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Dynamic Changes in Ultrastructure of the Primary Cilium in Migrating Neuroblasts in the Postnatal Brain

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New neurons, referred to as neuroblasts, are continuously generated in the ventricular-subventricular zone of the brain throughout an animal's life. These neuroblasts are characterized by their unique potential for proliferation, formation of chain-like cell aggregates, and long-distance and high-speed migration through the rostral migratory stream (RMS) toward the olfactory bulb (OB), where they decelerate and differentiate into mature interneurons. The dynamic changes of ultrastructural features in postnatal-born neuroblasts during migration are not yet fully understood. Here we report the presence of a primary cilium, and its ultrastructural morphology and spatiotemporal dynamics, in migrating neuroblasts in the postnatal RMS and OB. The primary cilium was observed in migrating neuroblasts in the postnatal RMS and OB. The primary cilium was observed in migrating neuroblasts in the postnatal RMS and rostral migration toward the OB. Serial section transmission electron microscopy revealed that each migrating neuroblast possesses either a pair of centrioles or a basal body with an immature or mature primary cilium. Using immunohistochemistry, live imaging, and serial block-face scanning electron microscopy, we demonstrate that the localization and orientation of the primary cilium are altered depending on the mitotic state, saltatory migration, and deceleration of neuroblasts. Together, our results highlight a close mutual relationship between spatiotemporal regulation of the primary cilium and efficient chain migration of neuroblasts in the postnatal brain.

Key words: electron microscopy; live imaging; neuronal migration; olfactory bulb; primary cilium; rostral migratory stream

Significance Statement

Immature neurons (neuroblasts) generated in the postnatal brain have a mitotic potential and migrate in chain-like cell aggregates toward the olfactory bulb. Here we report that migrating neuroblasts possess a tiny cellular protrusion called a primary cilium. Immunohistochemical studies with zebrafish, mouse, and monkey brains suggest that the presence of the primary cilium in migrating neuroblasts is evolutionarily conserved. Ciliogenesis in migrating neuroblasts in the rostral migratory stream is suppressed during mitosis and promoted after cell cycle exit. Moreover, live imaging and 3D electron microscopy revealed that ciliary localization and orientation change during saltatory movement of neuroblasts. Our results reveal highly organized dynamics in maturation and positioning of the primary cilium during neuroblast migration that underlie saltatory movement of postnatalborn neuroblasts.

Introduction

The ventricular-subventricular zone (V-SVZ), which lines the lateral walls of the lateral ventricles in the postnatal mammalian brain, harbors neural stem cells that continuously produce immature new neurons (neuroblasts) (Obernier and Alvarez-Buylla, 2019). These neuroblasts are able to divide and migrate in chain-like aggregates toward the olfactory bulb (OB) through the rostral migratory stream (RMS) (Luskin, 1993; Lois and Alvarez-Buylla, 1994; Lois et al., 1996), and toward the lesioned cortex or striatum following brain injury (Arvidsson et al., 2002; Parent et al., 2002; Yamashita et al., 2006; Kreuzberg et al., 2010). Furthermore, the morphology of migrating neuroblasts is highly dynamic and controlled by multiple extrinsic and intrinsic cues (Khodosevich et al., 2009; Belvindrah et al., 2011; Nakamuta et al., 2017; Fujikake et al., 2018; Sawada et al., 2018). However, how ultrastructural features, such as the fine morphology of neurites and the position of internal organelles, contribute to neuroblast migration is not yet fully understood.

The primary cilium is a tiny cellular protrusion that contains signaling receptors on its membrane (Guemez-Gamboa et al., 2014; Malicki and Johnson, 2017), and regulates neural stem cell proliferation (Breunig et al., 2008; Han et al., 2008; Amador-Arjona et al., 2011; Tong et al., 2014) and neuronal maturation (Kumamoto et al., 2012; Luo et al., 2015) in the postnatal brain. Primary cilia are reportedly present in cultured medial ganglionic eminence (MGE)-derived interneurons and cerebellar granule neurons (Baudoin et al., 2012; Trivedi et al., 2014). Both MGEderived interneurons and V-SVZ-derived neuroblasts show saltatory movement, in which the neuron extends a leading process, forms a swelling, and then translocates its soma toward the swelling (Bellion et al., 2005; Schaar and McConnell, 2005). The primary cilium extends from the basal body (Baudoin et al., 2012) and moves forward into the proximal leading process before somal translocation during saltatory movement (Trivedi et al., 2014). KO mice for the ciliary intraflagellar transport (IFT) molecules Kif3a or Ift88, or the ciliary small GTPase Arl13b, showed defective migration, morphogenesis, and connectivity of MGEderived interneurons (Baudoin et al., 2012; Higginbotham et al., 2012; Guo et al., 2017). Kif3a knockdown (KD) also disrupted migration and differentiation of embryonic pyramidal neurons (Chen et al., 2019). However, the primary cilium in postnatal-

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born neuroblasts migrating in chains in the RMS has not been investigated.

Primary cilium ultrastructure and dynamics in mature cells have been intensively studied. Transmission electron microscopy (TEM) analyses have revealed ultrastructural features of the primary cilium, including nine pairs of microtubules (MTs) (the axoneme) and a basal body with transition fibers and basal feet (Sorokin, 1962). Time-lapse imaging studies using fluorescent protein-fused Arl13b have demonstrated the dynamic behavior of the primary cilium (Higginbotham et al., 2012; Trivedi et al., 2014; Ford et al., 2018). In addition to these methodologies, new EM techniques that acquire high-resolution ultrastructural images in three dimensions have recently been used for primary cilium analysis (Paridaen et al., 2013; David et al., 2014; Buskin et al., 2018; Insinna et al., 2019). Serial block-face scanning electron microscopy (SBF-SEM) is one of the 3D-EM techniques that enable analysis of both cell morphology and subcellular organelles simultaneously at the ultrastructure level (Briggman and Bock, 2012; Denk et al., 2012; Ohno et al., 2015). SBF-SEM has been applied to the study of morphological changes in the primary cilium during mitosis in embryonic neuroepithelial cells (Paridaen et al., 2013). Therefore, SBF-SEM may also be useful for studying the morphology of the primary cilium in the V-SVZderived neuroblasts migrating in the RMS.

In this study, we examined the morphology and dynamics of the primary cilium in V-SVZ-derived neuroblasts migrating in the postnatal RMS and OB. Using time-lapse imaging, TEM, and SBF-SEM, we found that the localization and orientation of the primary cilium in migrating neuroblasts are altered during proliferation and saltatory movement. Our results suggest that the spatiotemporal regulation of the primary cilium is related to the efficient chain migration of postnatal-born neuroblasts.

Materials and Methods

Animals. WT male and female C57BL/6J mice were purchased from Japan SLC (RRID:IMSR_JAX:000664). Male and female *GFP::Centrin2* (*GFP::Cent2*) mice (RRID:MGI:3793423) were described previously (Higginbotham et al., 2004), as were male *5HT3A-EGFP* mice (Inta et al., 2008). Male *R26-tdTomato* mice (stock #7914, The Jackson Laboratory; RRID:IMSR_JAX:007914) (Madisen et al., 2010) were provided by Dr. Masahiro Yamaguchi (Kochi Medical School). WT male and female AB (RRID:ZIRC_ZL1) zebrafish (*Danio rerio*) (3–8 months old) were obtained from the Zebrafish International Resource Center, and maintained with standard procedures as reported previously (Kishimoto et al., 2011, 2013; Ogino et al., 2016). The 1-year-old male rhesus monkey brain was obtained from EUPRIM-Net biobank (www.euprim-net.eu). All experiments involving live animals were performed in accordance with the guidelines and regulations of Nagoya City University and Heidelberg University.

Viral vectors and plasmids. To generate pENTR-GFP::Centrin2-IRES-Cre, a GFP::Centrin2 fragment was amplified from pEGFP-Cent2 (provided by Dr. Shaoyu Ge, Stony Brook University) by PCR and inserted into the SalI site of pENTR-IRES-Cre (Jinnou et al., 2018; Sawada et al., 2018). To generate pENTR-D-TOPO-dnKif3A, a dnKif3A fragment was amplified from pTetOn-dTomato-F2A-dnKif3A (Kumamoto et al., 2012) by PCR and inserted into the pENTR-D-TOPO vector. Gateway cloning technology (Invitrogen) was used to generate the following expression vectors: CSII-EF-GFP::Centrin2-IRES-Cre, CSII-EF-dnKif3A-IRES-Venus, and pCAGGS-dnKif3A. To generate CSII-CMV-Arl13b:: Venus, an Arl13b::Venus fragment was inserted into the Eco47III site of CSII-CMV-MCS-IRES2-Bsd. CSII lentiviral vectors were provided by Dr. Hiroyuki Miyoshi (RIKEN Tsukuba BioResource Center). To generate lentiviral particles, lentiviral vectors and packaging vectors (pCAG-HIVgp and pCMV-VSV-G-RSV-Rev) were transfected into HEK293T cells. Three days after transfection, the culture supernatants were concentrated by centrifuging at 8000 rpm at 4°C for 16 h using an MX-307 refrigerated microcentrifuge (Tomy).

To obtain pCDH-CMV-5HT6::GFP and pCDH-CMV-ACIII::GFP lentiviral vectors, 5HT6::GFP (Addgene, #35624) and ACIII::GFP (kindly provided by Dr. Joshua J. Breunig) (Guadiana et al., 2013) cassettes were subcloned into the lentiviral vector pCDH-CMV (the original pCDH plasmid was from System Bioscience). Recombinant viruses were produced and titrated as done previously (Khodosevich et al., 2012, 2013). Briefly, HEK293 cells were transfected with the viral backbone vector together with lentiviral helper plasmids (pMDLgpRRE, pRSV-Rev, and pMD2.G), and viral particles were collected by ultracentrifugation (~120,000 × g; Beckman) 48–60 h after transfection. The viral pellet was resuspended in D-PBS, aliquoted, and stored at -80° C until use. Viral titers were determined by infecting HEK293 cells in 96-well plates across 5 orders of dilutions. Before injection, viral titer was adjusted to 2 × 10⁷ to 2 × 10⁸ fluorescent plaque-forming units (fpfu)/ml.

For an IFT88-KD experiment using retrovirus, pU6-shRNA IFT88-EF1a-EGFP (provided by Dr. Shaoyu Ge, Stony Brook University) was used as reported previously (Kumamoto et al., 2012). To generate retroviral particles, pU6-shRNA IFT88-EF1a-EGFP and packaging vectors (pCAG-HIVgp and pCMV-VSV-G-RSV-Rev) were transfected into HEK293gp cells. Three days after transfection, the culture supernatants were concentrated by centrifugation as described above. To visualize 5HT3A localization in migrating neuroblasts, a retrovirus expressing 5HT3A::GFP (MMLV-RSV-EGFP-5HT3A) was used as reported previously (García-González et al., 2017). Recombinant retroviruses were produced by transfection of HEK293 cells with retroviral vectors and helper plasmids (pGP and pMD2.G), and viral particles were collected by ultracentrifugation as described above. The viral pellet was resuspended in D-PBS, and viral titer was adjusted to 2×10^7 to 2×10^8 fpfu/ml before injection.

