Anxiety- and Depression-Like States Lead to Pronounced Olfactory Deficits and Impaired Adult Neurogenesis in Mice.
Eleni Siopi, Marie Denizet, Marie-Madeleine Gabellec, Fabrice De Chaumont, Jean-Christophe Olivo-Marin, Jean-Philippe Guilloux, Pierre-Marie Lledo, Françoise Lazarini

To cite this version:

HAL Id: pasteur-01300311
https://hal-pasteur.archives-ouvertes.fr/pasteur-01300311
Submitted on 10 Apr 2016

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

Distributed under a Creative Commons Attribution - NonCommercial 4.0 International License
Anxiety- and Depression-Like States Lead to Pronounced Olfactory Deficits and Impaired Adult Neurogenesis in Mice

Eleni Siopi, Marie Denizet, Marie-Madeleine Gabellec, Fabrice de Chaumont, Jean-Christophe Olivo-Marin, Jean-Philippe Guilloux, Pierre-Marie Lledo, and Françoise Lazarini

1Institut Pasteur, Unité Perception et Mémoire, F-75015 Paris, France, 2Centre National de la Recherche Scientifique, Unité Mixte de Recherche 3571, F-75015 Paris, France, 3Institut Pasteur, Unité d’Analyse d’Images Quantitative, F-75015 Paris, France, 4Centre National de la Recherche Scientifique, Unité de Recherche Associée 2582, F-75015 Paris, France, and 5Institut National de la Santé et de la Recherche Médicale UMR-S 1178, Faculté de Pharmacie, Université Paris-Sud, Châtenay-Malabry F-92296, France

Numerous clinical reports underscore the frequency of olfactory impairments in patients suffering from major depressive disorders (MDDs), yet the underlying physiopathological mechanisms remain poorly understood. We hypothesized that one key link between olfactory deficits and MDD lies in hypercortisolemia, a cardinal symptom of MDD. Corticosterone (CORT) is known to negatively correlate with hippocampal neurogenesis, yet its effects on olfactory neurogenesis and olfaction remain unknown. Here we used a rodent model of anxiety/depression-like states, which is based on chronic CORT administration and studied the effects of the antidepressant fluoxetine (FLX) on behavior, olfaction, and adult neurogenesis in the dentate gyrus (DG), olfactory bulb (OB), and the olfactory epithelium (OE). Chronic CORT had no effect on cell proliferation in the OE or on olfactory sensory neurons projecting to the OB, but induced pronounced deficits in olfactory acuity, fine discrimination of odorants and olfactory memory. These alterations were accompanied by a significant decrease in the number of adult-born neurons in both the DG and OB. Remarkably, FLX not only reversed depression-like states as expected, but also improved olfactory acuity, memory, and restored impaired adult neurogenesis. However, fine olfactory discrimination was not restored. Morphological analysis of adult-born neurons in both the DG and the OB showed that dendritic complexity was not significantly affected by CORT, but was increased by FLX. These findings demonstrate an essential role for glucocorticoids in triggering olfactory impairments in MDD and highlight a novel therapeutic effect of FLX.

Key words: corticosterone; dentate gyrus; olfactory bulb; olfactory epithelium; serotonin; subventricular zone

Significance Statement

Increasing clinical reports show that major depression is characterized by pronounced olfactory deficits, yet the underlying mechanisms remain unknown. In this work, we used an endocrine model of depression to study whether hypothalamic-pituitary-adrenal axis perturbation could be sufficient to provoke olfactory impairments. We found that chronic corticosterone not only induces marked deficits in olfactory acuity, fine discrimination and olfactory memory, but also significantly decreases bulbar and hippocampal neurogenesis. Importantly, the antidepressant fluoxetine restores both adult neurogenesis and depressive states, and improves most olfactory functions. Our data reveal that impairment of hypothalamic-pituitary-adrenal axis during depression can lead to olfactory deficits and that the neurogenic effects of selective serotonin reuptake inhibitor antidepressants can successfully restore certain olfactory functions.

Introduction

Major depressive disorders (MDDs) are highly prevalent and debilitating conditions, whose pathophysiological mechanisms and response to treatments remain poorly understood (Krishnan and Nestler, 2008). Increasing clinical reports underscore the frequency of olfactory impairments in patients suffering from MDDs (Pause et al., 2001; Negoias et al., 2010), yet, it remains...
unclear how mood states and olfactory processes interact in psychiatric disorders such as MDDs.

Olfactory deficits in MDDs could be partly due to endocrine dysfunction, which is suspected to play a key role in anxiety/depressive disorders. In fact, ~80% of severely depressed patients exhibit chronic hyperactivity of the hypothalamic-pituitary-adrenal (HPA) axis, mainly reflected by an excess of plasma cortisol (Anacker et al., 2011). This phenomenon is modeled in rodents, which show HPA axis abnormalities and increased corticosterone (CORT) levels upon unpredictable chronic mild stress, early life stress, or chronic social stress (Lehmann et al., 2013). Both the hippocampus and the olfactory bulb (OB) are densely populated with receptors for glucocorticoids (Morimoto et al., 1996). Although the effects of glucocorticoids in the adult OB remain relatively unknown, they play an important role in the hippocampus by negatively regulating the HPA axis, resulting in stress resilience. In conditions of chronic stress, when there is a sustained overproduction of glucocorticoids, this inhibitory feedback is impaired, creating a cascade of hippocampal damage and resulting in maladaptive stress responses (Selye, 1950; Mirescu and Gould, 2006).

Abundant evidence suggests that chronic stress is highly regulated by adult hippocampal neurogenesis (Dranovsky and Hen, 2006). In rodents, adult neurogenesis is a key mechanism to maintain plasticity in response to both internal and environmental challenges (Lledo et al., 2006), and its functional relevance ranges from perceptive to cognitive functions, including mood regulation, pattern separation, learning, memory, and behavioral responses to olfactory cues (Breton-Provencher et al., 2009; Lazarini and Lledo, 2011; Alonso et al., 2012). Abnormal hippocampal neurogenesis is likely to participate in psychiatric disorders, such as depression (Sahay and Hen, 2007), and the production of adult-born hippocampal neurons is influenced by stress and glucocorticoids (Murray et al., 2008; David et al., 2009; Snyder et al., 2011). In MDD patients and mouse models of chronic stress, there is evidence for reduced hippocampal volume and impaired adult neurogenesis, both of which can be restored by antidepressant treatments (Campbell et al., 2004; Boldrini et al., 2009).

Moreover, adult hippocampal neurogenesis is required for the response to antidepressants (Santarelli et al., 2003; David et al., 2009), and adult-born neurons are essential in restoring hippocampal inhibitory control over the HPA axis and in normalizing glucocorticoid levels following chronic stress (Snyder et al., 2011).

