Nwd1 Regulates Neuronal Differentiation and Migration through Purinosome Formation in the Developing Cerebral Cortex

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HIGHLIGHTS

STAND protein Nwd1 interacts with Paics to regulate the purinosome formation

Dysregulated expression of Nwd1 induced the premature differentiation of NSPCs

Nwd1 KD repressed the neuronal migration, causing the periventricular heterotopia

Tightly regulated assembly of purinosome components is crucial for corticogenesis
Nwd1 Regulates Neuronal Differentiation and Migration through Purinosome Formation in the Developing Cerebral Cortex

Seiya Yamada,1 Ayaka Sato,1 and Shin-ichi Sakakibara1,2,*

SUMMARY
Engagement of neural stem/progenitor cells (NSPCs) into proper neuronal differentiation requires the spatiotemporally regulated generation of metabolites. Purines are essential building blocks for many signaling molecules. Enzymes that catalyze de novo purine synthesis are assembled as a huge multienzyme complex called “purinosome.” However, there is no evidence of the formation or physiological function of the purinosome in the brain. Here, we showed that a signal transduction ATPases with numerous domains (STAND) protein, NACHT and WD repeat domain-containing 1 (Nwd1), interacted with Paics, a purine-synthesizing enzyme, to regulate purinosome assembly in NSPCs. Altered Nwd1 expression affected purinosome formation and induced the mitotic exit and premature differentiation of NSPCs, repressing neuronal migration and periventricular heterotopia. Overexpression/knockdown of Paics or Fgams, other purinosome enzymes, in the developing brain resulted in a phenocopy of Nwd1 defects. These findings indicate that strict regulation of purinosome assembly/disassembly is crucial for maintaining NSPCs and corticogenesis.

INTRODUCTION
The spatiotemporal differentiation of neural stem/progenitor cells (NSPCs) into immature neurons and neuronal migration are necessary for the proper development of the central nervous system (CNS). The cerebral cortex of embryonic mice contains two distinct types of NSPCs: paired box 6-positive (Pax6+) apical progenitor cells (radial glia), located in the ventricular zone (VZ), and T-box brain protein 2-positive (Tbr2+) basal progenitor cells (intermediate progenitor cells), which are located in the subventricular zone (SVZ) (Englund et al., 2005). In the neocortex, newborn neurons generated from NSPCs migrate radially toward the cortical plate, accompanied by sequential changes in cell shape. Neurite outgrowth and ensuing polarity formation in immature neurons are also required for cortical layer stratification, and defects in neuronal migration not only cause brain malformation but also various psychiatric disorders, including epilepsy and mental retardation (Hansen et al., 2017; Represa, 2019).

Purines, compounds containing a pyrimidine ring fused with an imidazole ring, are found in all living species and include the nucleobases adenine and guanine (Traut, 1994). Apart from their critical function as the building blocks of DNA (deoxyadenosine and deoxyguanosine) and RNA (adenosine and guanosine), purines work as components of essential biomolecules and as a source of second messenger molecules (cyclic AMP and cyclic GMP), cofactors coenzyme A and nicotinamide adenine dinucleotide (NADH), cellular energy substrate ATP, and GTP, which is essential for the signal transduction of a large number of G-proteins. Other purine derivatives contain hypoxanthine, xanthine, and uric acid. Specifically, purines function as neurotransmitters in the brain by acting upon purinergic receptors. Purine metabolites, including ATP and GTP/GDP, are crucial for polarity formation in postmitotic cortical neurons (Raman et al., 2018). During brain development, purinergic signaling is essential for NSPC maintenance and neuronal migration in the neocortical SVZ (Lin et al., 2007; Liu et al., 2008).

In mammalian cells, purine content is regulated by a coordinated balance between the de novo and salvage biosynthetic pathways. Although the cellular purine pool is usually supplied by the recycling of degraded bases via the salvage pathway, the de novo pathway is upregulated under cellular conditions demanding higher levels of purines and their derivative nucleotides, such as tumor growth and cell proliferation (Yamaoka et al., 1997).
De novo purine synthesis comprises a series of 10 enzymatic reactions and is mediated by six evolutionarily conserved enzymes (phosphoribosyl pyrophosphate amidotransferase [PPAT], phosphoribosylglycinamidine formyltransferase [GART], formylglycin-amidine ribonucleotide synthase [FGAMS], phosphoribosylaminomimidazole carboxylase phosphoribosylaminomimidazole succinocarboxamide synthetase [PAICS], adenylosuccinate lyase [ADSL], and 5-aminimidazole-4-carboxamide ribonucleotide formyltransferase inosine monophosphate [IMP] cyclohydrolase [ATIC]), to produce IMP from phosphoribosylpyrophosphate (Baresova et al., 2018). The enzymes that catalyze de novo purine synthesis are assembled near mitochondria and microtubules as a huge multienzyme complex called “purinosome” (An et al., 2008, 2010; French et al., 2016). Purinosome is a dynamic and functional giant protein complex that emerges during high levels of cellular purine demand in mammalian cultured cells (An et al., 2008). Purinosome formation is linked to cell division (Chan et al., 2015). Furthermore, the dynamic assembly and disassembly of purinosomes in vivo might be crucial for the proper development of the human brain. Mutations in ADSL and ATIC genes cause severe developmental brain defects, such as mental retardation, autistic features, epilepsy, microcephaly, and congenital blindness (Jurecka et al., 2015; Marie et al., 2004). The bifunctional enzyme PAICS, another component of the purinosome, is associated with prostate and breast cancer metastasis and proliferation (Barrfeld et al., 2015; Chakravarthi et al., 2018; Meng et al., 2018). PAICS deficiency in humans was recently reported. A missense mutation in Paics causes the severe phenotype with multiple malformations, including a small body, short neck, and craniofacial dysmorphism, resulting in early neonatal death (Pelet et al., 2019). To date, however, there is no direct evidence of the localization or physiological function of purinosomes during brain development.

Previously, we identified the NACHT and WD repeat domain-containing protein 1 (Nwd1) gene and showed that the Nwd1 protein is expressed in NSPCs and immature neurons in the cerebral cortex of embryonic mice (Yamada and Sakakibara, 2018). The Nwd1 protein contains a NACHT domain, which is predicted to have nucleoside-triphosphatase (NTPase) activity, in the central region and a cluster of WD40 repeats at the C terminus. Based on the domain structure, Nwd1 is designated as a member of signal transduction ATPases with numerous domains (STAND) protein superfamily and is conserved across species, including zebrafish, mice, rats, monkeys, and humans (Yamada and Sakakibara, 2018). Other members of the STAND protein family often mediate ligand-induced self-oligomerization to form the giant multiprotein complex critical for various important cellular responses; e.g., the apoptotic peptidase activating factor 1 (Apaf1) and nucleotide-binding oligomerization domain-like receptors (NLRs) induce the assembly of large multiprotein complexes, the “apoptosome” and “inflammasome,” and play central roles in cell death and innate immune responses, respectively (Cai et al., 2017; Dorstyn et al., 2018; Leipe et al., 2004). Although the cellular function of Nwd1 remains unclear, its domain structure is analogous to Apaf1, an essential molecule for apoptosome assembly, which is required for apoptosis initiation (Dorstyn et al., 2018; Yamada and Sakakibara, 2018). Our current study shows that Nwd1 regulates NSPC proliferation and neuronal migration through the control of purinosome formation during cortical development. These findings would shed light on a machinery governing the purine metabolism in nervous system.

RESULTS

Nwd1 Overexpression In Vivo Increases the NSPCs Fraction and Delays the Radial Migration of Immature Neuron

To investigate the role of Nwd1 in the developing cerebral cortex, we overexpressed the Nwd1 gene in vivo using in utero electroporation. Full-length Nwd1 or control EGFP was electroporated into NSPCs in the developing dorsal neocortex at E14.5, a stage at which extensive neurogenesis and neuronal migration occurs. Electroporated embryos were harvested and analyzed after 48 h (at E16.5). To visualize the electroporated cells, the EGFP reporter plasmid was co-electroporated with the Nwd1 plasmid into the same embryos. Figures 1A–1C show that Nwd1 overexpression significantly suppressed neuronal migration from VZ, causing the accumulation of Nwd1-overexpressing cells in VZ/SVZ (control, 16.5 ± 4.2%, n = 6; Nwd1,
Figure 1. Dysregulation of Nwd1 Disturbs the Radial Migration of Neurons and Directs NSPCs to Reside in the VZ/SVZ
(A–K) Nwd1 overexpression in the embryonic neocortex. (A–C) In utero electroporation of control EGFP (A) or Nwd1 together with EGFP (B) was performed on E14.5, and the neocortex was analyzed at E16.5. (C) Distribution of electroporated EGFP+ cells in the indicated areas. *p < 0.05, ***p < 0.001. Welch’s t test followed by Holm-Bonferroni correction. (D–H) EGFP-Nwd1 (E) or control EGFP (D) was electroporated at E14.5, and brains were immunostained for Nestin (red) at E16.5. (F and G) Higher magnification of the VZ cells expressing EGFP-Nwd1 (F) and Nestin (G). (H) Ratio of EGFP+ or EGFP-Nwd1+ cells to the total number of Nestin+ cells in the VZ. ***p < 0.001, Welch’s t test. (I–K) Nwd1 or control EGFP was electroporated at E14.5, and the brains were collected on E18.5. (K) Distribution of EGFP+ cells in the indicated areas. *p < 0.05, **p < 0.01, Welch’s t test followed by Holm-Bonferroni correction. (L–S) Nwd1 knockdown represses neuronal migration. An Nwd1 shRNA (shRNA #1 or shRNA #2) was co-electroporated with EGFP at E14.5 and cortices were analyzed at E16.5 (L–O) or E18.5 (P–S). (O) Distribution of EGFP+ cells in the indicated areas at E18.5. *p < 0.05, **p < 0.01, ***p < 0.001, Welch’s t test followed by Holm-Bonferroni correction. All data are presented as means ± SEM. Scale bars, 50 μm in (A), (B), (D), (E), and (I)–(N); 5 μm in (F) and (G); 100 μm in (I), (J), and (P)–(R).

