Polarization of migrating cortical neurons by Rap1 and N-cadherin
Revisiting the model for the Reelin signaling pathway

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Neuronal migration is essential for the development of the cerebral cortex. Mutations leading to defective migration are associated with numerous brain pathologies. An important challenge in the field is to understand the intrinsic and extrinsic mechanisms that regulate neuronal migration during normal development and in disease. Many small GTPases are expressed in the central nervous system during embryonic development. Recent findings have shown that Rap1 and its downstream partners Ral, Rac and Cdc42 are involved in the maintenance of N-Cadherin at the plasma membrane which is necessary for the correct polarization of migrating neurons. The activation of Rap1 is triggered by Reelin, an extracellular protein known for its role in the organization of the cortex into layers of neurons. In the absence of Reelin, neurons exhibit a broader and irregular pattern of positioning. The prevailing model suggests that Reelin signals to neurons during the last step of their migration, a notion that is inconsistent with new data describing an effect of Reelin on early steps of migration.

In regard to these recent findings I suggest a revised model, which I call the “polarity model,” that further refines our understanding of the developmental function played by Reelin and its downstream small GTPases.

Introduction

Small GTPases are guanine nucleotide binding proteins that function as molecular switches by cycling between active GTP-bound and inactive GDP-bound states. They are activated by Guanine nucleotide Exchange Factors (GEFs) that induce GTP loading and inhibited by GTPase Activating Proteins (GAPs) that return them into their GDP loaded inactive form. The large superfamily of small GTPases (also named the Ras superfamily) is comprised of the Rho/Rap/Ral, Rho, Ral, Arf and Ran families. They regulate a wide variety of essential cellular processes such as cell division, adhesion, polarity, migration and differentiation. It is thus not surprising that many of their members are involved directly or indirectly in numerous pathological conditions including cancer and brain developmental diseases.

This extra view will focus on the Rap subfamily of small GTPases. It is composed of five related proteins, Rap1 (A and B) and Rap2 (A, B and C) which have overlapping functions and patterns of expression. Rap1 in particular has been intensively studied for its role in the regulation of integrin-mediated cell adhesion and the control of endothelial and epithelial cadherin-based cell-cell junction integrity. We have recently demonstrated a new function of Rap1 in regulating the polarity of migrating neurons in the developing mouse brain cortex through the control of N-Cadherin (NCad). We found that Reelin, an extracellular matrix protein which has an important function in the organization of the cortex, triggers the activation of Rap1 in cortical neurons when they are midway through their migration path, at a stage where re-polarization occurs. These new findings do not fit with the current model in which Reelin affects neurons at the end of their migration. Here, I suggest a revised model of action for the Reelin signaling pathway with a central function for small GTPases.
I will start this extra view with a description of the development of the cerebral cortex and the recently discovered role played by Rap1. Then I will discuss the downstream proteins involved in this function, followed by the upstream Reelin signal.

**Cortical Development and Rap1**

**Small GTPases**

The development of the brain requires that neurons migrate away from their birth place in order to perform their functions properly. In addition, neurons have to extend neurites and ultimately differentiate and communicate with each other. Characterization of the molecular signaling pathways involved in cerebral cortex development is important for the understanding of brain pathologies such as lissencephaly, microcephaly, periventricular heterotopia, epilepsy, dyslexia, mental retardation, schizophrenia, bipolar disorder, and many others resulting from defective cortical architecture, connectivity and function. The majority of neurons in the cortex are the excitatory glutamatergic neurons that are generated from progenitor cells located at the ventricular zone (VZ) which lines the ventricle (Fig. 1).

