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Oxytocin stimulates hippocampal neurogenesis via oxytocin receptor expressed in CA3 pyramidal neurons

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In addition to the regulation of social and emotional behaviors, the hypothalamic neuropeptide oxytocin has been shown to stimulate neurogenesis in adult dentate gyrus; however, the mechanisms underlying the action of oxytocin are still unclear. Taking advantage of the conditional knockout mouse model, we show here that endogenous oxytocin signaling functions in a non-cell autonomous manner to regulate survival and maturation of newly generated dentate granule cells in adult mouse hippocampus via oxytocin receptors expressed in CA3 pyramidal neurons. Through bidirectional chemogenetic manipulations, we also uncover a significant role for CA3 pyramidal neuron activity in regulating adult neurogenesis in the dentate gyrus. Retrograde neuronal tracing combined with immunocytochemistry revealed that the oxytocin neurons in the paraventricular nucleus project directly to the CA3 region of the hippocampus. Our findings reveal a critical role for oxytocin signaling in adult neurogenesis.

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Oxytocin (OXT) is a nine amino acid neuropeptide that is primarily synthesized in magnocellular neurons of the hypothalamic paraventricular (PVN) and supraoptic nuclei (SON)1, 2. Apart from its release into the systemic circulation via the posterior pituitary, OXT is also transported axonally from hypothalamic parvocellular neurons to numerous extra-hypothalamic OXT receptor (OXTR)-expressing brain regions, including the hippocampus, amygdala, lateral septum, striatum, and bed nucleus of the stria terminals, acting as either a neurotransmitter or neuromodulator to regulate neurotransmission within these regions3–5. OXT has both peripheral and central functions. Peripheral OXT promotes uterine contractions during parturition and milk ejection during lactation6–9. Centrally acting OXT mediates its biological activities by binding to the OXTR, which belongs to the superfamily of G-protein-coupled receptors10. Growing evidence suggests that the hippocampus is one of the brain structures particularly vulnerable to the effects of OXT. For example, microinjection of OXT into the dorsal hippocampus has been shown to attenuate stress-induced neuroendocrine and behavioral responses in rats11. In addition, we and others have previously shown that OXT can promote the maintenance of long-term potentiation (LTP) in hippocampal CA1 region and enhance spatial memory during motherhood12, 13. Despite these study points toward a crucial role for OXT in facilitating hippocampal plasticity, and function, exogenous application of OXT has been reported to exert a neurotrophic effect to increase adult neurogenesis even when experienced to stressful situations14. Nonetheless, how OXT regulates adult neurogenesis remains an unresolved problem. Furthermore, it is not yet clear whether endogenous OXT signaling may also play a role in regulating adult hippocampal neurogenesis. In this study, we addressed the following three questions. First, is OXTR expressed in adult hippocampal neural progenitor cells? Second, does endogenous OXT signaling regulates specific stages of adult hippocampal neurogenesis? Finally, does endogenous OXT signaling regulates adult neurogenesis through a cell autonomous or non-cell autonomous mechanism? Using a Cre/loxP recombinase-based strategy to delete Oxtr, we report here that endogenous OXT signaling controls adult hippocampal neurogenesis through an indirect non-cell autonomous mechanism by OXTR expressed in CA3 pyramidal neurons.

Results

OXTR is not expressed in the neural progenitor cells. Adult neurogenesis is a multistep process that comprises the proliferation of neural progenitor cells, newborn neuron differentiation, maturation, and functional integration into the preexisting neural networks16, 17. We first examined OXTR expression in developing dentate granule cells (DGCs) in adult mouse hippocampus by staining for proteins expressed at distinct stages of cell differentiation. Because an immunofluorescent staining using antibody is not a reliable technique for identifying OXTR-expressing neurons in mice18, we therefore used OXTR-Venus knock-in (OxtrVenus-Neo<sup>fl/fl</sup>) mice to characterize OXTR-expressing neurons in the hippocampus. In 10-week-old mice, unexpectedly, we found few or no Venus-positive cells in the dentate gyrus (DG) expressing progenitor markers, such as nestin (Fig. 1a), Ki67 (Fig. 1b), or neuroblast and immature neuronal marker doublecortin (DCX; Fig. 1c). However, in virtually all Venus-positive cells coexpressed the mature neuronal-specific nuclear protein (NeuN; Fig. 1d). Immunofluorescence staining was also performed with antibodies against numerous known marker proteins expressed in the mature granule cell (calbindin) and the subtypes of GABAergic interneurons (calretinin and parvalbumin). In the hilus, 77.4 ± 2.3% of Venus-positive cells expressed calretinin immunoreactivity (Fig. 1e) and 25.0 ± 2.9% Venus-positive cells expressed parvalbumin immunoreactivity (Fig. 1f). However, no Venus-positive cells were found that expressed calbindin (Fig. 1g). To label proliferating cells, we administered a synthetic analog of the nucleoside thymidine 5-bromo-2′-deoxyuridine (BrdU) intraperitoneally and analyzed BrdU incorporation 2 or 4 weeks later. We detected no Venus-positive cells expressing BrdU immunoreactivity (Fig. 1h, i). We further characterize hilar and CA3 OXTR distribution regarding to overlap with GABA immunoreactivity. In the hilus, 27.8 ± 3.8% of Venus-positive cells were immunoreactive for the GABAergic marker glutamic acid decarboxylase (GAD)67. In the CA3, 3.8 ± 0.7% of Venus-positive cells expressed GAD67 immunoreactivity (Supplementary Fig. 1a, b). These results indicate that OXTR is not expressed in the neural progenitor cells that reside within the subgranular zone or mature granule cells of adult DG and a small subset of hilar GABAergic neurons express OXTR.