In this work, we determine whether such a link could be also drawn between HPA axis dysfunction and olfactory neurogenesis. In rodents, newborn neurons from the lateral wall of the subventricular zone (SVZ) migrate en route to the OB, where they differentiate into bulb interneurons. The few existing studies on the impact of chronic stress on olfactory neurogenesis show mixed results. While prolonged mild stress or CORT treatment has been shown to decrease cell proliferation in the SVZ (Hitoshi et al., 2007; Lau et al., 2011), there is contradictory evidence that the latter is not sensitive to repeated stress (Chen et al., 2006; Surget et al., 2008). Nevertheless, clinical studies have reported that adults with a history of childhood stress or MDD can have reduced OB volume (Negoias et al., 2010; Croy et al., 2013), suggesting a potential impact of chronic stress on OB functioning.

To investigate whether defects in the HPA axis could alter olfactory neurogenesis and olfaction, we used a neuroendocrine mouse model of anxiety/depression (David et al., 2009). We found that chronic CORT affects adult neurogenesis and induces both anxiety/depression-like states and olfactory impairments. We then report that coadministration of the antidepressant fluoxetine (FLX), a selective serotonin reuptake inhibitor, restores adult neurogenesis and attenuates the olfactory deficits.

Materials and Methods

**Subjects**

Experiments were performed using adult male C57BL/6Nac mice (8 weeks old, 20–25 g), purchased from Taconic Farms. They were housed in groups of five and maintained in standard conditions (controlled room temperature and humidity, 12 h light/dark cycle, with lights on at 8:00 a.m., ad libitum access to dry food pellets and water) at the Pasteur Institute animal care facility, officially registered for experimental studies on rodents (approval number for animal care facilities: A 75-15-01-6 2014-720). These studies were performed in compliance with the French legislation (Decree of Feb 1, 2013, 2013-118) and the European Communities Council Directive of Sept 22, 2010 (2010/63/EEC). Experimental designs and procedures were approved by the local ethics committee (no. 2011-0020, CETEA 59 and no. 2013-0081, CETEA 89).

**Drugs**

The dose of CORT and FLX treatments was selected upon previous studies (David et al., 2009). CORT (4-pregnen-11b-diol-3,20-dione 21-hemisuccinate, from Sigma-Aldrich) was dissolved in its vehicle (0.45% hydroxypropyl-β-cyclodextrin, β-CD, Sigma-Aldrich) and was available ad libitum in drinking water (35 µg/ml/d, equivalent to ~5 mg/kg/d). FLX (Astra Trading, 18 mg/kg/d) was delivered either with the vehicle or with CORT. During the period of water restriction, CORT was administered intraperitoneally (i.p.) as previously described (Harris et al., 2011).

**Design and treatment groups**

Mice were divided in a total of 11 treatment groups (total n = 105) for four separate experimental sets.

**Experimental set 1. Impact of CORT and FLX on behavior, olfaction, and adult neurogenesis (Figs. 1, 2, 4–6).** One group received the vehicle (n = 12) for 20 weeks and the second group received CORT (n = 20) for 20 weeks. The third group received CORT during 6 weeks and then CORT + FLX during 14 weeks (n = 35). Using standard behavioral tests, we confirmed the anxiogenic depressive phenotype of mice at 4 weeks of CORT treatment (Fig. 1). One test per day was performed in the following order: open field, elevated plus maze, novelty suppressed feeding and splash test. Then, olfactory tests were conducted at the eighth week of CORT treatment, using fewer mice in each group (vehicle group: n = 8; CORT group: n = 8; CORT + FLX group: n = 35). The choice of mice was random and the remaining animals were eliminated from the study. Following 1 week of training in the olfactometers, olfactory tests lasted 5 weeks (Fig. 3). Olfactory perception was assessed 4 weeks following the onset of FLX administration. Simple and difficult olfactory discrimination (carvones and carvone mixtures) was assessed at 6 and 7 weeks, respectively, following the onset of FLX treatment. Finally, olfactory memory was assessed at 12 weeks following the onset of FLX treatment. All mice (n = 51) were subjected to behavioral tests at week 16 of CORT onset (1 test per day, as follows: open field, elevated plus maze, novelty suppressed feeding, splash test, tail suspension test, and forced swim test; Fig. 2) and to an olfactory memory test at week 18 (Fig. 3F). Immunohistochemistry was performed at week 20 using 10 mice from the CORT + FLX group, five mice from the vehicle group and five mice from the CORT-treated group, that were randomly chosen (see Figs. 5, 6).

**Experimental set 2. Impact of chronic FLX on behavior (Fig. 3).** One group received the vehicle (n = 10) and the other group received FLX...
Chronic CORT treatment induces anxiety and depression-like states. Following 4 weeks of CORT administration, anxiety, and depression-like behaviors were assessed using the open field, elevated plus maze, novelty suppressed feeding test, and splash test. CORT decreased the time spent in the center of the open field and in the open arms of the elevated plus maze. The third group received CORT during 4 weeks and then FLX during 7 weeks. The fourth group received the vehicle during 4 weeks and then FLX during 7 weeks. The latency to feed was measured during 10 min. CORT did not affect feeding drive. The total number of grooming sessions within the 6 min trial was significantly decreased by CORT. Values are plotted as mean ± SEM (n = 12 for vehicle-treated and n = 55 for the CORT group). *p < 0.05, ***p < 0.001 compared with vehicle-treated.

Experimental set 3. Impact of CORT on cell proliferation in the olfactory epithelium (see Fig. 7). Immunohistochemistry was done using two groups of mice, one group that was treated with CORT during 4 weeks (n = 5) and a group that received the vehicle (n = 5).

Experimental set 4. Impact of FLX on the morphology of adult-born OB neurons in CORT-treated and naive mice (see Fig. 8). The first group (n = 4) received the vehicle and the second group (n = 4) received CORT during 11 weeks. The third group received CORT during 4 weeks and then CORT + FLX during 7 weeks (n = 4). The fourth group (n = 4) received the vehicle during 4 weeks and then FLX during 7 weeks (see Fig. 8A). A lentiviral vector was injected in the SVZ at week 8 to label adult newborn neurons (see Fig. 8C). Immunohistochemistry was performed at week 11.

Behavioral assessment
A test battery of behavioral tests was used for the mice included in experimental sets 1 and 2, to assess locomotor activity, anxiety, and depression-like behavior and olfaction.

Open-field. Animals were placed in white Plexiglas containers (43 × 43 cm$^2$) and their behavior was recorded by a video camera during 30 min. A tracking system (Noldus Ethovision 3.0) was used to map center and periphery zones and to calculate the time spent in each zone.

Elevated plus maze. The test was conducted using a plus-cross-shaped apparatus made of black Plexiglas that was elevated 58 cm above the floor and comprised two open and two closed arms (30 × 6 cm) that extended from a central platform (7 × 7 cm). The Noldus Ethovision 3.0 tracking system was used to record behavior for 6 min. The time spent in center, open arm, and closed arm zones were calculated.

Novelty suppressed feeding. The test container used was a white plastic box (50 × 50 × 20 cm) whose floor was covered by wooden bedding. Twenty-four hours prior testing, all food was removed from the home cage. A single food pellet (regular chow) was adhered on a piece of Whatman paper and positioned in the center of the container that was brightly illuminated. The latency to eat was measured during 10 min as previously described (David et al., 2009). At the end of each session, the animals were immediately transferred into their home cage and the amount of food consumed over the subsequent 5 min period, as well as their body weight, were measured as a control of feeding drive.

