Development of the cerebral cortex depends partly on neural activity, but the identity of the ion channels that might contribute to the activity-dependent cortical development is unknown. KCNK channels are critical determinants of neuronal excitability in the mature cerebral cortex, and a member of the KCNK family, KCNK9, is responsible for a maternally transmitted mental retardation syndrome. Here, we have investigated the roles of KCNK family potassium channels in cortical development. Knockdown of KCNK2, 9, or 10 by RNAi using in utero electroporation impaired the migration of late-born cortical excitatory neurons destined to become Layer II/III neurons. The migration defect caused by KCNK9 knockdown was rescued by coexpression of RNAi-resistant functional KCNK9 mutant. Furthermore, expression of dominant-negative mutant KCNK9, responsible for the disease, and electrophysiological experiments demonstrated that ion channel function was involved in the migration defect. Calcium imaging revealed that KCNK9 knockdown or expression of dominant-negative mutant KCNK9 increased the fraction of neurons showing calcium transients and the frequency of spontaneous calcium transients. Mislocated neurons seen after KCNK9 knockdown stayed in the deep cortical layers, showing delayed morphological maturation. Taken together, our results suggest that dysfunction of KCNK9 causes a migration defect in the cortex via an activity-dependent mechanism.

Keywords: activity dependent, calcium, channelopathy, electroporation, mental retardation

Introduction

Neural activity plays roles in the formation of neural circuits (Katz and Shatz 1996). In addition to its well-documented role in the refinement of circuits, some studies have shown that neural activity is also required for early neuronal development, including proliferation (Weissman et al. 2004; Liu et al. 2010), differentiation (Borodinsky et al. 2004), and migration (Komuro and Rakic 1993; Nakanishi and Okazawa 2006; Manent and Represa 2007; Bortone and Polleux 2009). In the developing cerebral cortex of rodents, studies using extracellular recording and calcium imaging have demonstrated that the immature cortical neurons show characteristic patterns of activity (Khazipov and Luhmann 2000; Allen and Cossart 2010). Such activity is thought to influence cortical development. Hypo- or hyperactivity during development may induce abnormality in the cortical architecture, which might be the basis for some developmental disorders (Ross and Walsh 2001; Ben-Ari 2008). These roles of neural activity in multiple developmental stages suggest that controlling neuronal activity is vital for the proper development of immature neurons.

The pattern of neural activity is regulated by ion channels. Developing cortical neurons express various ion channels, such as voltage-gated sodium channels, potassium channels, and calcium channels (Luhmann et al. 2000; Picken Bahrey and Moody 2003). KCNK family leak potassium channels, which have 4 transmembrane segments and 2 P-domains, are one of the major determinants of membrane potential (Goldstein et al. 2001). They conduct potassium current at resting membrane potential with little voltage dependence and are therefore called leak potassium channels. KCNK channels are critical determinants of neuronal excitability in the cortex (Goldstein et al. 2001; Day et al. 2005). In addition, a member of the KCNK family, KCNK9 (TASK3), is responsible for a maternally transmitted developmental disorder, Birk Barel mental retardation dysmorphism syndrome (Barel et al. 2008), and KCNK9 knockout mice show abnormalities in some cognitive functions such as working memory and spatial learning (Linden et al. 2007). Several members of the KCNK family (KCNK9, KCNK2/TREK1, and KCNK10/TREK2) are expressed in the developing cerebral cortex (Talley et al. 2001; Aller and Wisden 2008). These facts imply that KCNK9 is involved in cortical function and might also be involved in the formation of cortical neural circuits. However, whether dysfunction of KCNK9 affects cortical development has not yet been explored. In this study, we addressed this issue and found that knockdown of KCNK9 expression by RNAi affected migration of cortical neurons. Functional blockade of KCNK9 by overexpression of a dominant-negative, disease-associated mutant channel (Barel et al. 2008) also disturbed this migration. KCNK9 knockdown neurons showed more frequent calcium transients than control neurons. These results suggest that KCNK potassium channels contribute to the migration of cortical pyramidal neurons through stabilization of intracellular calcium concentration.

Materials and Methods

Animals

Wild-type ICR mice were used in all experiments. The experimental procedures were performed in accordance with the guidance for animal experimentation by the United States National Institutes of Health and Kyoto University and were approved by the local committee for handling experimental animals in the Graduate School of Science, Kyoto University.

Plasmid Construction

KCNK2, 9, and 10 shRNAs were obtained from Open Biosystems. Each shRNA was inserted into a pCAGmir30 vector (Matsuda and Cepko 2007). The sequences of KCNK9 shRNA2 and 3 were designed using the Hannon lab web site (http://cancan.cshl.edu/RNAi_central/main2.cgi) and were inserted into a pSuper vector under the H1

Dysfunction of KCNK Potassium Channels Impairs Neuronal Migration in the Developing Mouse Cerebral Cortex

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promoter (OligoEngine). The sequences of shRNAs used in this study are shown in Supplementary Table S1. The numbers of shRNAs initially obtained were 1 for KCNK2, 4 for KCNK9, and 1 for KCNK10. One KCNK9 shRNA did not effectively reduce the expression of KCNK9 protein in HEK293T cells, and therefore, was not used for further experiments. Mouse cDNA encoding KCNK9 was isolated from a cDNA library of adult mouse cortex by polymerase chain reaction. Mouse cDNAs encoding KCNK2 and 10 were purchased from Open Biosystems. Mutant and N-terminal EGFP-tagged KCNK channels were cloned into a pCAGplay vector (Kawaguchi and Hirano 2006). For western blot, EGFP was fused to N-terminus of each KCNK channel with a linker (Gly-Ser-Ala). The pCAsalEGFP vector (Horikawa and Takeichi 2001) was used for expression of enhanced green fluorescent protein (EGFP). GCaMP3 cDNA was purchased from Addgene (plasmid 22692) and inserted into pCAGplay vector. The sequences of primers to obtain mutant and EGFP-tagged KCNK channels are shown in Supplementary Table S3.

