The impact of age on number and distribution of proliferating cells in subgranular zone in adult mouse brain

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**ABSTRACT**

The mouse brain retains an ability to produce hippocampal granule neurons during the mouse’s entire lifespan. The neurons are produced in the subgranular zone (SGZ) located on the inner surface of the granule cell layer in the dentate gyrus (DG). In our study, we used a point cloud approach to characterize how the production and distribution of neural precursors for new hippocampal neurons change in the mouse brain relative to age. We found that the production of neural precursors decreases 64 fold from the age of 30 days to the age of 2.5 years. Within the SGZ the decline of cell proliferation continues during entire mouse life. We reconstructed the distribution of proliferating cells along the longitudinal axis of the SGZ and found that the highest number of proliferating cells are located approximately 0.75 mm from the dorsomedial end of the SGZ that corresponds to the most dorsal part of the DG in the mouse brain. The distribution of proliferating cells in the SGZ showed no apparent aggregations, periodicity or any other readily identifiable spatial characteristics. Proliferating cells in the SGZ tended to be located separately from other proliferating cells. About two thirds of them have no closely located other proliferating cells, and the remaining third is located in small clusters comprised of 2 or 3 proliferating cells. Based on our measurements, we calculated that from the age of 30 days to the age of 2.5 years 1.5 million neural precursors are produced in the SGZ.

**Introduction**

New hippocampal neurons are produced in the adult mouse brain. The production is strictly limited to new granule cells in the dentate gyrus (Balu and Lucki, 2009; Gonçalves et al., 2016; Ihunwo et al., 2016). Precursors for new neurons are produced in the SubGranular Zone (SGZ) located on the inner surface of the dentate gyrus granule cell layer. After production, the precursors migrate a short distance from the SGZ into the granule cell layer where they differentiate into neurons and are incorporated into existing neural circuitry in the hippocampus (Balu and Lucki, 2009; Ming and Song, 2011; Gonçalves et al., 2016).

The SGZ contains neural stem cells (NSCs). The identity of these cells is still not entirely clear, but the radial glia-like (RGL) cells are plausible candidates for this role (Seri et al., 2001). These cells resemble the radial glia cells that serve as embryonic NSCs in the developing mouse brain. The RGL cells produce neural precursors (Kriegstein and Alvarez-Buylla, 2009; Bonaguidi et al., 2011). These precursors may go through additional rounds of cell division and after that they differentiate and incorporate into the DG as new neurons or astrocytes (Bonaguidi et al., 2011; Kempermann et al., 2015). Only a small portion of progenitors complete this transition, and the majority of them are eliminated via programmed cell death (Bieble et al., 2000). New neurons show increased activity and plasticity and are thought to play an important role in hippocampal functioning (Gonçalves et al., 2016).

Hippocampal neurogenesis in mice responds to many environmental stimuli. Physical exercise stimulates neural precursor production (van Praag et al., 1999), and an enriched environment increases survival of new neurons (Kempermann et al., 1997a). In contrast, chronic stress decreases precursor production (Mirescu and Gould, 2006). These changes show that adult neurogenesis could be playing a role in the mouse adaptation to new environmental conditions. New neurons contribute to several hippocampal functions. The best studied are spatial learning and memory (Deng et al., 2010; Aimone et al., 2011). This could open a possibility to adjust these functions by altering adult neurogenesis. Since hippocampal neurogenesis is found in humans, studies in mice are relevant to human health (Ihunwo et al., 2016).

In contrast to significant progress in our understanding of molecular mechanisms of the regulation of adult neurogenesis, there is limited...
information available about the spatial organization of neural precursor production in the SGZ. Therefore, we report how neural precursor production is distributed in the SGZ and how it changes with mouse aging increase.

Experimental procedures

Ethics statement

All experiments with mice, including euthanasia, meet AVMA guidelines and were conducted following NIH and international guidelines and with veterinarian supervision. All experimental procedures were approved by The Institutional Animal Care and Use Committee (Department of Veterans Affairs, ENRM VA Hospital IACUC, Protocol SE-08-13-96).

Animals and tissue collection

C57BL/6 J male mice were purchased from The Jackson Laboratory (Bar Harbor, ME, USA) and the 2.5 year old mice were provided by the Institute of Aging (NIH) from the aging mice collection. The two mice per age group are listed in Table 1. Mice were injected intraperitoneally with 50 mg/kg 5-Ethynyl-2´-deoxyuridine (EdU) (Invitrogen, CA, USA), euthanized one hour after injection, and transcardially perfused first with 20 ml of cold PBS with 10 U/ml of heparin and then with 120 ml of cold PBS with 4% formaldehyde. Brains were extracted and incubated in PBS with 4% formaldehyde at + 4 °C for a day and then transferred into 100 mM phosphate buffer, pH 7.4, with 20% glycerol and 2% DMSO for at least two days before cutting.

