INTRODUCTION

Adult neurogenesis, the process whereby new neurons are born, survive and integrate into adult neural circuits, offers the potential for plasticity and repair in the mature brain. It is a prominent feature of the olfactory bulb (OB), olfactory epithelium, and dentate gyrus, however, the regulation of this process and its functional significance to the organism remains unclear.

Adult born neurons destined for the OB originate from a population of astrocyte stem cells lining a portion of the subventricular zone (SVZ) and rostral migratory stream (RMS) (Alvarez-Buylla and Garcia-Verdugo, 2002; Bonfanti and Peretto, 2007). Neuroblasts migrate from their germinal niche through the RMS into the OB where a most of them migrate toward lamina of the granule cell layer (GCL) to become GCs (Carleton et al., 2003), and a minority migrate towards the glomerular layer to become peri-glomerular interneurons (Belluzzi et al., 2003). Recent work on the maturation of adult born GCs indicates that they first form basal synaptic connections with centrifugal fibers (Whitman and Greer, 2007; Kelsch et al., 2008), followed by formation of a dendro-dendritic synapse with mitral cells (MC) (Carleton et al., 2003). Once fully integrated into the circuit, GCs integrate olfactory input from the periphery with diverse inputs from higher brain areas (Shepherd, 1998), and transmit this activity towards MC through dendro-dendritic GABA release (Jahr and Nicoll, 1980).

Theories regarding the computational function of OB adult neurogenesis often posit that by providing a continuously refreshed population of synaptic connections, young neurons may enable the system to better adapt to new olfactory environments (Cecchi et al., 2001; Lledo et al., 2006). There are three lines of evidence in support of this. First, the rate of neural proliferation in the SVZ appears to be sensitive to conditions with unique olfactory demands such as pregnancy (Shingo et al., 2003), and pheromone exposure (Mak et al., 2007). Next, between 2 and 3 weeks after genesis, adult-born neurons undergo massive cell death that is negatively correlated with olfactory activity (Petreanu and Alvarez-Buylla, 2002; Rochefort et al., 2002; Saghatelayan et al., 2005; Yamaguchi and Mori, 2005) and is sensitive to afferent activity from higher brain regions (Alonso et al., 2006; Mouret et al., 2008). This may reflect a mechanism that selects which young neurons survive, and thus which new connectivity patterns are added to the circuit. Finally, adult born GCs in the OB are functionally distinct from the GCs born during development (Lemasson et al., 2005), and young cells at any age express unique plasticity in their synapses with centrifugal fibers (Gao and Strowbridge, 2009; Nissant et al., 2009). Together, these data suggest that adult neurogenesis is not merely a deregulated form of post-natal development, but likely serves a critical function.

Recent work has shown that after ablation of SVZ adult neurogenesis and subsequent cell loss in the GCL, there are no behavioral deficits in olfactory discrimination, odor detection thresholds, or in the retention short-term olfactory memory (Imayoshi et al., 2008; Lazarini et al., 2009). However, there are conflicting reports regarding the importance of adult neurogenesis to long-term olfactory memory (Imayoshi et al., 2008; Lazarini et al., 2009). In general, the functional impact of adult neurogenesis in the OB is still unclear.

In this study we find that a chronic loss of adult neurogenesis impacts both OB structure and olfactory behavior, but not OB physiology. We ablated SVZ neurogenesis and characterized a consequent gradual reduction in the size of the GCL. We also...
observed that the survival of young GCs is insensitive to the rate of neurogenesis, suggesting a general decoupling between neuronal survival in the OB and neurogenesis in the SVZ. Functionally, we observed a behavioral disruption in olfactory-cued fear conditioning, but surprisingly this behavioral phenotype did not correlate with any alteration in GCL physiology. This suggests that adult neurogenesis functions in a behavioral system extending beyond the OB.

**MATERIALS AND METHODS**

**ANIMALS**
All mice were C57/BL6J purchased from the Jackson Laboratories, and housed at Columbia University in accordance with institutional I.A.C.U.C standards for animal care. Animals were group housed, and left to recover between 3 days and 2 weeks after arrival, before beginning the x-ray irradiation. All animals were between 8 and 12 weeks old at the time of irradiation. Therefore, all mice in this study were between 3 and 8 months old at the time of testing. For behavioral testing, mice were male C57Bl/6J irradiated at 6–8 weeks of age and were naïve to behavioral testing prior to fear conditioning. Animals were handled for approximately 1 min once daily for 2 days prior to experimentation to minimize handling induced stress during the experiment. Mice were housed in groups of five and sustained on a 12-h light/dark cycle with ad lib access to food and water.

**X-RAY IRRADIATION**
The X-ray source for all experiments was a Siemens Stabilopan operated at 12 mA and 300 kVp with a 0.2-µm copper filter to harden the beam, and emitted 30 cm above the head of the mouse. The ionization dosage to water was measured by the Department of Radiation Oncology Physics, and the corrected real dosage was calculated previously to be approximately 1.8 Gy per minute (Santarelli et al., 2003). Therefore, a full 15 Gy dosage consisted of a 2:40 min exposure given three times within a week. The mouse was anesthetized with an intraperitoneal injection of ketamine (100 mg/kg) and xylazine (7 mg/kg), and mounted in a stereotaxic frame (Kopf, Tujunga, CA, USA). A lead box was placed over the mouse, and aligned to the ear bars of the stereotaxic frame. One-half inch thick lead shielding covered the entire length of the mouse, except for a window 5 mm (rostral-caudal) by 13 mm (medial-lateral) that could be moved along the rostral-caudal axis of the animal. This sliding window could be positioned relative to the ear bars of the stereotaxic frame. For sham irradiated animals, cagemates were anesthetized at the same time and locations as their irradiated counterparts.

