalamocortical Projection

PRG-2/Lppr3 is prominently expressed in the thalamus (as shown by in situ hybridization by http://www.genepaint.org; Figure S1B, available online) and in developing thalamocortical fibers (Figures 1A and S1A; for antibody specificity see Figures S1G–S1I), where PRG-2 co-localized with the thalamic axonal marker L1 (Robichaux et al., 2014). According to this expression pattern, we hypothesized that PRG-2 plays a role in the developing thalamocortical projection. We therefore performed biocytin tracing studies in thalamocortical brain slices of wild-type (WT) and PRG-2−/− mice at embryonic day (E)17 (Figures 1D, 1E, 1H, and 1I), a time point when thalamocortical axons accumulate below the subplate (Molnár et al., 1998). Slices were injected with Alexa 488-labeled biocytin into the VB complex of (G1) and at higher magnification (G 2).

Using the Cre-driven RFP reporter, the full extent of the thalamocortical projection. We therefore performed biocytin tracing studies in thalamocortical brain slices of wild-type (WT) and PRG-2−/− mice at embryonic day (E)17 (Figures 1D, 1E, 1H, and 1I), a time point when thalamocortical axons accumulate below the subplate (Molnár et al., 1998). Slices were injected with Alexa 488-labeled biocytin into the VB complex of (G1) and at higher magnification (G 2).

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CP, while PRG-2-expressing fibers were restricted to the IZ (Figures 1B and 1C). Interestingly, in PRG-2−/− animals, the subplate had a normal appearance (Figures S1J and S1K) and the cortical organization, as well as the synaptic connectivity, was not altered (Figures S1M–S1T), ruling out possible subplate or cortical defects that could induce mistargeting of thalamocortical fibers (Ghosh et al., 1990).

Since thalamocortical projection defects observed at prenatal stages might be a temporary feature that eventually disappears after formation of thalamocortical connections, we analyzed the thalamocortical fiber tract at postnatal day (P)5, a time point at which the connections of these fibers are established (Molnár et al., 1998). We applied lipophilic tracers in different brain areas (motor cortex, somatosensory cortex, and visual cortex) at P5 and found that retrogradely labeled thalamocortical fibers derived from their corresponding thalamic nuclei grossly maintained their termination pattern (Figures S2A–S2C). To analyze whether the thalamocortical projection to the somatosensory cortex follows the well-established somatotopic termination pattern to the appropriate barrels, we inserted lipophilic tracers in the VB and visualized the thalamic projection in the corresponding barrels in tangential cut cortical slices. As shown in Figure 1N, in the somatosensory cortex of WT animals lipophilic tracer-stained axons were restricted to their corresponding barrels. However, in PRG-2−/− mice, thalamic axons readily crossed barrel borders aberrantly invading neighboring barrels (Figures 1O, 1P, and S1L). To further assess misrouting in PRG-2−/− mice, we performed retrograde tracing of thalamocortical fibers by paired injection of retrogradely transported fluorescent latex microbeads into adjacent cortical barrels with 400 μm separation between injection sites. As shown by Agmon et al. (1995), even injections placed in close proximity (75–120 μm apart) resulted in segregated neuronal clusters of retrogradely labeled cells in the thalamic VB (Agmon et al., 1995). While in WT animals, even nearby injections of differently labeled fluorescent beads (red and green fluorescence; see Figure S2D) resulted in labeling of distinct subpopulations in the corresponding VB thalamic nuclei (Figures 1Q and 1S), in PRG-2−/− mice, injections even at distant sites (see Figure S2E) resulted in double labeling of the corresponding subpopulations (Figures 1R and 1T; exemplarily retrograde stained neurons in the thalamus are shown in Figures S2F and S2G). Quantitative assessment of double-labeled neurons revealed a significantly higher number in PRG-2−/− animals, while the total number of retrogradely traced neurons did not differ between genotypes (Figures 1U and 1V). Thus, PRG2 depletion in mice leads to a misrouting in thalamocortical projections.

PRG-2 Function Is Required in Thalamocortical Afferents for Correct Targeting

To test whether the aberrant thalamic projection in PRG-2−/− mice results from absence of PRG-2 in thalamocortical axons, we used thalamocortical slice cultures of E15 WT and PRG-2−/− mice at the peak of thalamocortical outgrowth. The thalamocortical projection was visualized by co-electroporation of a GFP-expressing construct into the thalamic VB at E15.5. After 48 hr, slice cultures were fixed and the axonal projection was analyzed (Figure 2A). In WT slices, thalamocortical fibers were restricted to the IZ (Figure 2B), while PRG-2−/− thalamocortical axons aberrantly invaded the CP (Figure 2C). However, when PRG-2 was re-expressed (together with GFP) in thalamocortical axons, these fibers were again restricted to the IZ and did not show obvious differences to WT fibers (Figures 2D and S3A). Quantitative analysis of thalamocortical cultures confirmed the significant amount of aberrant thalamocortical projections and the rescued targeting of these axons after PRG-2 re-expression (Figure 2E). In sum, re-expression of PRG-2 in thalamocortical axons was sufficient to completely abrogate aberrant targeting of the thalamocortical fiber projection in slices from PRG-2−/− mice.

