A restricted period for formation of outer subventricular zone defined by Cdh1 and Trnp1 levels

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The outer subventricular zone (OSVZ) is a germinal layer playing key roles in the development of the neocortex, with particular relevance in gyrencephalic species such as human and ferret, where it contains abundant basal radial glia cells (bRGCs) that promote cortical expansion. Here we identify a brief period in ferret embryonic development when apical RGCs generate a burst of bRGCs that become founders of the OSVZ. After this period, bRGCs in the OSVZ proliferate and self-renew exclusively locally, thereby forming a self-sustained lineage independent from the other germinal layers. The time window for the brief period of OSVZ bRGC production is delineated by the coincident downregulation of Cdh1 and Trnp1, and their upregulation reduces bRGC production and prevents OSVZ seeding. This mechanism in cortical development may have key relevance in brain evolution and disease.
A hallmark of neocortical development in gyrencephalic mammals is the formation of the outer subventricular zone (OSVZ). This is a specialized germinal layer that emerges at mid-corticogenesis, contains vast numbers of progenitor cells and is crucial for the evolutionary expansion of the mammalian cerebral cortex in neuron number and surface area, thus promoting the emergence of gyrencephaly. To date, the mechanisms regulating the developmental emergence of the OSVZ remain largely unknown.

The OSVZ contains a variety of basal progenitor cells, namely, basal radial glial cells (bRGCs) and intermediate progenitor cells (IPCs). IPCs are multipolar cells expressing the t-box transcription factor Tbr2, and are a major source of neocortical neurons in rodents. Basal RGCs resemble the apical RGCs (aRGCs) of the ventricular zone (VZ), including a radially extended basal process and expression of Pax6, but lack attachment to the apical adherens junction (AJ) belt. Basal RGCs are scarce in lissencephalic species, but very abundant in gyrencephalic mammals, where they are highly neurogenic and accumulate prominently in the OSVZ.

Although lissencephalic mammals may develop a vestigial form of an OSVZ, it is in gyrencephalic species where this is largest, most complex and mitotically active. In mouse, video microscopy studies of brain slices in vivo have revealed that IPCs and bRGCs have very poor self-renewing capacity, and their pool can only be maintained by the continuous production from aRGCs in the VZ. This process is finely regulated by the action of Trnp1, a DNA-binding protein that limits IPC and bRGc production. Similar analyses have shown that this process is much more complex in gyrencephalic species such as ferret, macaque and human, where IPCs and bRGcs in the OSVZ have been reported to proliferate and self-renew to some extent locally.

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that decreased by P6 to then remain similar until P14 (Figs 1y,z and 2b), demonstrating that OSVZ bRGCs are abundantly generated in the postnatal OSVZ. To investigate the dynamics of this bRGC production in OSVZ, we injected rv::Gfp at various stages and analysed after only 2 days of survival in each case. At P3 and P8, bRGCs represented 60–52% of cells born in OSVZ (Fig. 2c), indicating their net increase (see Methods). Video microscopy in slice cultures demonstrated self-amplification of bRGCs, including de novo generation of a basal process (Fig. 2d) as in primates8,10. Although OSVZ injections also labelled cells in ISVZ and VZ via their radial process (see Methods and Fig. 1w), our above analyses showed that these layers never generate OSVZ MN MP DN StC.
bRGCs, and therefore these were being generated only locally by rv::Gfp-infected progenitors within the postnatal OSVZ.

At all these stages, IPCs (defined as the subset of MP cells that are Ki67+, Tbr2+) in ISVZ and OSVZ were a very small minority of cortical progenitors, as opposed to the mouse SVZ (Fig. 1p,u,z; Supplementary Fig. 3; Supplementary Note 1). Moreover, the few IPCs in OSVZ were not generated by VZ or ISVZ progenitors, but mainly locally within OSVZ similar to bRGCs (Supplementary Fig. 2). Taken together, our findings demonstrated that the OSVZ is a unique niche of progenitor cell production at postnatal stages, following a lineage independent from progenitors in VZ and ISVZ, and where the vast majority of progenitors are bRGCs that expand by self-amplification.

Embryonic initiation of OSVZ by founder bRGCs seeded from VZ. In spite of our above findings, the key question of where and when OSVZ progenitors (mainly bRGCs) arise initially remained open. Given that all cortical cells derive from the early embryonic VZ at some point, we focused on earlier stages to determine the origin of OSVZ bRGCs. In ferret, the OSVZ is first distinguishable at E36 (equivalent to mouse E13), 6 days before birth (E42/P0; Fig. 3a), so we traced VZ progenitors with injections of rv::Gfp into the lateral telencephalic ventricle at E34 (equivalent to mouse E12.5) and E36 in utero (Fig. 3b). Two days after injection, numerous bRGCs expressed GFP in ISVZ, but none in OSVZ (Fig. 3c,d). Because 2 days of survival might be insufficient time for new bRGCs to migrate from VZ to OSVZ, we repeated the injections at E34, but now analysing at E38. Four gestational days after injection (E34–E38), 21.7% of bRGCs were in OSVZ (representing 8.0 ± 1.8% of all GFP + cells; Fig. 3d). These results demonstrated that bRGCs in the postnatal OSVZ had been originally produced by aRGCs at embryonic stages, though not anymore at postnatal stages.