**HISTOLOGY**
Mice were transcardially perfused with 4% paraformaldehyde in PBS, and their dissected brain was post-fixed in 4% paraformaldehyde for an additional 24 h at 4°C. Brains were cryoprotected in 40% sucrose solution, and frozen in a cutting mold in OCT. Twenty micrometer thick sections were made using a freezing cryostat and then were mounted directly to glass slides (Fischer Superfrost Plus, Pittsburgh, PA, USA).

**IMMUNOHISTOCHEMISTRY**
Cryostat frozen section cut at 20-µm thickness were treated for antigen retrieval by incubating in 0.01 M Na-citrate for 10 min in a conventional kitchen steamer. The following primary antibodies were used: mouse-anti-BrdU (1:100, BD Biosciences, San Josa, CA, USA), rat-anti-BrdU (1:1000, Accurate Chemical, Westbury, NY, USA), donkey anti-doublecortin (1:1000, Dcx C-18, Santa Cruz Biotechnology, Santa Cruz, CA, USA). Secondary antibodies raised in donkey were used at 1:750 (Alexa 488 and Alexa 594, Invitrogen, Eugene, OR, USA), or biotinylated secondary antibodies were used (donkey anti-rat and biotinylated donkey anti-goat, 1:1000, Jackson ImmunoResearch, West Grove, PA, USA) followed by streptavadin conjugated Cy-3 (1:1000, Zymed, San Franciscso, CA, USA). Counterstain was performed using DAPI or Hoescht dyes.

**CELL COUNTING**
BrdU counts of the SVZ were made from coronal sections and involved complete counting of all cells along the lateral wall of the SVZ. Sections of the SVZ were cut every 120 µm with the rostralmost slice aligned to the rostral-tip of the SVZ. Eighty slices caudal to this point were counted, and the total number of cells across all sections was taken to represent the proliferative state of the tissue. Counting was done under 20× epifluorescence, and was done blind to sample identity.

For BrdU counts in the main OB, complete counts were made from coronal sections made every 120 µm anterior to the closure of the AOB. The area of each section was measured, and the cell count was then normalized to this area to obtain the surface density of BrdU nuclei. We chose to make this normalization to eliminate variability caused by variations in the plane of sectioning.

**VOLUME ESTIMATION**
Olfactory bulbs were sectioned into 60-µm sagittal sections using a cryostat. On chilled slides, these sections were suspended in a drop of PBS to prevent folding and compression artifacts, and were then dehydrated on a heating plate. Cell nuclei were stained using hoescht or DAPI stain, and photographed for off-line analysis. Total volume was calculated by multiplying the total section thickness (60 µm) by the surface area of every section (Cavaleri’s principle).

**DENSITY ESTIMATION**
We developed a method to use the optical dissector off-line, eliminating the need to count cells while on a confocal microscope. Confocal z-stacks of DAPI-stained nuclei were taken every 1 µm over the middle 16 µm of a 20 µm cryosection on a laser-scanning confocal microscope (Zeiss LSM510, Thornwood, NY, USA). Care was taken to maintain similar brightness for images at each depth. The middle 12 µm of each z-stack was then false-colored off-line using the program ImageJ, whereby the top two slices were colored red, and the bottom 10 µm slices were colored green to blue (see Figure 3E). In this way we could easily define cells within an exclusion volume by identifying all red, yellow or orange cells. The remaining cells, if not occupying this exclusion volume, would be colored blue-green. For each anatomical counting region, the number of cells in the exclusion volume and in the counting volume was calculated at three randomly chosen slices anterior to the closure of the AOB.
ODOR DELIVERY
A custom made olfactometer was adapted from a previous design (Lorig et al., 1999) to deliver odors during electrophysiology. Compressed air was humidified and passed by the animal's nose at a rate of 1 l/min (carrier stream). The odor amyl acetate, to which the animals were naïve, was dissolved in 1 ml DMSO to a final concentration of 0.5 mM, and a 20 ml vapor headspace was kept saturated with a cotton or paper wick to maximize liquid surface area. Amyl acetate was used as a representative odor because it evoked highly reproducible and stereotyped spectral-temporal patterns across multiply sampled positions within the GCL. Odor puffs, 2 s in duration, were diverted into the carrier stream, and puffs were separated by a minimum of 40 s. To avoid pressure changes, a compensatory open air channel was switched off during odor presentation. No decrease of odor response was measured in the LFPs after repeated stimulation over hours (unpublished observations). Odor concentration reported in the text does not take into account the dilution by a minimum of 40 s. To avoid pressure changes, a compensatory open air channel was switched off during odor presentation. No decrease of odor response was measured in the LFPs after repeated stimulation over hours (unpublished observations). Odor concentration reported in the text does not take into account the dilution of the saturated headspace into the carrier stream, therefore final concentrations are much less. For every mouse, odor was delivered at least seven times, spaced apart with pulses of solvent headspace.

