Gpr158 mediates osteocalcin’s regulation of cognition

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INTRODUCTION

Osteocalcin (OCN) is a multipurpose bone-derived hormone (Karsenty and Olson, 2016). In particular, OCN crosses the blood–brain barrier, binds to neurons in specific brain regions, prevents neuronal apoptosis in the hippocampus, favors the synthesis of all monoamine neurotransmitters, and inhibits that of gamma-aminobutyric acid. These cellular and molecular functions may explain why OCN is necessary for hippocampal-dependent memory and to reduce anxiety-like behaviors (Oury et al., 2013). The severity of the cognitive deficit noted in any mouse model lacking a peripheral delivery of exogenous OCN is sufficient to improve memory and decrease anxiety-like behaviors in 16-mo-old mice. A second question is to identify a receptor transducing OCN signal in neurons. Genetic, electrophysiological, molecular, and behavioral assays identify Gpr158, an orphan G protein–coupled receptor expressed in neurons of the CA3 region of the hippocampus, as transducing OCN’s regulation of hippocampal-dependent memory in part through inositol 1,4,5-trisphosphate and brain-derived neurotrophic factor. These results indicate that exogenous OCN can improve hippocampal-dependent memory in mice and identify molecular tools to harness this pathway for therapeutic purposes.

Abbreviations used: ACSF, artificial cerebrospinal fluid; AP, action potential; BDNF, brain-derived neurotrophic factor; BW, body weight; EPMT, elevated plus maze test; eEPSP, field excitatory postsynaptic potential; GPCR, G protein–coupled receptor; IP3, inositol 1,4,5-trisphosphate; LTP, long-term potentiation; MF, mossy fiber; NOR, novel object recognition; OCN, osteocalcin; OFT, open field test; TIMP, tissue inhibitor of metalloproteinases.
second question, we provide expression, genetic, electrophysiological, molecular, and functional evidence that the orphan class C G protein–coupled receptor (GPCR) Gpr158 mediates OCN’s regulation of cognitive function. Together, these data pave the way to test whether OCN signaling in the brain could be harnessed to combat cognitive decline.

RESULTS AND DISCUSSION

To determine whether OCN is a necessary component of plasma from young mice, contributing to its beneficial effect on cognition in older mice, we relied on the novel object recognition (NOR) test, which evaluates hippocampal-dependent memory because this task is altered in Ocn−/− mice (Oury et al., 2013). 16-mo-old WT mice had a significant defect in hippocampal-dependent memory compared with 3-mo-old (young) WT mice. This cognitive defect was significantly improved when plasma from young mice was injected into 16-mo-old mice but not when plasma from other 16-mo-old WT mice was injected in 16-mo-old WT littermates (Fig. 1 A). Because OCN limits anxiety-like behavior (Oury et al., 2013), we also subjected young and 16-mo-old mice to the elevated plus maze test (EPMT), which assesses anxiety-like behavior as a shorter time spent in the open arms and fewer number of entries in the open arms (Crawley, 1985). We found that plasma from young mice, when injected into 16-mo-old mice, significantly reduced anxiety-like behaviors (Fig. 1 B).

In contrast, when plasma from young Ocn−/− mice was injected into 16-mo-old WT mice, there was no improvement of hippocampal-dependent memory or anxiety-like behaviors in the recipient mice (Fig. 1, A and B; and Fig. S1 A). The inability of the plasma from young Ocn−/− mice to improve cognition in older mice was not explained by an increase in β2 microglobulin accumulation, a progeronic molecule in this plasma (Smith et al., 2015; Fig. S1 B). To further demonstrate that the presence of OCN is necessary for the beneficial effect of plasma from young mice on cognition when injected into 16-mo-old WT mice, we performed two experiments. First, we used plasma obtained from young Ocn−/− mice that had been supplemented with mouse recombinant uncarboxylated OCN (spiked plasma). We first verified that injections of spiked plasma increased circulating OCN levels in 16-mo-old WT mice (Fig. S1 C). Remarkably, injections of spiked plasma in 16-mo-old WT mice resulted in the same improvement in hippocampal-dependent memory that is achieved by injections of plasma from 3-mo-old WT mice (Fig. 1 A and B). This beneficial effect of the spiked plasma on cognitive function was not explained by a change in the accumulation of tissue inhibitor of metalloproteinases 2 (TIMP2) in the hippocampus, a molecule implicated in the beneficial effect of young plasma on cognition (Castellano et al., 2017; Fig. 1 C). Consistent with the beneficial effect of young plasma injections into older mice, injections of spiked plasma significantly reduced anxiety-like behaviors in 16-mo-old WT mice (Fig. 1 A and B). Second, we used an anti-OCN antibody to deplete OCN from the plasma of young WT mice. We verified before injection of this immune-depleted plasma that OCN content was decreased by 90% (Fig. S1 D). In a mirror image of what was observed when using spiked plasma, immune-depleted plasma from WT mice failed to increase hippocampal memory and to reduce anxiety to the same extent as control plasma (Fig. 1, D and E). Together, these experiments indicate that OCN is a necessary contributor to the beneficial effect of plasma from young WT mice on hippocampal-dependent memory and anxiety-like behaviors in older WT mice.

The steep decrease in circulating OCN levels before 1 yr of age (Fig. S1 E), and the crucial contribution of OCN to the beneficial effect of plasma from young mice on cognitive function of older mice, prompted us to test whether exogenous OCN would be sufficient to improve cognition in older mice. This was addressed by delivering OCN peripherally and continuously through minipumps for 2 mo in 14-mo-old WT mice because this hormone crosses the blood–brain barrier (Oury et al., 2013). This mode of administration substantially increased circulating OCN levels (Fig. S1 F). When tested for NOR, 16-mo-old WT mice that had received OCN continuously for 2 mo displayed a hippocampal-dependent memory that was similar to that of 3-mo-old WT mice (Fig. 1 F). The same improvement was observed when memory was analyzed through the Morris water maze test, which assesses spatial learning and memory in rodents (Nakazawa et al., 2002; Fig. 1 G). Similar results were observed in 12-mo-old WT mice (Fig. S1, G and H). In agreement with the notion that OCN regulates anxiety-like behaviors, chronic delivery of OCN improved anxiety-like behaviors as assayed by the EPMT in 12- and 16-mo-old WT mice (Oury et al., 2013; Fig. 1 H and Fig. S1 I).