Splash test. The test consisted in squirting ~200 µl of a sucrose solution on the dorsal coat of the mouse, as previously described (Surget et al., 2008). Grooming latency, frequency, and duration were assessed during 6 min.

Tail suspension test. Mice were suspended by the tail using adhesive tape on a metal rod of an automated tail suspension apparatus (Bioseb), as previously described (Cryan et al., 2005). Immobility time and power of movement were recorded during a 6 min period.

Forced swim test. The test was performed using an automated forced swim test software (Bioseb), as previously described (Garza et al., 2012). Total immobility time, swimming, and climbing states were measured during a 6 min session.

Olfactometry
Mice were partially water-deprived for 1 week and were then trained using an operant conditioning go/no-go procedure in computer-controlled olfactometers (Fig. 4A) as described previously (Alonso et al., 2008).
2012; www.olfacto-meter.com). In this paradigm, mice were trained to respond to the presence of a rewarded stimulus odorant (S⁺) by licking the water delivery tube, and to refrain from responding to the presence of non-rewarded stimulus odorant (S⁻). A response in an S⁺ trial and an absence of response in an S⁻ trial were scored as correct. Accuracy was scored for each block of 20 trials. Mice were subjected to a daily session of eight blocks. Monomolecular odorant compounds and binary odorant mixtures (Sigma-Aldrich), were diluted either in water or in mineral oil (MO), and 10 ml of the resulting solution was used as the odorant source.

Olfactometry was performed in the following order: (1) training and discrimination of isoamyl-acetate (IAA) versus MO (1st and 2nd weeks), (2) odorant (N-butanol) detection threshold (3rd and 4th weeks), and (3) carvone and carvone mixture discrimination (5th and 6th weeks). Olfactory memory was assessed in one session following a 4 week interval.

In the odorant detection threshold test, mice were tested for their ability to detect successively descending decimal concentrations of N-butanol (S⁻) diluted in water (S), starting with a concentration of 0.01%. The session ceased at the achievement of criterion performance (≥75% of correct responses in at least one block). The odorant detection threshold test ceased when the criterion performance was not achieved after three successive sessions with the same odorant dilution. The accuracy criterion (≥75% of correct responses in at least 1 block) was used to calculate odorant detection threshold.

In the odorant mixture discrimination tasks, mice were trained to distinguish between two carvone enantiomers diluted in MO: 0.1% carvone⁺ (S⁺) and 0.1% carvone⁻ (S⁻). Carvones belong to the terpenoid family and are food scents. S⁺-carvone smells like caraway (Carum carvi) and Anethum graveolens, whereas S⁻-carvone smells like spearmint. Mammals, including humans, are able to spontaneously discriminate between the two carvone enantiomers (Linner et al., 2002). Mice were then tested in two discrimination tasks including carvone mixtures. In the first task, S⁻ was attributed to a solution of 80% carvone⁻ and 20% carvone⁺, and S⁺ to a solution of 20% carvone⁻ and 80% carvone⁺. In the second task, the S⁻ odorant was a solution of 60% carvone⁻ and 40% carvone⁺, whereas S⁺ was a solution of 40% carvone⁻ and 60% carvone⁺).

For the olfactory memory task, mice underwent eight blocks of 20 trials every day for 5 consecutive days, to recall distinguishing between...
carvone− and carvone+. Then, mice were tested on the same task after a rest period of 4 weeks, in one block of 20 trials, but with no reward given for a correct response.

Bromodeoxyuridine injections. Mice were intraperitoneally injected with a DNA synthesis marker, bromodeoxyuridine (BrDU; 100 mg/kg, Sigma-Aldrich). They received one injection 2 h before perfusion for the analysis of cell proliferation, or four injections, at 2 h intervals, on a single day, 4 weeks before perfusion, for the analysis of cell survival.

Stereotaxic lentiviral injections
To study the morphology of adult-born neurons, a replication-deficient lentiviral vector expressing green fluorescent protein (GFP) under control of the cytomegalovirus (CMV) promoter (LV-CMV-GFP with a viral titer of 109 units/ml) was used as previously described (Lazareni et al., 2014). The concentration of lentiviral vector particles used for injection was 13.8 ng of viral antigen per microliter assayed by ELISA (10–12 transduction units per milliliter). The LV-CMV-GFP vector was prepared at the ENP vector platform (www.paris-neuroscience.fr). Mice were deeply anesthetized by a mixture of ketamine and xylazine (100 mg/kg ketamine, 16 mg/kg xylazine). Small craniotomies were performed above the injection sites using a dentist drill. We performed bilateral virus injections (200 nl per site) into the rostral migratory stream (RMS) at +3.30 mm anteroposterior and −0.82 mm mediolateral from bregma, and −2.90 mm dorsoventral from skull surface, to label SVZ-generated neuroblasts that migrate toward the OB. Mice were then transferred to their home cage for recovery.

Immunohistochemistry
Mice were deeply anesthetized with sodium pentobarbital (i.p., 100 mg/kg, Sanofi) and perfused transcardially with a solution containing 0.9% NaCl and heparin (5 × 103 U/ml, Sanofi-Synthelabo) at 37°C, followed by 4% paraformaldehyde in phosphate buffer, pH 7.3. Olfactory epithelia (OE) and brains were removed and postfixed by incubation in the same fixative at 4°C overnight. OE were cryoprotected by incubation in 30% sucrose in PBS for 24 h, and then embedded in Tissue-tek (Sakura). OE and brain slices and fixed sections for OE. Nonspecific staining was blocked by 0.2% Triton, 4% bovine serum albumin (Sigma-Aldrich). Immunoperoxidase-labeled cells were detected with a peroxidase-conjugated secondary antibody (ABC system, Vector Laboratories) at room temperature. Primary antibodies used in this study and their working dilutions are listed in Table 1.

For doublecortin (DCX) immunolabeling, brain sections were preincubated in citrate buffer (10 mM, pH 9.0) at 80°C for 20 min. For Ki67 immunostaining, OE sections were preincubated in citrate buffer (10 mM, pH 6.0) at 80°C for 20 min. For BrDU staining, DNA was denatured with 2N HCl at 37°C for 30 min. Fluorescent sections were stained with the nuclear dye 4′,6′-diamidino-2-phenylindole (DAPI) and then mounted in a solution of 1,4-diazabicyclo[2.2.2]octane (Sigma-Aldrich). Immunoperoxidase-labeled cells were detected with a peroxidase-conjugated secondary antibody (ABC system, Vector Laboratories), using biotinylated donkey anti-rabbit, goat anti-rabbit or horse anti-goat IgG (1:200, Vector Laboratories) and 3,3′-diaminobenzidine (0.05%) as a chromogen (Sigma-Aldrich).

For morphologic analysis of adult-born neurons in the OB, the labeling was enhanced with a GFP immunostaining (rabbit anti-GFP primary antibody, 1/1000, and goat anti-rabbit secondary antibody, 1/500, Invitrogen). DAPI (1/5000) was used as a nuclear stain.