ELECTROPHYSIOLOGY
Anesthesia was induced with an intraperitoneal injection of ketamine (100 mg/kg) and xylazine (7 mg/kg) in mammalian Ringer’s solution and supplemental injections were given every ~ 45 min. The animal was mounted in a stereotaxic frame (David Kopf Instruments, Tujunga, CA, USA) and body temperature was maintained at 37°C with a feedback controlled heating pad (CT-1000, CWE, Ardmore, PA, USA). The animal’s nose was inserted into an air-tight gas mask through which humidified air from the olfactometer passed at a rate of 1 l/min. Two small craniotomies were performed at the left hemisphere at stereotaxic coordinates corresponding to the region above the LOT, and to the central dorsal surface of the OB. A custom made tungsten recording electrode (Hubel, 1957) with an impedance of 0.7–2 MΩ was inserted 1 mm deep with a hydraulic microdrive into the GCL of the MOB, and a custom made bipolar tungsten stimulating electrode was driven between 3.5–4.5 mm deep into the LOT. We recorded evoked field potentials (Figure 5) from the same location in the GCL as for the LFP recordings (Figure 6). Correct placement of the stimulating electrode was confirmed by recording the antidromic evoked waveform in the GCL (Shepherd and Haberly, 1970). If the evoked GCL waveform did not contain three distinct phases corresponding to (1) back-propagating spikes (2) field GC excitatory postsynaptic potentials (EPSPs) and (3) field MC inhibitory postsynaptic potentials (IPSPs), we assumed the stimulation electrode was reaching too many centrifugal afferents, and not enough LOT fibers. In this case the stimulating electrode was inserted to a new position. In some animals the position of the stimulating and recording electrode was confirmed histologically after burning the tip position, and visualized in 100-µm thick vibratome slices with cresyl violet.

Local field potentials were amplified (DP-301, Warner Instruments, Hamden, CT, USA), filtered (1 Hz highpass, 3 kHz lowpass), and digitized at 5–10 Khz (Power 1401, CED, UK). Breathing was continuously monitored with a piezoelectric force-transducer (Stoelting, Wood Dale, IL, USA) in contact with the animal’s back, and this signal was used to trigger odor delivery upon crossing a threshold corresponding to the transition of inhalation to exhalation (I/E transition). In the breathing trace, inhalation occurs during positive values; exhalation during negative values (see Figure 6B). All on-line event detection and triggering was done using the software Spike2 (CED, UK). Similarly, LOT stimulation was triggered by the phase of the breathing cycle, and current was created using a pulse generator and stimulus isolation unit (Grass S88, W. Warwick, RI, USA). For every mouse, the stimulus-intensity response relationship was recorded with the GCL recording electrode. All subsequent stimulation in the paired-pulse paradigm was given at saturating intensity (1–1.5 mA over 100 μs). Experiments that involved LOT stimulation were always performed after the completion of odor delivery protocols.

LFP SIGNAL PROCESSING AND ANALYSIS
All signal processing was performed off-line using custom scripts in Spike2, and in Matlab using a combination of custom scripts and the program eeglab 6.01b (Delorme and Makeig, 2004). LFP spectra for Figure 5 were made using the multi-taper fft algorithm PMTM, applied to 1024 samples (205 ms) with a time-bandwidth product of 4, over 500 ms windows of data surrounding the inhalation/exhalation (I/E) point of a breath. No significant spectral contribution is observed in the DMSO containing pulses, indicating that observed spectral changes around odor stimulation are not due to artifacts from the stimulation apparatus.

To characterize the response properties in both time and frequency domains we have employed wavelet transformation analysis of individual breaths. Wavelet transformations used the Morlet basis with an initial window size of 446 ms at 2 cycles, and tapered to 43 cycles. 400 analysis windows were placed within the middle 5.5 s of each epoch, producing a time bin size of 14 ms. Because analog filtering was not applied before digitization, raw LFP traces contained breath locked theta oscillations with asymmetrical rise and fall times. The rise phase of the theta oscillation had a fundamental frequency in the beta range, so beta power in wavelet spectrograms reflects both beta and theta oscillations.

Spectral power at every frequency band (94 bands from 5 Hz to 120 Hz) was normalized to the average band-power contained in the baseline region (1.5 s after I/E of breath 1). Spectral power above baseline is expressed in decibels [10 × log[10](signal/baseline)]. In this way, each odor response was normalized to its own spectral baseline, and the normalized power of repeated odor stimulations for each animal (7–9 per animal) was averaged. Because of this normalization, wavelet powers will not reflect absolute difference in spectral power between strains and conditions, but rather the flux in spectral power over time. Additional normalization to bring the pre-stimulus baseline to zero dB was done relative to the first three baseline breaths. Compound spectrograms were made by averaging across all animals in a group after the data had been corrected to baseline and normalized. All spectrograms are colored using the same LUT (−12 dB to +12 dB).

PAIRED-PULSE ANALYSIS
For short ISI (<100 ms), we observed that both the peak (GC fEPSP) and trough (MC fIPSP) of the second pulse occur on top of a moving baseline created by the first pulse. We assumed the two waveforms overlapped additively, and corrected for this bias by subtracting the value of a single pulse at the time of the event (fEPSP or fIPSP) from
the second pulse. This bias factor was taken from an average of multiple single stimulations, and the baseline of these single stimulations was set to zero before averaging. Therefore in all paired stimulations with ISI <100 ms, it is necessary to compare the bias factor to the specific baseline preceding the first stimulus (baseline 1).