The fact that OCN is necessary for the maintenance, and sufficient for the restoration, of cognitive function in older mice underscores the importance of identifying a receptor for this hormone in the brain because Gprc6a, OCN’s receptor in peripheral organs (Karsenty and Olson, 2016), does not mediate OCN’s functions in the brain (Oury et al., 2013). The bell-shaped curve of OCN signaling in brainstem explants (Oury et al., 2013) suggested that, as in the case of Gprc6a, the receptor for OCN in the brain might be a GPCR. We used three criteria to identify an orphan GPCR capable of transducing OCN signal in the brain.

The first criterion was that this orphan GPCR should belong to the same subfamily of GPCRs as Gprc6a, the class C subfamily. Second, this GPCR should be expressed in the CA3 region of the hippocampus where OCN binds (Oury et al., 2013), and third, this GPCR should not be expressed in cell types where OCN’s signal is transduced through Gprc6a (Fig. 2 A). Using this approach, we identified one orphan class C GPCR, Gpr158, as being robustly expressed in neurons of the CA3 region of the hippocampus but not in any Gprc6a-expressing cells in peripheral tissues (Fig. 2, B and D; and Fig. S2, A and C). Gpr158 is present in Map2-expressing
Figure 1. OCN is sufficient to improve cognitive function and anxiety-like behaviors. (A and B) NOR (A) and EPMT (B) performed in 3-mo-old (young) WT mice treated with plasma from young WT mice (n = 8–10) or 16-mo-old (aged) WT mice treated with plasma from WT mice, either aged (n = 11–12) or young (n = 11–16), or from young Ocn−/− mice (n = 11–16), or plasma from young Ocn−/− mice supplemented with 90 ng/g BW of OCN (spiked; n = 12–13). For NOR, discrimination index was measured; for EPMT, number of entries and time spent in the open arms were scored (one-way ANOVA followed by a Bonferroni post hoc test compared with WT young). (C) TIMP2 accumulation (representative Western blot, left) and quantification of band intensities (right) in hippocampi of older WT mice receiving plasma from Ocn−/− mice supplemented with 90 ng/g BW of OCN (Student’s t test, n = 4 mice per group). α-Tubulin is used as a loading control. (D and E) NOR (D) and EPMT (E) performed in 18-mo-old mice treated with IgG (n = 9–10) or anti-Ocn immunodepleted plasma (n = 8–9). For NOR, discrimination index was measured. For EPMT, number of entries and time spent in the open arms were scored.
pyramidal neurons from the CA3 region of the hippocampus but not in Glap-expressing astrocytes in vivo (Fig. 2 F and Fig. S2 D). Similar results were obtained in cultured hippocampal cells using a specific antibody against Gpr158 (Fig. 2 G and Fig. S2 E). Adding credence to the notion that Gpr158 may be an OCN receptor, its accumulation was higher in Ocn−/− than in WT hippocampi, circulating OCN levels were twofold higher in Gpr158−/− compared with WT littermates, and local delivery of OCN in the anterior hippocampus induced c-Fos accumulation in neurons of the CA3 region in WT but not in Gpr158−/− mice (Fig. 2, H–J; and Fig. S2 F). A systematic analysis of its expression by in situ hybridization showed that Gpr158 is also expressed in the somatosensory, motor, and auditory areas of the cortex, the piriform cortex, the retro-splenial area, and the ventral tegmental area (Fig. 2 E). A signaling pathway recruited by GPR158 was identified by a pull-down assay performed on solubilized membranes obtained from Ocn−/− hippocampi. In the conditions of this assay, biotin-labeled OCN bound a complex containing Gpr158 and Gαq (Fig. 2 K). The amount of Gαq pulled down was greatly reduced when membranes from Gpr158−/− hippocampi were used to perform this assay (Fig. 2 K). Consistent with the presence of Gαq in this complex, OCN increased the production of inositol 1,4,5-trisphosphate (IP3) significantly more in WT than Gpr158−/− hippocampal neurons (Fig. 2 L) and failed to increase in cAMP production (Fig. S2 H).

We used behavioral and electrophysiological assays to assess whether Gpr158 mediates OCN’s regulation of cognitive function. Using NOR, we observed that mice lacking Gpr158 in the forebrain only had a deficit in hippocampal-dependent memory, as Gpr158−/− mice did. This deficit, however, was less severe in Gpr158−/− mice (Fig. 3 A and Fig. S3, A–D), most likely because of the low efficiency of Gpr158 deletion using CamkIIa-Cre mice (Fig. S3 E). A signaling pathway recruited by GPR158 was identified by a pull-down assay performed on solubilized membranes obtained from Ocn−/− hippocampi. In the conditions of this assay, biotin-labeled OCN bound a complex containing Gpr158 and Gαq (Fig. 2 K). The amount of Gαq pulled down was greatly reduced when membranes from Gpr158−/− hippocampi were used to perform this assay (Fig. 2 K). Consistent with the presence of Gαq in this complex, OCN increased the production of inositol 1,4,5-trisphosphate (IP3) significantly more in WT than Gpr158−/− hippocampal neurons (Fig. 2 L) and failed to increase in cAMP production (Fig. S2 H).