LPA Signaling Is Critical for Thalamocortical Axon Guidance in PRG-2−/− Mice

Since PRG-2 is a member of the family of LPA-interacting molecules (Trimbuch et al., 2009), we analyzed the presence of autotaxin (ATX, ENPP-2, LysoPLD), the LPA-synthesizing enzyme, in the somatosensory cortex at E16. ATX is expressed by subplate neurons, as has been shown on the mRNA level (Hoerder-Suabedissen et al., 2013), and by a GFP reporter at corresponding age (Figures S4A and S4B). Due to its secreted and diffusible nature, we found strong ATX expression at the lower border of the CP and at the upper border of the IZ (Figure 3A). Here, ATX immunoreactivity overlaps with that of LPA and PRG-2, which was expressed in fasciculated thalamocortical fibers in the IZ (Figures 3B–3D; higher magnification in Figures 3E–3H). These findings are in line with a putative role of ATX/LPA in restricting PRG-2-expressing thalamocortical fibers to the IZ. Moreover, LPA-degrading enzymes like LPP1 and LPP3 are strongly expressed in the IZ (Figures S6N and S6O), which suggests that proper regulation of phospholipids allows thalamocortical axons to occupy a corridor in the IZ, possibly contributing to the fasciculated appearance of the thalamocortical projection. We therefore perturbed phospholipid signaling and analyzed the effect of inhibited LPA synthesis on the correct targeting of thalamocortical axons. We applied a recently characterized ATX blocker (PF8380) (Gierrse et al., 2010), which was shown to effectively decrease LPA concentrations (Vogt et al., 2015), to the medium of WT thalamocortical slice cultures and traced thalamocortical fibers using biocytin. After ATX inhibition (0.1 μM), WT thalamocortical fibers aberrantly invaded the CP (Figure 3J), showing a similar phenotype as observed in slices from PRG-2−/− mice (Figures 1E and 1I). Quantitative analysis revealed that this phenotype was observed in more than 92% of all PF8380-treated slices (Figure 3L). To further study whether this effect was due to cortical LPA production, PF8380 was locally injected into the CP. Here, inhibition of ATX in the CP was sufficient to induce an aberrant targeting of thalamocortical fibers (Figure 3K). In contrast, PF8380 injection into the thalamus did not significantly alter the thalamocortical targeting, supporting a role for cortical-derived LPA in thalamocortical guidance (Figure 3L). Taken together, our data provide strong evidence that correct guidance of thalamic fibers to the cortex depends on subplate-derived LPA, produced by ATX, and on the presence of PRG-2 on thalamic fibers.
The role of LPA in GC collapse is well described (Campbell and Holt, 2003); however, the molecular mechanism transducing LPA action from the membrane to the cytoskeleton is not well understood. While specific, G protein-coupled LPA receptors were reported not to affect LPA-induced collapse of retinal GCs (Birgubauer and Chun, 2010), authors of the same group showed that downstream inhibition of the G12/13-rho-ROCK pathway was capable of inhibiting LPA-induced retinal GC collapse (Fincher et al., 2014). This is in line with reports of downstream LPA signal transduction acting on the cytoskeleton (Campbell and Holt, 2003). To analyze the role of axonal PRG-2 in LPA-mediated thalamic axon guidance, we embedded thalamic explants in matrigel with a polarized concentration of LPA (distance to TF-LPA-containing matrigel was 401.6 ± 53.63 μm for WT explants and 475 ± 146.80 μm for PRG-2/−/− explants) and analyzed axonal outgrowth. On one side, the matrigel contained 10 μM fluorescent-labeled LPA (TF-LPA), while the adjacent side did not contain LPA (see Figures 4A, 4B, SSA–SSE, and 4D2 for the border of the TF-LPA-containing matrigel region at higher magnification). The 3D matrigel structure, where fibers are able to radially expand, allows analysis of fiber growth into the LPA-rich zone and the adjacent control zones. However, while WT axons were not able to invade this LPA-rich zone, PRG-2/−/− axons grew far inside it (Figures 4C and 4D1). Quantitative analysis of fibers entering the LPA-rich zone at different distances revealed significantly higher invasion by fibers from PRG-2/−/− thalamic slices (Figure 4E), while no difference between WT and PRG-2/−/− axon fiber outgrowth was observed in the adjacent, non-LPA-containing control side (Figure 4F). Direct comparison of LPA-containing and control sides revealed that WT axons were significantly repelled by the LPA-rich zone, while PRG-2/−/− axons were not affected (Figure 4G). Since PRG-2 deficiency allowed axons to enter the LPA-rich zone, which is repulsive for WT axons, these data suggest that PRG-2 is critically involved in mediating the repulsive LPA sensitivity in thalamic axons.