In Situ Hybridization
In situ hybridization was performed as previously described (Tagawa et al. 2005; Watakabe et al. 2007). KCNK2, 9, and 10 cDNAs were cut by restriction enzymes (KCNK2, BamHI/HincII; KCNK9, PstI; KCNK10, BamHI/PstI), and subcloned into pBluescript. Digoxigenin (DIG)-labeled RNA probes were transcribed by T3 or T7 RNA polymerases. Embryonic and neonatal mouse brains were cryoprotected in OCT compound (Sakura), and coronal sections of 20 μm were cut on a cryostat CM1850 (Leica). The sections were fixed with 4% paraformaldehyde for 10 min at room temperature, followed by washing twice in 2× SSC for 5 min at room temperature. The sections were incubated with 0.1 M triethanolamine for 3 min, then acetic anhydride was added, and the sections were incubated for 10 min at room temperature. The sections were hybridized with DIG-labeled RNA probes for 40 h at 65 °C in hybridization solution (50% formamid, 5× SSC, 5× Denhardt’s solution, 50 ng/ml yeast RNA), followed by washing with 2× SSC for 30 min at 65 °C, 0.5× SSC for 15 min at 65 °C, and 0.5× SSC for 15 min at room temperature. The sections were incubated with alkaline phosphatase-conjugated anti-DIG (Roche) at 4 °C overnight. The hybridization signal was detected with TSA-plus Fluorescein System (Perkin Elmer) for 10 min at room temperature. The sections were mounted with Vectashield containing DAPI (Vector).

In Utero Electroporation
In utero electroporation was performed as reported previously (Saito and Nakatsuji 2001; Tabata and Nakajima 2001; Mizuno et al. 2007). In brief, ICR mice at E15 were anesthetized with Somnopentyl (50 mg/kg; Kyoritsu Pharmacy), and their uteruses were exposed. Plasmid vectors were prepared at 0.2 mg/ml (KCNK9*, G95E*, 0.5 mg/ml (GCaMP3) or 1.0 mg/ml (others) with Trypan Blue (Sigma), and 1–2 μl DNA solution was injected into the right lateral ventricle. Next, 5 electric pulses of 50 V for 50 ms at 1 Hz were applied to the embryos using an electroporator CUY21EDiT (NepaGene).

Immunohistochemistry
Neonatal (P0–P5) and juvenile (P10, P15, and P35) mice were deeply anesthetized. Brains were fixed with 4% paraformaldehyde in phosphate buffer at 4 °C overnight, transferred to 30% sucrose in phosphatebuffer saline (PBS), and incubated at 4 °C overnight. Neonatal brains were cryoprotected in OCT compound (Sakura), and coronal sections of 50 μm were cut on a cryostat CM1850 (Leica). Fifty-micrometer-thick coronal sections of juvenile brains were cut on a freezing microtome LM2000R (Leica).

Immunohistochemistry was performed using a free-floating procedure. The sections were incubated with blocking solution (5% normal goat serum (Sigma), 0.2% Triton X-100 in PBS) for 1 h at room temperature, followed by incubation with primary antibodies in blocking solution at 4 °C overnight. Subsequently, the sections were incubated with secondary antibodies for 1 h at room temperature. The sections were mounted with Vectashield containing DAPI (Vector).

The following primary and secondary antibodies were used: anti-GFP (chick, polyclonal, 1:600, Millipore), anti-Cux1 (rabbit, polyclonal, 1:100, Santa Cruz Biotechnology), Ctip2 (rat, monoclonal, 1:600, Abcam), anti-Thr1 (rabbit, polyclonal, 1:600, Abcam), anti-Ki67 (rabbit, monoclonal, 1:1000, Thermo Scientific), anti-Nestin (mouse, monoclonal, 1:600, BD Pharmingen), anti-KCNK3 (1 mg/ml, Alomone Labs), Alexa 488-conjugated anti-chick IgG, Alexa 568-conjugated anti-rabbit IgG, anti-mouse IgG, and anti-rat IgG (made in goat, 1:1000, Molecular Probes).

Detection of Apoptosis with TUNEL Assay
Apoptosis was detected with an ApoTag® Red In Situ Apoptosis Detection Kit (Chemicon). The experimental procedure was based on the protocol that was provided by the manufacturer. Briefly, 50-μm-thick coronal cortical sections were incubated with ethanol-acetic acid for 5 min at room temperature. Sections were incubated with TdT enzyme and DIG-labeled dNTP for 30 min at 37 °C. Subsequently, sections were incubated with rhodamine-conjugated anti-DIG antibody for 30 min at room temperature. Finally, sections were mounted with Vectashield containing DAPI (Vector).

