In Vitro Differentiation of Mouse Embryonic Stem Cells into Neurons of the Dorsal Forebrain

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Abstract

Pluripotent embryonic stem cells (ESCs) are able to differentiate into all cell types in the organism including cortical neurons. To follow the dynamic generation of progenitors of the dorsal forebrain in vitro, we generated ESCs from D6-GFP mice in which GFP marks neocortical progenitors and neurons after embryonic day (E) 10.5. We used several cell culture protocols for differentiation of ESCs into progenitors and neurons of the dorsal forebrain. In cell culture, GFP-positive cells were induced under differentiation conditions in quickly formed embryoid bodies (qEBs) after 10–12 day incubation. Activation of Wnt signaling during ESC differentiation further stimulated generation of D6-GFP-positive cortical cells. In contrast, differentiation protocols using normal embryoid bodies (nEBs) yielded only a few D6-GFP-positive cells. Gene expression analysis revealed that multiple components of the canonical Wnt signaling pathway were expressed during the development of embryoid bodies. As shown by immunohistochemistry and quantitative qRT-PCR, D6-GFP-positive cells from qEBs expressed genes that are characteristic for the dorsal forebrain such as Pax6, Dach1, Tbr1, Tbr2, or Sox5. qEBs culture allowed the formation of a D6-GFP positive pseudo-polarized neuroepithelium with the characteristic presence of N-cadherin at the apical pole resembling the structure of the developing neocortex.

Keywords

Mouse embryonic stem cell · Embryonic forebrain · Stem cell differentiation · Wnt signaling

Introduction

The formation of the neocortex (corticogenesis) starts at embryonic day (E) 10 from undifferentiated neural progenitors of the dorsal telencephalon. Proliferating progenitors reside in the ventricular and subventricular zone (VZ and SVZ) while newly generated neurons migrate from the germinal VZ towards the outer margin. At birth, the cortical plate consists of six layers that were generated by inside-first and outside-last succession, i.e., early-born neurons settle in the deepest layer while late-born neurons migrate along radial glial cells towards the superficial layers (Rallu et al. 2002; Wilson and Houart 2004; Fishell and Kriegstein 2005; Gotz and Huttner 2005; Rash and Grove 2006). Genetic mechanisms that control the
regionalization of the telencephalon have been partly elucidated showing the involvement of the regulatory pathways Wnt, Bmp, Fgf and Shh (Shimogori et al. 2004).

Wnt signaling is important for both cell specification and cell proliferation during development of neural tissue. At early developmental stages, the expression gradient of Wnts in the neural tube increasing from anterior to posterior is capable of inducing a caudal character, i.e., the midbrain and hindbrain (Wilson and Edlund 2001; Nordstrom et al. 2002). In accordance, inhibition of Wnt signaling at early developmental stages is required for the specification of an anterior character in the forebrain (Houart et al. 2002; Lagutin et al. 2003). At later developmental stages, experiments with chicken tissue explants and conditional β-catenin knock-out mice showed that canonical Wnt signaling determines dorsal telencephalic character at the expense of the ventral telencephalon (Gunhaga et al. 2003; Backman et al. 2005). Furthermore, Wnt-3a controls cell proliferation in the hippocampal primordium (Lee et al. 2000b). At the level of transcription control, two nuclear factors, Pax6 and Emx2, that are expressed in opposing gradients along the anterior-posterior axis play a crucial role in the specification of the cortex (Muzio et al. 2002).

