Hbp1 regulates the timing of neuronal differentiation during cortical development by controlling cell cycle progression

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Competing interests statement
The authors declare no competing financial interests.

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**Author contributions statement**

Naoki Watanabe: Developed the concepts, Performed experiments, Performed data analysis

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SUMMARY
In the developing mammalian brain, neural stem cells (NSCs) initially expand the progenitor pool by symmetric divisions. NSCs then shift from symmetric to asymmetric division and commence neurogenesis. Although the precise mechanisms regulating the developmental timing of this transition have not been fully elucidated, gradual elongation in the length of the cell cycle and coinciding accumulation of determinants that promote neuronal differentiation may function as a biological clock that regulates the onset of asymmetric division and neurogenesis. We conducted gene expression profiling of embryonic NSCs in the cortical regions and found that expression of high mobility group box transcription factor 1 (Hbp1) was upregulated during neurogenic stages. Induced conditional knockout mice of Hbp1 generated by crossing with Nestin-CreERT2 mice exhibited a remarkable dilatation of the telencephalic vesicles with a tangentially expanded ventricular zone and a thinner cortical plate containing reduced numbers of neurons. In these Hbp1-deficient mouse embryos, neural stem/progenitor cells continued to divide with a shorter cell cycle length. And downstream target genes of the Wnt signaling, such as cyclin D1 and c-jun, were upregulated in the germinal zone of the cortical regions. These results indicate that Hbp1 plays a critical role in regulating the timing of cortical neurogenesis by elongating the cell cycle and is essential for normal cortical development.
INTRODUCTION
During mammalian cortical development, neural stem cells (NSCs) gradually alter their characteristics and generate a variety of cell types, which establishes the functional complexity of the brain (Temple, 2001). In early developmental stages, a sheet of NSCs vigorously expands through repeated symmetric divisions, and the neural tube distends like a balloon with a thin wall composed of short neuroepithelial cells. After expanding the progenitor pool for several cycles, NSCs in the cortical regions begin to produce neurons by converting from symmetric to asymmetric division, which initiates neurogenesis (Takahashi et al., 1995, 1999). The timing of this transition is crucial for determining the overall number of NSCs and the size of ventricles, which largely determines the eventual size and morphology of the brain. However, the precise mechanisms that regulate the switch from symmetric to asymmetric division and the onset of neurogenesis have not been fully elucidated.

It has been reported that the length of the G1 phase of the cell cycle (T_{G1}) in neural stem/progenitor cells gradually rises over the course of cortical development. This elongation of T_{G1} is accompanied by downregulation of cyclin E and p21 along with upregulation of p27, cdk2, and cyclin B (Delalle et al., 1999; Caviness et al., 2003), and the forced reduction of T_{G1} by manipulation of cyclin D1 led to an expansion of neural progenitor cells in the developing and adult brain (Lange et al., 2009; Pilaz et al., 2009; Artegiani et al., 2011). One proposed mechanism for the transition from symmetric proliferative to asymmetric neurogenic division is that an elongation of T_{G1} allows fate determinants that promote neuronal differentiation to accumulate during the G1 phase and exert their neurogenic functions (Calegari and Huttner, 2003; Calegari et al., 2005; Götz and Huttner, 2005; Dehay and Kennedy, 2007).

To identify temporal alterations in the transcriptional properties of embryonic NSCs, we previously carried out DNA microarray-based gene expression profiling of embryonic NSCs prepared from the cortical regions at different developmental stages by using pHes1-d2EGFP transgenic mice, which express enhanced green fluorescent protein (EGFP) in NSCs (Ohtsuka et al., 2006, 2011). Among a variety of genes that were differentially expressed during the course of development, we hypothesized that high mobility group box transcription factor 1 (Hbp1) might be an important regulator of neurogenesis, given that it was upregulated during neurogenic stages around embryonic day 13.5 (E13.5) through embryonic day 15.5 (E15.5). Previous studies
demonstrated that Hbp1 acts as a transcriptional repressor that functions as a cell cycle inhibitor by repressing downstream targets of the Wnt signaling and cell cycle-related genes such as \textit{cyclin D1}, \textit{c-jun}, \textit{N-myc}, and \textit{Cdkn1a}, also known as \textit{p21} (Gartel et al., 1998; Sampson et al., 2001; Kim et al., 2006; Elfert et al., 2013; Yan et al., 2014).

Here, we analyzed the molecular function of Hbp1 in neuronal differentiation during cortical development. We found that Hbp1 controls the length of the cell cycle in neural stem/progenitor cells by modulating the expression levels of cyclin D1, thereby regulating the timing of neuronal differentiation during early cortical development.

\textbf{MATERIALS AND METHODS}

\textbf{In situ hybridization}

Preparation of Digoxigenin-labeled antisense RNA probes and \textit{in situ} hybridization were performed as described previously (Ohtsuka et al., 2011). We used partial-length cDNA of \textit{Neurog2} (NM_009718.2) (0.87 kb) and full-length cording sequence of \textit{Hbp1} (NM_153198.2), \textit{Neurod1} (NM_010894.2), \textit{Ascl1} (NM_008553.4), \textit{cyclin D1} (NM_007631.2), \textit{cyclin E1} (NM_007633.2) and \textit{N-myc} (NM_008709.3) as templates of the RNA probes.

\textbf{Quantitative real-time RT-PCR}

Total RNA samples were extracted from the FACS-sorted embryonic NSCs prepared from the cortical regions of \textit{pHes1-d2EGFP} transgenic mice as described previously (Ohtsuka et al., 2011). The reverse transcription of the total RNA and real-time RT-PCR were performed as previously described (Tan et al., 2012) using the primers listed in Table S1. \textit{β-Actin} was used as an internal control.

