**Pax6 regulates granule cell polarization during parallel fiber formation in the developing cerebellum**

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**SUMMARY**

The molecular mechanisms that govern the coordinated programs of axonogenesis and cell body migration of the cerebellar granule cell are not well understood. In Pax6 mutant rats ((rSey)/rSey), granule cells in the external germinal layer (EGL) fail to form parallel fiber axons and to migrate tangentially along these fibers despite normal expression of differentiation markers. In culture, mutant cells sprout multiple neurites with enlarged growth cones, suggesting that the absence of Pax6 function perturbs cytoskeletal organization. Some of these alterations are cell-autonomous and rescuable by ectopic expression of Pax6 but not by co-culture with wild-type EGL cells. Cell-autonomous control of cytoskeletal dynamics by Pax6 is independent of the ROCK-mediated Rho small GTPase pathway. We propose that in addition to its roles during early patterning of the CNS, Pax6 is involved in a novel regulatory step of cytoskeletal organization during polarization and migration of CNS neurons.

Key words: Pax6, Cerebellum, Cell polarization, Granule cell, Parallel fiber, Cytoskeleton

**INTRODUCTION**

Neural circuitry of the vertebrate CNS develops through the coordinated program of neuronal migration, neurite outgrowth and synaptic interconnections between constituent neurons. The mammalian cerebellum provides a favorable system with which to study the formation of neural circuitry because of its relative simplicity and clear cytoarchitecture. Early steps that are crucial for the establishment of synaptic connectivity in the cerebellar cortex are axon formation and dynamic migration of granule cell neurons (Hatten and Heintz, 1995; Altman and Bayer, 1997). Granule cells are generated in the external germinal layer (EGL), which covers the outer surface of the cerebellar cortex. Then, postmitotic neurons in the deep EGL extend bipolar axons horizontally to the pial surface, and subsequently move their cell bodies along the axis of the developing axons. During postnatal development, granule cells extrude a third process inwardly and make a 90° turn to descend into the internal granular layer (IGL). Migrating neurons leave trailing processes attached to the horizontal beams of the bipolar axons, forming the characteristic T-shaped axons, the parallel fibers. These parallel fibers ultimately innervate the Purkinje cells, the principal neurons of the cerebellar cortex.

A variety of molecules are known to act during axon formation and migration (Goodman and Shatz, 1993; Goodhill, 1998). These include long-range and short-range guidance molecules, signaling molecules that interpret and decode these cues, and cytoskeletal components that regulate cell morphology. Several signaling molecules, including ROCK (Bito et al., 2000), Unc51.1 (Tomoda et al., 1999) and DDR1 (Bhatt et al., 2000), have been implicated in the early steps of axonogenesis of granule cells in vitro. ROCK is a serine/threonine kinase activated downstream of Rho GTPase. Intensive studies have elucidated the importance of regulation by Rho family small GTPases in cytoskeletal reorganization during neuronal morphogenesis and migration (Luo et al., 1996; Threadgill et al., 1997; Zipkin et al., 1997; Steven et al., 1998; Albertinazzi et al., 1999; Nakayama et al., 2000). Inhibition of ROCK activity in cultured granule cells triggers excessive sprouting of axons, suggesting that the Rho/ROCK pathway negatively controls axon outgrowth probably through organizing actin dynamics, especially during the initial step of polarity induction (Bradke and Dotti, 1999; Bito et al., 2000). However, Unc51.1 and DDR1 have been shown to act as positive regulators of axon outgrowth as revealed by dominant-negative inhibition of these molecules (Tomoda et al., 1999; Bhatt et al., 2000). It remains elusive, however, how the...
activities of these molecules are orchestrated during cellular morphogenesis of the granule cell.

Differentiation of granule cells is thought to depend on both genetic programs intrinsic to the granule cells and epigenetic influences of adjacent siblings as well as of other cell types in the developing cerebellum (Trenkner et al., 1984; Dahmane and Ruiz-I-Altaba, 1999; Wechsler-Reya and Scott, 1999). Studies using culture systems have demonstrated that granule cells autonomously develop polarity to form parallel fibers in the absence of spatial cues (Nagata and Nakatsuji, 1991; Powell et al., 1997). One candidate molecule implicated in such an intrinsic mechanism is a paired domain-containing homeobox gene, \textit{Pax6}. In the \textit{Pax6} mutant mouse, small eye (\textit{Sev}/\textit{Sev}), the EGL is malformed and the horizontal array of newly forming parallel fibers is disorganized (Engelkamp et al., 1999). Besides this anomaly in the EGL, multiple neurological defects are also seen in the mutant cerebellum, including attenuated foliation, misplacement of a subset of EGL cells and deformity in the vermis-forming territory (Engelkamp et al., 1999). It remains unknown which of these defects are direct or indirect consequences of lack of \textit{Pax6} function.

