Ontogeny of NADPH-diaphorase positive neurons in guinea pig neocortex

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INTRODUCTION

Nitric oxide (NO) plays important roles in the nervous system via modulation of neurotransmission, synaptic plasticity and coupling of vasodilation with neuronal activity (Belvís et al., 1995; Saleem et al., 1996; Guix et al., 2005; Sunico et al., 2005; Melikian et al., 2005; Toda et al., 2009; Hardingham et al., 2013). Normal NO activity is required for a broad range of physiological processes including neurogenesis (Gibbs, 2003; Matarredona et al., 2005), learning and memory (Wultsch et al., 2007; Maielser et al., 2008; Paul and Ekambaram, 2011), sleep (Cudeiro et al., 2000; Gautier-Sauvigné et al., 2005; Gerashchenko et al., 2008; Greene, 2013) and neuroendocrine (Ceccatelli, 1997; Riedel, 2000; Garrel et al., 2002; Givalois et al., 2002). Aberrant NO signaling may be involved in some pathophysiological or disease conditions such as brain aging (McCann et al., 2005), neural stress and inflammation (Cherian et al., 2004; Gadek-Michalska et al., 2013), neuropathological pain (Zimmermann, 2001), neurodegenerative diseases (Togo et al., 2004; Aquilano et al., 2008) and psychiatric disorders (McLeod et al., 2001; Reif et al., 2006; Colvin and Kwan, 2014; Weber et al., 2014).

In mammalian cerebrum there exist two distinct types of interneurons expressing nitric oxide synthase (NOS). Type I neurons are large in size and exhibit heavy nicotinamide adenine dinucleotide phosphate diaphorase (NADPH-d) histochemical reaction, while type II cells are small with light NADPH-d reactivity. The time of origin of these cortical neurons relative to corticogenesis remains largely unclear among mammals. Here we explored this issue in guinea pigs using cell birth-dating and double-labeling methods. Bromodeoxyuridine (BrdU) pulse-chasing (2 doses at 50 mg/kg, 12 h apart) was given to time-pregnant mothers, followed by quantification of NADPH-d/BrdU colocalization in the parietal and temporal neocortex in offspring at postnatal day 0 (P0), P30 and P60. Type I neurons were partially colabeled with BrdU at P0, P30 and P60 following pulse-chasing at embryonic day 21 (E21), E28 and E35, varied from 2–11.3% of total population of these neurons for the three time groups. Type II neurons were partially colabeled for BrdU following pulse-chasing at E21, E28, E35 and E42 at P0 (8.6%–16.5% of total population for individual time groups). At P60, type II neurons were found to co-express BrdU (4.8–11.3% of total population for individual time groups) following pulse-chasing at E21, E28, E35, E42, E49, E56 and E60/61. These results indicate that in guinea pigs type I neurons are generated during early corticogenesis, whereas type II cells are produced over a wide prenatal time window persisting until birth. The data also suggest that type II nitrinergic neurons may undergo a period of development/differentiation, for over 1 month, before being NADPH-d reactive.

Keywords: corticogenesis, GABAergic, interneuron, neuronal development, nitric oxide
while a few studies also demonstrate type II cells in these small rodents (Freire et al., 2012; Magno et al., 2012). Colocalization studies with early cell lineage markers have established that in mice the majority of type I neurons arise from the medial ganglionic eminence (MGE), whereas type II cells have multiple origins including the MGE, lateral/caudal ganglionic eminences (LGE/CGE) and the preoptic area (Jaglin et al., 2012; Magno et al., 2012).

Guinea pig brain develops via a relatively longer period of prenatal morphogenesis and has a greater encephalization quotient in the adult, relative to mice and rats (Rice et al., 1985; Herculano-Houzel, 2007). Also unlike the latter small rodents, but similar to cats and monkeys, type II NADPH-d and doublecortin expressing cells are prominent in adult guinea pigs neocortex (Yan and Garey, 1997a; Cai et al., 2009; Xiong et al., 2008; Bonfanti and Nacher, 2012). Thus, guinea pigs might be suited for addressing certain phylogenetically-related issues about mammalian cortical neurogenesis, such as interneuron formation (Cahalane et al., 2014; Hladnik et al., 2014). Here we characterized the ontogenesis of NADPH-d neurons in guinea pig neocortex using 5-bromodeoxyuridine (BrdU) birth-dating and double-labeling histological methods.