These neurons undergo different phases of migration starting with a polarized migration within the VZ, which is followed by a more complex movement within the multipolar morphology zone (MMZ), which is made up of the sub-ventricular zone (sVZ) and the lower part of the intermediate zone (IZ), where they undergo more divisions. Importantly, at this stage, the neurons lose their bipolar morphology and elongate several neurites which is why they have been referred to as having a multipolar morphology. This occurs along with a slower migration and a few switches in their direction of movement. The neurons then migrate within the radial morphology zone (RMZ), comprising the upper part of the IZ and the cortical plate (CP), to reach the top of the CP. During the transition from MMZ to RMZ, they change morphology to become bipolar once again. Within the CP, new neurons migrate past the older ones already installed resulting in “inside-out” layering, that is, a gradient of cells with younger neurons in the outermost field of the CP and older neurons more inside. During the multipolar stage, and in spite of the multiple changes of direction of migration, the net movement is still directed toward the CP. Rap1 has emerged as a critical regulator of this polarization. Similar to all small GTPases, Rap proteins are regulated by specific GAPs or GEFs. Important clues for their involvement in brain development came from the phenotype of a hypomorphic mutant mouse for C3G, a Rap-specific GEF, which exhibits an accumulation of neurons within the MMZ. This arrest in cell migration is mainly due to the highly disorganized radial glia fibers (the main migration substrate for this type of cell) and disintegration of the basement membrane. Moreover, these mice die around E14.5 making it difficult to study the migration of later born neurons. C3G is not the only GEF regulating Rap during the development of the cortex. In fact, a dorsal telencephalon-specific knockout of PDZ-GEF-1, another activating protein for Rap1 and Rap2 enzymes, results in the accumulation of neurons underneath a normally developed cortex. The involvement of Rap activators in cortical development suggested that Rap enzymes might also play an important role. Indeed, the neuron-specific inhibition of Rap (i.e., without affecting radial glial cells) in vivo induces an ectopic accumulation of neurons within the MMZ. Time lapse video-microscopy revealed that this
phenotype is not the result of defective neuronal motility of the affected multipolar cells, but is rather due to a defect in their polarization toward the RMZ. This is because the movement of the Rap-inhibited neurons is randomized with a decreased net movement toward the RMZ. This phenotype is not due to a complete arrest of invasion of the RMZ, because many cells migrate out of the MMZ, albeit with a significant delay when compared with control cells. Surprisingly, the subsequent radial bipolar migration along glia fibers (also called glia-guided locomotion) is not affected by the absence of Rap activity. This absence of effect on locomotion has been confirmed in an in vitro lattice culture system where dissociated neurons move along glia fibers. Together, these observations suggest that Rap is important for the initial polarization of neurons but not migration per se.

Rap1 Polarizes Neurons Through Its Regulation of N-Cadherin, with a Potential Involvement ofRal, Rac and Cdc42

Signaling through the small GTPase Rap1 has been implicated in both integrin-mediated and cadherin-mediated adhesion events. To date, studies examining neuron-specific deletion of β1 Integrin[16] or the Integrin downstream effector FAK (focal adhesion kinase) did not observe any defect in glia-guided migration. On the other hand, both Rap1 and cadherins and their interaction are emerging as important regulators during the brain development. Classical cadherins are single-pass transmembrane adhesion receptors involved in cell-cell contact and epithelial polarity through calcium dependent homophilic binding. Intracellularly, cadherins interact with catenin family members. p120-catenin binds to the juxtamembrane region of cadherin to stabilize it at the plasma membrane, while α- and β-catenin serve a dynamic role in linking cadherin to the actin cytoskeleton. Many members of the cadherin family are expressed in the central nervous system and one of them, N-Cad, has recently attracted the interest of neuroscientists.

The prevailing view is that cadherin functions to mediate adhesion between stationary cells, thereby maintaining tissue integrity and segregation of different cell populations. Indeed, throughout development, N-Cad is highly exposed in the vertebrate central nervous system and its conditional deletion in the dorsal telencephalon results in disruption of the adhe-

erens junctions localized at the apical end of neuroepithelial cells, where N-Cad is most highly concentrated. This results in a general disruption of neuroepithelial integrity and aberrant radial glia fibers that do not expand toward the pial surface. However, evidence is emerging that cadherins also regulate cellular motility. In the rat caudal hindbrain, classic cadherins regulate tangential migration of precere-

