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Highlights
- Human microglia renew at a median rate of 28% per year
- Microglial cells are on average 4.2 years old
- Most of the microglia population (>96%) is renewed throughout life

In Brief
Taking advantage of the decreasing level of atmospheric $^{14}$C since the Cold War, Réu et al. show that human microglia, unlike most other hematopoietic lineages, slowly turn over at a yearly median rate of 28%. The absence of a large quiescent subpopulation indicates that most microglia will renew throughout life.

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The Lifespan and Turnover of Microglia in the Human Brain

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SUMMARY

The hematopoietic system seeds the CNS with microglial progenitor cells during the fetal period, but the subsequent cell generation dynamics and maintenance of this population have been poorly understood. We report that microglia, unlike most other hematopoietic lineages, renew slowly at a median rate of 28% per year, and some microglia last for more than two decades. Furthermore, we find no evidence for the existence of a substantial population of quiescent long-lived cells, meaning that the microglia population in the human brain is sustained by continuous slow turnover throughout adult life.

INTRODUCTION

Microglia are the resident macrophages of the CNS, which dynamically survey their surrounding for signs of infection or cell distress (Casano and Perl, 2015). In mice, microglial progenitor cells derive from an early myeloid branch of the hematopoietic lineage in the embryonic yolk sac and enter the CNS before the blood-brain barrier is formed (Ginhoux et al., 2010; Schulz et al., 2012). There is no further contribution from the peripheral hematopoietic system under physiological conditions, and this is a self-sustaining population within the CNS (Ajami et al., 2007; Askew et al., 2017; Bruttger et al., 2015; Hoeffel et al., 2015; Mildner et al., 2007). The self-contained nature of this population makes it vulnerable to local disturbances. Additionally, as the blood-brain barrier is formed, microglial cell numbers are stably maintained, because having reduced numbers results in behavioral and learning deficits (Parkhurst et al., 2013).

Based on [3H]thymidine incorporation, heavy water (D2O) labeling, and 5-ethyl-2′-deoxyuridine (EdU) and 5-bromo-2′-deoxyuridine (BrdU) incorporation, 0.075%–1.04% of microglia in adult mice of different strains and 2.35% of microglia in the young adult macaque were estimated to enter the cell cycle each day (Askew et al., 2017; Lawson et al., 1992; Shankaran et al., 2007; Tay et al., 2017; Tonchev et al., 2003). It is difficult to infer cell turnover dynamics in humans from data in experimental animals, which may have very different requirements or lifespans. Assessing the turnover of human immune cells is notably important because it is particularly hard to deduce renewal rates from laboratory animals, which live in pathogen-free barrier facilities (Beura et al., 2016). A recent study demonstrated that 2% of microglia in the adult human brain are in cell cycle at any given time based on Ki-67 labeling, but the authors noted the limitations of this approach and acknowledge the need for more precise measurements (Askew et al., 2017). It is difficult to estimate cell turnover dynamics based on cell cycle markers, because it rests on assumptions of cell cycle length. Moreover, it is not possible to know whether the cell will proceed through the cell cycle to mitosis or whether the potential progeny will survive.

RESULTS

We analyzed the frontal and occipital cortices from two subjects (17 and 41 years old) who had received IdU (5-Iodo-2′-deoxyuridine) as a radiosensitizer for cancer treatment (Table S1). On average, 0.8% of Iba1+ parenchymal microglia in the cortex were IdU+ after 4 days (donor 1) or 10 days (donor 2) of IdU administration (Figures 1A and 1B; Table S1). Accounting for the labeling period, it averages at 0.14% labeling per day (Figure 1C). These observations do, however, come with the following caveats: (1) the sample size is small and additional samples are not available; and (2) the subjects studied are not healthy individuals, and thus may exhibit aberrant turnover of different populations of cells. Still, our observations provide us with a general estimate of what to expect in our downstream analysis based on retrospective 14C measurements.