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To add support to the notion that Gpr158 mediates OCN’s regulation of cognitive functions, we performed electrophysiological studies in WT and Gpr158−/− cells. In whole-cell current-clamp recordings, we found that OCN enhanced action potential (AP) frequency in pyramidal cells of the CA3 region of WT but not Gpr158−/− hippocampi (Fig. 3 G and Fig. S3 G). Consistent with the observation that Gpr158 forms a complex with Gαq, 2-aminoethoxydiphenyl borate, an IP3 receptor antagonist (Maruyama et al., 1997), nearly abolished OCN’s ability to enhance AP frequency in pyramidal cells of the CA3 region (Fig. 3 H). Next, we examined long-term potentiation (LTP) expression in synapses from the mossy fiber (MF) to CA3 neurons using hippocampal slices from WT and Gpr158−/− mice. The field excitatory postsynaptic potentials (fEPSPs) recorded in the CA3 region by stimulating the MF cells could be blocked by the application of DCG-IV (Fig. S3 H). To verify LTP at the MF-CA3 pyramidal synapses (Kamiya et al., 1996), two trains of tetanic stimulation were applied to the MF-CA3 synapses after a stable baseline of fEPSPs were recorded. The tetanic stimulation induced a brief post-tetanic potentiation and LTP in WT (baseline, 106.5 ± 3.4%; 50 min after tetanus, 126.7 ± 10.1%; P < 0.05) but not in Gpr158−/− hippocampal slices (baseline, 100.9 ± 1.5%; 50 min after tetanus, 104.9 ± 6.8%; P > 0.05; Fig. 3 J). Of note, tetanic stimulation induced a brief depression but no potentiation right after the completion of the tetanus in Gpr158−/− slices.

Because OCN regulates anxiety-like behaviors, we subjected Gpr158−/− and WT littermates to the EPMT and the dark-to-light transition test, which measures anxiety-like behaviors on the basis of the innate aversion of rodents to brightly illuminated areas (Crawley, 1985). As in the case of Ocn+/− and Ocn−/− mice, anxiety-like behavior was increased in 3-mo-old Gpr158+/− and Gpr158−/− mice.
Figure 2. Identification of a receptor for OCN in the brain. (A) Strategy used to identify an OCN brain receptor. (B) In situ hybridization of Gpr158 in E14.5 WT mouse embryos. Bar, 0.5 mm. (C) In situ hybridization of Gpr158, Gpr156, Gpr179, Gprc5a, Gprc5b, Gprc5c, and Gprc5d in the brain of 10-d-old WT mice. Bar, 200 µm. (D) Expression of Gpr158 and Gprc6a in various tissues of 3-mo-old WT mice (Student's t test, n = 4 mice per group). In the brain-derived tissues, expression is relative to Gpr158 expression, and in peripheral tissues, expression is relative to Gprc6a. (E) In situ hybridization of Gpr158 in the brain of 3-mo-old WT mice. For the ventral tegmental area (VTA), Th was used as a positive control. Bar, 200 µm. (F) Immunofluorescence of Lacz and...
In a third test, the open field test (OFT), which measures anxiety as a decrease in total ambulation and time spent in the center of the box, Gpr158+/− and Gpr158−/− mice displayed anxiety-like behaviors similar to Ocn−/− and Ocn−/− mice (Oury et al., 2013; Fig. 3, L and M). Additionally, the month-long peripheral delivery of OCN to Gpr158−/− did not reduce anxiety-like behavior in these mice (Fig. S3 I).

In trying to identify molecular events triggered by OCN through Gpr158, we asked whether OCN may act through brain-derived neurotrophic factor (BDNF) signaling, a molecular pathway critical for hippocampal-dependent memory, whose regulation by bone-derived signals has not yet been reported (Hall et al., 2000; Yamada and Nabeshima, 2003). For that purpose, we monitored the transport of BDNF-containing dense-core vesicles in embryonic rat neurons seeded in microchambers, and transduced with lentivirus, allowing the expression of BDNF-mCherry (Fig. 4 A). These neurons were left to differentiate for 9 to 11 d. Axons were imaged every 200 ms in microchannels to quantify kinetics of anterograde (from soma to synapse) and retrograde (from synapse to axon) transport of BDNF vesicles under control conditions or after 4 h of OCN treatment. OCN thus stimulates the global trafficking of BDNF vesicles and its specific transport to synapses, as shown by the increase in transport velocity of both anterograde and retrograde BDNF vesicles and the decrease in time spent in pause (Fig. 4 B). Moreover, the number of motile anterograde vesicles was larger (Fig. 4 B).

Several lines of evidence obtained using WT or Gpr158-null cells support the notion that BDNF is a mediator of OCN’s functions in the brain. First, Bdnf expression was decreased in hippocampi and midbrains of Gpr158−/− mice (Fig. 4 C and Fig. S3 J) and OCN increased Bdnf expression significantly more in WT than in Gpr158−/− cultured hippocampal neurons (Fig. 4 D). Second, delivery of OCN increased the accumulation of BDNF in mice that had received scrambled shRNA but not in those in which Gpr158 abundance had been efficiently decreased by shRNA targeting Gpr158 (Fig. 4, E and F). Third, peripheral delivery of OCN increased Bdnf expression in hippocampi of 16-mo-old WT mice (Fig. 4 G). Fourth, BDNF accumulation was increased in hippocampi of older WT mice receiving plasma from young WT mice but not in hippocampi of older mice receiving plasma from young Ocn−/− or from older WT mice (Fig. 4 H). Last, BDNF accumulation in the hippocampus was higher in 16-mo-old WT mice that had received plasma from Ocn−/− mice supplemented with OCN (Fig. 4 H).

The identification of extracellular cues favoring hippocampal-dependent memory, and the elucidation of their mechanism of action, is of utmost importance to understand how cognition is regulated and can be altered under pathological circumstances. At the same time, the identification of such molecules raises questions regarding their signaling in the brain. By showing that the bone-derived hormone OCN is sufficient to improve hippocampal-dependent memory in 16-mo-old mice, the results presented here underscore the importance of OCN in modulating these functions throughout life. That these effects of OCN signaling in the brain are independent of β2 microglobulin and TIMP2 indicates that multiple circulating molecules, some of which are hormones, affect cognitive function (Villeda et al., 2014; Castellano et al., 2017). As a critically important molecular extension of this work, we show through several types of assays that OCN mediates its influence on cognitive function through its interaction with an orphan class C GPCR, Gpr158. The identification of a receptor, expressed in neurons of the hippocampal CA3 region that transduces OCN signal, paves the way to potentially use this pathway for therapeutic purposes. As important, because Gpr158 is expressed in other regions of the brain besides the hippocampus, its identification as a receptor for OCN provides a much-needed tool to embark on a more extensive analysis of the functions OCN may have in the brain.

