Calorie restriction activates new adult born olfactory-bulb neurones in a ghrelin-dependent manner but acyl-ghrelin does not enhance subventricular zone neurogenesis


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Abstract
The ageing and degenerating brain show deficits in neural stem/progenitor cell (NSPC) plasticity that are accompanied by impairments in olfactory discrimination. Emerging evidence suggests that the gut hormone ghrelin plays an important role in protecting neurones, promoting synaptic plasticity and increasing hippocampal neurogenesis in the adult brain. In the present study, we investigated the role of ghrelin with respect to modulating adult subventricular zone (SVZ) NSPCs that give rise to new olfactory bulb (OB) neurones. We characterised the expression of the ghrelin receptor, growth hormone secretagogue receptor (GHSR), using an immunohistochemical approach in GHSR-eGFP reporter mice to show that GHSR is expressed in several regions, including the OB but not in the SVZ of the lateral ventricle. These data suggest that acyl-ghrelin does not mediate a direct effect on NSPC in the SVZ. Consistent with these findings, treatment with acyl-ghrelin or genetic silencing of GHSR did not alter NSPC proliferation within the SVZ. Similarly, using a bromodeoxyuridine pulse-chase approach, we show that peripheral treatment of adult rats with acyl-ghrelin did not increase the number of new adult-born neurones in the granule cell layer of the OB. These data demonstrate that acyl-ghrelin does not increase adult OB neurogenesis. Finally, we investigated whether elevating ghrelin indirectly, via calorie restriction (CR), regulated the activity of new adult-born cells in the OB. Overnight CR induced c-Fos expression in new adult-born OB cells but not in developmentally born cells, whereas neuronal activity was absent following re-feeding. These effects were not present in ghrelin−/− mice, suggesting that adult-born cells are uniquely sensitive to changes in ghrelin mediated by fasting and re-feeding. In summary, ghrelin does not promote neurogenesis in the SVZ and OB; however, new adult-born OB cells are activated by CR in a ghrelin-dependent manner.

Keywords
calorie restriction, ghrelin, neurogenesis, olfactory bulb, subventricular zone
1 | INTRODUCTION

The generation of new adult-born neurones in the olfactory bulb (OB) continues throughout life and contributes to olfactory memory. The adult OB receives new neurones that originate from divided neural stem/progenitor cells (NSPCs) residing in the subventricular zone (SVZ) adjacent to the lateral ventricles. Following NSPC division, the cells differentiate into immature neuroblasts and migrate along the rostral migratory stream prior to integration with local OB circuitry. This process of adult OB neurogenesis (AOBN) is regulated by several intrinsic and extrinsic factors, including age, exercise, inflammation and glucocorticoids. However, the underlying mechanisms mediating this process are poorly understood.

Within the OB, new adult-born neurones promote olfactory memory and enhance the ability to discriminate distinct odours. AOBN is also important for OB granule cell replacement and tissue maintenance. Olfactory impairment has been reported as a prodromal indicator of several neurodegenerative diseases. For example, deficits in olfactory discrimination (i.e., the ability to distinguish odours) have been described in experimental neurodegenerative animal models and human Parkinson’s disease. Ghrelin, an orexigenic gut hormone produced in response to calorie restriction, acts on the hypothalamus to stimulate the release of growth hormone (GH) and promote meal initiation and food intake. Emerging evidence suggests that acyl-ghrelin may also have important extra-hypothalamic functions, such as increasing olfactory sensitivity and regulating activity in brain regions involved in olfaction and appetitive behaviour.

In the neurogenic niche of the hippocampus, acyl-ghrelin has been shown to increase cell proliferation and the number of new adult-born neurones in adult rodents. The ghrelin receptor, GH secretagogue receptor (GHSR), which is expressed within the dentate gyrus of the hippocampus, mediates the pro-neurogenic effect of calorie restriction (CR), as well as the increase in hippocampal neurogenesis and antidepressant-like effect following P7C3 treatment. Moreover, ghrelin-deficient mice are reported to have impaired cell proliferation in the SVZ that is normalised to wild-type (WT) levels with exogenous acyl-ghrelin treatment. GHSR is the only molecularly identified receptor for ghrelin, mediating the central effects of this hormone on appetite, body weight and energy metabolism. However, it is not known whether GHSR is expressed within the neurogenic niche of the SVZ or whether acyl-ghrelin modulates AOBN. Here, we aimed to determine the expression pattern of GHSR within the SVZ and whether ghrelin modulates AOBN.