The formula for the paired-pulse ratio of two fEPSPs for an ISI less than 100 ms was:

\[
\frac{\text{peak}_2 - (\text{bias} - \text{baseline}_1)}{\text{peak}_1 - \text{baseline}_1}
\]

where “peak” is the magnitude of the fEPSP peak, “bias” is the value of the average waveform at the time of the peak, and “baseline” is the value prior to the stimulus artifact. For measurements of fIPSPs, trough 1 was measured from the average trace that was already set to baseline prior to averaging, therefore it was not necessary to compare it to baseline 1. Thus, the formula for fIPSPs (ISI<100 ms) was:

\[
\frac{\text{trough}_2 - (\text{bias} - \text{baseline}_1)}{\text{trough}_1}
\]

For ISI greater than 100 ms, each peak and trough was set to their respective pre-stimulus baseline, so for fEPSPs this was calculated by the following formula:

\[
\frac{\text{peak}_2 - \text{baseline}_2}{\text{peak}_1 - \text{baseline}_1}
\]

\[
\frac{\text{fIPSP paired-pulse ratios (ISI>100 ms) were also calculated with formula 3.}
\]

ODOR-CUED FEAR CONDITIONING

All testing occurred during daylight hours and, with few exceptions, at least 2 h before or after the colony lights cycle. Prior to conditioning the mice were subjected to three 30-min acclimation sessions, each on 3 consecutive days in the fear conditioning chamber during which no CS or US were presented. On day 3 a fear conditioning session consisting of eight CS + US pairing trials with a 4 to 6-min inter-trial-interval was conducted for each mouse. Live observations were made to measure total freezing time for 2 min after each CS-US trial (Stopwatch+, David A. Brown, http://www.cbn-atl.org/research/stopwatch.shtml). 24, 48 and 72 h after conditioning mice were tested in a different context, the probe chamber, for 20 min: a null stimulus (mineral oil for the odor CS animals) was delivered at 2, 4, 6, and 8 min and the CS was delivered at 10, 12, 14, 16, and 18 min. Total freezing was measured throughout the 20-min session. Mice were tested at 6, 21, and 26 weeks after SVZ irradiation or sham treatment; in a parallel and balanced study half of the 26-week cohort was conditioned to an audio tone CS instead of an odor CS. The CS consisted of either a 3 to 5-s odor pulse or a 3-s audio tone followed immediately by a 2 s shock US.

The fear chamber (interior 30 cm × 24 cm × 25 cm H) has clear acrylic walls providing containment of the shock-grid floor apparatus (Med Associates, model ENV-005-02A) and ventilation sufficient to administer odors as discrete cues. The US consisted of a 0.5 mA, 2-s duration shock provided by a Med Associates ENV-414 shocker scrambler to coincide with administration of odor cues. The probe chamber (29 cm × 18 cm × 23 cm H) also has clear acrylic walls and uses the same ceiling fan for exhaust as the fear chamber, but has a smooth acrylic floor instead of the shocking grid floor. Odor is prepared as a 100 mM solution of pentanal (Sigma #110132) in light mineral oil (Sigma #330779). 2.5 mL of odor solution is sealed in a 16 × 125 mm culture tube (VWR #47729-578) with a section of filter paper to act as a wick. A 50 mL syringe is used to push 50 mL volume of air through the odor vial (or mineral oil vial for null stimulus trials) and into the testing chamber over 3-5 s. Audio CS was provided via PC speakers to deliver a 2 kHz square wave tone, 3 s duration, with 75 ± 5 dB intensity measured at the floor of the testing chamber.

STATISTICS

A one-tailed Students t-test was used to calculate statistical significance in all comparisons between two distributions. For repeated measures comparisons we used repeated measures ANOVA. All analysis was done in Statview 5.0.1 (SAS, Cary, NC, USA).

RESULTS

X-RAY IRRADIATION OF THE MOUSE SUB-VENTRICULAR ZONE PRODUCES A DOSE-DEPENDENT AND LONG-LASTING REDUCTION IN LOCAL CELL PROLIFERATION

X-ray irradiation is often clinically used to inhibit cancer cell proliferation and brain metastasis. Experimentally it has been successfully applied to ablate adult neurogenesis in the dentate gyrus of the hippocampus (Santarelli et al., 2003), and in the SVZ (Panagiotakos et al., 2007; McGinn et al., 2008; Lazarini et al., 2009). We targeted x-rays to specific regions of the mouse brain by using lead shielding aligned to stereotaxic coordinates. This strategy permitted accurate and reproducible bilateral targeting of the SVZ and a majority of the RMS without exposing the hippocampus or the olfactory epithelium (Figure 1A). Within the window of x-ray exposure, the SVZ was largely absent of cell proliferation as measured by 5-bromo-3'-deoxyuridine (BrdU) incorporation measured 24 h after a single injection (Figure 1B). Targeted irradiation did not alter adult neurogenesis in the dentate gyrus verified by doublecortin (dcx) staining (Figure 1C). The dosage of irradiation was calibrated by changing the duration of exposure on each of 3 days and measuring BrdU incorporation in the SVZ 2 weeks after the last day of irradiation. The most effective dose we tried was calculated to be equivalent to 15 Gy of ionization (Figure 1D, see methods, BrdU + nuclei per SVZ ± SEM; 0 Gy 550 ± 38.1; 1 Gy 640 ± 85.7; 5 Gy 450 ± 37.8; 10 Gy 136 ± 10.7; 15 Gy 45 ± 8.7; n = 3–9 per group, significance between 0 Gy and 15 Gy and 0 Gy and 10 Gy, unpaired t-test p < 0.001), which is similar to what has been previously reported (Mizumatsu et al, 2003; Santarelli et al., 2003; McGinn et al., 2008). In addition, a significant depression of cell proliferation in the SVZ persisted for over 4 months following exposure (Figure 1E, BrdU + nuclei per 15 Gy SVZ ± SEM; 3-days post x-ray 7 ± 2.4; 7-days post x-ray 45 ± 8.7; 14-days post x-ray 67 ± 8.9; 28-days post x-ray 34 ± 4.0; 120-days post x-ray 51 ± 15.6; pooled 0 Gy sham controls 525 ± 18.8; n = 3–10 per group, unpaired t-test p < 0.001).