Following CD11b magnetic bead selection, we isolated by fluorescence-activated cell sorting (FACS) CD45+CD11b+ microglia from the adult human postmortem cerebral cortex in order to perform retrospective 14C dating (Figures 2A–2C) (Olah et al., 2012). We used CD20, a B cell marker, to evaluate the presence of blood-borne cells in dissociated cortical preparations. The percentage of B cells in brain samples prior to MACS purification (Figure 2D) is very low relative to the circulation...
often cells need to divide within an individual of a given age in or-
2010). In simple terms, this is accomplished by calculating how
shots of cell dynamics for each individual donor (Bernard et al.,
is replaced at a fixed rate, it is possible to obtain robust snap-
ure 3C). Using a homogeneous turnover model in which each cell
it is possible to calculate the average cell age of the sample (Fig-
Based on the year of collection and date of birth of the sample,
we performed retrospective birth dating of microglia isolated
population (Spalding et al., 2005)( Figure 3A). Using this strategy,
we generated cell age distributions stochastically, showing that
have no evidence to support the existence of a significant subpopulation of quiescent very long-lived cells (Figure 3D). We have employed a conservative mathematical model, with few assumptions, which does not preclude that an extended dataset or other variables could explain the heteroge-
Cells that became postmitotic soon after birth have a unique
14C signature, corresponding to the atmospheric level at that
moment in time. This allowed us to directly investigate the
possible existence of a non-dividing subpopulation, using a het-
erogeneous turnover model (Figure 3D). The model that best fits
our data are one where the majority of the population (>96%)
is renewed and we have no evidence to support the existence of
a significant subpopulation of quiescent very long-lived cells
(Figure 3D). We have employed a conservative mathematical model, with few assumptions, which does not preclude that an extended dataset or other variables could explain the heteroge-
Based on the average cell age and cell division rate observed,
we generated cell age distributions stochastically, showing that
within an individual there is a wide range of microglia ages (Fig-
3E). Simply because not all cells are dividing at the same
time, some are quite young due to recent cell division and others
can be more than 20 years old (Figure 3E).

The lower rate of microglia renewal compared with most other
immune cells is probably a manifestation of the immune-privi-
eged status of the CNS (Figure 3F) (Busch et al., 2007; Macallan
et al., 2005). On the other hand, in comparison with other cells of

Figure 1. IdU Incorporation
(A) Confocal image with orthogonal projections, from human cortex, revealing microglia positive for Iba1.
(B) Co-staining of Iba1 and the thymidine analog IdU.
(C) Percentage of microglia incorporating IdU per day (mean ± SD). Each data point represents a glass slide. Nuclei are labeled with antibodies to histone H3. Scale bars, 10 μm.
the CNS, microglia show a high exchange rate (Figure 3F) (Spalding et al., 2005, 2013; Yeung et al., 2014). Thus, a constant basal renewal is likely necessary for the maintenance of a cohort of young and healthy microglial cells.

**DISCUSSION**

Administering nucleotide analogs for a short time period introduces a bias to label the cells with the highest proliferation rate within a potentially heterogeneous population. Also, labeled cells that continue to proliferate after the labeling period will give rise to additional positively labeled cells that lead to overestimations of cell proliferation (Neese et al., 2002). Nevertheless, samples of human brain labeled with nucleotide analogs are very valuable, not only due to their rarity but also as a confirmatory tool of in vivo renewal in humans (Eriksson et al., 1998; Ernst et al., 2014; Yeung et al., 2014).

Analysis of the integration of atmospheric $^{14}$C, derived from nuclear bomb tests, in genomic DNA is cumulative and gives a more comprehensive view of cell age and cell division history (Spalding et al., 2005), and it is possible to analyze tissue from subjects without previous serious illness. There is little exchange of carbon atoms in genomic DNA in non-dividing cells, and the effect of DNA repair and methylation is well below the detection limit of this retrospective birth dating strategy (Bergmann et al., 2012; Ernst et al., 2014; Spalding et al., 2005).

Most immune cells do not live longer than a few days or weeks (Busch et al., 2007; Macallan et al., 2005), making microglia one of the slowest dividing immune cells described to date. An extreme exception is plasma cells in the intestine, where the subset with the slowest renewal rate has a median age of 22 years (Landsvir et al., 2017). The turnover rate of microglia in humans is substantially lower than in mice. There is a much higher exchange of oligodendrocytes in mice compared with humans (Yeung et al., 2014), and it is possible that clearance of myelin and cell debris calls for a higher exchange rate of microglia in mice.

Based on Ki-67 staining, a recent study estimated that, at any given moment, 2% of microglia are proliferating in the
human brain (Askew et al., 2017). Similarly to short-term labeling experiments using BrdU or IdU, Ki-67 staining detects dividing cells, but it provides limited information for cells that may exhibit slow renewal rates. Importantly, Ki-67 does not measure proliferation directly, and cells blocked in G1 or destined to enter apoptosis can accumulate in the population as Ki-67+ events because of a G1/S block (Busch et al., 2007). An additional explanation for the results of the different techniques resides in the fact that newborn microglia are more likely to die than the resident microglia (Askew et al., 2017). Hence, many of the cells detected by Ki-67 and, to a lesser extent, by IdU may be destined to die and not to replace existing ones.

In conclusion, 14C analyses reveal that microglia as a whole turn over slowly, and that individual cells can potentially be decades old. Complementary, IdU and Ki-67 (Askew et al., 2017) measurements suggest heterogeneity within the population possibly induced by the presence of a subpopulation of fast dividing cells. Finally, our data predict a nearly complete renewal of the microglia population in the human cortex during the lifespan of an individual, similar to what has been seen in mice (Askew et al., 2017).