\textit{Pax6} is known as a morphogenetic gene with a myriad of activities in patterning (Stoykova et al., 2000; Toresson et al., 2000; Yun et al., 2001) and cell-type specification (Burrill et al., 1997; Ericson et al., 1997; Osumi et al., 1997) during early development of the CNS. It has also been shown that \textit{Pax6} is involved in various aspects of axon pathfinding and migration of CNS neurons. These include radial migration of cortical neurons (Schmahl et al., 1993; Caric et al., 1997), formation of thalamo-cortical projections (Kawano et al., 1999; Pratt et al., 2000), and formation of longitudinal and medial-ventral tracts in the forebrain (Mastick et al., 1997; Vitalis et al., 2000). However, disruption of these processes in the \textit{Pax6} mutant CNS appears to be an indirect consequence of the loss of a prerequisite role of \textit{Pax6} in conferring regional identity to the neurons, which either express \textit{Pax6} or interact with \textit{Pax6}-expressing cells. In the forebrain, for example, \textit{Pax6} expression defines the midbrain-forebrain boundary and provides local guidance information to the post-optic commissure axons (Mastick et al., 1997).

In this study, we present evidence for a novel role for \textit{Pax6} as a critical regulator in cell polarization in the cerebellar granule cell. \textit{Pax6} mutation caused severe deficits in initial polarization of axons and the cytoskeletal reorganization in growth cones. Furthermore, cell body migration was aberrant in \textit{Pax6} mutant granule cells. These phenotypes appeared to be due to the absence of intrinsic activity of \textit{Pax6} in the granule cell. It is thus suggested that \textit{Pax6}-mediated transcriptional control is essential to achieve proper granule cell polarization during parallel fiber formation both in vivo and in vitro.

**MATERIALS AND METHODS**

**Animals**

small eye rats (\textit{rSev}/\textit{rSev}) allele on a Sprague-Dawley (SD) background; Osumi et al., 1997) were maintained in laboratory colonies at the National Center for Neurology and Psychiatry and Tohoku University Graduate School of Medicine. Homozygous embryos derived from intercrosses of \textit{rSev}/\textit{rSev} were distinguishable by their external morphology. Wild-type embryos were obtained from intercrosses of heterozygous or of wild-type SD rats. The day of vaginal plug was considered as E0.

**Microexplant and reaggregate cultures**

Microexplant cultures of wild-type and \textit{rSev}/\textit{rSev}-\textit{EGL} at E21 were prepared as described by Nagata and Nakatsuji (Nagata and Nakatsuji, 1991). Reaggregate cultures were basically performed as described (Nagata and Nakatsuji, 1994). Briefly, the EGLs from E21 wild-type and \textit{rSev}/\textit{rSev} embryos were dissociated, and fractions containing small neurons were collected and labeled with PKH26 (Sigma). Unlabeled host cells were prepared from P2 EGL and mixed with labeled cells at a ratio of 20:1. The cell mixture was centrifuged in an Eppendorf tube and incubated for 1 hour to make reaggregates that were then cut into 300-400 \mu m pieces and placed on culture dishes coated with poly-D-lysine/laminin. Cultures were maintained at 37°C in 5% CO2 and analyzed after 1-3 days.

**Immunohistochemistry and fluorescent microscopy**

Cultures and fresh-frozen cryosections (10 \mu m) were fixed with 4% paraformaldehyde (PFA) for 15 minutes at room temperature. Primary and secondary antibodies used for staining were as follows: mouse monoclonal antibody (mAb) against TAG-1 (4D7: 1:4, DSHB); mAb against Zic1 (1:50, kindly provided by J. Aruga); mAb against MAP2 (1:200, Sigma); mAb against Tau1 (1:200, Boehringer Mannheim); mAb against α-tubulin (1:200, Sigma); rabbit polyclonal antibody (pAb) against Pax6 (Inoue et al., 2000); pAb against 440-kD Ankyrin (1:200, kindly provided by M. Kunimoto); and goat anti-rabbit or anti-mouse secondary antibodies conjugated with FITC, AMCA (Chemicon) or Alexa568 (Molecular Probes). F-actin was stained with Oregon Green 488-phalloidin (1:100, Molecular Probes). After washing, slides were mounted in a glycerol-based medium SlowFade (Molecular Probes) and analyzed using a Nikon E800 microscope with \times 20 and \times 40 PlanFluor objectives.

For quantitative analysis of the number and length of axons, the trace from the contour limit of the cell soma to the tip of the major process of the neuron was defined as its axon length. Processes longer than 3 \mu m emerging from the soma, or those longer than 10 \mu m bifurcating from the shaft were counted as axons. To quantify the relative size of the growth cones, live cells labeled with PKH26 were kept at 37°C and observed using water-immersed 40× Fluor objective (0.80W). The size of growth cones was measured using the graphic image of IpLab (Scananalytic).

**DiI labeling of EGL cells**

E21 cerebella fixed with 4% paraformaldehyde (PFA) were embedded in 3% agar and cut coronally at a thickness of 200 \mu m using a vibratome. Small crystals of the lipophilic dye DiI were placed on the EGL and left for 10 days at 4°C to allow diffusion of the dye. After washing away the crystals, sections were mounted and analyzed using a fluorescent microscope.