Image acquisition and quantification analysis
Immunofluorescence was analyzed using a confocal laser-scanning microscope (LSM 710; Zeiss) with Zen Imaging software (Zeiss) and a confocal spinning disk Cell Voyager (CV1000, Yokogawa). For the analysis of immunoperoxidase-labeled cells, reconstructed images of the dentate gyrus (DG), SVZ, and OB were taken using an Olympus BX51 microscope (20× objective) and the Compix Imaging software (Hamamatsu Photonics). Quantification was performed using the Icy open source platform (http://www.icy.bioimageanalysis.org; Olivo-Marín, 2002; de Chau-mont et al., 2012). The internal and external borders of the RMS, the glomerular layer (GL), and granule cell layer (GCL) of the OB were

Figure 3. Chronic FLX affects anxiety/depression-like states and impairs olfaction. A, Diagram showing experimental groups and study design. B, When administered alone, chronic FLX generated some anxiety and depression-related behaviors in otherwise naive mice, mainly an increase in feeding latency in the novelty suppressed feeding paradigm. C, Chronic FLX impaired olfactory acuity since FLX-treated mice required significantly higher concentrations of n-butanol for odorant detection compared with vehicle-treated animals. Mice treated with FLX displayed a poorer learning rate compared with vehicle-treated animals in the 80/20 vs 20/80 carvone mixture task, albeit reaching the same final score. D, In the more difficult discrimination task of 60/40 versus 40/60 carvone mixtures, FLX-treated mice displayed a weaker learning rate as well as a lower final score. E, To assess olfactory memory, mice were trained to distinguish between carvone− and carvone+ during 5 consecutive days. Mice were then retested for the carvone discrimination task at 30 d (t 30) following the end of the training session (t 0). FLX-treated animals exhibited an olfactory memory deficit. Values are plotted as mean ± SEM (n = 8–10/group). *p < 0.05, **p < 0.01.
delineated and DCX/H11001 and BrdU/H11001 cells were counted manually using the plugin “Manual counting” (http://icy.bioimageanalysis.org/plugin/manual_counting). Regions-of-interest were designed with the ring plug-in (http://icy.bioimageanalysis.org/plugin/Ring ROI). Values are expressed as the mean of total DCX/H11001 or BrdU/H11001 cell count in six OB slices per animal.

For the quantification of DCX immunoreactivity (IR) in the SVZ and olfactory marker protein (OMP) IR in glomeruli of the OB, pixel density was computed using the plugin “Thresholded pixel density.” This plugin computes the number of pixels over a threshold in a given region of interest over the total number of pixels present in the region of interest (http://icy.bioimageanalysis.org/plugin/Thresholded_pixel_density). For OMP

Table 1. Detailed information on the primary antibodies used in this study

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Manufacturer</th>
<th>Marker</th>
<th>Working dilutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>BrdU</td>
<td>Abcsy, ABC 117-7513</td>
<td>Proliferating cells</td>
<td>1:1000</td>
</tr>
<tr>
<td>DCX</td>
<td>Abcam, ab 18723</td>
<td>Neuroblasts</td>
<td>0.5 μg/ml</td>
</tr>
<tr>
<td>GFP</td>
<td>Molecular Probes</td>
<td>LV-CMV-GFP</td>
<td>1:1000</td>
</tr>
<tr>
<td>K67</td>
<td>Abcam, ab16667</td>
<td>Proliferating cells</td>
<td>1:500</td>
</tr>
<tr>
<td>NeuN</td>
<td>Millipore Biscience Research Reagents,</td>
<td>Neurons</td>
<td>5 μg/ml</td>
</tr>
<tr>
<td>OMP</td>
<td>MAB377</td>
<td>Olfactory sensory neurons</td>
<td>1:2000</td>
</tr>
<tr>
<td></td>
<td>Wako, 544-10001</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 4. Chronic CORT induces multiple olfactory deficits most of which are restored by FLX. A, Go/no-go procedure. In response to a positive odor stimulus (S+), the trained mouse licks the water port to receive the water reward. B, Graph showing the mean percentage of correct responses in each block of the discrimination task between 1% IAA and 1% MO. All groups of mice reached criterion performance, which corresponds to 90% of correct responses in at least one block. A score of 50% corresponds to the success rate at chance level. C, Graph depicting the mean percentage of correct responses in each block of the discrimination task between two carvone enantiomers and their mixtures. To further evaluate olfactory discrimination, we trained mice to discriminate mixtures of increasing complexity. Mice were first trained to distinguish 1% carvone− and 1% carvone+. Once mice reached this criterion, they were subjected to the more difficult task of mixture discrimination. The mixture ratio of carvone− and carvone+ is indicated on the graph. All groups showed similar performance in the simple discrimination task of carvone− versus carvone+. However, the ability to discriminate between carvone mixtures was reduced by CORT and not restored by FLX. D, CORT-treated mice required significantly higher concentrations of N-butanol for odorant detection compared with vehicle-treated and CORT + FLX-treated animals. E, To assess olfactory memory, mice were trained during 5 consecutive days to recall distinguishing carvone− and carvone+. Mice were then retested in the simple carvone discrimination task at 30 d following the end of the training session (t0). CORT-treated animals exhibited a memory deficit that was attenuated by FLX. Values are plotted as mean ± SEM (n = 8 for vehicle and CORT-treated; n = 35 for CORT + FLX). *p < 0.05, **p < 0.01.
IR, six glomeruli of the dorsomedial part of OB coronal sections were selected and assessed for each animal. The batch mode of this analysis is available as a protocol [http://icy.bioimageanalysis.org/protocol/Pix-el__density_](http://icy.bioimageanalysis.org/protocol/Pix-el__density_ [batch_mode]). All the codes and implementation of these programs are available online. For the quantification of Ki67+ cells in the OE, serial sections of the olfactory mucosa situated through the anterior end of the OB were analyzed blindly. Four to six serial sections were chosen per animal (5 animals analyzed per group, 6–7 zones analyzed per OE section).

For hippocampal analysis, values are expressed as the mean total number of DCX+ or BrdU+ cells counted manually in 8 DG sections per animal. For BrdU and neuronal nuclei (NeuN) double-staining analysis, positive cells were counted in the GCL (6 slices per animal, 4 mice analyzed per group, 20–30 BrdU+ cells analyzed) and in the DG (8 slices per animal, 4 mice analyzed per group, 20–30 BrdU+ cells analyzed). Data are expressed as the number of double-labeled cells in the GCL and DG.