SURVIVAL OF YOUNG ADULT-BORN NEURONS IN THE OLFACTORY BULB IS UNCHANGED BY REDUCING NEUROGENESIS

Young adult-born neurons undergo activity-dependent apoptosis during a critical period 2–4 weeks after cell birth (Yamaguchi and Mori, 2005; Mouret et al., 2008). We wanted to measure if SVZ
irradiation altered neuronal survival throughout this process. One month after irradiation, a persistent loss of adult-born neurons was qualitatively visualized by a reduction in dcx staining, a label for migrating progenitor cells (Figure 2A). We quantified the loss of neurogenesis to the OB by acutely injecting the animal with 5-bromo-2′-deoxyuridine (BrdU), and measuring the number of BrdU positive cells that migrated into the GCL. To observe adult-born GCs during their critical period, we counted BrdU positive nuclei 14 and 28 days after BrdU injection, in animals irradiated 3 months prior (Figure 2B). A 15 Gy dose reduced the number of adult born cells of the same age by approximately 96% 2 or 4 weeks after injection (Figure 2C, BrdU nuclei per mm² ± SEM 2-weeks post injection: 0 Gy 69.1 ± 5.1 (white); 7.5 Gy 29.5 ± 1.9 (grey); 15 Gy 2.5 ± 0.8 (red); BrdU nuclei per mm² ± SEM 4-weeks post injection: 0 Gy 42.0 ± 7.0 (white); 7.5 Gy 16.4 ± 0.7 (grey); 15 Gy 1.7 ± 0.2 (red)). Interestingly, when we compared the loss of BrdU labeled cells between 2 and 4 weeks at full or intermediate doses of irradiation, we found that this loss was not affected by the number of similarly aged cells integrating into the circuit (Percent decrease between 14 and 28 days; 0 Gy 39.2%; 7.5 Gy 44.4%; 15 Gy 33.6%). Thus, to our surprise, we found no compensatory increase in survival of adult born GCs when fewer of them migrate into the OB.

THE NUMBER OF CELLS IN THE Olfactory Bulb Granule Cell Layer IS REDUCED MONTHS FOLLOWING SVZ IRRADIATION

Considering that young adult-born GCs continue to die at the normal rate even under conditions of reduced neurogenesis, we next looked to see if GCs of any age showed increased survival. To estimate the total number of cells in the GCL, we combined measurements of density and volume over a period of months after irradiation.

We estimated the volume of the GCL by applying the Cavalieri principle to serial sectioning of the entire tissue (see Materials and Methods). Sagittal sections of the OB of irradiated mice showed a clear depletion of the RMS compared to sham controls (Figure 3A, arrows, compare left to right). The surface area encompassed by the border of the GCL was measured from animals that had received SVZ irradiation 1, 2, 3.5 or 5 months prior. In irradiated animals we found a uniform reduction in the medial to lateral cross-sectional area compared to sham irradiated animals (Figure 3B). This isotropic loss of volume along the medial-lateral axis suggests cell loss in the superficial layers of the GCL (which is present in all sections), rather than collapse of the RMS (present only in the middle sections). Reduced GCL volume in irradiated conditions was significant at all time points compared to sham controls (Figure 3C and Table 1). The time course of this reduction in volume peaked at 2 months; at that point the GCL was 18% smaller in the X-ray treated group compared to controls. A slight recovery of volume was observed from 2 months to the last measurement at 5 months. This recovery is likely due to a recovery of SVZ proliferation from neurogenic regions at the fringes of the irradiation window, and thus outside the counting window (data not shown).

We next measured the cell density in different regions in the main OB by combining biased placement of counting windows (Figure 3D) with unbiased counting (the optical dissector, Figure 3E). In this way we could detect changes in known sub-regions of the GCL with accurate stereological estimation. The
density of DAPI-stained nuclei in the GCL was measured in the medial, ventral or dorsal GCL in mice irradiated 1 and 4 months prior. No significant differences were observed between age groups or between anatomical regions (ANOVA with post hoc PLSD, \( p > 0.05 \)), so data were pooled (Figure 3F). The density of cells in the GCL of irradiated animals (1.102 ± 36.5 nuclei × 10³ per mm³) was not significantly different from that of non-irradiated animals (1.128 ± 43.7 nuclei × 10³ per mm³, unpaired \( t \)-test \( p > 0.05 \)).

From these measurements of volume and density, we could calculate the change in the total number of cells in the GCL after irradiation. With an average density of cells of 1.1 million per mm³, the reduction in GCL volume indicates that 1 month after irradiation the tissue lost \( \sim 368,000 \) cells, and 2 months after irradiation it lost \( \sim 590,000 \) cells. At both time points, this indicates a similar rate of loss (\( \sim 12,000 \) per day over 1 month; \( \sim 10,000 \) per day over 2 months).

**IRRADIATED MICE HAVE DIMINISHED CONDITIONED RESPONSE IN AN ODOR-CUED FEAR CONDITIONING TASK**

The proliferation and survival of adult born GCs is strongly linked to the animal’s environment. We therefore sought to determine if the reduction of granule cell proliferation would produce an observable behavioral effect in a challenging environment.