\textbf{Plasmid construction}

For overexpression of \textit{Hbp1}, full-length cording sequence of \textit{Hbp1} (NM_153198.2) was cloned into the \textit{pEF} (human elongation factor \textit{1α} promoter)-\textit{MM} expression vector, which was modified from \textit{pEF-BOS} vector (Mizushima et al., 1990). \textit{pCAGGS-HA-hRb1} vector, in which HA (hemagglutinin)-tagged human \textit{Rb1} was inserted into \textit{pCAGGS} vector, was kindly provided by Dr. Chiaki Takahashi and Dr. Nobunari Sasaki.
pCAG-HA-Hbp1 vector was made by inserting HA-tagged Hbp1 into pCAGEN (Matsuda et al., 2004). For reporter assay, cyclin D1 promoter region (from -3281 to +232) was inserted into pGL4.10 (Promega), and then we replaced the Luc2 in pGL4.10 for the ubiquitinated luciferase fused with two nuclear localization sequences (NLS2-Ub-Luc2). For knockdown experiments, expression vectors were generated from annealed oligonucleotides for shRNA targeting Hbp1 (shHbp1) or cyclin D1 (shCcnd1) (Hbp1:
5'-acctcGGACTCTTCTGCAGTGAACACTTcaagagACATAGACCAGACGATCTCCtt-3' and
5'-caaaaaGGACTCTTCTGCAGTGAACACTttcttgaACATAGACCAGACGATCTCCg-3'; cyclin D1:
5'-acctcGTGCATCTACACTGCAACTTcaagagAGTTGTGACGTAGATGCACtt-3' and
5'-caaaaaGTGCATCTACACTGCAACTttcttgaAGTTGTGACGTAGATGCACg-3') inserted into the BbsI site of psiRNA-h7SKneo G1 plasmid vector (InvivoGen) with the 20-21 nucleotide target sequence shown in uppercase letters, as described previously (Ohtsuka et al., 2011). Randomly scrambled sequence of the target was used for the negative control. The knockdown efficiency was confirmed in HEK293T cell lines.

In utero electroporation
Pregnant ICR mice were obtained from Japan SLC, Inc. (Shizuoka, Japan) and CLEA Japan, Inc. (Tokyo, Japan). In utero electroporation was performed with E13.5 ICR pregnant mice as described previously (Ohtsuka et al., 2011).

Immunohistochemistry
Immunohistochemistry was performed as described previously (Ohtsuka et al., 2011). The primary antibodies used were listed in Table S2. Primary antibodies were detected with Alexa Fluor-conjugated secondary antibodies (1:200; Molecular Probes). To visualize the cell nuclei, DAPI (4′,6-diamidino-2-phenylindole) (Sigma-Aldrich) was added on slides. Fluorescent staining was analyzed with LSM510 and LSM780 confocal microscopes (Zeiss).
**Analysis of cell cycle length**

To estimate the cell cycle length, we conducted a dual pulse labeling of DNA synthesis using BrdU (5-bromo-2’-deoxyuridine) (Sigma-Aldrich) and EdU (5-ethynyl-2’-deoxyuridine) (Molecular Probes), referring to the previous methods (Martynoga et al., 2005; Mairet-Coello et al., 2012). Previous reports used EdU first followed by BrdU, but here we administered BrdU first followed by EdU, and confirmed that the two reverse orders of injection produced the same results by using the anti-BrdU antibody MoBU-1 (Molecular Probes). BrdU (50 μg BrdU/g of body weight) and EdU (12.5 μg EdU/g of body weight) were injected intraperitoneally to the pregnant mice 2 hours and 30 minutes before sacrifice, respectively, and the ratios of cells that incorporated either or both BrdU and EdU were analyzed to estimate the cell cycle length. The detection of EdU labeled cells was performed based on a fluorogenic click reaction (Salic et al., 2008). Length of the S phase ($T_S$) and cell cycle length ($T_C$) were calculated as follows: $T_S = 1.5 \times S_{cells}/L_{cells}$ and $T_C = T_S \times P_{cells}/S_{cells}$ [$L_{cells} =$ the cells leaving the S-phase (identified as BrdU$^+$;EdU$^-$), $S_{cells} =$ the cells in the S-phase (EdU$^+$ only and double BrdU$^+$;EdU$^+$) and $P_{cells} =$ total proliferating cells (Ki67$^+$)].

**Reporter assay**

0.1 μg of the luciferase reporter of *cyclin D1* and 1.0 μg of the expression plasmids were transfected into HEK293T cells. 1 ng of *pRL-SV40* vector (Promega) was cotransfected to normalize the transfection efficiency. The cells were harvested after 48 hours and the reporter assay was performed as described previously (Sakamoto et al., 2003).

**Statistical analysis**

Each experiment was performed with at least 3 independent samples. Results are shown as mean ± standard error of the mean. Statistical differences were examined with Student’s $t$-test.
RESULTS

Hbp1 is expressed in the germinal zone during cortical neurogenesis

By evaluating our gene expression profiling data for embryonic NSCs in the developing cortex (Ohtsuka et al., 2011), we selected candidate genes expected to be involved in the regulation of neuronal differentiation based on their upregulation during the neurogenic period. Among these genes, we observed that *Hbp1* was prominently expressed in the germinal zone which comprises the ventricular zone (VZ) and subventricular zone (SVZ) in the developing telencephalon (Fig. 1A). The expression pattern of *Hbp1* was characterized by a lateral high-dorsal low gradient at E11.5 corresponding to the propagation of neurogenesis, and that was similar to the expression pattern of *Neurog2*, a proneural (neurogenic) basic helix-loop-helix (bHLH) gene (Fig. 1B). As it was difficult to accurately estimate alterations in *Hbp1* expression levels over the course of development by *in situ* hybridization, we performed real-time RT-PCR using total RNAs prepared from NSCs in the cortical regions of *pHes1-d2EGFP* transgenic mice at different embryonic stages and confirmed the temporal dynamics of *Hbp1* expression indicated by microarray analysis (Fig. 1C).