To generate pEGFP-N1-SSTR3::GFP, an SSTR3::GFP fragment was inserted into the HindIII/KpnI sites of pEGFP-N1 (Clontech). To generate pdTomato-C1-Cent2 (dTomato::Cent2), EGFP cDNA in pEGFP-Cent2 (Kumamoto et al., 2012) was replaced with dTomato cDNA. pcDNA3.1-EB3-mEGFP (EB3::GFP) was provided by Dr. Kozo Kaibuchi (Nagoya University) (Watanabe et al., 2015).

In vivo viral infection and transplantation. Injection of lentiviral suspension into the V-SVZ of adult WT and R26-tdTomato mice was performed as described previously (Sawamoto et al., 2006) with modification. A 2 μ l volume of lentiviral suspension was stereotaxically injected into the anterior V-SVZ (1.0 mm anterior, 1.0, 1.1, 1.2 mm lateral to bregma and 1.8–2.3 mm deep) of male adult WT and R26-tdTomato mice. For injection of retrovirus encoding Ift88 shRNA, 2 μ l of retroviral suspension was stereotaxically injected into the anterior V-SVZ (1.0 mm anterior, 1.0, 1.1, and 1.2 mm lateral to bregma and 1.8–2.3 mm deep) of male adult WT mice. For injection of retrovirus expressing 5HT3A::GFP, 1 μ l of retroviral suspension was stereotaxically injected into the anterior V-SVZ (0.6 mm anterior, 1.2 mm lateral to bregma, and 1.7 mm deep) of P20 male WT mice, as reported previously (García-González et al., 2017).

Transplantation of neonatal V-SVZ cells into the V-SVZ of adult WT mice was performed as described previously (Sawada et al., 2018) with modification. The V-SVZ tissues from postnatal day 0–1 (P0-P1) WT mice were dissected and dissociated with trypsin-EDTA (Invitrogen). The cells were washed with L-15 medium (Invitrogen) containing 40 μ g/ml DNase I (Roche Diagnostics) and transfected with 2.0 μ g pEGFP-N1-SSTR3::GFP using the Amaxa Nucleofector II system (Lonza Walkersville). The transfected cells were resuspended in L-15 medium and stereotaxically injected into the anterior V-SVZ (1.0 mm anterior, 1.0, 1.1, 1.2 mm lateral to bregma, and 1.8–2.3 mm deep) of adult male WT mice.

In vitro time-lapse imaging of migrating neuroblasts. In vitro culture of V-SVZ-derived neuroblasts was performed as described previously (Ota et al., 2014; Sawada et al., 2018). The V-SVZ tissues were dissected from P0-P1 male and female WT mice and dissociated with trypsin-EDTA (Invitrogen). The cells were washed in L-15 medium (Invitrogen) containing 40 μ g/ml DNase I (Roche Diagnostics) and transfected with plasmids using the Amaxa Nucleofector II system (Lonza Walkersville). The transfected cells were recovered in RPMI-1640 medium (Thermo Fisher

Scientific), allowed to aggregate, and then embedded in 60% Matrigel (BD Biosciences) in L-15 medium. The cell aggregates were cultured in Neurobasal medium containing 2% NeuroBrew-21 (Invitrogen), 2 mM L-glutamine (Invitrogen), and 50 U/ml penicillin-streptomycin (Invitrogen) for 2 d.

Four-dimensional time-lapse images of the basal body and primary cilium in migrating neuroblasts were captured at 2 min and 0.6 μ m intervals using an LSM880 laser-scanning confocal microscope (Carl Zeiss) and a 63× oil-immersion objective lens. Time-lapse images of DsRed-expressing migrating neuroblasts in chains were captured at 1 min intervals using an inverted light microscope (Axio-Observer, Carl Zeiss) equipped with a Colibri light-emitting diode light system and a 40× objective lens, as reported previously (Sawada et al., 2018). Behaviors of migrating neuroblasts and GFP::Cent2 ⁺ basal body were tracked using the ImageJ manual tracking tool. The resting phase was defined as a migration speed <12 μ m/h (Jinnou et al., 2018). Three-dimensional reconstructions of migrating neuroblasts were performed using Imaris (Carl Zeiss) as reported previously (Hikita et al., 2014), based on the faint cytosolic GFP ⁺ (see Fig. 2B) or dTomato ⁺ (see Fig. 2G) signals.

For EB3::GFP imaging, time-lapse images of EB3::GFP⁺ dots in migrating neuroblasts were captured at 4 s intervals using an LSM880 laserscanning confocal microscope (Carl Zeiss) and a $63 \times$ oil-immersion objective lens, as reported previously (Sawada et al., 2018) with modification. All the EB3::GFP⁺ dots pursuable for 8 s (three sequential frames) were tracked using the ImageJ manual tracking tool, and their moving speed was measured using the ImageJ manual tracking tool.

Immunohistochemistry and image acquisition. Immunohistochemistry in mouse brain sections was performed as described previously (Ota et al., 2014; Sawada et al., 2018). Briefly, the adult brains were fixed by transcardiac perfusion with 4% PFA in 0.1 M PB, and postfixed overnight at 4°C in the same fixative. Fifty-micrometer-thick floating coronal and sagittal sections were made using a vibratome (VT-1200S, Leica Microsystems) and incubated for 30 min at room temperature (RT) in 10% normal donkey serum and 0.2% Triton X-100 in PBS (blocking solution). These sections were incubated with primary antibodies in blocking solution overnight at 4°C, and with AlexaFluor-488/-568/-647conjugated secondary antibodies (1:1000, Invitrogen) for 2 h at RT in blocking solution. The following primary antibodies were used: rabbit anti-adenylate cyclase III (ACIII) (1:200, sc-588, Santa Cruz Biotechnology; RRID:AB_630839); mouse IgG anti-Arl13b (1:1000, 75-287, NeuroMab; RRID:AB_2341543); rabbit anti-doublecortin (Dcx) (1:1000, 4604S, Cell Signaling Technology; RRID:AB_10693771); rabbit anti-DsRed (1:1000, 632496, Clontech; RRID:AB_10013483); rabbit anti-GFP antibody (1:1000, A6455, Invitrogen; RRID:AB_221570); rat anti-GFP (1:1000, 04404-84, Nacalai Tesque; RRID:AB_10013361); rabbit anti-calbindin (CB) (1:1000, CB38, Swant; RRID:AB_2721225); rabbit anti-calretinin (1:1000, 7697, Swant; RRID:AB_2619170); rabbit anti-Ki67 (1:1000, Leica Microsystems, NCL-Ki67p; RRID:AB_442102); rabbit anti-parvalbumin (PV) (1:1000, PV235, Swant; RRID:AB_10000343); mouse IgG anti-phospho-histone H3 (pHH3) (1:1000, 9706, Cell Signaling Technology; RRID:AB_331748); and sheep anti-tyrosine hydroxylase (TH) (1: 1000, AB1542, Millipore; RRID: AB_90755) antibodies. Nuclei were stained with Hoechst 33342 (1:5000, Invitrogen).

Immunohistochemistry in zebrafish brain sections was performed as described previously (Kishimoto et al., 2011; Ogino et al., 2016). The fish were anesthetized with tricaine. The brain was dissected and fixed with 4% PFA in 0.1 M PB overnight at 4°C. Fifty-micrometer-thick floating coronal sections were made using a vibratome (VT-1200S, Leica Microsystems) and incubated for 30 min at RT in 10% normal donkey serum and 0.2% Triton X-100 in PBS (blocking solution). These sections were incubated with primary antibodies in blocking solution overnight at 4°C, and with AlexaFluor-488/-568-conjugated secondary antibodies (1:500, Invitrogen) for 2 h at RT in blocking solution. The following primary antibodies were used: mouse IgM anti-polysialylated form of neural cell adhesion molecule (PSA-NCAM) (1:1000, a kind gift from Dr. Tatsunori Seki, Tokyo Medical University) (Seki and Arai, 1991) and mouse IgG anti-acetylated tubulin (1:11000, Sigma-Aldrich; RRID:AB_477585). Nuclei were stained with Hoechst 33342 (1:5000, Invitrogen).

Table 1. Numbers of neuroblasts examined in the SBF-SEM analyses^a

		Ciliary types					
	Data	Centrioles	Ciliary vesicle	Procilium	Nonextended primary cilium	Extended primary cilium	Total
Results	Neuroblast orientation	_	—	_	_	_	418 (88)
	Leading-process extension phase	22 (0)	23 (0)	10 (0)	82 (0)	6 (0)	143 (0)
	Swelling-formation phase	10 (0)	14 (0)	28 (0)	33 (0)	22 (0)	107 (0)
Fig. 5 <i>F—H</i> (Fig. 7 <i>B,M</i>)	Somal translocation phase	9 (0)	11 (0)	8 (0)	44 (0)	8 (0)	80 (0)
	Leading-process extension phase	—	—		—	—	15 (0)
	Swelling-formation phase	—	—	—	—	—	15 (0)
Fig. 5/	Somal translocation phase	—	—	_	—	—	15 (0)
Results	Mitochondrial length	—	—		—	—	18 (0)
Fig. 5 <i>Q</i>	Mitosis	—	—		—	—	1 (0)
	RMS	41 (0)	48 (0)	46 (0)	159 (0)	36 (0)	330 (0)
Fig. 7 <i>A</i>	OB	1 (1)	2 (2)	1 (1)	7 (7)	14 (14)	25 (25)
	RMS	—	—		159 (0)	36 (0)	195 (0)
Fig. 7C	OB	_	_		7 (7)	14 (14)	21 (21)
Fig. 7 <i>D,E</i>	Neuroblast	_	_		_	28 (0)	28 (0)
Fig. 7 <i>F,G</i>	Astrocyte	—	—		—	8 (0)	8 (0)
Fig. 7 <i>H,I</i>	Granule cell		_		—	3 (3)	3 (3)
Fig. 7 <i>J,K</i>	Presynaptic structure		—	—	—	11 (11)	11 (11)

"Values in parentheses indicate the numbers of neuroblasts with partial cell morphology (partial cells).

Immunohistochemistry in rhesus monkey brain sections was performed as described previously (García-González et al., 2017). Briefly, the brain was dissected, fixed with 4% PFA in 0.1 M PB 24 h at 4°C, and cut into 0.5 cm slabs. These slabs were postfixed with 4% PFA in 0.1 M PB, cryoprotected with 30% sucrose for 72 h. Fifty-micrometer-thick coronal sections were prepared by cryostat (Leica Microsystems). These sections were incubated for 30 min at RT with 5% BSA and 0.5% Triton X-100 in blocking solution, primary antibodies in blocking solution for 48 h at 4°C, and AlexaFluor-488/-568-conjugated secondary antibodies in blocking solution for 24 h at 4°C. The following primary antibodies were used: goat anti-Dcx antibody (1:250, sc-8066, Santa Cruz Biotechnology; RRID:AB_2088494) and mouse IgG anti-Arl13b (1:1000, 75-287, NeuroMab; RRID:AB_2341543). Nuclei were stained with DAPI (1:5000, Thermo Fisher Scientific; RRID:AB_2307445).