Immunohistochemistry

The processing and staining of mouse brain sections were performed as described in our previous publication (Bordiuk et al., 2014). Mouse brains were cut serially at 50 μm through the entire extent of the brain using a freezing sledge microtome. Sections were collected in a 24-well plate with wells filled with tris-buffered saline (TBS, pH7.5). Transverse sectioning was used because the shape of the section is better maintained during the staining and mounting procedures which simplified the virtual reconstruction using the images of all sections of the mouse brain. Sections were first permeabilized using free-floating incubation in a TBS solution with 0.5% triton-X100 for one hour at room temperature on a rocker table with gentle agitation. Sections were then transferred directly into the EdU Alexa 647 staining solution containing EdU staining buffer, CuSO₄, Alexa Fluor 647 azide and ascobic acid in a proportion recommended by the manufacturer (Thermo Fisher Scientific, catalog# C10340) and incubated in the dark for one hour. Sections were then well washed in PBS (3 x 10 min), and all sections from each well were mounted on one gelatin coated microscope slide (25 mm x 75 mm) and air dried. Slides were coverslipped using a 5% propyl gallate/glycerol mounting media with DAPI and sealed with black nail polish.

Microscopy and image analysis

Image acquisition and analysis were performed as described previously (Bordiuk et al., 2014). Each microscope was scanned with the Zeiss AxiosImagerZ2 microscope with an EC Plan-NEOFLUAR 5X/0.16 objective. We used a 5X objective for image acquisition because its focal depth (54.84 μm at 690 nm) exceeds the thickness of brain sections (after air drying, mounted 50 μm brain sections become approximately 25–27 μm thin) and the spatial resolution (1.29 μm X 1.29 μm per pixel) is sufficient for unambiguous detection of EdU-labeled nuclei. Each scan produced a 16-bit composite image of the entire microscope slide consisting of 600 individual images. Composite images were stitched, images for individual section were cut out, arranged in the order according to their position in the brain and manually registered (Bordiuk et al., 2014). We counted the number of all EdU-labeled nuclei in all brain sections using the Find Maxima Process with the tolerance parameter set to 1500 (Fiji image processing package) and obtained coordinates for each EdU-labeled nucleus. The tolerance parameter was set to 1500 because this allowed us to automatically identify all visually detectable EdU-labeled nuclei and at the same time had a low rate of false positive identifications that we manually removed. EdU-labeled nuclei identified on each brain section were placed in one horizontal plane. The dorsoventral distance between the planes produced for two neighbouring sections was set to be equal to the section thickness, 50 μm. Thus, each reconstructed SGZ was presented as a distribution of EdU-labeled nuclei in about 80 horizontal planes which corresponds to the number of brain sections used for the reconstruction.

We used brain sections of a 120 day old mouse that was not injected with EdU as the negative control. The staining of these sections for EdU did not reveal any EdU stained nuclei and the Find Maxima Process had not identified any EdU positive nuclei on images of these sections.

To distinguish proliferating cells located in the SGZ we manually selected all EdU-labeled nuclei located within 20 μm from the inner surface of the granular cell layer. We selected and counted EdU-labeled nuclei in the SGZ on all mouse brain sections that cut across the SGZ.

Data analysis

Microsoft Excel was used for all data analysis and chart drawing. Adobe Photoshop and Microsoft PowerPoint were used for figure preparation.

To calculate the local number density for each EdU-labeled nucleus in the SGZ, we calculated the distances between each EdU-labeled nucleus and all other EdU-labeled nuclei in the SVZ using the nuclei coordinates and then counted how many of them were located closer than 200 μm to this nucleus. Such an approach is applicable to our data even if all EdU-labeled nuclei on each brain section is located in one plane. The difference is that in our case we are counting the number of EdU-labeled nuclei located in a series of cylinders stacked on top of each other rather than inside the sphere with a 200 μm radius. The height of all cylinders is equal to the section thickness, 50 μm, and the radius is equal to the radius of the circle that the sphere creates when it crosses each plane with the nuclei. The total volume of all these cylinders produced by the intersection of the sphere with the planes is only 1.5% smaller than the volume of the sphere. Thus, by using this approach we are counting the number of EdU-labeled nuclei in a volume that is practically equal to the volume of the sphere. To calculate the number of EdU-labeled nuclei inside the 50 μm sphere surrounding each nucleus, we calculated the number of EdU-labeled nuclei located closer than 52.5 μm to each nucleus. This allows us to count the number of EdU-labeled nuclei in the volume that is only 2% smaller than the volume of 50 μm sphere.

To virtually transform the distribution of proliferating cells in the SGZ and represent it as a flat surface, we established a three-dimensional tracing of the hilus apex for the left and right dentate gyrus in the brain of 60D-A mouse. Next we placed points on these tracings with 50 μm intervals and determined their coordinates. Then we found the perpendicular distance (the shortest distance) from each proliferating cell to the placed points. To visualize the distribution of proliferating cells we showed the hilus tracing as a straight line and each proliferating cell as a black dot. The position of each proliferating cell on the flattened SGZ was defined by its location on the external or internal SGZ arm and the position along the hilus tracing and the distance from the tracing.