**Transmission and scanning electron microscopies (TEM and SEM)**

Cultured cells were fixed with 4% PFA and 5% glutaraldehyde in phosphate buffer (PB; 0.1M, pH 7.4) at room temperature overnight. They were washed with PB and post-fixed with 1% osmium tetroxide in PB for 1 hour. For TEM, the samples were washed well with distilled water (DW) and stained en bloc with 1% uranyl acetate for 1 hour. They were then dehydrated through a graded ethanol series, cleared with hydroxypropyl metacrylate and embedded in Epon 812. They were cut with a ultramicrotome and observed under a TEM (JEM-1200EX, JEOL; or EM-002B, Topcon, Japan) at an accelerating voltage of 80 kV. For SEM, the samples washed with DW were subsequently immersed in 2% tannic acid for 1 hour, and then 1% osmium tetroxide in DW for 1 hour (Katsumoto et al., 1981). They were dehydrated and immersed in t-butyl alcohol. The samples were
dried at critical point (JFD-300, JEOL, Japan), coated with platinum/palladium and observed under an SEM (H-800, Hitachi, Japan) at an accelerating voltage of 15 kV.

In situ hybridization
In situ hybridization on cryosections was performed by a method described previously (Lin and Cepko, 1998). The Pax6 cDNA used as a probe was described previously (Matsuo et al., 1993).

Transfection of plasmid cDNA
Plasmid cDNAs used for transfection were as follows: mouse Pax6 and Pax6-EnR cDNAs subcloned in a pCAX expression vector (full-length cDNA of mouse Pax6 was kindly provided by P. Gruss); ROCK-Delta3 (Ishizaki et al., 1997); C3-GFP (Watanabe et al., 1999); and EGFP-N1 (Clontech).

For expression in a microexplant culture, a modified calcium phosphate procedure developed by A. Ghosh was used (Threadgill et al., 1997). Transfections were performed 6-12 hours after plating when emigrating cells were observed. Plasmids used were either 2 μg GFP-N1 alone or a mixture of 0.8 μg GFP-N1 and 2.4 μg plasmid of interest per well on multi-well plates (Nunc Multidish four wells, Nune). Analysis was carried out 24-36 hours after transfection. Under these conditions, usually 10-40 neurons expressed GFP. The efficacy of co-transfection was >80%.

Transfections in low-density cultures of dissociated cells were performed as previously described (Bito et al., 2000). Small cerebellar neurons were labeled with PKH26 and plated at 2×10^5 cells per well in polylysine/laminin-coated wells. Cells were fixed 20 hours after transfection for fluorescent microscopy. Some cells were stained with F-actin or with anti-Tau instead of PKH26, all of which gave basically the same results.

Construction and infection of recombinant retrovirus
The Pax6-EnR chimera was created by fusing the N-terminal domain of Pax6, which contains two DNA binding sites (amino acids 3-306), with the Engrailed repressor domain. Using PCR-based mutagenesis, NcoI and EcoRI sites were introduced in the Pax6 cDNA at the first methionine and at the amino acids 307-308, respectively. Primers used were as follows; 5’-GACTCGAGCCATGGAGAACAGTCACAGCCG-3’, 5’-CTGATATCCACAGGTGTGGTGGCTG-3’. The NcoI-EcoRV fragment of the PCR product was ligated into a pBluescript SK+.

The insert was cut with EcoRI and then ligated into a NcoI-EcoRI-digested EnR-pSLAX21 (kindly provided by T. Furukawa). The entire Pax6-EnR fusion and EnR were cloned into a retroviral vector pLIA as described by Furukawa et al. (Furukawa et al., 1997). The viruses were produced in a Phoenix cell line (generously provided by G. Nolan), and were concentrated at 1×10^5-1×10^8 pfu/ml according to the procedure described by Cepko (Cepko, 1998).

To inject virus, neonatal ICR mice were anesthetized on ice. A Hamilton microliter syringe (Hamilton, Whitter, CA) was inserted through the skin and the skull to a position above the EGL and 1 μl of virus solution was injected over 2 minutes. The pups were revived at 36°C and returned to the litter. At 10 days post-transfection, infected cerebella were dissected, sectioned at 50 μm using vibratome and stained for AP.

RESULTS
Parallel fiber formation is disrupted in the Pax6 mutant cerebellum
Granule cell progenitors form the EGL by the end of gestation. During postnatal development, the EGL is subdivided into two sublayers; the granule cell progenitors continue mitosis in the outer sublayer, whereas postmitotic neurons start to extend bipolar axons and move horizontally in the inner sublayer of the EGL (Hatten and Heintz, 1995; Altman and Bayer, 1997). Previous studies have documented that, while proliferation remained unaffected, the postmitotic granule cells fail to segregate in the inner premigratory layer of the EGL in the Pax6 mutant mice small eye (Sey/Sey; Engelkamp et al., 1999).