bellar neurons.[17] In the zebrafish, NCad concentrates transiently at the front of cerebellar granule cells during the initia-
tion of their chain-migration and is required for them to polarize prior to migrate.[18] In the early chick embryo PDGF signaling controls NCad expression in mesoderm cells, which is required for efficient migration.[19] Interestingly, C3G and PDZ-GEF1, two of the Rap-specific GEFs known to be important for mammalian brain development (see above), have also been linked to the cadherins. In epithelial cells, C3G directly interacts with E-cadherin and is important for the initial steps of adherens junction forma-
tion.[20,21] While PDZ-GEF1 is recruited by MAGI-1 at VE-cadherin-mediated endothelial cell-cell adhesions.[22] Our recent findings demonstrated that in the mammalian cerebral cortex NCad has an important function in polarizing cortical neurons before they are able to start migrating into the RMZ. The inhibition of cadherins in post-mitotic neurons without affecting progenitor cells and their radial glia fibers, recapitulates the phenotype induced by inhibition of Rap1 i.e., loss of polarity during the multipolar migration with no effect on the speed of migration as multipolar or bipolar neu-
ones. Several experiments confirmed that NCad functions downstream from Rap1. First, inhibition of Rap1 in vivo and in vitro reduced the presence of NCad at the plasma membrane with a concomitant decrease in intracellular NCad. Second, a functional assay demonstrated that inhibi-
tion of Rap1 reduced the binding of neurons to the NCad extracellular domain.

And finally, overexpression of NCad in the cortex is able to partially rescue the cell positioning defect due to inhibition of Rap1. These data suggest that Rap1 activity is important in migrating neurons in order to maintain the high level of NCad at the plasma membrane necessary to allow cells to polarize correctly. Yet we do not know whether other cadherins, also expressed in the MMZ, might have some redundant function with NCad. In addition, how NCad allows the polariza-
tion of cortical neurons is still under investigation. Nevertheless, hypotheses might be suggested. NCad may be activated locally in order to increase the binding to radially-oriented processes on other neurons or glial fibers. This adhesion could stabilize the position of the centrosome. A similar model has been suggested for the directional chain migration of cerebellar granule neurons in the zebrafish with NCad transiently accumulating at the front of the cells.[13] However, in other cell types, it is the cadherin-free cell edge that shows the polarity of migration. Indeed, cadherin-mediated cell-cell inter-

actions induce the centrosome and Golgi apparatus to move toward the free cell edges in cultured astrocytes and stimulate protrusions at the free edge in Xenopus neural crest cells.[23] Alternatively, NCad may be a regulator for other cell surface receptors that respond to directional signals from the CP. For example, NCad modulates FG-2 signaling in MCF-7 breast cancer cells and VE-cadherin regulates TGFβ signaling in endothelial cells.[24] This model has parallels with the migration of Drosophila border cells, where Drosophila E-Cad (Decad) is required in the migrating cells as well as in the cells they migrate between.[25] The border cells extend a long leading process, whose direction is specified by a growth factor gradient but whose formation requires Decad.[26] The induction of the long extensions on border cells may be analogous to the induction of a radial leading process on cortical neurons, and in both cases surface cadherin expression may be key to developing the polarity needed for the process (Fig. 2).
The Rap effector RalGDS activates RalA, which docks secretory vesicles to the exocyst complex and recycles E-Cadherin to epithelial cell-cell junctions. It has been shown that Rap1, Rac1 and Cdc42, activated by nectins, are able to trigger the formation of adherens junctions in epithelial cells and fibroblasts. This positions them upstream of cadherin function. However, they also can be activated downstream of cadherins, suggesting a potential positive feedback loop. In epithelial cells, Rap1 is important for the recruitment of E-cadherin into nascent cell-cell contact sites and in this process it functions upstream of Rac1 and Cdc42: by recruiting their GEF Vav2. Cdc42 has been reported to regulate the trafficking of basolateral membrane proteins as well as to modulate the association of cell-cell contacts with the actin cytoskeleton. Finally, activation of Rac1 and Cdc42 by Rap1 could also stabilize cadherins at the membrane through their target IQGAP1. Studies have suggested two possibilities as to how they work. First, it has been proposed that IQGAP1 destabilizes cadherins by inducing their dissociation from α-catenin and IQGAP1 is negatively regulated by Rac1 and Cdc42. Second, another study suggested that IQGAP stabilizes cadherins through the reorganization of the actin cytoskeleton and is positively regulated by Rac1 and Cdc42. Regardless of which of these pathways downstream of Rap1 are predominant, perhaps depending on cell types or experimental conditions, all those results indicate that, in epithelial cells, stimulation of Rap1 may induce the activation of Rac1 and Cdc42, which in turn might facilitate the directional vesicle transport of E-cadherin and/or organize the actin cytoskeleton, enabling neighboring cells to contact one another. Other regulatory pathways have also been proposed such as the interaction of Rap1 with AF6, increasing AF6 association with p120 catenin which in turn strengthens the interaction of p120 catenin with E-cadherin, reducing its internalization and/or degradation.