**MATERIALS AND METHODS**

**ANIMALS AND PRENATAL BrdU ADMINISTRATION**

Animal use was in accordance with the National Institute of Health Guide for the Care and Use of Laboratory Animals. All experimental procedures in the present study were approved by the Ethics Committee of Central South University Xiangya School of Medicine for animal care and use. Efforts were also made to minimize stress and pain, and to avoid unnecessary use of experimental animals.

Mating pairs of Hartley guinea pigs aged 4–6 months were purchased from the animal center of Xiangya School of Medicine for animal care and use. Efforts were also made to minimize stress and pain, and to avoid unnecessary use of experimental animals.

Two sets of sections from each brain were processed in parallel by incubation in 0.05 M Tris–HCl buffered saline (pH 8.0, TBS) containing 0.3% Triton X-100, 1 mM nitroblue tetrazolium (NBT, N6639, Sigma-Aldrich) and 5% dimethyl sulfoxide for 45 min at 37°C (Yan et al., 1994). The reaction was stopped by rinsing sections with PBS in room temperature. The sections were then mounted for histological examination or processed further for BrdU immunolabeling using DAB as a chromogen (see below).

**BrdU IMMUNOHISTOCHEMISTRY**

A set of new sections and selected sections stained with NADPH-d histochemistry described above were simultaneously processed for BrdU immunolabeling with the DAB-peroxidase method. Sections were pre-treated in 1 × SSC and 50% formamide for 1 h at 65°C, in 2N HCl for 30 min at 37°C, and in PBS containing 1% H2O2, 5% normal rabbit serum and 0.3% Triton X-100 for 45 min. Sections were next incubated overnight at 4°C with rat anti-BrdU (AbD Serotec, Raleigh, NC, USA, MCA2060, 1:2000) diluted in PBS containing 5% rabbit serum, reacted with biotinylated rabbit anti-rat IgG at 1:400 for 2 h, and subsequently with the ABC reagents (1:400) (Vector Laboratories, Burlingame, CA, USA) for an additional hour. Immunoreactivity was visualized in 0.003% H2O2 and 0.05% diaminobenzidine.

**IMAGING, CELL COUNT AND DATA PROCESSING**

An Olympus microscope (BX53) equipped with a digital imaging system (CellSens Standard, Olympus, Japan) was used for tissue observation and imaging. Sections from the levels of the striatum to mid-hippocampus were used for morphometry of NADPH-d and BrdU/NADPH-d colocalized neurons. Images were taken from 5 equally-spaced sections in each brain at 20×, continuously along the pia for ~3 mm distance (6 image wide) above and below (~1 mm away from the sulcus, where the cortex become flat) the lateral sulcus, respectively, covering both the cortex and white matter regions. The above region in an adjacent Nissl stain section was also imaged at 10× for laminar reference including areal measurement. The images from the same section were montaged, with cortical laminar borders in BrdU/NADPH-d preparation marked by referring to the adjacent Nissl stain images. Cell count was carried out on-screen at high resolution in the montaged BrdU/NADPH-d images, while laminar areas were obtained in the Nissl stain montages using Image I. Layer 1 derives from the embryonic marginal zone (MZ), whereas layers 2–6 arise.
from the cortical plate (CP). Considering this corticogenesis feature as well as the differential laminar distribution of type I and type II nitrigeneric neurons in adult guinea pigs (Yan and Garey, 1997a), cell identification and count were carried out according to four laminar sectors, layer 1, layers 2–4, layers 5/6 and the white matter. Cells were classified and marked, with the numbers of NADPH-d labeled and BrdU/NADPH-d colabeled neurons in each laminar sector recorded. The percentages of double-labeled relative to total NADPH-d cells (type I and type II respectively) were then calculated for each animal. In a given animal, at least 300 type I and 1000 type II NADPH-d positive neurons (with and without BrdU colocalization) were counted. The areal densities of NADPH-d neurons were calculated using the sums of cells and the areas measured for corresponding laminar sectors.