Fig. 1. Disrupted formation of the EGL and the parallel fibers in Pax6 mutant (rSey2/rSey2) cerebella. (A,B) HE-stained cross-sections of E21 cerebella. The wild-type EGL (A) is subdivided in the outer germinall layer (g) and the inner premigratory layer (m). Spindle-shaped cells (arrows) in the premigratory layer are presumably migrating along the young parallel fibers. Pax6 mutant EGL (B) is thicker and uniform accumulation of round cells is observed. (C,D) Dil-labeled parallel fibers. Horizontal beams of parallel fibers in the deep EGL of the wild-type cerebellum (C) are not formed in the Pax6 mutant (D). Pial surface of the cerebella is indicated by asterisks. (E,F) Expression of TAG-1 in E21 cerebella. TAG-1 signal delineates the boundary between the EGL and the molecular layer in the wild-type cerebellum (E). TAG-1 is diffusely expressed throughout the EGL in the Pax6 mutant cerebellum (F). Some background autofluorescence can be observed in the vascular cells on the pial surface of the EGL. Scale bars: 25 μm in A-D; 40 μm in E,F.
Using a Pax6 mutant rat rSey2/rSey2 (Osumi et al., 1997), we confirmed that this phenotype was a general feature of Pax6-deficiency (Fig. 1A,B). We next analyzed the formation of parallel fibers in the wild-type and the mutant cerebella by DiI labeling and immunofluorescent studies. In the E21 wild-type rat, DiI particles implanted in the EGL labeled multiple long neurites aligned parallel to the pial surface in the deeper portion of the EGL (Fig. 1C). The deep EGL was also delineated by expression of TAG-1, a transient marker for early parallel fibers (Fig. 1E; Furley et al., 1990; Yamamoto et al., 1990). These data suggest that the postmitotic EGL cells are already forming the premigratory layer and extending parallel fibers by late gestation. In contrast, DiI incorporation by several round EGL cells resulted in labeling of very few neurites in the rSey2/rSey2 EGL (Fig. 1D). Additionally, TAG-1-positive cells were randomly dispersed in the rSey2/rSey2 EGL and never observed to form a sharp band, as seen in the wild-type cerebellum (Fig. 1E,F). These results indicate that the EGL cells of Pax6 mutants fail to form the premigratory layer and to extend the parallel fibers, in accordance with the previous study (Engelkamp et al., 1999).

**Pax6 is expressed in the granule cells during development**

To explore Pax6 activity during granule cell differentiation, we examined Pax6 expression in developing rat cerebella. Pax6 transcripts were detected in the EGL throughout pre- and postnatal development (Fig. 2A-D). At E20, intense expression of Pax6 mRNA was observed in the EGL cells and in dispersed cells in the cerebellar cortical primordia (Fig. 2A). After P5, Pax6 expression was seen in postmigratory granule cells in the emerging IGL at reduced levels compared with premigratory EGL cells (Fig. 2B-D; Stoykova and Gruss, 1994). Observation at higher magnification revealed that a subset of cells in the molecular layer (ML) also expressed Pax6 mRNA (arrows in Fig. 2E). Expression of Pax6 protein showed basically the same profile (Fig. 2F); immunoreactivity was prominent in the EGL cells and declined in the IGL cells. Cells in the ML that expressed Pax6 protein had fusiform nuclei, suggesting that they were granule cells migrating from the EGL to the IGL. Thus, Pax6 mRNA and protein are expressed in granule cells throughout the course of their differentiation and migration.

**Pax6 mutant cells differentiate as granule cells but fail to form bipolar axons and migrate in random orientation**

The expression pattern of Pax6 in developing cerebella prompted us to examine its potential role during neurite outgrowth and migration of granule cells. As the homozygous mutants die at birth during the early phase of granule cell differentiation, we performed microexplant cultures of E21 EGL in which cell-autonomous migration and cellular morphogenesis of the granule cells can be reconstituted (Nagata and Nakatsuji, 1991). By 1 day in culture (1 DIV), wild-type cells extruded long neurites radially and emigrated out from the EGL explant. The radial neurites formed parallel bundles after 2 DIV. TEM and time-lapse tracing showed that most neurons maintained close contact with the bundles of preexisting long neurites and migrated along straight trajectories (Fig. 3A,C). In contrast, only a few radial fibers were seen in the Pax6 mutant EGL culture. TEM revealed that many round cells failed to form long leading processes in the mutant cultures (Fig. 3B). Active migration was observed, although the cells dispersed randomly (Fig. 3D). The radial migration of cultured granule cells has been shown to be dependent on laminin substrate (Nagata and Nakatsuji, 1990). We therefore checked the expression of putative binding partners of laminin in granule cells, L1 and β1-integrin. Both are expressed equally in wild-type and mutant cells, precluding the possibility that the disorganized appearance of Pax6 mutant cells was due to mere non-responsiveness to laminin substrate (data not shown).