Images of stained RMS and OB sections were acquired by scanning at 0.4 μ m (see Figs. 1, 3*C*,*F*) or 2 μ m (see Fig. 3*A*,*B*,*D*,*E*,*G*–*I*) intervals using an LSM 700 confocal laser-scanning microscope (Carl Zeiss) with a 40× water-immersion objective lens. To investigate the positional relationship of the basal body and primary cilium in detail (see Fig. 1*B*), images of GFP::Cent2⁺ signals and Arl13b⁺ signals were acquired by scanning at 0.4 μ m intervals using an LSM 880 Airyscan confocal laser-scanning microscope with a 63× oil-immersion objective lens. For quantification of labeled cells in viral experiments (see Fig. 3), all the labeled cells observed in the RMS and OB in every section (dnKif3A) or every third section (IFT88-KD) were counted.

TEM. Sample preparation for TEM was performed as described previously (Kaneko et al., 2018). Adult male mouse brains were fixed by transcardiac perfusion with 2% PFA and 2.5% glutaraldehyde in 0.1 M PB, pH 7.4, at 4°C, and postfixed overnight at 4°C in the same fixative. These brains were cut into 200-µm-thick coronal sections using a vibratome (VT-1200S, Leica Microsystems) (anterior 2.0-2.3 mm from bregma). The sections were treated with 2% osmium tetroxide (OsO_4) Electron Microscopy Sciences) in 0.1 м PB, pH 7.4, at 4°C, dehydrated with a graded series of ethanol, placed in propyleneoxide (Nisshin EM), and embedded in Durcupan resin for 72 h at 60°C. Semithin sections (1.5- μ m-thick) were sequentially cut using an ultramicrotome (UC6, Leica Microsystems) with a diamond knife (histo, DiATOME), and stained with 1% toluidine blue, and sections of interest were chosen under a light microscope CX23 (Olympus). Ultrathin sections (60- to 70-nm-thick) were sequentially cut from the embedded semithin sections using an ultramicrotome (UC6, Leica Microsystems) with a diamond knife (SYM2045, SYNTEK), and stained with 2% uranyl acetate in distilled water for 15 min and with modified Sato's lead solution for 5 min. TEM images of the primary cilium in neuroblasts were acquired using a JEM-1400plus (JEOL) fitted with a digital camera. Thirty-six neuroblasts were examined in this study.

SBF-SEM. Sample preparation, observation with SBF-SEM, and analyses of acquired data were performed as described previously with slight modifications (Nguyen et al., 2016; Thai et al., 2016; Kaneko et al., 2018; Sawada et al., 2018). Adult WT brains were fixed by transcardiac perfusion with 2.5% glutaraldehyde and 2% PFA in 0.1 M PB, pH 7.4, at 4°C, and postfixed overnight at 4°C in the same fixative. Fixed RMS tissues (anterior 2.0-2.3 mm from bregma) were treated with 2% OsO4 and 1.5% potassium hexacyanoferrate in PBS for 1 h at 4°C, 1% thiocarbohydrazine for 20 min at RT, 2% aqueous OsO₄ for 30 min at RT, and lead aspartate solution for 30 min at 65°C. The RMS tissues were then dehydrated in a graded ethanol series, treated with dehydrated acetone, and embedded in Quetol 812 epoxy resin containing Ketjen black powder for 3 h at 95°C to ensure polymerization. SBF-SEM observation of the RMS and OB was performed using a Merlin and Sigma scanning electron microscope (Carl Zeiss) equipped with a 3View in-chamber ultramicrotome system (Gatan). Serial image sequences were $40.96 \times 40.96 \,\mu\text{m}$ wide (5.0 nm/pixel) and >80 μ m deep with 80 nm steps. Sequential images were processed using Fiji. In the image stacks, $\sim 2\%$ of the total images were replaced by neighboring images because some parts of the images were contaminated by debris or corrupted during image acquisition. Our SBF-SEM analyses contained neuroblasts with whole and partial morphology ("whole cells" and "partial cells," respectively). "Partial cells," containing part of the nucleus, Golgi apparatus, and centrosome or basal body with ciliary structures, and having a leading process >6µm, were used for analyses of neuroblast orientation in the RMS (described in Results), ciliary classification (see Fig. 7A), and ciliary length (see Fig. 7C). Numbers of analyzed whole and partial cells in the SBF-SEM analyses (see Figs. 5, 6, 7) are summarized in Table 1. Segmentation of the cell membrane, nuclei, centrioles, mitochondria, Golgi apparatus, and primary cilium were performed using Microscopy Image Browser (Belevich et al., 2016). Manual segmentation was done in every 2nd-10th and interpolation tool was used for the other slices. Three-dimensional reconstruction of migratory neuroblasts and their subcellular organs was performed using Amira software (Maxnet) as reported previously (Kaneko et al., 2018; Sawada et al., 2018). Three-dimensional object files (.obj) obtained from Amira software were further processed by Blender software (https://www.blender.org/) and visualized as interactive 3D models by Sketchfab (https://sketchfab.com).

Experimental design and statistical analysis. Statistical analyses were performed using EZR (Kanda, 2013), as reported previously (Fujikake et al., 2018). The experiments were not randomized. Sample sizes were not predetermined but were chosen based on previous studies. All the nu-

merical data, except for Figure 7M, were two-sided, and are shown as the mean \pm SEM. The normality of the data was analyzed by Kolmogorov– Smirnov test. For normally distributed data, equality of variances was examined by F test, and a comparison of means between two independent groups was performed by unpaired t test or Welch's t test. For comparisons of means among three or more independent groups, oneway ANOVA followed by Tukey-Kramer test was applied. To analyze the dataset from multiple observations of individual samples, two-way repeated-measures ANOVA followed by post hoc Tukey-Kramer test or unpaired t test between groups with Bonferroni correction was performed. For not normally distributed data, a comparison of medians between groups was performed by Mann–Whitney U test, Kruskal–Wallis test followed by Steel-Dwass test, or Friedman's test followed by Wilcoxon signed rank test with Bonferroni correction. In Figure 7M, all the data indicate the probability, and they were compared by Fisher's exact test. p < 0.05 was considered to be statistically significant.

Results

Expression of ciliary markers in migrating neuroblasts in the postnatal vertebrate RMS

To examine whether migrating neuroblasts in the postnatal RMS and OB present a primary cilium, we first stained brain sections for primary cilium marker Arl13b (Caspary et al., 2007) (Fig. 1A1-B'). To observe the basal body and a ciliary marker simultaneously, we studied Dcx⁺ neuroblasts in transgenic mice expressing GFP-tagged Centrin-2 protein (GFP::Cent2) (Fig. 1A1-A2') or V-SVZ-derived neuroblasts in WT mice infected with the GFP::Cent2-expressing lentivirus (Fig. 1B,B') (Higginbotham et al., 2004; Kumamoto et al., 2012). An Arl13b-positive (Arl13b⁺) structure was observed in close proximity to one of the centrioles in the Dcx⁺ neuroblasts in the RMS and OB, suggesting that it is associated with the basal body. Another marker for the primary cilium, ACIII (Bishop et al., 2007), is reported to be expressed in mature granule cells in the OB (Luo et al., 2015). ACIII was also detectable in migrating neuroblasts in the RMS (Fig. 1C) and mature interneurons in the external plexiform layer and glomerular layer of the OB (Fig. 1D-G). Moreover, fluorescent protein-tagged ciliary markers, such as Arl13b::Venus (Fig. 1H), ACIII::GFP (Fig. 1I), and 5HT6::GFP (Fig. 1J) (Brailov et al., 2000), accumulated in the perinuclear region of Dcx⁺ neuroblasts in the RMS. These results suggest that the migrating neuroblasts possess a primary cilium in the postnatal RMS and OB.

A functional primary cilium expresses signaling receptors on its membrane (Malicki and Johnson, 2017; Wheway et al., 2018). To investigate the subcellular localization of ciliary receptors in migrating neuroblasts in the RMS, we introduced GFP-tagged serotonin receptor 3A (5HT3A::GFP) or somatostatin receptor-3 (SSTR3::GFP) into migrating neuroblasts. These ciliary receptors colocalized with a ciliary marker (ACIII or Ar113b) (Fig. 1K-M'), suggesting that V-SVZ-derived migrating neuroblasts possess a functional primary cilium in the postnatal RMS.

Furthermore, we examined whether the expression of ciliary markers in migrating neuroblasts in the RMS is conserved in other vertebrates. We observed an acetylated tubulin-positive primary cilium in neuroblasts in the adult zebrafish RMS (Fig. 1N, N'), and an Arl13b-positive primary cilium in the 1-year-old rhesus monkey RMS and OB (Fig. 1O-P'). Collectively, these results suggest that the V-SVZ-derived migrating neuroblasts possess a primary cilium in the postnatal vertebrate brain.

Dynamic changes in primary cilium localization during neuronal saltatory migration

Previous EM studies have shown that the primary cilium is located at the cell surface at a distance from the nucleus, or in the perinuclear cytosol, in cultured MGE-derived interneurons (Baudoin et al., 2012). Furthermore, time-lapse imaging of Arl13b::Venus in cultured cerebellar granule neurons revealed that the primary cilium dynamically moves with the basal body (Trivedi et al., 2014). To study the subcellular localization of the primary cilium during saltatory movement in chain-forming neuroblasts, Arl13b::Venus was introduced into V-SVZ cells dissected from P0-P1 GFP::Cent2 mice (Fig. 2A). Saltatory movement of neuroblasts was divided into three phases: leadingprocess extension phase, swelling-formation phase, and somal translocation phase, as reported previously (Sawada et al., 2018). Three-dimensional confocal time-lapse imaging revealed that the primary cilium is submerged in the cytosol or extended from the cell surface (Fig. 2B-E'). The proportion of primary cilium extension from the cell surface was significantly higher in swellingformation phase than in other phases (Fig. 2C-F), suggesting that the extension of the primary cilium is promoted transiently before somal translocation in cultured chain-forming neuroblasts.

During neuronal deceleration in the postnatal OB, V-SVZderived neuroblasts transiently form a filopodium-like lateral protrusion (FLP) from the proximal leading process in the swelling-formation phase, which coincides with the pausing of somal translocation (Sawada et al., 2018). We found that both chain-forming and FLP-bearing neuroblasts extend a primary cilium from the cell surface during the swelling-formation phase (Fig. 2B-E', G-J'). The basal body with extended primary cilium was observed to be continuously migrating forward during the swelling-formation phase in chain-forming neuroblasts (Fig. 2*B*; 10-14 min), but not in FLP-bearing ones (Fig. 2*G*), suggesting that FLP formation is related to the migration termination of the primary cilium in the leading process.

Together, these results suggest that the primary cilium localization changes spatiotemporally during saltatory movement in cultured V-SVZ-derived migrating neuroblasts.