Overexpression of Hbp1 suppresses cell proliferation, but inhibits terminal neuronal differentiation

We next conducted overexpression experiments by *in utero* electroporation. *Hbp1* expression vectors (*pEF-Hbp1*) and control vectors (*pEF-EGFP*) were cointroduced into ventricular cells in the cortical regions of E13.5 mouse embryos, and the fates of transfected cells were subsequently examined at E14.5 and E16.5. As shown in our previous paper (Ohtsuka et al., 2011), cells transfected with *pEF-Hbp1* mainly remained in the SVZ and the intermediate zone (IZ) at E16.5. Noticeably, a majority of transfected cells differentiated into neurons positive for neuronal markers such as Tuj1 and Map2 and negative for Ki67, a marker of proliferating cells, or phospho-histone H3 (pH3), a marker of dividing cells in the M phase, but stagnated in the IZ and failed to migrate into the cortical plate (CP) (Fig. 2A; Fig. S1A-D), indicating that neuronal migration was inhibited. The proportions of transfected cells positive for Pax6, a specific marker of NSCs during this period of development, and Tbr2, a marker of intermediate progenitor cells (IPs) were significantly lower than in the control at E16.5. Transfected cells that incorporated BrdU administered 4 hours after *in utero*
electroporation at E13.5 exited the VZ earlier than control cells at E15.5 but the most cells still remained in the IZ at E16.5 (Fig. S1E,F), suggesting that overexpression of Hbp1 promoted initial neuronal differentiation but impaired terminal neuronal differentiation and migration.

Next, we evaluated the rate of cell proliferation by administering the thymidine analogue EdU intraperitoneally to the pregnant mice 30 minutes before sacrifice at E14.5 to mark cells in the S phase. In cells transfected with \textit{pEF-Hbp1}, EdU incorporation was suppressed compared to control (Fig. 2B), indicating that overexpression of Hbp1 caused an attenuation of cell proliferation or premature cell cycle exit. We then carried out a calculation of cell cycle kinetics by BrdU/EdU double labeling protocol modified from a previously described IdU/BrdU double labeling method (Martynoga et al., 2005; Mairet-Coello et al., 2012), and estimated the cell cycle length. Cells transfected with \textit{pEF-Hbp1} showed a slightly longer cell cycle length compared to control (Fig. 8F; Table S3); however, the effect on the cell cycle was minimal in comparison to \textit{Hbp1} knockdown or \textit{Hbp1}-deficient cells.

\textbf{Coexpression of Rb1 derepresses terminal neuronal differentiation and migration}

Hbp1 has been shown to be a target of retinoblastoma 1 (Rb1) and p130, also known as Rb2 (Tevosian et al., 1997). Shih et al. reported that muscle cell differentiation was blocked when Hbp1 was overexpressed in C2C12 cells without interfering with cell cycle exit, and the expression of MyoD and myogenin, but not Myf5, was inhibited in Hbp1-expressing cells; however, full differentiation occurred when Rb1 was coexpressed with Hbp1 and the ratio of Rb1/Hbp1 was elevated, suggesting that the relative ratio of Rb1 to Hbp1 is important as a determinant of whether cell cycle exit or full differentiation occurs (Shih et al., 1998). Thus, we introduced a mixture of \textit{pEF-Hbp1} and \textit{pCAGGS-HA-human Rb1 (hRb1)} vectors at ratios of 3:1 or 1:3 by \textit{in utero} electroporation, and found that transfected cells were released from the stagnation and significantly more transfected cells reached the CP, when Hbp1 and Rb1 were coexpressed (Fig. 2C,D).

Next, we examined the expression levels of proneural bHLH genes by \textit{in situ} hybridization 24 hours after \textit{in utero} electroporation. We found that \textit{Neurod1} expression was downregulated in the regions transfected with \textit{pEF-Hbp1} (Fig. 2E), and the repression was rescued to some extent by coexpression of Rb1, while expression of
Neurog2 was not significantly affected. Taken together, these results supported findings from a previous study of muscle cell differentiation (Shih et al., 1998) and suggested that overexpression of Hbp1 promoted cell cycle exit but impaired terminal neuronal differentiation required for full migration owing to the low Rb1/Hbp1 ratio, and conversely, a higher Rb1/Hbp1 ratio obtained by coexpressing Rb1 promoted terminal differentiation and neuronal migration.

Knockdown of Hbp1 inhibits neuronal differentiation and activates cell cycle progression

Next, we carried out knockdown experiments using expression vectors of shRNA targeting Hbp1 (shHbp1), in addition to scrambled shRNA control vectors (scrambled) (Fig. S2A,B). When we introduced shHbp1 into ventricular cells together with pEF-EGFP by in utero electroporation at E13.5, most cells transfected with shHbp1 remained in the VZ/SVZ/IZ with only a few cells observed in the CP at E16.5 (Fig. S2C). However, neuronal differentiation occurred later and most transfected cells in the IZ were Tuj1 and Map2-positive and negative for Ki67 or pH3 (Fig. S1A,C; Fig. S2C), although the number of pH3\(^+\) cells increased in the VZ and SVZ in regions transfected with shHbp1 (Fig. S1C,D). Knockdown cells that incorporated BrdU administered 4 hours after electroporation exhibited slower exits from the VZ than in the control at E15.5 (Fig. S1E,F), indicating that Hbp1 knockdown inhibited and delayed neuronal differentiation. While most cells transfected with scrambled shRNA differentiated into Cux1\(^+\) neurons (layer II-IV) and settled in the CP at E18.5, a majority of cells transfected with shHbp1 still remained in the VZ/SVZ/IZ (Fig. S2D). The number of Cux1\(^+\) cells in the superficial layers of the CP decreased markedly in the regions transfected with shHbp1, while the number of Ctip2\(^+\) cells (layer V) was comparable to the control.