The developmental transition of undifferentiated cells in the early embryo into committed neurons has been partially mimicked by in vitro differentiation of embryonic stem cells (ESCs). Lee et al. (2000a) cultured ESCs to form embryoid bodies that contained multi-potent cells present in three germ layers. From this poorly defined pool of tissue-specific cells, neural stem cells (NSCs) can be expanded in the presence of EGF and FGF2 and their subsequent exposure to Shh with Fgf8b triggers differentiation into neurons of midbrain/hindbrain character including dopaminergic neurons. Differentiation of ESCs into neural precursors in adherent culture yielded a remarkably high number of neural precursors labeled with Sox1-GFP without the requirement of exogenous factors (Ying et al. 2003) suggesting that neural fate may be a default differentiation program for pluripotent stem cells (Munoz-Sanjuan and Brivanlou 2002; Levine and Brivanlou 2007). Naïve neural cells which arise from differentiating ESCs can respond to inductive signals that can further influence their fate decisions. For instance, inhibition of canonical Wnt signaling with the antagonist Dkk1 promoted telencephalic character by suppressing caudal fate in neural cells (Watanabe et al. 2005). In addition, the same authors showed that subsequent treatment with recombinant Wnt3a increased the proportion of cortex-specific neurons while Shh induced the expression of markers typical of the basal (ventral) telencephalon. In agreement with these data, inhibition of the Shh pathway with cyclopamine remarkably increased the expression of dorsal markers such as Pax6 or Vglut in cultured neuronal cells that were generated by controlled differentiation of ESCs (Gaspard et al. 2008, 2009). The role of Wnts in ESC differentiation towards neurons appears more complex because Wnt1 and Lrp6 are dispensable for generation of dopaminergic neurons and their absence rather elevates their numbers (Cajanek et al. 2009). Interestingly, in vitro differentiated neurons do not only show cell fate specific markers but also resemble the cellular structure of mature organs. Eiraku et al. (2008) showed that differentiating neural cells self-arranged in a manner that resembled the developing cortex with an apical neuroepithelium and a basal margin when they were cultured in tight contact within large cell aggregates.

We previously created transgenic reporter mice D6-GFP where GFP was specifically expressed in cortical progenitor cells from E10.5 onwards and in mature neuron descendant from these cells in the adult hippocampus and cortex (Machon et al. 2002). Thus D6-GFP serves as a highly specific marker for neural progenitors and neurons that originate in the dorsal telencephalon and that are predominantly excitatory glutamatergic. In this study, we used ES cells derived from transgenic mice containing the D6-GFP marker to directly observe differentiation of ESCs into cortical neurons in tissue culture. GFP + cortical neuronal progenitors were efficiently generated in vitro as clusters within qEBs. In contrast normal embryoid bodies (nEBs) yielded GFP + cortical neuronal progenitors only with an extremely low efficiency. Transcription factors that are specific for the dorsal forebrain were activated in D6-GFP + cells, further confirming correct fate determination. After cell sorting, GFP + cells derived from qEBs formed neurospheres in cell culture that were indistinguishable from normal neural stem cells derived from the developing telencephalon. Finally, additional stimulation of the canonical Wnt signaling pathway during the later period of ESC differentiation increased the percentage of D6-GFP + cells but was not essential for their appearance. These results suggest that the in vitro generation of cells with a specific dorsal forebrain fate can be directed by signals that are present within qEBs while increased stimulation of canonical Wnt signaling will further enhance dorsal fate specification in qEBs.

Materials and Methods

ESC Culture

ESCs were derived from the inner cell mass of the blastocyst of transgenic mice carrying a D6-driven GFP reporter (Machon et al. 2002). ESCs were expanded on feeder cells (primary mouse embryonic fibroblasts) and were subsequently adapted to feeder-free culture conditions on 0.1% gelatin-coated plastic. The ESC medium consisted of...
knock-out Dulbecco’s Modified Eagle Medium (DMEM, Invitrogen) supplemented with 10% ES-grade fetal calf serum (FCS, Speciality Media), ESGRO (1000 U/ml) (Millipore), non-essential amino acids (NEAA, Invitrogen), 0.1 mM 2-mercaptoethanol (2-ME, Sigma), 2 mM l-glutamine (Invitrogen), and 100 unit/ml penicillin/streptomycin (Invitrogen). Cells were plated at the density of $0.5 \times 10^5$ per ml and were passaged before reaching confluence.

Differentiation of ESCs

All cell culture protocols are schematically illustrated in Fig. 4a.

For nEBs culture, adherent ESCs were dissociated in trypsin–EDTA and plated in uncoated 100-mm plastic dishes at a density of 500,000 ESCs per 10 ml EB media. This day was set as the differentiation day 0 (d0). For qEBs culture, dissociated ESCs were grown in 96-well plates with a round bottom at the density of 2000 cells per well in the EB medium (qEBs culture 1, see Fig. 4a) or in the Neuro-medium from d0 (qEBs culture 2). qEBs were cultured in round-bottom wells for the whole culture period without adhering to the bottom and the medium was changed every third day. The EB medium was composed of GMEM (Invitrogen) with 5% knock-out serum replacement (KSR, Invitrogen); NEAA (Invitrogen), 0.1 mM 2-mercaptoethanol (2-ME, Sigma) and 2 mM l-glutamine (Invitrogen). Neuro-medium consisted of the Neurobasal-A medium (Invitrogen) supplemented with B27 (Invitrogen), l-glutamine, penicillin/streptomycin and growth factors bFGF 10 ng/ml, EGF 20 ng/ml (R&D Systems). For additional treatment, Wnt3a was applied to the EB medium at 10 ng/ml (qEBs culture 3), Dkk1 50 ng/ml (qEBs culture 4) between d6 and d12. 0.5 μM retinoid acid (RA) was applied from d5 to d9 in the EB medium (qEBs culture 5). In growth assays, diameters of qEBs were measured from d1 to d20. The images of growing clusters were taken every day and their diameter was set as a function of the pixel size using Zeiss Axiovision software.