During normal development, thalamocortical axons do not collapse at the border to the CP but rather turn away from this...
border and continue to grow in the IZ until reaching their final target. This indicates that LPA provided by the subplate induces the turning of axon GCs when thalamocortical fibers approach LPA-rich areas. To mimic this situation in our in vitro outgrowth assay, we exposed thalamocortical explants to lower LPA concentrations (1 μM LPA; Figure 4H). Here, WT axons displayed a typical turning behavior in front of the LPA-rich zone (Figures 4I and 4K). To confirm this observation, we turned to live imaging of cultured thalamocortical axons. Axon GCs were exposed to a low, but continuous, LPA concentration (500 nM LPA). To avoid bias by perfusion flow, which may repel axons, GCs were exposed to an LPA-covered pipette tip at 40 μm distance (see Figure S5F for experimental overview). While WT axons stopped growing and turned away from the LPA source, this behavior was not observed for PRG-2−/− axons, which even seemed to be attracted by LPA (Figures 4L–4N; Movies S1 and S2). Quantitative analysis revealed a significant repulsion of WT axons by 500 nM LPA, which was not observed in PRG-2−/− axons (Figure 4O).
However, \(^{PRG-2^-}\) axons were still repelled by other guidance cues like Sema3A (Figures S5G and S5H), which are well-described repulsive cues (Mintz et al., 2008). In sum, these data indicate that \(^{PRG-2^-}\) at the GC mediates the repellent effect of LPA for axonal guidance.

**Radixin Is a Downstream PRG-2-Interacting Molecule in the Axon GC**

PRG-2/Lppr3 has an intracellular 400 aa-long C-terminal tail, which is putatively involved in signal transduction. To determine interaction partners of this intracellular domain, we first performed a Y2H screen using this PRG-2 C-terminal tail as a bait and found a positive interaction with the FERM domain of radixin (RDX) (Fehon et al., 2010). Using immunoprecipitation (IP) on E17 brain lysate, we confirmed that PRG-2 interacts with RDX (Figures 5A and S6E). This interaction is specific for PRG-2 since only PRG-1 and PRG-2 possess an intracellular C terminus, and PRG-1 is not expressed at embryonic ages (Figure S6J).

To further elucidate the role of this interaction in proper guidance of thalamocortical fibers to the developing cortex, we performed co-localization studies at E16 and found co-expression of PRG-2 and RDX in the thalamocortical tract (Figures 5B, 5C, S6K, and S6L). Expression analysis of PRG-2 and RDX showed a similar expression pattern, where expression for both molecules increased from E11.5, a time point where thalamic neurons are already present and start to differentiate (Figure S6F). To prove that the RDX/PRG-2 interaction in fact occurs in the axon GC, we performed subcellular immunofluorescent co-localization studies of RDX and PRG-2 and found a strong signal in GCs (Figure 5D). Here, PRG-2 is expressed at the GC membrane and co-localizes with RDX, which shows an asymmetric distribution, as described by others (Mintz et al., 2009).

RDX belongs to the ezrin, radixin, and moesin (ERM) protein family. However, due to their embryonal expression pattern at E14.5, ezrin and moesin do not seem to be involved in the outgrowth of the thalamocortical projection (Figures S6L and S6M). The C terminus of RDX can directly bind to filamentous actin (F-actin), thereby acting as a crosslinker between the cortical F-actin cytoskeleton and the plasma membrane (Fehon et al., 2010). The activation of RDX involves the phosphorylation of a critical C-terminal threonine residue (T564), which is highly conserved in the ERM protein family. Therefore, we analyzed the role of PRG-2 in RDX activation using a specific antibody that recognizes the phosphorylation of T564 in RDX, T567 in ezrin, and T558 in moesin (Hausrat et al., 2015). Interestingly, we found significantly lower phosphorylated ERM (pERM) levels in \(^{PRG-2^-}\) neurons (Figures 5E and 5F), indicating diminished activation of ERM proteins. To prove the role of LPA in ERM activation, we assessed the effect of different LPA concentrations. While 100 nM LPA already revealed an increase in ERM phosphorylation, 1 \(\mu\)M LPA induced a significant and robust increase in pERM levels (Figures 5G and 5H). We further analyzed pERM levels over time and found a fast and constant increase in ERM phosphorylation in the course of LPA stimulation (Figures 5I, 5J, and S6A–S6C). To understand the functional significance of the LPA-dependent pERM expression, we analyzed the RDX/PRG-2 interaction under LPA-free conditions and after 1 \(\mu\)M LPA stimulation for 15 min. Here, we used RDX-GFP transfected, stable PRG-2-expressing HEK cells, which have comparable PRG-2 expression, and performed co-IPs of RDX and PRG-2 using an anti-PRG-2 antibody (Figure 5K). Quantitative analysis (RDX levels were adjusted according to the PRG-2 IP signal) revealed a significantly increased RDX/PRG-2 interaction upon 1 \(\mu\)M LPA stimulation (Figure 5L). Interestingly, analysis of the IP product revealed that phosphorylation of the PRG-2-interacting RDX was significantly higher upon LPA stimulation when compared to serum-starved cells (Figures 5M and 5N). These data suggest that higher RDX phosphorylation upon LPA stimulation augmented PRG-2/RDX association. Since we could show that RDX binds to the intracellular C terminus of PRG-2 (Figure S6D), to further analyze whether RDX phosphorylation is a prerequisite for PRG-2 binding and to determine the PRG-2 binding sites, we performed a systematic peptide scan using a Celluspot \(\mu\) Array format (\(\mu\)SPOT) and mapped the RDX/PRG-2 interaction (Figure 5O). Here, PRG-2 intracellular C-terminal overlapping peptides (69 overlapping peptide fragments, 14 aa in length) were analyzed regarding their binding affinity to RDX-GST or to RDX\(^{T564A}\)-GST, its non-phosphorylated form. We detected six putative interaction sites that showed significant difference in their binding to RDX-GST or to RDX\(^{T564A}\)-GST, pointing to a lower binding affinity of the non-phosphorylated RDX\(^{T564A}\)-GST, its non-phosphorylated form. In line with data from the peptide-microarray-based mapping, although binding of RDX\(^{T564A}\)-GST was present at lower levels when compared to RDX\(^{WT}\) binding could be clearly detected (Figures 5P and 5Q). This points to the fact that RDX phosphorylation, although not a prerequisite for RDX/PRG-2 interaction, significantly increases RDX/PRG-2 interaction. To answer the question of whether RDX/PRG-2 association at the membrane is important for LPA-dependent RDX phosphorylation in neurons, we analyzed the effect of LPA stimulation on pERM expression in WT and \(^{PRG-2^-}\) neurons. Here, LPA stimulation of WT neurons resulted in a significant increase in pERM levels, while \(^{PRG-2^-}\) neurons failed to show any significant change (Figures 5R and 5S). These results point to a yet unknown, but critical role of PRG-2 in LPA-induced RDX activation.