To evaluate the specificity of the Hbp1 knockdown, we performed rescue experiments by coelectroporation of shHbp1 and pEF-Hbp1 at ratios of 3:1, 1:1, or 1:3, and found that pEF-Hbp1 rescued the phenotype of Hbp1 knockdown (inhibition of neuronal differentiation and migration) at a 1:1 ratio (Fig. S2E,F). However, the neuronal migration defect was observed again at a 1:3 ratio, similarly to our observations in the case of Hbp1 overexpression.
We performed immunostaining with anti-phosphorylated vimentin (p-Vim) antibody but did not observe a significant increase in p-Vim+ mitotic radial glial cells outside the VZ (Fig. S1C,D). In addition, we found that cell death (cleaved-caspase 3+) was not enhanced by either overexpression or knockdown of Hbp1 3 days after electroporation (Fig. S1G,H). Estimation of the cell cycle length revealed that Hbp1 knockdown cells showed a considerably shorter cell cycle length compared to control (Fig. 8F; Table S3), indicating that knockdown of Hbp1 shortened the cell cycle length, in particular the duration of the G2-M-G1 phase, and activated cell cycle progression.

**Deficiency of Hbp1 in neural stem/progenitor cells impairs cortical morphogenesis**

To analyze the function of Hbp1 over the course of brain development, we generated Hbp1 induced conditional knockout (icKO) mice, in which tamoxifen-inducible Cre recombinase is expressed specifically in neural stem/progenitor cells in the embryonic brain (see supplementary materials and methods; Fig. S3A,B). We confirmed by real-time RT-PCR that expression of Hbp1 in the telencephalon of Hbp1 icKO mice was downregulated by E11.5 when tamoxifen was administered at E9.5 (Fig. S3C).

The whole brain size of Hbp1 icKO embryos was comparable to that of the negative controls (NC), but the telencephalic wall appeared to be thinner (Fig. 3A). The telencephalic vesicles of mutant embryos were remarkably dilated, although the ventricular dilatation was not prominent at E12.5 (Fig. 3A,B). The volume of the ventral telencephalon was severely diminished and the wall of dorsal telencephalon was significantly thinner than that of the control after E14.5 (Fig. 3B). Notably, the ventricular surface area in the dorsolateral telencephalon (cortical region) was substantially enlarged; thus, the VZ appeared to be stretched in the tangential direction. The thickness of both the VZ (Pax6+) and the neuronal layer (Tuj1+) were thinner than in the control. The midline and medial structures of the telencephalon, such as hippocampus and corpus callosum, were hypoplastic or missing at E18.5. Results of the measurement of each structure were summarized in Fig. 3C (see supplementary materials and methods).

**Hbp1-deficiency leads to delayed neuronal differentiation and tangential expansion of neural stem cells**

Next, we investigated the dynamics of cortical development in the Hbp1-deficient mice.
We determined the number of distinct cell types through immunolabeling with various cell- and layer-specific markers within a radial column of constant width (200 μm) in the cortical regions (Fig. 4A-C; Fig. S4A,B). At E12.5, the thickness of the VZ (Pax6+ and Hes1+) was similar to that in the control mice. Noticeably, the number of Tbr2+ cells and early born neurons (Tuj1+ and Tbr1+) was reduced in the icKO mice at E12.5, indicating that the onset of neurogenesis was delayed. Instead, the NSC population was still expanding by symmetric divisions, thus leading to the tangential extension of the VZ and ventricular dilatation that became prominent after E14.5. We determined the number of Pax6+ NSCs throughout the cortical regions in multiple coronal sections at constant intervals along the antero-posterior axis of lateral ventricle. Through this method, we estimated and compared the total number of NSCs in the hemisphere of control and mutant brain, and found that the total number of NSCs was increased in the icKO at E14.5 (Fig. 3C).

As the generation of Tbr2+ IPs and neurons commences after the transition from symmetric to asymmetric division mode of NSCs (Sessa et al., 2008), these results suggested that the Hbp1-deficiency delayed this transition and the onset of neurogenesis. Thus, we analyzed the fates of cells after cell divisions by administering EdU 12 hours before sacrifice (Fig. 5A). A majority of cells that incorporated EdU remained Pax6+ NSCs in the cortical regions of icKO embryos at E12.5, while the number of EdU+ cells that differentiated into Tbr2+ IPs was much higher in the control (Fig. 5B). Given the significantly higher frequency of cell divisions in Pax6+ cells than in Tbr2+ cells, as demonstrated by pH3 staining (Fig. 5C), these results indicated that the symmetric proliferative division of Pax6+ NSCs were maintained in the icKO at a population level, although it was not revealed at a single cell level, whereas many NSCs had shifted to asymmetric neurogenic division in the control at E12.5. Estimation of the ratios of Ki67+;EdU+/EdU+ (cell cycle re-entry) and Ki67+;EdU+/EdU+ (cell cycle exit) also indicated that more cells re-entered the cell cycle in the icKO, while more cells exited the cell cycle in the control at E12.5 and E14.5 (Fig. 5D).

Cortical neurons are reduced in number proportionately throughout all layers
Immunostaining with Pax6 and Hes1 antibodies revealed that the thickness of the VZ was reduced in the icKO mice compared to control at E14.5 and thereafter (Fig. 4A-C; Fig. S4A,B). Intriguingly, the number of Tbr2+ cells remarkably increased in the mutant
cortex by E14.5 (Fig. 4A-C), and EdU incorporation experiments revealed that those Tbr2+ cells were highly proliferative at E14.5 (Fig. 5A,B); however, Tbr2+ IPs rapidly reduced in number between E14.5 and E16.5. These results indicated a drastic thinning of the germinal zone composed of NSCs and IPs at later stages. We found that cell death was enhanced in Pax6+, Tbr2+, and Tuj1+ populations in the icKO (Fig. S5A,B), suggesting that the cell death somewhat masked the expansion of NSCs through continued symmetric divisions and caused the reduction of neural stem/progenitor cells at later stages. Enhanced cell death was also observed in the ventral telencephalon (Fig. S5C,D). However, in the cortical regions, no significant difference was observed in the number of GABA (γ-aminobutyric acid)-positive GABAergic interneurons, which are generated in the ventral telencephalon (data not shown).