Immunofluorescence

To prepare samples for staining, ESCs were plated onto gelatin-coated round-shape coverslips and incubated for 2 days in the ESC medium. The whole nEBs were placed on coverslips coated with retinectin (AH Diagnostics) in the EB medium. NSCs from neurosphere cultures of FACS-sorted qEBs were attached on poly- l-lysine (Sigma)-coated coverslips and incubated for 24 h in Neuro-medium without EGF. For cryosections, qEBs were collected and fixed in suspension in 4% paraformaldehyde in PBS and dehydrated overnight in 30% sucrose in PBS. Subsequently, qEBs were embedded in OCT (Sakura) and 8 μm cryosections were prepared using a cryostat (Leica).

For immunostaining, cells were fixed in 4% paraformaldehyde for 30 min. Specimens were permeabilized in 0.1% TritonX-100 in PBS for 15 min. After saturation in 5% goat serum, 5% bovine serum albumin (BSA, Invitrogen), 0.1% TritonX-100 in PBS for 30 min, cells were incubated overnight at 4°C with a primary antibody diluted in PBS containing 0.5% goat serum, 0.5% BSA, and 0.1% TritonX-100. After washing with PBS, incubation with a secondary antibody followed for 1 h at room temperature in the same buffer. 1 μg/ml 4,6-diamidino-2-phenylindol (DAPI, Roche) in PBS was used to visualize cell nuclei.

Primary antibodies: rabbit anti-Pax6 (1:200) (Covance), rabbit anti-Tbr1 (1:200) (Chemicon), rabbit anti-Tbr2 (1:200) (Chemicon), mouse anti-N-cadherin (1:400) (BD Biosciences), rabbit anti-Oct4 (1:100) (Santa Cruz, rabbit anti-Sox1 (1:200) (Chemicon), rabbit anti-Sox2 (1:200) (Cell Signaling), mouse anti-Nestin (1:400) (Millipore), mouse anti-Tuj1 (1:400) (R&D), mouse anti-GFAP (1:400) (Sigma). Secondary antibodies: anti-mouse or anti-rabbit IgG conjugated with Alexa 488/594 (1:400) (Invitrogen). Stained sections were examined using a fluorescent microscope Axioskop2 (Zeiss) and images were taken with Axiovision software (Zeiss).

Quantitative qRT-PCR

RNA was extracted using a total RNA isolation kit (Sigma). cDNA was synthesized from 1 μg RNA using Oligo primer p(NT)$_6$ (Roche) and reverse transcriptase kit (Stratagene) according to manufacturer’s instructions. Real-time polymerase chain reaction (qRT-PCR) was performed using the SYBR green PCR master mix kit (Stratagene). Amplification: initial denaturation for 5 min at 95°C, then 40 cycles: denaturation 95°C 30 s, annealing 55°C 40 s, elongation 72°C 30 s. Primer sequences for genes are listed in Supplementary Table 1. Statistical analysis: The values show relative values of expression that was normalized to GAPDH as the average of three independent with standard deviations.

Flow Cytometry Analysis and FACS Sorting

For SSEA-1 labeling, dissociated ESCs were suspended in PBS containing 0.2% BSA with mouse anti-SSEA-1 antibody (1:200, Millipore) and incubated on ice for 30 min. Anti-mouse IgM-Alexa594 in the same buffer was applied for another 30 min after washing. Cells were analyzed by flow cytometry after washing in PBS.

qEBs (with or without D6-GFP$^+$ cell clusters) were dissociated with trypsin/EDTA at 37°C, and trypsinization was stopped using trypsin neutralizer solution (Gibco). Single cells thus obtained were washed and re-suspended in...
PBS. Cells were immediately measured using a FACS Aria flow cytometer (BD Biosciences). Both GFP− and GFP+ cells were collected and data were analyzed with FACS-Diva software.

