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Citation for published version:
Kazanis, I & Ffrench-Constant, C 2012, 'The Number of Stem Cells in the Subependymal Zone of the Adult Rodent Brain is Correlated with the Number of Ependymal Cells and Not with the Volume of the Niche' Stem Cells and Development, vol 21, no. 7, pp. 1090-6., 10.1089/scd.2011.0130

Digital Object Identifier (DOI):
10.1089/scd.2011.0130

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Publisher's PDF, also known as Version of record

Published In:
Stem Cells and Development

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The Number of Stem Cells in the Subependymal Zone of the Adult Rodent Brain is Correlated with the Number of Ependymal Cells and Not with the Volume of the Niche

Ilias Kazanis1 and Charles ffrench-Constant2

The mammalian subependymal zone (SEZ; often called subventricular) situated at the lateral walls of the lateral ventricles of the brain contains a pool of relatively quiescent adult neural stem cells whose neurogenic activity persists throughout life. These stem cells are positioned in close proximity both to the ependymal cells that provide the cerebrospinal fluid interface and to the blood vessel endothelial cells, but the relative contribution of these 2 cell types to stem cell regulation remains undetermined. Here, we address this question by analyzing a naturally occurring example of volumetric scaling of the SEZ in a comparison of the mouse SEZ with the larger rat SEZ. Our analysis reveals that the number of stem cells in the SEZ niche is correlated with the number of ependymal cells rather than with the volume, thereby indicating the importance of ependymal-derived factors in the formation and function of the SEZ. The elucidation of the factors generated by ependymal cells that regulate stem cell numbers within the SEZ is, therefore, of importance for stem cell biology and regenerative neuroscience.

Introduction

In mammalian species investigated thus far, some level of constant neurogenic activity driven by neural stem cells (NSCs) occurs within niches in the mature, adult brain. Such activity has been identified in the subependymal zone (SEZ) of the lateral walls of the lateral ventricles and in the subgranular zone of the dentate gyrus, within the hippocampal formation [1]. These neurogenic niches are facilitators of stem cell activity, offering protection to stem and daughter precursor cells and allowing their regulated proliferation, migration, and differentiation. The discovery of neurogenic niches in the adult brain has opened new possibilities for the treatment of degenerative diseases of the central nervous system (CNS) [2]. The level of contribution of these niches to tissue repair depends on their cell generation capacity; and the elucidation of the factors that support stem cells within niches such as the SEZ is, therefore, of great importance in regenerative medicine.

Previous experimental work has suggested that neurogenesis within the mouse SEZ is structurally and functionally correlated with the vasculature and with the cerebrospinal fluid (CSF)/ependymal cell layer interface, with both of these elements associated with neurogenic activity in their proximity [3–6]. Adult NSCs are directly connected to each element (CSF/ependymal cell and vessel) by their apical and basal processes, respectively. These attachments represent persistence of those to ventricular surface and basal lamina of the embryonic radial glial cells from which the adult NSC are generated. However, the relative contribution of signals from each element to the regulation of adult NSCs remains poorly defined. One way to gain further evidence as to the roles of the blood vessels and ependymal cells in the formation of the niche is to investigate how neurogenic activity and the structure of the niche change in different conditions; for example, during the early postnatal development of the SEZ [7] or in response to degenerative insults [3,8]. An alternative approach, however, is to explore evolutionary traits by directly comparing the neurogenic activity and the structure of the niche in different animal species. Here, we take the latter approach by comparing the SEZ of the mouse and the rat. These 2 species are the most widely used experimental animals; they are evolutionary [9], ecologically, and behaviorally related, but importantly, differ significantly in their brain size. By analyzing this naturally occurring example of scaling of the SEZ, we were able to show a close correlation between the number of NSCs and the size of the ependymal cell layer, thus suggesting an important role of ependymal cells in generating signals that control stem cell behavior.
Materials and Methods