Microscopy and Morphological Analysis
Fluorescent images were acquired on a confocal laser-scanning microscope FV1000 (Olympus). For acquisition of images of in situ hybridization and analysis of neuronal migration, a x10 dry objective was used. Images of a single confocal plane were acquired for in situ hybridization. Confocal images of 50-μm-thick z-stacks were acquired for analysis of neuronal migration, and the z-series of images were projected onto 2D representations. The fraction of EGFP-positive neurons in each layer was calculated by dividing the number of EGFP-positive neurons in each cortical layer by that of total EGFP-positive neurons in the cortex. At least 10 sections (1 section per mouse) for each experimental condition were analyzed. For acquisition of images to analyze the cell fate, proliferation, apoptosis, and dendritic morphology, a x20 dry objective was used. Images of a single confocal plane were acquired for analysis of cell fate, proliferation, and apoptosis, and z-series of images were obtained for morphological analysis. z-Series of images were projected onto 2D planes. Dendrites were traced, and total length and branch points were analyzed with NeuroJ software (Meijering et al. 2004) (http://rsb.info.nih.gov/ij/). At least 5 neurons from 1 mouse were analyzed. Statistical analysis was performed with Student’s t-test or the Dunnett test. All results were expressed as mean ± SEM.

Western Blotting
Plasmids encoding EGFP-tagged KCNK2, 9, 10, KCNK9*, G95E*, or G236R* with or without shRNA expression vectors (2 μg each for) were transfected into HEK293T cells with Lipofectamine 2000 (Invi- trogen), based on the protocol that was provided by the manufacturer. Twenty-four hours after transfection, the cells were lysed in sample loading buffer (50 mM Tris–HCl, 4% SDS, 20% glycerol, 10 mM dithiothreitol, 0.001% bromophenol blue) and were denatured for 30 min at 37 °C. The samples were separated by 10% SDS-PAGE and transferred onto nitrocellulose membranes. The membranes were incubated with blocking solution (5% skim milk and 0.1% Tween-20 in TBS) for 1 h at room temperature, and incubated with primary antibody (mouse monoclonal anti-EGF, 1:1000, Nacalai Tesque; mouse monoclonal anti-β-actin, 1:1000, Sigma) in blocking solution overnight at 4 °C. The membranes were incubated with HRP-conjugated secondary antibody (donkey monoclonal anti-mouse IgG, 1:1000, Molecular Probes) in TBS for 1 h at room temperature. Signals were detected with an ECL Chemiluminescent kit (GE Healthcare). Images were acquired using LAS 3000 (Fuji Film).

Electrophysiology
Electrophysiological experiments were performed as previously described (Ohtsuki et al. 2004). In brief, neonatal mouse brains were removed, and coronal 300-μm-thick cortical slices were cut on a...
vibratome (linear slicer PRO 7, Dosaka EM) in chilled RDS solution (in mM; 130 NaCl, 4.5 KCl, 2 CaCl₂, 33 glucose, and 5 HEPES). Slices were recovered for longer than 1 h in Krebs’ solution saturated with 95% O₂ and 5% CO₂ (in mM; 124 NaCl, 26 NaHCO₃, 1.8 KCl, 1.24 KH₂PO₄, 2.5 CaCl₂, 1.3 MgCl₂, and 10 glucose) at room temperature. Oxygenized Krebs’ solution was used as extracellular bath solution. The glass patch pipettes of 4–7 MΩ were used for recording. The patch pipettes were filled with the internal solution (in mM; 140 n-glucuronic acid, 7 KCl, 155 KOH, 5 EGTA, 10 HEPES, 2 Mg-ATP, and 0.2 Na-GTP). The junction potential was offset. Whole-cell recordings from migrating neurons in intermediate zone of P1 cortex were performed at room temperature (22–24 °C) with a patch-clamp amplifier (EPC 9 or 10, HEKA). In voltage-clamp experiments, the membrane potential was held at −60 mV. Input resistance was measured by applying −20-mV voltage pulses, and voltage-dependent inward and outward currents were recorded by applying positive voltage pulses with a 10-mV step. Resting potential was recorded in a current-clamp condition. Statistical analysis was performed with Student’s t-test. All results were expressed as mean ± SEM.

**Calcium Imaging**

GCaMP3 cDNA with control, KCNK9 shRNA, or KCNK9 G236R were transfected by in utero electroporation. Coronal 500-μm-thick cortical slices of P1 mice were cut on a vibratome in chilled RDS solution. Slices were allowed to recover for longer than 1 h in Krebs’ solution saturated with 95% O₂ and 5% CO₂ at room temperature. Krebs’ solution was used as extracellular solution. Optical recording was performed at room temperature, using a fluorescent microscope BX50WI (Olympus) with a ×20 water-immersion objective and an EM-CCD camera C9100 (Hamamatsu Photonics). The sampling rate was 1 Hz. The mean fluorescent signal of a single cell was calculated, and divided by that at the start using Aquacosmos software (Hamamatsu Photonics). A fluorescence change of 5% above the baseline within 30 s was regarded as a calcium transient. Statistical analysis was performed with the χ²-test or Dunnett test. All results were expressed as mean ± SEM.

**Results**

**Expression Pattern of KCNK Potassium Channels in Developing Cortex**

Previous studies showed the expression of KCNK2, KCNK9, and KCNK10 in the developing cortex (Aller and Wisden 2008), but their laminar distributions were unclear. As a first step to study the roles of KCNK potassium channels in cortical development, we examined the expression patterns of these potassium channels in the developing cortex by in situ hybridization.