The morphology and behavior of the mutant cells raised the possibility that differentiation of the granule cells might be disturbed or delayed in the absence of Pax6 expression. To test this, we examined the expression of several neuronal and granule cell markers by immunofluorescence. We first confirmed that 70-80% of emigrating cells from the wild-type and mutant explants were granule cells and/or granule cell progenitors, as revealed by Zic1 expression (data not shown; Aruga et al., 1994). These cells strongly expressed Pax6...
Pax6 controls parallel fiber formation

protein in wild-type cultures, while no Pax6 antigen was detectable in mutant cultures (Fig. 3E,F). A subset of cells that emigrated from the wild-type explant expressed TAG-1, an early marker for parallel fiber axons, on their long bipolar neurites (Fig. 3G). In contrast, although equal numbers of Pax6 mutant cells showed strong expression of TAG-1 around their cell bodies, extension of TAG-1-positive long neurites was rarely seen (Fig. 3H). The expression of the microtubule-associated protein, MAP2, was mostly confined to the leading processes at 2 DIV in both wild-type and mutant explants (Ono et al., 1994). While wild-type cells regularly aligned long leading processes distally to the explant, mutant cells randomly misoriented short processes regardless of the orientation of other cells (Fig. 3J). Pax6 mutant cells occasionally had multiple processes that expressed MAP2, suggesting an abnormal control of the leading process formation. After 3 DIV, many cells in both the wild-type and the mutant cultures had ceased expressing TAG-1, suggesting they had exited the early phase of granule cell differentiation (data not shown). These cells indeed strongly expressed 440 kDa AnkyrinB, a specific marker for the parallel fiber axons (Kunimoto, 1995), together with Tau1, a general marker for axons (Fig. 3K,L and data not shown). In the wild-type culture, the massive radial neurites appeared to differentiate into AnkyrinB-positive parallel fibers. Pax6 mutant cells also synthesized Ankyrin but formed large meshes of irregular neurites rather than organized parallel bundles. Little difference was observed in the relative numbers of GFAP-positive glia, most of which were found in close proximity to the explant, between the wild-type and the mutant cultures (around 10%; data not shown). After 3 DIV, we observed apoptotic cell death more frequently in mutant cultures, although it affected neither the apparent density nor the total number of surviving cells (data not shown). Taken together, these results suggest that Pax6 mutant cells can recapitulate the typical cascade of gene expression during granule cell differentiation, but fail in correctly polarizing their parallel fiber axons and subsequently migrate in a disorganized manner.

Pax6 regulates morphogenesis and migration of granule cells in a cell-autonomous manner.

The specific expression of Pax6 in developing granule cells suggests that abnormal polarization of rSey2/rSey2 granule cells might be a direct consequence of loss of intrinsic Pax6 activity within the differentiating granule cells. Alternatively, it may indicate that Pax6 is involved in cell-cell interaction of granule cells with the same and/or other cell types in the cerebellum. To distinguish between these possibilities, we created a mixed culture of rSey2/rSey2 mutant EGL cells with wild-type cells.
Cellular morphology was visualized by labeling cells with a fluorescent membrane-soluble dye PKH26 (Fig. 4A). The control wild-type cells labeled with PKH26 extended long neurites and migrated radially along the fibers (Fig. 4B). We occasionally observed cells with characteristic T-shaped axons resulting from vertical reorientation (Fig. 4D). On the other hand, Pax6 mutant cells rarely elongated long neurites and scattered randomly in spite of the regular radial fibers formed by surrounding wild-type cells (Fig. 4C,E). These results suggest that the abnormal morphology and migration of Pax6 mutant cells are cell-autonomous, as they could not be rescued by co-culture with wild-type cells.

To assess whether Pax6 activity is sufficient to correct the anomalies of the mutant cells, we transfected Pax6 cDNA into mutant cells. The number and the length of neurites of Zic1-positive cells were quantified by co-transfecting green fluorescent protein (GFP) cDNA in explant cultures (Fig. 5). When GFP cDNA was transfected alone, the proportion of Zic1-positive cells with longer neurites was significantly higher in wild-type cells than in mutant cells. Notably, we found a considerable number of neurons with more than two neurites in mutant cultures. In contrast, expression of Pax6 in mutant cells restored wild-type phenotypes; most of the cells co-transfected with Pax6 and GFP cDNAs became bipolar and extended neurites to lengths comparable with wild-type cells.

These results strongly suggest that Pax6 functions intrinsically in the granule cell upstream of the formation of long bipolar neurites and orderly migration. To verify that Pax6 function is truly involved in the genetic cascade of granule cell differentiation in normal developing cerebellum, we misexpressed a dominant-negative form of Pax6 in granule cell precursors in situ as well as in vitro. The DNA-binding domain of Pax6 was fused to the repressor domain of Drosophila Engrailed (EnR), producing a fusion protein that should block transcriptional activation by intrinsic Pax6 (Furukawa et al., 1997; Matsunaga et al., 2000). We first confirmed that misexpression of the Pax6-EnR cDNA in the microexplant culture of wild-type cells resulted in multipolar morphology similar to rSey2/rSey2 cells (Fig. 5). Thus, perturbation in the maintenance of Pax6 function to activate transcription of the downstream target gene(s) was sufficient to cause the morphological deficits observed in Pax6 mutant EGL cells even within the wild-type genetic background.