The thickness of the cortical neuronal layers was reduced in the icKO throughout embryonic stages, as estimated by Tuj1 expression in the IZ/CP and expression of NeuN, a neuronal marker, in the CP (Fig. S4A,B). We counted the number of cells positive for each marker and then normalized to the number of all DAPI+ nuclei. We found that the subsets of layer-specific neurons were proportionately reduced in the icKO at all stages analyzed (Fig. 4C). It is likely that the significant cell death and rapid depletion of neural stem/progenitor cell pool contributed to the reduced thickness of cortical layers.

*In situ* hybridization and real-time RT-PCR revealed that the expression levels of *Neurog2* and *Neurod1* were prominently upregulated around E14.5–16.5 in the mutant cortex (Fig. 6A,B), while expression of *Ascl1* (*achaete-scute complex homolog 1*) was not significantly affected. We further assessed whether neurogenesis terminated earlier and gliogenesis began precociously in the icKO. Although aberrant generation of astrocytes (GFAP+) or oligodendrocytes (Olig2+) was not observed at either E16.5 or E18.5 (data not shown), the number of oligodendrocyte precursor cells (PDGFRα+) decreased in the mutant cortex (Fig. S4C,D), probably due to enhanced cell death and shrinking of the ventral telencephalon. It is possible that the sustained expression of neurogenic bHLH factors prevented precocious gliogenesis in the mutant brain (Sun et al., 2001). Neither did we observe any significant increase of p-Vim+ basal radial glial cells in the SVZ/OSVZ (outer subventricular zone) of mutant cortex at either E16.5 or E18.5 (Fig. S4C,D).
Neural stem/progenitor cells are actively dividing in Hbp1-deficient mice

It was previously reported that Hbp1 is involved in regulation of the Wnt signaling by inhibiting the function of TCF4-β-catenin complex via physical blockade of TCF/LEF-mediated DNA binding, and functions as a growth suppressor by repressing downstream genes of the Wnt signaling and cell cycle-related genes (Tevosian et al., 1997; Gartel et al., 1998; Sampson et al., 2001; Shih et al., 2001; Kim et al., 2006; Elfert et al., 2013; Yan et al., 2014).

We therefore assessed cell proliferation activity in neural stem/progenitor cells in the developing cortex. The number of cycling cells that incorporated EdU for 30 minutes before sacrifice, proliferating cells (Ki67\(^+\)), and mitotic cells (pH3\(^+\)) was significantly higher in the icKO at E14.5 (Fig. 7A,B). We then calculated the ratios of proliferating and mitotic Pax6\(^+\) cells and Tbr2\(^+\) cells separately, and determined that both cell types were more proliferative than in the control (Fig. 7C). The ratios of Ki67\(^+\) cells that coexpressed Pax6 or Tbr2, and the ratios of Pax6\(^+\) or Tbr2\(^+\) cells that coexpressed Ki67 were shown in Fig. S6.

Furthermore, we found that the cell cycle length was much shorter in neural stem/progenitor cells (Ki67\(^+\)) in the mutant cortex at each developmental stage (Fig. 7D; Table S3). Intriguingly, the length of the overall cell cycle (Tc) was gradually elongated as development proceeded in both the control and the mutant cortex. The length of the S phase (Ts) was also shortened in the mutant cortex, although it was less prominent than the reduction in Tc and relatively constant during this period. These results indicate that the duration of the G2-M-G1 phase was significantly shortened in the Hbp1 icKO mice.

We then analyzed the cell cycle length separately in either Pax6\(^+\) NSCs or Tbr2\(^+\) IPs and found that both populations exhibited a shorter cell cycle (Fig. 7E,F; Table S3).

Growth regulatory genes are upregulated in Hbp1 knockdown and Hbp1-deficient cells

We next searched for genes that prospectively lay downstream of Hbp1. Immunohistochemistry and in situ hybridization revealed that cyclin D1 expression was remarkably upregulated in Hbp1 knockdown cells (Fig. 8A,B; Fig. S7A). In addition, we estimated the cyclin D1 expression levels and found that the ratio of cyclin D1-expressing GFP\(^+\) cells was not significantly changed, but the signal intensity of cyclin D1 staining in the GFP\(^+\) area was lower in Hbp1 overexpression and higher in
In line with the above findings, expression levels of cyclin D1 were strikingly upregulated and c-Jun expression was also upregulated in the cortical regions of *Hbp1* icKO mice at E12.5 and E14.5 (Fig. 8C,D). Double labeling with Pax6 or Tbr2 revealed that enhanced cyclin D1 expression was observed in most NSCs and a subset of IPs (data not shown). Real-time RT-PCR confirmed the significant upregulation of *cyclin D1* expression in the icKO at E12.5 and E14.5 (Fig. 8E). The expression levels of *cyclin E1* and *N-myc* were slightly upregulated and sustained until later stages in the icKO (Fig. S7B). Immunohistochemistry using antibodies against phosphorylated Rb (p-Rb) (Ser 780, Ser 807/811), a major target of cyclin D1/Cdk4/6 complexes (Kitagawa et al., 1996; Zarkowska and Mittnacht, 1997; Ely et al., 2005), revealed that the ratios of M phase cells that were strongly positive for p-Rb (Ser 780) were significantly higher in both Pax6⁺ NSCs and Tbr2⁺ IPs in the mutant cortex at E12.5 and E14.5, and the signal intensity of p-Rb (Ser 807/811) was slightly higher throughout the VZ of icKO at E14.5 (Fig. S8A,B). Western blot analysis revealed the increased levels of p-Rb (Ser 807/811) in the mutant cortex at E14.5 (Fig. S8C,D).