In addition, as fasting and feeding increase and decrease olfactory sensitivity, respectively, we aimed to determine whether ghrelin modulates the fasting-induced activation of both new adult-born and developmentally born OB neurones.

2 | MATERIALS AND METHODS

2.1 | Animals and procedures

All experiments involving animals were performed with appropriate ethical approval. Mouse studies were performed at Cardiff University (GHSR-null, ghrelin−/−) and Monash University (GHSR-eGFP). Studies involving rats were performed at the University of Cambridge.

2.2 | Mice

2.2.1 | GHSR-eGFP mice

Adult male GHSR-eGFP reporter mice were housed at room temperature under a 12:12 hour light/dark cycle (lights on 7.00 am) with access to food and water available ad lib. GHSR-eGFP reporter mice (n = 6) were obtained from the Mouse Mutant Regional Resource Center at the University of California Davis (Davis, CA, USA) and the hemizygous mice were back-crossed to C57BL/6J mice. GHSR-eGFP reporter mice were terminally anaesthetised and transcardially perfused with 4% paraformaldehyde (PFA) in 0.1 mol L−1 phosphate-buffered saline (PBS). Whole brains were rapidly removed and post-fixed in ice cold 4% PFA for 24 hours at 4°C before being sunk in 30% sucrose. Finally, brains were transferred to PBS + 0.1% sodium azide (Sigma-Aldrich, St Louis, MO, USA) and stored at 4°C prior to analysis. Brains were frozen using a fine powder of ground-up dry ice and mounted on a sliding sledge freezing microtome (Microm HM 450; Carl Zeiss, Oberkochen, Germany) using Jung’s freezing medium. The thermostat was set to −30°C to ensure brains remained frozen. Thirty-micrometer thick coronal sections cut along the entire rostral-caudal axis (bregma +5.345 mm to −4.08) were collected in a 96-well plate (nunclon surface; Nunc, Rochester, NY, USA) filled with PBS + 0.1% sodium azide and stored at 4°C until required. GHSR-eGFP mouse brains were also collected in a sagittal orientation (bregma +3.925 to −0.20).

2.2.2 | Immunofluorescence for GHSR-GFP

All experiments were performed on free-floating tissue sections at room temperature, unless stated otherwise. A one in six series of coronal or sagittal brain sections was selected (minimum of 10 sections per mouse), transferred into a 24-well culture plate (nunclon surface; Nunc) and washed in PBS (Sigma-Aldrich) three times for 5 minutes each. Tissue sections were then permeabilised in methanol (Thermo Fisher Scientific, Waltham, MA, USA) at −20°C for 2 minutes and washed (as described before) in PBS. Non-specific binding sites were blocked with 5% normal goat serum (NGS) (Sigma-Aldrich) in PBS + 0.1% Triton X-100 (Sigma-Aldrich) (PBS-T) for 1 hour. Excess block was removed and tissue sections incubated with chicken anti-GFP (chicken polyclonal; Ab13970; Abcam, MA, USA), diluted 1:1000 in PBS-T, for 24 hours at 4°C. Primary antibody was omitted from the negative control. Sections were washed...
and incubated in goat anti-chicken Alexa-fluor 488 (goat polyclonal; A11039; Life Technologies, Grand Island, NY, USA), diluted 1:500 in PBS-T, for 30 minutes in the dark. Finally, sections were washed, mounted onto Superfrost+ slides (Thermo Fisher Scientific) and cover-slipped with vectashield (containing 4’,6-diamidino-2-phenylindole) (Vector Laboratories, Inc., Burlingame, CA, USA) before being stored at 4°C. The slides were analysed by laser scanning confocal microscopy (LSM710; Carl Zeiss) and Zen software (2010 edition; Carl Zeiss) after 24 hours. Gimp, version 2.8 was used to prepare tiled images of coronal and sagittal sections (www.gimp.org).

2.2.3 | GHSR-null mice

To assess exogenous acyl-ghrelin regulation of SVZ cell proliferation, homozygous male loxTB-GHSR mice (GHSR-null) (a gift from Professor Jeffrey Zigman, University of Texas Southwestern Medical Center, Dallas, TX, USA) and WT (C57BL/6J; WT) controls (Harlan, Bicester, UK) (14 weeks old, n = 3 per group) were used.19 The methodological and metabolic aspects of the study have been described previously.20 Briefly, mice were prepared with jugular vein cannulae attached to osmotic minipumps (Alzet model 2001; DURECT Corp., Cupertino, CA, USA) under isofluorane anaesthesia. The minipumps delivered either vehicle or acyl-ghrelin (48 µg day−1; Phoenix Pharmaceuticals, Mannheim, Germany) for 7 days. This treatment protocol was shown to increase abdominal adiposity via GHSR, although it had no effect on body weight.20 Mice were killed by cervical dislocation and whole trunk blood was collected into heparinised tubes for plasma separation by centrifugation at 4000 g for 10 minutes at 4°C. Whole brain was removed and immediately snap frozen on dry ice and stored at −80°C prior to analysis.