**PRG-2 Mediates pERM Increase at GC Membranes Induced by Extracellular LPA**

RDX is concentrated below the plasma membrane, a critical position linking transmembrane signaling to the actin cytoskeleton. Since phosphorylation influences RDX activity in connecting the actin-cytoskeleton with the membrane (Fehon et al., 2010), transmembrane signaling events leading to RDX activation are optimally suited to act in extracellular cue mediated axon guidance. We therefore analyzed the effect of extracellular LPA in inducing pERM changes at the GC membrane in WT and \(^{PRG-2^-}\) neurons. In WT GCs, pERM was concentrated in a compartment directly adjacent to the GC membrane (Figures 6A and 6B) and strongly increased upon LPA stimulation (Figures 6E, 6F, and 6I). In contrast, no change of pERM was observed in \(^{PRG-2^-}\) GCs (Figures 6C, 6D, and 6G–6I). Moreover, direct
comparison of pERM levels in the center of the GC and in the periphery revealed a shift of the pERM signal toward the membrane of WT GCs, but not of PRG-2−/− GCs (Figure 6U). To prove whether RDX is dynamically recruited to the membrane via PRG-2 binding induced by a local extracellular LPA source, we transfected WT or PRG-2−/− neurons with GFP-tagged RDX and performed live imaging studies. While in WT neurons, local LPA stimulation rapidly induced a translocation of RDX-GFP to the membrane next to the LPA source (Figures 6K and 6L), PRG-2−/− neurons did not show such a change (Figures 6M and 6N). In sum, these data show that PRG-2 is the mediator of LPA-induced localization of RDX to the GC membrane, where it is present in its phosphorylated form, in turn increasing binding to PRG-2.

To prove that ERM phosphorylation and activation play a role in LPA-induced GC turning, we analyzed the subcellular localization of pERM in GCs of thalamic explants exposed to an LPA-rich environment (1 μM LPA), as shown in Figure 4I. Detailed analysis of turning GCs, exemplarily encircled in Figure 7A, revealed strong pERM signal co-localized with F-actin at the tip of GCs turning away from the LPA-rich zone (Figures 7B–7E). To confirm that PRG-2 interacts with RDX at the GC, we performed a proximity ligation assay (PLA), a method that allows the study of in situ protein-protein interactions at resolutions <40 nm (Rhett et al., 2011). Here, we found a strong signal at the GC tip, confirming the PRG-2/RDX interaction at the site of LPA action (Figures 7F and 7G). Taken together, these results confirm the critical role of PRG-2/RDX interaction for the induction of RDX activation in GCs by extracellular LPA, which eventually leads to axon turning.

**RDX Deficiency Phenocopies PRG-2 Thalamocortical Axon Defect**

To further substantiate the role of RDX in LPA-induced GC turning, we analyzed the thalamocortical projection in RDX−/− mice. Using biocytin tracing in living E17 thalamocortical slices, we observed misrouted thalamocortical axons in RDX−/− slices that aberrantly invaded the CP (Figures 7I–7K), while in slices from control littersmates, the thalamocortical projection was confined to the IZ (Figures 7H and 7K). In order to better understand the functional role of the PRG-2/RDX interaction in thalamocortical axon guidance, we analyzed the effect of partial reduction of these proteins in transheterozygous mice (PRG-2−/−/RDX−/−) in thalamocortical slices at E17. Here, the thalamocortical projection aberrantly invaded the CP, suggesting that it is indeed the interaction of PRG-2 and RDX that is important for correct guidance of these fibers by an LPA gradient (Figures 7L–7O). These data corroborate the idea that the LPA-PRG-2-RDX/pERM signal transduction axis is critical for correct guidance and restriction of thalamocortical axons to the IZ. Moreover, in line with decreased RDX expression after E16 (Figures S6G and S6H), these data could explain how thalamocortical axons are finally able to enter into their target cortical region.