The lineage tracing between E34 and E38 potentially allowed for several rounds of cell division, so we could not distinguish if bRGCs in OSVZ had been generated in a single step directly by aRGCs in VZ at E34 or indirectly following a second round of division in ISVZ between E36 and E38. To distinguish between these possibilities, we injected rv::Gfp at E34, administered BrdU at E36.0 and E36.5 to label cells cycling between E36 and E37 (S-phase ≈ 12 h)20,31, and analysed at E38 (Fig. 3e). BrdU administration was interrupted at E37 to avoid labelling bRGCs that might have reached the OSVZ early. Only 11.5% of all bRGCs in OSVZ contained BrdU (Fig. 3f–h), indicating that the majority had been directly generated by bRGCs in VZ before E36. In addition, rv::Gfp injection of VZ at E34 labelled some IPCs in OSVZ at E38 (MP, Ki67+; 0.46 ± 0.37% of GFP + cells), unlike their null labelling at postnatal stages (Supplementary Fig. 2). Hence, aRGCs in the embryonic VZ produce bRGCs and IPCs that migrate through the ISVZ en route to the OSVZ, a process not occurring after birth (E42) in the ferret cerebral cortex (Fig. 3i). Together, our results demonstrated that the OSVZ develops into two distinct phases: an early period of progenitor cell seeding from the VZ and a later period of self-amplification independent from other germinal layers.

A restricted period to massively produce OSVZ bRGCs from VZ. To precisely define the embryonic period during which the VZ produces bRGCs (for any germinal layer), we monitored the developmental dynamics of bRGC generation from aRGCs with rv::Gfp injections in utero and 2-gestational day survivals (Fig. 4a). At all ages, virtually all cells born from VZ (GFP +) were aRGCs or bRGCs (Fig. 4b,c), but their proportions changed significantly along development. At E32 and E34, 47–56% of GFP + cells were aRGCs, and only 29–34% bRGCs. Between E34 and E36, the production of bRGCs increased markedly to represent 66% of GFP + cells at E36, paralleled by the self-consuming of aRGCs (24%); aRGCs are not re-generated on division, but produce two daughter cells different from the mother aRGC; see Methods; Fig. 4b,c). This situation lasted only 2 days, and by E38 and P1, the generation of bRGCs was down to 53–45% of GFP + cells, respectively, and aRGCs were back up to 47–51% (Fig. 4b,c).

Given the short-survival time of these lineage experiments, and that the originally infected cells were aRGCs, we interpreted changes in the percentage of GFP + aRGCs as changes in cell-fate decisions at the population level: self-amplification (>50% of GFP + cells are aRGCs), self-renewal (50%) or consumption (<50% are aRGCs). Cell-fate decisions by aRGCs co-vary with their angle of mitotic cleavage plane, where vertical cleavage planes (60–90° to the VZ surface) occur in symmetric self-amplifying divisions and horizontal cleavage planes (0–30° to VZ surface) occur in asymmetric divisions generating bRGCs17,25,29,32,33. In agreement with this notion, we found that the proportion of horizontal mitotic cleavage planes in VZ peaked transiently at E34, following a temporal dynamics parallel to the depletion of aRGCs and the production of bRGCs, which are maximal for VZ progenitors infected at E34 (Fig. 4d–f; Supplementary Fig. 4). In summary, bRGCs are generated by aRGCs following self-consuming divisions with a burst between E34 and E36.

Unfortunately, these lineage tracings across embryonic development were done with only 2 days of survival, which is too short to determine if the bRGCs generated were destined to OSVZ or not (Fig. 3d). To investigate whether the embryonic production of OSVZ bRGCs from the VZ was narrowly restricted to the burst period between E34 and E36 (Fig. 4c), or it continued at later stages, we infected VZ progenitors in utero with rv::Gfp at
various stages and allowed 7–8 days of survival in each case, providing sufficient time for labelled cells to migrate and reach the OSVZ. We observed most OSVZ bRGCs in pups infected at E34, very few when infections were done at E36 and E38, and none in P1 injections (Fig. 4g,h). These experiments thus allowed us to identify and delineate the period of OSVZ bRGC production/seeding from the VZ with a prominent peak at E34 and very limited contributions at E36 and E38. As no OSVZ bRGCs are generated any longer at P1, their production is strikingly restricted to a rather brief period of cortical development (E34–P0), with a burst or maximum peak between E34 and E36.

Cdh1 and Trnp1 restrict the period for OSVZ generation. To identify candidate genes regulating the dynamics of bRGC production by aRGCs, we searched for differential gene expression in VZ between three key stages: E30, E34 and P1. Using a ferret-specific microarray34,35, we identified 1,852 differentially expressed genes (DEGs) between at least two stages (false discovery rate < 0.05, fold change > 2; Fig. 5a). Only 59 DEGs were found between E30 and E34 (27 upregulated and 32 downregulated), whereas 1,822 genes changed between E34 and P1 (871 upregulated and 951 downregulated). Because the production of bRGCs increased in the early period (E30–E34), but then decreased in the late period (E34–P1; Fig. 4c), we

Figure 2 | Postnatal aRGCs generate ISVZ bRGCs, while OSVZ bRGCs are generated by self-amplification. (a) Time-lapse imaging frames of an aRGC dividing apically, between 30 and 37 h of imaging time, to generate a bRGC (green arrowhead) retaining the basal process. The apical cell (red arrowhead) retained the apical process. (b) Abundance of GFP + bRGCs located in ISVZ after VZ and ISVZ infections at P1 or in OSVZ after OSVZ injections at P1, and analysed at the indicated ages (mean values ± s.e.m.; n = 420–5,544 cells, 2–5 animals per group; Supplementary Table 1). (c) Abundance of GFP + bRGCs in OSVZ at the indicated postnatal stages after injections of rv::Gfp in OSVZ and 2–3 days of survival. Data are mean ± s.e.m. (n = 420 cells, 3 animals, P1–P3; 1,288 cells, 4 animals, P6–P8; 819 cells, 2 animals, P14–P17); dashed line indicates 50%. (d) Time-lapse imaging frames of a bRGC in OSVZ with a basal process (black arrowheads), undergoing a near-horizontal division at t = 29:30 hours (dashed line) to generate two bRGCs. The basal daughter cell (top) retained the maternal basal process, whereas the apical daughter (bottom) grew a new basal process (solid red arrowheads) tipped with a small growth cone (red open arrowheads).
discarded DEGs whose expression levels changed in the same direction in both periods (first up- then again upregulated or first down- then again downregulated). Following the same reasoning, from the remaining DEGs we selected those whose expression levels changed in opposite directions between early and late phases (first up- then downregulated or down- then upregulated). Only one gene followed this ‘peak’ profile of expression, which encodes a synaptic protein (Sv2b) that is not expressed in mouse OSVZ.