Statistical analyses were performed using a one-way analysis of variance (ANOVA) with a post-hoc two-tailed Student's t-test with significance set at p < 0.05. The preliminary analysis shows that age has an extremely large effect on the number of proliferating cells in the SGZ with Cohen’s D routinely exceeding 10 between measurements for two
consecutive age points. In such circumstances a t-test can be reliably applied to evaluate the significance of the difference between measurements with sample size \( n = 2 \) (de Winter, 2013).

Data availability

All data generated or analysed during this study are included in this published article.

Results

We analysed the brains of 30, 60, 120, 240 day and 1 and 2.5 year old mice to find how the number and distribution of proliferating cells change in the SGZ relative to aging. For each age group we analysed two mice (Table 1). The average difference between the numbers of detected proliferating cells in the SGZ was about 2.5% showing good repeatability of our detection technique.

Table 1

<table>
<thead>
<tr>
<th>Mouse</th>
<th>Mouse age</th>
<th>SGZ</th>
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<tbody>
<tr>
<td></td>
<td>Number of EdU labeled nuclei</td>
<td>Average local number density</td>
</tr>
<tr>
<td>30D-A</td>
<td>30 days</td>
<td>4526</td>
</tr>
<tr>
<td>30D-B</td>
<td>30 days</td>
<td>4493</td>
</tr>
<tr>
<td>60D-A</td>
<td>60 days</td>
<td>2732</td>
</tr>
<tr>
<td>60D-B</td>
<td>60 days</td>
<td>2579</td>
</tr>
<tr>
<td>120D-A</td>
<td>120 days</td>
<td>1554</td>
</tr>
<tr>
<td>120D-B</td>
<td>120 days</td>
<td>1581</td>
</tr>
<tr>
<td>240D-A</td>
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</tr>
<tr>
<td>240D-B</td>
<td>240 days</td>
<td>613</td>
</tr>
<tr>
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<tr>
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<td>1 year</td>
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<tr>
<td>2.5Y-A</td>
<td>2.5 years</td>
<td>69</td>
</tr>
<tr>
<td>2.5Y-B</td>
<td>2.5 years</td>
<td>70</td>
</tr>
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</table>

We cut entire mouse brains transversely, stained all sections for EdU, and counted all cells in the SGZ with EdU stained nuclei in all sections of all analysed mouse brains. Some EdU labelled nuclei in the SGZ could be cut across in the process of brain sectioning, and consequently their fragments would be present on two consecutive sections. Counting both fragments would lead to the overcounting of the number of proliferating cells in the SGZ. To evaluate how many such cut nuclei are present on brain sections, we overlay images of consecutive sections of a 120 day old mouse (120D-A), count all EdU stained nuclei and identify among them pairs that are located in the same position on both sections (Fig. 1A). We analysed areas of the brain with a low density of proliferating cells to simplify identification of such nuclei. We counted 617 EdU labeled nuclei on 7 consecutive sections and identified only 4 pairs of nuclei that were located in same position on two consecutive sections and possibly representing two parts of one cut nucleus.

Granule cells in the DG form two symmetrical arches on the left and right side of the mouse brain in the posterior half of the telencephalon (Fig. 1B). Proliferating cells in the SGZ are located on the inner surface of the granule cell layer. To distinguish these cells we manually selected them (Bordiuk et al., 2014) and obtained their coordinates. Proliferating cells are quite abundant in the dorsal part of the SGZ in young mice. However, their number substantially decreases towards the middle portion of the SGZ and the most ventral part of the SGZ appears to be almost devoid of proliferating cells (Figs. 1B, 2). In whole, the distribution of proliferating cells in the SGZ shows no apparent aggregations, periodicity or any other readily identifiable spatial characteristics (Fig. 2).

We counted the number of EdU labelled nuclei in the SGZ and calculated their average local number density in each analysed mouse brain. The local number cell density for each EdU stained nuclei in the SGZ was calculated by counting how many other EdU stained nuclei are located closer than 200 μm to this nucleus. We detected 4500 proliferating cells with the average local number cell density of 67 in the SGZ of 30 day old mice (Table 1). These numbers decrease by 40%, from 4500 to about 2650 and from 67 to 37 correspondingly in the brain of the 60 day old mice. During the next two months the decrease becomes 20% per month, and during the following 8 months, about 12–15% per month. After that it becomes less than 10% per month. In total, the number of proliferating cells in the SGZ decreases 64 fold from 4500 to 70 and the local cell density decreases 38 fold from 67 to 1.8 (Table 1, Fig. 3).