Animal treatments and tissue processing

Two-month-old adult Sprague-Dawley rats and 129sv mice were used, and all experiments were performed in accordance to the United Kingdom Animals (Scientific Procedures) Act 1986. For Cytosine β-D-arabinofuranoside (AraC; Sigma) treatment, adult animals were anesthetized, and a cannula (Bik-II; Alzet) was fixed on the skull (1 mm lateral to bregma) connected to a subcutaneously implanted mini-osmotic pump (1007D; Alzet). 4% AraC or saline alone was infused for 4 and 7 days onto the surface of the brain of mouse and rats, respectively, and animals were sacrificed 2 days after the end of the infusion. Vibratome-cut (Leica) 70 μm thick brain sections were processed for immunohistochemistry using the rabbit antiphosphohistone 3 (anti-PH3; 1/500; Sigma), mouse anti-glial fibrillary acidic protein (anti-GFAP; 1/500; Sigma), rabbit-pan Laminin (1/200; Sigma) and rabbit anti-S100β (prediluted; Dako) antibodies, followed by incubation in appropriate Alexa-conjugated secondary antibodies (Invitrogen). Images were acquired using a Leica SP5 confocal microscope and were processed using Photoshop (Adobe) software.

Analysis of structure

Three rostro-caudal levels were chosen (Table 1, Section B and Fig. 1), and at least 5 sections were analyzed from each level. To assess GFAP and PH3 coexpression at least 3 confocal-generated optical sections from each mitotic cell (using the ×63 objective and a further ×4 digital zoom) were analyzed (Fig. 2, Supplementary Figs. S1–S3; Supplementary Data are available online at www.liebertonline.com/scd). For the calculation of the distance from the ventricular wall, measurements were performed as previously described [3] (and Supplementary Fig. S1) using the ImageJ software in at least 3 sections from each rostro-caudal level analyzed in each animal. ImageJ was also used to calculate the size of the nuclei of ependymal cells and the volume of the rat SEZ. For the latter, the volume estimated based on the complex

| Table 1. The Structural Characteristics of Neurogenic Activity (Sections A and C) and of the Anatomy of Different Brain Areas (Section B) Explored in This Study Are Presented, Focusing on the Comparison Between the Mouse and the Rat |
|-------------------------------------------------|----------|--------|
| (A) Positioning of PH3 immuno-positive cells | Mouse | Rat | Rat/mouse |
| Maximal distance of mitotic cells from the ventricular wall (mm) | 0.022±0.003 | 0.055±0.011* | 2.50 |
| Maximal distance of mitotic NSCs from the ventricular wall (mm) | 0.011±0.005 | 0.021±0.003 | 1.91 |
| Average distance of mitotic NSCs from the ventricular wall (mm) | 0.009±0.003 | 0.012±0.009 | 1.33 |
| (B) Indicative dimensions of areas and cells | | |
| Level 1: [0.02 and –0.26 mm from Bregma in mouse and rat, respectively] | 1.75±0.04 | 3.00±0.08 | 1.71 |
| Dorso-ventral dimension of the lateral ventricle (mm) | | |
| Level 2: [0.62 and 1.0 mm from Bregma in mouse and rat, respectively] | 1.75±0.02 | 3.40±0.05 | 1.94 |
| Dorso-ventral dimension of the lateral ventricle (mm) | | |
| Level 3: [1.94 and 2.70 mm from Bregma in mouse and rat, respectively] | 0.40±0.03 | 0.60±0.05 | 1.50 |
| Dorso-ventral dimension of the lateral ventricle (mm) | | |
| Overall rostro-caudal dimension [between level 3 and level 1] (mm) | 1.92 | 2.96 | 1.54 |
| Olfactory bulbs | | |
| Maximal rostro-caudal dimension of the olfactory bulb (mm) | 3.5±0.02 | 5.5±0.07 | 1.57 |
| Maximal dorso-ventral dimension of the olfactory bulb (mm) | 2.9±0.04 | 5.0±0.03 | 1.72 |
| Estimated surface of the ventricular wall between levels 1 and 3 (mm²) | 2.79 | 7.43 | 2.66 |
| Estimated volume of the SEZ between levels 1 and 3 (mm³) | 0.05 | 0.24 | 4.75 |
| Size of ependymal cells (maximal radius of nucleus) (μm) | 5.97±0.56 | 7.43±0.59 | 1.24 |
| Neurogenic area per ventricular wall length (in coronal sections using arbitrary units) | 1.67 | 2.75 | 1.65 |
| (C) Quantification of NSC and ependymal cell numbers | | |
| Number of mitotic NSCs per ventricular wall surface (cells/mm²): | | |
| normal SEZ | 13 | 14 | 1.06 |
| Number of mitotic NSCs (approximation of total NSCs) per ventricular wall surface (cells/mm²): regenerating SEZ | 119 | 131 | 1.10 |
| Number of ependymal cells per ventricular wall surface (cells/mm²) | 22,200 | 20,600 | 0.93 |
| Estimated total number of mitotic NSCs per SEZ at a random time during homeostasis | 35 | 105 | 3.00 |
| Estimated total number of mitotic NSCs (approximation of total number of NSCs) per SEZ during regeneration | 360 | 973 | 3.08 |
| Estimated ratio of ependymal cells per mitotic NSCs during homeostasis | 1460 | 1687 | 1.15 |
| Estimated ratio of ependymal cells per mitotic NSCs (approximation of total NSCs) during regeneration | 157 | 187 | 1.19 |