KCNK2 was expressed in the cortical plate (CP), intermediate zone (IZ), subventricular zone (SVZ), and ventricular zone (VZ) at embryonic day 15 (E15) and E17. At postnatal day 0 (P0), KCNK2 was strongly expressed in the marginal zone (MZ), IZ, SVZ, VZ, and lower CP, but not in upper CP. It was expressed in Layers I, V, and VI, but was not expressed in Layers II/III or IV at P3 (Fig. 1A). KCNK9 was expressed in MZ, CP, IZ, SVZ, and VZ at E15 and 17. At P0 and P3, KCNK9 was expressed in all cortical layers except Layer I. The expression level of KCNK9 in CP increased after birth (Fig. 1B). KCNK10 was most highly expressed at E15. KCNK10 was strongly expressed in MZ, IZ, SVZ, and VZ and was weakly expressed in CP. At P0, KCNK10 was expressed in MZ, lower CP, IZ, SVZ, and VZ. KCNK10 was weakly expressed in the lower cortical layers at P3 (Fig. 1C). Taken together, these findings indicate that KCNK2, 9, and 10 were expressed in SVZ and VZ at E15, and in IZ, SVZ, and VZ at E17 and P0, suggesting that developing cortical neurons, including Layer II/III neurons, express these KCNK potassium channels in the late embryonic and early postnatal stages.

**Neuronal Migration was Impaired by Knockdown of KCNK Potassium Channels**

To determine whether KCNK potassium channels are involved in the cortical development, we performed RNAi experiments using in utero electroporation. Short hairpin RNAs (shRNAs) targeting KCNK2, 9 and 10 effectively inhibited the expression of each KCNK channel protein in HEK293T cells, whereas control shRNA did not affect the translation of mRNAs for these genes (Fig. 2A). A mutant variant of KCNK9 shRNA was used as a control shRNA (Supplementary Table S1).

The control or KCNK channel shRNA was then transfected into cortical Layer II/III progenitor cells together with EGFP. Coronal cortical sections including visual, auditory, or somatosensory cortices were prepared at P0, P3, and P5, and the distribution of EGFP-positive neurons was examined. At P0, about half of EGFP- and control shRNA-expressing neurons were observed in IZ, SVZ, and VZ, and some EGFP-positive neurons were observed in the upper CP. The distributions of EGFP-positive neurons were not altered by knockdown of KCNK channels at P0 (Fig. 2B, Supplementary Table S2). Most neurons expressing control shRNA had arrived at the superficial layer by P3 (Fig. 2B, Supplementary Table S2). In contrast, only a fraction of neurons expressing KCNK channel shRNAs reached Layer II/III at P3 (Fig. 2B, Supplementary Table S2). KCNK2 shRNA, 8.3 ± 3.1%, n = 12, P < 0.001, Dunnett test; KCNK9 shRNA, 24.5 ± 2.1%, n = 10, P < 0.001; KCNK10 shRNA, 47.7 ± 4.5%, n = 11, P < 0.001). A decrease in the fraction of neurons in IZ/SVZ/VZ between P0 and P3 was small in KCNK knockdown mice (Fig. 2B, Supplementary Table S2, control shRNA P0, 52.3 ± 2.7%, n = 12; P3, 10.1 ± 3.6%, n = 11; KCNK2 shRNA P0, 50.3 ± 3.0%, n = 11; P3, 41.9 ± 2.3%, n = 11; KCNK9 shRNA P0, 46.2 ± 2.1%, n = 11; P3, 47.1 ± 4.3%, n = 10; KCNK10 shRNA P0, 47.5 ± 1.5%, n = 12; P3, 39.8 ± 3.8%, n = 11), suggesting that an invasion into the CP was decelerated by knockdown of KCNK channels. At P5, some neurons expressing KCNK9 shRNA still remained in deep cortical layers (Fig. 2B, Supplementary Table S2, control shRNA P0, 3.6 ± 1.3%, n = 14; KCNK2 shRNA, 1.9 ± 0.6%, n = 12, P = 0.698, Dunnett test; KCNK9 shRNA, 9.8 ± 1.8%, n = 10, P = 0.002; KCNK10 shRNA, 2.3 ± 1.1%, n = 12, P = 0.858) and white matter (Fig. 2B, Supplementary Table S2, Control shRNA, 10.4 ± 3.7%, n = 14; KCNK2 shRNA, 8.3 ± 3.1%, n = 12, P = 0.977, Dunnett test; KCNK9 shRNA, 38.0 ± 2.3%, n = 10, P < 0.001; KCNK10 shRNA, 8.2 ± 2.7%, n = 12, P = 0.975), but most EGFP-positive neurons expressing KCNK2 or 10 shRNA had reached superficial layer (Fig. 2B, Supplementary Table S2). Control shRNA P0, 86.0 ± 4.6%, n = 14; KCNK2 shRNA, 89.9 ± 3.5%, n = 12, P = 0.872, Dunnett test; KCNK9 shRNA 52.2 ± 2.4%, n = 10, P < 0.001; KCNK10 shRNA, 89.5 ± 2.6%, n = 12, P = 0.906). Thus, knockdown of KCNK2, 9, or 10 impaired neuronal migration at P3, but only KCNK9 knockdown significantly impaired neuronal migration at P5. Knockdown of these 3 members of KCNKs (KCNK2, 9, and 10) together impaired neuronal migration as severely as KCNK9 knockdown (Fig. 2B, Supplementary Table S2, Layer II/III,
54.8 ± 4.2%, P < 0.001; Layers IV–VI, 13.8 ± 1.7%, P < 0.001; WM, 31.5 ± 4.4%, P < 0.001, compared with control shRNA, n = 14). These results suggest that KCNK family leak potassium channels are involved in the migration of cortical Layer II/III neurons. We focused on the function of KCNK9 in neuronal migration and performed further analysis of it, because knockdown of KCNK9 most clearly impaired neuronal migration.