MATERIALS AND METHODS

Animals

For all experiments, we used females and littersmates as controls unless otherwise stated. Ocn−/− (osc−/−/osc−/−) mice were generated by our laboratory and previously described (Ducy et al., 1996). Gpr158−/− (Gpr158tm1(KOMP)Vlcg−/−) mice were purchased from the KOMP repository (VG10108). This mouse line was made by the insertion of a LacZ cassette with a stop codon in the first two exons of Gpr158 (Orlandi et al., 2015). Gpr158 conditional allele generation strategy is described in Fig. S3 A. Gpr158f/f were generated using 129/Sv embryonic stem cells. Chimeric mice harboring a mutant allele of Gpr158 were crossed with C57BL/6J WT, and Gpr158−/+ mice were then crossed with CamkHα-Cre (T29-1; Jax: B6.Cg-Tgf(Camk2α-cre)T29-1Stl/J) to obtain F2 generation. Analyses of Gpr158CamkHα-Cre were performed with Gpr158f/f as a control. Three−, 12−, and 16−mo-old 129S6/
Figure 3. Functional evidence that OCN signals through Gpr158. (A) NOR performed in 3-mo-old Gpr158/f/f (n = 9), Gpr158CamkIIa−/− (n = 5), Gpr158−/− (n = 10), and WT (n = 12) littermates. Discrimination index was measured (Student’s t test). (B) MWMT performed in 3-mo-old Gpr158−/− (n = 10) and WT (n = 9) littermates. The graph shows the time needed to localize a submerged platform in the swimming area (two-way repeated-measures ANOVA followed by Fisher’s least significantly different [LSD] post hoc test). (C) NOR performed in 4-mo-old WT (n = 7) or Gpr158−/− mice treated with saline (n = 7) or OCN (n = 8; 90 ng/hr) for 1 mo. Discrimination index was measured (one-way ANOVA compared with WT followed by Bonferroni’s post hoc test).
SvEvTac mice were purchased from Taconic Biosciences. Eighteen-mo-old C57BL/6J were obtained from the National Institute on Aging. Upon arrival at the animal facility, mice were housed at least 2 wk before experiments were performed. Gpr158+/−, Ocn+/−, and Gpr158+/−;Ocn+/− and Gpr158−/− were maintained on a 129-Sv/C57BL/6j mixed genetic background. Ocn−/− mice were maintained on a 129-Sv background. Mice were housed five animals per cage (polycarbonate cages, 35.5 × 18 × 12.5 cm), under a 12-h light/dark cycle with ad libitum access to food and water before experiment. All experiments involving animals were approved by the Institutional Animal Care and Use Committee of Columbia University Medical Center and by the European Communities Council Directive (2010/63/UE).

**Primary hippocampal culture**

Hippocampal neurons were isolated from mouse embryos (embryonic day 16.5). After dissection, hippocampi were digested with trypsin 0.05% and EDTA 0.02% for 15 min at 37°C. After three washes with DMEM (containing high glucose and sodium pyruvate) supplemented with 10% FBS, 100 U/ml penicillin-streptomycin and 1× GlutaMAX, cells were dissociated by pipetting up and down, and then plated. After the culture was established, culture medium (Neurobasal medium [21103-049]; Thermo Fisher Scientific) supplemented with B-27 supplement [17504044; Thermo Fisher Scientific] and 1× GlutaMAX [35050061; Thermo Fisher Scientific] was changed two times per week. Experiments were performed on cells after 15 d of culture.

**Plasma collection**

Pooled mouse plasma was collected from 3-mo-old WT or Ocn−/− mice or 16-mo-old WT mice by intracardial bleed at time of euthanasia. Plasma was prepared from blood collected with EDTA into Capiject (T-MQK; Terumo) tubes followed by centrifugation at 1,000 g for 10 min. All plasma aliquots were stored at −80°C until use. Before administration, plasma was dialyzed using 3.5-kD D-tube dialyzers (71508-3; EMD Millipore) in PBS to remove EDTA.

Mice were systemically treated with plasma (100 µl per injection) by injections into the tail vein eight times over 24 d before behavioral analysis. For Western blot analysis, mice were sacrificed to collect hippocampi 4 h after a single plasma injection.

**Immunodepletion of OCN from plasma**

To remove OCN from plasma derived from 6–12-wk-old mice, anti-C-terminal OCN antibody (Ferron et al., 2010b) or goat control IgG (R&D Systems) was conjugated to 2.8-µm superparamagnetic M-270 Epoxy Dynabeads according to the manufacturer’s instructions (Thermo Fisher Scientific), with a coupling ratio of 7 µg of antibody per 1 mg of beads. Covalent conjugation was done at 37°C for 24 h with end-over-end rotation. Coupled beads were washed with PBS before incubation with plasma to prevent detergent contamination. Plasma pooled from 6–12-wk-old female C57BL/6J mice was incubated with beads coupled either to anti-C-terminal OCN antibody or goat control IgG at 4°C overnight with end-over-end rotation, after which the plasma was dialyzed, aliquoted, and kept frozen at −80°C until use. An aliquot of plasma treated with either anti-C-terminal OCN antibody or goat control IgG was analyzed using an OCN ELISA to confirm significant reduction of OCN in the immunodepleted plasma.