**EXPERIMENTAL PROCEDURES**

**Tissue Collection**

Neocortical tissue was obtained from donors admitted for autopsy at the Department of Forensic Medicine in Stockholm from 2014 to 2016, after informed consent from the relatives. The ethical permit for this study was granted by the Regional Ethics Committee of Stockholm (2010/313-31/3). Formalin-fixed and paraffin-embedded sections of cortical frontal lobe and occipital lobe from cancer patients, who had received IdU for therapeutic purposes, were obtained from the National Heart, Lung and Blood Institute, NIH. Buffy coats were obtained from anonymous regular blood donors at Blodcentralen, Karolinska University Hospital.

**IdU Quantification**

The sections were immersed in xylene to remove the paraffin, and the tissue was rehydrated in descending ethanol series. Triton X-100 (0.2%) was used to permeabilize the tissue, and antigen retrieval was performed in 0.05% citraconic acid solution (pH 7.4) for 20 min in a domestic steamer. The sections were left for 20 min at room temperature and then immersed in 2.0 N HCl for 40 min. The slides were blocked (10% donkey normal serum in PBS with 0.2% Triton X-100) at room temperature for 1 hr. Following incubation with the primary antibodies (1:60 mouse anti-BrdU, BD347580; 1:100 goat anti-Histone H3, Abcam 12079; 1:100 rabbit anti-Iba1, Wako 019-19741), the sections were incubated with the secondary antibodies (1:200 donkey anti-mouse Cy3, Jackson ImmunoResearch 715-165-150; 1:200 donkey anti-goat A647, 

**Figure 3. Microglial Population Dynamics**

(A) Schematic illustration of the 14C atmospheric curve over time (Levin et al., 2010). The concentration of 14C in the genomic DNA of a cell population is dependent on the atmospheric 14C concentration (y axis). Thus, the birth date of the cell population can be read off the x axis.

(B) 14C content in the genomic DNA of human cortical microglia from donors born across six decades. Data points plotted along the x axis according to the date of birth of the donors. Close-up view of four nearly overlapping data points (gray square). The error bars represent the 14C concentration measurement error.

(C) Representation of the average age of microglia in each individual and linear regression (black line).

(D) Different models for distribution of data on the atmospheric 14C curve considering different frequencies of dividing cells. The model that best fits the data are where most cells (>96%) renew.

(E) Considering the turnover rate, the average cell age, and the fact that all microglial cells do not divide simultaneously, we created a stochastic cell age distribution model. Our model shows that within an individual there is a distribution of cells of different ages, with some having recently renewed and others not having divided in more than 20 years (donors with infinite turnover not included).

(F) The approximate rate of microglia turnover is 0.08% a day, a low turnover rate in comparison with other immune cells (granulocytes, monocytes, and naive B cells) but a high turnover rate relative to other CNS cells (neurons in the dentate gyrus, oligodendrocytes, and cortical neurons).
of Proteinase K (40 mg/ml) were added to the sorted cells and incubated at 12 mL glass vial. Ethanol 95% (6 mL) was added and the glass vial was manually 13,000 rpm for 6 min. The supernatant containing the DNA was transferred to the sample; then it was vortexed for 30 s. The solution was spun down at 2,000 rpm for 2 hr. To convert the samples into graphite, DNA samples suspended in 0.5 mL of water were lyophilized to dryness in a clean room (ISO8). The extraction protocol was modified from Miller et al. 2013a, 2013b, 2015. The measurement error was determined for each sample and ranged between ±8% and 40% (2 SD) Δ14C for the large (>100 mg C) and small samples (10 μg C), respectively. All 14C data are reported as Fraction Modern F 14C as defined in Reimer et al. (2004) or Δ 14C as defined in Stuiver and Polach (1977). All accelerator mass spectrometry (AMS) analyses were performed blind to the identity of the sample.

Tissue Dissociation
After careful removal of the meninges and all visible blood vessels, the tissue was cut into small pieces and thoroughly rinsed with PBS. The tissue was then homogenized in media A (1x HBSS, 150 mM HEPES, 2 mM EDTA, 5% BSA) with 2 U/mL papain ( Worthington) and 10 U/mL DNaseI (Roche) at 37°C for 1.5 hr. The homogenized tissue was mix with 3 volumes of sucrose media I (PBS, 0.7 M sucrose, 2 mM EDTA) and centrifuged for 20 min at 1,000 × g. The pellet was then resuspended in sucrose media II (PBS, 0.9 M sucrose, 2 mM EDTA) and centrifuged for 25 min at 800 × g. Finally the pellet was resuspended in blocking solution (PBS, 0.1% FBS, 2 mM EDTA) and filtered through a 40 μm cell strainer.