For analysis of Ki67, snap-frozen brains were sectioned at 10 µm thickness using a cryostat (Leica Microsystems, Wetzlar, Germany) and mounted directly onto Superfrost+ slides (VWR Scientific, Radnor, PA, USA). A one-in-fifteen series of 10 µm sections (150 µm apart) from each animal, a minimum of eight sections per mouse, was immunostained using rabbit anti-Ki67 (dilution 1:500, ab16667; Abcam) along with a biotinylated goat anti-rabbit for Ni-DAB based detection, as previously described.11 Cells were imaged by light microscopy (model 50i; Nikon, Tokyo, Japan) prior to quantification using ImageJ (NIH, Bethesda, MD, USA).

A separate cohort of 19-week old male GHSR-null mice, derived from crosses between animals that were heterozygous for the GHSR-null allele and that had been backcrossed >10 generations onto a C57BL/6J genetic background, and WT littermate mice were housed under normal laboratory conditions (12:12 hour light/dark cycle; lights on 7.00 AM) prior to the onset of the study. Mice were divided into six groups (n = 5-8 per group) that included ad lib. fed WT, calorie restricted (CR) WT, calorie restricted/re-fed (CR/RF) WT, ad lib. fed ghrelin−/−, CR ghrelin−/− and CR/RF ghrelin−/−. For the first 28 days of the study, mice were fed on an ad lib. diet with daily injections (from days 1-4) of the thymidine analogue, bromodeoxyuridine (BrdU) (50 mg kg−1, i.p.), to label actively dividing cells. On day 28, food was withdrawn at 5.30 PM from the CR and CR/RF mice. The next day, CR/RF mice were allowed to feed ad lib. for 1 hour prior to all animals undergoing cervical dislocation when under terminal anaesthesia (approximately 18 hours CR). Ghsr−/− mice have growth rates and appetite similar to WT littermates, with no impairment in hyperphagia after fasting.21,22 Similarly, adult ablation of ghrelin in mice does not impair growth, nor appetite.23 Whole brains were removed, immersed in ice cold 4% PFA for 24 hours and cryoprotected in 30% sucrose. Coronal sections (30 µm) cut in a 1:12 series along the entire rostral-caudal axis of the OB (bregma +5.345 mm to 2.445 mm) using a freezing stage microtome (MicroM; Thermo Fisher Scientific) and collected for immunohistochemistry (IHC). For DAB-immunohistochemical analysis of GHSR labelling, a minimum of six sections per mouse was washed in 0.1 mol L−1 PBS (2 × 10 minutes) and 0.1 mol L−1 PBS-T (1 × 10 minutes). Subsequently, endogenous peroxidases were quenched by washing in a PBS plus 1.5% H2O2 solution for 20 minutes. Sections were washed again (as above) and incubated in 5% normal donkey serum in PBS-T for 1 hour. Sections were incubated overnight at 4°C with rabbit anti-GHSR1a (H-001-62; Phoenix Pharmaceuticals), diluted 1:2000 in PBS-T and 2% NGS solution. Another wash step followed prior to incubation with biotinylated goat anti-rabbit (dilution 1:400; Vector Laboratories, Inc.) in PBS-T for 70 minutes. The sections were washed and incubated in ABC (Vector Laboratories, Inc.) solution for 90 minutes in the dark prior to another two washes in PBS, and incubation with 0.1 mol L−1 sodium acetate pH6 for 10 minutes. Immunoreactivity was developed in Ni-DAB solution followed by two washes in PBS. Sections were mounted onto Superfrost+ slides (VWR Scientific) and cover-slipped with vectashield containing 4’,6-diamidino-2-phenylindole (Vector Laboratories, Inc., Burlingame, CA, USA) prior to the onset of the study. Mice were di-
2.2.5 | Quantification of BrdU+/c-Fos*

All IHC was performed on free-floating sections at room temperature, unless otherwise stated. A one in six series of 30 μm sections (180 μm apart) were washed three times in PBS for 5 minutes, permeabilised in methanol at −20°C for 2 minutes, and washed as described before. DNA was denatured with 2 mol L−1 HCl for 30 minutes at 37°C prior to washing sections in 0.1 mol L−1 borate buffer (pH 8.5) for 10 minutes. Sections were washed, blocked with 5% NGS plus 5% bovine serum albumin in PBS-T for 60 min and incubated in a cocktail of primary antibodies that included rat anti-BrdU (dilution 1:400; MCA2060; ABD Serotec, Kidlington, UK) and rabbit anti-c-Fos (dilution 1:1000; SC-52; Santa Cruz Biotechnology, Santa Cruz, CA, USA) in PBS-T overnight at 4°C. The primary antibody was omitted from the negative control.