Inhibition of Kif3A and IFT88 impairs ciliogenesis and saltatory movement of V-SVZ-derived neuroblasts

The IFT molecules Kif3A and IFT88 control formation of the primary cilium and migration of pyramidal neurons (Chen et al., 2019) and interneurons (Baudoin et al., 2012) in the embryonic cerebral cortex. Loss of function of Kif3a or Ift88 causes impaired ciliogenesis, leading process morphology, and centrosomal movement in the MGE-derived cortical interneurons, leading to their abnormal directionality during migration and altered final distribution in the cerebral cortex (Baudoin et al., 2012). To examine the effects of Kif3A inhibition on V-SVZ-derived neuroblasts in the RMS, we injected a lentivirus expressing a dominant-negative form of Kif3A (dnKif3A) and Venus into the V-SVZ of adult WT mice. At 7 d postinjection (dpi), the proportion of neuroblasts with an Arl13b $^+$ primary cilium (Fig. 3A–C) and of those that had reached the OB (Fig. 3D) was decreased by dnKif3A expression, suggesting that ciliogenesis and migration toward the OB were disturbed. Unlike the deformation of leading process reported in Kif3A-deficient MGE-derived interneurons (Baudoin et al., 2012), the morphology of dnKif3A-expressing chain-forming neuroblasts was indistinguishable from the control (Fig. 3A, B). In the RMS, the proportion of neuroblasts pointing toward the OB was significantly decreased by dnKif3A expression (Fig. 3E). Similar defects in ciliogenesis and rostral migration in the V-SVZ-derived neuroblasts were also observed by IFT88-KD (Fig. 3F-H). At 14 dpi, a significantly larger proportion of V-SVZ-derived neuroblasts was observed to remain in the RMS in the dnKif3A-expressing group compared with the



Figure 1. Expression of ciliary markers in migrating neuroblasts in the RMS of the postnatal vertebrate brain. *A*–*A2'*, Representative images of RMS (*A1,A1'*) and OB (*A2,A2'*) sections of adult *GFP::Cent2* mice stained for GFP (green), Dcx (red), and Arl13b (white). *A1, A2*, Boxed areas are enlarged in *A1'* and *A2'*, respectively. Dcx ⁺ neuroblasts have Arl13b ⁺ structures (arrows) close to one of the pair of GFP::Cent2 ⁺ centrioles (arrowheads) in the RMS (*A1'*) and OB (*A2'*). *B, B'*, Representative image of tdTomato- and GFP::Cent2-labeled cells in the RMS. Lentivirus expressing GFP::Cent2 and Cre was injected into the V-SVZ of adult *R26-tdTomato* mice. A sagittal section of the RMS of *R26-tdTomato* mice at 6 dpi stained for GFP (green), DsRed (red), and Arl13b (white, arrows) is shown. *B*, Boxed areas are enlarged in *B'*. *B'*, An Arl13b ⁺ structure (arrows) is located close to one of the pair of GFP::Cent2 ⁺ centrioles (arrowheads). *C*, Representative image of an RMS section in *5HT3A-EGFP* mice (Inta et al., 2008) stained for GFP (green) and ACIII (red). Arrow indicates an ACIII ⁺ structure. *D*–*G*, Representative images of DB sections in WT mice stained for olfactory interneuron markers (green; *D*, PV; *E*, TH; *F*, calretinin [CR]; *G*, CB) and ACIII (red). Arrows indicate ACIII ⁺ structures in the olfactory interneurons. EPL, External plexiform layer; GL, glomerular layer. *H–J*, Localization of fluorescent protein-fused ciliary markers in migrating neuroblasts in the adult RMS. Ar113b::Venus (*H*), ACIII::GFP (*J*), or 5HT6::GFP (*J*) accumulated as spots (arrows) in the perinuclear region of Dcx ⁺ neuroblasts. Dashed lines indicate the boundaries of individual nuclei. Blue represents Hoechst 33342-stained nuclei. *K–M'*, Colocalization of GFP-fused 5HT3A and SSTR3 with ciliary markers in migrating neuroblasts in the adult RMS. SSTR3::GFP signal (*M*, *M'*, green) is localized in the Ar113b ⁺ structure (*M*, *M'*, red) in the leading process of migrating neuroblast



Figure 2. Primary cilium dynamics during saltatory movement of V-SVZ-derived cultured neuroblasts. *A*, Labeling of primary cilia and basal body in V-SVZ-derived cultured neuroblasts. Ar113b::Venus was introduced into V-SVZ cells dissected from P0-P1 *GFP::Cent2* mice. Two GFP::Cent2⁺ centrioles (green arrows) and an Ar113b::Venus⁺ primary cilium (yellow arrow) are visualized. *B*–*E*', Time-lapse imaging of a cultured migrating neuroblast expressing GFP::Cent2 and Ar113b::Venus. *B*, Extension or nonextension of an Ar113b::Venus⁺ primary cilium from the cell surface is indicated below by plus and minus signs, respectively. *B*, Boxed areas (10, 12, and 20 min) are shown orthogonally in *C*–*E*. *B*, Boxed area (0 min) is enlarged in *A*. *B*, Asterisks and dotted lines indicate leading process of a neighboring neuroblast and boundary between two neuroblasts, respectively. Three-dimensional reconstructions (*y*-*z* view) of the neuroblast in *B* (10, 12, and 20 min) are shown in *C'*–*E'*. *D*, *D'*, Yellow arrows indicate an extended primary cilium. *B*, Opened and filled arrowheads indicate spread and collapsed growth cones, respectively. *F*, Proportion of extension of a primary cilium from the cell surface during leading-process (LP) extension phase, swelling-formation phase, and somal translocation phase ($\chi^2_{(2)} = 30.679$, *p* = 0.00000022, Friedman's test; LP extension phase vs swelling-formation phase, *p* = 0.00011, LP extension phase vs somal translocation phase, *p* = 0.0017, Wilcoxon signed rank test, adjusted with Bonferroni correction; *n* = 20 cells from 11 independent cultures prepared on different days). *G*–*J'*, Time-lapse imaging of a cultured FLP-bearing neuroblast expressing dTomato::Cent2 and Ar113b::Venus⁺ primary cilium and dTomato::Cent2⁺ basal body. *G*, White arrows indicate an FLP. Boxed areas (10, 20, and 24 min) are shown orthogonally in *H*–*J*. Three-dimensional reconstructions (*y*-*z* view) of the neuroblast in *G* (10, 20, and 24 min) are shown. *H'*–*J'*, Yel

control (Fig. 31). Thus, loss of function of IFT molecules disrupted primary cilium formation and rostral migration in V-SVZ-derived neuroblasts in the RMS.

To study the effect of dnKif3A expression on neuronal saltatory movement in greater detail, we analyzed the migratory behavior of dnKif3A-expressing cultured neuroblasts dissected from the V-SVZ of P0-P1 *GFP::Cent2* mice (Fig. 3*J*–*V*; Movie 1). The dnKif3A-expressing neuroblasts spent longer in both the resting and migratory phases and showed a decreased migration

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speed (Fig. 3*J*–*N*). In the resting phase, while dnKif3A expression did not affect leading process extension (Fig. 3*O*,*P*), it slowed basal body speed (Fig. 3*Q*) without affecting its stride (Fig. 3*R*). Moreover, dnKif3A expression increased the frequency of backward basal body migration in the leading process (Fig. 3*K*; 0–24 min; Fig. 3*S*). In the migratory phase, dnKif3A expression decreased somal speed (Fig. 3*T*) but not stride (Fig. 3*U*). dnKif3A expression increased the frequency of reversal turn of neuroblasts (Fig. 3*K*, 24–54 min; Fig. 3*V*). Similar to MGE-derived interneurons (Baudoin et al., 2012), the phenotypes caused by the inhibition of IFT molecules suggest that the basal body and the primary cilium are involved in the saltatory migration of V-SVZ-derived neuroblasts.

Fine morphology of the primary cilium in migrating neuroblasts in the postnatal RMS

We investigated the ultrastructure of the primary cilium in migrating neuroblasts in the adult RMS using serial-section TEM (36 cells from 5 mice). Immature neuroblasts with migratory

⁽*Figure legend continued.*) PSA-NCAM (*N*,*N'*, red) and acetylated tubulin (*N'*, Ac-tubulin, green). *N*, Boxed area is enlarged in *N'*. Ac-tubulin $^+$ dots (*N'*, green, arrows) were observed in the PSA-NCAM $^+$ immature neuroblasts in the adult zebrafish RMS. V, Ventricle. *O–P''*, Representative images of coronal RMS (*O*, DAPI) and OB (*P*, DAPI) sections of an adult rhesus monkey stained for Dcx (*O'*, *P'*, *P''*, red) and Arl13b (*O'*, *P'*, *P''*, green). Arl13b $^+$ structures (*O'*, *P'*, *P''*, arrows) were observed in the Dcx $^+$ immature neuroblasts in the adult rhesus monkey RMS. Scale bars: *A1*, *A2*, *B*, *C*, *M*, *O'*, *P''*, *P''*, 5 μ m; *A1'*, *A2'*, *D–J*, *L*, *J*, *M'*, 2 μ m; *N*, 50 μ m; *P*, 500 μ m; *O*, 1 mm.



Figure 3. Defects in ciliogenesis and saltatory movement of V-SVZ-derived migrating neuroblasts caused by expressing a dominant-negative form of Kif3A or IFT88 shRNA. *A–I*, Defects in ciliogenesis, orientation, and final positioning of V-SVZ-derived migrating neuroblasts caused by expressing a dominant-negative form of Kif3A (dnKif3A) (*A–E,I*) or IFT88 shRNA. (*F–H*) in the adult RMS. Lentivirus encoding dnKif3A and Venus (*A–E,I*), or retrovirus encoding IFT88 shRNA and DsRed (*F–H*), was injected into the V-SVZ of WT mice. *A, B*, Representative images of control (*A*, green) and dnKif3A-expressing (*B*, green) cells in the adult RMS. An Arl13b ⁺ primary cilium (red, arrow in the enlarged box in *A*) was observed in the control but not the dnKif3A-expressing migrating neuroblasts. *C*, Proportion of Arl13b ⁺ primary cilium-bearing neuroblasts in the control and dnKif3A-expressing groups at 7 dpi ($t_{(4)} = 15.76$, $p = 9.5 \times 10^{-5}$, unpaired *t* test; n = 3 mice each). *D*, Distribution of labeled neuroblasts in the OB of control and dnKif3A-expressing groups at 7 dpi ($t_{(6)} = 3.62$, p = 0.011, unpaired *t* test; n = 4 mice each). *E*, Orientation of labeled neuroblasts in the control and dnKif3A-expressing groups at 7 dpi ($t_{(6)} = 3.62$, p = 0.011, unpaired *t* test; n = 4 mice each). *E*, Orientation of labeled neuroblasts in the control and dnKif3A-expressing groups at 7 dpi ($t_{(6)} = 3.62$, p = 0.011, unpaired *t* test; n = 4 mice each). *E*, Orientation of labeled neuroblasts in the control and dnKif3A-expressing groups at 7 dpi ($t_{(6)} = 3.62$, p = 0.011, unpaired *t* test; n = 4 mice each). *E*, Drientation of labeled neuroblasts in the OB of control and dnKif3A, $t_{(6)} = 5.0$, p = 0.099, *F*_{orientation(2,12)} = 79.8, *p*_{orientation} = 1.2 × 10⁻⁷, *F*_{group×orientation(2,12)} = 21.8, *p*_{group×orientation(2,12)} = 0.18, *p*_{group×orientation(2,12)} = 0.18, *p*_{group×orientation(2,12)} = 0.0034 in fGNS at 7 dpi ($t_{(4)} = 17.3$, $p = 6.6 \times 10^{-5}$, unp



Movie 1. Time-lapse imaging of control (red) and dnKif3A-expressing (red) neuroblasts (green) derived from the V-SVZ of *GFP*:: *Cent2* mice.