Note that although the width of the SEZ (estimated by the maximal distance from the ventricular wall where mitotic progenitors occur) is significantly higher in the rat, the periventricular distribution of mitotic NSCs is similar in the 2 species (Section A). Also, note that the distribution of mitotic cells per ventricular wall surface is similar among the 2 species (Section C).

*P<0.05 using the nonparametric Kruskal–Wallis test.

SEZ, subependymal zone; NSC, neural stem cell; PH3, phosphohistone 3.
trapeziums shape (similarly to the mouse SEZ, Fig. 1) had to be corrected for the observed irregularities. To achieve that, the surface of the neurogenic area was defined and measured in at least 4 domains of the SEZ per section and in serial confocal-generated optical sections (as shown in Fig. 3 and Supplementary Figs. S1, S3). Subsequently, the size of the neurogenic area was expressed per length of ventricular wall (Table 1, Section B), and this was used to adjust the initially calculated volume. For NSC number calculations, the total number of GFAP, PH3 double-positive cells was counted in each section; whereas for ependymal cells, S100β and GFAP immunostainings were performed in at least 3 sections from each rostro-caudal level analyzed in each animal, and the total number of ependymal cells (defined either as S100β+ cells or as GFAP− cells lining the ventricle; see Fig. 3 and Supplementary Fig. S2) was counted in at least 3 domains of the SEZ in each section and it was expressed per ventricular wall surface.

Results and Discussion

Initially, we sought to compare the size (volume) of the mouse and rat neurogenic niche by defining its dimensions and shape. The neurogenic area was defined using both histological and immuno-histological criteria, based on the observation that mitotic cells were positioned within a cell-dense zone, proximal to the ventricle, which was followed deeper in the tissue by a hypocellular, mitoses-free zone, that we considered to be outside the SEZ (Supplementary Fig. S4). Three indicative rostro-caudal levels (Fig. 1A and Table 1, Section B) were selected, and specific anatomical marks -such as the shape of the lateral ventricles and of the corpus callosum and the position of the anterior commissure were used to define the respective levels in the 2 species (n = 4 and 6, mice and rats, respectively). The dorso-ventral dimensions of the mouse and rat SEZ were calculated in sections collected from these 3 rostro-caudal levels (Table 1, Section B) and were found to be comparable to data obtained by other sources [10,11]. The third dimension of the niche (the width, in a coronal section perspective; Fig. 1A) was estimated by measuring the maximal distance from the ventricular wall where neurogenic activity occurs. Dividing cells (that comprise both NSCs and daughter transit amplifying precursors and neuroblasts) were identified by immunostaining serial 70 μm-thick sections of adult brains for PH3, a marker of cells proceeding through the M phase of cell cycle, and the distance of each mitotic nucleus from the ependymal cell layer was calculated (Supplementary Fig. S1). In addition, NSCs within these dividing (PH3-positive) cells were identified as had been previously done [3] based on co-expression of the astrocytic marker GFAP (expressed by NSCs and not by downstream precursors; Fig. 2 and Supplementary Figs. S2, S3).