73.7 ± 6.0%, n = 6). At E16.5, the majority of cells electroporated with the control EGFP plasmid had migrated and reached the intermediate zone (IZ) and cortical plate (CP), where they became positive for Tbr1, a marker for post-mitotic neurons in the deep cortical layers and subplate (IZ, 72.3 ± 2.5%; CP, 11.2 ± 3.3%) (Figures 1A and S2A–S2C). However, Nwd1-overexpressing cells were rarely observed within the CP (Figures 1B, 1C, and S2D–S2F). Many Nwd1+ cells remaining in the VZ/SVZ were positive for the neural stem cell marker Nestin (Figures 1D–1H) (control, 29.0 ± 6.0%, n = 4; EGFP-Nwd1, 73.8 ± 4.8%, n = 4), suggesting that they retained their NSPC nature and lined the ventricular wall for at least 2 days, without moving toward the pial surface. After 4 days (at E18.5), EGFP expression was observed in the Brn2+ upper cortical layers (II–IV) and was almost absent in the Brn2+ deep cortical layers (V and VI) or IZ in controls (Figures 1I and S2J–S2L). At this time, cells overexpressing Nwd1 remained in the lower layers of the neocortex, including IZ and SVZ (Figures 1J and 1K) (layers II–IV: control, 80.5 ± 1.2%, n = 4 versus Nwd1, 38.4 ± 6.1%, n = 4; layers V–VI: control, 8.3 ± 1.9% vs. Nwd1, 27.6 ± 2.7%; IZ: control, 11.3 ± 1.8% vs. Nwd1, 34.0 ± 5.7%). Within the Brn2+ lower cortical layers, Nwd1-overexpressing cells exhibited the elongated bipolar morphology of traveling immature neurons (Figures 1J and S2M–S2O). These observations indicated that Nwd1 overexpression causes a significant increase in the Nestin+ NSPC pool accumulating in the VZ/SVZ and delays the radial migration of immature neurons.

Nwd1 Knockdown Causes Premature Differentiation of NSPCs and Represses Neuronal Migration

We explored the effect of Nwd1 loss of function on cortical development using small hairpin RNA (shRNA) delivery in vivo via in utero electroporation. Two different shRNA constructs (shRNA #1 and shRNA #2) targeting the coding region of mouse Nwd1 significantly reduced Nwd1 protein expression levels (Figure S1A). shRNA specificity was further demonstrated by Nwd1 immunostaining. Endogenous Nwd1 protein expression in cultured NSPCs was silenced by shRNA constructs (Figures S1B–S1E). We co-electroporated one of the shRNA constructs with an EGFP-expression plasmid into the neocortex at E14.5 and harvested embryos after 2 or 4 days. Then, we assessed the distribution of EGFP+ cells among the discrete cortical zones. In control embryos, almost all EGFP+ cells were found in either IZ or CP, and only a small fraction of cells was observed in VZ/SVZ at E16.5 (Figure 1L). However, Nwd1 knockdown (KD) resulted in a drastically reduced cell migration into the Tbr1+ CP as a large number of cells remained in VZ/SVZ at E16.5 (% of cells in the VZ/SVZ: control, 15.7 ± 3.7%, n = 8; Nwd1 shRNA #1, 45.9 ± 4.2%, n = 4; Nwd1 shRNA #2, 42.2 ± 5.7%, n = 5) (Figures 1L–1O and S2G–S2I). We noticed that Nwd1 KD cells accumulated in IZ if they were unable to penetrate the boundary between IZ and CP (Figures 1M and 1N). This phenotype was more pronounced after a further 2 days of shRNA expression. At E18.5, Nwd1 KD caused a significant accumulation of cells in IZ (control, 5.4 ± 2.0%, n = 4; Nwd1 shRNA #1, 56.8 ± 7.8%, n = 5) (Figures 1P–1S). Consequently, fewer Nwd1 KD cells reached the upper cortical layers (layers II–IV, control, 89.4 ± 1.3%, n = 4; Nwd1 shRNA #1, 32.0 ± 7.1%, n = 5; layers V–VI: control, 5.2 ± 1.0%; Nwd1 shRNA #1, 11.3 ± 1.4%). Many Nwd1 KD cells were still observed as Brn2+ cells in the deep cortical layers (Figures 1P–1S and S2Q). These defects were rescued by overexpression of the human NWD1 homolog, which is resistant to targeting by the mouse Nwd1 shRNA. We co-electroporated Nwd1 shRNA and the full-length human NWD1 cDNA into the E14.5 cerebral cortex and performed analysis at E18.5. A large fraction of the electroporated cells reached the upper cortical layers through IZ (Figures 1Q and 1S) (% of cells in layers II–IV, 79.0 ± 2.4%; layers V–VI, 14.8 ± 1.3%; IZ, 6.3 ± 1.6%, n = 10), restoring the cellular distribution comparable with that of the non-targeting control (see above). This finding further supported the notion that the loss of function of Nwd1 causes a severe migratory defect in immature neurons in vivo.