Neurosphere Culture and Differentiation

GFP− and GFP+ cells were isolated by FACS, resuspended in Neuro-medium containing bFGF and EGF, and plated at a density of 40,000 cells per ml media. Neurospheres formed within 1–2 weeks after FACS and were passaged further using trypsinization. For differentiation, dissociated neurosphere cells were plated onto poly-ℓ-lysine-coated coverslips in Neuro-medium without EGF (seeding density 40,000 cells per cm²). After 24 h, differentiated cells were fixed in 4% paraformaldehyde and stained using immunofluorescence as described above.

Results

Neural Differentiation of ESCs in nEBs

To study differentiation of ESCs into cortex-specific neural cells, we generated ESCs from blastocysts derived from transgenic D6-GFP mice containing a GFP reporter that is driven by the forebrain-specific enhancer of the Dach1 gene D6 (Machon et al. 2002). During embryogenesis, D6 enhancer activity was detectable from embryonic day (E) 10.5 and expanded throughout the developing neocortex and the hippocampus by E12.5 and E15 (Fig. 1a–b). Immunohistochemistry with antibodies against several cortex-specific transcription factors was performed on tissue sections from E15 D6-GFP transgenic mice. An anti-Pax6 antibody was used to visualize neural progenitors along the intermediate zone (IZ) (Englund et al. 2005). Figure 1c and d shows an overlapping expression of Pax6 or Tbr2 with D6-GFP. Pax6 in the VZ and Tbr2 in the SVZ were co-labeled with the D6-GFP marker, which is, apart from the VZ and SVZ, also expressed in cortical postmitotic neurons (the outer layer of the cortex). D6-GFP therefore marks both neocortical VZ and SVZ progenitors and mature neurons in the cortical layers at critical stages of neurogenesis. This implies that ESCs derived from D6-GFP transgenic mice provide a simple and explicit model for in vitro cell differentiation into embryonic cortical neurons.

D6-GFP ESCs were maintained on a gelatin-coated surface in the ESC medium containing ESGRO. These ESCs showed the typical morphology and growth properties of pluripotent cells (Supplementary Fig. 1a). Immunofluorescence with antibodies recognizing the pluripotency markers Oct4 and Sox2 documented that D6-GFP ESCs strongly expressed these transcription factors (Supplementary Fig. 1c−d). Further, FACS quantification of the SSEA-1 surface marker, which is specific for mouse ESCs, showed a high percentage of SSEA-1 positive cells in the D6-GFP derived ESCs culture (Supplementary Fig. 1b). In contrast, the early neuroectodermal marker Sox1 and the neural progenitor marker Nestin were not detected (Supplementary Fig. 1e−f). In conclusion, ESCs derived from D6-GFP blastocysts displayed standard characteristics of pluripotency.

To test whether D6-GFP-derived ESCs could be differentiated into neurons of the cortex we tested several cell culture strategies. At first, we used a protocol based on generation of nEBs. At day 0 (d0), ESCs were dissociated with trypsin and seeded in uncoated plastic dishes in the EB medium (nEBs culture, see Fig. 4a). The absence of ESGRO, replacement of FCS with KSR, and the growth in suspension triggered cell differentiation. Floating cells formed middle-size aggregate nEBs (Fig. 3a–b). To examine whether a differentiation towards a neural lineage occurred, fluorescent immunohistochemistry of nEBs was performed after 12 days of incubation. As shown in Fig. 2a, many cells in D6-GFP ESCs derived nEBs were positive for the markers for neuronal precursors Nestin, the marker for differentiated neurons Tuj1 and for the astrocyte-specific protein GFAP, demonstrating a neural identity of many cells within nEBs.