Deletion of Oxtr from hippocampal excitatory neurons. Previous studies have shown that CA3 pyramidal neurons send reciprocal projections back to the DG and regulate neurogenesis in the DG of adult rats21. Since OXTR is enriched in the CA2 and CA3 of the hippocampus20, we therefore hypothesized that OXT may control adult hippocampal neurogenesis via OXTR expressed in CA3 pyramidal neurons. Consistent with previous findings20, our immunofluorescent staining data showed strong Venus immunoreactivity in the hippocampal CA3 region of Oxtr<sup>Venus-Neo<sup>fl/fl</sup></sup> mice. We found that more than 95% Venus-positive cells expressed calcium/calmodulin-dependent protein kinase II α (CaMKIIα) immunoreactivity (Fig. 2a), indicating that OXTR predominantly expressed in excitatory pyramidal neurons. To test whether OXTR expressed in CA3 pyramidal neurons plays a role in regulating adult DG neurogenesis, we used the Cre-loxP recombination approach to conditionally delete Oxtr from hippocampal excitatory neurons by crossing mice expressing CaMKIIα-Cre with mice in which Oxtr was floxed (Oxtr<sup>fl/fl</sup>). Polymerase chain reaction (PCR) screening of mouse genomic tail DNA confirmed heterozygous (Oxtr<sup>+/−</sup>) and homozygous Oxtr<sup>−/−</sup> conditional knockout mice (Fig. 2b). Quantitative real-time PCR analysis confirmed a reduction in Oxtr mRNA expression in the CA2, CA3, and hypothalamus, but not the hilus of the DG, in Oxtr<sup>−/−</sup> mice compared with wild-type (WT, Oxtr<sup>+/+</sup>) mice (Fig. 2c). In parallel, fluorescence in situ hybridization (FISH) with Oxtr gene probe also revealed that the numbers of Oxtr mRNA-positive cells in the CA2 and CA3 of Oxtr<sup>−/−</sup> mice were markedly reduced compared with WT mice (Fig. 2d, e), confirming the efficiency of Cre-loxP-mediated deletion of Oxtr. Dual-probe FISH also confirmed that the majority of Oxtr mRNA-positive cells were CaMKIIα mRNA-expressing cells in the CA3 (Supplementary Fig. 2a). Very few Oxtr mRNA immunoreactivity was detected in the granule cell layer of the dorsal and ventral DG (Supplementary Fig. 2b, c). Although Oxtr mRNA immunoreactivity was detected in dorsal...
and ventral hilus of the DG, we did not observe colocalization of Oxtr mRNA with CaMKIIα mRNA. We also demonstrated that Oxtr−/− mice specifically displayed a reduction of Oxtr mRNA expression in the CA3, but not the hilus of the DG, compared with WT mice (Supplementary Fig. 2).

**Effects of Oxtr deletion on newly generated DGCs.** To determine whether Oxtr deletion from hippocampal excitatory neurons may affect the proliferation of newly generated DGCs, WT and Oxtr−/− mice were given a single injection of BrdU, and double fluorescent labeling for detection of BrdU-positive (BrdU+) proliferating cells (Ki67+) was performed on both dorsal and ventral hippocampal sections 2 h later (Fig. 3a, c). Stereological analysis revealed no differences in the total number of either BrdU+ (Fig. 3d) or Ki67+ cells (Fig. 3e) between WT and Oxtr−/− mice at 2 h after the BrdU injection in both dorsal and ventral hippocampus (Supplementary Fig. 3a–d). The percentage of BrdU+Ki67+ cells to total BrdU+ cells was not different between the two groups (Supplementary Fig. 3i). To determine whether conditional deletion of Oxtr from hippocampal excitatory neurons may affect the survival of newly generated DGCs, WT and Oxtr−/− mice were subjected to multiple BrdU injections, and double fluorescent labeling for detection of BrdU+ immature (DCX+) or mature neuronal cells (NeuN+) was performed on hippocampal sections 14 or 28 days later (Fig. 3b, c). Interestingly, we found a significant reduction in the total number of BrdU+ cells in Oxtr−/− mice at 14 and 28 days after BrdU injection compared with WT mice (Fig. 3f). Likewise, a significant reduction in the total number of DCX+ cells in Oxtr−/− mice at 14 days after BrdU injection compared with WT mice in both dorsal and ventral hippocampus (Fig. 3g and Supplementary Fig. 3e–h). However, the percentage of BrdU+DCX+ cells or BrdU+NeuN+ cells to total BrdU+ cells was not different between two groups (Supplementary Fig. 3j, k). These results suggest that OXTR plays a non-cell autonomous role in controlling the survival of newly generated DGCs, whereas the progenitor cell proliferation and early differentiation into immature neurons were unaltered by Oxtr deletion.