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We previously reported the substantial expression levels of Nwd1 in the VZ/SVZ and immature neurons (Yamada and Sakakibara, 2018). Accordingly, a larger number of cells overexpressing Nwd1 remained within VZ/SVZ (Figure 1). Thus, we examined whether Nwd1 KD affected the nature of the NSPC pool in VZ/SVZ. At E18.5, i.e., 4 days after shRNA electroporation, double immunostaining revealed that the Nwd1 KD cells remaining in VZ/SVZ were positive for doublecortin (Dcx) (control, 13.6 ± 3.7%, n = 7; Nwd1, 68.7 ± 9.2%, n = 5) (Figures 2A–2F and 2S) and β-tubulin III (Figures S3A–S3D), which are markers of newborn immature neurons. Interestingly, Nwd1 KD drove many VZ cells to ectopically and prematurely express Tbr2, a marker of the SVZ basal progenitor cells (intermediate progenitor cells) (control, 6.0 ± 3.9%, n = 4; Nwd1, 24.7 ± 4.7%, n = 4) (Figures 2G–2L and 2T). Concurrently, we observed a decreased density of Pdx6* apical progenitors in the VZ region, where Nwd1 shRNA was expressed (control, 47.2 ± 3.0%, n = 5; Nwd1, 25.6 ± 4.2%, n = 6) (Figures 2M–2S). Immunostaining for the mitotic marker Ki67 revealed that the proliferation rate of NSPCs was significantly reduced by Nwd1 KD (control, 27.1 ± 1.7%, n = 6; shRNA #1, 14.4 ± 2.3%, n = 6) (Figures 2U and 2V). Terminal deoxynucleotidyl transferase dUTP nick-end labeling (TUNEL) staining of E16.5 brains revealed that Nwd1 KD did not induce apoptosis (control shRNA, 0.7 ± 0.3%, n = 8; Nwd1 shRNA, 1.2 ± 0.3%, n = 8) (Figures S3E–S3I). To directly assess the effect of Nwd1 KD on NSPC proliferation, Nwd1 shRNA or control shRNA was transferred into primary cultured NSPCs then labeled with bromodeoxyuridine (BrdU). Following in vitro BrdU administration for 24 h, NSPCs were fixed after 0 or 24 h and the BrdU+ cells were counted. Nwd1 KD significantly decreased the number of BrdU+ cells in relation to the total number of EGFP+ cells. Data are presented as means ± SEM. **p < 0.01, control shRNA (green), which remained in ZV/SVZ areas at E18.5. The VZ surface is outlined by the dashed line. (S) Quantification of EGFP+ Dcx* or EGFP+ Pax6* cells to total EGFP+ cells in VZ/SVZ. Data are presented as means ± SEM. *p < 0.05, Welch’s t test. (T) Quantification of EGFP+ Tbr2+ cells to total EGFP+ cells in VZ. Data are presented as means ± SEM. **p < 0.01, Welch’s t test. (U and V) An Nwd1 shRNA (U, right panel) or control shRNA (U, left panel) was co-electroporated with EGFP at E14.5, and the cortices were harvested at E18.5, i.e., 4 days after shRNA electroporation, double immunostaining revealed that the Nwd1 KD cells remaining in VZ/SVZ were positive for doublecortin (Dcx) (control, 13.6 ± 3.7%, n = 7; Nwd1, 68.7 ± 9.2%, n = 5) (Figures 2U and 2V). Terminal deoxynucleotidyl transferase dUTP nick-end labeling (TUNEL) staining of E16.5 brains revealed that Nwd1 KD did not induce apoptosis (control shRNA, 0.7 ± 0.3%, n = 8; Nwd1 shRNA, 1.2 ± 0.3%, n = 8) (Figures S3E–S3I). To directly assess the effect of Nwd1 KD on NSPC proliferation, Nwd1 shRNA or control shRNA was transferred into primary cultured NSPCs then labeled with bromodeoxyuridine (BrdU). Following in vitro BrdU administration for 24 h, NSPCs were fixed after 0 or 24 h and the BrdU+ cells were counted. Nwd1 KD significantly decreased the number of BrdU+ cells (labeling index at 0 h: approximately 45%; 24 h: approximately 35%) relative to the control (0 h: approximately 80%; 24 h: approximately 70%) (Figures 2W and 2X). Conversely, the silencing of Nwd1 in NSPCs resulted in an accelerated commitment to the neuronal lineage. A differentiation assay of NSPCs indicated that Nwd1 KD increased the fraction of β-tubulin III+ neurons (Figures 2Y and 2Z). Based on the early onset of lineage markers and decreased cell division rate in vivo and in vitro, we concluded that the loss of function of Nwd1 induced the cell-cycle exit and premature neuronal differentiation of NSPCs. Abnormally produced progenies might follow neuronal differentiation near their place of birth, without proper cell migration, leading the apparent accumulation of EGFP+ cells within the VZ/SVZ in addition to the i2 (Figures 1O and 2F).
cell density than the neighboring SVZ region and brighter nuclei, evidenced by Hoechst and hematoxylin staining (Figures 3A–3D). Most heterotopia-forming cells had large round cell bodies with few fine processes (Figures 3E and 3H), resembling neurons. Indeed, double immunostaining revealed that they were Dcx+ neuron (Figures 3E–3J). Notably, these neurons expressed the vesicular glutamate transporter 1 (VGlutT1), and their somata were also closely surrounded by multiple VGlutT1+ presynaptic terminals (Figures 3K–3P), implying the formation of excitatory circuits of glutamatergic neurons within a heterotopia. Conversely, these cells never exhibited labeling of the astrocyte marker GFAP (Figures 3Q–3V). Cellular architecture of this malformation was similar to the human periventricular nodular heterotopia composed of hyperexcitable neurons, which is a developmental cortical dysgenesis frequently characterized by focal drug-resistant epilepsy (Battaglia et al., 2006).

Expression Levels of Nwd1 Is Crucial for Neurite Outgrowth and Axon Formation of Cortical Neurons

To understand Nwd1 cellular function in postmitotic differentiating neurons, loss-of-function and gain-of-function experiments were performed using primary cultured cortical neurons (Figure S5A). Nwd1 shRNA constructs were transferred into dissociated neurons prepared from E16.5 embryos and cultured for 3 days in vitro (div). In the control, a large fraction (~75%) of cells extended a single long axon immunostained for...
Figure 4. Tightly Regulated Nwd1 Expression Is Required for the Induction of Neuronal Identity

(A–G) A non-targeting shRNA (A–C) or Nwd1 shRNAs (D–F) were electroporated together with EGFP into primary cultured cortical neurons. Neurofilaments were stained with an anti-SMI312 antibody (red) at 3 div. Nuclei were stained with Hoechst dye (blue). (G) Number of SMI312+ axons extending from a single neuron. **p < 0.01 (chi-square test); control shRNA, n = 101; shRNA #1, n = 108; shRNA #2, n = 99.
Nwd1 Protein Interacts with Paics

We attempted to understand the molecular mechanism by which Nwd1 regulates cortical development. We used a Y2H screen to identify proteins interacting with Nwd1. Based on its structural similarity to other STAND-family proteins (Leipe et al., 2004), we hypothesized that the N-terminal region of Nwd1 serves as an effector domain by which the protein binds signaling molecule(s) to trigger self-oligomerization mediated by the NACHT domain and WD40 repeats. The N-terminal region of Nwd1 contains a DUF4062 motif, a functionally uncharacterized motif found in bacteria and eukaryotes (Yamada and Sakakibara, 2018). The principal function of this gene in the morphological transformation of neurons during neurogenesis remains unclear. The defects in neuronal migration caused by manipulating Nwd1 might reflect a principal function of this gene in the morphological transformation of neurons during neurogenesis.

Nwd1 and Paics Are Localized in Purinosomes

We investigated the localization of Paics in the embryonic and postnatal mouse brain. An immunostaining analysis using an anti-Paics antibody showed high levels of Paics expression in the developing brain. At E10.5, Paics immunoreactivity was uniformly detected in the undifferentiated NSPCs in the telencephalon, extending from the VZ to the outer part of the neural tube (Figure 5E). We also noticed elevated Nwd1

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**Figure 4. Continued**

(H–N) Cortical neurons transfected with EGFP-Nwd1 (K–M) or control EGFP (H–J) were stained for SMI312 (red) at 3 div. Nuclei (blue) (N) Number of SMI312+ neurons extending from a single neuron. ***p < 0.001 (chi-square test); EGFP, n = 149, EGFP-Nwd1, n = 149. (O–U) EGFP-Nwd1 or control EGFP were electroporated into cortical neurons and cultured for 1 div (O and R), 2 div (P, S), and 3 div (Q and T). To visualize fine immature neurites, a dsRed expression plasmid was co-electroporated into the cells. (U) Number of neurites extending from a single neuron. Data are presented as means ± SEM. ***p < 0.01, Welch’s t test; EGFP, n = 150, 2 div, n = 150; 3 div, n = 200; EGFP-Nwd1 1 div, n = 150, 2 div, n = 150; 3 div, n = 300. Scale bars, 20 μm.

the SMI312 neurofilament marker (Figures 4A–4C). At this time, neurons transfected with Nwd1 shRNA #1 and shRNA #2 exhibited fewer SMI312+ axons (Figures 4D–4F), and a notable number of cells lacked axons (Figure 4G). We assessed whether Nwd1 overexpression affected axonal extension in each cortical neuron. Control cells electroporated with EGFP usually had a single axon after culture for 3 div (Figures 4H–4J). In contrast, EGFP-Nwd1 expression inhibited axonal extension (Figures 4K–4M). These cells occasionally had few short neurites that were devoid of SMI312 immunoreactivity. To visualize all immature neurites extending directly from soma, a plasmid encoding the red fluorescent protein dsRed was co-electroporated into the cortical neurons and the total number of neural processes was counted as neurites. We found that EGFP-Nwd1 overexpression reduced the number of neurites by almost half at 1, 2, and 3 div (Figures 4O–4U) (1 div: control, 4.0 ± 0.2; EGFP-Nwd1, 2.4 ± 0.2; 2 div: control, 4.6 ± 0.2; EGFP-Nwd1, 2.6 ± 0.2; 3 div: control, 5.3 ± 0.1; EGFP-Nwd1, 2.6 ± 0.2). Compared with the control, each neurite appeared thinner and unbranched, suggesting early stage of neurite development. Consistently, in the embryonic cortex electroporated with Nwd1 shRNAs in utero, we observed a significant number of apolar cells within the IZ, which had round cell bodies, and it appeared as if they had failed to transform into migratory spindle-shaped neurons (arrows in Figure 5L). These results indicated that Nwd1 plays a vital role in axon formation in newborn neurons and that the highly controlled and just sufficient level of Nwd1 may be essential for axon and neurite outgrowth. Recent studies have indicated that dynamic changes in cell shape is closely coupled with the neuronal migration and cortical layer formation (Hirota and Nakajima, 2017). In the developing mammalian neocortex, newborn neurons transiently become multipolar cells with multiple neurites inside SVZ and lower IZ; thereafter, they undergo a change in morphology to a bipolar state before the onset of radial migration to the CP (Ohtaka-Maruyama and Okado, 2015); however, its molecular mechanism remains unclear. The defects in neuronal migration caused by manipulating Nwd1 might reflect a principal function of this gene in the morphological transformation of neurons during neurogenesis.
that a significant proportion of the endogenous Paics protein colocalizes in these clusters (Figures 7E, 7F, rinosomes (HSP90 in the human prostate cancer cell line, LNCaP (Correa et al., 2014). Taken together, these results strongly indicate that Nwd1 is a component of purinosomes. Conceivably, Nwd1 may act in cooperation with the chaperone machinery in purinosome assembly or stabilization.