**Ion Channel Function of KCNK9 was Required for Neuronal Migration**

We next confirmed that the impairment of migration by KCNK9 shRNA expression was not caused by an off-target effect. Two other shRNAs for KCNK9, whose target sequences were different from that of KCNK9 shRNA, also inhibited the expression of KCNK9 (Fig. 3A). These shRNAs impaired neuronal migration (Fig. 3B, C, Supplementary Table S2, Layer II/III, pSuper, 94.1 ± 1.3%, n = 12 sections; KCNK9 shRNA2, 18.3 ± 1.4%, n = 11, P < 0.001, Dunnett test; KCNK9 shRNA3, 28.8 ± 1.6%, n = 11, P < 0.001). Next, we attempted to rescue the phenotype by expressing mutant KCNK that was resistant to shRNA. We introduced mutations that disrupted the target sequence of KCNK9 shRNA but that did not alter the amino acid sequence. Indeed, the expression of this mutant KCNK9 (KCNK9*) was not inhibited by KCNK9 shRNA (Fig. 3D). The impairment of neuronal migration was partially rescued by coexpression of KCNK9* with KCNK9 shRNA (KCNK9 rescue) at P3 (Fig. 3E, F, Supplementary Table S2, Layer II/III, KCNK9 shRNA, 35.9 ± 2.7%, n = 11 sections; KCNK9 rescue, 65.4 ± 3.0%, n = 11, P < 0.001, Dunnett test). These results suggest that the impairment of neuronal migration was caused by knockdown of KCNK9.

To examine further the role of KCNK9 in neuronal migration, we used 2 KCNK9 mutant channels. Previous studies showed that glycine 95 at the potassium-selective filter

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**Figure 1.** (A–C) Expression of KCNK potassium channels during cortical development. Representative images of in situ hybridization showing laminar expression patterns of KCNK2 (A), 9 (B), and 10 (C) in the developing mouse cortex from E15 to P3. (D) DAPI staining shows the location of nuclei. Scale bar: 100 μm. MZ, marginal zone; CP, cortical plate; IZ, intermediate zone; SVZ, subventricular zone; VZ, ventricular zone.
and glycine 236 at the ion channel pore are crucial for the ion channel function of KCNK9. The mutants G95E and G236R are nonfunctional when expressed alone, and act as dominant-negative proteins when expressed with the wild-type channel (Pei et al. 2003; Barel et al. 2008) (Fig. 4A). We performed electroporation with these mutant channels. Neuronal migration was impaired at P3 by the expression of these dominant-negative mutants (Fig. 4B, C, Supplementary Table S2, Layer II/III, EGFP, 84.6 ± 3.6%, n = 10; G95E, 54.6 ± 3.4%, n = 12, P < 0.001, Dunnett test; G236R, 38.5 ± 1.1%, n = 11, P < 0.001). This result supported our findings from RNAi experiments and suggested that the expression and/or function of KCNK9 were critical for neuronal migration.

The crucial role of channel function of KCNK9 in the migration was suggested by another set of experiments. As mentioned above, migration defect caused by KCNK9 knockdown could be rescued by coexpression of RNAi-resistant functional KCNK9 (KCNK9*) channel (Fig. 3E, F, Supplementary Table S2). However, migration defect caused by KCNK9 knockdown was not recovered by coexpression of RNAi-insensitive nonfunctional G95E or G236R mutant (Fig. 4E, F, Supplementary Table S2). Thus, the ion channel function of KCNK9 is necessary for the normal neuronal migration.
The requirement for ion channel function of KCNK9 for neuronal migration suggests that knockdown of KCNK9 might have altered electrophysiological properties of migrating neurons. It was shown that KCNK9 is critically involved in the regulation of resting membrane potential in cerebellar granule neurons (Brickley et al. 2007). We performed whole-cell patch-clamp recording from migrating cortical neurons expressing control shRNA or KCNK9 shRNA. Knockdown of KCNK9 significantly increased input resistance (Fig. 5A, B, control shRNA, 3.3 ± 0.7 GΩ, n = 16 neurons; KCNK9 shRNA, 9.7 ± 2.0 GΩ, n = 17 neurons, P = 0.004, Student’s t-test). KCNK9 knockdown also significantly depolarized the resting membrane potential (Fig. 5C, control shRNA, −58.2 ± 5.1 mV, n = 16 neurons; KCNK9 shRNA, −40.9 ± 3.1 mV, n = 17 neurons, P = 0.007, Student’s t-test). On the other hand, KCNK9 knockdown did not affect the voltage-dependent transient inward current (control shRNA, −90.1 ± 24.6 pA, n = 16 neurons; KCNK9 shRNA, −91.8 ± 21.3 pA, n = 17 neurons at −10 mV, P = 0.960, Student’s t-test) or sustained outward current (control shRNA, 197 ± 27.8 pA, n = 16 neurons; KCNK9 shRNA, 159 ± 11.2 pA, n = 17 neurons at 0 mV, P = 0.227, Student’s t-test) induced by depolarizing voltage steps (Fig. 5D–F). These results are consistent with the known roles of KCNK channels (Goldstein et al. 2001) and suggest that KCNK9 contributes to the resting potassium permeability of membranes in migrating cortical neurons.