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(*Figure legend continued.*) migratory phases (*N*; p = 0.002, Welch's *t* test), leading process (LP) extension speed (*O*), maximum LP length (*P*), basal body (BB) speed (*Q*; W = 6669.5, $p = 6.82 \times 10^{-6}$, Mann–Whitney *U* test), BB stride (*R*), frequency of backward BB migration (*S*; W = 73, $p = 3.17 \times 10^{-5}$, Mann–Whitney *U* test), somal speed (*T*; p = 0.00012, Welch's *t* test), somal stride (*U*), and frequency of polarity reversal (*V*; W = 93, $p = 2.9 \times 10^{-5}$, Mann–Whitney *U* test) in control and dnKif3A-expressing cultured V-SVZ-derived neuroblasts (*L*-*V*; control, n = 21 cells; dnKif3A, n = 24 cells; five independent cultures prepared on different days). Scale bars: *A*, *B*, *J*, *K*, 5 μ m. *p < 0.05; ***p < 0.01; ***p < 0.005; ****p < 0.001; adjusted with Bonferroni correction in *E*, *H*, and *I*. Error bars indicate mean \pm SEM.

features in the RMS were identified based on their electron-dense nucleus with multiple nucleoli, dark cytoplasm with many free ribosomes, and the presence of extracellular spaces between neuroblasts, as described previously (Doetsch et al., 1997). Serial TEM images of migratory neuroblasts in the RMS showed either a pair of centrioles (Fig. 4A; 5 of 36 cells), or a basal body with an immature or mature primary cilium (Fig. 4*B*–*E*'; 31 of 36 cells). MTs were radially extended from the basal body of the primary cilium (Fig. 4*D*; *z* = 1 and 2), suggesting that the basal body acts as a MT-organizing center in migrating neuroblasts. The immature or mature primary cilium attached to the basal body (31 cells) was further classified into four types: ciliary vesicle, procilium, nonextended primary cilium, and extended primary cilium (Fig. 4*B*–*E*; see below).

In previous studies on primary cilium development in immature neurons, two types of immature forms of the primary cilium have been described: ciliary vesicles (Baudoin et al., 2012) and procilium (Arellano et al., 2012). Ciliary vesicles are attached to the basal body in the cytosol (Baudoin et al., 2012). The procilium is a membrane protrusion from the plasma membrane containing no axonemal structures (Arellano et al., 2012). We found immature primary cilia similar to ciliary vesicles (Fig. 4B; 6 of 31 cells) and to the procilium (Fig. 4C; 4 of 31 cells) in migratory neuroblasts in the RMS. In addition, elongated ciliary structures with a 9 + 0 ciliary axoneme and a ciliary pocket, a typical morphology of the mature primary cilium (Sorokin, 1962), were observed in migratory neuroblasts in the RMS (Fig. 4D-F; 21 of 31 cells). The tip of the primary cilium of these cells (21 cells) was observed to be either nonextended (Fig. 4D,D'; 18 cells) or extended (Fig. 4E, E'; 3 cells) from the cell surface, which we defined as a "nonextended primary cilium" and an "extended primary cilium," respectively. Interestingly, the tip of the nonextended primary cilium appeared to have access to the extracellular environment, even under its submerged state (Fig. 4D', yellow arrows). Collectively, these observations suggest that the primary cilium of immature migratory neuroblasts showed diverse morphologies in the postnatal RMS.

Three-dimensional ultrastructural features of migrating neuroblasts in the postnatal RMS analyzed by SBF-SEM

SBF-SEM is a powerful tool to study both whole cell morphology and its subcellular organs at an ultrastructural level (Briggman and Bock, 2012; Denk et al., 2012; Ohno et al., 2015). Since chain-forming neuroblasts align along the rostrocaudal axis in the anterior RMS, repeated coronal sectioning and image acquisition of RMS tissues by SBF-SEM efficiently include migrating neuroblasts. To establish criteria for celltype identification, ultrastructural features of adult mouse RMS cells observed under SBF-SEM (Figs. 5, 6; Movie 2) were compared with those observed under TEM, which have been reported previously (Doetsch et al., 1997).

Immature migratory neuroblasts could be identified in the SBF-SEM images by a smooth cell contour, dark cytoplasm with small Golgi apparatus, and a few short endoplasmic-reticulum fragments, and the presence of extracellular spaces, similar to the TEM images, as we reported previously (Kaneko et al., 2018; Sawada et al., 2018) (Fig. 5A). While the TEM images clearly showed adherens junction-like structures and an MT network in migratory neuroblasts (Doetsch et al., 1997; Fujikake et al., 2018; Sawada et al., 2018), our SBF-SEM images could not clearly visualize these structures. By performing 3D reconstruction of neuroblasts in the RMS, we examined 330 cells with whole morphology and 88 cells with partial morphology, which lack



Figure 4. *In vivo* identification of the primary cilium in migrating neuroblasts in the adult RMS. *A*–*E*['], TEM images of centrioles (*A*), ciliary vesicles (*B*, pink arrows), a procilium (*C*, magenta arrow), a nonextended primary cilium (*D*, magenta arrow), and an extended primary cilium (*E*, magenta arrow) in neuroblasts in the adult RMS. Green filled arrowheads indicate centrioles (*A*) or mother centrioles (*B*–*E*). *D*, Green open arrowheads indicate daughter centrioles. *D*, *E*, Boxed areas are enlarged in *D*['] and *E*['], respectively. *D*['], *E*['], Blue and white arrows indicate ciliary pockets and the ciliary axoneme, respectively. *D*['], *F*['], Blue and white arrows indicate ciliary pockets and the primary cilium in neuroblasts in the adult RMS. Coronal images of the basal body (*z* = 1), transition zone (*z* = 2–5), and doublet zone (*z* = 6–11) of the primary cilium are shown. The primary cilium shows 9 + 0 axoneme structure (*F*, *z* = 6; *F*['], red). Scale bars, 200 nm.

part of the leading process and/or soma (for a detailed definition of "partial cells," see Materials and Methods). The majority of identified neuroblasts were oriented rostrally (87.1 \pm 1.4%, 292 of 330 whole cells and 72 of 88 partial cells from 5 mice) and

tightly attached to their neighboring neuroblasts, forming chains in the RMS (Fig. 5*A*).

In addition to migratory neuroblasts, tunnel-forming astrocytes, oligodendrocytes, and microglia were observed in and



Figure 5. Ultrastructural features of V-SVZ-derived neuroblasts during proliferation and saltatory movement analyzed by SBF-SEM. *A*, Representative SBF-SEM images of migratory neuroblasts in the adult RMS. The migratory neuroblast consists of the growth cone, distal/proximal part of the leading process (LP), soma, and trailing process (blue). Migratory neuroblasts are tightly attached with their neighbors. Red represents all the neighbors. Between neuroblasts, extracellular spaces were also observed. *B–D'*, Representative SBF-SEM images of astrocytes (*B–B''*), oligodendrocytes (*C,C'*), and microglia (*D,D'*) in the adult RMS. Astrocytes showed long endoplasmic reticulum (*B'*, black arrows) and intermediate filaments (*B''*, white arrows), and were tightly attached to a blood vessel (*B,B'*). Oligodendrocytes showed a round cell contour with short and dilated endoplasmic reticulum (*C'*, back arrows), and direct contacts to myelinated and nonmyelinated axons (*C*). Microglia showed an irregular cell contour with long processes, dark cytosol with long endoplasmic reticulum (*D'*, black arrows) and a lysosome (*D'*, asterisk), and extracellular spaces (*D'*, white arrowheads). *E*, Time-lapse images for chain migration of DsRed-expressing cultured neuroblasts. Black and white arrowheads indicate spread and (*Figure legend continues*.)

around the RMS. Tunnel-forming astrocytes were identified by an irregular contour, light cytoplasm with long endoplasmic reticulum, and abundant intermediate filaments (Fig. 5B-B"), similar to the TEM images that have been reported previously (Doetsch et al., 1997). These astrocytes enwrapped the blood vessels and/or neuronal chains by their processes to form a glial tube in the RMS. Oligodendrocytes were located around the RMS and could be identified by a smooth and round contour, short and dilated endoplasmic reticulum, and direct contacts with myelinated and nonmyelinated axons (Fig. 5C,C') (Menn et al., 2006). Microglia were identified by an irregular contour with elongated processes, dark cytoplasm with long endoplasmic reticulum and lysosomes, and the presence of extracellular spaces (Fig. 5D,D') (Capilla-Gonzalez et al., 2014). Thus, SBF-SEM successfully classified all major cell types in the postnatal RMS based on their ultrastructural features.