The SGZ spans 4 mm along the dorso-ventral axis in the mouse brain. The majority of proliferating cells are located in the dorsal part of SGZ where they also have a higher cell density (Fig. 4A, D). Their number and density steadily decreases towards the ventral part of the SGZ with very few proliferating cells in the most ventral part of SGZ, even in the brain of the 30 day old mice (Fig. 4A, D). We do not find any part of the SGZ along the dorso-ventral axis where the number or density of proliferating cells is maintained with age. It appears that the number and density of proliferating cells decrease in all parts of SGZ with age (Fig. 4A, D). To confirm this observation, we divided the SGZ into four equal 1 mm thick slices along the dorso-ventral axis. The most
dorsal slice contains 77% of all proliferating cells in the SGZ in 30 day old mice. The next slices contain 13%, 8% and 2% correspondingly (Fig. 5A). The average proliferating cell density also is highest in the most dorsal slice of SGZ and decreases in the more ventral slices from 79 to 32, 20 and 9 correspondingly (Fig. 5B). Despite the significant difference in the number and density of proliferating cells in different SGZ slices, the rate of their decrease with age is very similar in all SGZ slices (Fig. 5C, D).
The SGZ has a wedge-like shape (Fig. 6A), and we can distinguish an external arm of the SGZ that underlines the external arm of the granule cell layer that faces the cortex. There is also an internal arm that underlines the internal arm of the granule cell layer that faces the mesencephalon or diencephalon (Altman and Das, 1966). The tip of the wedge can be identified as the apex of the DG hilus (Fig. 6A). We virtually transformed the distribution of proliferating cells in the SGZ of 60 day old mouse and represented it as a flat surface with traces of the hilus apex as a line that divides the flattened internal and external arms of the SGZ (Fig. 6B). The approximate shape of the internal and external SGZ arms were obtained by tracing the margins of both arms. The distribution of proliferating cells on the flattened SGZ appears to be random with no apparent empty spaces. Some aggregation of proliferating cells could be seen, but the aggregating appears in different parts of SGZ on the left and right side of the brain showing that no apparent pattern exists in their distribution. The ventrolateral end of the SGZ has a significantly lower density of proliferating cells than the dorsomedial end and the medial part (Figs. 6B, 7A). This trend can be observed in the internal and external SGZ arms (Figs. 6B, 7B, C). By smoothing the graphs showing the distribution of proliferating cells along the apex of the DG hilus (Fig. 7A), we found that the highest number of proliferating cells are located approximately 0.75 mm from the dorsomedial end of the SGZ (Fig. 7D) which corresponds to the most dorsal part of the DG.

The distribution of proliferating cells across the SGZ shows that the highest number of proliferating cells is located near the hilus apex mostly on the external SGZ arm (Fig. 7E). This peak is surrounded by shoulders with a fairly even distribution of proliferating cells that extend in both arms 0.4 mm from the hilus apex. The steady decline of the number of proliferating cells is observed beyond the shoulders (Fig. 7E). The steep decline in the number of proliferating cells can be observed at the position of the hilus apex (Fig. 7E). This decline is observed because the proliferating cells in this part of SGZ are located at some distance from the granule cell layer in contrast to other parts of SGZ where they are located immediately on the surface of the granule cell layer. The distribution across SGZ also shows that the number of proliferating cells located in the external arm is larger than that in the internal arm (Fig. 7E, F).

The local density of proliferating cells varies greatly in the SGZ of 30 day old mice from 180 to 1 (Fig. 8A). The distribution shows a steady increase in the number of proliferating cells with the local density increasing from 1 to 30. The number of proliferating cells with local cell densities from 30 to 110 remains fairly constant, and after that it steadily declines (Fig. 8A). The distribution does not show any peaks or other irregularities that could indicate the presence of two or more separate populations of proliferating cells in the SGZ. The distribution in the brains of 60 day old mice resembles the distribution in 30 day old mice, but with the highest density about 100. The distribution in older mice continues to show single-peak with the progressive decrease of the highest density with age (Fig. 8A). The highest density reaches only 5 in the 2.5 year old mice which is 36 times less than in 30 day old mice.

Proliferating cells in the SGZ could be randomly distributed, or they could form clusters where dense groups of proliferating cells are surrounded by the areas without proliferating cells. To evaluate these two possibilities, we calculated a ratio for each proliferating cell between the number of other proliferating cells located closer than 50 μm and the number of proliferating cells located closer than 200 μm. If cells are distributed randomly one would expect that in a planar structure the number of proliferating cells located closer than 50 μm be 16 fold smaller than the number of cells located closer than 200 μm. This is because the area of the circle with a 50 μm radius is 16 times smaller than the circle with a 200 μm radius. On the other hand, one could expect this ratio to be 1/64 for proliferating cells distributed randomly in the volume because the volume of the sphere with a 50 μm radius is 64 times smaller than with a 200 μm radius. Any deviation of the ratio
Fig. 4. Distribution of proliferating cells in the SGZ along the dorso-ventral, antero-posterior, and left-right axis in a mouse brain. 

(A, B, C). Distribution of proliferating cells. The number of EdU-labeled nuclei in a 50 μm brain section is shown along the vertical axis of the charts. (D, E, F). Distribution of average volume number density of proliferating cells. The average number of neighboring EdU-labeled nuclei located closer than 200 μm to each EdU-labeled nucleus in a 50 μm brain section is shown along the vertical axis of the charts. 