Following primary antibody treatment, sections were washed, incubated with biotinylated goat anti-rat (dilution 1:400; BA-9400; Vector Laboratories, Inc.) in PBS-T for 60 minutes in the dark and then washed as described before. Similarly, secondary antibodies were also applied as a cocktail that included goat anti-rabbit (dilution 1:400; BA-1000; Vector Laboratories, Inc.) and streptavidin AF-594 (dilution 1:500; S11227; Life Technologies) in PBS-T for 30 minutes. Following another wash, including one containing Hoechst nuclear stain, sections were mounted onto Superfrost* slides (VWR Scientific) and cover-slipped with prolong gold anti-fade solution (Life Technologies).

2.2.6 | Quantification of immunolabelled cells

A one in six series of 30 μm sections (180 μm apart) from each animal was analysed for immunoreactivity using an epifluorescence microscope system (Imager M1 with Axiocam MRm; Carl Zeiss). Immunolabelled cells were manually counted bilaterally using a 40× objective through the Z-axis of the entire rostral-caudal extent of the dorsal granule cell layer (GCL), glomerular layer (GL), subependymal zone (SEZ) and the lateral olfactory tract body (LOT). Resulting numbers were divided by the total area measurement to give a count per pixel, which was converted into mm² and averaged for each brain. All analyses were performed blind to both genotype and treatment.

2.3 | Rats

Adult male Lister hooded rats (n = 10 per 11 per group, weighing 250-300 g; Harlan) were housed in groups of four and maintained at room temperature under a 12:12 h light/dark cycle; lights on at 7.00 AM. These experimental procedures have been described previously. Briefly, from days 0-14, rats received daily i.p. injections of acyl-ghrelin (031-31; Phoenix Pharmaceuticals) or saline (10 μg kg⁻¹ body weight) with BrdU injections (50 mg kg⁻¹) on days 5-8. On day 29, rats were terminally anaesthetised, transcardially perfused with 4% PFA and brains were removed for immersion fixation and cryoprotection (as described before). Analysis of adult hippocampal neurogenesis (AHN) in these rats demonstrated that acyl-ghrelin significantly increased the number of new adult born neurones.

2.3.1 | Double immunofluorescence for BrdU/neuronal nuclei (NeuN)

A one in six series of coronal OB brain sections (bregma +5.345 mm to +2.445) was transferred into a 24-well culture plate and washed in PBS, permeabilised in methanol at −20°C for 2 minutes and washed in PBS as before. DNA was denatured using 2 mol L⁻¹ HCl (Thermo Fisher Scientific) and incubated at 37°C for 30 minutes. Excess HCl was removed and the sections washed in 0.1 mol L⁻¹ borate buffer (pH 8.5) for 10 minutes to neutralise the remaining HCl. Tissue sections were then washed, blocked with 5% NGS diluted in PBS-T for 1 hour and incubated with rat anti-BrdU (rat monoclonal; ABD Serotec), diluted 1:3000 in PBS-T for 24 hours at 4°C. The primary antibody was omitted from the negative control. Sections were washed and incubated with biotinylated goat anti-rat (goat polyclonal; BA-9400; Vector Laboratories, Inc.), diluted 1:400 in PBS-T for 1 h in the dark. Tissue sections were subsequently washed, incubated in streptavidin AF594 (S11227; Life Technologies), diluted 1:500 in PBS-T for 30 minutes and washed as described before. Sections were then incubated in mouse anti-NeuN (mouse monoclonal; MAB377; EMD Millipore, Burlington, MA, USA), diluted 1:1000 in PBS-T for 1 hour. The negative control contained PBS-T. Tissue sections were then washed and incubated in goat anti-mouse AF 488 diluted 1:500 in PBS-T for 30 minutes, prior to being washed with Hoescht, diluted 1:10 000 in PBS, for 5 minutes. Finally, sections were washed, mounted onto Superfrost* slides and cover-slipped with prolong gold anti-fade reagent, prior to storage at 4°C.