By utilizing a replication-incompetent retrovirus vector (pLIA), we next misexpressed the dominant-negative gene in neonatal EGL cells and assayed morphology and location of the infected cells after 10 days by staining for the marker gene AP (Fig. 6). During postnatal development, granule cell precursors represent almost the sole mitotic cells in the EGL infectable by retrovirus (Zhang and Goldman, 1996). Granule cell progenitors infected with control vector viruses or those carrying Engrailed repressor domain alone differentiated normally and correctly migrated to the IGL (Fig. 6B,F). They elongated bipolar parallel fibers in the molecular layer, formed glomeruli at the tip of dendrites and adopted a T-shape morphology, as visualized by AP staining (Fig. 6D). By contrast, most of the cells infected with viruses transducing the Pax6-EnR transgene remained in the EGL (Fig. 6C,F). Short, bushy processes were observed around their cell bodies in the EGL and few or no parallel fibers were formed in the ML (Fig. 6E). While descendant cells carrying the control viruses spread out over a broad area, those misexpressing Pax6-EnR tended to remain clustered around the injection site, possibly owing to the failure in tangential migration of the EGL cells in the absence of the leading process formation. It was also notable that a significantly smaller number of infected cells were observed with Pax6-EnR viruses, suggesting that Pax6 might also contribute towards maintaining survival of granule cells during differentiation.

Taken together, these results indicate that Pax6 functions as an intrinsic factor required for elongation of bipolar axons and subsequent orderly migration of granule cells in the developing EGL.

**Aberrant sprouting of axons accompanies enlargement of growth cones in the Pax6 mutant granule cells**

The next obvious question was which step of cellular morphogenesis and migration was disturbed in the Pax6 mutant granule cells. As shown in Fig. 3, Pax6 mutant EGL cells expressed a series of differentiation markers, precluding the possibility that the onset of molecular switches in granule cell differentiation was simply retarded in the absence of Pax6 function. Taking into account that Pax6 is a transcription factor,
Pax6 controls parallel fiber formation

and that its action is cell-autonomous in the granule cell, we hypothesized that Pax6 regulates transcription of cytoplasmic and/or transmembrane protein(s) implicated in cellular morphogenesis of granule cells during axon formation and migration.

We noticed that Pax6 mutant cells had multipolar processes with occasional bifurcated branches or lamellipodia (Fig. 7A,B). Growth cones at the tips of axons were also found to be larger in the mutant cells compared to the wild-type (Fig. 7C-G). Double-immunofluorescence staining revealed entry of actin and microtubule networks into those growth cones (Fig. 7E,F). These phenotypes were reminiscent of those seen following treatment with an inhibitor of p160 ROCK, a Rho GTPase-associated kinase (Bito et al., 2000).

The above observation raised an intriguing possibility that Pax6 might regulate the expression of factor(s) implicated in the Rho/ROCK pathway. To assess this possibility, we first examined initial outgrowth of axons in dissociated EGL cells (Fig. 8). By 20 hours after plating, most wild-type cells extruded bipolar neurites, presumably forming future parallel fiber axons (Fig. 8A,G). In contrast, a significant proportion of Pax6 mutant cells possessed an increased number of processes (Fig. 8D,G). Differences in neurite lengths between wild-type and mutant cells were not prominent at 20 hours of culture. Wild-type cells treated with Y27632, a specific inhibitor of ROCK, exhibited multipolar morphology similar to Pax6 mutant cells (Fig. 8B). The only noticeable difference in their phenotypes was a slight increase in axon length in Y27632-treated cells (Fig. 8H). Overexpression of a RhoA-inhibiting enzyme, C3, resulted in further enhancement of similar phenotypic changes (Fig. 8C). We also tested modulators of other small GTPases, including dominant-negative and constitutively-active forms of Cdc42 and Rac1, but they exhibited phenotypes apparently distinct from Pax6 mutant cells (data not shown). As these observations suggested that Pax6 might be involved in the Rho/ROCK pathway, we blocked the Rho/ROCK pathway in Pax6 mutant cells by treatment with Y27632 or by misexpression of C3, and quantified the number and the length of neurites in comparison with untreated mutant cells. We hypothesized that if the phenotype of Pax6 mutant cells was caused by loss of expression of factor(s) implicated in the Rho/ROCK pathway, there should be little additional change in the phenotypes of Pax6-deficiency by treatment with Y27632 or C3. However, the effects of Y27632 and C3 were additive in the Pax6 mutant cells; the number and the length of neurites were significantly augmented by treatment with Y27632 or C3 (Fig. 8D-H). Furthermore, misexpression of the constitutively-active form
of ROCK (Delta 3) had a comparable inhibitory effect on neurite formation in both wild-type and mutant cells (data not shown). These results suggested that the phenotypes caused by the absence of Pax6 function was unlikely to be mediated by a modification of endogenous ROCK activity.