(A, D). Distribution along the dorso-ventral axis. The distance from the furthest dorsal part of the SGZ is shown along the horizontal axis of the charts. (B, E). Distribution along the antero-posterior axis. The distance from the furthest anterior part of the SGZ is shown along the horizontal axis of the charts. (C, F). Distribution along the left-right axis. The distance from the furthest right part of the SGZ is shown along the horizontal axis of the charts.
from 1/16 or 1/64 should indicate that proliferating cells are distributed in a non-random manner. The distribution of proliferating cells in the SGZ follows the 1/16 line as expected for randomly distributed cells in a planar structure for proliferating cells with the local number cell density from 1 to 40 (Fig. 8B). This ratio is lower for proliferating cells with the higher cell density (Fig. 8B).

The distribution of the ratio appears to be slightly above 1/16 line at cell densities between 1 and 20 (Fig. 8B). This might indicate that a fraction of the proliferating cells in the SGZ is located in small clusters. We calculated the fraction of proliferating cells that have 1, 2, 3, 4, or 5 proliferating neighbours located closer than 10 μm and found that two thirds of the proliferating cells have no close neighbours, 25% of them have one other proliferating cell located closer than 10 μm, 6% have two neighbours and 1% have three (Fig. 8C). These numbers are very close for 30, 60, 120 and 240 day old mice, somewhat lower in the 1 year old mice and even lower in the 2.5 year old (Fig. 8C).

Discussion

The labelling of proliferating cells with Bromodeoxyuridine (BrdU) or EdU (Chehrehasa et al., 2009; Zeng et al., 2010; Benjamin et al., 2017) is a commonly used technique. These thymidine analogs are able to incorporate into replicating chromosomal DNA during the S phase of the cell cycle which allows detection of all proliferating cells. The length of the S phase of the cell cycle in neural precursors does not appear to change with age (Olariu et al., 2007) thereby allowing to obtain a consistent measurement of neural precursor production using BrdU or EdU at different ages. The neural stem and progenitor cells...
account for 98% of all dividing cells in the SGZ in the mouse brain (Fig. 1 in Encinas et al., 2011) and a majority of them differentiate into new neurons (Brown et al., 2003; Snyder et al., 2009a). Therefore, in this study, we have not used any additional markers to distinguish neural progenitor cells from any other proliferating cells in the SGZ.

Cells labelled with BrdU or EdU continue to divide and produce progeny that are also labelled (Encinas et al., 2011). Therefore, for an accurate measurement of the rate of cell proliferation in the SGZ, the labelling time should be sufficiently short to prevent division of these labelled cells. We used a 1 h labelling time in this study because an extension of labelling time to 2 h leads to the appearance of divided labelled cells in the mouse brain (Bordiuk et al., 2014). This is in contrast to many publications that use either a 2 h labelling time or even a longer multiday and multi-injection labelling protocol to address cell proliferation in the SGZ (van Praag et al., 1999; Tapia-González et al., 2013) and makes it rather impossible to directly compare results obtained in these studies and our study.

We examined publications using a single injection of BrdU and found that the number of proliferating cells found in the SGZ varied greatly. Walker et al. found 2100 labelled nuclei in the SGZ of 8 week old mice after 2 h labelling (Walker et al., 2013); DeCarolis et al. found 3200 labelled nuclei in the SGZ of 9–14 week old mice after 24 h labelling (DeCarolis et al., 2014); Walter et al. found 1025 labelled nuclei in the SGZ of 3 month old mice after 2 h labelling (Walter et al., 2011); Tapia-González et al. found 734 labelled nuclei in the SGZ of 3–4 month old mice after 30 min labelling (Tapia-González et al., 2013); Sui et al. found 260 labelled nuclei in the SGZ of 16–20 week old mice after 2 h labelling (Sui et al., 2013); and Contet et al. found 2500 labelled nuclei in the SGZ of 4–6 month old mice after 2 h labelling. We list results in the order of mouse age increase and expect that with the age increase the number of proliferating cells counted in the SGZ will decrease. To our surprise the numbers increase and decrease in what appears to be a random manner. The difference could reach almost ten times in mice of similar age showing that differences in labelling and counting protocols could result in the detection of profoundly different numbers of proliferating cells in the SGZ.

One notable feature of the majority of the above-mentioned studies is the use of mice of no particular age but rather within the age interval such as 4–6 month old or 9–14 week old. It is notable because the difference between the number of proliferating cells in the youngest and the oldest mice in these groups could reach 30%. Just using mice in one experimental group comprising mostly from younger mice and in another mostly from older mice could be result in a difference of 20%–30% between these experimental groups even if experimental conditions have no effect on cells proliferation in the SGZ. On the basis of our data we could calculate the difference in cells proliferation in the SGZ that one day makes. This difference is about 2% between age of 30 and 60 days; about 1% between age of 60 and 120 days; about 0.8% between age of 120 and 240 days; about 0.6% between age of 240 days and 1 year; and about 0.35% between age of 1 and 2.5 years. Thus, mice in any experiment studying neurogenesis in the SGZ should be exactly the same age or as close as possible and in the former case the