Downregulation of KCNK9 Increased the Frequency of Spontaneous Calcium Transients

Neuronal migration is partly regulated by intracellular calcium (Zheng and Poo 2007). We thought that KCNK9 knockdown or functional blockade might alter the pattern of spontaneous calcium transients. To test this, we transfected a genetically encoded calcium indicator, GCaMP3 (Tian et al. 2009), with control, KCNK9 shRNA or a dominant-negative mutant, G236R, and performed calcium imaging from migrating neurons at P0/P1 (Fig. 6). A small fraction (ca. 10%) of neurons expressing control shRNA showed calcium transients at a frequency of about 1 event/10 min (Fig. 6A, B). We compared the pattern of spontaneous calcium transients between control and KCNK9 knockdown neurons in both IZ and lower CP. The fraction of neurons expressing KCNK9 shRNA that showed calcium transients was more than that of control neurons at P1 (P0 IZ, control shRNA, 7.1%, 18 of 254 neurons; KCNK9 shRNA, 11.7%, 15 of 128 neurons, P = 0.128, χ²-test; P0 lower CP, control shRNA, 10.7%, 15 of 140 neurons; KCNK9 shRNA, 12.6%, 27 of 214 neurons, P = 0.588; P1 IZ, control shRNA, 11.0%, 19 of 173 neurons; KCNK9 shRNA, 25.5%, 39 of 153 neurons, P < 0.001; P1 lower CP, control shRNA, 9.2%, 7 of 76 neurons; KCNK9 shRNA, 25.2%, 37 of 147 neurons, P = 0.005). At P0, the frequency of spontaneous calcium transients in KCNK9 knockdown neurons was higher than that in control neurons in IZ, but not in
lower CP (Fig. 6A, B, IZ, control shRNA, 1.2 ± 0.1 events/10 min, \( n = 18 \) neurons; KCNK9 shRNA, 2.7 ± 0.7 events/10 min, \( n = 15 \) neurons, \( P = 0.024 \), Student’s \( t \)-test; lower CP, control shRNA, 1.3 ± 0.1 events/10 min, \( n = 15 \) neurons; KCNK9 shRNA, 1.6 ± 0.2 events/10 min, \( n = 27 \) neurons, \( P = 0.191 \)). At P1, the frequency of spontaneous calcium transients in
KCNK9 knockdown neurons was higher than that in control neurons both in IZ and lower CP (Fig. 6A, B, IZ, control shRNA, 1.3 ± 0.1 events/10 min, n = 19 neurons; KCNK9 shRNA, 2.9 ± 0.5 events/10 min, n = 39 neurons, P = 0.002; lower CP, control shRNA, 1.1 ± 0.1 events/10 min, n = 7 neurons; KCNK9 shRNA, 2.1 ± 0.2 events/10 min, n = 37 neurons, P = 0.001, Student’s t-test). Thus, KCNK9 knockdown increased the frequency of spontaneous calcium transients in migrating cortical neurons.

Is there any correlation between an increase in calcium transients and the migration defect? To test this, we carried out another set of experiments in which we compared the pattern of calcium transients in neurons expressing control shRNA, KCNK9 shRNA, or KCNK9 rescue with RNAi-insensitive KCNK9 (KCNK9*) (Fig. 6C, D). The increase in the frequency of calcium transients caused by KCNK9 knockdown was partially “rescued” by coexpression of KCNK9* (Fig. 6C, D, KCNK9 rescue, fraction of neurons, 47 of 138 neurons; frequency, 2.0 ± 0.2 events/10 min, n = 47 neurons, P = 0.012, compared with KCNK9 shRNA, P = 0.471, compared with control shRNA, Dunnett test). Thus, coexpression of RNAi-insensitive KCNK9 with KCNK9 shRNA partially restored both the pattern of calcium transients (Fig. 6C, D) and the migration phenotype (Fig. 3E, F).

We also examined effects of the expression of a disease-associated, dominant-negative mutant (G236R) of KCNK9. The fraction of neurons showing calcium transients was significantly higher in dominant-negative mutant
transfected mice (GCaMP3, 4.7%, 13 of 279 neurons, G236R, 33.9%, 57 of 168 neurons, P < 0.001, χ²-test). Furthermore, the frequency of calcium transients was higher in neurons expressing G236R than in control neurons showing calcium transients (Fig. 6E,F, GCaMP3, 1.4 ± 0.2 events/10 min, n = 13 neurons, G236R, 3.1 ± 0.3 events/10 min, n = 57 neurons, P < 0.001, Student’s t-test). These results together suggest that dysfunction of KCNK9 influences the pattern of calcium transients in migrating neurons.

**Cell Fate, Neuronal Proliferation, and Apoptosis Were not Affected by Knockdown of KCNK9**

Next, we performed immunohistochemistry to examine whether the cell fate of neurons that failed to migrate was changed or not. Previous studies identified several genes expressed in a layer-specific manner, and some genes have been reported to be determinants of the neuronal subtype specification in the cerebral cortex (Molyneaux et al. 2007). Tbr1 is a determinant of corticothalamic neurons in Layer VI and preplate (Hevner et al. 2001). Ctip2 promotes the differentiation of Layer V corticospinal motor neurons (Arlotta et al. 2005). Cux1 is expressed in Layers II/III and IV, but is not expressed in the other layers (Nieto et al. 2004).

Coronal cortical sections were stained with antibodies against Cux1, Ctip2, or Tbr1. Neurons expressing EGFP and control shRNA were Cux1-positive, consistent with previous studies (Fig. 7A). KCNK9 shRNA-expressing neurons that reached Layer II/III were also Cux1-positive (Fig. 7B). KCNK9 shRNA-expressing neurons that failed to migrate also expressed Cux1, but did not express Ctip2 or Tbr1 (Fig. 7B). These results suggest that cell fate specification is not affected by KCNK9 knockdown.