The mouse SEZ was found to be a compact zone, homogeneously extending 3–4 cell layers adjacent to the ependymal cell layer (Fig. 3 and Supplementary Fig. S3), with the PH3+ immunohistochemistry data showing that all dividing cells (GFAP+NSCs and GFAP− precursors) were positioned up to 22 μm from the ventricular wall (Table 1, Section A). In the rat, the PH3 immunostaining data showed that neurogenic activity occurred up to 55 μm from the ventricular wall; and so, the SEZ was predicted to extend 12–14 cell layers adjacent to ependymal cells. However, the histological analysis revealed the rat SEZ to be irregular in its width; comprising areas formed by 3–4 cell layers (similarly to the mouse SEZ) and areas consisted of up to 12–14 cell layers (Fig. 3 and Supplementary Figs. S3, S4). Another difference between the mouse and rat SEZs was that mouse ependymal cells were flat, elongated in parallel to the ventricular wall, whereas rat ependymal cells were more cuboid, in some cases bearing processes perpendicular to the ventricular wall (Fig. 3). These observations indicate that the morphology of the ependymal cell layer is modified in association with the widening of the SEZ, and rat ependymal cells (some of which bear thin and short basal processes) are morphologically intermediate between the flat mouse ependyma and the ependyma of the primate and human SEZ that exhibits long basal processes [12,13].

Based on the observations regarding the width of the SEZ as well as on the shape of the ventricular wall in the rostro-caudal and dorso-ventral axes, the mouse and rat SEZs were modeled to resemble 3-dimensional (3D) structures composed by 2 trapeziums (Fig. 1B). In addition, the volume of the rat SEZ was corrected for the irregularities of its width, as calculated from tissue sections (see Materials and Methods section). These calculations revealed that the rat SEZ was approximately 4.75 times bigger than the mouse SEZ. This estimate was very close to the previously reported difference in the size of the whole brain (the rat brain being approximately 4.33 times bigger than the mouse brain) [14], in concert with the theory that during evolution the scaling in the size of the brain between 2 species is reflected in analogous scaling in the size of brain subdivisions [15]. To further validate our result, we also made volumetric calculations for the olfactory bulbs (OBs), areas that are functionally related to the SEZ, as they are the targets of SEZ-generated cells, based on the assumption that OBs resemble cylinders (Supplementary Fig. S5, Table 1, Section B). Rat OB were estimated to be 4.67 times bigger from the mouse OB. Taken together, these results suggest that the volume of the rat SEZ neurogenic niche is scaled, as compared with the mouse SEZ, in proportion to the scaling of the whole brain.

This expansion of the rat SEZ would be predicted to have different effects on the availability of the 2 potential sources of factors that regulate NSCs. Any factors released by sources present within the entire 3D space of the SEZ, for example, those derived from the vasculature, would expand in proportion to volume, whereas those released from the ependymal cells lining the ventricle and, therefore, forming a 2-dimensional sheet would show a much smaller expansion (just as the area of each side of a cube only expands as a square of the length of each side, whereas the volume of the entire cube expands as a cube of this length). Next, therefore, we calculated the correlation between NSC numbers, SEZ volume, and area of the ependymal layer. The total number of PH3-positive NSCs was counted in each rat and mouse tissue section, and it was revealed that the rat SEZ was 4.75 times larger but contained only 2.7 times more mitotic NSCs (resulting in a 42% decrease in their density). However, when the increase in the number of mitotic NSCs in the rat was compared with the increase in the ventricular wall surface (2.66 times larger than that of the mouse), the ratio was similar in both species (Table 1, Section C).