Cell Isolation
The cell suspension was incubated for 5 min with human Fc-gamma receptor (FcR)-binding inhibitor (1:100; eBioscience) and for 30 min with CD11b antibody-conjugated microbeads (1:25, 130-093-634; Miltenyi Biotec). The magnetic isolation was performed according to the manufacturer. The samples were next incubated for 20 min with PE-CD11b (1:20, clone ICRF44, BioLegend) and Alexa 647 CD45 (1:20, clone H30; BioLegend), and finally FACS sorted in an Influx flow cytometer (BD Biosciences). Blood cells were isolated from buffy coats by density gradient lymphoprep). Peripheral blood mononuclear cells (PBMCs) were positively selected with CD11b antibody-conjugated microbeads (1:25, 130-093-634; Miltenyi Biotec) according to the manufacturer.

DNA Isolation
In order to prevent carbon contaminations, we performed the DNA isolation in a clean room (ISO8). The extraction protocol was modified from Miller et al. (1988). The glassware was prebaked for 4 hr at 450°C. 1 mL of lysis buffer (100 mM Tris [pH 8.0], 200 mM NaCl, 1% SDS, and 5 mM EDTA) and 12 μL of Proteinase K (40 mg/ml) were added to the sorted cells and incubated at 65°C overnight. The samples were further incubated at 65°C for 1 hr after 6 μL of RNase cocktail (Ambion) was added. 600 μL of NaClI (5 M) was added to the sample; then it was vortexed for 30 s. The solution was spun down at 13,000 rpm for 6 min. The supernatant containing the DNA was transferred to a 12 mL glass vial. Ethanol 95% (6 mL) was added and the glass vial was manually agitation. The DNA precipitate was washed three times in DNA washing buffer (70% ethanol [v/v] and 0.5 M NaCl), dried at 65°C overnight, and resuspended in 0.5 mL DNase/RNase free water (GIBCO/Invitrogen). The DNA purity and concentration were verified by UV spectroscopy (NanoDrop).

Accelerator Mass Spectrometry
DNA samples suspended in 0.5 mL of water were lyophilized to dryness in a vacuum centrifuged at 2,000 rpm for 2 hr. To convert the samples into graphite, we added excess CuO to each dry sample, and the quartz tubes were evacuated and sealed with a high temperature torch. The tubes were placed in a furnace set at 900°C for 3 hr to combust all carbon to CO2. The gas was then purified by freezing the residual water at ~60°C, as well as cryogenically trapping the CO2 at ~196°C and discarding all the other gases. The CO2 was chemically reduced to graphite in the presence of zinc powder and iron catalyst in individual miniaturized reactors at 550°C for 6 hr. Thorugh laboratory protocols are exercised to minimize the introduction of stray carbon into the sample (Salehpour et al., 2013a). Graphite targets were pressed into individual cathodes and are measured at the Department of Physics and Astronomy, Ion Physics, Uppsala University (Salehpour et al., 2013a, 2013b, 2015) using the 5 MV Pelletron Tandem accelerator. Large CO2 samples (~100 μg) may be split, and Δ14C are measured by stable isotope ratio mass spectrometry, which established the Δ14C correction to ~24.1% ± 0.5% (1 SD), which was applied to the samples. Corrections for background carbon introduced during sample preparation were made as described previously (Salehpour et al., 2013a, 2013b, 2015). We are grateful to Marcelo Toro and Sarantis Giatrellis for help with flow cytometry, the staff at the Swedish National Board of Forensic Medicine for procuring tissue, and Karl Håkansson and Peter Senneryd for AMS sample preparation. This study was supported by grants from the Swedish Research Council, the Swedish Cancer Society, the Karolinska Institute, Tobias Stiftelsen, the ERG, Knut och Alice Wallenberg’s Stiftelse, and Torsten Söderbergs Stiftelse. P.R. was supported by the Portuguese Foundation for Science and Technology (grant SFRH/BD/33465/2008).

SUPPLEMENTAL INFORMATION
Supplemental Information includes Supplemental Experimental Procedures and two tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2017.07.004.

AUTHOR CONTRIBUTIONS
P.R. and J.F. designed the study. P.R. performed most of the experiments. A.K. performed the IdU analyses. S.B. did all of the mathematical analyses. J.E.M. performed additional experiments. M.S. and G.P. did the AMS measurements. K.A. and H.D. collected and classified the samples for 14C. S.P. and J.T. collected and classified the IdU samples. P.R., J.E.M., and J.F. wrote the manuscript.

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