**Functional and Behavioral Alterations in PRG-2−/− Mice**

To assess the functional outcome of altered thalamocortical projections in PRG-2−/− mice, we first measured intrinsic optical responses in the corresponding barrel upon single-whisker back-and-forth deflection. This approach measures blood-flow changes in the representation of single whiskers in the barrel cortex upon activation. Using this method, we found that the response area upon single-whisker stimulation, which corresponds to the stimulated barrel (for morphological barrel assessment, see Figure S7A), was larger in PRG-2−/− mice. These data point to a more diffuse and broader cortical response extending beyond the border of the activated barrel (Figures 8A–8C). Moreover, it reflects our findings using retrograde tracing of thalamocortical fibers in PRG-2−/− mice showing a clear overlap in the termination pattern of neurons from neighboring thalamic barreloids in the VPM (Figures 1R, 1T, and 1U). This is in agreement with enlargement of functional barrel columns in mice due to

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**Figure 4. PRG-2 Deficiency Abolishes Thalamic Axon Sensitivity to LPA**

(A and B) Thalamic explants from WT (A) and PRG-2−/− (B) mice exposed to 10 μM TF-LPA for 40 hr. TF-LPA-containing zone is located in the lower left part of the image. The opposite, LPA-free right side was regarded as control (C). Lines delineate regions located 100 or 200 μM within the LPA-containing or the control (C) region, respectively. Outgrowing axons stained for TuJ1 were color coded in green.

(C and D) Higher magnification of WT axons at the TF-LPA interface shows a repulsive effect on outgrowing axons (C, arrows). In contrast, PRG-2−/− axons were able to enter the LPA-containing region (D1). Higher magnification shows the border of the TF-LPA-containing region colored in red (D2).

(E) Quantitative analysis of fibers protruding 100 and 200 μM into the LPA-rich region (normalized to the number of fibers 100 μM before the TF-LPA-rich matrigel). (Mann-Whitney test; n = 8 WT and 8 PRG-2−/− thalamic explants).

(F) On the control side (C; lower right corner in A and B, containing no TF-LPA), no difference was observed when WT and PRG-2−/− axons were analyzed at the same distances (as measured to the LPA front). (Mann-Whitney test for 100 μM and t test for 200 μM; n = 8 WT and 9 PRG-2−/− thalamic explants).

(G) Comparison of axon numbers at different depths on the LPA-rich side and on the control side. (Kruskal-Wallis test with Dunn’s multiple comparisons test; n = 8 WT and 8 PRG-2−/− thalamic explants).

(H and I) When exposed to lower LPA concentrations (1 μM; H; thalamic WT axons displayed a turning behavior (I; red arrows pointing to turning axons) in front of the LPA-rich region. Border of the LPA-rich zone is marked by dotted line and visible by addition of red fluorescent beads.

(J) Thalamic explants from PRG-2−/− mice at the border of the LPA-rich area displayed no turning behavior and entered the LPA-rich zone in high numbers.

(K) Quantitative analysis of the ratio of turning axons to the total number of axons revealed that WT axons displayed a turning behavior while PRG-2−/− axons were not affected by 1 μM LPA (one-way ANOVA with Bonferroni correction for multiple comparisons; n = 8 WT thalamic explants exposed to 1 μM LPA and to control conditions, 11 PRG-2−/− thalamic explants exposed to 1 μM LPA, and 8 to control conditions).

(L and M) Live imaging of WT (L) and PRG-2−/− GCs (M) exposed to low LPA concentrations (500 nM) at a distance of 40 μM for 60 min (see also Movies S1 and S2; Figure S3F). Line intersection represents GC starting point used for graphic display of axon growth shown in (N).

(N) Schematic diagram showing starting point and end point of GCs during live imaging. Axons extending during live imaging (60 min) into the quadrant facing the LPA source were assigned positive values; axons growing in other quadrants were assigned negative values.

(O) Analysis of GC behavior (unpaired t test; n = 9 WT and 9 PRG-2−/− GCs).

*p < 0.05, **p < 0.01, ***p < 0.001. Bars represent mean ± SEM.

Scale bars, 500 (A and B), 100 (C and D1), 500 (H), 50 (I and J), and 10 μM (L–N).
impaired precision of the thalamocortical projection, as reported in other mouse models (Lokmane et al., 2013).