Fig. 6. Distribution of proliferating cells in the flattened SGZ of a 60 day old mouse. (A). Schematic drawing of a coronal section of the dorsal part of DG showing the location of the external and internal SGZ arms and the apex of DG hilus. (B). Distribution of proliferating cells in the flattened SGZ of a 60 day old mouse. The line that traces the apex of DG hilus was established for the left and right DG in the 3D reconstruction of a 60 day old mouse brain, and the perpendicular (shortest) distances from this line to all EdU labelled cell nuclei in the SGZ was calculated using the line and nuclei coordinates. To generate the flattened image, we straightened the apex lines and showed EdU labelled nuclei as black dots according to the position of these nuclei along the apex lines and the distance from the apex lines. EdU labelled nuclei located in the internal and external arms of the SGZ are shown on different sides of straightened apex lines as indicated in the figure. The distance from the dorsomedial end of the SGZ is shown at the bottom. The distance from the apex lines is shown on the left.
Fig. 7. Distribution of proliferating cells along and across the flattened SGZ of a 60 day old mouse.

(A). Distribution of EdU labelled nuclei in the left and right SGZ along the straightened apex of the DG hilus line. (B). Distribution of EdU labelled nuclei in the left and right internal arm of SGZ along the straightened apex of the DG hilus line. (C). Distribution of EdU labelled nuclei in the left and right external arm of the SGZ along the straightened apex of DG hilus line. (D). Location of the highest number of proliferating cells in the SGZ is shown by arrowheads along traces of the apex of the DG hilus. Coronal view. (E). Distribution of EdU labelled nuclei across the flattened left and right SGZ. (F). Number of EdU labelled nuclei in the internal and external arm of SGZ. Error bars show standard deviation. The t-test shows that means are significantly different from each other (P < 0.05).
experimental groups should be balanced for the mouse age. In our experiment we use mice born on the same day in all experimental groups but 2.5 year group. The 2.5 year old mice were obtained from the aged mice collection of the Institute of aging (NIH) that provides only the month of mice birth.

One of our goals was to study the spatial distribution of proliferating cells in the SGZ. Consequently, we could not use stereological methods for section analysis, as stereology does not provide spatial information. It can be only used to estimate the number of proliferating cells in the SGZ. Therefore, we opted out for the use of point cloud approach (Bordiuk et al., 2014). In this approach we directly count all EdU stained nuclei in the SGZ and obtain their coordinates on all sections in each analysed mouse brain. Counting all proliferating cells in the SGZ on mouse brain sections was done before for Ki67 (Ben Abdallah et al., 2010; Amrein et al., 2015) and BrdU staining (De Carolis et al., 2014).

The major concern about such counting is a possibility that some nuclei of proliferating cells could be cut across during section preparation and their fragments will be present on two consecutive sections. Consequently, counting all stained nuclei on these sections should lead to the counting of each fragment as positive nucleus that may lead to the overestimation of proliferating cells number in the SGZ. We tested this assumption by counting EdU stained nuclei located at the same position on two consecutive brain sections. We estimated that around 10% of all EdU stained nuclei should be cut during section preparation based on the assumption that nuclei of proliferating cell in the SGZ is about 5 μm in diameter. However, we found only 4 probable pairs of EdU stained nuclei among 617 counted. Thus, the majority of cut nuclei fragments are likely lost and direct counting should lead to the undercount of proliferating cells in the SGZ. This undercount could reach 10% if during section preparation all fragments of cut nuclei are lost.

We found that the numbers of proliferating cells obtained by direct counting of all proliferating cells in the SGZ on all brain sections are consistent in mice of the same age (Fig. 3A), and despite a small sample size (n = 2), a one-way ANOVA shows that the measurements we obtained for mice at different ages are significantly heterogeneous (P = 7.61E-10). Originally, the Student’s t-test was developed using sample size n=4 (Student, 1908). Recently it was shown that the Student’s t-test could be reliably used for sample size n = 2 especially for cases with the effect size D bigger that 10 Cohen units (de Winter, 2013). The smallest effect size in our study between two experimental groups is 13.2. We also used the Tukey HSD as an alternative post hoc test and found again that all measurements are significantly different from each other (x-crit = 193.4).

The decrease of hippocampal neurogenesis in rats with age was noticed by Dr. Joseph Altman in 1965 (Altman and Das, 1965). Later this decrease was reported in many other mammalian species (Seki and Arai, 1995; Kuhn et al., 1996; Drapeau and Nora Abrous, 2008; Morgenstern et al., 2008; Lee et al., 2012; Apple et al., 2017; Mosher and Schaffer, 2017; Smith et al., 2017). Despite many reports of the decline of neurogenesis with age, there is no published detailed analysis of the age effect on neurogenesis in the SGZ in mice (Table 1 in (Drapeau and Nora Abrous, 2008)) similar to what is presented in our study. The cell proliferation decline in the SGZ with age which we are reporting in this study is similar to the reported decline of the number of BrdU labelled cells (Bondolfi et al., 2004), Ki67 (Ben Abdallah et al., 2010; Gil-Mohapel et al., 2013), PCNA (Gil-Mohapel et al., 2013) and DCX (Ben Abdallah et al., 2010) expressing cell in the SGZ. Despite the similarity in the extent of the reported decrease, the numbers of proliferating cells reported in these studies are substantially different from what we are reporting in our study. This is due to use 5-day labelling protocol in the study with BrdU labelling (Bondolfi et al., 2004) and the difference in the length of S phase of the cell cycle and periods when Ki67, PCNA, and DCX are expressed in proliferating neural progenitors.