Although migrating neurons are different from static epithelial cells making contacts, Rap1, Ral, Rac and Cdc42 may function similarly to regulate cell to cell contact of cortical neurons through the regulation of NCad. Our inhibition and rescue experiments in the animal and in vitro suggested that Rap1, Ral, Rac and Cdc42 may also play a role influencing Rap1’s effect on the presence of NCad at the plasma membrane and the resulting function on polarity of cortical neurons. This process is likely to be very dynamic and control of the amount of cadherins at the cell surface must be tightly regulated by a balance between endocytosis and recycling to the sites of new contact formation. Of note, a recent study demonstrated the involvement of Rab5-dependent endocytic and Rab11-dependent recycling pathways in the regulation of NCad in cortical neurons, while, in epithelial cells, Rap1 has been shown to co-localize with E-cadherin at the Rab11-positive recycling endosome compartment. It would be of interest to determine whether Rap1 and the Rab pathways work in parallel or cooperatively in the regulation of NCad and neuronal migration. Another important next step would also be to investigate whether the exocyst, downstream of Ral, or IQGAP and other proteins downstream of Rac and Cdc42, play roles in the polarization of cortical neurons through the Rap1/NCad pathway.

Reelin Polarizes Multipolar Neurons Toward the RMZ Through Rap1 and NCad: A New Model of Action

An important question is what stimulates Rap1 and its downstream effectors in the polarization of migrating neurons.
Reelin is an extracellular protein secreted by Cajal-Retzius cells present in the marginal zone above the RMZ and is required for the correct organization of the CP. The prevailing model suggests that Reelin acts on migrating neurons when they arrive at the top of the RMZ during their final somal translocation (named "detach and go" because cells would detach from the radial glia then proceed through the final somal translocation). This model explains why, in the absence of Reelin, the inside-out layering of the cortex is inverted. However, it is important to point out that the reeler cortical phenotype is not simply an inversion of the neuronal layering. Even though the earliest neurons shift from a deep laminar position to form a superficially located superplate in the reeler brains, the later born neurons exhibit a broader and irregular distribution which is far more than just an inversion of laminar fate. This suggests that the current model may not fully explain the phenotype. Earlier studies already indicated that Reelin may signal the neurons before they start their radial migration within the RMZ. First, the active cleavage fragment of Reelin has been shown to diffuse from the marginal zone into the deep tissue. Second, cells in the MMZ express the highest level of functional Reelin receptors. Recently, we

![Diagram of neuronal migration](image)