To track the differentiation process in nEBs, qRT-PCR on mRNA obtained from days 6 and 12 of differentiation was carried out and the expression profile was compared to the parental ESCs. As a control, mRNA from E13 mouse cortex derived neurospheres was used. Our results showed that cells grown under conditions to form nEBs lost their pluripotency character between d6 and d12 as reflected by the reduced expression of Oct4, Nanog, and Klf4 (Fig. 2b). Sox2, which was highly expressed in ESCs, decreased during the first days of differentiation but its expression almost recovered at d12. As Sox2 expression is abundant both in ESCs and in neural progenitors (Zappone et al. 2000), the temporary decrease in the expression of Sox2 may reflect a spontaneous differentiation into multiple cell types at earlier stages of differentiation while the neural fate starts to dominate later. Further, the decrease of pluripotency was accompanied with an increase of markers for neuroectoderm, Nestin and Sox1 (Fig. 2b), which is in line with other reports (Pevny and Nicolis 2010). We also measured remarkably increased expression the forebrain-specific gene FoxG1 (Ahlgren et al. 2003; Hanashima et al. 2002; Wataya et al. 2008); and genes expressed in the developing the cortex such as Pax6 (Bishop et al. 2000; Englund et al. 2005) and Sox5 (Kwan et al. 2008; Lai et al. 2008; Hanashima et al. 2002) (Fig. 2b).
In addition, nEBs mRNA analysis showed an upregulation of several other neural genes such as the ventral forebrain marker \(Nkx2.1\) (Sussel et al. 1999), the hindbrain marker \(Hoxb9\) (Irioka et al. 2005), and the astrocyte-specific marker GFAP indicated a presence non-cortical cell types (Hanashima et al. 2002). The elevation of Brachyury (\(Bra\)) and Vimentin suggested a presence of non-neural cell identity. Altogether, qRT-PCR data reflected an expected cell heterogeneity in nEBs that includes dorsal, ventral, anterior, and posterior cell types of the neural tissue and other cell types. All the genes related to neural character that were tested in nEBs approached levels of expression close to the levels obtained from NSCs that were directly derived from E13 mouse forebrains.

Along with induction of cell-type specific genes in nEBs, genetic components of the Wnt pathway were activated and showed dynamic changes in their expression (Fig. 2c). For instance, \(Wnt1\), \(Dkk1\), and \(sFrp2\) were induced upon ESC differentiation whereas \(Wnt3a\), which was expressed in pluripotent ESCs, was downregulated in nEBs. Induction of components of the canonical Wnt signaling during differentiation led to activation of \(Axin2\) (Fig. 2c), a well-known Wnt target gene (Fujimura et al. 2007; Jho et al. 2002), indicating activation of the canonical Wnt pathway in spontaneously differentiating cells.

During more than a 12-day culture period, cells in nEBs were monitored for the expression of GFP that would indicate differentiation of ESCs into neural cells of the
neocortical fate. Although the Dach1 mRNA was induced in 12d nEBs (Fig. 2b), live D6-GFP signal was barely visible. Only a few GFP-positive cells were found in selected nEBs, and immunostaining with an anti-GFP antibody was required to visualize those cells (Fig. 2a, right). In conclusion, an ESC differentiation protocol using nEBs was prone to generate neural tissue but the telencephalic/neocortical character as marked by the D6-GFP reporter was established with low efficiency.

Forebrain-Specific D6-GFP+ Cells were Generated in qEBs

To find culture conditions that would yield neural cells of the dorsal forebrain with high efficiency, we adapted a culture protocol from Eiraku et al. (2008). In short, dissociated ESCs were allocated in round-bottom 96-well plates where they aggregated in large clusters within 1 day. These aggregates are called qEBs (Fig. 3c–d). Within qEBs as well as in nEBs, cell differentiation is thought to be a stochastic process that is influenced by a close cell–cell contact.

In contrast to nEBs in which few D6-GFP+ cells were found during the culture period, qEBs culture yielded frequently fluorescent cells (Fig. 3e–h). Commonly, one or two clusters of D6-GFP+ cells were present within one qEB. Images of several D6-GFP+ qEBs are shown in Supplementary Fig. 2 with several clusters of fluorescent cells within a qEB.

Two different culture medias were tested for the frequency of D6-GFP+ induction in qEBs: (i) EB medium containing knock-out serum replacement (KSR) and (ii) Neuro-medium which was serum-free but contained EGF and bFGF. The size of qEB was significantly larger in Neuro-medium (up to 700 μm in diameter) at day 18 whereas those in EB media were smaller (around 400 μm in diameter) (Fig. 4b). Clusters from 8 parallel wells were measured every day during a 20-day incubation period.

Generally, the first D6-GFP+ cells in qEBs emerged around d10 to d12, in rare cases as early as at d7. After
18 days, no further clusters with D6-GFP<sup>+</sup> cells emerged although the GFP signal persisted for at least another week. The incidence of D6-GFP-positive clusters (regarded those containing at least one fluorescent cell) was higher in Neuro-medium. On average 75% qEBs contained D6-GFP-positive clusters.