DISCUSSION

In this study, we have demonstrated that Pax6 controls a genetic program involved in the polarization of cerebellar granule cells. In the Pax6 mutant rat rSey2/rSey2, granule cells displayed severe morphological abnormalities including multipolar neurites and random migration, despite apparently normal expression of a series of granule cell axonal differentiation markers. These results suggest that Pax6 acts at the level of cellular morphogenesis and migration rather than in regulation of the early steps of granule cell determination or differentiation. We also showed that cytoskeletal dynamics of mutant granule cells were apparently disorganized in a manner reminiscent of the inhibition of the ROCK pathway. To our knowledge, this is the first indication that Pax6 functions as an intrinsic factor regulating neuronal morphogenesis and migration via cytoskeletal control.

Cell-autonomous function of Pax6 during parallel fiber formation and tangential migration of granule cells

Pax6 mutant animals show a wide variety of neurological abnormalities in migration and axonal pathfinding of CNS neurons. However, most of the phenotypes characterized so far result from indirect effects of Pax6 mutation. In sharp contrast, granule cells require the intrinsic activity of Pax6 to properly form bipolar axons and for subsequent cell body migration, implicating Pax6-involvement in the transcriptional control required for polarization of axons. Disrupted formation of long bipolar neurites in mutant cells was rescued by forced expression of Pax6, but not by co-culturing with wild-type cells, indicating cell-autonomy of Pax6 function during granule cell morphogenesis. Combinatorial expression of different markers for granule cells and parallel fiber axons ensured that Pax6 mutant EGL cells were competent to differentiate as granule cells and to form parallel fiber axons. It is thus plausible that Pax6 is involved in polar elongation of axons after the initial stage of their formation.

Another striking feature of Pax6 mutant EGL cells is their unsteady zigzag migration path in culture. While wild-type EGL cells elongated long leading processes along preexisting radial axons, Pax6 mutant cells extended short leading processes in random orientations that often detached from the axons, as visualized by immunofluorescence against MAP2 and by TEM studies. Our co-culture experiments demonstrated that this disorganized migration of Pax6 mutant cells was not due to the absence of a radial neurite scaffold, but rather to a cell-autonomous defect of migrating cells. Thus, besides its role in axon elongation, Pax6 might also be involved in regulation of tangential migration of granule cells guided by leading processes that contact with preformed axon fibers. In contrast to the mutant cells in culture, the EGL cells that misexpress a dominant-negative Pax6 failed to move away from the site of their birth in developing cerebella. The appearance of the thickened EGL of mutant animals also suggests that tangential migration of premigratory EGL cells is disturbed in the absence of Pax6 function. This is not surprising if one considers that tangential migration of postmitotic EGL cells is guided by leading processes via contact-mediated association with pre-existing parallel fibers. Hence, densely packed EGL cells might offer massive...
resistance to the passage of Pax6-deficient cells that randomly move on the free surface of a culture dish.

Which step of parallel fiber formation – formation of the bipolar leading and trailing processes, tangential migration, or axon elongation – does Pax6 specifically regulate? As the leading and trailing processes of bipolar EGL cells are later transformed into parallel fiber axons (Ono et al., 1994), it is difficult to draw a sharp distinction between these continuous steps in granule cell morphogenesis. From our data, it is likely that Pax6 function is involved in axon elongation as well as during the initial outgrowth of bipolar neurites, as expression of Pax6 after 1 DIV was sufficient to rescue attenuated elongation of bipolar neurites in mutant cells (Fig. 5). Thus, the cell-autonomous function of Pax6 may be involved in sustaining the formation of bipolar processes that guide tangential migration and later elongate to form parallel fiber axons.

**Cellular mechanisms downstream of Pax6 function**

We also searched for the cellular mechanisms regulating granule cell morphogenesis impaired in Pax6 mutant cells. Formation of multiple axonal processes, abnormal lamellipodia and increase in the growth cone size in mutant cells strongly suggest an important role for Pax6 in the control of cytoskeletal dynamics during neurite formation. Apparent similarities in the phenotypes of Pax6 mutant cells with those evoked by inhibition of the Rho/ROCK activities prompted us to examine a functional interaction of Pax6 and the Rho/ROCK pathway. Quantitative analysis of the action of Y-27632 on Pax6 mutant cells excluded direct involvement of Pax6 in the ROCK pathway. In other words, it is unlikely that Pax6 regulates expression of signaling molecule(s) in direct association with ROCK. Some differences in the phenotypes of mutant cells and ROCK-inhibited cells were observed: a mild increase in axon length, presumably owing to early initiation of axon outgrowth, was seen in Y27632-treated cells but not in Pax6-mutant cells (Fig. 8). In fact, unlike Y27632 treatment, initiation of neurite formation was not advanced in mutant cells (data not shown). Nevertheless, several common morphological features between Pax6- and ROCK-deficient cells suggested that Pax6 might regulate cytoskeletal dynamics via a mechanism closely related to the ROCK pathway. One possibility to be tested in future studies is that Pax6 may be involved in a step maintaining neuronal polarity downstream of ROCK-mediated initial polarization of granule cells. The enhanced phenotypes induced by the Rho-inhibiting enzyme C3 are consistent with the idea that multiple co-existing Rho-dependent mechanisms regulate actin and/or other cytoskeletal dynamics (Kaibuchi et al., 1999; Bito et al., 2000; Bradke and Dotti, 2000). C3 expression yielded alterations in the number and the length of axons of wild-type and mutant cells to a similar plateau, implying that loss of Pax6 function might cause inactivation of a ROCK-independent Rho pathway. Alternatively, Pax6 might be involved in a cascade independent of Rho/ROCK activity, as we cannot rule out that phenotypes evaluated by the parameters we used in this study were saturated in C3-expressing cells.