(Figure legend continued.) collapsed growth cones, respectively. Collapsed growth cones were only observed during somal translocation. Neuroblasts in the LP extension phase showed a spread growth cone with no swelling in their leading process (0, 5 min). Neuroblasts in the swellingformation phase showed a spread growth cone with swelling in their leading process (7, 9 min). Neuroblasts in the somal translocation phase showed a collapsed growth cone with swelling in their leading process (12 min). Arrows indicate swelling. *F–H*, Representative 3D constructions of migratory neuroblasts (blue) under LP extension phase (F), swelling-formation phase (G), and somal translocation phase (H). Magenta and orange represent all the mitochondria and Golgi apparatus, respectively. Yellow represents nuclei. F-H, Interactive 3D models of neuroblasts in (without the Golgi apparatus) are shown at https://sketchfab.com/3d-models/399a425539e44087a9ba4a5c3dc5f361 (F), https://sketchfab.com/3d-models/fbaa32c3b1b34fdb9af1569cb3973475 (G), and https:// sketchfab.com/3d-models/ee40d9f934e840d18ba61a653fa2f7fc (H). I-K, Distribution and morphology of mitochondria in migrating neuroblasts. The number of mitochondria in LP extension phase, swelling-formation phase, and somal translocation phase (I; soma, $F_{(2,42)} = 6.44$, p =0.0036, one-way ANOVA; LP extension phase vs swelling-formation phase, p = 0.0032, LP extension phase vs somal translocation phase, p = 0.048, Tukey–Kramer test, proximal leading process, $F_{(2,42)} = 7.89$, p = 0.0012, one-way ANOVA; LP extension phase vs swellingformation phase, p = 0.0035, LP extension phase vs somal translocation phase, p = 0.004, Tukey–Kramer test, growth cone, $\chi^2_{(2)}=$ 19.2, p= 6.9 imes 10 $^{-5}$, Kruskal–Wallis test; LP extension phase vs somal translocation phase, $t(\infty) = 4.19$, $p = 8.3 \times 10^{-5}$, swellingformation phase vs somal translocation phase, $t(\infty) = 2.88$, $p = 1.1 \times 10^{-2}$; Steel–Dwass test; n = 3 mice). Proximal leading process contains swelling in swelling-formation phase and somal translocation phase (I). Most of the mitochondria in the proximal leading process are located in the swelling in swelling-formation phase (I, 94.3 \pm 1.6%) and somal translocation phase (I, 96.9 \pm 1.5%). J, K, Images are magnified views of boxed areas in G. L–O, Arl13b expression in proliferating and nonproliferating Dcx ⁺ cells in the adult RMS. Representative images of RMS sections of adult GFP::Cent2 mice stained for GFP (green, dot-like structures), pHH3 (L) or Ki67 (M,N) (green, nucleus), Arl13b (white), and Dcx (red). L–N, Arrows and arrowheads indicate the Arl13b⁺ primary cilium and centrioles, respectively. Proportions of Arl13b ⁺ primary cilium-bearing neuroblasts in proliferating and nonproliferating Dcx ⁺ neuroblasts are shown in **0** (left) ($F_{(3,8)} = 23.15 p = 0.00027$ one-way ANOVA; proliferating in M phase [pHH3⁺] vs proliferating in $G_1/S/G_2$ phases, p = 0.047, proliferating in M phase $[pHH3^+]$ vs nonproliferating, p = 0.00086, proliferating in M phase $[Ki67^+]$ vs proliferating in $G_1/S/G_2$ phases, p = 0.012, proliferating in M phase [Ki67⁺] vs nonproliferating, p = 0.00031, proliferating in $G_1/S/G_2$ phases vs nonproliferating, p = 0.047, Tukey–Kramer test; n = 3mice). Arl13b⁺ ciliary length in proliferating and nonproliferating neuroblasts is shown in **O** (right) (proliferating neuroblasts, 342.1 ± 25.8 nm, n = 42 cells from 3 mice; nonproliferating neuroblasts, 737.4 \pm 40.3 μ m, n = 90 cells from 3 mice; W = 3365.5, $p = 5.7 \times 10^{-13}$, Mann–Whitney U test). L–N, Bottom, Illustrations of the Arl13b⁺ primary cilium (pink, black arrows) in proliferating (green nuclei) and nonproliferating (white nucleus) Dcx ⁺ cells. Pairs of green dots represent pairs of centrioles. P, P', SBF-SEM images of dividing neuroblasts in the adult RMS. Black and green arrowheads indicate a disrupted nuclear membrane and a centriole without a primary cilium, respectively. P, Boxed area is enlarged in P'. Q, Three-dimensional construction of dividing neuroblasts in the adult RMS. Magenta, orange, and green represent mitochondria, Golgi apparatus, and centrioles, respectively. Yellow represents divided nuclei. An interactive 3D model of dividing neuroblasts (without the Golgi apparatus) is shown at https://sketchfab.com/3d-models/528050289d9143caab1c2b35c7fa1b3c. Raw SBF-SEM images in the adult RMS are shown in Movie 2. Scale bars: A–D, P, 1 μm; E, 5 μm; F–H, L–N, Q, 2 μ m. Error bars indicate mean \pm SEM. *p < 0.05, **p < 0.01, ***p < 0.005.

We next compared the SBF-SEM images of neuroblasts with the time-lapse images of cultured chain-forming neuroblasts during saltatory movement (Fig. 5E-H). The tip of the leading process showed a spread growth-cone-like structure during leading-process extension and swelling-formation phases (Fig. 5*E*, black arrowheads) that collapsed during somal translocation phase (Fig. 5E, white arrowhead). Swelling was observed after the extension of the leading process and before the ending of somal translocation, as reported previously (Ota et al., 2014; Sawada et al., 2018) (Fig. 5E, arrows). Consistent with these findings, 3D reconstruction of cell contour based on the SBF-SEM images visualized a long leading process, with or without swelling, which has a spread or collapsed growth cone at its tip, in addition to an elongated cell body and trailing process (Fig. 5F-H; Movies 3, 4, 5, and 6). Thus, by analyzing the morphology of the swelling and growth cone, we could identify the migration phase of each neuroblast in SBF-SEM images.

Three-dimensional reconstruction also revealed that mitochondria and the Golgi apparatus were mainly distributed in the proximal leading process or swelling, as reported previously (Bellion et al., 2005; Lin-Hendel et al., 2016) (Fig. 5F–H). While there was no significant difference in the numbers of mitochondria in neuroblasts that showed leading process extension, swelling formation, and somal translocation, their distribution was altered depending on saltatory movement (Lin-Hendel et al., 2016) (Fig. 5F-H, magenta, Fig. 5I). Mitochondria located in the proximal leading process were longer than those in the distal leading process (Fig. 5*J*,*K*; distal leading process, 0.58 \pm 0.02 μ m [n = 166from 18 cells]; proximal leading process, 1.04 \pm 0.03 μ m [n = 417 from 18 cells]; W = 19,594, $p = 2.6 \times 10^{-16}$, Mann–Whitney U test). Together, these findings suggest that mitochondria and the Golgi apparatus move in a saltatory movement-dependent manner in migratory neuroblasts in vivo.

In contrast to immature neurons migrating in the developing brain, V-SVZ-derived neuroblasts have remarkable mitotic potential during their migration in the RMS (Ikeda et al., 2010; Ponti et al., 2013). To examine the relationship between mitotic states and ciliogenesis in migrating neuroblasts in the RMS, we classified Dcx⁺ cells into proliferating or nonproliferating neuroblasts based on expression of the cell cycle marker Ki67. Proliferating neuroblasts were further divided into two subgroups: M phase (with a mitotic nucleus) or $G_1/S/G_2$ phases (Fig. 5L–N). In proliferating neuroblasts, an Arl13b⁺ primary cilium was observed less frequently, and its length was shorter, compared with these features in nonproliferating neuroblasts (Fig. 5O). Among these proliferating neuroblasts, the proportion of cilium-bearing neuroblasts was lower in M phase than in G₁/S/G₂ phases (Fig. 50), suggesting that ciliogenesis is transiently suppressed during mitosis. In the SBF-SEM analysis, dividing neuroblasts in the adult RMS possessed two pairs of centrioles, divided nuclei, no apparent Golgi apparatus (Danilov et al., 2009), and no primary cilium (Fig. 5P, Q; Movie 7). Together, these results suggest that mitotic states affect ciliogenesis in migrating neuroblasts in the postnatal RMS.

Changes in subcellular localization and orientation of the primary cilium during saltatory movement of migrating neuroblasts in the postnatal brain

To visualize the fine whole-cell morphology of migrating neuroblasts in the adult RMS and primary cilium *in vivo*, we reconstructed their images three-dimensionally (Figs. 6, 7; 330 whole cells from 5 mice). Consequently, a pair of centrioles (Fig. 6A-B'; Movie 8) or one of four types of primary cilium (Fig. 6C-J'; Movies 9, 10,



Figure 6. Three-dimensional ultrastructural analysis of the primary cilium in migrating neuroblasts in the adult RMS by SBF-SEM. *A–J'*, Sequential SBF-SEM images and 3D constructions of centrioles (*A–B'*), a ciliary vesicle (*C–D'*, pink arrows), a procilium (*E–F'*, magenta arrows), a nonextended primary cilium (*G–H'*, magenta arrows), and an extended primary cilium (*I–J'*, magenta arrows). Green filled arrowheads indicate centrioles (*A*) or mother centrioles (*C,E,G,I*). Green open arrowheads indicate daughter centrioles (*C,G,I*). *D'*, *F'*, *H'*, *J'*, Pink arrows indicate primary cilium orientation in the cells. *K*, *K'*, Three-dimensional construction of an extended primary cilium (*K'*) in an FLP-bearing neuroblast in the adult 0B. The arrow indicates an FLP. *B*, *D*, *F*, *H*, *J*, *K*, Interactive 3D models of neuroblasts are shown at https://sketchfab.com/3d-models/62e9a69dec414b31a64e8d0158bf7945 (*B*), https://sketchfab.com/3d-models/1c9eb432686748e8ac537afab59e335a (*D*), https://sketchfab.com/3d-models/5a5ba2836417475a9bb7513056824055 (*F*), https://sketchfab.com/3d-models/73abab8f28364a3c99277f7086c19c85 (*H*), https://sketchfab.com/3d-models/46cff34b6a9947bfa95fbd12b0696f4 (*J*), and https://sketchfab.com/3d-models/0e3963794ca74fc2a15d8cdde4e3c0ee (*K*). Scale bars: *A*, *C*, *E*, *G*, *I*, 200 nm; *B*, *D*, *F*, *H*, *J*, *K*, 5 µcm.

11, 12) was observed adjacent to the Golgi apparatus (100 \pm 0%, 330 whole cells from 5 mice) in migratory neuroblasts.

In the adult RMS, a mature primary cilium was observed in 59.8 \pm 3.3% of migratory neuroblasts, most of which (81.8 \pm 1.7%) were classified as the nonextended type (Fig. 7*A*). The remaining 40.2 \pm 3.3% had centrioles, ciliary vesicles, or a procilium (Fig. 7*A*). Consistent with the results of live imaging of cultured neuroblasts (Fig. 2*A*–*F*), the proportions of extended primary cilium and procilium were larger in the swelling-formation phase than in other phases of saltatory movement (Fig. 7*B*). The primary cilium length was not significantly different between nonextended and extended forms in the RMS (Fig. 7*C*). These results suggest that extension of the mature and immature

primary cilium is promoted transiently during the swellingformation phase of saltatory movement in chain-forming migrating neuroblasts *in vivo*.

While immature neuroblasts migrate in chains at a high speed in the RMS, these neuroblasts detach from chains to migrate individually, and transiently form FLPs during migration termination in the OB (Sawada et al., 2018). In contrast to the chainforming neuroblasts in the RMS, those in the granule cell layer of the OB, which are radially dispersed and have an elongated leading process, could not be included in our imaging volume ($40 \times$ $40 \times 80 \ \mu$ m). Therefore, we analyzed 25 partial OB cells from 3 mice. We investigated the primary cilium morphology in immature neuroblasts in the OB and found a mature primary cilium in



Movie 2. Sequential SBF-SEM images in the adult RMS. Representative sequential SBF-SEM images (1156 sections) in the adult RMS are shown. Neuroblasts shown in Figure *5F*, *G*, *H* appear in yellow, red, and blue, respectively. Dividing neuroblasts shown in Figure *5Q* appear in green.



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Movie 4. Three-dimensional reconstruction image of a migratory neuroblast during leading process extension. Green, yellow, orange, and magenta represent centrioles, nucleus, Golgi apparatus, and mito-chondria, respectively.





Movie 5. Three-dimensional reconstruction image of a migratory neuroblast in the swelling-formation phase. Green, yellow, orange, and magenta represent centrioles, nucleus, Golgi apparatus, and mitochondria, respectively.





Movie 6. Three-dimensional reconstruction image of a migratory neuroblast during somal translocation. Green, yellow, orange, and magenta represent centrioles, nucleus, Golgi apparatus, and mitochondria, respectively.





Movie 3. Three-dimensional reconstruction of a migratory neuroblast in the adult RMS. Blue represents the cell contour of the migratory neuroblast. Green, yellow, orange, and magenta represent centrioles, nucleus, Golgi apparatus, and mitochondria, respectively.