For the analysis of dendritic morphology of adult newborn OB neurons, 8 coronal sections of a 180 μm distance were selected, taking the accessory OB as a landmark. Fifteen neurons (located ≥40 μm in the GCL) were selected for each animal. These neurons were imaged using a confocal microscope (Zeiss LSM 700, 25× objective, 0.5–1 numerical zoom). The dendritic tree was manually traced using the Filament tracer module of the Imaris Bitplane software v7.6. Total dendritic length, number of branches and Sholl data were collected. For Sholl analysis, virtual concentric circles were traced at every micrometer starting from the beginning of the dendrite, and the number of intersections between the dendritic tree and each circle were counted as previously described (Palotto et al., 2012). For the analysis of dendritic morphology of hippocampal adult-born neurons, the maturation index was calculated as previously described (Mendez-David et al., 2014). DCX+ cells were subcategorized according to their dendritic morphology as DCX+ cells without and DCX+ cells with tertiary (or higher order) dendrites. The maturation index was defined as the ratio of DCX+ cells possessing tertiary dendrites to the total number of DCX+ cells.

**Statistical analysis**

All data were expressed as mean ± SEM. Statistical analyses were performed using the Prism software (GraphPad v6), with p < 0.05 considered statistically significant. Data were analyzed using unpaired two-tailed Student’s t test, one-way ANOVA or two-way ANOVA followed by Bonferroni post hoc test or Mantel-Cox log-rank test when appropriate.

**Results**

**Chronic CORT treatment induces anxiety- and depression-like states in mice**

Using standard tests for anxiety and depression, we confirmed that chronic treatment with CORT triggers an anxiety/depression-like phenotype that is responsive to chronic FLX treatment. At 4 weeks of treatment, CORT decreased the time spent in the open field (185 ± 12 s vs 333 ± 43 s, t = 4.53, p < 0.001, unpaired t test; Fig. 1A), and in the open arms of the elevated plus maze (39 ± 3 s vs 58 ± 11 s, t = 2.19, p = 0.03, unpaired t test; Fig. 1B). In the novelty suppressed feeding test, CORT significantly increased feeding latency (359 ± 23 s vs 97 ± 12 s, p < 0.001, Kaplan–Meier survival analysis, Mantel-Cox log-rank test; Fig. 1C) without affecting feeding drive (Fig. 1D), whereas in the splash test CORT increased grooming latency (207 ± 17 s vs 55 ± 12 s, p < 0.001, Kaplan–Meier survival analysis, Mantel-Cox log-rank test; Fig. 1E) and decreased grooming frequency (2 ± 0.2 s vs 5 ± 0.8, t = 4.78, p < 0.001, unpaired t test; Fig. 1F).

At 16–17 weeks of CORT treatment (Fig. 2A), feeding latency in the novelty suppressed test remained significantly increased (470 ± 52 s vs 80 ± 15 s, p < 0.001, Kaplan–Meier survival analysis, Mantel-Cox log-rank test; Fig. 2B) without changes in feeding drive (vehicle: 0.29 ± 0.05; CORT: 0.25 ± 0.06; CORT + FLX: 0.23 ± 0.07; F(2,44) = 0.19, p = 0.1, one-way ANOVA, Bonferroni post hoc test; Fig. 2C). Moreover, changes in grooming behavior persisted, namely decreased grooming frequency (3 ± 0.3 vs 6 ± 0.7; F(2,44) = 7.09, p = 0.03, one-way ANOVA, Bonferroni post hoc test; Fig. 2D) and increased latency to groom (116 ± 18 s vs 29 ± 7 s, p < 0.001; Kaplan–Meier survival analysis, Mantel-Cox log-rank test; Fig. 2E). CORT also increased immobility in the forced swim test (119 ± 18 s vs 230 ± 37 s; F(2,44) = 4.13, p = 0.02, one-way ANOVA, Bonferroni post hoc test; Fig. 2F) and in the tail suspension test, albeit in a non-significant manner (Fig. 2G). Chronic FLX exerted antidepressant effects that were mainly prominent in the novelty suppressed feeding and splash tests. More specifically, FLX restored feeding latency (176 ± 34 s, p < 0.001 vs CORT, one-way ANOVA, Bonferroni post hoc test), as well as grooming frequency (6 ± 0.5, p = 0.002 vs CORT, one-way ANOVA, Bonferroni post hoc test) and latency to groom (27 ± 4 s, p < 0.001 vs CORT, Kaplan–Meier survival analysis, Mantel-Cox log-rank test; Fig. 2B,D,E). When administered alone for 10–11 weeks (Fig. 3A), chronic FLX generated some anxiety-related behaviors in otherwise naive mice, reflected mainly by an increase in feeding latency in the novelty suppressed feeding test (vehicle: 81 ± 13 s; FLX: 212 ± 42 s, t = 3.3, p = 0.005, unpaired t test; Fig. 3B). Together, these data indicate that chronic CORT exposure induces an anxiety/depression-like state in mice that is reversed by prolonged coadministration of FLX.

**Olfactory discrimination is impaired by chronic CORT and not restored by FLX**

To test whether the observed anxiety/depression-like state was accompanied by olfactory deficits, we first evaluated olfactory discrimination by using an automated operant conditioning procedure (a go/no-go test; Fig. 4A). Mice were trained to recognize IAA as the stimulus rewarded with water (S+) and MO as the non-rewarded stimulus (S−; Fig. 4B). In the next step, mice were trained to distinguish between two carvone enantiomers diluted in MO; carvone+ (S+) and carvone− (S−); and we then assessed fine olfactory discrimination using binary mixtures of the same carvone enantiomers (Fig. 4C). We observed that CORT and CORT + FLX did not affect acquisition and success rates in the simple discrimination task of IAA versus MO and in the carvone enantiomer task (Fig. 4B,C). However, in the “difficult” mixture discrimination task, the performance of CORT-treated animals was significantly poorer compared with controls and was not restored by FLX (two-way repeated measures ANOVA, 8/2 mixture ratio task: treatment factor F(2,44) = 5.04, p = 0.01; 6/4 mixture ratio task: treatment factor F(2,44) = 5.43, p = 0.01; Fig. 4C).

**Olfactory acuity is impaired by chronic CORT and restored by FLX**

Mice were trained to detect successively descending decimal concentrations of N-butanol (S−) diluted in water (S+), starting with a concentration of 0.01%. CORT decreased the detection threshold by two orders of magnitude (Fig. 4D; mean odorant dilution (−log); vehicle: 8.2 ± 0.4; CORT: 6.5 ± 0.3; F(2,44) = 4.23, p = 0.02, one-way ANOVA, Bonferroni post hoc test). Co-treatment with FLX attenuated the CORT-induced deficit (Fig. 4D; mean odorant dilution (−log); 7.7 ± 0.3, p = 0.03, one-way ANOVA, Bonferroni post hoc test).
Olfactory memory is affected by chronic CORT and improved by FLX
We assessed long-term olfactory memory by using the same pair of carvone isomers (Fig. 4E). Thirty days following task consolidation, the ability of mice to recognize carvone+ and carvone− enantiomers was re-evaluated, using one block of 20 trials that was devoid of water reward. Four weeks following task consolidation, the performance of CORT-treated mice was near chance level, revealing that chronic CORT impaired olfactory memory [mean correct response; vehicle: 72.5%; CORT: 50.6%; p = 0.002, two-way ANOVA; F(2,44) (treatment × memory) = 7.22]. Treatment with FLX was able to improve performance (mean correct response; 61.3%, p < 0.05 compared with CORT).