**Osmotic pumps**

Alzet micro-osmotic pumps (model 1002) were loaded with saline or uncarboxylated OCN (30 ng/hr for 12-mo-old and 90 ng/hr for 16-mo-old mice). Mice were anesthetized with isoflurane, and osmotic pumps were surgically installed subcutaneously in the backs of the mice for 28 d.

**Gene expression analysis by quantitative PCR**

All dissections were performed in ice-cold PCR 1× under a Leica MZ8 dissecting light microscope. All parts of the brain isolated were snap frozen in liquid nitrogen and kept at −80°C until use. Total RNA was isolated from brain tissue and hippocampal cells using TRIzol (Invitrogen). Total RNA were
Figure 4. BDNF as a mediator of OCN function in the brain. (A) Representation of the microfluidic device (top) composed of somatic and synaptic chambers connected by 500-µm-long microchannels. BDNF-mCherry-positive vesicles were imaged in a 100-µm-long distal region of axons for 30 s. Kymographs (bottom) extracted from the movies. Examples of traces of anterograde (green), retrograde (orange), and pausing (blue) vesicles are depicted on the kymograph. Bar, 10 µm. (B) Quantification of BDNF-mCherry transport in axons of hippocampal neurons cultured in microchambers treated with OCN (n = 108) or vehicle (n = 110) for 4 h (Mann-Whitney U test) compared with control condition. Anterograde and retrograde mean velocities of vesicles per axon, the number of anterograde and retrograde vesicles that are motile in a 100-µm length of axon during 30 s, the percentage of time spent by vesicles in pause, and the directional flux of vesicles within axons were measured. (C and D) Bdnf expression (quantitative PCR [qPCR]) in midbrain and hippocampi
first incubated with DNase I for 30 min at room temperature to remove any genomic DNA. DNase I-treated total RNA were converted to cDNA by using M-MLV reverse transcription (28025013; Thermo Fisher Scientific) and random hexamers (N8080127; Thermo Fisher Scientific). Quantitative PCR was performed on a CFX96 Touch Real-Time PCR Detection System (Bio-Rad), and analyses were done using specific quantitative PCR primers and expressed relative to Gapdh levels. Quantitative PCR primer sequences for Gpr158 were as follows: forward, 5′-GCTCATCCTGTGGGAAAACATT-3′; reverse, 5′-TGGCTCAAGTACAAGATAACGAC-3′; Bdnf: forward, 5′-AGTCTCCAGGAGCCAAGC-3′; reverse, 5′-TGCAACGCCAGTATGAAATAACC-3′; Gapdh: forward, 5′-AACATCCATCGGCGGCTC-3′; reverse, 5′-CCATTTTGTCTACGGGACGA-3′; and Gpcre6a: forward, 5′-GGGGTTCCTAAAATACCGGACTGC-3′; reverse, 5′-TGCGATGCTTACATACACTCCA-3′.

In situ hybridization

In situ hybridization was performed on cryosections from tissue fixed with 4% PFA and using 35S-labeled riboprobe as previously described (Ducy et al., 1997). The sequence of Th, Gpr158, Gpcre6a, Gpr156, Gpcre5a, Gpcre5b, and Gpcre5d probe are 500-bp to 1-kb fragments of each gene’s 3′ UTR. Hybridizations were performed overnight at 58°C, and washes were performed at 63°C.

Hormonal and second messenger measurement

Circulating levels of OCN in mouse serum were determined with a specific ELISA previously described (Ferron et al., 2010a,b). Total levels of OCN and carboxylated OCN were estimated using two different specific antibodies. Bioactive OCN was determined by subtracting the carboxylated OCN levels from total OCN levels (Ferron et al., 2010a,b). cAMP accumulation was measured by using the cAMP Parameter Assay kit (50-900-648; R&D Systems) and performed in primary hippocampal neurons following the manufacturer’s instructions. IP1 accumulation was determined in primary hippocampal neurons by using the IP-One ELISA assay kit (KGE002B; Cisbio) following the manufacturer’s instructions. Results are calculated by dividing the optical density of the treated cells by that of the untreated cells and expressed as a percentage of stimulation according the manufacturer’s recommendations.

Recombinant OCN

Mouse uncarboxylated OCN was purified from BL21 transformed with pGEX2TK-mOCN as previously described (Lee et al., 2007; Ferron et al., 2010a; Oury et al., 2011, 2013; Mera et al., 2016). In brief, GST-OCN fusion protein was bacterially produced in BL21 pLysS transformed with pGEX-2TK-mOCN after induction with IPTG. Cells were collected in lysis buffer (PBS 1×, 10 mM Tris, pH 7.2, 2 mM EDTA, 1% Triton, and 1× protease and phosphatase inhibitor cocktail; 78443; Thermo Fisher Scientific). Following four freeze-thaw cycles and sonication, lysates were cleared by centrifugation. The supernatant was incubated with glutathione-Sepharose 4B (17075601; GE) for 4 h at 4°C. Following six washes with washing buffer (PBS 1× and 1% Triton) and with PBS 1×, OCN was then cleaved out from the GST moiety by using thrombin (27-0846-01; GE). Four fractions were collected, and each of them was incubated with benzamidine Sepharose (17-5123-10; GE) for 30 min at room temperature to remove thrombin.