It is noteworthy that in culture, Pax6 mutant neocortical radial glia and cerebellar granule cells exhibit similar morphological abnormalities (Götz et al., 1998): they form multipolar short processes instead of long bipolar processes of

**Fig. 7.** Aberrant formation of cytoskeletal structures in Pax6 mutant (rSey2/rSey2) EGL cells. (A,B) PKH26-labeled cells emigrating from a microexplant of E21 EGL observed at 1 DIV. Wild-type cells (A) have long straight neurites, while mutant cells (B) protrude short neurites, occasionally bifurcated and/or extruding lamellipodia in the midst of the shafts (arrow). (C,D) Scanning electron micrographs of microexplant cultures (3 DIV). Wild-type cells (C) possess thin growth cones at the tip, most of which simply taper off. Pax6 mutant cells (D) expand large growth cones with massive lamellipodia. (E,F) Distribution of F-actin (green) and microtubules (red) in growth cones. In comparison with wild type (E), an excess amount of F-actin accumulates in the large growth cones of Pax6 mutant EGL cells, accompanied by invasion of microtubules (F). (G) Quantification of growth cone size. Live cells labeled with PKH26 were observed at 1 DIV. The growth cone area was measured as the number of pixels in the graphic image. Data are indicated as mean±s.e.m. (P<0.005). Scale bars: 10 μm.
wild-type cells. This raises an intriguing possibility that Pax6 might have a general role in regulating cytoskeletal dynamics during polarization of neural cells in the CNS. Pax6 may activate transcription of the genes required for organization of cytoskeletal proteins in granule cells, which should act independently of the ROCK pathway. Identification of downstream target genes for Pax6 is in progress to reveal the molecular cascades that control the dynamic change of granule cell polarity during development.

Contribution of Pax6 in other aspects of cerebellar development

Pax6 expression is first detected in the EGL stem cells in the upper rhombic lip at as early as E14.5 in normal rat embryos (K. K. and M. K., unpublished). The EGL stem cells move rostrally from the upper rhombic lip along the surface of the cerebellar primordium to form the EGL by E19 in rodents. This raises the possibility that Pax6 is involved in the early phase of cerebellar development, including migration of the stem cells to populate the EGL. Previous studies have indeed shown that a subset of EGL cells is mislocated in the inferior colliculus in the Pax6 mutant mouse Sey/Sey (Engelkamp et al., 1999). This ectopic extension of the EGL may be the result of loss of Unc5h3 expression in the Sey/Sey mouse. However, in the rat rSey2/rSey2 mutants, we found no difference in the level of expression of netrin and its receptors, including Dcc, Unc5hs and neogenin, in the EGL (data not shown). Consistently, the rostral migration of EGL stem cells appeared normal in rats, suggesting that neither loss of Unc5h3 expression nor mislocation of EGL cells necessarily occur in the context of Pax6 deficiency. That the EGL was completely covered by granule cells in Pax6 mutant animals supports the notion that the initial migration of EGL stem cells does not require Pax6 function. Histological analysis could distinguish rostral migration of the EGL stem cells from tangential migration of premigratory EGL cells, in that the former is not guided by leading processes but resembles passive dispersal of round cells (Ryder and Cepko, 1994; Altman and Bayer, 1997; Komuro et al., 2001). It is intriguing that leading process-guided migration of lower rhombic lip cells to form precerebellar nuclei is also disrupted in Pax6 mutant animals (Engelkamp et al., 1999; Yee et al., 1999). Thus, the ability of Pax6 to regulate migration may be a mechanism common to neurons that use leading processes in the cerebellar system.

What is the role of early expression of Pax6 in the upper rhombic lip cells? Besides abnormal behavior of EGL cells, multiple morphological defects are observed in Pax6 mutant cerebellum, including mediolateral regionalization of the cerebellar primordium. Early expression of Pax6 might regulate these aspects independently of its action on postmitotic EGL cells.

Concluding remarks

Pax6 plays critical roles in numerous aspects of early patterning and cell-type specification in the CNS. Our present report pinpoints a novel function of Pax6 in the cell-autonomous control of cytoskeletal networks during the polarization of the CNS neuron. As could be speculated from its pleiotropic actions during development, downstream targets of Pax6 might well be context dependent within individual Pax6-expressing cells. It would thus be necessary to identify the specific target genes in the granule cell to clarify the molecular mechanisms regulating formation of their bipolar axons. Further studies will be needed to understand how widely
this Pax6-dependent mechanism is used in various cell types during polarization in the developing CNS.

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