84.9 ± 11.4% of these neuroblasts (Fig. 7*A*). The proportion of extended primary cilium was significantly larger in FLP-bearing neuroblasts than in neuroblasts without FLP (Fig. 6*K*,*K*'; FLP-bearing neuroblasts, 95.2 ± 4.8%; neuroblasts without FLP, 22.2 ± 14.7%; $t_{(4)} = -4.73$, p = 0.0091, paired *t* test; n = 3 mice). Furthermore, the extended primary cilium was longer than the



Movie 7. Three-dimensional reconstruction image of dividing neuroblasts. Blue and yellow represent the cell contour and divided nuclei, respectively. Green, orange, and magenta represent centrioles, Golgi apparatus, and mitochondria, respectively.



nonextended primary cilium in the OB, and both were longer than their counterparts in the RMS (Fig. 7C). Together, these results suggest that the elongation and extension of primary cilium are promoted in individual decelerating neuroblasts in the OB.

To understand the extracellular microenvironment of the extended primary cilium in migrating neuroblasts in the RMS and OB, we identified neural structures contacting the extended primary cilium in the RMS and OB. In the RMS, each extended primary cilium made contact with a neighboring neuroblast (28



Figure 7. Changes in subcellular localization, length, and orientation of the primary cilium during saltatory movement of migrating neuroblasts in the adult RMS. *A*, The proportions of immature migratory neuroblasts that have centrioles, a ciliary vesicle, a procilium, a nonextended primary cilium, and an extended primary cilium in the adult RMS ($F_{(4,20)} = 56.59$, $p = 1.3 \times 10^{-10}$, one-way ANOVA; centrioles vs nonextended primary cilium, p < 0.000001, ciliary vesicle vs nonextended primary cilium, p < 0.000001, procilium vs nonextended primary cilium, p < 0.000001, ciliary vesicle vs nonextended primary cilium, p = 0.00034, rone-way ANOVA; centrioles vs extended primary cilium, p = 0.00034, ciliary vesicle vs extended primary cilium, p = 0.00034, ciliary vesicle vs extended primary cilium, p = 0.00034, ciliary vesicle vs extended primary cilium, p = 0.0005, procilium vs extended primary cilium, p = 0.00034, nonextended primary cilium, sextended primary cilium, p = 0.0005, procilium vs extended primary cilium, p = 0.00034, nonextended primary cilium, and an extended primary cilium, p = 0.0005, procilium vs extended primary cilium, p = 0.00034, nonextended primary cilium, and an extended primary cilium, p = 0.0005, procilium vs extended primary cilium, p = 0.00034, nonextended primary cilium, and an extended primary cilium during leading-process (LP) extension phase, the swelling-formation phase, and somal translocation phase in the adult RMS ($F_{phase(2,12)} = -0.004$, $p_{phase} = 1$, $F_{class}(4,48) = 75.52$, $p_{class} < 2.2 \times 10^{-16}$, $F_{phase \times class}(8,48) = 9.69$, $p_{phase \times class} = 7.1 \times 10^{-8}$, two-way repeated-measures ANOVA; $F_{(2,12)} = 20.31$, p = 0.00014, one-way ANOVA; LP extension phase vs swelling-formation phase, p = 0.00045, in procilium; $F_{(2,12)} = 13.02$, p = 0.00099, one-way ANOVA; LP extension (*Figure legend continues*.)



Movie 8. Three-dimensional reconstruction image of a neuroblast having centrioles. Blue and green represent the cell contour and centrioles, respectively.



of 36 cells) or astrocyte (8 of 36 cells) (Fig. 7*D*–*G*). In the granule cell layer of the OB, the extended primary cilium of neuroblasts directly contacted neurites of mature neurons (14 of 14 cells). Close observation shows that each primary cilium of these neu-

(Figure legend continued.) phase vs swelling-formation phase, p = 0.0071, somal translocation phase vs swelling-formation phase, p = 0.02, in nonextended primary cilium; $F_{(2,12)} =$ 18.55, p = 0.00021, one-way ANOVA; LP extension phase vs swelling-formation phase, p =0.00088, somal translocation phase vs swelling-formation phase, p = 0.023, in extended primary cilium; Tukey–Kramer test; *n* = 5 mice). **p* < 0.05; ***p* < 0.01; ****p* < 0.005; adjusted with Bonferroni correction. C, Length of the nonextended and extended primary cilium in immature neuroblasts in the adult RMS and OB (RMS vs OB, W = 869.5 p = 0.012, in nonextended primary cilium; Mann–Whitney *U* test; $t_{(48)} = 6.68$, $p = 2.3 \times 10^{-8}$, in extended primary cilium; nonextended primary cilium vs extended primary cilium, $t_{(19)} = -2.7$, p = 0.014, in OB; unpaired t test; n = 3 mice each). **D–K**, Sequential SBF-SEM images and 3D constructions of extended primary cilium contacting a neighboring neuroblast (**D**,**E**, red), astrocyte (**F**,**G**, green), granule cell (H,I, orange), and presynaptic structure (J,K, yellow). J', White arrows and arrowheads indicate postsynaptic density and synaptic vesicles, respectively. L, EB3::GFP speed in the growth cone, distal/proximal leading process, and soma during saltatory movement of cultured chain-forming migrating neuroblasts ($\chi^2_{(3)} =$ 14.67, p = 0.0021, Kruskal–Wallis test; distal leading process vs soma, $t(\infty) = 3.66$, p = 0.0014, in LP extension phase; $\chi^2_{(3)} = 17.77$, p =0.00049, Kruskal–Wallis test; distal leading process vs soma, $t(\infty) = 3.13$, p = 0.0094, growth cone vs proximal leading process, $t(\infty) = 2.84$, p = 0.023, proximal leading process vs soma, $t(\infty) = 3.12, p = 0.0097$, in swelling-formation phase; $\chi^2_{(3)} = 53.72, p = 1.3 \times 10^{-11}$ Kruskal–Wallis test; distal leading process vs soma, $t(\infty) = 6.87$, $p = 4.0 \times 10^{-11}$, growth cone vs soma, $t(\infty) = 4.19$, $p = 1.6 \times 10^{-4}$, proximal leading process vs soma, $t(\infty) = 3.45$ $p = 3.1 \times 10^{-3}$, in somal translocation phase; $\chi^2_{(2)} = 8.61$, p = 0.0014, Kruskal–Wallis test; LP extension phase vs somal translocation phase, $t(\infty) = 2.71$, p = 0.0018, swelling-formation phase vs somal translocation phase, $t(\infty) = 2.4$, p = 0.043, in growth cone; $\chi^2_{(2)} = 7.21$, $p < 10^{-10}$ 2.2×10^{-16} , Kruskal–Wallis test; LP extension phase vs somal translocation phase, $t(\infty) =$ 7.46, $p = 2.7 \times 10^{-13}$, swelling-formation phase vs somal translocation phase, $t(\infty) = 6.55$, p= 1.8 imes 10 $^{-10}$, in distal leading process; $\chi^2_{(2)}=$ 8.8, p= 0.0012, Kruskal–Wallis test; LP extension phase vs swelling-formation phase, $t(\infty) = 2.43$, p = 0.04, swelling-formation phase vs somal translocation phase, $t(\infty) = 2.69$, p = 0.019, in proximal leading process; $\chi^2_{(3)}$ = 22.18, $p = 1.5 \times 10^{-5}$, Kruskal–Wallis test; LP extension phase vs somal translocation phase, $t(\infty) = 4.1$, p = 0.00012, swelling-formation phase vs somal translocation phase, $t(\infty) = 4.11, p = 0.00012$, in soma; Steel–Dwass test). *p < 0.05; **p < 0.01; ***p < 0.005; $p^{*}p < 0.05$; versus LP extension phase and swelling-formation phase in growth cone. $p^{*}p < 0.05$; versus LP extension phase and swelling-formation phase in growth cone. 0.005, versus LP extension phase and swelling-formation phase in distal leading process. $p \leq 1$ 0.05, versus LP extension phase and somal translocation phase in proximal leading process. $\pm p < 0.005$, versus LP extension phase and swelling-formation phase in soma. Values in parentheses indicate the number of events from eight cells in three independent cultures. *M*, The proportions of immature and mature primary cilium that orient posteriorly under LP extension phase, the swelling-formation phase, and somal translocation phase in the adult RMS (p = 0.00015, Fisher's exact test, somal translocation phase vs swelling-formation phase, p =0.0064, LP extension phase vs somal translocation phase, p = 0.00098, in ciliary vesicle; p =0.00015, Fisher's exact test; somal translocation phase vs swelling-formation phase, p = 0.0078, LP extension phase vs somal translocation phase, p = 0.00075, in procilium; p = 6.2×10^{-8} , Fisher's exact test, somal translocation phase vs swelling-formation phase, p =0.00012, LP extension phase vs somal translocation phase, $p = 9.4 \times 10^{-7}$, in nonextended primary cilium; $p = 9.5 \times 10^{-5}$, Fisher's exact test, somal translocation phase vs swellingformation phase, p = 0.00039, LP extension phase vs somal translocation phase, p = 0.014, in extended primary cilium; n = 5 mice). *p < 0.05; ****p < 0.005; versus LP extension phase. p < 0.05; p < 0.005; versus swelling-formation phase. Error bars indicate mean \pm SEM. Scale bars, 1 μ m.



Movie 9. Three-dimensional reconstruction image of a neuroblast having a ciliary vesicle. Blue, green, and magenta represent the cell contour, centrioles, and ciliary vesicle, respectively.





Movie 10. Three-dimensional reconstruction image of a neuroblast having a procilium. Blue, green, and magenta represent the cell contour, centrioles, and ciliary membrane, respectively.





Movie 11. Three-dimensional reconstruction image of a neuroblast having a nonextended primary cilium. Blue, green, and magenta represent the cell contour, centrioles, and ciliary membrane, respectively.





Movie 12. Three-dimensional reconstruction image of a neuroblast having an extended primary cilium. Blue, green, and magenta represent cell contour, centrioles, and ciliary membrane, respectively.



roblasts was making contact with the soma of granule cells (3 of 14 cells) or with presynaptic structures containing synaptic vesicles that had formed on the neurites (11 of 14 cells) (Fig. 7H–K). These results suggest that the extended primary cilium in a neuroblast interacts with its surrounding cells during migration in the RMS and OB.

Finally, we analyzed the dynamics of MT polymerization in each cellular domain during saltatory movement of cultured chain-forming neuroblasts, and compared the data with the orientation of the primary cilium analyzed using SBF-SEM. The highest polymerization levels of MTs labeled with EB3::GFP (Sawada et al., 2018) were observed in the distal leading process during leading-process extension phase, in the proximal leading process during the swelling-formation phase, and in the soma during somal translocation phase (Fig. 7L; Movie 13). On the other hand, quantification of ciliary orientation based on the SBF-SEM images (Fig. 6D', F', H', J') revealed that the majority



Movie 13. Time-lapse imaging of EB3::GFP and dTomato::Cent2 during saltatory movement of a cultured V-SVZ-derived migrating neuroblast. The migratory behavior of an EB3::GFP- (white) and dTomato:: Cent2 (magenta)-expressing V-SVZ-derived neuroblast was recorded at 4 s intervals by superresolution microscopy. Leading-process extension phase, 0 - 440 s; swelling-formation phase, 444 - 1280 s; somal translocation phase, 1284 - 1920 s. Scale bar, 2μ m.



of migrating neuroblasts (80%–100%) of the leading-process extension and swelling-formation phases had a posteriorly oriented primary cilium regardless of the ciliary type, whereas migrating neuroblasts of the somal translocation phase had an anteriorly oriented primary cilium (Fig. 7M). The orientation of the primary cilium in the direction opposite to the cellular domain with the highest levels of MT polymerization suggests that ciliary orientation is related to the difference in the local MT polymerization levels in each migration phase rather than to ciliary maturation states.