To get a layer-specific resolution of neuronal activity, we performed in vivo electrophysiological recordings using multi-channel electrodes. We observed in PRG-2 /– mice a slight decrease in multi-unit activity (MUA) upon single-whisker deflection in layer IV and a significant decrease in layers II/III for PRG-2 /– mice compared to WT controls (Figures 5D–5F), which are the main target layers of the thalamocortical projection (Meyer et al., 2010). Recent data suggest that layers II/III contain neurons specifically responsive for texture coarseness (Garion et al., 2014). We therefore assessed the ability of PRG-2 /– mice to distinguish texture coarseness using a modified radial arm maze, where arms were covered with sandpaper of two slightly different grades (60 grade [grain size, 265 μm] and 80 grade [grain size, 190 μm]; Figure 5G). Only arms covered with the coarser (60 grade) sandpaper were rewarded by accessible food pellets and counted as correct entrance. Since experiments were performed in darkness and no other cues were presented, mice had to orient by sensing sandpaper roughness. While WT mice showed a significant improvement in their ability to choose the correct arm, PRG-2 /– mice did not improve beyond chance levels even after 9 days of continuous training (Figure 5H). To exclude bias by an impaired learning ability, we tested PRG-2 /– mice in the Morris water maze (MWM). Escape latency for visible and hidden platforms was indistinguishable between PRG-2 /– and WT littermates (Figures S7E–S7G). Since learning per se was not impaired in PRG-2 /– mice, we conclude that their performance at chance level in the texture discrimination task is a direct result of impaired thalamocortical axon projection. In line with unaltered learning ability, motor performance was also unaltered, as shown by analysis in the MWM or supporting limb responses (Figures S7H–S7K). To exclude an effect of PRG-2 deletion outside the thalamus, e.g., in the cortex, which could affect cortical processing of somatosensory information, we also tested the above described thalamus-specific PRG-2−/− deficient mouse line (PRG-2ΔE12/ΔE12). As shown in Figures S8L–S8O using retrograde tracing of thalamocortical fibers projecting to the cortical barrels, these animals display a similar phenotype as described for the constitutive PRG-2 /– animals, where we could show that different barrels were aberrantly innervated by the same neuron in the thalamic VPM. Furthermore, staining for PRG-2 revealed that fibers originating from Gbx-2−/− expressing thalamic neurons (identified by the RFP reporter) did not express PRG-2 (Figures S7P–S7S). Behavioral analysis of the PRG-2ΔE12/ΔE12 mice in the sandpaper radial maze revealed that these mice displayed the same phenotype as the constitutive PRG-2 /– mice and were not able to distinguish between the two sandpapers of different coarseness (Figure 5H). Together with our in vivo imaging and electrophysiological data, these findings strongly indicate that misguided thalamocortical axons in PRG-2−/− mice alter cortical information processing, resulting in a specific deficit in sensory discrimination at adult ages.

DISCUSSION

An intriguing phenomenon in cortical development is the specific outgrowth of thalamocortical fibers that reach the appropriate cortical area before their target cells migrate to their final position in the CP (Kostovic and Rakic, 1990). As shown in other systems, thalamocortical axons accumulate in the IZ below the CP (Ghosh et al., 1990). Here, they interact with a transient neuronal population, the subplate neurons, which keep them in place and are thereby critically important for correct patterning of the thalamocortical connections. Although the role of subplate neurons has long been recognized and gene expression patterns of subplate neurons have been described (for review, see Hoerder-Suabedissen and Molnár, 2015), molecular mechanisms regulating the subplate neurons/thalamocortical fiber interaction are far from understood.
from being understood. Here, we analyzed the role of PRG-2/Lppr3, a phospholipid-interacting molecule strongly expressed in thalamocortical axons during development (Figure 1A).

Using a constitutive PRG-2 KO mouse line as well as a thalamus-specific PRG-2 KO line (PRG-2 fl/fl/Gbx2CreER/R26tdTomato), where tamoxifen application at E12.5 leads to specific deletion of PRG-2 in the thalamus (PRG-2 DE12/E12; Normand et al., 2013), we discovered that PRG-2-deficient thalamocortical axons prematurely and aberrantly entered the CP. This altered the precise somatotopic termination of thalamocortical fibers in the somatosensory barrel field cortex and led to a deficit in sensory discrimination. Molecular analysis revealed a signal transduction axis in the GC leading from the LPA-synthesizing molecule ATX to LPA, PRG-2, and finally to phosphorylated RDX, a well-described linker between the GC membrane and the actin cytoskeleton. We demonstrate this ATX/LPA/PRG-2/pRDX signaling axis on thalamocortical axon GC turning and suggest that this mechanism is responsible for keeping thalamocortical axons in the IZ before entering the CP.