We developed a model of odor-cued fear conditioning to assess olfactory processing in mice with diminished adult neurogenesis. In our odor-cued fear conditioning paradigm we paired an olfactory conditioned stimulus (CS) with an aversive footshock unconditioned stimulus (US). After one session of conditioning, animals formed a fearful association to the CS exhibited by their freezing behavior. We probed this association in a novel context by measuring the animal's freezing conditioned response (CR) to the CS in the absence of the US (Figure 4A). Using cued rather than contextual conditioning allows us to assess the CR as it pertains to the CS specifically, and assess the animal’s behavior in the absence of the CS. This is illustrated (Figure 4B) for a 20 min long probe session with data from all mice pooled into either irradiated (XR) or control (Sham) groups: the CS, in the form of a brief odor pulse, results in a dramatic increase in freezing behavior during the 1-min time bins following CS delivery, the "post-CS minute". During the 10 min prior to CS delivery the odor vehicle, mineral oil, was pulsed into the probe chamber. While the XR and Sham mice did not differ in level of locomotor activity (data not shown) or in the amount of baseline freezing in the probe session (data not shown), it was observed that irradiated mice freeze less in response to the odor CS during the post-CS minute (XR \( n = 17 \) and sham \( n = 18 \); repeated measures ANOVA, \( p < 0.05 \)) (Figure 4B).
following odor-cued conditioning (repeated measures ANOVA \( p < 0.05 \); Figure 4E). Importantly, this effect was observed in three cohorts of mice tested 6, 21, and 26 weeks after SVZ irradiation; their data is pooled here (\( XR = 17 \), sham \( n = 18 \)).

**DENDRO-DENDRITIC SYNAPTIC PLASTICITY IN-VIVO IS UNCHANGED IN THE ABSENCE OF YOUNG ADULT-BORN NEURONS, AND AFTER GRANULE CELL LAYER CELL LOSS**

To probe the OB for electrophysiological changes that may mirror the behavioral phenotype, we first measured the properties of the synapse formed between GCs and mitral/tufted (MT) cells. Antidromic stimulation of the MT cell axon bundle in the lateral olfactory tract (LOT) excites this synapse, producing a field potential reflecting glutamatergic granule cell excitation followed by GABAergic M/T cell inhibition (Rall and Shepherd, 1968; Nicoll, 1969; Mori and Takagi, 1978) (Figure 5A). We used the magnitude of the field EPSP (fEPSP) and field IPSP (fIPSP) to reflect aggregate synaptic strength (Figure 5B).

We first sought to determine if, in the absence of young adult-born neurons, the population of GCs showed less excitability. This may arise if young adult-born neurons are intrinsically hyper-excitatory, as previously suggested (Belluzzi et al., 2003; Saghatelyan et al., 2005). No significant differences were observed in the stimulus response relationship at any age in either the magnitude of the GC response (fEPSP magnitude; Figure 5C, left, data grouped across all time points, unpaired \( t \)-test \( p > 0.05 \)) or in the input-output function of the response (stimulus strength at half-maximal fEPSP; Figure 5C, right, data grouped across all time points, unpaired \( t \)-test \( p > 0.05 \)).

We next looked at the short-term plasticity of the dendrodendritic synapse at different times after irradiation to probe for changes to pre or post-synaptic function. For this we stimulated the LOT using a paired-pulse paradigm which reveals facilitation of the GC fEPSP (Figures 5D–F, left) and depression of the reciprocally evoked mitral and tufted cell fIPSP (Figures 5D–F right) at short inter-stimulus intervals (ISI). Between 1 and 5 months after irradiation, no systematic changes were observed in the plasticity profiles of this synapse, nor were differences observed within any individual time point (repeated measures ANOVA \( p > 0.05 \)).
LOCAL FIELD POTENTIAL RESPONSES TO ODORANT STIMULATION REMAIN UNPERTURBED MONTHS FOLLOWING IRRADIATION

Having observed no changes to the dendro-dendritic synapse in a bulb largely absent of new adult-born neurons, we reasoned that functional changes may only emerge if odors are used as stimuli. To gain a system-wide perspective of how odors are processed in the OB, we measured local field potential (LFP) oscillations which, when measured in the deep GCL, largely reflect aggregate coherent synaptic activity in the external plexiform layer (Rall and Shepherd, 1968). In addition, distinct frequency bands of LFP oscillations can be tied to specific mechanisms; for instance, gamma oscillations are produced by the mitral to granule cell dendro-dendritic synapse (Schoppa, 2006; Lagier et al., 2007; Kay et al., 2009).

In freely breathing anesthetized mice, we recorded the LFP oscillations that emerge around an odor stimulus. During normal respiration we observed prominent slow oscillations locked to the breathing rate (Figure 6B), with embedded higher frequency oscillations emerging and decaying within the course of a breath. We measured the relationship between odor concentration and evoked LFP power in beta frequency oscillations (Figure 6C, 10–40 Hz), gamma frequency oscillations (Figure 6D, 40–70 Hz), and high gamma oscillations (Figure 6E, 70–100 Hz) from mice 1 and 2 months after irradiation. No significant differences emerged at either time point, or when the data were pooled (Figures 6C–E, unpaired t-test, p > 0.05).