2.3.2 | Quantification of BrdU*

IMAGEL, version 1.47 (NIH) was used to quantify the number of new adult-born cells in the dorsal and ventral GCL of the OB. Images taken by the fluorescence microscope were inverted and unsharpmasked, using a radius of 10.0 pixels and a mask weight of 0.60. The polygon tool was then used to draw around the granular cell layer and the total area measured. Each image’s threshold was individually optimised, typically ranging from 0.100 to 0.180. The particle size was set to 20-300 pixel² and circularity at 0.0-1.0. Resulting numbers were divided by the total area measurement to give a count per pixel, which was then averaged for each brain.

2.3.3 | Quantification of BrdU*/NeuN*

To quantify the number of new adult-born neurones in the dorsal and ventral granular cell layer of the OB, BrdU*/NeuN* immunoreactive cells were manually counted through the Z-axis of the entire rostral-caudal extent of the OB. Resulting numbers were divided by the area of the Z-stack to give a count per pixel, which was then averaged for each brain.
2.4 | Microscopy

Tissue sections were analysed using a fluorescence microscope (Imager M1 with Axiocam MRm; Carl Zeiss) with axiovision, version 4.6 software and a laser scanning confocal microscope (LSM 710; Carl Zeiss) with Zen software (ZEN 2010 edition; Carl Zeiss). Images were collected using 4×, 10× and 40× objectives. BrdU+/NeuN+ immunoreactive newborn adult neurones, in the dorsal and ventral GCL, were imaged using a 40× oil immersion objective. A Z-stack consisting of 21-25 tissue slices at 0.7 μm intervals (14.0-16.8 μm range) were taken throughout the rostral-caudal extent of the OB.

All experiments and analyses were performed blind to genotype and treatment.

2.5 | Statistical analysis

Statistical analyses were performed using Prism, version 6.0 for Mac (GraphPad Software Inc., San Diego, CA, USA). Statistical significance was assessed by an unpaired two-tailed Student’s t test or one-way ANOVA with Bonferroni’s post-hoc test. Where there was more than one variable, a two-way ANOVA with Tukey’s multiple comparisons test was used or by a Kruskal-Wallis test followed by a Dunn’s multiple comparisons test when a normal distribution of data could not be assumed. Data are presented as the mean ±SEM. P < 0.05 was considered statistically significant.

3 | RESULTS

3.1 | GHSR is expressed in the adult OB but not in the SVZ

The expression of GHSR was assessed to determine whether ghrelin could directly influence the proliferation of NSPCs in the SVZ. Accordingly, we used GHSR-eGFP reporter mice to show that eGFP immunoreactivity was present within the anterior olfactory nucleus and orbital and motor orbital cortex in the caudal OB (Figure 1i). Immunoreactivity was observed within the anterior cingulate cortex, motor cortex and lateral septal nucleus (Figure 1ii). Sagittal sections revealed strong immunoreactivity in the anterior amygdala area, granule cell layer of the hippocampal dentate gyrus and the medial amygdala nucleus (Figure 1iii). Notably, staining was absent within the lateral lining of the SVZ in tissue sectioned in both a coronal and sagittal orientation (Figure 1iD,iiC). To determine whether the GHSR-eGFP immunoreactivity was similar to that observed with GHSR1a antisera, we performed IHC using a rabbit anti-GHSR1a antibody on adult WT and GHSR-null mouse brain tissue. These analyses revealed a similar pattern of immunoreactivity on both WT and GHSR-null tissues.
including the SVZ (see Supporting information, Figure S1), suggesting a lack of binding specificity for the GHSR1a antigen. These data suggest that ghrelin may be involved in olfactory function but not via direct modulation of NSC's lining the lateral wall of the SVZ.

3.2 | Acyl-ghrelin does not increase cell proliferation in the adult SVZ

The proliferative effect of ghrelin and GHSR-agonists have been widely reported within the central nervous system and peripheral tissues. A recent study reported that ghrelin promoted proliferation of cells within the SVZ. Here, we took advantage of genetically modified mice to analyse the effect of acyl-ghrelin treatment on SVZ cell proliferation in adult WT and GHSR-null mice, where GHSR is transcriptionally silenced. Using the mitotic marker, Ki67, we report that acyl-ghrelin treatment had no effect on the number of proliferating cells within the SVZ niche in WT mice ($P \geq 0.99$) (Figure 2). Similarly, transcriptional silencing of GHSR did not affect the rate of SVZ cell division in vehicle (WT vehicle vs GHSR-null vehicle, $P = 0.6388$) or acyl-ghrelin treated mice (GHSR-null vehicle vs GHSR-null acyl-ghrelin, $P = 0.0944$). The low number of replicates means that the statistical analysis is of low power; however, the data suggest that acyl-ghrelin does not regulate cell proliferation in the adult mouse SVZ and also that genetic silencing of GHSR does not decrease cell division in this niche.