Olfaction is impaired by chronic FLX
Although FLX restored most CORT-induced olfactory deficits, it significantly impaired olfaction when administered alone chronically (Fig. 3C–F). FLX induced a substantial decrease in olfactory acuity [mean odorant dilution (−log); vehicle: 6.5 ± 0.2; CORT: 4.1 ± 0.8; t = 3.28, df = 16, p = 0.003, unpaired t test; Fig. 3C]. Fine discrimination was also affected (Fig. 5D,E), and this effect was most prominent in the 6/4 versus 4/6 carvone mixture task (two-way ANOVA; F(1,16) treatment = 8.59, p = 0.01; Fig. 3E). Long-term olfactory memory was also significantly impaired by FLX (two-way ANOVA; F(1,32) interaction = 16.45, p = 0.003; Fig. 3F).

Adult neurogenesis is impaired by chronic CORT and restored by chronic FLX
We then investigated whether adult neurogenesis was affected by CORT. We measured the production of new neurons in the SVZ and DG and their survival rate in the OB and DG. Quantitative analysis of the population of adult-born neurons at 20 weeks of CORT and 14 weeks of CORT+FLX treatment showed a significant twofold decrease in the number of DCX+ neurons in the DG that was reversed by FLX (mean cell number; vehicle: 4782 ± 115; CORT: 1879 ± 286; CORT+FLX: 7439 ± 910 cells; F(2,18) = 7.06; CORT vs vehicle: p = 0.04; CORT vs CORT+FLX: p = 0.002; one-way ANOVA, Bonferroni post hoc test; Fig. 5A,D). On the other hand, neither CORT nor CORT+FLX altered the number of DCX+ in the SVZ (Fig. 5B,F). These results indicate that CORT affects differentially the two stem cell niches of the adult brain, reducing neural production in the DG but sparing SVZ neurogenesis. Yet, we found that the number of DCX− neurons in the core of the OB, the RMS,oh, was decreased in the CORT group, and this effect was reversed by FLX (vehicle: 15,834 ± 1049; CORT: 10,316 ± 310; CORT+FLX: 15,590 ± 1332 cells; F(2,18) = 5.78; CORT vs vehicle: p = 0.04; CORT vs CORT+FLX: p = 0.03, one-way ANOVA, Bonferroni post hoc test; Fig. 5C,H). Moreover, the number of DCX+ cells was notably reduced by CORT and restored by FLX in the GCL (vehicle: 29,292 ± 1159; CORT: 15,876 ± 355; CORT+FLX: 29,545 ± 4557 cells; F(2,18) = 4.02; CORT vs vehicle: p = 0.05; CORT vs CORT+FLX: p = 0.05, one-way ANOVA, Bonferroni post hoc test; Fig. 5C,G) and in the GL (vehicle: 2491 ± 64; CORT: 1387 ± 158; CORT+FLX: 2476 ± 282; F(2,18) = 5.72; CORT vs vehicle: p = 0.04; CORT vs CORT+FLX: p = 0.03, one-way ANOVA, Bonferroni post hoc test; Fig. 5I). Therefore, it is likely that CORT affects olfactory adult neurogenesis by altering neuronal survival in the OB, while leaving intact cell production in the SVZ.

To address this question more directly, we examined both cell proliferation and survival in the DG and OB, by measuring BrdU+ cells 2 h and 1 month following BrdU injection, respectively (Fig. 6A). BrdU+ cell counting confirmed that CORT significantly decreases cell proliferation in the DG, whereas FLX completely restores it (mean BrdU+ cell number; vehicle: 723 ± 60; CORT: 414 ± 58; CORT+FLX: 985 ± 126 cells; F(2,18) = 5.76; CORT vs vehicle: p = 0.05; CORT vs CORT+FLX: p = 0.02, one-way ANOVA, Bonferroni post hoc test; Fig. 6B,F). However, neither CORT nor CORT+FLX altered the number of BrdU+ in the SVZ (Fig. 6G).

Moreover, CORT induced a twofold decrease in the survival of newborn cells in the DG (BrdU+ cell number; vehicle: 997 ± 123 cells; CORT: 523 ± 60 cells; F(2,18) = 1.17, p = 0.05, one-way ANOVA, Bonferroni post hoc test; Fig. 6I) and the GCL (BrdU− cell number; vehicle: 9082 ± 1212 cells; CORT: 4978 ± 475 cells; F(2,18) = 4.91, p = 0.02, one-way ANOVA, Bonferroni post hoc test; Fig. 6F). FLX reversed this effect albeit in a nonsignificant manner. Likewise, cell survival was reduced by CORT in the OB, both in the GL (BrdU+ cell number; vehicle: 97 ± 123 cells; CORT: 523 ± 60 cells; F(2,18) = 1.17, p = 0.05, one-way ANOVA, Bonferroni post hoc test; Fig. 6J) and the GCL (BrdU− cell number, 985 ± 101 cells, p = 0.03 vs CORT; one-way ANOVA, Bonferroni post hoc test). Finally, BrdU and NeuN colabeling showed that the vast majority of BrdU+ cells were also colabeled with the neuronal biomarker NeuN in all treatment groups tested (Fig. 6E,K). In the GCL of OB, analysis of 80–120 cells per animal (n = 4 mice) in the vehicle group showed that ~90% BrdU+ cells coexpressed NeuN and thus had a neuronal phenotype. We found that the percentage of BrdU+ cells labeled with NeuN was not modified by CORT and CORT+FLX treatment (Fig. 6K; vehicle: 92.5 ± 1.5%; CORT: 92.8 ± 0.9%; CORT+FLX: 91.1 ± 0.6%). Moreover, we found no group difference in BrdU/NeuN double staining in the DG (20–30 cells per animal; vehicle: 91.1 ± 0.5%; CORT: 87.7 ± 3.1%; CORT+FLX: 91.8 ± 0.7%). Together, these data indicate that chronic CORT exposure affects both adult hippocampal and olfactory neurogenesis and that chronic FLX can stimulate the recruitment and/or survival of newborn neurons within the DG and OB.