**STATISTICAL TESTING AND FIGURE PREPARATION**

Means of density or percentage values were calculated for each animal using the data from the above systemically sampled sections and laminar regions, and entered into the Prism spreadsheet (Prism GraphPad 4.1, San Diego, CA, USA) for graph preparation. The means and standard derivations (SD) were also calculated for individual groups using this set of data. Kruskal-Wallis Test with Dunn’s Multiple Comparison was used for statistical analysis. The minimal significant level of difference was set at $p < 0.05$. Figures were assembled with Photoshop 7.1.

**RESULTS**

**CORTICAL NADPH-d NEURONS IN NEONATAL AND YOUNG ADULT GUINEA PIGS**

Details of corticogenesis, including the establishment of cortical layers, in guinea pigs still remain largely uncharacterized (to our knowledge). However, a few previous studies have indicated that the basic lamination of the neocortex is readily established at birth (Schütz, 1981; Rice et al., 1985). Here we briefly note that the 6 layered basic architecture of the cerebral neocortex as seen in Nissl stain was established in the newborn (Figure 1A), as compared to young adults (Figure 1I).

Consistent with an early report (Yan and Garey, 1997a), we observed type I and type II NADPH-d neurons in guinea pig cerebral cortex in the present study, with both types readily present in the newborns (Figures 1B–D). The laminar distribution pattern of type I cells were comparable at the three postnatal time points examined, localizing primarily to layers 5 and 6 and the subcortical white matter, while a few of them scattered in the superficial layers, including layer 1 (Figures 1B–G). Also, there was no significant difference in the numerical density of these cells between the three age groups as quantified over layer I, layers 2–4 and layers 5/6 and subcortical white matter (data not shown).

Type II cells were distributed over layers 1–6 and rare in the white matter in the postnatal guinea pigs, denser in layers 1–4 than 5/6 in all postnatal animals. The numerical density of type II cells appeared to be increased at P60 relative to P0 (Figures 1C,F). Using the cell count data from the NADPH-d/BrdU preparations, the mean areal densities of type II cells at P0 ($n = 5$) were estimated to be $136.6 \pm 15.4$, $70.5 \pm 9.8$ and $42.2 \pm 8.1$ per square mm in layers 1, 2–4 and 5/6, respectively, while those at P60 ($n = 5$) were estimated to be $354.7 \pm 45.9$, $156.9 \pm 28.1$ and $85.4 \pm 23.9$ per square mm in layers 1, 2–4 and 5/6, respectively. The medians of the densities were significant different ($P < 0.0001$, Kruskal-Wallis test), with a significant increase of density in layer 1 and layers 2–4 at P60 relative to P0 (Figure 1H).

**LAMINAR DISTRIBUTION OF BrdU/NADPH-d COLABELED NEURONS IN NEONATAL GUINEA PIGS**

Mother guinea pigs give birth following a pregnancy of approximately 9 weeks. In order to determine the lamination pattern of NADPH-d positive neurons during corticogenesis, we gave BrdU injection to pregnant guinea pigs at weekly intervals from E21 to just before birth (E60/61) for the expected offspring, and attempted to quantify the distribution of BrdU/NADPH-d colabeled cells against cortical laminar portions (Figures 1J).

In the newborns received BrdU on E21 (the E21-P0 group), BrdU immunoreactive cells were distributed mostly in the middle portion of the gray matter, i.e., layers 4/5, in the neocortex. A smaller amount of labeling occurred over layer 6 and the white matter, while layers 1–3 contained the least amount of BrdU immunoreactivity (Figures 2A,B). BrdU/NADPH-d colabeled neurons, evidently the type I cells but some type II cells as well, were observed across the cortex and white matter (Figures 2C–E). A similar pattern in the distribution of BrdU single labeling was seen in the neocortex in neonatal guinea pigs received BrdU on E28 (E28-P0) (image not shown). Also, both type I and type II cells were partially colocalized with BrdU in the E28-P0 group (Figures 2F–H). Quantitative data for the E21-P0 group indicated that $17.3 \pm 2.96%$ of the colabeled cells occurred in layer 1, $38.8 \pm 4.0\%$ in layers 2–4, $33.8 \pm 4.5\%$ in layers 5/6, and $10.3 \pm 2.9\%$ in the white matter, with an overall difference among the laminar portions ($P = 0.004$) (Figure 2I). The cell count data from the E28-P0 group showed that $23.3 \pm 1.5\%$ of the colabeled cells occurred in layer 1, $40.0 \pm 2.2\%$ in layers 2–4, $29.0 \pm 2.2\%$ in layers 5/6, and $7.7 \pm 1.3\%$ in the white matter, with a differential laminar localization ($P = 0.016$) (Figure 2J).