3.3 | Acyl-ghrelin does not increase the number of new adult-born OB neurones

We recently showed that treatment with acyl-ghrelin increased AHN in adult rats. To determine whether acyl-ghrelin treatment modulates AOBN in a similar way to AHN, we quantified the number of new adult-born neurones in the OB of adult rats from the same study. The use of OB tissue from the same rats where acyl-ghrelin increased AHN provides an valuable experimental control. We show there was no significant difference in the number of new adult-born cells (BrdU+/NeuN+) in the GCL of the OB following acyl-ghrelin treatment compared to saline treatment ($P = 0.8482$) (Figure 3G). Similarly, no differences were observed in the number of new adult-born neurones (BrdU+/NeuN+) ($P = 0.7388$) (Figure 3H) or in the rate of neurone differentiation ($P = 0.6870$) (Figure 3I).

3.4 | Calorie restriction induces activation of new adult-born OB cells in a ghrelin-dependent manner

To determine whether a CR-mediated increase in endogenous acyl-ghrelin was able to increase the expression of the proto-oncogene, c-Fos, in new adult-born OB cells, we analysed the number of active c-Fos+ cells within the GCL, GL, SEZ and LOT in WT and ghrelin−/− mice. A two-way ANOVA revealed a statistically significant main effect of treatment on BrdU+/c-Fos+ ($P = 0.0031$ and BrdU+/c-Fos+ ($P = 0.0487$) cells within the GCL. Comparatively, genotype and the interaction (treatment and genotype) showed a significant effect on c-Fos+ ($P = 0.0001$ and 0.017, respectively) and BrdU+/c-Fos+ ($P = 0.0002$ and 0.00021, respectively) cells within the GCL. Outside of the GCL, a significant main effect of treatment was reported in BrdU+/c-Fos+ cells of the SEZ ($P = 0.0118$) and BrdU+ cells of the LOT ($P = 0.0075$). No increase was observed within the GL (see Supporting information, Table S1).

Tukey's post-hoc test revealed a reduction in the number of new adult-born cells (BrdU+) in CR/re-fed (CR/RF) ghrelin−/− mice.
FIGURE 3  Exogenous acyl-ghrelin does not increase the number of new adult born neurones in the granule cell layer of the rat olfactory bulb. A, Experimental paradigm. IHC, immunohistochemistry. B, Collage image of the rat olfactory bulb. Representative images of bromodeoxyuridine (BrdU) (red) and neuronal nuclei (NeuN) (green) in (C) dorsal granule cell layer (GCL) and (D) ventral GCL of the olfactory bulb (OB). Scale bar = 200 μm. Representative images of new adult-born neurones co-expressing NeuN+ and BrdU+ (yellow) in (E) dorsal GCL and (F) ventral GCL. Scale bar = 50 μm. Quantification of new adult-born OB cells (G) ($P = 0.8482$), new adult-born neurones (H) ($P = 0.7388$) and percentage neuronal differentiation (I) ($P = 0.6870$) after acyl-ghrelin or saline treatment. Data are the mean ± SEM. Statistical analysis was performed by a two-tailed unpaired Student’s t test. $P < 0.05$ was considered statistically significant; NS, not significant; n = 11 rats per group.
New adult-born olfactory bulb (OB) cells are activated by calorie restriction in a ghrelin-dependent manner. A, Schematic of experimental paradigm. IHC, immunohistochemistry. B, New adult-born active neuron (yellow; scale bar = 25 μm) co-expressing bromodeoxyuridine (BrdU) (green) and c-Fos (red) in the granule cell layer (GCL) of the OB. Scale bar = 50 μm. Quantification of (C) new adult-born cells (BrdU+), (D) active cells (c-Fos+), (E) active new adult-born cells (BrdU+/c-Fos+) and (F) active developmentally born cells (BrdU−/c-Fos+) in the GCL of the OB. G, Representative images of new adult-born cells (BrdU+; green), active cells (c-Fos+; red) and active new adult-born cells (BrdU+/c-Fos+; yellow in merged image). Arrows correspond to new BrdU+/c-Fos+ cells, whereas arrowheads represent active new adult-born BrdU+/c-Fos+ cells. Scale bar = 50 μm. Statistical analysis was performed by two-way ANOVA with Tukey’s post-hoc test. *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001. All data shown are the mean ± SEM; n = 5-8 rats per group. AL, ad lib.; CR, calorie restriction; CR/RF, calorie restriction/re-fed; WT, wild-type; GKO, ghrelin−/−.
compared to CR ghrelin−/− mice (P = 0.0086) (Figure 4C) within the GCL. Furthermore, there was an increased number of activated cells (c-Fos+) in CR WT mice compared to ad lib. WT (P = 0.0258) and CR/RF WT (P = 0.0043) mice. Notably, CR also increased activated cells in WT relative to CR ghrelin−/− mice (P < 0.0001) within the GCL (Figure 4D). Further analysis revealed that the number of active new adult-born cells (BrdU+/c-Fos+) was increased in CR WT mice compared to CR/RF WT (P = 0.0169) and CR ghrelin−/− mice (P < 0.0001) within the GCL (Figure 4E). However, there were very few active developmentally born cells (BrdU+/c-Fos+) and these cells were not significantly affected by treatment or genotype (Figure 4F). Outside of the GCL, the number of active new adult-born cells (BrdU+/c-Fos+) was reduced in CR ghrelin−/− mice compared to ad lib. fed ghrelin−/− mice (P = 0.0169) within the SEZ. No significant differences were reported in the other regions tested (see Supporting information, Table S1). There was no significant difference in body weight change in ghrelin−/− mice relative to WT mice in either of the groups (two-way ANOVA; main effect of genotype, P = 0.9335; main effect of feeding pattern, P = 0.0049; main effect of interaction [feeding pattern vs genotype], P = 0.3469). Collectively, these data suggest that CR increases the activation of new adult-born cells in a ghrelin-dependent manner.