A previous study that used co-IP with FGAMS followed by a proteomics analysis demonstrated that heat shock protein 90 (Hsp90) and Hsp70 colocalize in the purinosome (French et al., 2013). Knockdown of these chaperones leads to the disruption of purinosomes, implying the involvement of Hsp90/Hsp70 chaperone machinery in the protein complex assembly (French et al., 2013). The molecular chaperones Hsp70 and Hsp90 are ubiquitously expressed proteins that have many functions, including assisting in protein folding and stabilizing protein complexes (Makhnevych and Houry, 2012); however, their exact function in purinosome formation remains unclear (Pedley and Benkovic, 2017). To examine the colocalization of Nwd1 and Hsp90 in purinosomes, the distribution of EGFP-Nwd1 and HA-tagged Hsp90 (Hsp90-HA) was assessed in purine-depleted cells; we observed overlapping localization of Nwd1 in purinosomes (arrows in Figures 6E and 6K).

A previous study that used co-IP with FGAMS followed by a proteomics analysis demonstrated that heat shock protein 90 (Hsp90) and Hsp70 colocalize in the purinosome (French et al., 2013). Knockdown of these chaperones leads to the disruption of purinosomes, implying the involvement of Hsp90/Hsp70 chaperone machinery in the protein complex assembly (French et al., 2013). The molecular chaperones Hsp70 and Hsp90 are ubiquitously expressed proteins that have many functions, including assisting in protein folding and stabilizing protein complexes (Makhnevych and Houry, 2012); however, their exact function in purinosome formation remains unclear (Pedley and Benkovic, 2017). To examine the colocalization of Nwd1 and Hsp90 in purinosomes, the distribution of EGFP-Nwd1 and HA-tagged Hsp90 (Hsp90-HA) was assessed in purine-depleted cells; we observed overlapping localization of Nwd1 in purinosomes (arrows in Figures 6E and 6K). A co-IP assay of HEK293 cells expressing FLAG-Nwd1 and Hsp90-HA demonstrated an interaction between Nwd1 and Hsp90 (Figure 6Q). Consistent with this, Correa et al. showed that NWD1 binds to HSP90 in the human prostate cancer cell line, LNCaP (Correa et al., 2014). Taken together, these results strongly indicate that Nwd1 is a component of purinosomes. Conceivably, Nwd1 may act in cooperation with the chaperone machinery in purinosome assembly or stabilization.

**Purinosome Assembly Is Regulated by Nwd1 in NSPCs**

To date, there is no evidence of the induction of purinosome assembly in nervous tissues. Therefore, next, we investigated whether NSPCs are capable of forming purinosomes and whether Nwd1 localizes in these structures in NSPCs. NSPCs isolated from the E12.5 cerebral cortex and cultured as a monolayer frequently exhibit Nestin+ fine unipolar or bipolar processes, resulting in a morphology that resembles that of neuroepithelial cells in the embryonic VZ (Figures S6B and S6E). The expression of Fgams-EGFP distinctly emerged as a granular structure (Figures 7A–7D). Immunostaining using the anti-Paics antibody indicated that a significant proportion of the endogenous Paics protein colocalizes in these clusters (Figures 7E, 7F, and S6G–S6R). The colocalization of Fgams and Paics, which are two enzymes that are essential for de novo purine biosynthesis, strongly suggested that these clusters are functional purinosomes in the NSPCs.
Purinosomes were often observed within the cellular processes, in addition to the cell body, under the plasma membrane of NSPCs (Figure 7F). Immunostaining with an anti-Nwd1 antibody revealed the localization of the endogenous Nwd1 protein in Fgams-EGFP+ purinosomes in NSPCs (Figures 7G–7L). The purinosome localization of Nwd1 became more evident after the introduction of EGFP-Nwd1 into NSPCs (Figure 7N). These data showed the presence of purinosomes in NSPCs.

To examine the role of Nwd1 in purinosome assembly, Nwd1 expression was suppressed by shRNA in NSPCs. Nwd1 shRNA constructs were electroporated into NSPCs expressing Fgams-EGFP. At 2 div, we counted the number of cells containing the functional purinosomes that are defined as the granules simultaneously labeled with Fgams-EGFP and endogenous Paics. As shown in Figure 7V, compared with the non-targeting shRNA (Figures 7P–7R), Nwd1 shRNAs reduced the number of cells containing Fgams-EGFP+ Paics+ purinosomes considerably (control, 58.3 ± 3.0%; shRNA #1, 35.0 ± 1.5%; shRNA #2, 34.7 ± 4.8%). We noticed that Nwd1 KD resulted in a decreased number of Paics+ granules in NSPCs, whereas the total number of exogenously introduced Fgams-EGFP+ granules remained
unchanged (Figure S6S). Consistent with this, the fraction of cells that were labeled with Fgams-EGFP alone (Fgams-EGFP + Paics–) was increased in NSPCs upon treatment with the shRNAs (arrowheads in Figure 7U). Because a protein complex lacking Paics no longer functions as a purinosome, we concluded that Nwd1 is required for the assembly of the functional purinosome in NSPCs.

Purinosome Enzymes Are Essential for Cortical Development

To clarify the involvement of the purinosome in brain development, we examined the loss-of-function or gain-of-function phenotypes of Paics and Fgams. First, E14.5 embryos were electroporated in utero with

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**Figure 7. Nwd1 Regulates Purinosome Assembly in NSPCs**

(A–C) NSPCs derived from E12.5 telencephalons underwent electroporation with Fgams-EGFP (A) and were immunostained with an anti-Paics antibody (B, red) at 2 div. (C) Merged view. Nuclei (blue).

(D–F) Higher magnification of the boxed area depicted in (C), showing the clustered signals of Fgams+ (D) Paics+ (E) purinosomes (arrows) in NSPCs. (F) Merged view.

(G–L) Fgams-EGFP-expressing (G) NSPCs were immunostained with an anti-Nwd1 antibody (H). (I) Merged view. Nuclei (blue). (J–L) Higher magnification of the boxed area depicted in (I), demonstrating the localization of endogenous Nwd1 (K) in Fgams-EGFP+ (J) purinosomes. (L) Merged view.

(M) EGFP-Nwd1-expressing NSPCs were immunostained with an anti-Paics antibody.

(N and O) Higher magnification of the boxed area depicted in (M), showing the colocalization of EGFP-Nwd1 (N) and endogenous Paics (O) in purinosomes.

(P–V) NSPCs were electroporated with the control shRNA (P–R) or Nwd1 shRNAs (S–U) together with Fgams-EGFP, followed by immunostaining with an anti-Paics antibody at 2 div. The arrows indicate the Fgams-EGFP+ Paics+ functional purinosomes in NSPCs. The arrowheads indicate the Fgams+ Paics– cells. (V) Number of NSPCs containing Fgams-EGFP+ Paics+ purinosomes. Data are presented as means ± SEM. **p < 0.01, Welch’s t test followed by Holm-Bonferroni correction.

Scale bars, 20 μm in (A)–(C) and (G)–(I), (M); 4 μm in (D)–(F), (J)–(L), and (P)–(U); 2 μm in (N)–(O).
Nwd1 as a Novel Component of Purinosomes

Here, we described the induction of the formation of Fgams+ Paics+ purinosomes in NSPCs. We also revealed that Nwd1 interacts with Paics and is localized in purinosomes in NSPCs. Nwd1 functions as a component in the assembly of purinosomes. Nevertheless, Nwd1 has no enzymatic activity related to purine biosynthesis, unlike Paics and Fgams. It is possible that Nwd1 participates in the assembly of purinosomes as a member of the STAND family of proteins. The STAND proteins are a newly recognized family of proteins that mediate the energy-dependent remodeling of proteins and the translocation of macromolecules. Generally, STAND genes encode multidomain proteins, typically encompassing an N-terminal effector domain, a middle domain, and a C-terminal ligand-binding domain (Leipe et al., 2004). The binding of specific ligands onto the C-terminal domain elicits a conformational change in STAND proteins, which is dependent on ATP levels; this results in
the formation of the oligomeric ring-shaped superstructures of STAND proteins, which exhibit a central pore (Mermigka et al., 2020). Such superstructures serve as the tightly regulated molecular switch that controls diverse biological processes, including apoptosis and innate immune responses, in which the ring-like superstructures of STAND proteins drive the translocation or remodeling of the substrate proteins (Mermigka et al., 2020).