Neural activity in the OB is tightly constrained by the breathing cycle (Bathellier et al., 2008; Wesson et al., 2008), so finer spectral analysis was needed to extract how LFP frequencies emerge and decay within a breath. We turned to wavelet techniques that can estimate the power of LFP oscillations with high temporal resolution (see Materials and Methods). This data can be visualized by frequency-time spectrograms which we centered on the time when the mouse transitions from inhalation to exhalation (Figure 6F). Analysis of this data at different times after irradiation failed to reveal any fine temporal disruptions in the emergence of neural activity within a breath, illustrated by the profile of oscillatory power in the gamma (Figure 6G, top) and beta bands (Figure 6G, bottom) measured 1–2 months after irradiation (repeated measures ANOVA, p > 0.05).

DISCUSSION

Adult neurogenesis in the OB has been known and investigated for more than two decades, but a clear function for this robust cellular production, nearly unique in the vertebrate brain, has been elusive. Here we provide new evidence for the function of adult neurogenesis in the OB granule cell population. We produced a 20-fold reduction in the number of incoming adult-born neuroblasts, and observed a behavioral deficit in odor-cued fear conditioning. However, in spite of a reproducible behavioral effect, we were unable to identify any corresponding electrophysiological change measured from the altered GCL. Surprisingly, a variety of electrophysiological tests showed normal GCL circuit function as the standing population of old GCs declined in number by up to 18% compared to the non-irradiated condition. This suggests robust functional homeostasis in a circuit with a declining proportion of interneurons.

A further unexpected finding was that, in parallel with the slow attrition of the GCL population, survival of young adult-born GCs was unchanged after SVZ irradiation. Specifically, survival of young neurons during their “critical period” between 2 and 4 weeks of cell age (Petreanu and Alvarez-Buylla, 2002; Yamaguchi and Mori, 2005) remained constant regardless of the level of SVZ proliferation. As a result, survival of young adult-born GCs in this circuit is likely not determined by competition for scarce trophic resources contrary to hippocampal adult neurogenesis (Sairanen et al., 2005) and classical...
neurotrophic regulation of survival in the peripheral nervous system (Levi-Montalcini, 1987). Thus, the extent of neurogenesis from the SVZ had no effect on either the physiology or the dynamics of neuronal turnover within this tissue. Nevertheless, the robust behavioral phenotype in irradiated mice is the first evidence that adult SVZ neurogenesis has functional relevance to an extensive behavioral system beyond the OB.

The fear conditioning paradigm probes the interaction between olfactory processing and the amygdala (Sah et al., 2003). In rats, lesions of the amygdala are known to disrupt the association of CS and US regardless of stimulus modality or response measure (Blanchard and Blanchard, 1972). The basic mechanism involves association within the amygdala of the US, a foot shock, with a CS, here an odor pulse that is paired with foot shock. Projections from the amygdala control the fear response and target sensory areas including the olfactory cortex which, along with the OB, return projections to amygdalar nuclei (Scalia and Winans, 1975; Shepherd, 1998).

We observed a diminished fear response of SVZ irradiated mice to a fear-conditioned odor, despite successful formation of the association. This phenotype did not appear when an auditory stimulus was used, and it did not reflect any disruption in their ability to exhibit fear during the formation of the association. In addition, recent work has shown that SVZ irradiated mice possess unaltered detection thresholds and a normal capacity for fine odor discrimination, indicating that our phenotype was not the result of disrupted sensitivity to the conditioned stimulus (Lazarini et al., 2009). Intriguingly, the aforementioned study also found disrupted long-term recall of a positive-valence association, which, when taken together with our observed phenotype, may generally implicate young adult-born GCs in the recall or maintenance of both positive and negative valence associations.

This phenotype could result from the acute loss of young adult born neurons in the MOB, or after months due to the slow decline in interneuron population size. As we are unable to rescue adult SVZ proliferation in the x-ray model we cannot directly test if an acute loss of adult born cells alone is sufficient to generate this phenotype. Nonetheless, we observed this behavioral effect at three time points after irradiation, each with a different amount of GCL cell loss, suggesting our behavioral phenotype is the result of an acute loss of young adult born neurons. In addition, the phenotype was consistently observed 6–26 weeks after irradiation, ruling out the off-target effects of cranial irradiation such as inflammation (Monje et al., 2002).

**FIGURE 5** | Loss of neurogenesis does not change the activation and plasticity of the dendro-dendritic synapse in-vivo. (A) Schematic of the stimulation paradigm, with a diagram of the dendro-dendritic circuit between mitral cells (MC) and granule cells (GC). (B) The waveforms following stimulation have three distinct phases corresponding to the mitral cell compound action potential (cAP), the field EPSP (fEPSP) originating in the granule cells, and the reciprocal field IPSP (fIPSP) in mitral cells.

(C) Input-output function relating intensity of 100 µs LOT stimulation to fEPSP magnitude from all animals. The current that evokes a half-maximal response is depicted on the right. (D–F) Paired-pulse stimulation of the synapse at maximal intensity, and over a range of inter-stimulus intervals (ISI) from 10 ms to 1 s. This paradigm reveals minor facilitation (values > 1) in the granule cell fEPSP (left) and large depression (values < 1) in the mitral cell fIPSP (right).
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In searching for physical loci for the behavioral phenotype, we made measurements that reflect general olfactory processing across a wide region of the OB. While these tests are necessarily an incomplete characterization of OB function, they nonetheless reflect two critical aspects of granule cell function: the generation of oscillatory potentials, and the dendro-dendritic inhibition of mitral and tufted cells. In a previous report, locally generated field potentials in the OB were sensitive to physiological correlates of olfactory behavior (Le Pichon et al., 2009). Specifically, odor-evoked LFP measurements assay the aggregate current evoked during the processing of a stimulus and thus reflect the magnitude and synchronization of neural activity in a local volume. With their temporal and spectral complexity these oscillations are informationally rich, yet, using the