In the newborns received BrdU on E35 (E35-P0), BrdU immunoreactive neurons were predominantly localized to layer 2 (Figure 3A). BrdU/NADPH-d colabeled neurons were found in the gray matter but rarely in the white matter (Figures 3B–D), with the great majority being type II cells (to be addressed quantitatively later). Quantitative data for this group revealed that $28.5 \pm 3.0\%$ of the colabeled cells occurred in layer 1, $50.3 \pm 3.6\%$ in layers 2–4, $17.8 \pm 1.5\%$ in layers 5/6, and $3.5 \pm 1.1\%$ in the white matter, exhibiting an overall difference between the laminar sectors ($P = 0.003$) (Figure 3I). In the E42-P0 group, BrdU immunolabeled cells were distributed across the entire cortical gray matter without apparent laminar preference (images not shown, similar to Figure 3E). Some type II cells, but none of type I cells, were found to colocalize with BrdU (images not shown). Cell count data revealed that $31.5 \pm 4.0\%$ of the colabeled cells occurred in layer 1, $51.3 \pm 3.9\%$ in layers 2–4, $12.38 \pm 0.8\%$ in layers 5/6, and $5.0 \pm 0.7\%$ in the white matter, with an differential laminar pattern ($P = 0.003$) (Figure 3J).
In the neonatal guinea pigs subjected to BrdU pulse-chasing on E49 (E49-P0 group) (Figures 3E–H), E56 (E56-P0 group, image not shown) and E60/61(E60/61-P0 group, image not shown), BrdU immunoreactive cells were scattered across the gray matter without apparent laminar preference (Figure 3E). No NADPH-d reactive neurons were found to co-express BrdU in the E49-P0 (Figures 3F–H), E56-P0 or E60/61-P0 groups (images not shown).

LAMINAR DISTRIBUTION OF BrdU/NADPH-d COLABELED NEURONS IN 2 MONTH-OLD GUINEA PIGS

A partial colocalization of BrdU and NADPH-d was observed during microscopic examination of the sections from animals surviving 1 and 2 months following prenatal BrdU pulse-chasing. Overall, the pattern and extent of BrdU colocalization in types I and II NADPH-d neurons in 1 month-old animals (n = 2 for most groups receiving prenatal BrdU administration, except n = 3 for the E49-P30 and E60/60-P30 groups) appeared largely similar to that described above for the P0 groups. Specifically, BrdU localization was found in type I cells in P30 animals received pulse-chasing on E21, E28 and E35, and in type II cells received the chasing on E21, E28, E35 and E42. For simplicity, data from the 1-month old animals are not presented here. In the 2-month-old animals, type I NADPH-d neurons partially co-expressed BrdU in the groups received pulse-chasing by but not later than E35, while a subset of type II NADPH-d neurons exhibited colocalization with BrdU following pulse-chasing at all prenatal time points (Figures 4, 5).
FIGURE 2 | Representative images and quantitative laminar analyses showing bromodeoxyuridine (BrdU) labeling and colocalization in NADPH-d neurons in neonatal guinea pig neocortex following pulse-chasing at embryonic day 21 (E21) and E28. Panel (A) shows the distribution of BrdU immunoreactive nuclei largely in the middle layers of the neocortex in the E21-P0 group. Panel (B) illustrates a low power view of BrdU and NADPH-d dual staining. The remaining panels are high power views of BrdU/NADPH-d colocalized neurons in different cortical layers as marked, from the E21-P0 (C–E) and E28-P0 (F–H) groups. Examples of double labeled type I and type II cells are pointed by arrows and arrowheads, respectively. Dot graphs (I,J) show the laminar distribution of colabeled type I and type II neurons among the groups, with each dot representing the mean of an individual animal. The colabeled cells are present differentially (P = 0.004 by Kruskal-Wallis) between the laminar sectors. Arab numbers: cortical layers, WM: white matter. Scale bar = 100 µm in (A) applying to (B), equivalent to 10 µm for other image panels.