**Cyclin D1 is a crucial factor involved in the regulation of cell cycle length by Hbp1**

To address whether the upregulation of cyclin D1 is causative of the phenotype of *Hbp1* deficiency, we performed rescue experiments by cotransfecting *shHbp1* and knockdown vectors against *cyclin D1 (shCcnd1)* by in utero electroporation (Fig. S9A,B). We introduced a mixture of *shHbp1* and *shCcnd1* at ratios of 3:1, 1:1, or 1:3, and found that cotransfection of *shCcnd1* rescued the shortening of cell cycle caused by *Hbp1* knockdown (Fig. 8F). However, the coexpression of *shCcnd1* could not rescue the effects of *Hbp1* knockdown on neuronal differentiation/migration (Fig. S9C,D), suggesting that Hbp1 has critical functions not only in the regulation of cyclin D1 but also in the regulation of neuronal differentiation.

We then performed real-time RT-PCR using total RNAs prepared from NSCs in the embryonic cortex of *pHes1-d2EGFP* transgenic mice and found that *cyclin D1* expression was downregulated by 3-fold between E11.5 and E13.5 (Fig. 8G). Intriguingly, the temporal dynamics of *cyclin D1* expression exhibited a striking contrast to *Hbp1* expression (see Fig. 1C). We further performed a reporter assay using a *cyclin D1* promoter (3.3 kb)-luciferase construct. Expression of Hbp1 by
cotransfection of $pCAG-HA-Hbp1$ vectors in HEK293T cells significantly repressed the cyclin D1 promoter activity (Fig. 8H).

Collectively, these data suggest that upregulation of cyclin D1 caused active proliferation with a short cell cycle length in neural stem/progenitor cells in the developing cortex of $Hbp1$-deficient mice, and that cyclin D1 is a crucial factor involved in the regulation of cell cycle length by Hbp1 during cortical development.

**DISCUSSION**

**Rapid proliferation of neural stem cells delays the onset of neurogenesis**

In the developing embryonic cortex, NSCs gradually alter their characteristics and give birth to distinct cell types in a precise temporal order (McConnell, 1989; Temple, 2001; Ohtsuka et al., 2011). In particular, the timing of the transition from symmetric proliferative to asymmetric neurogenic division is crucial in determining the onset of neurogenesis and the size of the initial stem cell pool. One possible mechanism underlying regulation of the timing of this transition is that the number of cell divisions intrinsically functions as a biological clock to determine their competence and mark developmental steps in NSCs. If so, rapid proliferation of NSCs with a shorter cell cycle will result in a precocious transition from symmetric to asymmetric division and an early-onset of neurogenesis. An alternative possibility is that the gradual elongation of cell cycle length in NSCs during cortical development allows determinants for neuronal differentiation to accumulate to the threshold level that initiates asymmetric neurogenic division and thus acts as a regulator of the biological clock (Calegari and Huttner, 2003; Calegari et al., 2005; Götz and Huttner, 2005; Dehay and Kennedy, 2007). If this is the case, rapid cycling of NSCs inhibits these determinants from reaching the threshold needed to drive neuronal differentiation.

Our data support the latter hypothesis, because rapidly proliferating NSCs in the cortex of $Hbp1$ icKO mice were maintained as cycling NSCs and the generation of Tbr2$^+$ IPs and neurons was delayed (Fig. 4A-C; Fig. 5A,B). Our results suggest that Hbp1 is required to elongate the cell cycle length and facilitate the attainment of the threshold length required to commence neurogenesis, as illustrated in Fig. 7F.

As observed in the mutant cortex at later stages, when NSC proliferation shifted to the asymmetric neurogenic division mode, the aberrantly rapid cell cycle...
accelerated the production of Tbr2+ IPs and their proliferation in the SVZ. This resulted in the premature exhaustion of the neural stem/progenitor cell pool, rendering the neurogenic period shorter in the Hbp1-deficient mice. This raises the possibility that the number of cell divisions in neural stem/progenitor cells is intrinsically limited after the transition to the neurogenic mode.

**Hbp1 inhibits cell cycle progression by suppressing cyclin D1 expression**

Our observations revealed that Hbp1 knockdown and Hbp1-deficiency led to cyclin D1 upregulation and accelerated cell cycle progression in neural stem/progenitor cells in the cortical regions, where reduction in Tc was more substantial than reduction in Ts (Fig. 7C-E; Fig. 8E). Given that cyclin D1 is required for the G1/S transition, and the durations of the G2 and M phases are relatively constant in distinct cell types, it is likely that T_{G1} was mainly reduced via upregulation of cyclin D1. Indeed, the primary phenotypes observed in the Hbp1-deficient mice were consistent with the cyclin D1 overexpression phenotype, in which T_{G1} was shortened, and inhibition of neurogenesis and expansion of IPs were observed (Lange et al., 2009; Pilaz et al., 2009).

It has been reported that Tc in neural progenitors is gradually elongated as development proceeds and it is approximately doubled over the period between E11 and E15, for the most part due to the elongation of T_{G1} (Miyama et al., 1997). Here, we found the contrasting expression dynamics between Hbp1 and cyclin D1 in NSCs over the course of cortical development (Fig. 1C, Fig. 8F), and revealed that Hbp1 repressed the cyclin D1 promoter activity (Fig. 8G). These findings together support our model that Hbp1 is involved in the elongation of T_{G1} in NSCs during early cortical development by regulating the expression of growth regulatory genes including cyclin D1. Future genome-wide expression analysis comparing gene expression in the wild-type and Hbp1-deficient cortex will provide new insights into novel targets or effectors of Hbp1.