**Figure 3.** Regulation of neuronal migration within the mammalian cerebral cortex by the Reelin/Rap1/N-cadherin pathway and other small GTPases. (A) Reelin signals through Rap1 and its downstream enzymes Ral, Rac and Cdc42) and N-cadherin on neurons at the MMZ to polarize them toward the RMZ. Rab5 and Rab11 might also be involved. The subsequent glia-guided migration within the RMZ is independent of Reelin, Rap1 or N-cadherin. During this migration, the Reelin signal is downregulated (degradation of phosphorylated Dab1, downregulation of functional receptors) and N-cadherin is downregulated by the Rab7 pathway which may be a consequence of the downregulation of the Reelin pathway. When neurons reach the top of the RMZ, they undergo a final somal translocation. This final somal translocation could depend on the reduction of the Reelin signal or, alternatively, Reelin may induce a second set of intracellular signals in cells performing the final somal translocation as they are more mature than when they encountered Reelin for the first time in the RMZ and are in a different biological context. (B) In the absence of Reelin, Rap1 or N-cadherin, neurons are disoriented within the MMZ. Rab5 or Rab11 inhibition induces a similar phenotype. In the absence of Reelin or Rab7, the final somal translocation is also affected and could be, at least in part, a consequence of defective N-cadherin downregulation.
showed that inhibition of Reelin by in utero electroporation delays the cells at the RMZ and affects the localization of NCad at their plasma membrane, mimicking the phenotype caused by Rap inhibition. Rescue experiments in vivo confirmed that Rap1 is involved in this phenotype downstream of Reelin. These data suggest that Reelin is, at least in part, responsible for the Rap1/N-cadherin-mediated polarization function in neurons migrating within the MMZ. In agreement with this, NCad protein levels are decreased in the embryonic reeler mutant cerebral cortical cells exclusively at the MMZ. A function of Reelin through Rap1 and NCad on polarity of neurons before they enter the RMZ may better explain the disorganized positioning of late-born neurons. Indeed, in the absence of a polarizing signal, neurons would exist the MMZ in a disorderly manner disregarding their date of birth. It is surprising that Reelin affects the polarization of neurons because previous experiments showed that the localization of the source of Reelin is not important for its function in the CP. After the polarization mediated by Reelin may thus be indirect. Reelin would rather act as a permissive signal allowing neurons to respond to another cue which still remains to be discovered.

After neurons have received the polarizing cue triggered by Reelin, the signal is downregulated. Previous works have shown that neurons downregulate the signal and/or become less responsive to Reelin once they commence migration within the RMZ. For example, Reelin induces the downregulation of its functional receptors at the time neurons migrate within the RMZ. Also, Reelin induces the phosphorylation of the intracellular adaptor Dab1 which is consequently degraded. It is therefore likely that membrane-associated NCad levels are downregulated because the Reelin signal is not any longer there to maintain it at the membrane. Indeed, NCad also exhibits a downregulation of its expression in the wild-type RMZ, which is much less pronounced in the reeler brain. Interestingly, the down-regulation of the Reelin signal seems to be an important event for a correct neuronal positioning. A testable prediction would be that the downregulation of NCad depends on the decreased Reelin signal. Indeed, a recent study showed that Rab7-dependent degradation of NCad is important for the final phase of migration when neurons reach the top of the RMZ. Therefore, I propose a model (Fig. 3) in which Reelin first triggers the polarization of neurons when they are at the MMZ by activating Rap1 and stabilizing NCad at the plasma membrane but, at the same time as phosphorylated Dab1 and the Reelin signal are downregulated, NCad is also degraded through a Rab7 pathway when neurons migrate as bipolar cells. This downregulation is a consequence of their migration at the multipolar stage. In this view, the final somal translocation is a consequence of the downregulation of the Reelin signal that was initiated when the cells were in the MMZ.

In the above model, Reelin signals only in multipolar cells. However, it is also conceivable that Reelin stimulates migrating neurons twice: first at the MMZ and a second hit later during the final somal translocation. Neurons might be refractory to Reelin signaling after the first stimulation but become sensitive again when they reach the top of the RMZ.

Future Directions

The next challenge in the field is to determine exactly how Rap1 and its client GTPases RapA/B, Rac and Cdc42 affect the polarity of cortical neurons. Other small GTPases such as members of the Rab family might also come into play. How all those small GTPases finely tune neuronal migration needs to be investigated in more details. How NCad, and maybe other cadherins, regulate the polarized movement of cortical neurons is also ground for future work. Reelin certainly does not work alone and it will be exciting to uncover the interconnection of the multiple signals that regulate the development of the cerebral cortex.

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