**Fig. 3** Differentiation of ESCs into neocortical progenitor cells in quick embryoid bodies (qEBs). EB size comparison during ESC differentiation: a Normal EBs after 2 days (nEBs 2d). b Normal EBs after 12 days (nEBs 12d). c Quick EBs after 2 days (qEBs 2d). d Quick EBs after 12 days (qEBs 12d). e-h qEBs after 12 days with corresponding D6-GFP fluorescence in right panels (f, h). Scale bar: 100 μm

**Fig. 4** Quantification of the generation of D6-GFP neuronal progenitors in different media. a A schematic illustration of the cell culture protocols that were used in this work. b A growth curve of qEB during 20 days (d1–d20). qEBs were incubated in EB medium or in Neuro-medium, and the size of qEBs was measured daily for 20 days (diameter in μm). On d18, clusters stopped growing. The average diameter of qEBs in Neuro-medium was 800 μm while only half the size was reached in the EB medium. c Incidence of GFP<sup>+</sup> qEBs shows higher percentage of fluorescent clusters in Neuro-medium (%) compared to EB medium (40%). d Flow cytometry analysis showing a relatively lower percentage of D6-GFP<sup>+</sup> cells in qEBs grown in Neuro-medium (19.6%) compared to qEBs grown in EB medium (35%). n number of independent experiments.
cells in 2 independent experiments \((n = 2)\). On the other hand in EB medium, on average 40% qEBs contained positive cells in six experiments \((n = 6)\) (Fig. 4c). Nevertheless, the relative number of GFP\(^+\) cells in pooled qEBs was lower in Neuro-medium and fluorescence was relatively weaker. The peak of the GFP intensity in the FACS histograms was around \(10^4\) in Neuro-medium and \(5 \times 10^3\) in EB medium (Fig. 4d). This indicates that EB medium is more suitable for generation of D6-GFP\(^+\) cells in qEBs while Neuro-medium, containing the growth factors EGF and bFGF, may support growth of noncortical neuronal types at the expense of the cortex-specific ones.

**ESC-Derived D6-GFP\(^+\) Cells Grown in qEBs Expressed Typical Markers of Neural Progenitors and Neurons of the Dorsal Telencelphalon**

To further characterize qEBs-derived D6-GFP\(^+\) cells, the presence of markers that are specifically activated in the developing neocortex was examined by immunofluorescence (Englund et al. 2005). Cryosections of qEBs containing clusters of D6-GFP\(^+\) cells were analyzed with antibodies against Sox2, Pax6, Tbr1, and Tbr2. As shown in Fig. 5a–d, a substantial number of D6-GFP\(^+\) cells expressed the markers Sox2, Pax6, Tbr2, and Tbr1, further confirming the telencephalic/neocortical character of the D6-GFP\(^+\) cells.

Next, we isolated mRNA from GFP\(^+\) and GFP\(^-\) cells and real-time qRT-PCR analysis of other genes that are specifically expressed in the telencephalon was performed. Some qEB contained up to one third of fluorescent cells as quantified by FACS analysis (Fig. 6a). However, viable cells were not easily collected by FACS and mRNA was isolated instead from manually separated fluorescent cells under the UV-light microscope. Results in Fig. 6b confirmed that FoxG1, Dach1, Pax6, and Sox5 showed increased expression in D6-GFP\(^+\) cells.

The cortical neuroepithelium in the mouse embryo is polarized, i.e., cells at the apical pole of the ventricular zone express N-cadherin that is concentrated in adherens junctions (Fig. 1e). N-cadherin is essential for the proper function of the adherens junctions since loss-of-function mutant mice exhibit a disrupted neuroepithelial polar structure (Kadowaki et al. 2007). Labeling of qEBs with N-cadherin revealed that some clusters self-assembled in rosette-like structures. In these structures, D6-GFP\(^+\) cells surrounded a cavity with a central circle of apical N-cadherin which was similar to the organization of the neuroepithelium with adherens junctions (Fig. 5e). Clusters of Pax6\(^+\) cells around a central N-cadherin-positive rosette showed an arrangement similar to the developing neocortical wall (Fig. 5f) indicating that differentiating cells within qEBs may be able to establish a cellular architecture reminiscent of the developing neocortex.