Figure 3 | Embryonic aRGCs generate directly bRGCs for the OSVZ. (a) Nissl stains of the ferret embryonic neocortex illustrate the developmental progression of germinal layers and the first histological recognition of OSVZ between E34 and E36. (b-d) Ferret embryos received a ventricular injection of rv::Gfp at E34 or E36, and developed in utero until E36 or E38. GFP + bRGCs were found in OSVZ after 4 days of survival (E34–E38, detail), but only in ISVZ after 2 days (E34–36, arrow; n = 931 cells, 8 embryos; E36–38, n = 104 cells, 2 embryos; E34–38, n = 1,830 cells, 7 embryos; mean values + s.e.m.). (e-h) VZ progenitors were labelled with rv::Gfp at E34, BrdU was administered 2 days later and were analysed at E38 (f, g). Arrow in f points at the bRGC magnified in inset. Very few GFP + bRGCs in OSVZ contained BrdU (f, right; f, g; n = 229 cells, 5 embryos), indicating that most were directly generated from VZ. In c and f, solid arrowheads indicate aRGCs and open arrowheads indicate basal process of bRGCs. (i) A RGCs generate bRGCs that reach the ISVZ 2 days later (E36) and OSVZ 4 days later (E38). Few bRGCs in OSVZ are generated between E36 and E38 (likely from ISVZ), the majority being generated between E34 and E36 directly from VZ. Postnatally, aRGCs generate bRGCs for the ISVZ, but neither generates bRGCs for the OSVZ. Scale bar, 30 μm (E30 and E34; a), 100 μm (E36; a), 150 μm (E38; a), 200 μm (E42/P0, a); 25 μm (c); 150 μm (f).
cortical progenitors (www.genepaint.org), and has no known function in cell proliferation, cell fate or delamination\(^3\). At this point, we considered that the early phase (E30–E34) and late phase (E34–P1) might be regulated by different genes, so we distinguished between genes changing only in the early phase (‘early-change’ profile) or the late phase (‘late-change’ profile; DAPI   GFP   Pax6

Figure 4 | Transient peak of bRGC production from aRGCs at the onset of OSVZ. (a-c) Tracing of bRGC production from aRGCs along development. Ferret embryos aged E30, E32, E34, E36 and E43/P1 kits received an intraventricular injection of rv::Gfp and were analysed 2 days later. (b) GFP labelling in embryos injected as indicated; solid arrowheads indicate ventricular end feet of aRGCs, arrows indicate bRGCs and open arrowheads indicate basal process. (c) Abundance of GFP + aRGCs and bRGCs at the indicated ages. Production of bRGCs peaked at E36, when aRGCs self-consumed (<50%); ***\(P<0.001\), \(\chi^2\)-test; \(n=170\) cells, E32; 304 cells, E34; 931 cells, E36; 104 cells, E38; 1,094 cells, E45/P3; 2–8 embryos per group; mean values ± s.e.m. (d–f) Analysis of cleavage orientation plane of VZ mitoses with respect to ventricular surface (dashed lines; \(n=91\) cells, E30; 90 cells, E34; 124 cells, E36; 69 cells, E38; 2–8 embryos per group). Each dot in f represents 2% of mitoses. Data in e show that developmental variations in horizontal (0–30\(^\circ\), red) and oblique (30–60\(^\circ\), blue) cleavage planes were paralleled by bRGC and aRGC production, respectively, after rv::Gfp infection of VZ at those ages (grey curves). (g,h) Long-term GFP labelling of bRGCs in OSVZ (arrows) after intraventricular injection of rv::Gfp and analysis at the indicated ages. High magnifications show details of these bRGCs in OSVZ exhibiting a long basal process (open arrowheads) and expressing Pax6 (single confocal plane). (h) Abundance of GFP + cells corresponding to bRGCs in OSVZ at the indicated ages (E34–P0, \(n=513\) cells, 2 kits; E36–P1, \(n=958\) cells, 3 kits; E38–P3, \(n=906\) cells, 3 kits; P1–P10, \(n=1,499\) cells, 5 kits; mean values ± s.e.m.). Note that embryonic VZ progenitors generate bRGCs that populate the postnatal OSVZ, in contrast to postnatal VZ progenitors that do not. Scale bar, 40 \(\mu\)m (E32; b), 75 \(\mu\)m (E36 and 45; b); 15 \(\mu\)m (d); 100 \(\mu\)m (g).
Some of the early genes are related to axonal navigation (Slit and EphA5), whereas others are implicated in human diseases (LYPD1 and FAM167A). Late genes also include human disease genes such as BATF2 (acute stress disorder), GJB6 (sensorineural deafness and a variety of skin disorders) or APOE (lipoproteinemias), as well as a variety of biological functions such as cell adhesion (NDNF and THBS1), ribonucleotide binding (NABP1 and MSH3) or interaction with various organelles (FKBP4 and MARS2).