Previous studies have shown a role of KCNK9 in the development of cerebellar granule neurons, in which this channel is highly expressed (Talley et al. 2001; Han et al. 2002) and involved in potassium-dependent apoptosis (Lauritzen et al. 2003). In addition, KCNK9 has been implicated in cell division in tumor cells (Mu et al. 2003; Pei et al. 2003). We performed immunostaining against Ki67, which is a marker for cell division, and TUNEL staining for detection of apoptotic cells at E17. The fraction of EGFP/Ki67-double-positive cells (Fig. 7C,D, control shRNA, 9.3 ± 1.2%; KCNK9 shRNA, 10.2 ± 1.9%, n = 12 sections for each, P = 0.691, Student’s t-test) and the number of TUNEL-positive cells (Fig. 7E,F, control shRNA, 5.3 ± 0.7 cells; KCNK9 shRNA, 5.6 ± 0.8 cells, n = 12 sections for each, P = 0.755, Student’s t-test) were not significantly different between control and KCNK9 shRNA (Fig. 7C–F). In addition, immunostaining against Nestin showed that radial glial scaffold was not affected by KCNK9 knockdown (Fig. 7G). These results suggest that knockdown of KCNK9 does not affect neuronal proliferation, apoptosis, and radial glial scaffold.

**KCNK9 Knockdown Impeded Morphological Maturation**

What is the fate of neurons showing a migration defect during the neonatal period? Do they stay in deep cortical layers and...
mature, or eventually disappear? To answer these questions, we examined the laminar distribution of EGFP-positive neurons expressing KCNK9 shRNA at P10, P15, and P35. At P35, the fraction of EGFP-positive neurons in WM was slightly smaller than that at P10 (Fig. 8A,B, Supplementary Table S2, P10, 12.1 ± 1.8%, n = 11 sections; P15, 8.7 ± 2.0%, n = 15, P = 0.322, Dunnett test; P35, 4.1 ± 1.3%, n = 10, P = 0.014), but the fraction of EGFP-positive neurons in Layer II/III and Layers IV–VI did not significantly differ among these ages (Fig. 8A,B, Supplementary Table S2, Layer II/III, P10, 77.9 ± 2.7%, n = 11; P15, 80.7 ± 2.4%, n = 15, P = 0.635, Dunnett test; P35, 83.6 ± 2.4%, n = 10, P = 0.243, Layers IV–VI, P10, 10.0 ± 1.4%; P15, 10.6 ± 1.1%, P = 0.926; P35, 12.6 ± 1.3%, P = 0.291). At P35, some of neurons in WM may have moved into the cortex, not have died, because the number of EGFP-positive cells did not significantly change among ages (P10, 146.0 ± 20.0 neurons; P15, 133.0 ± 10.7 neurons P = 0.786, Dunnett test; P35, 152.5 ± 20.1 neurons, P = 0.948). These results suggest that many of the mislocated neurons resulting from KCNK9 knockdown stay in deep cortical layers during development. At P10, many EGFP-positive neurons in deep cortical layers had apical protrusions with few branches, reminiscent of the leading process of typical migrating neurons (Fig. 8C–F, branch points, 1.2 ± 0.3; total length, 287.3 ± 25.7 μm, n = 34 neurons). At P15, these neurons had some branched dendrites (Fig. 8D–G, branch points, 3.3 ± 0.7, P = 0.019, Dunnett test; total length, 384.7 ± 42.5 μm, P = 0.348, n = 23 neurons), and at P35, they had complex dendritic arbors, similar to those of mature neurons (Fig. 8E–G, branch points, 10.9 ± 0.8, P < 0.001; total length, 2081.0 ± 133.0 μm, P < 0.001, n = 27 neurons). These observations suggest that dendritic maturation is delayed in neurons that failed to migrate due to KCNK9 knockdown, and that these neurons somehow mature by P35 in deep cortical layers.

**Discussion**

We have demonstrated that KCNK potassium channels, especially KCNK9, play an important role in neuronal migration in the developing cerebral cortex. Knockdown of KCNK9 in late-born cortical neurons by RNAi resulted in a migration defect. Expression of the dominant-negative mutant channel G236R, found in patients with an inherited mental retardation (Barel et al. 2008), also caused a migration defect in the cortex, suggesting that ion channel function is required for neuronal migration. Electrophysiological experiments showed that knockdown of KCNK9 depolarize resting membrane potential. Knockdown or functional blockade of KCNK9 with the dominant-negative mutant channel G236R, increased the frequency of spontaneous calcium transients. Taken together, these results suggest that dysfunction of KCNK9 impairs migration of cortical excitatory neurons in an activity-dependent mechanism.

**Dysfunction of KCNK9 can Cause a Migration Defect in the Cortex**

Our data, obtained from RNAi experiments using 3 independent shRNAs, rescue experiments, and functional blockade experiments using dominant-negative mutants, indicate that a deficit in the expression/function of KCNK9 can cause a migration defect in the cortex. Earlier studies reported that KCNK9 knockout mice showed abnormality in some cognitive functions and in the regulation of cellular excitability in mature cerebellar granule neurons (Brickley et al. 2007; Linden et al. 2007). It is unclear whether migration of cortical excitatory neurons is impaired in KCNK9 knockout mice, because these studies did not focus on neuronal migration in the cortex. There are several cases where the extent of migration defects caused by genetic knockout is not comparable to that by in utero knockdown (Bai et al. 2003; Chen et al. 2008; Ip et al. 2011), and in the genetic knockout mice,
compensatory mechanisms by other family members have been suggested (Corbo et al. 2002; Deuel et al. 2006; Koizumi et al. 2006). In utero knockdown can be a useful approach for identifying genetic factors responsible for abnormality in cortical development, and for further investigation on how loss-of-function of target molecules affects cortical neuron development.