**Wnt Signaling Pathway Enhances the Efficiency of Corticogenesis In Vitro**

Wnt signaling plays an important role in embryonic patterning of the telencephalon, since it promotes dorsal telencephalic identity while suppressing ventral fate (Backman et al. 2005; Guillemot 2007). We noticed that during the differentiation of ESCs in nEBs many genes of the canonical Wnt signaling pathway were induced (Fig. 2c). To test a direct impact of canonical Wnt signaling on cell differentiation, qEBs were treated with soluble Wnt3a and the percentage of D6-GFP\(^+\) cells in qEBs was quantified by FACS. Representative histograms are depicted in Fig. 7. Wnt3a treatment \((10 \text{ng/ml})\) during d6–d12 increased the number of fluorescent cells from \((35.8 \pm 2.9)\%\) (untreated samples) to \((57.5 \pm 3.6)\%\) (treated samples). In contrast, treatment with the antagonist Dkk1 \((50 \text{ng/ml})\) decreased the number of positive cells to \((18.1 \pm 4.2)\%.\) Finally, qEBs were treated with 0.5 \(\mu\text{M}\) retinoic acid (RA) between d5 and d9. Although RA has been shown to trigger neural induction (Bain et al. 1995; Kim et al. 2009) it completely hindered the generation of D6-GFP\(^+\) cells in our assays. This is in line with previous reports showing that RA imposes a caudal character of neural cells at the expense of anterior cell fates including the dorsal forebrain (Irioka et al. 2005).

**ESC-Derived D6-GFP\(^+\) Cells Formed Neurospheres with Properties of NSCs**

In our previous work (Machon et al. 2002, 2005), we showed that D6-GFP\(^+\) cells from E13 and neonatal forebrains could be propagated and differentiated into neurons and astrocytes. We therefore tested these properties in D6-GFP\(^+\) NSCs cells derived from qEBs. After FACS isolation of d15 qEBs, GFP\(^+\) cells formed neurospheres of various sizes when grown in standard Neurobasal A-based medium supplemented with EGF and bFGF (Fig. 8a). During expansion, the inner core of the neurospheres remained positive for GFP while the surface cells gradually lost fluorescence (rectangles in Fig. 8a, a’). NSCs in neurospheres lost fluorescence after repeated passages and thus behaved similar to NSCs isolated from the developing neocortex under the caudalizing effect of bFGF or EGF (Prajero et al. 2010; Machon et al. 2002, 2005).

Neural stem cells meet two major criteria: the capability to proliferate and the ability to differentiate into various cell types of the nervous system. We therefore tested the differentiation potential of 2nd passage neurospheres derived from FACS-sorted D6-GFP\(^+\) cells. Dissociated neurosphere cells were plated onto polylysine-coated coverslips and grown in Neuro-medium without EGF for 24 h. As shown in Fig. 8b, most of the cells were stained positive.
Fig. 5 Characterization of D6-GFP\textsuperscript{*} cells in cryosections of qEBs by immunofluorescence.

\textbf{a–d} Immunofluorescence of D6-GFP\textsuperscript{*} cell clusters in qEBs.

\textbf{a’–d’} Immunostaining of the markers Sox2, Pax6, Tbr2, Tbr1 on the same sections. a’–d’ A higher magnification of merged images shows overlapping expression of D6-GFP and differentiation markers, indicated by \textit{rectangles} in \textbf{a–d}. Sox2\textsuperscript{+} and Pax6\textsuperscript{+} cells are scattered within the GFP\textsuperscript{*} cluster but Tbr2\textsuperscript{+} and Tbr1\textsuperscript{+} cells are organized in radial layers around a presumptive center (\textit{arrows} in e’ and d’).

Scale bars 100 \(\mu\text{m}\) in \textbf{a–d} and a’–d’ and 50 \(\mu\text{m}\) in a”–d” panels.

\textbf{e–e'} Immunostaining of N-cadherin (Ncad, red) with D6-GFP showing a neuroepithelial organization of qEB with N-cadherin in the apical pole of the rosette (\textit{arrows} in e’–e”). e–e’ Immunostaining of N-cadherin (Ncad, red) with Pax6 (green) showing a rosette with neuroepithelial structure with apical N-cadherin and basal Pax6.
for the pan-neural progenitor marker Nestin, while some cells expressed the Tuj1 marker of postmitotic neurons or the astrocyte-specific marker GFAP. Quantification of immunostaining documented that (87.7 ± 1.7)% cells were positive for Nestin, (9.6 ± 1.8)% for Tuj1 and (4.2 ± 1.1)% for GFAP (Fig. 8c). Thus, qEB derived D6-GFP-positive cells corresponded well to their counterparts from the mouse embryonic brain when grown under neurosphere culture conditions.