Illustrated as examples, in the E49-P60 group BrdU immunoreactive cells were present across the CP with some cells also in layer 1 and white matter (Figure 4A). BrdU/NADPH-d double-labeled type II, but not type I, cells were found in the cortex predominately in layer 1 and the supragranular layers (Figures 4B,C). Quantitative analysis indicated that 37.7 ± 5.5% of the colabeled cells occurred in layer 1, 54.0 ± 4.4% cells in layers 2–4, 6.7 ± 2.8% in layers 5/6, and 1.7 ± 1.5% in the white matter, with an differential laminar pattern (P = 0.016) (Figure 4G). In the E60-P60 group, BrdU immunoreactive cells occurred in all cortical layers and white matter (Figure 4D), noticeably less abundant relative to the E49-P60 group (Figure 4A). BrdU/NADPH-d colabeled cells, all morphologically featured as type II neurons, were present mostly over layer 1 and supragranular layers (Figures 4E,F). Cell count analysis showed that 35.8 ± 3.6% of the colabeled type II neurons were located in layer 1, 52.0 ± 4.9% in layers 2–4, 9 ± 1.6% in layers 5/6, and 3.3 ± 0.8% in the white matter (P = 0.003) (Figure 4H).

RATE OF BrdU COLABELEING IN NADPH-d NEURONS IN NEONATAL AND 2 MONTH-OLD GUINEA PIGS
To determine the relative proportion and timing of type I and type II NADPH-d cells produced at various stages of prenatal development, we calculated the percentage of BrdU colabeled cells against the total population of the cells measured over the parietotemporal neocortical areas. The calculation was based on identification of BrdU reactivity (DAB reaction) in NADPH-d positive neurons. A colabeled cell was defined if there existed at 3 small granules or a large chunk of brown DAB deposit
clearly present in the nucleus by high resolution. All colabeled and the NADPH-d single labeled cells present across the entire analyzed image covering layer 1 to the white matter were summed. For type I cells, the judgment was made in some cases with a lowering the contrast of the image if the blue NADPH-d reaction was too intense (which could mask the brown DAB product).

For type I NADPH-d neurons, 11.3 ± 1.6% of the total population were colabeled with BrdU in the E21-P0 group. The colocalization rates were 8.9 ± 2.2% and 2 ± 1.4% for the E28-P0 and E35-P0 groups, respectively. No type I NADPH-d neurons were colabeled for BrdU at P0 following BrdU pulse-chasing at E42, E49, E56 and E60/61 (Figure 5A). There was an overall statistically significant difference between the groups (P = 0.001), with Post Test indicating difference for the E21-P0 group relative to E42-P0, E56-P0 and E60/61-P0 groups. In the E21-P60 group, 9.3 ± 1.9% of type I NADPH-d neurons were colabeled with BrdU. In the E28-P60 and E35-P60 groups, 8.9 ± 2.2% and 2 ± 1.4% of type I neurons co-expressed BrdU, respectively. BrdU pulse-chasing at E42, E49, E56 and E60/61 did not yield BrdU colabeling in type I neurons in the 2 month-old offspring (Figure 5B). There was an overall difference between the groups (P = 0.001).