Discussion

SBF-SEM analyses of migrating neuroblasts and the primary cilium *in vivo*

Time-lapse imaging of V-SVZ-derived migrating neuroblasts reveals each step of their saltatory movement (Schaar and McConnell, 2005; Shinohara et al., 2012; Ota et al., 2014; Sawada et al., 2018). Here, by analyzing the presence/absence of swelling and growth cone morphology, we could identify the migration phases of immature neuroblasts both in time-lapse imaging and in SBF-SEM analyses. Moreover, 3D reconstruction of mitochondria showed changes in their distribution in migration phases, which is consistent with a previous study in MGE-derived cortical interneurons (Lin-Hendel et al., 2016). Since dynamic changes in cellular morphology in migrating neuroblasts are supported by mitochondrial ATP production (Lin-Hendel et al., 2016), it is possible that mitochondria accumulate in subcellular regions with high-energy demands, such as the swelling.

SBF-SEM or another type of 3D-SEM, focused ion beam-SEM, has previously been used to analyze the fine morphology of the primary cilium (Paridaen et al., 2013; Insinna et al., 2019) and the molecular mechanisms of ciliogenesis (David et al., 2014; Buskin et al., 2018). Although our SBF-SEM analyses could not clearly visualize MT structures, we successfully identified primary cilium structures by detecting their basal bodies and associated membrane structures, whose electron density was strengthened by the reduced osmium-thiocarbohydrazideosmium method (Seligman et al., 1966; Willingham and Rutherford, 1984). Our ultrastructural SBF-SEM images for different stages of ciliogenesis should provide a basis for analysis of the primary cilium in other cells in the brain or in other tissues under physiological and pathological conditions.

Unique morphology and dynamics of the primary cilium in V-SVZ-derived migrating neuroblasts

In contrast to other immature neurons migrating in the developing brain, the V-SVZ-derived neuroblasts are unique in that they are still mitotic and migrating in a chain in the RMS. Our observations suggest that ciliogenesis in migrating neuroblasts is transiently suppressed during mitosis (Fig. 8A), as reported in other mitotic cells (Malicki and Johnson, 2017). In addition, a small portion of proliferating Dcx⁺ cells has an Arl13b⁺ primary cilium, even in the M phase. It is possible that the status of the primary cilium in migrating neuroblasts is influenced by dynamic changes in their cellular circumstances, including those in the cell cycle and saltatory movement.

Our live imaging and EM analyses revealed that migrating neuroblasts in the postnatal RMS possess a primary cilium in the cytosol, which is transiently extended from the cell surface before somal translocation (Fig. 8*B*). Similar subcellular translocation of the primary cilium during migration has been reported in MGEderived interneurons (Baudoin et al., 2012), suggesting that ciliogenesis in interneurons is regulated by common mechanisms in the embryonic and postnatal brain. Interestingly, the "nonextended" primary cilium in immature neuroblasts appeared to have access to the extracellular environment, which has been



Figure 8. Spatiotemporal regulation of the primary cilium in V-SVZ-derived migrating neuroblasts in the postnatal brain. *A*, The primary cilium in proliferating neuroblasts in the postnatal RMS. Ciliogenesis is suppressed during mitosis (green nuclei) of V-SVZ-derived migrating neuroblasts in the RMS. *B*, The primary cilium in migration of chain-forming neuroblasts in the RMS. The primary cilium (pink) is submerged in the cytosol and orients posteriorly during leading process extension, when a spread growth cone is observed. During the swelling-formation phase, the primary cilium is extended from the cell surface and moves forward within the swelling. The primary cilium is not extended when neuroblasts translocate their soma. The orientation of the primary cilium is related to active MT polymerization (bold blue arrows) in migration phases. *C*, The primary cilium in migration of FLP-bearing neuroblasts in the postnatal 08. The FLP is transiently formed from the proximal part of the leading process before somal translocation (Sawada et al., 2018). FLP formation coincides with active and local MT reorganization in the proximal leading process (bold blue arrows), which maintains migration termination in neuroblasts. The elongation and extension of the primary cilium are promoted in individual decelerating neuroblasts in the 08.

reported in other types of mature cells (Sorokin, 1962; Dahl, 1967; Smith et al., 1969; Mazo et al., 2016).

In MGE-derived migrating interneurons, extracellular extension of the primary cilium could be regulated by fusion and fission of Golgi-derived ciliary vesicles (Baudoin et al., 2012). A recent study suggests that the submerged state of the primary cilium in the cytosol is maintained by association of daughter centriole with the Golgi apparatus in retinal pigmental epithelial cells (Mazo et al., 2016). It remains unknown whether these mechanisms are involved in maintenance of the nonextended state of the primary cilium in postnatal-born migrating neuroblasts.

The FLP-bearing individual neuroblasts decelerating in the OB restrict forward movement of the primary cilium in the leading process and promote cilium extension from the cell surface (Fig. 8*C*), which is distinct from chain-forming neuroblasts in the RMS. These changes in primary cilium dynamics may be related to termination of migration and initiation of differentiation in the OB, which are suppressed by homophilic chain formation and/or efficient migration in the RMS.

Possible functions of the primary cilium in V-SVZ-derived migrating neuroblasts

Our SBF-SEM results showed that 88% of immature neuroblasts in the adult RMS possess ciliary structures on their mother centrioles. Moreover, MTs were observed to be extended radially from the basal body of the primary cilium in migrating neuroblasts. Similar relationships between MTs and the basal body of the primary cilium have also been reported in fibroblasts (Albrecht-Buehler and Bushnell, 1980; Schrøder et al., 2011) and enteric neurons (Luesma et al., 2013). These findings suggest that the basal body of the primary cilium acts as an MT-organizing center in the V-SVZ-derived migrating neuroblasts, which is consistent with previous studies (Tsai et al., 2007; Baudoin et al., 2012).

We found that the primary cilium orients posteriorly when neuroblasts actively polymerize MTs in the leading process, whereas it orients anteriorly when neuroblasts actively form a perinuclear MT cage (Fig. 8*B*). Thus, it is possible that active MT polymerization generates traction force that pulls the basal body, which would affect the ciliary orientation during saltatory movement. During neuroblast deceleration, the FLP induces local MT reorganization in the proximal leading process, which can decrease the MT polymerization-based traction force and sustain extension of the primary cilium in FLP-bearing neuroblasts. Together, these results suggest that the dynamics of the basal body and primary cilium are regulated by MT dynamics in saltatory movement of migrating neuroblasts.

Kif3A promotes ciliogenesis by regulating axoneme elongation and subdistal appendage formation in the basal body (Kodani et al., 2013). However, Kif3A also has nonciliary functions, such as transport of N-cadherin (Teng et al., 2005; Ichinose et al., 2015) and modulation of Wnt signaling (Corbit et al., 2008). Therefore, defective neuroblast migration caused by dnKif3A expression does not directly indicate the function of the primary cilium in neuroblast migration in the RMS. The KD of IFT88, which contributes to ciliogenesis independently of Kif3A (Pazour et al., 2000), also disrupted migration of V-SVZ-derived neuroblasts, similar to MGE-derived cortical interneurons (Baudoin et al., 2012). However, since the V-SVZ-derived migrating neuroblasts are mitotic, nonciliary functions of IFT88 during mitosis, including the regulation of spindle orientation (Robert et al., 2007; Delaval et al., 2011; Borovina and Ciruna, 2013), may affect neuronal distribution. Thus, our results suggest that disrupted neuroblast migration resulting from inhibition of Kif3A and IFT88 is, at least in part, attributable to the defects of primary cilium function in the V-SVZ-derived migrating neuroblasts.

The primary cilium contains signaling receptors, such as Smoothened and 5HT6, on its membrane, which act as chemical sensors to regulate migration and maturation of cortical interneurons (Riccio et al., 2009; Baudoin et al., 2012; Higginbotham et al., 2012; Guo et al., 2017). Shh signaling promotes neuroblast migration in the V-SVZ (Angot et al., 2008). Moreover, a recent study suggests that serotonin controls speed and directionality of the V-SVZ-derived migrating neuroblasts (García-González et al., 2017). Whereas 5HT6::GFP localized only in the primary cilium, 5HT3A::GFP was detectable not only in the cilium but also in the cell body (Fig. 1*J*,*L*). Furthermore, 5HT6 and 5HT3A are a GPCR and a ligand-gated ion channel, respectively (Derkach et al., 1989; Monsma et al., 1993; Ruat et al., 1993). Therefore, it is possible that these two types of serotonin receptors coordinately regulate neuroblast migration in the adult RMS and OB via distinct signaling pathways. Serotonin fibers extending along the RMS-OB pathway (García-González et al., 2017) and the presence of a primary cilium in migrating neuroblasts are widely conserved among vertebrates. Since the postnatal-born neuroblasts must migrate over a long distance within the mature neuronal circuitry, which is distinct from embryonic neurons, they may need to continuously sense extracellular cues through the primary cilium for their efficient migration.

Migration and maturation of neuroblasts in the OB are affected by both olfactory (Petreanu and Alvarez-Buylla, 2002; Saghatelyan et al., 2004; Yamaguchi and Mori, 2005) and centrifugal (Kaneko et al., 2006; García-González et al., 2017) inputs. Interestingly, the extended primary cilium in individually migrating neuroblasts made direct contact with presynaptic structures in the OB (Fig. 7J,K). If the extended primary cilium has the potential to transduce electrical activity, as in the renal collecting duct epithelium (Liu et al., 2018), it may contribute to activity-dependent neuroblast migration and maturation in the OB.

Although the primary cilium has also been proposed to act as a mechanosensor (Malicki and Johnson, 2017), it remains unknown whether the primary cilium in migrating neuroblasts mediates mechanosensing. In retinal pigment epithelial cells, whereas the submerged primary cilium was not affected by fluid flow, the surfaced primary cilium, induced by *C-Nap1/CEP128* deficiencies, bent in response to fluid flow (Mazo et al., 2016). If the primary cilium in migrating neuroblasts is able to transduce mechanostimulation, its activity could be altered between nonextended and extended states, and affected by cell–cell contact that changes dynamically during chain migration.

In conclusion, this study suggests that primary cilium dynamics is related to the proliferation and saltatory movement of V-SVZ-derived migrating neuroblasts. Our detailed description of the ultrastructural features and dynamics of primary cilium in migrating neuroblasts will be useful for future studies, which will clarify the functional significance of the primary cilium and novel regulatory mechanisms for neuroblast migration.

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