Among the STAND-family proteins, Nwd1 shares a similar domain structure with Apaf1 (Dorstyn et al., 2018; Leipe et al., 2004; Yamada and Sakakibara, 2018). During apoptosis, the C-terminal WD40 domain of Apaf1 binds to the cytochrome c molecules that leaked from damaged mitochondria. This ligand binding induces the energy-dependent self-oligomerization of Apaf1. Subsequently, the ring-like superstructure of Apaf1 tethers caspase 9 through the N-terminal CARD domain of Apaf1 to form the macromolecular complex named apoptosome, which triggers the apoptotic caspase cascade (Zou et al., 1999). Similarly, Paics is assembled as a homo-octameric structure in purinosomes (Li et al., 2007), similar to the caspase

Figure 9. Models of Nwd1 Function in Corticogenesis and in Purinosome Formation
(A) Purinosome assembly/disassembly regulates cortical development. Overexpression of purinosome components (Nwd1 or Fgams) disturbs the radial migration of neurons and causes a significant increase in the Nestin+ NSPC pool in the VZ/SVZ. Knockdown of purinosome components (Nwd1 or Paics) accelerates mitotic exit and premature differentiation of NSPCs and represses neuronal migration leading to periventricular heterotopia. EGFP+ cells represent cells harboring the transgene.
(B) Hypothetical molecular model of purinosome formation by Nwd1. Undiscovered signals trigger the interaction of Nwd1 with Paics via the DUF4062 and AIRc domains, mediating the formation of the multimeric structure of Nwd1–Paics. The Nwd1–Paics complex systematically tethers other core enzymes, including Fgams, Ppat, and Gart, to form a functional purinosome. Nwd1, NACHT and WD repeat domain-containing protein 1; Ppat, phosphoribosyl pyrophosphate amidotransferase; Gart, phosphoribosylglycinamidine formyltransferase; Fgams, formylglycin-amidine ribonucleotide synthase; Paics, phosphoribosylaminomimidazole carboxylase phosphoribosylaminoimidazole succinocarboxamide synthetase; Adsl, adenylosuccinate lyase; Atic, 5-aminomimidazole-4-carboxamide ribonucleotide formyltransferase imosine monophosphate cyclohydrolase; PRPP, phosphoribosyl diphosphate; IMP, inosine monophosphate; Hsp90, heat shock protein 90.
9 heptamer in apoptosomes. Based on these observations, we postulated that Nwd1 undergoes an ATP-dependent conformational change upon binding to the ligand(s), via which Paics proteins are triggered to be recruited to a purinosome (Figure 9B). Primary complex of Nwd1/Paics and several chaperones including Hsp90 might systematically tether other core enzymes, including Fgams, Ppat, and Gart to form an integrated purinosome (Figure 9B). This complicated molecular machinery may explain why the in vitro reconstitution of a functional purinosome has been unsuccessful thus far. Nwd1 may act as a sensor protein that drives the assembly and disassembly of purinosomes and activates the de novo purine biosynthesis pathway during CNS development.

**Purinosome Components Regulate the Maintenance of NSPCs and Neuronal Migration during Cortical Development**

We reported previously the strong expression of Nwd1 in NSPCs and immature neurons during the development of the rodent brain (Yamada and Sakakibara, 2018). The present study revealed a similar distribution of Paics and Nwd1 in the developing neocortex. The gain and loss of function of Nwd1, Paics, and Fgams, which were achieved using in utero electroporation, demonstrated that these purinosome components are essential for proper cortical development and that their dysregulation leads to a severe delay in the migration of immature neurons (Figure 9A). In addition, in vivo knockdown of Nwd1 resulted in a decrease in the number of Pax6+ apical progenitors, in conjunction with the ectopic emergence of Tbr2+ basal progenitor cells in the embryonic VZ. A previous study reported that the forced expression of the Tbr2 transcription factor directs the conversion of radial glia into basal progenitor cells (Sessa et al., 2008). Thus, we assumed that the altered expression level of Nwd1 caused the premature differentiation of NSPCs, suggesting a vital role for this protein in the maintenance of NSPC pools, including CNS stem cells (Figure 9A). Consistently, a previous study suggested a possible role for Nwd1 in tumor cells endowed with stem-cell-like properties; i.e., the proliferative and self-renewing properties. The expression of Nwd1 is strikingly upregulated by Sox9, a transcription factor in malignant prostate tumor cells (Correa et al., 2014). A gain- and loss-of-function study indicated that Sox9 plays a central role in the specification and maintenance of CNS stem cells that reside in the embryonic VZ and adult SVZ (Scott et al., 2010). As a downstream target of Sox9, Nwd1 may have a function in the maintenance of CNS stem cells. Interestingly, it was also demonstrated that Paics is necessary for the proliferation and invasion of prostate cancer cells and that the silencing of Paics expression abrogates the progression of several types of prostate tumors (Chakravarthi et al., 2018). Taken together with this evidence, our findings imply that the formation of the purinosome machinery is crucial for the maintenance of somatic stem cells and tumor cells, which commonly require a large amount of de novo purine production.

In addition, we demonstrated that a tight control of the level of expression of Nwd1 is crucial for neurite extension and axon formation and that altered levels of expression of the Nwd1 gene caused migration defects in cortical neurons in vivo. Considering that the spatiotemporally controlled outgrowth of neurites is needed for the establishment of neuronal polarity and neuronal migration (Hansen et al., 2017), purinosome formation might be closely linked to the dynamic morphological transformation of migrating neurons that occurs during corticogenesis. Purines affect many aspects of neuronal differentiation. For example, the activation of Rac, which is a small GTP-binding protein, is required for the formation of the leading process in radially migrating neurons in the embryonic cerebral cortex (Konno et al., 2005). Nwd1 might affect discrete aspects of neural development, including neuronal migration and the maintenance of the NSPC pool, via the regulation of the assembly/disassembly of purinosomes.

However, de novo purine synthesis is energy intensive and required for numerous substrates; therefore, it has been thought that terminally differentiated neurons place greater reliance on the purine salvage pathway than the de novo pathway to achieve prompt repair of damaged DNA and prevent neurodegeneration. A previous in vitro quantitative analysis of purines using neuroblastoma cell lines demonstrated that the intracellular purine content increases as neuronal differentiation proceeds, whereas de novo purine synthesis decreases during neuronal differentiation (Gottle et al., 2013), suggesting that a regulated balance between the de novo and purine salvage pathway is critical for coordinated neuronal differentiation. However, the switching molecules that determine this balance remain unidentified. Our current study provides an insight into the machinery governing purine metabolism during nervous system development.

**Implication of Nwd1 and Purinosome Components in Neurological Disorders**

Downregulation of Nwd1 by shRNA expression in the embryonic cerebral cortex often caused the cortical dysgenesis similar to human periventricular nodular heterotopia (PH), a cortical malformation that is
characterized by the formation of ectopic aggregates of neurons that line the lateral ventricle. These nodules exhibited a rosette-like structure and were filled with the glutamatergic neurons innervated by VgulT1+ excitatory terminals, indicating the formation of abnormal excitatory circuit. In humans, PH is associated with intractable epilepsy and intellectual disability (Cossu et al., 2018). Previous studies using a genetic animal model showed that PH is caused by the failure of the radial migration of newborn neurons from the VZ in addition to the abnormal proliferation of NSPCs (Li et al., 2015; Lian and Sheen, 2015); however, the molecular mechanisms underlying the development of PH are not fully understood. Thus, the disturbance of the purine de novo synthesis pathway may be associated, at least in part, with the mechanism underlying the pathogenesis of PH.

In addition to PH, purinosome-related genes are responsible for certain neurological disorders. Deficiency of ADSL in humans causes atrophy of distinct regions of the brain, including the cerebral cortex, in addition to hypomyelination and lissencephaly (Jurecka et al., 2015). Patients with ATIC mutation exhibit neurological symptoms, including profound mental retardation and epilepsy accompanied by various dysmorphic features (Marie et al., 2004). A previous study that used cultured fibroblasts from these patients demonstrated that ATIC and ADSL mutations destabilize the assembly of the purinosome to various degrees and that the ability to form purinosomes is correlated with the severity of the phenotype of individual patients (Baresova et al., 2012). Recently, PAICS deficiency was reported in humans. Patients carrying a homozygous missense mutation in the PAICS gene exhibit multiple severe malformations, including a small body and craniofacial dysmorphism, resulting in early neonatal death (Pelet et al., 2019). Although inactivating mutations in the human NWD1 gene have not been reported to date, it was recently shown that the neuronal expression of NWD1 is upregulated in patients with temporal lobe epilepsy (Yang et al., 2019). Using a mouse model of acute epileptic seizures, it was suggested that Nwd1 regulates the neuronal hyperexcitability of glutamatergic synaptic transmission in the adult brain (Yang et al., 2019). Therefore, Nwd1 might be involved in a mechanism of regulation of the synaptic transmission via the formation of purinosomes or other macromolecular complexes.