4 | DISCUSSION

The generation of new OB neurones in the adult brain is important for olfactory discrimination, which is a process that is impaired in ageing and several neurodegenerative disorders. In the present study, we tested the hypothesis that ghrelin is an important regulator of AOBN. First, we characterised the expression of GHSR in the adult mouse brain. Numerous studies have attempted to characterise the expression pattern of GHSR in several species, including mouse, rat and lemur,27–30 although a lack of reliable anti-GHSR antibodies has limited progress. More recently, a report using the GHSR-eGFP mouse and in situ hybridisation histochemistry demonstrated GHSR expression within the OB, hippocampus and hypothalamic nuclei.15 Furthermore, Cre-activity in Ghstr-IRESCre/ROSA26-ZsGreen reporter mice was also reported in the main and accessory OB.31

In the present study, using adult GHSR-eGFP mice, we report GHSR immunoreactivity in the MCL, anterior olfactory nucleus and orbital and motor orbital cortices of the OB, as well as within the anterior cingulate cortex, motor cortex, lateral septal nucleus, entopeduncular nucleus, hippocampus and medial amygdaloid nucleus. However, GHSR was not expressed within the neurogenic niche of the SVZ in GHSR-eGFP reporter mice. Indeed, this finding is consistent with previous studies that do not report GHSR immunoreactivity within the SVZ niche. Taken together, these findings suggest that ghrelin does not mediate direct effects on NSPC proliferation.

Because eGFP immunoreactivity in this transgenic model may correspond to two structurally different receptors, GHSR1a, which encodes the functional receptor, and the truncated GHSR1b, generated from alternative splicing of GHSR, we aimed to identify GHSR1a expressing cells using antisera raised against GHSR1a. The specificity of polyclonal antibodies used to characterise GHSR within the adult brain remains unclear. Li et al14 reported GHSR expression within the adult mouse neurogenic niche of the SVZ using immunofluorescence with the rabbit anti-GHSR1a antibody (H-001-62; Phoenix Pharmaceuticals), diluted 1:500. In the present study, using the same antibody, IHC in brain tissue from adult GHSR-null and WT mice revealed detectable immunoreactivity in tissue from both genotypes. Our data suggest that the rabbit anti-GHSR antibody resulted in non-specific staining within the SVZ and cingulate cortex (see Supporting information, Figure S1), preventing its use for determining GHSR1a expression in this context. Taken together, these studies suggest that the ghrelin receptor is not expressed in the SVZ, and thus does not directly modulate NSPC proliferation.