For type II NADPH-d neurons, 8.6 ± 2.0% of the total population showed BrdU colabeling in the E21-P0 group, while 11.5 ± 2.2%, 16.3 ± 3.8% and 13.3 ± 2.5% exhibited BrdU immunoreactivity for the E28-P0, E35-P0 and E42-P0 groups, respectively. In contrast, no type II NADPH-d neurons were colabeled for BrdU at P0 following BrdU pulse-chasing at E49, E56 and E60/61 (Figure 5B) (P = 0.006, with Post Test indicating difference for the E35-P0 relative to E42-P0 and E60/61-P0 groups). Among the 2-month-old groups, 8.6 ± 1.7% of total type II NADPH-d neurons displayed BrdU immunoreactivity in the E21-P60 group, while 11.3 ± 1.9%, 9.5 ± 1.6% and 8.3 ± 1.2% of
FIGURE 4 | Examples of images and laminar quantifications for bromodeoxyuridine (BrdU) labeling and colocalization in type II NADPH-d neurons in 2 month-old guinea pigs following prenatal BrdU pulse-chasing. Shown are the E49-P60 (A–C,G) and E60-P60 (D–F,H) groups. BrdU immunoreactive cells are present across the cortex without apparent laminar preference in both groups, with noticeably less amount cells in the latter group (A,D). High magnification images illustrate BrdU colabeling in a few type II NADPH-d neurons (arrowheads and inserts) (B,C,E,F), with no colocalization in type I neurons. Over 50% and 30% of the colabeled type II cells occur in layers 2–4 and layer I, respectively for both groups (G,H). Scale bar = 100 µm in (A) applying to (D), equivalent to 20 µm for (B,E) and 10 µm for (C,F).

DISCUSSION

Compared to other species including nonhuman primates, little information about prenatal cortical development and neurogenesis in guinea pigs is currently available. Specifically, the time frame for the genesis of neocortical GABAergic interneuron subgroups remains unknown. In the present study we use BrdU birth-dating and double-labeling methods to explore the ontogenesis of type I and type II NADPH-d reactive interneurons in guinea pig neocortex. Following BrdU injections at selected embryonic time points, incorporation of this exogenous cell division marker can be identified in a subgroup of NADPH-d reactive neurons in postnatal cerebral cortex. Analyses of BrdU/NADPH-d colabeled neurons allow us to gain the information about the timing of overall prenatal cell proliferation relative to cell lamination in postnatal neocortex, duration of the
FIGURE 5 | Dot graphs summarizing the percentages of bromodeoxyuridine (BrdU) labeled relative to the total population of type I (A,B) and type II (C,D) NADPH-d neurons among animal groups examined at postnatal day 0 (P0) (A,C) and P60 (B,D) following BrdU pulse-chasing at different embryonic (E) days (E21, E28, E35, E42, E49, E56 and E60/61). Each dot represents the mean from an individual animal. The mean is designated as zero for the animal if no BrdU colocalization in type I or type II NADPH-d neurons can be found by examination of multiple sections. BrdU colocalization in type I NADPH-d neurons can be detected in P0 and P60 animals received BrdU injections at E21, E28 and E35, variably from 2–11.3% group-wise (A,B). BrdU colocalization in type II NADPH-d neurons exists in P0 animal groups received BrdU injections at E21, E28, E35 and E42, variably from 8.6%–16.5% among individual groups. In P60 animals, BrdU colocalization in type II NADPH-d neurons is found in animal groups received BrdU injections at all prenatal time points i.e., E21, E28, E35, E42, E49, E56 and E60/61, ranging from 4.8–11.3% among individual groups. P values represent comparison by Kruskal-Wallis test, with differences (per Dunn’s comparison test) between individual groups marked by lines and asterisks.

production of type I and type II NADPH-d neurons, and an estimate of time delay for type II neurons being differentiated phenotypically following BrdU incorporation.

PRENATAL CELL GENESIS AND LAMINATION IN GUINEA PIG NEOCORTEX

Mammalian corticogenesis involves an initial formation of the germinal ventricular zone (VZ) and the preplate (PP). The PP contains the early appearing Cajal-Retzius cells and some GABAergic pioneer neurons (Marin-Padilla, 1978; Raedler and Raedler, 1978; Luskin and Shatz, 1985a; Molnár et al., 2006). This is followed by the development of the CP, which splits the PP into the MZ, the future layer 1, and the subplate (SP). The CP is differentiated sequentially into layer 6–2 by the arrival of the radially migrating excitatory pyramidal neurons generated in VZ/SVZ, with later-born neurons bypassing and residing superficially to earlier-born neurons. This cortical genesis framework is established by [3H]-thymidine autoradiographic studies in mouse (Angevine and Sidman, 1961), rat (Bayer and Altman, 1990; Ignacio et al., 1995), cat (Luskin and Shatz, 1985b), ferret (Jackson et al., 1989) and nonhuman primates (Rakic, 1974), and is supported by later cell birth-dating and genetic studies, while more complex pattern of cell proliferation and migration appear to occur in phylogenetically high relative to low mammals (for reviews, see Florio and Huttner, 2014).