Limitations of the Study
In the present study, we revealed that Nwd1 interacts with Paics to regulate purinosome assembly in NSPCs, whereas the formation of purinosomes in terminally differentiated neurons has not been demonstrated in vitro and in vivo. We are currently analyzing the function of purinosomes in differentiated neurons. These findings will be published in our future study. In addition, there is no clear evidence for the mechanism underlying the premature differentiation of NSPCs induced by Nwd1 gene silencing. Because the de novo purine synthesis pathway is generally upregulated under cellular conditions that demand higher levels of purines, such as tumor growth, it might be reasonable to suppose that the exhaustion of the purine pool in dividing NSPCs leads them to exit the cell cycle. Such an unexpected quiescent status among NSPCs may provoke the induction of fate-determining genes, such as Ascl1 and Nkx2-1, resulting in premature neuronal differentiation.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101058.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

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AUTHOR CONTRIBUTIONS
All authors had full access to the data in this study and take responsibility for the integrity of the data and accuracy of the analysis. Study concept and design: S.Y. and S.S. Acquisition of data: S.Y., A.S., and S.S. Analysis and interpretation of data: S.Y., A.S., and S.S. Drafting of the manuscript: S.Y. and S.S. Obtained funding: S.S.
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Supplemental Information

Nwd1 Regulates Neuronal Differentiation and Migration through Purinosome Formation in the Developing Cerebral Cortex

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Figure S1. Validation of the *Nwd1* shRNAs. Related to Figure 1.

(A) The non-targeting control or *Nwd1* shRNA (shRNA #1 or shRNA #2) was transfected into HEK293 cells that expressed EGFP-*Nwd1* exogenously. The expression level of *Nwd1* was evaluated by immunoblotting using an anti-*Nwd1* antibody. Immunoblotting using an α-tubulin antibody was performed to ensure the equal loading of cell lysates.

(B–E) The control shRNA or *Nwd1* shRNA #1 was transfected into the primary culture of NSPCs, together with EGFP. Cells were immunostained with an anti-*Nwd1* antibody at 2 div, which showed that the expression of endogenous *Nwd1* (red) was silenced by the *Nwd1* shRNA. Scale bars, 20 μm.
Figure S2
**Figure S2. Dysregulated expression of Nwd1 suppresses the neuronal migration.**

**Related to Figure 1.**

E14.5 embryos were electroporated in utero with non-targeting control (A–C, J–L), EGFP-Nwd1 (D–F, M–O) or Nwd1 shRNA #1 (G–I, P–R) or, and brains were harvested at E16.5 (A–I) and E18.5 (J–R). The shRNA constructs were co-electroporated with EGFP. (A–I) E16.5 brain sections were immunostained with anti-Tbr1 antibody (red), a marker for the deep cortical layer and subplate. Dashed lines represent the border between CP and subplate. Arrow in (B) indicates the Tbr1+ subplate. (J–R) E18.5 brain sections were immunostained with anti-Brn2 antibody (red), a marker for the upper cortical layers. Dashed lines denote the borders of upper layers (II–IV), deep cortical layers (V–VI) and IZ. Scale bars, 100 μm.
Figure S3
Figure S3. Nwd1 knockdown causes premature differentiation of NSPCs. Related to Figure 2.

The non-targeting control or Nwd1 shRNA #1 were electroporated into E14.5 brains along with EGFP, and embryos were harvested at E16.5 and E18.5. (A–D) Confocal images of a neocortex at E18.5 stained with an anti-β-tubulin III (red) antibody. The areas surrounded by dashed line denote the distribution of cells electroporated with the Nwd1 shRNA (green) within the VZ. (E–I) Apoptotic cells were detected by TUNEL staining (red) at E16.5. (I) Number of EGFP+ TUNEL+ apoptotic cells. Control shRNA, n=8; Nwd1 shRNA, n=8. Data are presented as means ± SEM. NS, not significant, Welch’s t-test. (J–M) Confocal projection images of E18.5 cerebral cortex. Note that the significant number of Nwd1 KD cells exhibited an apolar and round morphology (arrows) within the IZ. Nuclei are counterstained with Hoechst dye (blue). Scale bars, 50 μm.
Figure S4

Figure S4. Periventricular heterotopia caused by Nwd1 knockdown. Related to Figure 3.

The Nwd1 shRNA was electroporated into the neocortex at E14.5 together with EGFP, and brains were collected at P7. Representative confocal image of a coronal section, showing the ectopic formation of periventricular heterotopia composed of densely packed EGFP+ cells (A) located under the neocortex (arrow) in the electroporated hemisphere. Nuclei were stained with Hoechst dye (B). Scale bar, 500 μm.
Figure S5. Nwd1 expression is required for axonal extension, Related to Figure 4

(A–C) Primary cortical neurons prepared from E16.5 embryos were cultured for 3 days (3 div) and double-immunostained with a dendritic marker, MAP2 (green), and an axon marker, SMI312 (red), confirming dendritic arborization and the extension of a single axon from differentiated neurons. (D–U) Non-targeting shRNA or Nwd1 shRNA were electroporated together with EGFP into the cortical neurons, and each cell was immunostained with an SMI312 antibody (red) at 3 div. Inhibition of Nwd1 expression impaired axonal extension from immature neurons. Scale bars, 20 μm.
Figure S6

![Figure S6](image_url)
Figure S6. Purinosome formation in NSPCs. Related to Figure 7.

(A–F) Primary cultured NSPCs prepared from E12.5 telencephalons were electroporated with non-targeting shRNA or Nwd1 shRNA, together with EGFP and then immunostained with Nestin antibody (red). (G–R) NSPCs electroporated with Fgams-EGFP were immunostained with anti-Paics antibody (red). Each panel shows the high-power view of individual Fgams-EGFP+ cells. (J–L) The higher magnification of the boxed area in (I) represents the formation of functional purinosomes that emerged as four clusters positive for Fgams-EGFP+ and endogenous Paics+ (arrows) in the distal region of the NSPC process. (M–R) Arrows indicate Fgams-EGFP+ Paics+ purinosomes emerged in cell process of NSPCs. (S) NSPCs were electroporated with Fgams-EGFP together with control or Nwd1 shRNA, followed by immunostaining with anti-Paics antibody at 2 div. The graph shows the number of Fgams-EGFP+ or Paics+ granular signals in NSPCs. Data are presented as means ± SEM. NS, not significant, *p<0.05, Welch’s t-test followed by Holm–Bonferroni correction. Scale bars, 20 μm in A–F and G–I, M–R; 4 μm in J–L.
Figure S7. Validation of Paics shRNAs. Related to Figure 8.
N2a cells were transfected with three different mouse Paics shRNA constructs (shRNA #1, shRNA #2, and shRNA #3) or a non-targeting control shRNA, followed by immunoblotting with an anti-PAICS (A) or anti-α-tubulin (B) antibody.
Figure S8

**A**

Bar graph showing the percentage of positive cells for each marker in control shRNA and Paics shRNA conditions in VZ/SVZ.

**E16.5 (electroporation E14.5)**

- **B**
  - EGFP/Nestin

- **C**
  - EGFP

- **D**
  - Nestin

**E**

Bar graph showing the ratio of EGFP+ cells to Nestin+ cells in the VZ.

- EGFP
- Fgams-EGFP

Significance levels: NS, *p < 0.05, **p < 0.01, ***p < 0.001.
Figure S8. Purinosome components regulate the cortical development. Related to Figure 8.

(A) Control shRNA or Paics shRNA was delivered into the brain on E14.5, together with EGFP, and the cortices were stained with anti-Dcx, anti-Pax6, anti-Tbr2, and anti-Ki67 antibodies at E16.5. Graph shows the quantification of the positive cells for each marker to total EGFP+ cells in the VZ/SVZ. NS, not significant, *p<0.05, **p<0.01, ***p<0.001. Note that the knockdown of Paics gene accelerates the mitotic exit of Pac6+ or Tbr2+ NSPCs and induces the premature expression of neuron marker Dcx. (B–E) Fgams-EGFP was electroporated into the E14.5 neocortex and embryos were analyzed at E16.5. The cerebral cortex was stained with an anti-Nestin antibody (red). (C, D) Magnified view of the boxed area depicted in (B), showing that Fgams-EGFP+ cells were persistently located in the VZ/SVZ as Nestin+ NSPCs. (E) Quantification of Fgams-EGFP+ cells to total Nestin+ cells in VZ. *p<0.05. All data are presented as means ± SEM. Statistical significance value was determined using the Welch’s t-test. Scale bars, 100 μm in B; 10 μm in C and D.
Table S1