**Role of Phospholipids in Subplate/Thalamocortical Axon Interaction**

The role of subplate cells was concluded from ablation experiments, where thalamocortical fibers, no longer able to contact subplate cells, escaped the subcortical waiting period and extended beyond their target cortical regions. Here, we found that PRG-2−/− thalamocortical axons lose their guidance by the subplate. This specific phenotype relies mechanistically on...
the lost sensitivity of thalamocortical axons toward extracellular LPA gradients synthesized by ATX, a diffusible molecule expressed in subplate neurons (Hoerder-Suabedissen et al., 2013). Eventually, this leads to loss of proper axon guidance and to an invasion of inappropriate cortical compartments. Since we also analyzed thalamus-specific PRG-2^−/−/RDX^−/− KO animals (Normand et al., 2013), we could rule out that this misrouting is due to defects, e.g., derived from cortical PRG-2 deficiency. Interestingly, the same phenotype was detected when the catalytic function of ATX was blocked, thereby impeding LPA...
RDX, an F-Actin-Interacting Molecule, Is the Downstream Executer of LPA/PRG-2-Dependent Axon Guidance

The critical importance of PRG-2-mediated extracellular LPA effects on thalamic axon guidance was further clarified by single-axon analysis. While downstream mechanisms mediating LPA GC collapse have been described (Campbell and Holt, 2003), direct and specific LPA-interacting molecules involved in this mechanism remain unclear. In line with live imaging data, WT axons were repelled/collapsed when in contact with 10 μM LPA, while lower LPA concentrations (1 μM) induced a turning behavior in GCs. However, PRG-2−/− axons were not affected even by high LPA concentrations (10 μM). Searching for downstream interacting partners, we found RDX to directly interact and be co-expressed with PRG-2 in thalamocortical axons.

Figure 8. PRG-2−/− Mice Have Altered Somatosensory Cortical Processing and a Deficit in Somatosensory Discrimination

(A and B) Color-coded images represent changes in reflection of hemodynamic response (ΔR/R0) after single-whisker stimulation using intrinsic optical imaging (see also Figures S7C and S7D) in WT (A) and PRG-2−/− animals (B). Threshold set at 5% ΔR/R0 reflecting steepest drop of hemodynamic response is delineated. High hemodynamic response to single-whisker stimulation in WT animals (A) typically corresponds to one barrel with a sharp delineation toward the neighboring barrels.

(C) PRG-2−/− mice revealed a significantly broader hemodynamic response (Mann-Whitney test; n = 7 mice per group). *p < 0.05 at 5% threshold.

(D) Averaged traces of evoked multi-unit activity (MUA) responses recorded in six WT (black trace) and six PRG-2−/− (red trace) mice. Blue dashed line indicates time point of whisker deflection and blue braces indicate period from 0 to 100 ms after whisker deflection used for calculation of MUA shown in (E) and (F).

(E and F) MUA in layers II/III (E) and layer IV (F), both marked with green star in (D). Unpaired t test; n = 17 barrel columns from 6 animals per genotype).

(G) Image of the eight-arm maze designed for somatosensory perception where arms are covered with sandpaper of different grain sizes.

(H) Constitutive PRG-2−/− mice and thalamus-specific PRG-2-deficient mice (PRG-2 ΔE12/ΔE12) show significant differences in correct performance when compared to WT mice (two-way ANOVA [genotype, time]; n = 8 WT, 20 PRG-2−/−, and 11 PRG-2ΔE12ΔE12 animals).

*p < 0.05, ****p < 0.0001. Bars represent mean ± SEM. Scale bar, 200 μm (A).
Moreover, subcellular analyses revealed RDX to interact with PRG-2 in GCs as shown by PLA. Importantly, IP experiments revealed that the PRG-2/RDX binding was highly increased upon LPA stimulation. The role of this PRG-2/RDX interaction was functionally confirmed, showing that PRG-2 is critically important for RDX phosphorylation, in PRG-2−/− neurons, RDX phosphorylation did not increase, even upon 15 min of 1 μM LPA stimulation. Our data showing RDX phosphorylation upon LPA stimulation in full-length PRG-2-expressing HEK293 cells suggest that PRG-2/RDX association increased RDX phosphorylation upon LPA stimulation. However, when analyzing binding to PRG-2 to either WT RDX-GFP or its non-phosphorylated form (RDX1564A-GFP), we detected binding under both conditions, albeit with a significant reduction of binding to PRG-2 of the non-phosphorylated form. These data show that RDX phosphorylation is not a prerequisite for PRG-2/RDX binding; however, it influences this binding.

Our finding that transheterozygous loss of PRG-2 and RDX already results in altered thalamocortical fiber projection indicates an important role of the LPA/PRG-2/RDX signal axis. Since the active form of ERMs (pERM) is critical for GC guidance (Mintz et al., 2008), we investigated the role of this signal axis at the GC, the critical location for axon guidance. Here, LPA stimulation induced strong ERM phosphorylation at the GC membrane of WT axons, while PRG-2−/− GCs failed to show these changes. In line, RDX was rapidly translocated to the membrane directly next to a local LPA source, indicating an active role of LPA/PRG-2/RDX signaling in GC guidance. Together with the detection of pERM adjacent to F-actin at the tip of turning GCs, this signaling axis appears to directly control LPA-induced axon turning.