Using BrdU birth-dating method the present study extends the first preliminary set of data of prenatal neocortical cell production and lamination in guinea pigs. Thus, BrdU pulse-chasing around the third embryonic week reveals labeled cells largely in layers 3–5 in neonatal and young adult neocortex, while cells generated by the 5th embryonic week are localized predominantly to layer 2. These data are consistent with the general inside-out lamination order of corticogenesis in mammals, representing the formation and final settling down of cortical principal neurons (Florio and Huttner, 2014). However, it should be noted that a substantial population of cells appears to be continuously generated in guinea pig neocortex after the cells destined to layer 2 are born, from the 6th embryonic week to at least until birth. These late-born-cells reside across the cortical gray matter without a particular laminar preference. While the precise fates of the late-born cells remain to be characterized, some of them would contribute to the population of type II NADPH-d neurons (to be discussed further).

PRENATAL GENESIS OF TYPE I NADPH-d NEURONS IN GUINEA PIGS

Type I NADPH-d neurons are large in size with long range dendritic and axonal processes capable of modulating cortical neuronal activity in wide areas (Higo et al., 2007; Tamamaki and Tomioka, 2010). In all mammals studied so far these cells are present largely in the subcortical white
matter (Estrada and DeFelipe, 1998; Judas et al., 1999), an area related to the early-formed SP (Luskin and Shatz, 1985a; Kostovic and Rakic, 1990; Aboitiz et al., 2005). In the human cerebrum, type I NADPH-d neurons are readily detectable in the subcortical area of the developing neocortex in the first trimester of gestation (Yan et al., 1996b; Yan and Ribak, 1997b; Judas et al., 1999). Thus, it appears that type I NADPH-d cells may be one of the earliest produced or chemically-differentiated neuronal phenotypes in the mammalian cerebrum.

The present BrdU birth-dating study has verified that type I NADPH-d neurons are a group of early-born neurons in guinea pig neocortex. Specifically, these neurons appear to be generated no later than the 5th embryonic week in guinea pigs. In other words, they are produced essentially before the cortical principal neurons destined to layer II are born. It is also of note that type I NADPH-d neurons in the superficial cortical layers, including layer I, are produced during the same embryonic period as with the bulk of cells in the subcortical white matter. This finding indicates that early-born cortical interneurons do not appear to follow the inside-out lamination pattern typical for cortical principal neurons. Given their early-born feature and their final destinations being the subcortical white matter with a few also in layer 1, some of these cells could be produced probably around the time when the cells in the PP are generated.

PRENATAL GENESIS OF TYPE II NADPH-d NEURONS IN GUINEA PIGS

Previous studies have shown that type II NADPH-d neurons are present in the prenatal human cerebral neocortex with a fairly low density (Yan et al., 1996b; Yan and Garey, 1997a; Judas et al., 1999). In contrast, in adult human or monkey neocortex, type II cortical neurons are considerably abundant especially in the supragranular layers (Hashikawa et al., 1994; Barone and Kennedy, 2000; Garbossa et al., 2005; Cruz-Rizzolo et al., 2006). Therefore, one may speculate that type II neurons could belong to a subpopulation of late-born GABAergic interneurons.

The present BrdU birth-chasing study reveals that type II NADPH-d neurons in guinea pig neocortex could be generated during a surprisingly long period of prenatal corticogenesis. Thus, BrdU colabeled type II neurons in neonatal and 1–2 month old guinea pigs are detected following birth-dating as early as the third embryonic to just before birth. Although it remains debated whether there exists constitutive neurogenesis in the adult mammalian neocortex, neurogenesis was described in the adult guinea pig visual cortex a half century ago with the [3H]-thymidine autoradiographic method (Altman and Das, 1967). The finding of type II NADPH-d neurons being produced until birth raises a question as to whether they could be produced in postnatal life.