Immunoblotting

Frozen hippocampi were pulverized in lysis buffer (20 mM Tris-HCl, pH 7.65, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, and 1% Triton) supplemented with 1× protease and phosphatase inhibitor cocktail (78443; Thermo Fisher Scientific). Lysates were clarified by centrifugation, and protein concentration was determined using a Bradford assay. Clarified samples were heated at 65°C in Laemmli buffer for 5 min. Proteins were run on an SDS-PAGE gel and transferred to nitrocellulose membrane (1620112; Bio-Rad). Membranes were blocked with TBST-5% BSA for 1 h, incubated with primary antibody overnight at 4°C, and probed with HRP-conjugated antibodies for 1 h at room temperature. The following antibodies were used in these studies: anti-TIMP2 (1:1,000; D18B7; Cell Signaling), anti-α-tubulin (1:5,000; T6199; Sigma-Aldrich), anti-b2 microglobulin (1:1,000; ab75853; Abcam), antitransferrin (1:1,000; ab82411; Abcam), anti-Gpr158 (1:1,000; ABIN486340; Antibodies-online), anti-Na,K ATPase (1:1,000; 3010S; Cell Signaling), anti-SR2C (1:1,000; sc-17797; Santa Cruz), anti-Gaq (1:1,000; D5V1B; Cell Signaling), anti-BDNF (1:100; sc-65514; Santa Cruz). Band intensities were quantified using ImageJ software. Dashed lines separating two bands indicate that these bands are on the same membrane but are not adjacent.
Immunohistochemistry
Primary cultured neurons were fixed with 4% PFA and 4% sucrose for 20 min and blocked in PBS containing 3% BSA and 0.2% Tween 20 for 2 h at room temperature. Primary antibodies, anti-MAP2 (MAB364; EMD Millipore), anti-glia fibrillary acidic protein (SAB2500462; Sigma-Aldrich), and anti-Gpr158 (ABIN486340; Antibodies-online) were diluted (1:400) in blocking solution and incubated overnight at 4°C. Coverslips were washed three times with PBS 1× containing Triton 0.2%, and secondary antibodies (1:200; Jackson ImmunoResearch) diluted in blocking solution were incubated for 1 h 30 min at room temperature. After three washes, coverslips were mounted with ProLong antifade mountant (Life Technologies). Images were obtained with an inverted Zeiss APO Tome 2 microscope. Immunofluorescence of adult brains was performed on 35-µm coronal vibratome slices of tissue fixed with 4% PFA in 0.1 M phosphate buffer. Sections were permeabilized and blocked with 0.5% Triton in TBS, pH 7.4, and 10% normal donkey serum. After blocking, sections were incubated with primary antibodies overnight at 4°C. The following antibodies were used for this assay: anti-c-Fos (1:400; 2250; Cell Signaling), anti–β-galactosidase (1:400; ab4761; Abcam), anti-MAP2 (1:1,000; ab5392; Abcam), and anti–glial fibrillary acidic protein (SAB2500462; Sigma-Aldrich). Sections were washed three times in 0.3% Triton in TBS, pH 7.4, followed by an incubation with secondary antibodies (1:200; Invitrogen) diluted in blocking solution and incubated for 1 h at room temperature. After three washes, sections were mounted with Fluorogel (Electron Microscopy Sciences) and visualized using a Nikon Ti Eclipse microscope. Images were analyzed using ImageJ version 2.0.

Pull-down assay
Pulldown of Gpr158 was performed in solubilized membranes from Ocn−/− or Gpr158−/− hippocampi. Hippocampi were dissected on ice and homogenized in buffer A (10 mM Tris-HCl, pH 7.4, 320 mM sucrose, 1× protease, and phosphatase inhibitor cocktail [78443; Thermo Fisher Scientific]) with a Glass/Teflon Potter Elvehjem homogenizer (20 strokes). Homogenized hippocampi were centrifuged at 3,000 g for 10 min at 4°C. Then, supernatants were ultracentrifuged at 40,000 g for 20 min at 4°C. Pellets were resuspended in buffer A supplemented with 150 mM NaCl and 1% n-Octyl β-D-thioglycopyranoside (28310; Thermo Fisher Scientific). Solubilized membranes were diluted in buffer A supplemented with 150 mM NaCl and 0.2% n-Octyl β-D-thioglycopyranoside. Recombinant uncarboxylated OCN was biotinylated using the EZ-Link NHS-PEO4 biotinylation kit (21455; Thermo Fisher Scientific) according to the manufacturer’s instructions. For the pulldown, biotinylated OCN (7 µg) was incubated for 2 h at 4°C. Thirty microliters of Dynabeads M-280 Streptavidin (11205D; Thermo Fisher Scientific) were added for 30 min at room temperature followed by three PBS 1× washes. Purified proteins were eluted from the beads by adding Laemmli protein buffer and heated at 65°C for 15 min and analyzed by Western blot.

Behavioral tests
All animals of the same batch were born within an interval of 2 wk and were kept in mixed genotype or groups of two to five females in the same cage, at standard laboratory conditions (12-h dark/light cycle, constant room temperature and humidity, and standard laboratory chow and water ad libitum). For each test, mice were transported a short distance from the holding mouse facility to the testing room in their home cages. Behavioral testing was performed between 3 and 16 mo of age, and mouse weight was between 22 and 32 g. The tests were performed by an experimentalist blind to the genotypes or treatment of the mice under study.

EPMT. This test takes advantage of the aversion of rodents to open spaces. The elevated plus maze apparatus comprises two open and two enclosed arms, each with an open roof, elevated 60 cm from the floor (Holmes et al., 2000; Lira et al., 2003). Testing takes place in dim ambient light conditions. Animals are placed into the central area facing one closed arm and allowed to explore the elevated plus maze for 6 min. The total number of arm entries and time spent in open arms is recorded. An increase in anxiety is indicated by a decrease in the proportion of time spent in the open arms (time in open arms/total time in open or closed arms) and a decrease in the proportion of entries into the open arms.

Light-to-dark transition test. This test is based on the innate aversion of rodents to brightly illuminated areas and on their spontaneous exploratory behavior in response to the stressor that light represents. The test apparatus consists of a dark, safe compartment and an illuminated, aversive one. Mice were tested for 6 min, and two parameters were recorded: time spent in the lit compartment and the number of transitions between compartments, indices of anxiety-related behavior and exploratory activity.

OFT. This test takes advantage of the aversion of rodents to brightly lit areas (David et al., 2009). Each mouse is placed in the center of the OFT chamber (a white 43 × 43 cm chamber) and allowed to explore for 30 min. Mice were monitored throughout each test session by video recording and analyzed using autotyping (Patel et al., 2014). The overall motor activity was quantified as the total distance traveled (ambulation). Anxiety was quantified by measuring the time spent in the center of the OFT chamber.