| Gene name                                      | Gene Symbol | NCBI Reference Sequence | Identified protein region (amino acids) | Protein function                                         |
|-----------------------------------------------|-------------|-------------------------|----------------------------------------|---------------------------------------------------------|
| Abhd3 domain containing 3                    | Abhd3       | NP 598891.1             | 350–411 aa                             | unknown, paralog of Abhd1                                |
| ATP-binding cassette, sub-family D (ALD), member 3 | Abcd3       | NP 033017.2             | 530–659 aa                             | peroxisomal import of fatty acids and/or fatty acyl-CoA   |
| chymotrypsin-like elastase family, member 1   | Celi1       | NP 291090.2             | 14–155 aa                              | protease associated with elastin remodeling               |
| clavematin 2                                  | Cls2        | NP 001346068.1          | 35–181 aa                              | recycling of synaptic vesicles                           |
| Kin17 DNA and RNA binding protein             | Kin         | NP 079556.1             | 1–188 aa                               | DNA/RNA binding protein                                  |
| phosphoribosylaminomimidazole carboxylase     | Piacs       | NP 080215.1             | 323–425 aa, 312–425 aa, 305–425 aa     | de novo purine synthesis enzymes                         |
| phosphoribosylaminobethylaminomimidazole, succinocarboxamide synthetase | Piacs       | NP 080215.1             | 323–425 aa, 312–425 aa, 305–425 aa     | de novo purine synthesis enzymes                         |
| opaline                                      | Opk         | NP 001152988.1          | 120–319 aa                             | RNA-binding protein                                      |
| serine (or cysteine) peptidase inhibitor, clade E, member 2 | Serpin2     | NP 033281.1             | 307–397 aa                             | inhibitor for serine proteases                           |
| serine palmitoyltransferase, small subunit A  | Sptnas      | NP 598815.2             | 10–71 aa                               | serine palmitoyltransferase isozymes                     |
| E26 avian leukemic oncogene 1, 5' domain      | Eto1        | NP 001359463.1          | 1–111 aa                               | transcription factor                                     |
| tripeptidyl peptidase II                     | Tpp2        | NP 033444.1             | 1171–1261 aa                           | serine exopeptidase                                      |
| WD repeat domain 74                          | Wdr74       | NP 598900.1             | 114–378 aa                             | regulator of exosome complex formation                   |

**Table S1.** Nwd1-binding partners identified by yeast two-hybrid screen. Related to Figure 5.

Each protein entry is shown with the respective NCBI accession number and the identified protein region. The respective protein functions are also listed.
Transparent Methods

Animals

ICR male mice were purchased from Japan SLC Inc. (Shizuoka, Japan). The date of conception was established by the presence of a vaginal plug and recorded as embryonic day zero (E0). The day of birth was designated as P0. Mice were housed under temperature- and humidity-controlled conditions on a 12/12 hr light/dark cycle, with ad libitum access to food and water. All protocols were approved by the Committee on the Ethics of Animal Experiments of Waseda University.

Tissue Preparation

Embryos at E16.5 and E18.5 were perfused through the cardiac ventricle with 4% paraformaldehyde (PFA) in 0.1 M phosphate buffer (pH 7.4), followed by post-fixation overnight at 4ºC (Yamada and Sakakibara, 2018). Fixed embryo brains were cryoprotected in 30% sucrose in phosphate-buffered saline (PBS) overnight at 4ºC and embedded in optimal cutting temperature compound (Sakura Finetek). Frozen sections were cut at a thickness of 14 µm using a cryostat and were collected on MAS coated glass slides (Matsunami Glass).

Plasmid Vectors

Mouse Nwd1 cDNAs were subcloned into the pEGFP-C2 vector (Clontech Takara Bio) to express the Nwd1 protein fused with EGFP. The Nwd1 and EGFP-Nwd1 cDNAs were subcloned into the pCAGGS vector (a gift from Dr. Jun-ichi Miyazaki, Osaka University, Japan). For the yeast two-hybrid (Y2H) screening, Nwd1 cDNAs corresponding to the N-terminal portion of the protein (accession number BC082552; 4 bp–1026 bp) were
subcloned into pGBKT7 (Clontech Takara Bio) to express the N-terminal domain of Nwd1 fused with the GAL4 DNA-binding domain. The Flag-tagged human NWD1 expression plasmid was provided by Dr. Correa (Sanford-Burnham Medical Research Institute, Canada) (Correa et al., 2014). pFGAMS-EGFP and pPAICS-EGFP (# 99108) were gifts from Dr. Stephen Benkovic (Addgene plasmids # 99107 and # 99108, respectively) (An et al., 2008). HSP90-HA was a gift from Dr. William Sessa (Addgene plasmid #22487) (Garcia-Cardena et al., 1998). pCAG-DsRED was a gift from Dr. Connie Cepko (Addgene plasmid # 11151) (Matsuda and Cepko, 2004).

**shRNA Expression Vectors**

We purchased a MISSION shRNA vector library encoding the microRNA-adapted shRNA targeting mouse *Nwd1* (Sigma-Aldrich). Among five shRNA clones (TRCN0000257630, TRCN0000247062, TRCN0000257635, TRCN0000257616, and TRCN0000179877), TRCN0000247062 and TRCN0000257635 yielded efficient knockdown of the exogenous Nwd1 and EGFP-Nwd1 expressed in cultured cells; these clones were designated as shRNA #1 and shRNA #2, respectively. The targeting sequences of shRNA #1 and shRNA #2 were: 5\'-TACGACTGTGCATGCTCTAAA-3' and 5\'-CAGGTAATCCAAGTTCGATAT-3', respectively. The two constructs targeted the coding region of the Nwd1 mRNA. We also used MISSION shRNA plasmids for mouse *Paics* (Sigma-Aldrich). Among five clones (TRCN0000076100, TRCN0000076101, TRCN0000076102, TRCN0000076098, and TRCN0000076099), TRCN0000076101, TRCN0000076102, and TRCN0000076098 were designated as shRNA #1, shRNA #2, and shRNA #3, respectively. The targeting sequences for shRNA #1, shRNA #2, and shRNA #3 were 5\'-CTGCTCAGATATTGGTTAA-3', 5\'-
GCTGATGTCATTGATAA and 5’-GCACCTGCCTTCAATACTAT-3’, respectively. shRNA #1 and shRNA #2 targeted the coding region, whereas shRNA #3 targeted the 3’ untranslated region (3’-UTR) of the Paics mRNA. A non-targeting shRNA (# SHC202) was also purchased from Sigma-Aldrich.

**Primary Antibodies**

The following primary antibodies were used: anti-Nwd1 (affinity-purified rabbit polyclonal antibody used previously (Yamada and Sakakibara, 2018), 1:200 for immunostaining, 1:2000 for immunoblotting), anti-Nwd1 (rabbit polyclonal antibody generated by immunizing the recombinant mouse Nwd1 protein; 1:500 for immunostaining, 1:5000 for immunoblotting), anti-Nestin (chicken polyclonal IgY, Aves Labs, NES; 1:4000 for immunostaining), anti-Nestin (rabbit polyclonal, IBL, 18741; 1:250), anti-α-tubulin (rabbit polyclonal, MBL, PM054; 1:2000 for immunoblotting), anti-GFP (chicken polyclonal IgY, Aves Labs, GFP-1010; 1:2000 for immunostaining), anti-GFP (rabbit polyclonal, GeneTex , GTX113617; 1:2000 for immunoblotting), anti-doublecortin (DCX) (goat polyclonal, Santa Cruz, sc-271390; 1:200 for immunostaining), anti-β-tubulin III (chicken polyclonal IgY, AVES Labs, TUJ; 1:1000 for immunostaining), anti-Pax6 (rabbit polyclonal, MBL, PD022; 1:1000 for immunostaining), anti-Tbr2 (chicken polyclonal, Merck Millipore, 633572; 1:1000 for immunostaining), anti-Ki67 (rabbit monoclonal clone SP6, Lab Vision, RM-9106; 1:1000 for immunostaining), anti-Paics (rabbit polyclonal, Proteintech, 12967-1-AP; 1:200 for immunostaining), anti-GFAP (mouse monoclonal clone G-A-5, Sigma-Aldrich, G3893; 1:400 for immunostaining), anti-SMI312 (mouse monoclonal, Biolegend, 837904; 1:1000, for immunostaining), anti-HA (rabbit polyclonal, MBL, 561; 1:200 for
immunostaining, 1:2000 for immunoblotting), and anti-DDDDK (Flag) (mouse monoclonal, MBL, FLA-1; 1:10000 for immunostaining and immunoblotting), anti-VGluT1 (rabbit polyclonal, GeneTex, GTX133148; 1:250 for immunostaining), anti-Tbr1 (rabbit polyclonal, GeneTex, GTX117615; 1:250 for immunostaining), anti-Brn2 (rabbit polyclonal, GeneTex, GTX114650; 1:250 for immunostaining), anti-MAP2 (chicken polyclonal IgY, Biolegend, 822591, 1:1000 for immunostaining), anti-BrdU (sheep polyclonal, Abcam, 1:1500 for immunostaining).