Unfortunately, functional gene annotation analysis of these gene sets showed that none of the ‘early’ DEGs were associated to biological processes related to bRGC production, such as mitotic spindle orientation or cell delamination from the apical junction belt\textsuperscript{2,19,37} (Fig. 5b; Supplementary Fig. 5), and hence were not mechanistically promising. As microarray analysis always includes many false negatives due to the stringent statistical criteria, we scrutinized the array for AJ-related genes and found a trend of regulation for Cdh1. Therefore, we re-screened our samples by qRT–PCR to analyse AJ-related genes\textsuperscript{30}. Among our tested candidates, \textit{Cadherin1 (Cdh1)} was the only gene with significantly lower messenger RNA levels in VZ at E34 compared with E30 (Fig. 5c). Low \textit{Cdh1} expression decreased >2.5-fold between these two ages, whereas \textit{βCat} increased 1.6-fold and other apical complex protein genes remained unchanged. *\(P<0.05\), **\(P<0.01\), t-test.

Figure 5 | Differential gene expression in VZ across developmental stages. (a) Heatmap of unsupervised hierarchical clustering of gene probes differentially expressed in VZ between E30, E34 and P1 (FC, fold change), and dendrogram of similarity between samples. Each lane is a biological replica. The vast majority of probes were differentially expressed between E34 and P1, but not between embryonic stages. (b) Examples of developmental expression profiles of differentially expressed genes, and for each profile the functional gene annotation terms as analysed and clustered using DAVID. The five clusters with the highest enrichment scores for up- and downregulated genes are shown. The vast majority of DEGs (FC >2, \(P<0.05\)) had ‘late-change’ profile (increase, red axis; decrease, blue axis; only 10 genes with highest FC in each axis are plotted), whereas much fewer had ‘early’ change profiles (41 probes). Red axes are for data represented in red, and blue axes for data in blue. (c) Quantitative RT–PCR data for candidate gene expression levels in VZ, expressed as FC between E34 and E30 (mean values ± s.e.m.). Negative values indicate lower expression at E34 and positive values indicate higher expression at E34. \textit{Cdh1} expression decreased >2.5-fold between these two ages, whereas \textit{βCat} increased 1.6-fold and other apical complex protein genes remained unchanged. *\(P<0.05\), **\(P<0.01\), t-test.
Of note, due to the high density of GFP + cells in SVZ on electroporation (Fig. 6e), bRGCs could not be identified by morphology, unlike with rv::Gfp; hence, aRGCs and bRGCs were identified here as GFP + cells expressing Pax6 + in VZ and SVZ, respectively. Overexpression of DN-Cdh1 caused a 2.5-fold increase in GFP + bRGC abundance and a significant decrease in GFP + aRGCs by E32 (Fig. 6e,f). Because only a proportion of aRGCs were electroporated, this decrease in GFP + aRGCs did not cause a general depletion of the VZ. Conversely, electroporation to overexpress Cdh1 at E34 was sufficient to abrogate the massive bRGC production normally occurring between E34 and E36, down to 33% of control embryos (Fig. 6f). Thus, reduced Cdh1 function is instrumental for the switch in mitotic cleavage plane orientation and loss of cell adhesion.

Regarding 'late' DEGs that might be responsible for the decrease in bRGC production between E34 and P1, Gene ontology analysis highlighted the terms cell adhesion, response to hormones, ions and steroids, organelle lumen, ribonucleotide binding and DNA repair (Fig. 5b). This list of 'late' genes included Trnp1, which was expressed twofold more at P1 than at E34 (Supplementary Table 3; Supplementary Fig. 5; Supplementary Note 1). Trnp1 is a protein previously involved in bRGC generation, but only in mouse (without OSVZ), where acute downregulation of Trnp1 causes overproduction of bRGCs25. Thus, our expression data suggested that low levels of endogenous Trnp1 in ferret VZ at E34 may favour massive bRGC production for the OSVZ, while higher Trnp1 levels at birth may close this period (Fig. 7a–c). To test the potential involvement of Trnp1 in regulating bRGC production for the OSVZ, we first used retroviruses to overexpress Trnp1 in the VZ of E34 embryos and

However, these changes did not perfectly phenocopy those observed between E32 and E34 in normal development (Fig. 4e), indicating that this process may be co-regulated by additional factors. These results demonstrated that the developmental loss of Cdh1 leading to increased bRGC production may act both on the switch in mitotic cleavage plane orientation and loss of cell adhesion.