Morris water maze test. Animals were transported to the testing room in their home cages and left undisturbed for at least 30 min before the first trial. The maze was comprised of a large swimming pool (150 cm in diameter) filled with water (23°C) made opaque with nontoxic white paint. The pool was located in a brightly lit room filled with visual cues, in
including geometric figures on the walls of the maze demarking the four fixed starting positions of the trials, at 12 o’clock, 3 o’clock, 6 o’clock, and 9 o’clock. A 15-cm round platform was hidden 1 cm beneath the surface of the water at a fixed position. Each daily trial block consisted of four swimming trials, with each mouse starting from the same randomly chosen starting position. The starting position varied between days. On day 1, mice that failed to find the platform within 2 min were guided to the platform. They were required to remain on the platform for 15 s before they were returned to their home cages. Mice were not guided to the platform after day 1, and the time it took them to reach the platform over repeated trials (three trials each day for the next 10 d) was recorded as a measure of spatial learning.

**NOR test.** The NOR paradigm assesses a rodent’s ability to recognize a novel object in the environment. The NOR task was conducted, as previously described (Broadbent et al., 2010), in an opaque plastic box using two different objects: (a) a clear plastic funnel (diameter 8.5 cm, maximal height 8.5 cm) and (b) a black plastic box (9.5 cm³). These objects elicit equal levels of exploration as determined in pilot experiments (Denny et al., 2012; Oury et al., 2013). The NOR paradigm consists of three exposures over the course of 3 d. On day 1, the habituation phase, mice are given 5 min to explore the empty arena, without any objects. On day 2, the familiarization phase, mice are given 10 min to explore two identical objects, placed at opposite ends of the box. On day 3, the test phase, mice are given 15 min to explore two objects, one novel object and a copy of the object from the familiarization phase. The object that serves as the novel object (either the funnel or the box) as well as the left/right starting position of the objects is counterbalanced within each group. Mice are placed in the center of the arena at the start of each exposure. Between exposures, mice are held individually in standard cages, the objects and arenas are cleaned, and the bedding is replaced. Preference for the novel object is assessed on the basis of the fraction of time that a mouse spends exploring the novel object compared with the familiar object. Exploration is scored from video recordings of each exposure and scored using the Stopwatch program. Climbing or sitting adjacent to the object were not counted as exploration. An equal exploration time for the two objects, or a decreased percentage of time spent with the novel object compared with WT controls indicates impairment in hippocampal memory. For this test, discrimination index (time spent with novel object – time spent with old object)/total exploration time) and preference index (time spent with novel object/total exploration time) were measured.

**Contextual fear conditioning.** The conditioning apparatus consisted of sound-attenuating conditioning boxes (Bioseb), and mice were run individually in the conditioning boxes. The floor of the chamber consisted of stainless steel bars wired to a shock generator with a scrambler for the delivery of foot shock. Signal generated by mouse movement was recorded and analyzed by a high-sensitivity weight transducer system. The fear-conditioning procedure took place over two consecutive days. On day 1, mice were placed in the conditioning chamber and received three foot shocks (1 s, 0.5 mA), which were administered at time points of 60, 120, and 180 s after the animals were placed in the chamber. They were returned to their home cage 60 s after the final shock. Contextual fear memory was assessed 24 h after conditioning by returning the mice to the conditioning chamber and measuring freezing behavior during a 4-min retention test. Freezing was scored and analyzed automatically using Packwin software version 2.0 (Bioseb). Freezing behavior was considered to occur if the animals froze for a period of at least 2 s.

**Electrophysiology**

Coronal brain slices containing the hippocampus were prepared from WT and Gpr158⁻/⁻ mice (males, 3–4 wk old), as previously reported (Rao et al., 2007). In brief, mice were anesthetized with isoflurane and then decapitated to harvest brains, which were rapidly removed and immersed in an oxygenated cutting solution at 4°C containing 220 mM sucrose, 2.5 mM KCl, 1 mM CaCl₂, 6 mM MgCl₂, 1.25 mM NaH₂PO₄, 26 mM NaHCO₃, and 10 mM glucose and adjusted to pH 7.3 with NaOH. Coronal slices containing the hippocampus (350 μm thick) were cut with a vibratome and trimmed to contain just the hippocampus. After preparation, slices were stored in a holding chamber at room temperature (22–25°C) with oxygenated (5% CO₂ and 95% O₂) artificial cerebrospinal fluid (ACSF) containing 124 mM NaCl, 3 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 1.23 mM NaH₂PO₄, 26 mM NaHCO₃, and 10 mM glucose, pH 7.4, with NaOH. The slices were eventually transferred to a recording chamber continuously perfused with ACSF at 33°C at a rate of 2 ml/min for at least 1 h to allow recovery before the start of patch-clamp recording. Whole-cell current clamp was performed to observe spontaneous APs in visually identified pyramidal neurons in the CA3 area of the hippocampus, as reported by Kim and Connors (2012), with a Multiclamp 700 A amplifier (Molecular Devices). The patch pipettes with a tip resistance of 4–6 MΩ were made of borosilicate glass (World Precision Instruments) with a pipette puller (Sutter P-97) and back filled with a pipette solution containing 135 mM K-glucuronate, 2 mM MgCl₂, 10 mM Hepes, 1.1 mM EGTA, 2 mM Mg-ATP, 10 mM Na₂-phosphocreatine, and 0.3 mM Na₂-GTP, pH 7.3, with KOH. After a stable base of APs were recorded for 10 min, OCN was applied to the recorded cells through bath application at a concentration of 10 ng/ml for 10 min and then washed out with ACSF. Both input resistance and series resistance were monitored throughout the experiments. Only recordings with stable series resistance were accepted (the change in series resistance was less than 20%). In LTP experiments, fEPSPs were recorded in the CA3 area with a glass pipette filled with ACSF (with a tip resistance of 2–4 MΩ) by stimulating the MF in the dentate gyrus area...
with a bipolar electrode. The amplitude of fEPSPs was always adjusted to 40% of the maximum value before the starting of LTP experiments. Two trains of high-frequency stimuli (100 Hz, 1-s duration, 10-s interval) were delivered to the MF after a stable baseline recording of fEPSPs was achieved, as routinely reported by others (Spillane et al., 1995). The amplitude of fEPSPs was normalized to the mean amplitude of fEPSPs recorded before high-frequency stimulations, as reported by others (Spillane et al., 1995). All data were sampled at 10 kHz and filtered at 6 kHz with an Apple Macintosh computer using Axograph X (AxoGraph Scientific). APs were detected and analyzed with AxoGraph X and plotted with Igor Pro software (WaveMetrics), as described previously (Rao et al., 2007). 2-Aminoethoxydiphenyl borate (1224; Tocris Biosciences) was used to inhibit IP3 receptor.