Chronic CORT treatment does not alter adult neurogenesis in the olfactory epithelium
Given the prominent effects of CORT on olfactory function, we checked whether the olfactory sensory organ could be impaired. Olfactory sensory neurons located in the OE project via the olfactory nerve to glomeruli in the GL of the OB, where they synapse on the dendrites of output mitral/tufted cells. Olfactory sensory neurons are continually replaced throughout life by the progeny of dividing basal cells located in the OE. We quantified the level of cell proliferation in the OE by counting the number of Ki67− cells in this zone. Anti-Ki67 immunostaining is mainly localized in the basal area of the normal OE (Fig. 7A,B). We found no difference in the distribution and the number of Ki67− cells between vehicle and CORT-treated mice following 4 weeks of treatment [mean number of Ki67− cells (104) per mm; vehicle: 8.09 ± 0.23; CORT: 8.55 ± 0.65, unpaired t test; Fig. 7C]. Staining for OMP, a cell marker specific of mature olfactory sensory neurons, was quantified in the GL (Fig. 7E). We found no difference in the distribution and intensity of OMP between vehicle, CORT and CORT+FLX-treated mice (mean IR within glomeruli; vehicle: 0.1498 ± 0.02; CORT: 0.1803 ± 0.01; CORT+FLX: 0.1503 ± 0.01; one-way ANOVA, Bonferroni post hoc test; Fig. 7F). These results indicate that cell proliferation in the OE and the degree of sensory innervation in the OB are both insensitive to chronic CORT treatment.
Impact of CORT and FLX on the morphology of newborn interneurons

Dendritic complexity of adult-born DG neurons increases during the first month following neuronal birth, a stage during which cells express the DCX marker. To further examine the effects of CORT on dendritic morphology of newborn neurons, we characterized DCX+ cells in the DG according to the number of dendrites, and calculated the maturation index as the ratio of DCX+ cells with tertiary (or higher order) dendrites to the total number of DCX+ cells. Although we found no effect of chronic CORT on the maturation index of adult-born DG neuroblasts, we observed a significant threefold increase induced by CORT+FLX (mean maturation index; vehicle: 0.20 ± 0.02; CORT: 0.21 ± 0.03; CORT+FLX: 0.55 ± 0.09; $F_{(2,12)} = 0.89$; CORT vs CORT + FLX: $p = 0.007$, one-way ANOVA, Bonferroni post hoc test; Fig. 5E).

DCX is also expressed by adult-born OB interneurons for <10 d after birth but cannot be used for the assessment of dendrite complexity due to the large staining in the OB. Thus, to assess the impact of chronic CORT and FLX on the maturation of newborn OB interneurons, a lentiviral vector expressing the GFP was injected into the RMS following 8 weeks of CORT and 4 weeks of CORT+FLX or FLX alone (Fig. 8A, C). Three weeks after injection, we analyzed the apical dendritic tree of GFP+ adult-born granule cells in the OB and found that chronic CORT did not alter the number of dendrites (Fig. 8D) nor total dendritic length (Fig. 8E). However, CORT+FLX and FLX alone increased both the number of dendrites (vehicle: 4.93 ± 0.29; CORT: 4.92 ±
Figure 6. Adult-born neuron proliferation and survival is decreased by chronic CORT and restored by FLX. A, Diagram depicting BrdU injection protocols for the evaluation of cell proliferation and survival. B, C, Representative coronal images of BrdU + cells in (B) the DG of the hippocampus (scale bar, 100 μm), and (C) the SVZ (scale bar, 100 μm) 2 h following BrdU administration. D, Representative images showing BrdU + cells in the GCL of the OB of vehicle, CORT and CORT + FLX-treated mice, one month after BrdU administration. Scale bar, 15 μm. E, BrdU (red) and NeuN (green) colabeled cells in the GCL of vehicle and CORT-treated mice, one month after BrdU administration. Scale bar, 15 μm. F, G, Graphs depicting the effects of CORT and CORT + FLX on cell proliferation in (F) the DG and (G) the SVZ (n = 5–10/group). H–J, Graphs showing the effect of CORT and CORT + FLX on cell survival in (H) the DG, (I) the GL of the OB, and (J) the GCL of the OB (n = 5–10/group). K, Graph showing the percentage of BrdU + cells labeled with NeuN in the GCL of vehicle, CORT and CORT + FLX-treated mice 1 month following BrdU administration (n = 4–5 animals per group, 80–120 cells analyzed per group). Values are plotted as mean ± SEM. *p < 0.05.

0.25; CORT + FLX: 5.66 ± 0.36; FLX: 5.84 ± 0.34; F(3,16) = 0.46; vehicle vs FLX: p = 0.05; one-way ANOVA, Bonferroni post hoc test; Fig. 8D, E). Sholl analysis revealed that the most significant changes in the number of branches were obtained by CORT + FLX, suggesting a cumulative effect of the two treatments (two-way ANOVA; F(3,155) = 7.36, p < 0.001; Fig. 8F). Finally, we observed no effect of CORT and FLX on spine density and length (data not shown). Together, these results sug-
gest that FLX, but not CORT, facilitates the maturation of newborn neurons in both the adult hippocampus and OB circuits.

**Discussion**

This study reports that a mouse model that mimics states of HPA axis hyperactivity, a common trait of MDD, is associated with impaired adult neurogenesis and pronounced olfactory deficits, including acuity, discrimination, and memory. We found that prolonged antidepressant treatment with fluoxetine not only restored adult neurogenesis and mood, but also improved most olfactory functions. Our data identify a key role for the HPA axis in olfactory impairments associated with anxiety/depressive states.

**Deleterious impact of chronic CORT on adult neurogenesis and olfaction**

Our present findings demonstrate that the CORT model not only evokes a persistent fear-related response inhibition, reminiscent of anxiety/depression-like states, but also induces olfactory dysfunction. Importantly, similar olfactory deficits affecting odorant detection and discrimination have been reported in MDDs (Pause et al., 2001; Negoias et al., 2010). In turn, perturbations of the olfactory circuit and function can induce anxiety and depression-like behaviors in rodents (Glinka et al., 2012; Lazarini et al., 2012). Thus, we cannot rule out the possibility that CORT-induced olfactory deficits could be in part responsible for the mood states observed in this model. For instance, olfactory bulbectomy constitutes a model of depression and is used as a predictor of antidepressant drug response in rodents (Wren et al., 1977). Although the precise mechanisms remain elusive, it can be speculated that such sensory deficits likely affect the function of neural circuits involved in the control of mood and emotions.

Interestingly, we also found that chronic CORT significantly reduces neurogenesis in the OB. The number of new OB interneurons was decreased twofold following CORT treatment in all OB layers. This decrease could partly explain the CORT-induced olfactory deficits, because adult-born OB interneurons take part in odorant detection, discrimination, olfactory learning, and memory (Breton-Provencher et al., 2009; Moreno et al., 2009; Mouret et al., 2009; Belnoue et al., 2011; Lazarini et al., 2011; Alonso et al., 2012; Sakamoto et al., 2014). However, it is important to note that in humans, MDD-related hyposmia is unlikely to be due to reduced neurogenesis in the OB, as cells

**Figure 7.** Chronic CORT treatment does not affect sensory afferentation to the OB. A, B, Immunostaining with the Ki67 antibody in a normal 8-week-old mouse olfactory mucosa. B, A magnification of the region delineated by a red rectangle in A. Scale bars: A, 500 μm; B, 50 μm. C, Four weeks of CORT administration did not change the number of Ki67+ cells in the OE (n = 5 mice per group). D, OB coronal section of a vehicle-treated mouse. Scale bar, 100 μm. E, Representative images of OMP (green) and DAPI (blue) staining in the GL following 20 weeks of CORT and 14 weeks of CORT + FLX treatment. Scale bar, 50 μm. The dotted line delineates one glomerulus. F, CORT and CORT + FLX had no effect on OMP IR in the OB (values are plotted as mean ± SEM, n = 4 mice per group; 6 glomeruli in the dorsomedial GL analyzed per animal). au, Arbitrary units.
born in the adult SVZ are mostly directed to the striatum and not to the OB (Ernst et al., 2014). Moreover, one cannot rule out the possibility that the CORT-induced hyposmia revealed in this study could be partly linked to a deleterious effect of CORT on preexisting granule and periglomerular cells in the OB. This could also be the case in human patients suffering from MDD. Finally, because glucocorticoid receptors are widely distributed in the OB and olfactory cortex, as in other limbic areas, notably the hippocampus, amygdala, and hypothalamus (Morimoto et al., 1996), it is possible that CORT-induced changes in overall