Figure 6 | Onset of the restricted period for bRGC production and OSVZ generation depends on developmental variations of Cdh1 expression.
(a) Schematic of the early period of bRGC generation from aRGCs. (b,c) Candidate gene screening by qRT-PCR revealed Cdh1 as differentially expressed in VZ between E30, E34 and P1 (b, mean values ± s.e.m., **P < 0.01, ***P < 0.001, unpaired t-test), also shown by ISH (c). (d–f) In utero electroporation of DN-Cdh1 from E30 to E32 increased bRGC production (GFP + /Pax6 + cells in SVZ) and decreased aRGCs, whereas overexpression of Cdh1 from E34 to E36 had the opposite effect. In these experiments, GFP + cells in SVZ were at high density, making unreliable the identification of bRGCs by morphology. Instead, bRGCs were identified as Pax6 + /GFP + cells with the cell body in SVZ, and aRGCs as Pax6 + /GFP + cells with the cell body in VZ and a distinct apical process. Numbers of cells and embryos analysed are indicated in Supplementary Table 4; *P < 0.05, **P < 0.01, ***P < 0.001, χ²-test. (g–h) Analysis of cleavage orientation plane of VZ mitoses with respect to ventricular surface (dashed lines) in Gfp- and DN-Cdh1-ires-Gfp-overexpressing cells (n = 103 cells, Ctrl; 51 cells, DN-Cdh1; 2 embryos per group). Insets are details of each mitosis under 4,6-diamidino-2-phenylindole (DAPI) stain. Each dot in h represents 2% of mitoses. Scale bar, 150 μm (c); 75 μm (e); 15 μm (g).
analysed the cellular output at E36 (Fig. 7d). The abundance of GFP+ bRGCs was reduced to nearly half in Trnp1-overexpressing embryos compared with control embryos expressing GFP alone, and this was concomitant with a twofold increase in aRGC abundance (Fig. 7e,f). This demonstrated that low levels of endogenous Trnp1 expression are important for bRGC production between E34 and E36. To define if this decrease in bRGC generation indeed affects seeding of the OSVZ, we again overexpressed Trnp1 in VZ cells at E34, but now followed by long-term survival until E42/P0 (Fig. 7g). The OSVZ of Trnp1-overexpressing ferrets was nearly devoid of GFP+ bRGCs (these only represented 0.9% of GFP+ cells), concomitant with a twofold increase in aRGC abundance (Fig. 7e,f). This demonstrated that the dynamic temporal regulation of Cdh1 and Trnp1 expression is necessary and sufficient to control the variations in bRGC generation from aRGCs during cortical development. High expression of Cdh1 and Trnp1 is sufficient to limit bRGC generation before and after the restricted period, respectively, whereas low expression of both genes simultaneously is necessary for the massive self-consumption of aRGCs to produce bRGCs, and thus for the restricted period of OSVZ generation.

Discussion

Our study identifies a completely novel mechanism involved in cortical development, whereby the formation of a germinal layer (OSVZ) depends on the seeding of founder progenitor cells during a restricted period. We show that the OSVZ first emerges by the accumulation of large amounts of bRGCs generated directly by apical progenitors in the VZ. This seeding of progenitor cells from VZ to OSVZ is only transient, and eventually the OSVZ lineage becomes completely independent from the other germinal layers and thereon relies on the self-amplification of its progenitor cells for further expansion to its remarkable size at subsequent stages (Fig. 8). These are highly

Figure 7 | Closure of the restricted period for bRGC production and OSVZ generation depends on developmental variations of Trnp1 expression.

(a-c) Validation by qRT-PCR and ISH of differential expression of Trnp1 between E34 and P1 (later period of bRGC generation from aRGC; mean values ± s.e.m., ***P < 0.001, unpaired t-test). (d-f) Overexpression of Trnp1 by retroviral delivery from E34 to E36 decreased bRGC production and increased aRGC abundance, whereas expression of dominant-negative (DN) Trnp1 from P1 to P3 had the opposite effect (mean values ± s.e.m., ***P < 0.001, χ²-test). (g-i) Sustained overexpression of Trnp1 from E34 to P0 by retroviral delivery markedly blocked bRGC production for the OSVZ. Measures are relative to GFP+ RGCs, or all GFP+ cells in lineage (mean values ± s.e.m., *P < 0.05, **P < 0.01, ***P < 0.001, χ²-test). In all panels, solid arrowheads indicate apical end feet of aRGCs and open arrowheads indicate basal fibres of bRGCs. Scale bar, 150 μm (c); 75 μm (e,h).
unexpected findings, extending the complexity of the mechanisms that regulate cortical development and expansion3,4,26,43. We find that the period for bRGC seeding of the OSVZ depends on the combined temporal regulation of Cdh1 and Trnp1 expression in VZ (Fig. 8). These two genes were previously studied in mouse and shown to play key roles in the balance between self-renewal and delamination of aRGCs in the VZ25,40,44. Here we provide the first demonstration of a functional relationship between Cdh1 and Trnp1, and of their endogenous regulation to dynamically modulate bRGC production in the developing embryo. Importantly, we show that abrogation of bRGC production by maintaining elevated Trnp1 levels from E34 onward resulted in a severe reduction of OSVZ cell lineages in the long term (E42/P0). Therefore, given that this period of production of bRGC for the OSVZ is very brief (with a maximum peak between E34 and E36 and very limited contributions at E36 and E38), its timing is under tight genetic control, and the abrogation of bRGC production during this embryonic period results in their virtual absence in the postnatal OSVZ, this may be considered a critical period for the formation of the OSVZ. Given the relevance of the OSVZ in the development of gyrencephaly2,3,7,45, the temporally dynamic regulation of endogenous Trnp1 and Cdh1 expression during embryogenesis might have evolved as a mechanism generating cortical phenotypic diversity in mammals.

Previous video microscopy analyses of slice cultures from embryonic human and non-human primate cerebral cortex revealed a wide variety of progenitor cell types in the OSVZ and other germinal layers, and some of their complex lineage relationships8,10,29. However, the developmental origin of the OSVZ during gestation was never identified primarily due to the lack of appropriate in vivo models where this can be addressed. Likewise, the types of neurons generated by VZ, ISVZ and OSVZ cannot be studied in slice cultures due to survival time limitations, but only in vivo. One previous report in the early postnatal ferret demonstrated that the lineages of both VZ and OSVZ are neurogenic, producing layer 2/3 pyramidal neurons with no reported differences4. Here we use also the ferret to define the cellular substrates for the developmental origin and expansion of the OSVZ in vivo. The process we have identified in ferret is completely different than in the mouse cerebral cortex, where basal progenitors rarely self-renew and the SVZ is formed by transient populations of progenitor cells that are continuously seeded by the VZ and self-consumed shortly after13–16,24. Even in the mouse lateral ganglionic eminence, where the SVZ is largest and progenitors frequently self-amplify, there is continuous seeding from aRGCs26. Importantly, although rodents and primates are closer in phylogeny than with carnivores such as the ferret46, many of the typical features of OSVZ and bRGCs in ferret are strikingly similar to human and macaque, but different from mouse (that is, self-amplification, relative abundance of bRGCs and IPCs). Hence, we speculate that a similar critical period for the formation of the OSVZ, and its subsequent independent lineage, may also occur in human and other primates. Given that seeding of the OSVZ occurs only during a brief and transient period (E34–P0), the rate of self-amplification of its constituent progenitor cells becomes a major factor for its expansion and size. This notion is consistent with the dynamics of bRGC proliferation in the OSVZ observed in non-human primates, where this layer undergoes a massive increase in the size during a period of rapid bRGC self-amplification48. Intriguingly, the ISVZ seems to share features with the mouse SVZ, as it continuously receives bRGCs from the VZ while maintaining some self-renewing capacity. Expansive cell lineages are frequent in the mouse lateral ganglionic eminence (LGE)25, suggesting the evolutionary co-option of this strategy in forebrain development42.