**Lentivirus production and shRNA sequences**

The shRNA sequence 5′-GAGCGCCTCCACTGAGCGACGGACCATCTTGGA-3′ targeting mouse Gpr158 was purchased from Origene (TL50693B); shRNA was cloned into pGFP-C-Lentivector (Origene). For virus production, human embryonic kidney 293T cells were transfected with VSVG, helper, and pGFP-C-shLentiVector (containing Gpr158 or scramble shRNA) plasmids. After 2 d, viral particles were purified and concentrated.

**Stereotaxic injection**

Stereotaxic surgery was performed in 3-mo-old C57BL/6j male mice obtained from the Janvier Laboratory stock. Mice were anesthetized with intraperitoneal injection of a mixture of ketamine hydrochloride (100 mg/kg of body weight [BW]; 1000; Virbac) and 100 mg/ml BW xylazine (10 mg/kg of BW; Rompun 2%; Bayer) and placed in a stereotaxic frame (900SL-KOPF). Ophthalmic eye ointment was applied to the cornea to prevent desiccation during surgery. The fur in the area surrounding the incision was trimmed, and Vetedine solution (Vetiquinol) was applied. Lentiviruses expressing shRNA targeting Gpr158 or noneffective scramble shRNA in pGFP-C-shLentiVector were injected bilaterally into the anterior hippocampi ([from bregma] anteroposterior = −2.0 mm, dorsal vertical = ±1.4 mm, and medial lateral = ±1.33 mm) using a 10-µl Hamilton syringe (1701RN) over either 12 or 4 min (injection speed 0.25 µl/min). Three microliters of lentivirus expressing shRNA against Gpr158 (titer: 3.4 × 10^9 GC/ml) or scramble shRNA (1.4 × 10^9 GC/ml) were injected. To limit reflux along the injection track, the needle was maintained in place for 4 min between each 1-µl injection. Then, the skin was closed using silk suture and the mice were injected locally with surgical analgesic (ketoprofen). Two weeks after the lentivirus injection, bilateral stereotaxic injections of OCN (10 ng/µl) or NaCl was performed into the anterior hippocampi at the same stereotaxic coordinate. Twelve hours after the stereotaxic injections of OCN, mice were subjected to the training phase (NOR and contextual fear conditioning) and to the testing phase 24 h following the habitation phase.

**Live-cell imaging of BDNF axonal trafficking in microfluidic devices**

Primary hippocampal neurons were prepared from E17.5 WT rats. Dissociated neurons were resuspended in growing medium (Neurobasal medium supplemented with 2% B27, 2 mM Glutamax, and 1% penicillin/streptomycin) and seeded in the upper chamber of microchambers prepared according to Zala et al. (2013) at a final density of 7,000 cells/mm². Three hours after seeding, neurons were infected with 3.10^5 UI of lentivirus carrying pCMV- BDNF-mCherry constructs and cultured for 9 to 11 d at 37°C in a 5% CO2 incubator. Neurons were treated or not for 4 h with 10 ng/ml OCN added in the upper chamber and imaged. Videomicroscopy acquisitions of 100-µm axon length were done on the lower part of channels, far enough from dendrites, at 5 Hz for 30 s with an inverted microscope (AxioObserver Z1; Zeiss) coupled to a spinning-disk confocal system (CSU-W1-T3; Yokogawa) connected to an electron-multiplying charge-coupled device camera (ProEM’1024; Princeton Instrument) at 37°C and 5% CO2. BDNF-mCherry positive mean vesicle velocity per axon, vesicle number per 100 µm length axon over 30 s, percentage of vesicle pausing time, and directional net flux (mean velocity × number of anterograde vesicles per axon − mean velocity × number of retrograde vesicles per axon) were extracted from kymographs (x axis, 100 µm; y axis, 30 s) using the KymoTool Box ImageJ plugin as previously described (Zala et al., 2013). Each condition was tested using 3 microchambers per culture and three independent cultures. In each chamber, three fields containing at least four axons were analyzed to reach a minimum number of 100 axons (1,400 vesicles).

**Quantification and statistical analysis**

All values are expressed as mean ± SEM. Statistical parameters including the exact sample size (n), post hoc tests, and statistical significance are reported in every figure and figure legend. Number of mice were estimated to be sufficient on the basis of pilot experiments and previously published work (Oury et al., 2013). Data were estimated to be statistically significant when P ≤ 0.05 by nonparametric Mann-Whitney U test, Student’s t test, one-way ANOVA, or two-way ANOVA. In every figure, an asterisk denotes statistical significance (*, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001). Data were analyzed using GraphPad Prism 6.

**Online supplemental material**

Fig S1 shows bioactive OCN levels in various animal models, as well as the peripheral treatment of 12-mo-old mice with OCN. Fig. S2 shows the negative control for the expression pattern of all class C orphan family GPCRs in the hippocampal region and the stereotactic coordinates used for hip-
pocampal injections. Fig. S3 presents how the brain-specific deletion of Gpr158 was performed.

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