In the present study we observe that BrdU/NADPH-d colabeled neurons (including type I and type II, primarily the latter due to the difference in population size) produced during E21–E28 tend to distribute somewhat more evenly across the cortical gray matter relative to those generated later, especially after E35. Thus, with the advance of embryonic age, the colabeled neurons become more concentrated (80–90%) in layers 1–4, while very few of these cells (<5%) reside in the white matter. Notably, layer 1 contains a substantial subpopulation of the colabeled neurons following BrdU birth-dating at all of the examined embryonic time points. Interneurons in the mammalian cerebral cortex are reported to predominantly derive from the subpallial ganglionic eminences, including the MGE, LGE and CGE (Lavdas et al., 1999; Wichterle et al., 1999; Ang et al., 2003). However, studies including those from primates have also suggested pallial origins of cortical interneurons, including layer 1 (Soriano et al., 1992; Zecevic and Rakic, 2001; Hevner et al., 2004; Rymar and Sadikot, 2007; Xiong et al., 2010).

For instance, pallial production of calretinin neurons during the second trimester may substantially contribute to cortical GABAergic interneuron pool in monkeys (Hladnik et al., 2014). Comparing the laminar distribution of BrdU/NADPH-d type II neurons in different groups with increasing survival duration, it appears that these cells may follow an outside-in migratory pattern. As many type II cells in layer I are colabeled following BrdU injections at later embryonic time points (e.g., after E35), cell proliferation leading to interneuron formation clearly occurs in guinea pigs after the basic lamination of the neocortex is established.

POSTMITOTIC DIFFERENTIATION OF TYPE II NEURONS IN GUINEA PIGS

Neurogenesis consists of a continuing process whereby postmitotic cells develop into immature and then mature neurons, with specific markers being expressed at different stages (von Bohlen Und Halbach, 2007). BrdU birth-dating may allow an estimate of the time period during which the labeled postmitotic cells develop and mature into chemically differentiated neurons (Taupin, 2007). Such developmental time period for either the general population or specific subgroups of cortical GABAergic interneurons remains largely unclear to date in most mammals. As discussed above, our results suggest that type I NADPH-d neurons are produced fairly early relative to corticogenesis in guinea pigs. However, as BrdU pulse-chased brains are not examined prenatally in the present study, it is not feasible to estimate how long it would take for type I neurons to become NADPH-d reactive following the cell division that incorporates the injected BrdU.

The data obtained in the present study however allow a rough prediction that type II neurons would likely take more than 1 month following BrdU incorporation to become NADPH-d reactive. This estimate is based on whether and when BrdU/NADPH-d colabeled type II neurons are detected in postnatal neocortex in the groups received pulse-chasing at different embryonic time points. We find that no BrdU/NADPH-d colabeled type II neurons are detectable in neonatal and 1 month-old guinea pigs with BrdU injected on and after E49. In contrast, BrdU/NADPH-d colabeled type II neurons are present in 2 month-old animals following BrdU injections at E49, E56 and E60/61. Thus, the finding of BrdU/NADPH-d colocalization in type II neurons in the E49-P60 group, but not in the E49-P0 group, would implicate that type II neurons become NADPH-d positive between approximately 40 and 70 days following mitosis (roughly counting guinea pig pregnant time as ~60 days here).
By the same token, the lack of BrdU/NADPH-d colocalization in type II neurons in neonatal and 1 month-old, but the presence of which in 2 month-old, animals following BrdU injections on E60/61 would suggest that there is a delay between 30 and 60 days for postmitotic precursors to be detected by NADPH-d histochemistry.

In summary, the present study demonstrates that type I nitrinergic interneurons in guinea pig cerebral neocortex are generated during embryonic day 21–35 (potentially earlier). Type II NADPH-d neurons may be generated over a wide prenatal time window, from embryonic day 21 until birth. Overall, these results may imply that the formation of type I NADPH-d neurons could be affected by early embryonic insults, while certain late prenatal and postnatal factors could impact the development of type II nitrinergic neuronal population in mammalian cerebral cortex.

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