Although these results suggest that the NSC population expands in proportion to the area of the ependymal cell layer
rather than the volume of the SEZ, an alternative explanation would be that total NSC numbers do expand in proportion to SEZ volume but the balance between quiescent (non-mitotic) and activated (mitotic) NSCs changes in the larger rat SEZ in such a way that our analysis based on the use of a mitotic marker to quantify NSC significantly underestimates their numbers in the rat. To address this, we used an experimental paradigm in which AraC was infused on the surface of the brain to ablate the rapidly dividing transit amplifying and neuroblast populations in the SEZ. After this, NSCs are induced to re-enter the cell cycle to regenerate their daughter populations [16] and we have previously estimated that this involves the transient mitotic activation of almost the entire NSC pool [17]. The identification of mitotic NSCs at this stage, therefore, allows an accurateapproximation of the total number of NSCs residing in the SEZ. AraC was infused for 4 and 7 days (in the mouse and the rat, respectively) and animals (n = 3 per species) were killed 2 days after the end of the treatment. AraC induced an identical 9-fold increase in the numbers of mitotic NSCs in both species (Table 1, Section C), thereby confirming that the proportion of quiescent NSCs in the normal SEZ is not greater in the rat than in the mouse and that our conclusions as to the correlation between NSC numbers and ependymal cells were correct. Indeed, by calculating the number of ependymal cells in these experiments, where all NSCs are activated, we estimate that the SEZ contains 1 NSC per 187 and 157 ependymal cells in the mouse and the rat SEZ, respectively (Table 1, Section C).

If NSCs are primarily regulated by cues derived from the CSF/ependymal cell layer interface, or by direct contact with the ventricular wall, then we can make a further prediction that these cells will remain near the ependymal cells in the thicker rat SEZ. To test this, we investigated the topography of NSCs in the 2 species (Table 1, Section A). In the 20-µm-thick mouse SEZ, mitotic NSCs were found largely within 10 µm of the ventricular wall. In the rat, almost all mitotic NSCs (84.62%) were located within 20 µm of the ventricular wall even, though, as described earlier, the rat niche was of irregular width with both narrow domains of an average maximal width of 23.23 ± 2.12 µm and wide domains of an average maximal width of 55 ± 1.1 µm (the latter accounting on average for 41.14% of the total rat SEZ volume). This cell distribution revealed that, despite the finding just described that mitotic cells were distributed throughout the increased thickness of the rat SEZ, mitotic NSCs in both species were preferentially located close to ependymal cells, irrespective of the thickness of the SEZ (average distance of NSCs from the ventricular wall: 8.9 µm in the mouse and 12.2 µm in the rat; as shown in Table 1, Section A; and as illustrated in Fig. 4). Our results showing a correlation between NSC numbers and ependymal cell numbers (or surface area of the ependymal cell layer) and a close association between the cell bodies of the two cell populations even in a thickened SEZ argue that the ependymal cells might be providing short range cues that support the NSC. In this, they may be analogous to the hub cells present in the Drosophila gonadal niches where cell-cell interaction between hub and stem cell are required to maintain the latter in an undifferentiated state, in the SEZ providing signals that cannot be transmitted via the basal process [18,19]. However, our observations do not argue against conclusions from recent experimental work that the vasculature plays a critical role in supporting stem/precursor cell proliferation [4–6]. Indeed, in the proximal (NSC containing) region of the rat SEZ, we observed numerous large punctae immunopositive for laminin (Fig. 3). These are indicative of the presence of specialized basal membrane structures called fractones, originally described in the mouse SEZ as extensions of the laminin-rich basement membrane material surrounding blood vessels and contacting each stem cell [20]. More recently, these have been shown to have growth factor-binding properties that might regulate stem cell behavior [21], and their presence in the proximal to the ventricle SEZ emphasizes that both ependymal cells and vessel-associated signals likely regulates NSC behavior.