To determine whether ghrelin induces cell proliferation within the SVZ we treated GHSR-null and WT mice for 7 days with acyl-ghrelin. Subsequent analysis revealed no effect of genotype or treatment on the number of dividing Ki67+ cells in the SVZ. By contrast, a previous study reported that ghrelin−/− mice had a reduced number of proliferating NSPCs, migrating neuroblasts and OB interneurons, which could be restored to WT levels by i.p. administration of acyl-ghrelin.14 Several differences between the two experimental procedures might account for the contrasting results. For example, Li et al14 used 8-9-week-old WT and ghrelin−/− mice that received acyl-ghrelin (80 μg kg−1) via i.p. injection, once daily for eight consecutive days. In the present study, 14-week old WT and GHSR-null mice were given acyl-ghrelin (48 μg day−1) via an i.v. minipump. Therefore, inconsistencies between studies may be attributable to genetic background, the physiological dose or the route of administration of acyl-ghrelin.

Next, using a BrdU pulse-chase approach, we determined the effect of exogenous acyl-ghrelin treatment on the maturation and survival of new adult-born neurones in the rat OB. Consistent with our previous cell proliferation analysis in mice, acyl-ghrelin did not increase in the number of new adult-born BrdU+ cells or BrdU+/NeuN+ neurones in the GCL of the OB. Furthermore, no differences were observed in the rate of neuronal differentiation. Notably, we have previously reported that adult hippocampal neurogenesis was significantly increased by acyl-ghrelin in these rats.11 The high level of GHSR expression within hippocampal neurogenic niche12 and its absence in the SVZ niche is likely responsible for this effect. These data provide compelling evidence that acyl-ghrelin does not promote AOBN.

Numerous studies have suggested that ghrelin plays an important role in olfactory-related behaviours including odour discrimination and sensitivity.8,32,33 Loch et al8,32,33 reported an increased responsiveness of the mouse olfactory epithelium following nasal application of ghrelin. This resulted in a higher reactivity of olfactory sensory neurones within the olfactory epithelium, which in turn, increased the activity of receptor-specific glomeruli. GHSR expression on the surface of olfactory sensory neurones suggests that ghrelin and GHSR may play an important role in enhancing neuronal responsiveness and olfaction. However, the underlying mechanism...
by which acyl-ghrelin enhances olfaction remains elusive and it is unclear whether new adult-born OB neurones are involved in this physiology. Because acyl-ghrelin is known to regulate both olfaction and appetite, we aimed to determine whether new adult-born OB neurones are activated by CR in a ghrelin-dependent manner. Our data demonstrate that, overnight, CR activated new adult-born cells in the OB. Re-feeding for 1 hour returned the number of c-Fos positive cells to baseline, suggesting that the new adult-born cells are sensitive to feeding status. Notably, this CR effect was absent in ghrelin−/− mice, demonstrating that the activation of new adult-born cells was dependent upon intact ghrelin signalling. Furthermore, there was no CR-mediated activation of developmentally born cells (BrdU+/c-Fos+) in the GCL of the OB, indicating that adult-born neurones are uniquely responsive to acute changes in food intake. Therefore, we confirm that CR activates new adult-born OB cells in a ghrelin-dependent manner. This finding provides further support for ghrelin acting as a mediator of CR-associated physiology, including, neuroprotection, anti-anxiety, hippocampal neurogenesis and cognitive enhancement and glycaemic regulation.

Although the relationship between hunger stimulation and olfaction has been long recognised, a molecular mechanism relating the two processes has not been determined. Soria-Gómez et al observed that cortical feedback projections to the OB crucially regulate food intake, possibly through cannabinoid type-1 receptor (CB1R) signalling. The endocannabinoid system, in particular CB-1Rs, promotes food intake in fasted mice by increasing odour detection. Notably, the orexigenic effect of ghrelin is lost in CB-1R knockout mice. Although the relationship between ghrelin and the endocannabinoid system in the OB is unknown, both GHSR and CB-1R are GPCRs known to form homo- and heterodimers (or higher-order oligomers) as part of their normal trafficking and function. Therefore, heterodimerisation of CB-1R and GHSR may be important in linking ghrelin to adult-born OB neurones and olfaction. Several questions remain unanswered, including whether ghrelin alters the electrophysiological properties and/or directly activates GCs in the OB to enhance odour discrimination. Because new adult-born OB cells enhance the odour-reward association, further work is needed to determine whether the ghrelin-induced intake of rewarding foods requires signalling via new neurones in the OB. Similarly, it is not known whether ghrelin can increase appetite and improve olfaction in the absence of new adult-born OB cells.

In summary, these data demonstrate that, although ghrelin does not increase SVZ-OB neurogenesis, it does mediate the CR-induced activation of new adult-born OB cells. We speculate that ghrelin modulates new OB neurone activity to integrate olfactory responses with nutritional status.

**DATA AVAILABILITY**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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**REFERENCES**


SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.