One major difference between aRGCs and bRGCs is their epithelial nature, where aRGCs are anchored to the ventricular surface by AJs, while bRGCs are not. Thus, the generation of bRGCs from aRGCs may be favoured by oblique and horizontal mitotic cleavage planes, combined with an active loss of AJs, leading to the apical detachment of daughter cells17,25–27,29. Orientation of mitotic spindle is molecularly regulated by a protein complex including mInsc, LGN, Par3, aPKC and also Cdh1, the blockade of which favours oblique and horizontal cleavage planes and results in an excess of daughter cells delaminating from the ventricular surface33,41,47,48. Accordingly, we find that the onset of the restricted period with a massive generation of bRGCs from aRGCs is bookmarked by a significant decrease in Cdh1 expression and a shift in mitotic cleavage plane orientation, and we have demonstrated a causal relationship between these three events. However, the combined frequency of divisions with oblique or horizontal cleavage planes that we have observed at the critical period (E34: 50%) can only explain the delamination from VZ of 25% of daughter cells, but not 70% as we observed by lineage tracing (Fig. 4). The delamination of additional cells may require the loss of apical anchoring, which occurs on the loss of AJs proteins, their links to the cytoplasm such as α- and β-catenin or regulators of the F-actin belt linking AJs such as RhoA30,49. Among these factors, Cdh1 is a crucial component of AJs previously implicated in regulating apical cell delamination (though not bRGC production), and here we show that the loss of Cdh1 leads to the loss of aRGCs and increased bRGC production. Hence, our present findings point at Cdh1 as...
key in the production of bRGCs from aRGCs by regulating both mitotic cleavage plane orientation and delamination from the AJ belt. Repression of cadherins and cell delamination from VZ is promoted by the Snail family of transcription factors, and also by Robo receptor signalling. Unfortunately, current detection methods are insufficient to reveal differences in membrane-bound Cdh1 between individual aRGCs primed for delamination versus not. Finally, closure of the period of massive bRGC production at E36 is marked by an increase in expression of genes which favours vertical cleavage planes and has been shown to repress bRGC production in mouse.

Our results demonstrate that the combined regulation of endogenous Cdh1 and Trnp1 levels is necessary and sufficient to control the abundance of VZ cell depletion and bRGC seeding, and thus it defines the time window for our newly identified critical period of OSVZ formation. Our Cdh1 and Trnp1 downregulation experiments demonstrate that aRGCs have the intrinsic potential to generate OSVZ bRGCs before and after the critical period, so the onset and duration of this period could be easily modified in phylogeny or disease by changing the temporal expression of these genes. Given the relevance of the OSVZ in cortical expansion and folding, the evolution of mechanisms controlling the precise temporal regulation of critical period genes seems key to generate the extraordinary diversity of cortical phenotypes across mammals.

Critical periods are unique windows of opportunity during development, but also periods of high vulnerability to disease. Given the central role of the OSVZ in cortical development from ferrets to monkeys and humans, including neurogenesis and surface area expansion, the critical period of OSVZ seeding is a time of susceptibility to cortical disease, where subtle or acute defects may have magnified long-term consequences. Indeed, acute abrogation of cell proliferation in ferret embryos at the onset of this critical period leads to reduced OSVZ and lissencephaly. In humans, brain malformations due to acute insults in fetal development are rarely traced, but defects in VZ integrity, progenitor proliferation and precise gene expression regulation are emerging as critical malformations of cortical development. Our identification of the critical period for OSVZ formation in ferret brings a novel perspective to fundamental mechanisms of cortical development, and this may help to better frame the mechanistic effect of mutations perturbing cerebral cortex development in humans.

**Methods**

**Animals.** Pigmented ferrets (*Mustela putorius furo*) were obtained from Marshall Bioresources (North Rose, NY) and isoquimen (Barcelona, Spain), and kept on a 16:8 h light-dark cycle at the Animal Facilities of the Universidad Miguel Hernández. Wild-type mice were maintained in an Institute for Cancer Research: Harlan Inc. (ICR) background. The day of vaginal plug was considered as embryonic day (E) 0.5. All animals were treated according to Spanish and EU regulations, and experimental protocols were approved by the Universidad Miguel Hernández Institutional Animal Care and Use Committee (IACUC).

**Constructs.** For retroviral delivery, constructs encoding Gfp alone, Trnp1-Gfp, fusion protein (Trnp1-DN) or a bicistronic cassette encoding Trnp1-IRES-Gfp were subcloned into a murine melanoma virus (MMLV) retroviral-packaging vector downstream of the CAG promoter (generous gift of F.H. Gage). For electroporation constructs encoding Gfp, E-Cadherin (Addgene #28009) or DN-Cdh1 ref. 40) were subcloned into a pCAG promoter-containing vector. All plasmids were produced under endotoxin-free conditions (QIAGEN EndoFree Plasmid Maxi kit).