Developing cortical neurons express several members of KCNK potassium channels (Aller and Wisden 2008). When one of these (e.g., KCNK9) is knocked down, other members (e.g., KCNK2 and/or 10) may compensate for the reduced expression. However, we did not observe an additive effect of multiple knockdown: the distribution of EGFP-positive neurons in KCNK2/9/10 knockdown cortex was comparable with that in single knockdown of KCNK9 at P0, P3, and P5 (Fig. 2B,C, Supplementary Table S2). KCNK9 may play a dominant role in neuronal migration in the developing cortex.

It is reported that KCNK3 (TASK1) can form heterodimer with KCNK9 in cerebellar granule cells and motoneurons and modulate its ion channel function (Berg et al. 2004, Kang et al. 2004). There is a possibility that KCNK3 may be involved in KCNK9-dependent regulation of neuronal migration in the developing cerebral cortex. However, a previous report has shown no or low expression of KCNK3 mRNA in the developing cerebral cortex (Aller and Wisden 2008). Our immunohistochemical experiments using KCNK3-specific antibody (Toyoda et al. 2010) showed no apparent expression of KCNK3 in the developing cortex, and we did not observe compensatory upregulation of KCNK3 expression in KCNK9 knockdown cortex (data not shown).

**KCN9 Contributes to Neuronal Migration Through the Regulation of Spontaneous Calcium Transients**

How does dysfunction of KCNK9 impede migration? Our results suggest a possibility that downregulation of KCNK9 disturbed neuronal migration by increasing spontaneous calcium transients. Experiments using calcium imaging demonstrated that KCNK9 knockdown or expression of a disease-associated mutant (which behaves as a dominant-negative mutant) increased the frequency of calcium transients in migrating cortical neurons (Fig. 6). Coexpression of RNAi-insensitive KCNK9 with KCNK9 shRNA restored the pattern of calcium transients and the migration phenotype. It is known that neuronal migration is in part regulated by neural activity and calcium signaling (Komuro and Rakic 1992, 1993). Cortical interneuron migration is also sensitive to changes in the intracellular calcium concentration that are regulated by GABA_A and AMPA/NMDA receptors (Bortone and Polleux 2009). In both types of neuron, the frequency of calcium transients positively correlates with the rate of migration, and loss of calcium transients acts as a “stop” signal for migration (Komuro and Rakic 1996; Kamada and Komuro 2004; Bortone and Polleux 2009). Their results and ours suggest that cortical excitatory neurons may differ from cerebellar granule neurons and cortical interneurons, in that they require relatively low levels or low frequencies of calcium transients for proper migration. Similar cell-type-specific effect of calcium transients on the motility has been reported in growing axons, where the frequency of calcium transients in growth cones are inversely or directly proportional to the rate of axonal outgrowth (Gomez and Spitzer 1999; Hutchins et al. 2011). To further clarify the relationship between calcium transients and cell motility during migration of cortical excitatory neurons, a combination of calcium and time-lapse imaging would be useful in future.

**KCN9 and Developmental Disorders**

A dominant-negative mutant of KCNK9 (G236R) was found in patients suffering from an inherited developmental disorder, Birk Barel mental retardation dysmorphism syndrome (Barel et al. 2008). In addition, it has been reported that KCNK9 knockout mice show abnormities in some learning tasks (Linden et al. 2007). These studies support the notion that dysfunction of KCNK9 causes a developmental disorder. We found that dysfunction of KCNK9 disrupted neuronal migration in the developing cerebral cortex. This result suggests the possibility that suppressed neuronal migration by KCNK9 dysfunction could be a cause of mental retardation. Indeed, migration defects are closely coupled with some developmental disorders in humans (Ross and Walsh 2001; LoTurco and Bai 2006; Ben-Ari 2008). For instance, microtubule-associated proteins DCX and Lis1 are responsible for cortical malformations such as subcortical band heterotopia and lissencephaly, and their knockdown in an animal model disrupts neuronal migration (Bai et al. 2003; Tsai et al. 2005). Interestingly, in DCX knockdown cortex, heterotopic
neurons located in the deep cortical layer and normally migrated Layer II/III neurons form an abnormal neural network, and this abnormal network becomes a source of synchronous epileptogenic firings, reproducing some symptoms present in human patients (Ackman et al. 2009). We have shown that mislocated neurons, resulting from KCNK9 knockdown, stayed and slowly matured in the deep cortical layers in early postnatal period, which includes the period during which large-scale dynamic calcium transients and propagating waves have been observed in the cortex (Garaschuk et al. 2000; Khasipov and Luhmann 2006; Allene and Cossart 2010). Dysfunction of KCNK9 may alter the pattern of such network activities, possibly through the regulation of excitability of individual neurons or the mislocated neurons abnormally integrated into the cortical circuits.

In conclusion, we have demonstrated that KCNK potassium channels play roles in the migration of cortical excitatory neurons. KCNK potassium channels promote migration by suppressing the frequency of calcium transients to keep it within the appropriate range.

**Supplementary Material**

Supplementary material can be found at: http://www.cercor.oxfordjournals.org/.

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**References**