Figure 8. Dendritic length and complexity are increased by chronic CORT and FLX treatments. A, Experimental timeline of viral injections and analysis. B, Confocal images of vehicle, CORT, and CORT + FLX-treated OB slices showing GFP+ neurons and DAPI staining. Scale bar, 20 µm. C, Lentiviral vectors encoding GFP under the control of the CMV promotor were injected in the RMS. D, FLX treatment alone increased the number of dendrites in newborn OB neurons. E, CORT + FLX and FLX treatments significantly increased dendritic length. F, Graph depicting the effect of chronic CORT, CORT + FLX and FLX treatments on the number of dendritic branches, according to Sholl analysis. CORT + FLX treatment increased the number of branches. Values are plotted as mean ± SEM (n = 4/group). *p < 0.05, **p < 0.01, ***p < 0.001 in CORT versus vehicle; #p < 0.05, ##p < 0.01 in CORT versus CORT + FLX; §p < 0.05 in vehicle versus FLX.
cortical and limbic processing could be linked to the observed olfactory deficits.

Our results also showed that chronic CORT impairs both pro-
life and survival of progenitor cells in the DG, demonstrat-
ing a role for glucocorticoids in several steps of the neurogenic
process. Our data are in line with previous work reporting that
ablation of the adrenal glands abolishes the stress-induced de-
crease in cell proliferation in the DG (Tanapat et al., 2001). Al-
though CORT reduced cell proliferation in the DG, it did not
affect cell proliferation in the OE nor the number of newly gen-

erated immature neurons expressing DCX in the SVZ. Our re-

sults indicate that the early steps of adult neurogenesis are
differentially regulated by CORT in the DG, EO, and SVZ. How-
ever, given the decrease in the population of adult-born neurons
in the OB, it is possible that CORT treatment alters the survival
rate of adult-born OB neurons. Interestingly, their dendritic
morphology was not changed by CORT, suggesting that the sur-

viving neurons are functionally integrated in the OB circuit. This
hypothesis is also supported by the fact that the distribution and
degree of the sensory afferents reaching the OB was unchanged.

Concerning the adult olfactory system, the functional impor-
tance of CORT remains largely unknown. New experiments aim-
ning to assess firing activity, dendritic dynamics, and innate
responses to odorants are needed to decipher the precise role of
CORT in olfactory processing and odor-driven behaviors.

Chronic FLX treatment attenuates CORT-induced olfactory deficits and restores adult neurogenesis

Our findings show that the antidepressant FLX can restore olfa-
tory acuity and memory in CORT-treated mice. Interestingly, we
also showed that FLX restores olfactory neurogenesis and facili-
tates the maturation of adult-born neurons in the OB and hip-
pocampus. This proencephalic effect could partly support the
FLX-induced restoration of olfactory deficits. Similarly, recruit-
ment of newborn neurons was recently reported to selectively
restore circuits in the lesioned OB and resume olfaction (Lazarini et al., 2014).

FLX restored both proliferation and survival of progenitor
cells in the DG, as previously reported (David et al., 2009; Rainer et al., 2012; Guilloux et al., 2013). By restoring hippocampal neu-
rogenesis, FLX probably helps reestablish HPA axis function, re-
sulting in functional improvement. In fact, it is argued that newly
generated neurons may facilitate a dysfunctional hippocampus to
restore control on stress response systems, and that the ability of
FLX to restore hippocampal regulation of the HPA axis under
chronic stress occurs only in the presence of an intact neurogenic
niche. Further experiments aiming to selectively block either DG
or SVZ neurogenesis in our model are required to disambiguate
the role of each site of adult neurogenesis in the response to FLX
and their associated behavioral rehabilitation.

Finally, to explain the olfactory deficits induced by FLX alone,
one should keep in mind that the OB is one of the most densely
innervated serotonergic targets of the raphe nuclei, and fibers
from the latter are the densest centrifugal projections in the OB
(Dugué and Mainen, 2009). Serotonin alters synaptic transmis-
sion and information processing in the glomerular layer of the
OB, where the density of serotonergic fibers from the raphe nu-
clei is maximal. As it has been indicated by early experiments,
local application of serotonin receptor agonists attenuates odor-
evoked olfactory neuron activity (Bloom et al., 1964), whereas
serotonin receptor antagonists boost glomerular activation,
showing that endogenous serotonin suppresses odor input (Pet-
zold et al., 2009). These previous studies could explain the deficits
in olfactory acuity, discrimination, and long-term memory ob-
tained by chronic FLX treatment.

Together, our findings introduce novel effects of chronic
CORT and FLX on olfaction and olfactory neurogenesis in the
mouse. These observations bring further insights on the poorly
known biological underpinnings of anxiodepressive states and
help move toward more effective therapeutic ends targeting neu-
rogenesis, as well as useful diagnostic tools based on olfaction.

References


Anacker S, Zunszain PA, Carvalho LA, Pariente CM (2011) The glucocor-
ticoid receptor: pivot of depression and of antidepressant treatment? Psy-
choneuroendocrinology 36:415–425. CrossRef Medline

Belloue L, Grosjean N, Abrons DN, Koehl M (2011) A critical time window for the recruitment of bulbar newborn neurons by olfactory discrimina-

Bloom FE, Costa E, Salmoiraghi GC (1964) Analysis of individual rabbit olfactory bulb neuron responses to the microelectrophoresis of acetylcho-
line, norepinephrine and serotonin synergists and antagonists. J Pharma-
col Exp Ther 146:16–23. Medline


Breton-Provencher V, Lemasson M, Peralta MR 3rd, Saghatelian A (2009) Interneurons produced in adulthood are required for the normal func-
tioning of the olfactory bulb network and for the execution of selected olfactory behaviors. J Neurosci 29:15245–15257. CrossRef Medline

Campbell S, Marriott M, Nahmias C, MacQueen GM (2004) Lower hip-

Chen H, Pandey GN, Dwivedi Y (2006) Hippocampal cell proliferation reg-


Garza JC, Guo M, Zhang W, Ly XY (2012) Leptin restores adult hippocam-
pal neurogenesis in a chronic unpredictable stress model of depression and reverses glucocorticoid-induced inhibition of GSK-3β/b-catenin sig-
naling. Mol Psychiatry 17:790–808. CrossRef Medline


Siegli et al. • Olfactory Deficits in Depression
tidepressant and anxiolytic potential of the multimodal antidepressant vortioxetine (Lu AA21004) assessed by behavioral and neurogenesis outcomes in mice. Neuropharmacology 73:147–159. CrossRef Medline