Our observed behavioral phenotype may result from altered centrifugal modulation of the OB. Specifically, odor-cued fear conditioning may use the centrifugal systems that feed back onto young adult-born GCs. Anatomically, the medial amygdalar nuclei connect to the OB through the olfactory peduncles and piriform cortex (Shepherd, 1998). Recent work suggests that the proximal dendrites of GCs, which receive inputs from piriform cortex among other regions, function as the gatekeeper of information from different brain modalities into the OB (Balu et al., 2007), and exhibit unique plasticity in young adult-born neurons (Nissant et al., 2009). Our behavioral phenotype was also similar to changes in contextual freezing behavior observed after ablating hippocampal adult neurogenesis (Saxe et al., 2006). As suggested in the dentate gyrus, young adult-born olfactory GCs may be particularly influenced by high-valence stimuli or states of arousal, potentially through their muscarinic, cholinergic, (Castillo et al., 1999; Linster and Cleland, 2002) or noradrenergic synaptic inputs (Doucette et al., 2007). In high-valence olfactory environments these pathways may act on GCs in concert with proximal synaptic inputs from the cortex, recruiting specifically young adult born GCs into new patterns of functional connectivity.

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most sensitive techniques available we could find no alteration in spectral-temporal patterning in the GCL. With paired-pulse measurements, we monitored short-term synaptic plasticity specifically among the population of dendro-dendritic synapses. This synapse is the only output of the granule cell, and in a previous report was the locus for homeostatic compensation for olfactory deprivation (Saghatelyan et al., 2005). We found no evidence of altered short-term plasticity, or a change of input gain to either the M/T or GC side of the dendro-dendritic synapse.

In characterizing the x-ray lesion, we found that the survival of young adult-born GCs is uncoupled from the quantity of cells of the same age integrating into the circuit. This indicates that the survival of a neuroblast within its critical period is not regulated by competition with neurons of the same age. Instead, as is well established from the literature, neuroblast survival is sensitive to efferent activity (Petreanu and Alvarez-Buylla, 2002; Rochefort et al., 2002; Winner et al., 2002; Yamaguchi and Mori, 2005; Mandairon et al., 2006) and afferent activity (Alonso et al., 2006; Mouret et al., 2008). This leads to a quandary regarding the maintenance of the GCL: how does the tissue adjust to variations in the amount of SVZ neurogenesis? These variations may occur frequently and unpredictably. SVZ proliferation is modulated by many factors including pregnancy (Shingo et al., 2003), ischemia (Zhang et al., 2004) seizures (Parent et al., 2002) and a multitude of trophic factors such as EGF (Kuhn et al., 1997) and sonic hedgehog (Machold et al., 2003). In the absence of changes in the olfactory environment, from our data we would predict that downward fluctuations in SVZ proliferation deprive the GCL of young adult-born neurons, and upward fluctuations produce over-abundance.

The innate plasticity of GCs may be one way the GCL adapts to fluctuations in neurogenesis (Bonfanti and Theodosis, 2009). Adult born GCs, but not post-natal born GCs, are uniquely capable of long-term synaptic plasticity (Nissant et al., 2009). Furthermore, adult born GCs, but not post-natal born GCs, display increased excitability to compensate for decreased spine density following olfactory deprivation (Saghatelyan et al., 2005). Similar homeostatic mechanisms may emerge after changes in SVZ proliferation to adjust the functional properties of the GCL. These as yet unidentified mechanisms would constitute an important link between SVZ neurogenesis and OB physiology.

While this study did not address the properties of the two other main categories of SVZ-derived adult-born neurons, peri-glomerular cells, and the GCs of the accessory OB, it is quite possible that neurogenesis among these populations has a unique functional purpose. We also cannot rule out the possibility that the behavioral observations stem from changes to these circuits. Interestingly, the percentage of adult-born neurons in the GCL remains constant at approximately 10%, while the glomerular layer undergoes a net addition of adult-born neurons over time (Ninkovic et al., 2007), similar to neuronal addition in the hippocampus (Bayer et al., 1982), suggesting a unique role for adult neurogenesis in the peri-glomerular circuit over the life of the animal.

Further defining the functional significance of adult neurogenesis will require sophisticated genetic dissection in concert with multiple behavioral and physiological tests. Neurogenesis in specific subpopulations, such as those defined by protein expression (e.g Pax6), pharmacological profile (e.g adrenergic receptor subtypes), or morphology (e.g inter-glomerular vs. intra-glomerular) may be regulated according to independent mechanisms, and may subserve specific and independent circuit functions. A challenge for the future will be to uncover the range of functions imparted by adult neurogenesis onto these diverse microcircuits. This knowledge is crucial for a program in which these adult born neurons can be of value in other brain regions that have been affected by damage, disease or aging.

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AUTHOR CONTRIBUTIONS
Matthew T. Valley, Tanner R. Mullen and Stuart Firestein wrote the paper. Matthew T. Valley designed and analyzed, and Matthew T. Valley and Lucy C. Schultz conducted histology experiments. Matthew T. Valley and Botir T. Sagdullaev designed, and Matthew T. Valley conducted and analyzed electrophysiology experiments. Tanner R. Mullen designed, conducted, and analyzed behavior experiments.

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