In Vivo Effects of Altered Thalamocortical Guidance in Adult PRG-2−/− Mice

Using in vivo imaging and high-density multi-electrode recordings in the barrel cortex of PRG-2−/− mice and performing behavioral tests both in PRG-2−/− mice and PRG-2ΔE12/ΔE12 animals, we elucidated the functional consequences of PRG-2 deficiency and subsequent altered thalamocortical connectivity in cortical layers IV and II/III as well as in the characteristic discrimination abilities associated with these layers (Garion et al., 2014). It appears that rather specific deficits can be observed even upon drastic developmental defects resulting in grossly impaired cortical lamination and subsequent changes in the thalamocortical projection (Wagener et al., 2016), as shown by preserved functional sensory maps in the somatosensory barrel cortex observed in Reelin-deficient mice (Guy et al., 2015). Interestingly, Cossell et al. (2015) suggest microcircuits in layers II/III are important for specific stimulus orientation (Cossell et al., 2015), supporting the idea that layer II/III neurons mediate rather specific functions. This implies that functional changes in layers II/III seen in PRG-2−/− mice can best explain the sensory discrimination deficits of the animal.

Taken together, we describe a novel molecular guidance mechanism involving phospholipid signaling influencing thalamocortical axon guidance at a critical time point, at which axons exhibit a waiting period at the cortical subplate. The alteration of the LPA/PRG-2/pRDX signal transduction pathway resulted in distinct electrophysiological and behavioral deficits in adult animals and highlighted the importance of this pathway for proper thalamocortical connectivity.

EXPERIMENTAL PROCEDURES

Mouse Lines

All experiments were conducted in accordance with the national laws for the use of animals in research and with the European Communities Council Directive 86/609/EEC, and approved by the local ethical committee (Landesuntersuchungsamt Rheinland-Pfalz 23. 177-07/G 13-1-073). Details of the generation and breeding of PRG-2−/−, PRG-2ΔE12/ΔE12/Gbx2CreER/R26tdTomato and RDX−/− mice are provided in the Supplemental Experimental Procedures.

Immunofluorescent Staining and Tracing Experiments

Immunofluorescent (IF) stainings, PLA for detection of in situ protein interaction, and anterograde and retrograde tracing experiments were performed following standard protocols.

Slice Culture Preparation, Biocytin Labeling, and Slice Electroporation

Organotypic slices were prepared as described in the Supplemental Experimental Procedures. For biocytin tracing experiments, pressure injections were performed into E17 embryonic slices using glass capillaries attached to a Toohey Spritzer (Toohey Company) to induce biocytin Alexa Fluor 488 (Thermo Fisher Scientific). Slices were then incubated for 6 hr and fixed with 4% PFA (4%). For electroporation experiments, corresponding plasmids were injected into organotypic slices from E15.5 brains that were electroporated by Needle Tungsten Electrodes and an NEPA21 Super Electroporator (Nepa Gene).

Assessment of pERM Levels in Cortical Neurons

Cortical neurons were prepared as described (Vogt et al., 2012), cultivated for 2 days in vitro (DIV) and used for assessment of pERM levels in the GC. IF staining and pERM assessment were performed as described in the Supplemental Experimental Procedures.

IP Studies

IP studies were performed on E17 brain material or on cell lysate from stably PRG-2-expressing HEK293 cells according to standard protocols.

Intrinsic Signal Optical Imaging and In Vivo Electrophysiological Recordings

Intrinsic signal optical imaging (IOI) and in vivo electrophysiological recordings in the barrel cortex of adult mice were performed according to standard protocols as described in the Supplemental Experimental Procedures.

Animal Behavior

Sandpaper discrimination test and MWM were performed according to standard procedures.

Statistical Analysis

For animal experiments, mice from the same litter or of similar age were chosen. The investigator was blinded to the genotype of the animals. Following experiments, results were analyzed, animals were re-genotyped, and corresponding final statistical analyses were performed. Briefly, after assessing for normal distribution (using the Kolmogorov-Smirnov test), comparison between two groups, if not otherwise stated, was performed using a two-tailed unpaired t test for normal distributed data or a Mann-Whitney test for nonparametric data. When data were normalized to controls (set as 100%), a one-sample t test was used. For quantitative assessment of aberrant fibers in traced or electroporated slices, a Fisher’s exact test was applied. P value was adjusted for multiple comparisons. Comparison between more than two groups was performed using a one-way ANOVA for parametric data or a Kruskal-Wallis test for nonparametric data.
SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, eight figures, one table, and two movies and can be found with this article online at http://dx.doi.org/10.1016/j.neuron.2016.08.035.

AUTHOR CONTRIBUTIONS

Conceptualization, J.V. and R.N.; Methodology, J.V., J.C., K.R., and A.S.; Investigation, J.C., S.S., T.J.H., J.-W.Y., H.E., X.L., Y.L., R.B., H.J., N.S., H.M.M., A.H.-S., P.-H.P., W.F., and T.T.; B.V. and R.S. provided reagents; Writing – Original Draft, J.V.; Review and Editing, J.V., R.N., J.C., T.J.H., O.N., Z.M., H.J.L., and M.K.; Supervision, J.V., R.N., A.S., H.J.L., J.H., Z.M., K.S., M.K., and W.F.; Funding Acquisition, J.V., R.N., H.J.L., A.S., and M.K.

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