**Rb1 cooperates with Hbp1 to promote terminal differentiation**

It was speculated that Hbp1 induces cell cycle exit and promotes neuronal differentiation when overexpressed in neural stem/progenitor cells. However, overexpression of Hbp1 alone failed to promote terminal neuronal differentiation (Fig. 2A). We hypothesized that overexpression of Hbp1 alone would lead to cell cycle exit
without tissue-specific gene expression due to a low Rb1/Hbp1 ratio, and found that Neurod1 expression was downregulated by Hbp1 overexpression (Fig. 2D) and the repression was partly rescued by coexpression of Rb1, while expression of Neurog2 was not significantly affected. It has been shown that Neurod1 regulates terminal neuronal differentiation (Lee et al., 1995; Schwab et al., 2000; Gao et al., 2009), and that Neurod1 expression is activated by the Wnt signaling in adult hippocampal neurogenesis (Kuwabara et al., 2009). Therefore, it is possible that overexpression of Hbp1 repressed Neurod1 expression via inhibition of the Wnt signaling. We found that the expression levels of Neurod1 were significantly upregulated in the cortex of Hbp1-deficient mice (Fig. 6A,B), suggesting that Hbp1 has a role in repressing Neurod1 expression in neural stem/progenitor cells. Although it is likely that the cell cycle length was too short in the Hbp1-deficient mice for Neurog2 and Neurod1 to induce neuronal differentiation at earlier stages, the enhanced expression of these genes might have accelerated the production of Tbr2+ IPs after the transition to the asymmetric neurogenic division mode (Fig. 4A-C). Thus, our present study further corroborates the finding that Hbp1 promotes cell cycle exit, but blocks terminal differentiation if the Rb1/Hbp1 ratio is low. Confirming this hypothesis, an increased ratio of Rb1/Hbp1 promoted terminal neuronal differentiation, similar to observations for muscle cell differentiation (Shih et al., 1998).

**G1 elongation by Hbp1 is essential for cortical development**

We previously reported that expression of the bHLH transcriptional repressor Hes1 is oscillatory with a period of 2–3 hours in neural stem/progenitor cells in the developing cortex, and that expression of neurogenic bHLH genes such as Neurog2 and Ascl1 is also oscillating (Shimojo et al., 2008, 2011; Imayoshi et al., 2013). Thus, it is likely that some neurogenic bHLH factors or their downstream effectors gradually accumulate during the phase of oscillation when Hes1 expression is low. If the G1 phase is elongated, such determinants have a greater chance to reach the threshold necessary to exert their neurogenic functions. Therefore, it is presumed that the duration of the G1 phase is critical in determining the onset of cortical neurogenesis.

Increases in the number of cell divisions during the symmetric proliferative division phase will cause the exponential expansion of NSCs as neuroepithelial cells (a non-neurogenic form of NSCs) and the increase in radial units composed of each stem
cell and its progeny, thus leading to expansion of brain vesicles/ventricles and tangential extension of the VZ (Noctor et al., 2001). Once NSCs shift to the asymmetric neurogenic division mode, they transform into radial glial cells (a neurogenic form of NSCs), and the number of NSCs does not increase any more. Our findings demonstrated that in the absence of Hbp1 neuroepithelial cells continued to rapidly proliferate by symmetric proliferative divisions and expanded the neuroepithelial sheet, leading to the expansion of the ventricular surface and the VZ.

Taken together, our results indicate that the elongation of cell cycle length by Hbp1 mediated by repression of cyclin D1 is a key mechanism regulating the proper timing of neuronal differentiation. As such, Hbp1 is a key factor in determining the onset of cortical neurogenesis and the duration of neurogenic period, and thus being essential for normal cortical development.