**Virus injections and electroporation.** High-titre MMLV-based SVGV-pseudotyped retrovirus (5 x 10^5−5 x 10^6 p.f.u. ml^−1) encoding Gfp, Trnp1-IRES-Gfp or Trnp1-Gfp under the CAG promoter were prepared by transient transfection of HEK293 cells, concentrated by ultracentrifugation and viral titre estimated by clonal infection of HEK cell cultures. High-titre replication-incompetent adenovirus (9.3 x 10^11 p.f.u. ml^−1) encoding Gfp under the CMV promoter were obtained from QBiogene (Irvine, CA). Viral solutions were injected using pulled glass micropipettes. For postnatal injections, ferret kits were deeply anaesthetized and anaesthetized during surgery, and the entire period of massive bRGC production at E36 was marked by the telencephalic lateral ventricle, ISVZ or OSVZ by means of stereotaxic coordinates: lateral ventricle: antero-posterior (AP) = −0.5 mm, latero-medial (LM) = 2.0 mm, dorsal-ventral (DV) = 1.0 mm, with an AP inclination of 22.5°; ISVZ: AP = −1.3 mm, LM = 2.0 mm, DV = 2.0 mm, with an AP inclination of 22.5°; OSVZ: AP = 1.3 mm, LM = 1.0 mm, DV = 2.0 mm, with an AP inclination of 45°. These infections were very accurate, as confirmed a posteriori with short-survival animals where the bulk of GFP-labelled cells indicated the tip of the injection pipette (for ISVZ and OSVZ injections), and sparse cell labeling in VZ suggested that the cortex indicated in the injection was actually infected in the lateral telencephalic ventricle (Fig. 1). We only found 2 out of 11 animals where injections aimed at ISVZ actually labelled a sparse number of VZ cells, and thus had occurred in the ventricular cavity. On injection in ISVZ and OSVZ, aRGCs cells in VZ and bRGCs in ISVZ were also labelled immediately below the injection site, injected retrogradely via their basal process.

For in utero injections in both ferret and mouse, timed-pregnant females were deeply anaesthetized and maintained in 2% isoflurane during surgery. The abdominal cavity was open, the uterine horns exposed and retrovirus solutions were pressure injected into the telencephalic lateral ventricle of embryos through the uterine wall.

For electroporation, DNA plasmids encoded Gfp, E-Cadherin (Cdh1) or DN-Cdh1, under the CAG promoter. In utero and postnatal electroporation of ferret embryos and kits were performed as described. After the appropriate survival period, postnatal kits or pregnant females were overdosed with sodium pentobarbital (Nembutal), and further processed for immunohistochemistry or in situ hybridization (ISH).

**Immunohistochemistry and ISH.** For histological analysis, embryos were obtained by caesarean section of timed-pregnant females on day 16 or day 17. Animals were deeply anaesthetized with sodium pentobarbital, and perfused transcardially with 4% paraformaldehyde; postnatal ferrets were deeply anaesthetized with sodium pentobarbital before transcardial perfusion with paraformaldehyde. After perfusion, brains were extracted and sectioned. For immunohistochemistry, brain sections were incubated with primary antibodies overnight (anti-GFP, 1:1,000, Aves Labs; anti-Pax6 1:500, Millipore; anti-Tbr2 1:200, Abcam; anti-PhVim 1:1,000, Abcam; anti-PH3, 1:1,000, Upstate; anti-Ki67, 1:200, Novoceastra; anti-BrdU, 1:200, Abcam; anti-GFAP, 1:1,000, Dako; anti-Olig2, 1:200, IBL), followed by appropriately fluorescently conjugated secondary antibodies (1:200, Jackson), and counterstained with 4,6-diamidino-2-phenylindole (Sigma). For anti-BrDU staining, sections were pretreated with 2 N HCl for 30 min. For ISH, sense and anti-sense complementary DNA probes were synthesized and labelled with digoxigenin (Roche Diagnostics) according to the manufacturer’s instructions. ISH was performed as described previously. For ferret-specific ISH probes were cloned using the following primers: Trnp1, forward: 5'-TCTGACCTCCCTGGATGACG-3', reverse: 5'-TGGCGCTGTTGCTTTACGGAGC-3'; Cdh1, forward: 5'-CGAGGCCTGATGTTTCGGTC-3', reverse: 5'-GAAGGGCTGATGTTCTCTTCGG-3'. These ferret-specific primers were designed based on the same ferret-specific sequences as in the microarray. The PCR reaction was carried out using Go Taq Flexi DNA polymerase (Promega), and the resulting amplicons were purified with Wizard SV Gel and PCR Clean-Up System (Promega) and cloned into pGEM-T Easy Vector System I.

**Slice culture and time-lapse imaging.** Ferret brain slices were prepared and maintained in culture as described. To image aRGC behaviour, P1 ferrets were injected with rC-Gfp, and their brains obtained for slicing at P8. To image the behaviour of bRGCs in OSVZ, slices were prepared from unmanipulated P18 ferrets, and 1 h later, slices received various injections of Ad-GFP (3–6 nl per injection site, 2–8 injection sites per slice) in the cortical plate. One day after slice preparation, slices containing fluorescent cells were selected for time-lapse imaging. Images were obtained either under fluorescence optics (Filter 410/17, EX449-489/EM500-548; Chroma, Rockingham, VT, USA) through an air immersion ×20 lens in an inverted epifluorescence microscope or under two-photon optics (×40) through a water immersion ×20 lens in a Leica SP2 inverted microscope, both equipped with an incucubation chamber: 5% CO2, 37°C. Frames were obtained intermittently during 49–142 h in culture. Digital images were acquired, contrast-enhanced and analysed with Metamorph (Microbrightfield) or Imaris software (Bitplane).