In conclusion, we have investigated how the volume of the SEZ and the distribution of NSCs within it change in a naturally occurring example of enlargement of this neurogenic niche by comparing mouse and rat brains. These 2 species are very close in terms of evolution and ecology, the latter being reinforced by the use of laboratory animals exposed to similar environments (eg, ad libido food and water intake, similar light-dark cycles), thus ensuring that the analysis is not confound by potential differences in the functional utility of adult neurogenesis. The data we report lead to 3 major conclusions. First, the size of the neurogenic niche, as defined by the volume of the area in which neurogenic activity occurs, is determined by the size of the brain, thus following the evolution-conserved principle that brain subdivisions are scaled in proportion to the scaling of the whole brain [15]. Second, however, the number of NSCs contained in the niche...
FIG. 2. Identification of mitotic NSC and precursors. The images show parts of the SEZ from coronal sections of rat brains immunostained for GFAP (in red), to mark astrocytes and PH3 (in green), to mark cells in M phase. Panel A shows 1 optical section (obtained with a confocal microscope) focusing on a mitotic (A2) astrocyte (A3) (white star), considered to be a dividing NSC. Panel B shows an optical section from brain tissue of a mouse treated with AraC, again focusing on a mitotic (B2) astrocyte (B3). Panel C shows an optical section focusing on a dividing (C2) GFAP negative (C3) cell, thus considered to be a precursor. Insets show higher magnifications of each mitotic cell, respectively, with GFAP and PH3 immunostainings as well as DAPI staining having been merged. Scale bars = 10 μm. DAPI, 4′-6-diamidino-2-phenylindole; GFAP, glial fibrillary acidic protein; NSC, neural stem cell; PH3, phosphohistone 3.

FIG. 3. Comparison of the mouse and rat SEZ. (A, B) Coronal sections taken from normal adult mouse (A) and rat (B) brains, triple-immunostained for PH3 (in green, to mark mitotic cells, indicated by white arrows), GFAP (in red, to mark astrocytes) and laminin (in yellow, marking blood vessels, indicated by white arrowheads, and fractone-like structures, indicated by yellow arrows). The images focus on the SEZ neurogenic niche, and the lateral ventricle is at the bottom. Note that in the rat the width of the SEZ is not homogeneous; an enlargement of the niche is observed at the right side. (C–G) Coronal sections taken from normal adult mouse (C, F) and rat (D, E, G) brains, immunostained for S100β (in red, marking ependymal cells). Note that S100β stains the mono-cellular layer lining the wall of the lateral ventricle, which is formed by ependymal cells. Also, note that mouse ependymal cells are flat and oriented in parallel to the ventricular wall (C and F); whereas rat ependymal cells are of a cuboid shape, both in wide (D) and narrow (E) areas of the SEZ. Some rat ependymal cells show S100β+ processes running perpendicular to the ventricular wall (white arrow in G). Color images available online at www.liebertonline.com/scd
is proportional not to its volume but to the area of the ventricular wall. The differential scaling in the size of the niche and the number of NSCs resulted in lower density of NSCs in the larger rat SEZ, a type of tissue scaling called hypermetric (ie, the size of the tissue increases faster than the number of cells) that has also been observed in the rodent cortex [22]. Third, the distribution of NSCs within the SEZ is not random, with these cells constrained in areas proximal to the ventricular wall. Together, these lead to the overall conclusion that the ependymal cell layer generates signals that have a major role in regulating adult neurogenesis and in determining the size of the NSC pool, a conclusion consistent with 2 lines of recent experimental work. First, with reports documenting the importance of ependyma-derived factors such as regulators of bone morphogenetic proteins signaling, pigment epithelium-derived growth factor, and stromal cell-derived factor 1 in the control of NSC activity [23–25]. Second, with the evidence that adult NSCs can respond to ependymal cell loss by replacing the damaged cells [26]. They also corroborate observations in the early postnatal brain, when the maturation of adult NSCs follows the generation of ependymal cells [27], as well as in the ageing brain, where neurogenic activity is reduced in concert with the disappearance of ependymal cells [28]. Therefore, the data presented here emphasize that further studies of ependymal cell-derived signals could lead to important findings in CNS regeneration and ageing.

Acknowledgments

This work was funded by the National Institutes of Health—National Institute of Biomedical Imaging and Bioengineering Quantum Grant Project [1P20EB00706]. The authors would like to thank Dr. Jing-Wei Zhao (University of Cambridge, UK) and Dr. Vera Weisbecker (University of Cambridge, UK, and University of Queensland, Australia) for many stimulating discussions and critical comments.

Author Disclosure Statement
The authors have no commercial associations or conflicts of interest to disclose.

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Received for publication March 21, 2011
Accepted after revision July 15, 2011
Prepublished on Liebert Instant Online July 15, 2011