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Fig. 1. Temporal and spatial expression patterns of \textit{Hbp1} in the developing brain. (A,B) \textit{In situ} hybridization for \textit{Hbp1} (A) or \textit{Neurog2} (B) was performed on coronal and sagittal (rostral, left; caudal, right) sections of the telencephalon of mouse embryos at various developmental stages. (C) Real-time RT-PCR for \textit{Hbp1} using total RNAs prepared from NSCs in the cortical regions of \textit{pHes1-d2EGFP} transgenic mice. \textit{β-Actin} was used as an internal control, and the values were normalized to that of E11.5 sample. \(n=3\), error bars: s.e.m.
Fig. 2. Inhibition of terminal neuronal differentiation and migration by overexpression of Hbp1. In utero electroporation (ep) was performed with control vectors (pEF-EGFP) alone or with a combination of Hbp1 expression vectors (pEF-Hbp1) at E13.5. The fates of transfected cells were analyzed at E14.5 (B) or E16.5 (A,C) by immunohistochemistry (IHC). (A) Coronal sections of dorsolateral telencephalon were double-stained using anti-GFP (green) and anti-Tuj1/Tbr2/Pax6 (red) antibodies. (B) EdU was administered intraperitoneally to pregnant mice 30
minutes before sacrifice, and incorporated EdU (red) was detected by a fluorogenic click reaction. (C) *pCAGGS-HA-hRb1* vectors were cotransfected with *pEF-Hbp1* at ratios of 3:1 or 1:3 by *in utero* electroporation at E13.5, and the fates of transfected cells were analyzed at E16.5. (D) The proportions of transfected cells that migrated into the cortical plate were shown in bar graphs. (E) *In situ* hybridization for neurogenic bHLH genes such as *Neurog2* and *Neurod1* was performed on coronal brain sections 24 hours after *in utero* electroporation at E14.5. Brackets indicate regions transfected with expression vectors. *n*=3, error bars: s.e.m., *p*<0.05, **p**<0.01; Student’s *t*-test; N.S., not significant. Scale bars: 200 µm.
Fig. 3. Impaired cortical morphogenesis in Hbp1 icKO mice. Tamoxifen (Tam) was administered at E9.5 and embryos were sacrificed at the indicated stages. (A) Photos of whole brains at different embryonic stages comparing the gross shape and size of the brains (dorsal view) of the negative control (NC) and Hbp1 icKO mice. (B) Immunohistochemistry on coronal sections using anti-Tuj1 (green) and anti-Pax6 (red) antibodies. The boxed areas are magnified in the bottom panels. (C) The numbers of Pax6+ NSCs in hemispheres were estimated and compared between the control and the icKO mice at E14.5; the vesicle size of lateral ventricle, the length of brain surface and ventricular surface in cortical regions, the area of ventral telencephalon, and the thickness of each cortical layer (VZ, SVZ, and IZ+CP) were measured and compared between the control and the icKO mice at E14.5. $n=3$, error bars: s.e.m., **$p<0.01$, ***$p<0.001$; Student’s $t$-test; N.S., not significant. Scale bars: 2 mm in A; 1 mm in B.
Fig. 4. Perturbed cortical neurogenesis in Hbp1-deficient mice. (A) Immunohistochemistry on coronal sections of the cortical regions at different
developmental stages with markers specific to each cortical layer (Cux1 for layer II-IV, Ctip2 for layer V, and Tbr1 for layer VI), Tbr2, a marker for IPs, and Pax6, a marker for NSCs. (B) Graphs showing the proportion of cells positive for each marker. The number of cells immunoreactive for each antibody was counted within a radial column of 200-μm width in the middle part of dorsolateral telencephalon. (C) Graphs showing the proportion of cells positive for each marker of all DAPI+ cells within the radial column. Cux1+ cells in the VZ/SVZ/IZ are shown in yellow bars and those in the CP are shown in red. n=3, error bars: s.e.m., *p<0.05, **p<0.01, ***p<0.001; Student’s t-test. Scale bar: 100 μm.
Fig. 5. Prolonged symmetric cell division and delayed neuronal differentiation. (A) EdU was administered intraperitoneally to pregnant mice 12 hours before sacrifice, and the fates of cells that incorporated EdU were analyzed by immunolabeling using anti-Pax6 and Tbr2 antibodies. EdU+ cells are shown in grey and Pax6+ cells are shown...
in red. (B) Graphs showing the ratios of Pax6\(^+\) or Tbr2\(^+\) cells of EdU-incorporated cells. (C) Ratios of pH3\(^+\) cells of Pax6\(^+\) or Tbr2\(^+\) cells at E12.5. (D) Ratios of EdU/Ki67 double-positive (cell cycle re-entry) or EdU single-positive cells (cell cycle exit) of the total number of EdU\(^+\) cells 12 hours after administration of EdU. \(n=3\), error bars: s.e.m., \(*p<0.05\), \(*\ast p<0.001\); Student’s \(t\)-test; N.S., not significant. Scale bars: 200 \(\mu\)m.
Fig. 6. Upregulated expression of neurogenic bHLH genes. (A) *In situ* hybridization (ISH) for neurogenic bHLH genes such as *Neurog2*, *Ascl1*, and *Neurod1* on coronal brain sections. Dotted lines indicate the ventricular surface. (B) Real-time RT-PCR showing the temporal expression patterns of *Neurog2*, *Ascl1*, and *Neurod1*. β-Actin was used as an internal control, and the values were normalized to that of E12.5 NC sample. n=3, error bars: s.e.m., *p*<0.05, **p*<0.01; Student’s *t*-test; N.S., not significant. Scale bar: 100 μm.
Fig. 7. Enhanced cell cycle progression due to *Hbp1* deficiency. (A) Detection of incorporated EdU by a fluorogenic click reaction and immunolabeling using anti-Ki67
and pH3 antibodies on coronal sections of the cortical regions at E14.5. EdU was administered intraperitoneally to pregnant mice 30 minutes before sacrifice. The boxed areas are magnified on the right. (B) Graphs showing the numbers of EdU⁺, Ki67⁺, or pH3⁺ cells within a radial column of 200-μm width in the middle part of dorsolateral telencephalon. (C) Graphs showing the ratios of proliferating (EdU⁺) and mitotic (pH3⁺) cells in either Pax6⁺ or Tbr2⁺ population at E14.5. (D-F) Estimation of the cell cycle length in whole proliferating cells (Ki67⁺), NSCs (Pax6⁺), or IPs (Tbr2⁺) in the cortical regions by the BrdU/EdU double labeling method. (G) A schematic drawing demonstrating the relationship between the gradual elongation of cell cycle length and the timing of the transition from symmetric proliferative to asymmetric neurogenic division based on our observations. n=3, error bars: s.e.m., *p<0.05, **p<0.01, ***p<0.001; Student’s t-test; N.S., not significant. Scale bar: 200 μm.
Fig. 8. Altered expression of cell cycle-related genes under modified Hbp1 expression levels. (A,C) Immunohistochemistry (for cyclin D1 and c-Jun) and in situ hybridization (for cyclin D1) on coronal brain sections from mice subjected to overexpression or knockdown of Hbp1 (A) or in the Hbp1 icKO mice (C). The boxed areas are magnified on the right (A). Dotted line indicates the ventricular surface (C). (B) Graphs showing the ratios of cyclin D1^{+} cells of GFP^{+} cells and the signal intensity of cyclin D1 in the GFP^{+} area at E14.5. (D) Graphs showing the signal intensity of
cyclin D1 and c-Jun in DAPI+ nuclei in the dorsolateral telencephalon. (E) Real-time RT-PCR using total RNAs prepared from the telencephalon to compare the expression levels of cyclin D1 in the control and the icKO mice. β-Actin was used as an internal control, and the values were normalized to that of E12.5 NC sample. (F) pEF-EGFP, pEF-Hbp1, scrambled shRNA, shHbp1, and various combinations of shHbp1 with shCcnd1 were introduced into ventricular cells by in utero electroporation at E13.5, and the cell cycle length was estimated at E14.5 by the BrdU/EdU double labeling method. (G) Real-time RT-PCR using total RNAs prepared from NSCs in the cortical regions of pHes1-d2EGFP transgenic mice showing the temporal dynamics of cyclin D1 expression. β-Actin was used as an internal control, and the values were normalized to that of E11.5 sample. (H) Reporter assay using a cyclin D1 promoter (3.3 kb)-luciferase construct. This reporter construct and Hbp1 expression vectors (pCAG-HA-Hbp1) or control vectors (pCAG-mCherry) were cotransfected in HEK293T cells, and the luciferase activity was measured 48 hours later. n=3, error bars: s.e.m., *p<0.05, **p<0.01, ***p<0.001; Student’s t-test; N.S., not significant. Scale bars: 200 μm in A; 100 μm in C.