Discussion

In this study, we studied cell culture conditions that would yield the highest numbers of neocortex-specific neuronal cells. For this purpose, ESCs were derived from transgenic mice carrying the neocortex-specific D6-GFP reporter. This reporter turned out to be useful for tracking neocortical cells in differentiating ESCs using embryoid bodies. Interestingly, the cortical fate marked by D6-GFP was rarely reached in differentiation protocols using nEBs. In contrast, the qEBs culture, producing substantially larger cell aggregates, yielded a frequent induction of neocortex-specific cells. A related approach was used by Eiraku et al. (2008) for differentiation of ESCs into the neocortical tissue where the cell identity was monitored by immunohistochemistry with the standard cellular markers Pax6, Sox1, Bf1, or Tbr2. In addition, Eiraku employed a GFP knock-in into the Bf1 locus to directly observe cell differentiation. In comparison to Bf1-GFP, D6-GFP appears to be a more stringent marker for the neocortex as it visualizes with high specificity neocortical progenitors including radial glial cells and descendant neurons.

We think that cell communication plays a crucial role in acquirement of the neocortical cell identity. A larger mass of cells in qEBs probably resulted in proper cell signaling and self-organization that promoted cell differentiation towards the neocortical cell fate. This hypothesis is supported by the fact that GFP-positive cells were always found in large clusters rather than scattered single cells. Further, a longer time of in vitro culture is required for the induction of the neocortical cell fate compared to in vivo. The appearance of fluorescent cells peaked between 12 and

Fig. 6 D6-GFP+ cell quantification and gene expression analysis. a Flow cytometry analysis of qEBs without (left, negative control) and with D6-GFP+ cells (right). Selected samples from qEBs at d15–d18 show that up to one third of all cells are D6-GFP+. b Quantitative qRT-PCR of RNA from D6-GFP negative cells (gray columns) and positive cells (green columns) showing up-regulation of genes that are characteristic for the forebrain (FoxG1), or dorsal forebrain (Dach1, Pax6, Sox5)

Fig. 7 Activation of the canonical Wnt pathway during differentiation of qEBs promotes D6-GFP+ cell fate. Flow cytometry analysis of qEBs treated with recombinant Wnt3a (10 ng/ml) or Wnt-antagonist Dkk1 (50 ng/ml) at days 6–12 shows higher number of fluorescent cells in Wnt3a-treated qEBs and lower numbers in Dkk1-treated qEBs compared to untreated controls. As a control, retinoic acid (RA, 0.5 μM) was used to impose a caudal character in cultured neural cells which strongly inhibited neocortical fate as measured by D6-GFP+ cells
16 days whereas only 9 days are sufficient in the mouse embryo (from the blastocyst stage to E12).

Signals that directly control cell identity and organization of the neocortical tissue are not completely elucidated. Nonetheless, several reports have shown that signals including Fgf8 (Fukuchi-Shimogori and Grove 2001) and canonical Wnt signaling (Gunhaga et al. 2003; Backman et al. 2005) play an important role in this process. Our results suggest that the required inductive signals are produced within qEBs in a spatial and temporal manner that allows the induction of neocortical fate. The induction itself did not require supplementary signaling molecules but the addition of the recombinant Wnt3a in culture enhanced the process. Conversely, Dkk1, an inhibitor of the canonical Wnt signaling, reduced the number of D6-GFP-positive cells, which indicates that the Wnt pathway was activated at a physiological level in qEBs. Our experimental setup, however, was not able to unravel whether Wnt3a affected cell proliferation of cortical progenitors or an enhanced differentiation into cortical cells.

It is very intriguing that cell aggregates were organized in a way that resembled the polarized neuroepithelium in the embryonic brain (Abranches et al. 2009). It may well be that a few neuroepithelial cells with the polarized distribution of N-cadherin in the membrane induce a similar cell polarity in neighboring cells. Rosette-like structures can thus be formed via N-cadherin positive adherens junctions and this may provide a basis for the neocortical induction. Further experiments are necessary to examine the inner structure of the in vitro generated neuroepithelium.
example, genetic markers for all six layers of the cortex or various signaling molecules can be used to study its organization in more detail.

D6-GFP-positive neocortical tissue can be maintained in cell culture for more than 2 weeks. This rather broad time window can provide sufficient experimental time in tissue culture for studies on neuronal networks, discovery of new inductive signals in the neuroepithelium, or studies on small-molecule drugs that enhance or interfere with the process.

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