The decline of cell proliferation in the SGZ continues during the entire mouse life. We have not noticed periods in the mouse life when cell proliferation continues at a steady rate (Fig. 3). The continued decrease is observed in areas with high and low density of proliferating cells (Figs. 4 and 5). A visualisation of the distribution of proliferating cells (Fig. 2) does not reveal any local areas in the SGZ where cell proliferation is maintained for a considerable period of time at the similar level. The steady state cell production is the hallmark of cell renewal in skin, intestinal epithelia and blood. Such stable and continued cell production balances the continued and stable loss of worn-out cells.
in these tissues and maintains them in a homeostasis. The continued decrease of cells proliferation in the SGZ shows that the processes these new cells are intended to support do not require the steady supply of new neural progenitors or that this process is not homeostatic.

EdU is incorporated into chromosomal DNA only during the S phase of the cell cycle. The length of the S phase in neural precursors in the SGZ is approximately 7.6 h (Hayes and Nowakowski, 2002). Based on this number and our data about cell proliferation in the SGZ, we can calculate that from the age of 30 days to the age of 2.5 years 1.51 million new cells are produced in the SGZ if we assume that the number of proliferating cells in the SGZ decreases exponentially between measured points. The number becomes 1.60 million if we assume that the decrease is linear. Cell production changes considerably during mouse life. Thus, about 350,000 new cells are produced in the SGZ during the 30 days between ages of 30 and 60 days. In contrast, it takes 1.5 years to produce the same number of new cells toward the end of mouse life between ages of 260 days and 2.5 years. The number of granule cells in mouse hippocampus is about 500,000 (Kempermann et al., 1997a; Farrar et al., 2005; Ben Abdallah et al., 2010; Tapia-González et al., 2013). Given this, all granule cells could be replaced by new cells three times during the lifespan of the mouse. However, only a small fraction of granule cells are replaced because only some of the neural precursors differentiate into new neurons, and the majority of new neurons undergo programmed cell death (PCD) during processes of differentiation and integration into the granule cell layer (Bieble et al., 2000). The rate of cell proliferation in the SGZ in C27BL/6J mice appears to be almost twice as high as in other commonly used mouse strains (Kempermann et al., 1997b). This rate is also sensitive to housing conditions and access to exercise equipment (Kempermann et al., 1997a; van Praag et al., 1999). Therefore, neural precursor production in other mouse strains or mice hosted in different conditions could be different than reported in our study.

Our estimate of the production of new cells in the SGZ is vastly larger than the estimate presented in the recent publication by Pillez et al. (2018). Using chronic live imaging of neurogenesis in the SGZ in two month old mice, the authors estimated that only 45,000 new cells are produced in the SGZ during the entire mouse lifespan. For comparison, this number of new cells represents only 3% of our estimate and such a number of new cells in the SGZ should be produced in less than six days in two month old mice according to our data. The simplest explanation for this apparent discrepancy could be that the removal of the neocortex above the hippocampus, required for the direct neurogenesis observation, causes brain trauma which suppresses proliferation of RGL cells in the SGZ. Another plausible explanation is that the RGL cell labelling method relying on the activity of Achaete-scute homolog 1 (Ascl1) promoter selectively labels RGL cells with low proliferative potential. In this case we should expect that other RGL cells, that are not labelled by this method, have high proliferative potential and are responsible for the production of 97% new cells in the SGZ. We might also expect that 97% of new cells in the SGZ are produced by non-RGL cells or that such cells produce new RGL cells that, in turn, produce new neuronal precursors in the SGZ. All these possibilities need to be tested experimentally. The utility of our estimate is that we provide the total number of new cells produced in the SGZ. Thus, the comparison of this number with the number of new cells produced by some particular RGL or other cells in the SGZ, as we show above, will immediately reveal the relative contribution of such cells in neurogenesis in the SGZ.

Several experimental approaches have previously been used to study neurogenesis in different parts of the SGZ. One of them is to divide the SGZ along the anteroposterior axis of the mouse brain on the rostral SGZ and caudal SGZ (Olariu et al., 2007; Kanatsou et al., 2017). Another is to divide SGZ along dorsoventral axis (Banasr et al., 2006; Snyder et al., 2009b; Jinno, 2011; Nollet et al., 2012; Tanti et al., 2012); to study neurogenesis in the external and internal SGZ arms (Fig. 6A) (Kempermann et al., 2003; Olariu et al., 2007; Snyder et al., 2009b; Jinno, 2011; Kanatsou et al., 2017); or to use isolated and straightened hippocampi and divide it longitudinally (Wiget et al., 2017). In our study, we obtained coordinates of all proliferating cells in the SGZ and represented them as a point cloud. Using this approach it is possible to visualize the distribution of proliferating cells in the SGZ at different mouse ages (Fig. 2), show the distribution of the number and density of proliferating cells along dorso-ventral and other axis of the SGZ (Fig. 4), evaluate the randomness of distribution and clustering of proliferating cells in the SGZ (Fig. 8), and flatten the SGZ to evaluate distribution of proliferating cells along the longitudinal axis and across of the SGZ (Figs. 6 and 7). All these used in our study analyses allow a detailed characterization of proliferating cell distribution in the SGZ to better understand how their distribution changes with age. In contrast, the above mentioned approaches, with the exception of Wiget et al. study (Wiget et al., 2017), provide only a number of proliferating cells in some particular part(s) of the SGZ.