**Cell Culture**

HEK293T, HeLa, and Neuro2a (N2a) cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM) containing 10% fetal bovine serum (FBS), penicillin/streptomycin, and L-glutamine. In purine-depleted conditions, HeLa cells were cultured in RPMI 1640 medium supplemented with L-glutamine and 5% dialyzed FBS. FBS was dialyzed against 0.9% NaCl using a 10K MWCO SnakeSkin Tubing (Thermo Fisher Scientific) for 48 hr at 4°C (An et al., 2008; French et al., 2013). NSPCs were isolated from the E12.5 telencephalon, seeded onto dishes coated with fibronectin and polyethylenimine (PEI) (Sigma-Aldrich), and cultured in Advanced DMEM/F-12 (1:1) (Life Technologies) supplemented with 15 μg/mL insulin (Life Technologies), 25 μg/mL transferrin (Life Technologies), 20 nM progesterone (Sigma-Aldrich), 30 nM sodium selenite (Sigma-Aldrich), 60 nM putrescine (Sigma-Aldrich), 20 ng/mL basic fibroblast growth factor (FGF2) (Merck Millipore), and 10 ng/mL epidermal growth factor (Merck Millipore) (Yamada and Sakakibara, 2018; Yumoto et al., 2013). To assess cell proliferation, 10 μM BrdU (5-bromodeoxyuridine, Tokyo Chemical Industry) was administrated to the primary cultured NSPCs. After 24 hr of incubation, cells were cultivated in the medium
without BrdU for the indicated periods, and then processed for immunostaining using anti-BrdU antibody. For the differentiation assay of NSPCs, culture medium was changed from the proliferation medium to the EGF and FGF2-free differentiation medium containing 1% FBS. For the primary culture of cortical neurons, embryonic cerebral cortices at E16.5 were dissected and mechanically dissociated. After washing with Opti-MEM I (Life Technologies), cells were electroporated, seeded onto poly-D-lysine-coated dishes, and cultured in neurobasal medium containing 2% B27 (Life Technologies) and 1% GlutaMax (Life Technologies) for 1–3 days in vitro (div). For immunostaining, cultured cells were fixed with 4% PFA for 20 min at 4°C and permeabilized in 0.05% Triton X-100 in PBS for 10 min.

**Cell Transfection**

Cultured cell lines were transfected with plasmid DNA and PEI MAX (Polysciences) complexes (ratio of DNA to PEI MAX, 1:3, w/w) formed in Opti-MEM I by incubation for 15 min at room temperature. The DNA complexes were added to cell cultures together with Opti-MEM I for 3 hr, followed by cultivation with serum containing complete DMEM. Mouse NSPCs were expanded in vitro as described above, and primary cortical neurons were electroporated using a NEPA21 Electroporator (Nepagene) according to the manufacturer’s specifications (NSPCs: two pulses of 125 V for 5 ms with an interval of 50 ms; primary cortical neurons, two pulses of 275 V for 0.5 ms with an interval of 50 ms).

**In utero Electroporation**
Pregnant mice were anesthetized via intraperitoneal injection of a mixture containing medetomidine, midazolam, and butorphanol (Kawai et al., 2011). A DNA solution (5 μg/μL) in PBS with 0.01% Fast Green dye (Sigma-Aldrich) was injected into the lateral ventricle through the uterus wall, followed by electroporation (Yumoto et al., 2013). The following constructs were electroporated: pCAG-Nwd1, pCAG-EGFP, pCAG-EGFP-Nwd1, pFgams-EGFP, Nwd1 shRNAs, Paics shRNAs, and non-targeting shRNA. Electric pulses were generated by NEPA21 (Nepagene) and applied to the cerebral wall using a platinum oval electrode (CUY650P5, Nepagene), with four pulses of 35 V for 50 ms with an interval of 950 ms. An anionic electrode was placed on the lateral cortex, to ensure the incorporation of DNA into the VZ/SVZ. Embryos were perfused at E16.5, E18.5, and P7 with 4% PFA through cardiac perfusion.

**Immunostaining**

Frozen sections were blocked for 2 hr with 5% normal goat or donkey serum in PBST (0.1% Triton X-100 in PBS), followed by the incubation with primary antibodies in blocking buffer at 4°C overnight (Yamada and Sakakibara, 2018). After washing with PBST four times, sections were incubated for 2 hr with Alexa Fluor 488-, Alexa Fluor 555- (Life Technologies), or DyLight 488-, DyLight 549- (Jackson ImmunoResearch) conjugated secondary antibodies. After counterstaining with 0.7 nM Hoechst 33342 (Life Technologies), sections were mounted and imaged using a confocal (FV3000, Olympus) or fluorescence inverted microscope (Axio Observer, Zeiss). For BrdU staining, fixed cells were treated with 2 N HCl for 1 hr at 37°C to denature the DNA, followed by the treatment with 0.1 M sodium borate buffer (pH 8.5) for 10 min (Sakakibara et al., 2001). After washing with PBS, anti-BrdU immunostaining was performed.
Immunoprecipitation and Western Blotting

For immunoprecipitation (IP), cells were washed in ice-cold PBS and lysed in ice-cold lysis buffer containing 50 mM Tris-HCl (pH 7.5), 150 mM NaCl, 2 mM EDTA, 1% NP-40, and protease inhibitors (cOmplete Mini Protease Inhibitor Cocktail, Roche Diagnostics) for 30 min at 4°C. Lysates were centrifuged at 15,000 rpm for 10 min and the supernatants were precleared with TrueBlot anti-rabbit or anti-mouse IP beads (Rockland Immunochemicals) for 1 hr. After centrifugation at 15,000 rpm for 10 min, the supernatants were incubated with the primary antibody coupled to TrueBlot IP beads (Rockland Immunochemicals) overnight or 1 hr at 4°C. Rabbit or mouse IgG (Thermo Fisher Scientific) was used as a control. After brief centrifugation, beads were washed four times with 0.1% NP40 in PBS, and then the bound proteins were dissolved by treatment with 2×sample buffer (125 mM Tris-HCl, pH 6.8, 4% SDS, 10% sucrose, and 0.01% bromophenol blue). IP samples were resolved on 8% or 10% SDS-PAGE, and electroblotted onto Immobilon-P membranes (Merck Millipore) using a semidy transfer apparatus. After blocking with 5% skim milk in TBST (150 mM NaCl, 10 mM Tris–HCl, pH 7.4, 0.1% Tween 20), membranes were incubated with primary antibody for 1 hr, followed by incubation with the horseradish peroxidase (HRP)-conjugated secondary antibody (GE Healthcare). The signal was detected using the Immobilon western chemiluminescent HRP substrate (Merck Millipore) and visualized by Fusion Solo S (Vilber Lourmat).

TUNEL staining
TUNEL (Terminal deoxynucleotidyl transferase dUTP nick end labeling) assay was performed for detecting apoptosis using the In Situ Cell Death Detection Kit, TMR-red (Roche Diagnostics), according to the manufacturer’s instruction.

**Yeast Two-hybrid Screening**

To identify Nwd1-binding proteins in the CNS, a Y2H screening was performed using the Matchmaker Gold Yeast Two-Hybrid System (Clontech Takara Bio), according to the manufacturer’s instructions. The sequence encoding the 340 N-terminal amino acid residues of mouse Nwd1 (BC082552; 4–1026 bp) was subcloned in frame into pGBK7, to express the N-terminal region of Nwd1 (bait) fused with the GAL4 DNA-binding domain. As the prey, we used the normalized mouse brain library (Clontech Takara Bio, Normalized Mate & Plate Library, cat. # 630488), in which each clone was fused with the Gal4 DNA-activating domain (AD). Mated yeast clones were selected using minimal synthetic defined (SD) medium with double dropout (Leu− and Trp−) supplement (Clontech Takara Bio) containing Aureobasidin A and X-α-Gal as the blue-colored colonies; this was followed by a second screening using the SD quadruple dropout (Leu−, Trp−, Ade−, and His−) selective medium (Clontech Takara Bio). After the elimination of duplicates containing the same AD/library plasmid via yeast-colony PCR, plasmids were rescued from yeast using the Easy Yeast Plasmid Isolation Kit (Clontech Takara Bio). Protein interactions were confirmed by co-transformation of the Nwd1 bait with each candidate prey plasmid into Y2H Gold yeast host cells, followed by sequencing of the cDNA inserts.

**Statistical Analyses**
All numerical data are expressed as means ± SEM. In two-group comparisons, Welch’s $t$-test was used to assess the significance of the differences in cell distribution over the cortical layers, positive cells for each marker, the number of neurites, or the number of purinosomes between different groups. In multiple-group comparisons, analysis of variance followed by Welch’s $t$-test was used. The $p$ values obtained were corrected for multiple testing using the Holm–Bonferroni correction. The chi-squared test was used for comparison with the number of BrdU$^+$ or β-tubulin III$^+$ cells in the NSPCs proliferation/differentiation assays, and the number of SMI312$^+$ axons counted in the neuronal differentiation assay.

**Supplemental References**

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