**Microarray and qRT-PCR.** For RNA extraction, P1 ferret kits were deeply anaesthetized, and timed-pregnant ferret females were deeply anaesthetized and their living embryos were obtained by caesarean section. Subjects were then decapitated, their brains dissected and blocked in ice-cold ACSF (1:40 mM NaCl, 5 mM KCl, 1 mM MgCl2, 24 mM NaHCO3, 1 mM KH2PO4, 1 mM CaCl2, pH 7.2), and tissue blocks containing the occipital cortex were vibrotome-cut in 300-μm-thick slices. Living cortical slices were further microdissected with microscissors in ice-cold ACSF to isolate the VZ from the caudal pole of the cerebral cortex. Germinial layers were identified in living slices under the dissecting stereozoom. After the VZ was microdissected, a 250-μm-thick layer on the apical side of this VZ. Tissue pieces were fresh-frozen in Trizol for RNA extraction, with a post-mortem interval of <1 h.
Total RNA was extracted using RNeasy Mini kit (Qiagen), followed by treatment with RNase-Free DNase Set (Qiagen). RNA quality was confirmed using the RNA 6000 Nano kit on the Agilent Bioanalyzer platform, and then 200 ng of total RNA was labelled using the one-colour labelling kit from Agilent Technologies according to the manufacturer’s protocol. Labelled complementary RNA was then hybridized for 16 h on a custom-made microarray containing 43,692 ferret-specific probes covering 17,386 genes. Microarray slides were scanned on an Agilent High-Resolution C Scanner, and the raw image files were processed by the Agilent feature extraction software. Raw data files were normalized using quantile normalization in Partek Genomics Suite. Statistical analysis of microarray data was done in Multiexperiment Viewer44. To identify genes with significantly different expression levels, we used analysis of variance comparisons between samples, using P-values based on 500 permutations and Bonferroni’s false discovery correction, as in Ayoub et al.65. Functional gene annotation analysis was performed using the web-based DAVID v6.7 software (http://david.abcc.ncifcrf.gov)66. The microarray data from this publication have been submitted to the GEO database (http://www.ncbi.nlm.nih.gov/geo/) and assigned the identifier GSE63203.

For qRT–PCR, primers for ferret gene homologues were designed based on the same ferret-specific sequences. Template complementary DNA was generated using maximal First Strand cDNA Synthesis kit for quantitative real-time PCR (qRT–PCR; Thermo Fisher). Quantitative RT–PCR was performed using the One Step plus sequence detection system and the SYBR Green method (Applied Biosystems), with each point examined in triplicate. Transcript levels were calculated using the comparative Ct method normalized using actin. Primers used were: Tmp1, forward: 5′-TGGGCTGAGAATCTCCTGC-3′; reverse: 5′-CGCTGCTGTCCTCTGGCAGG-3′; Cdh1, forward: 5′-GCCGCAGAAATGGCAGGA-3′; reverse: 5′-AACAAATACACCAAAACGGA-3′; actin, primers as in ref. 67. In each experiment group, we analysed two to three samples, each consisting of a pool of two to three embryos/kits. Reactions were performed in triplicate per independent sample. Data were statistically analysed with SPSS software using t-test.

Cell count measurements. Cell types were identified according to the following criteria: aRGC, cell with the soma located within the VZ, a single apical process contacting the ventricular surface and a very long radially oriented basal process, expressing Pax6 and Ki67; bRGC, cell with the soma located outside the VZ, with a very long radially oriented basal process, without an apical process anchored to the ventricular Al belt, expressing Pax6 and Ki67; MP, cell with multiple short processes extended from the cell soma and without obvious polarity; IPC, MP cell expressing Ki67 and Tbr2; migrating neuron, cell with clear apical–basal polarity extending a relatively thin basal process much shorter than that of RGCs, and a much shorter and thinner apical process; DN, cell with a single basally-directed process highly branched at a distance from the cell body, frequently located at the top of the cortical plate; PC, multipolar cell with several very highly branched and short processes extending from the cell soma. In electroporation experiments, identification of GFP+ cell identity in SVZ on morphology was unreliable due to their high density. In those experiments, aRGCs were defined as Pax6+/GFP+ cells with the cell body in VZ and a distinct apical process, while bRGCs were identified as GFP+ cells with the cell body in SVZ. The angle of mitotic cleavage plane was measured using PH3 stains and considering only cells at telophase. Angles were measured with respect to the general trajectory of the radial fibre scaffold, or with respect to the ventricular surface. Images were considered horizontal if they occurred at 60–90° with respect to radial fibres or 0–30° with respect to the ventricular surface.

Double-labelling analyses. Quantification of cell co-staining was performed by confocal microscopy (Leica) through a ×40 lens and ×2–4 zoom. Images were acquired from cells in four sections per subject, two to three subjects per condition and age. Images were analysed using Imaris software (Bitplane) and Canvas X software.

Progenitor self-amplification. To determine whether the population of RGCs increased or not, we profited from a unique property of retroviruses: the GFP appearance of the neuronal phenotype in apical versus basal progenitor non-surface-dividing cortical progenitor cells. Live imaging at the onset of cortical neurogenesis reveals differential morphological features of the proliferative zones and postmitotic compartments of the neural epithelium giving rise to striate and extrastriate cortex in the monkey. Proc. Natl Acad. Sci. USA 101, 3683–3693 (2011).

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