The distribution of proliferating cells shows no periodicity, expansive empty spaces or abrupt changes along the SGZ (Figs. 2, 4–7). We found that proliferating cells are distributed throughout the entire SGZ at all analysed ages (Figs. 2, 4 and 5). This finding shows that the zone of proliferation in the SGZ does not shrink with increased age, but rather the rate of cell proliferation decreases in all parts of the SGZ (Fig. 5). The ratio between the number of proliferating cells located closer than 200 μm to the number located closer than 50 μm clearly shows that proliferating cells in the SGZ have a planar distribution (Fig. 8B). We show that this ratio becomes smaller than 1/16 for proliferating cells with high local density (Fig. 8B). Proliferating cells in the SGZ areas with a high density are located not only on the inner surface of the granule cell layer but also small distances into the hilus; which makes their distribution more like a three-dimensional distribution.

The dorsal hippocampus appears to be mostly involved in cognitive functions such as spatial learning and memory and the ventral hippocampus in emotional reactions. And, correspondingly it was speculated that neurogenesis in the dorsal part of the SGZ affects cognitive functions and in the lateral part emotional functions such as stress or depression (reviewed in Moser and Moser, 1998; Sahay and Hen, 2007; Fanselow and Dong, 2010; O’Leary and Cryan, 2014; Wu et al., 2015). The majority of proliferating cells are located in the dorsal part of the SGZ (Fig. 4A, 5A) mostly due to a higher density in the dorsal part of SGZ (Figs. 4B, 5B). This trend continues during the mouse lifespan (Figs. 4A, D, 5). We have not found any evidence of an abrupt transition in the distribution of proliferating cells between the dorsal and ventral parts of the SGZ (Figs. 1, 2, 6B and 7A). This could be explained by a possibility that functional difference of the dorsal and ventral DG is not reflected in the distribution of proliferating cells in the SGZ or that the boundary between dorsal and ventral parts of the DG is not abrupt. However, it is worthwhile to note that there is a profound difference in the number of proliferating cells between the dorsal and ventral parts of the SGZ. We found that the number of proliferating cells in the most
dorsal quarter of the SGZ is about 70-times higher than in the most ventral quarter (Fig. 5A).

We obtained the longitudinal distribution of proliferating cells in the SGZ by virtual transformation (Fig. 6B). This distribution looks very similar to the distribution observed in the straightened hippocampi (Fig.1A in (Wiget et al., 2017)) thereby showing validity of our approach. At the same time, our results, obtained by different experimental approach, confirm the finding by Wiget et al., (Wiget et al., 2017) that in C57BL/6 mice the highest rate of cell proliferation occurs about one third from the septal end of the SGZ that corresponds to the most dorsal part of the SGZ (Fig. 7D). The distribution of proliferating cells in DBA2/Crl house and wood mice and bank voles are very different from C57BL/6 mice. The cell proliferation in these species/mouse strains are shifted toward the temporal part of the SGZ (Wiget et al., 2017) showing that even in closely related species/mouse strains the distribution of proliferating cells in the SGZ could be remarkably different. This difference shows that the higher rate of neural precursor production in the dorsal SGZ in C57BL/6 mice might not represent a typical distribution of neurogenesis in the SGZ. It also tempers hypotheses suggesting that neurogenesis plays a more significant role in cognitive/spatial functions of the hippocampus compared to emotional reactions.

The presence of neurogenesis in the SGZ in the adult human brain is still under discussion. Even in very recent publications, one may find data showing that neurogenesis stops completely at the age of 3 years (Dennis et al., 2016; Cipriani et al., 2018; Sorrell et al., 2018), and data showing that neurogenesis stops completely at the age of 3 years (Dennis et al., 2016; Cipriani et al., 2018; Sorrell et al., 2018), and at the same time that neurogenesis is present at substantial levels even at a very old age (Spalding et al., 2013; Boldrini et al., 2018). One notable feature of adult neurogenesis studies in the human brain is that the human hippocampus is treated as a uniform structure. However, if one decides to study adult neurogenesis in the extreme ventral portion of the mouse hippocampus, there will be no detectable adult neurogenesis even in very young mice. Thus, in order to establish if a similar difference in the rate of neurogenesis between different parts of the human hippocampus also exists, the human SGZ needs to be analysed in a manner similar to our analysis of the mouse SGZ.

Conflict of interest

The authors declare no conflict of interest.

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