We next evaluated the impact of Oxtr deletion on the morphological maturation of newly generated DGCs by using a retrovirus-mediated birth-dating and labeling strategy. Engineered retroviruses expressing enhanced green fluorescent protein (EGFP) were stereotaxically microinjected into the DG hilus of WT and Oxtr−/− mice. Sholl analysis for dendritic complexity of EGFP-positive (EGFP+) DGCs was carried out 14
or 28 days post infection (dpi). Using confocal microscopy to reconstruct the dendritic arborization of EGFP+ DGCs, we found that newly generated DGCs from Oxtr−/− mice exhibited less elaborated dendritic arborization than those from WT mice at 14 dpi (Fig. 4a). Unpaired Student’s t test revealed a significant reduction in both total dendritic length (P < 0.001; Fig. 4b) and branch number (P < 0.01; Fig. 4c) in EGFP+ DGCs from Oxtr−/− mice compared to those from WT mice. Sholl analysis further revealed a significant decrease in the dendritic complexity of EGFP+ DGCs in Oxtr−/− mice compared to WT mice at 14 dpi, an effect more pronounced for the dendritic branches extended beyond 90–110 μm from the soma (Fig. 4d). However, we observed no significant differences in total dendritic length, branch number, or dendritic complexity of EGFP+ DGCs between Oxtr−/− and WT mice at 28 dpi (Fig. 4e–h).

To further confirm OXT controls adult hippocampal neurogenesis through an indirect non-cell autonomous mechanism, we therefore examined morphological maturation of newly generated DGCs by using Oxtrfloxed mice in combination with Cre recombinase-mediated gene deletion in a localized fashion through bilateral stereotaxic injections of Cre-expressing (Retro-Cre-EGFP) or control EGFP-expressing (Retro-EGFP) retroviral vectors targeting the hilus of the DG (Fig. 4i). We observed no significant differences in total dendritic length or dendritic complexity of EGFP+ DGCs between two groups at 14 dpi (Fig. 4j, k).
To assess whether the morphological changes that resulted from conditional Oxtr deletion may lead to alterations in physiological properties, we performed electrophysiological analysis of newly generated DGCs in acute slices from retrovirus-injected mice at 14 dpi. We first examined glutamatergic synaptic transmission by recording spontaneous excitatory postsynaptic currents (sEPSCs) in EGFP⁺ DGCs in the presence of GABA_A receptor antagonist gabazine (10 μM; Fig. 5a). A significant difference in cumulative interval interval distribution was observed between newly generated DGCs from Oxtr⁻/⁻ and WT mice (Fig. 5b). However, no significant difference in cumulative amplitude distribution was observed between two groups (Fig. 5c). The mean frequency of sEPSCs in EGFP⁺ DGCs from Oxtr⁻/⁻ mice was significantly less than those from WT mice (P < 0.05; Fig. 5d). No significant difference was observed between Oxtr⁻/⁻ and WT mice in the mean amplitude of sEPSCs (P = 0.88; Fig. 5e).

The newly generated DGCs exhibit initial depolarizing response to GABA, which gradually shift to hyperpolarizing responses over a period of 2–3 weeks of birth. The polarity of GABA actions depends on the chloride concentration gradient across the plasma membrane. We next used gramicidin-perforated patch-clamp recordings to determine the nature of GABA activation by recording evoked inhibitory postsynaptic currents (IPSCs) in EGFP⁺ DGCs from Oxtr⁻/⁻ and WT mice at 10, 14, and 28 dpi (Fig. 5f). In agreement with previous findings, we found that the reversal potential for GABA_A receptor-mediated IPSCs (E_GABA) in EGFP⁺ DGCs gradually decreased during maturation. Two-way ANOVA revealed a main effect of conditional Oxtr deletion on E_GABA (F{(1,18)} = 25.1, P < 0.001; Fig. 5g). Bonferroni’s post hoc tests showed that EGFP⁺ DGCs from Oxtr⁻/⁻ mice demonstrated more positive values for E_GABA compared to those from WT mice at both 10 (P < 0.01) and 14 dpi (P < 0.05). To determine whether more positive values for E_GABA could be due to higher intracellular chloride concentrations ([Cl⁻]), we used the Nernst equation to calculate [Cl⁻]. Two-way ANOVA revealed a main effect of conditional Oxtr deletion on [Cl⁻], F{(1,18)} = 23.7, P < 0.01; Fig. 5h). Bonferroni’s post hoc tests showed that GFP⁺ DGCs from Oxtr⁻/⁻ mice had higher [Cl⁻] than those from WT mice at both 10 (P < 0.0001) and 14 dpi (P < 0.01). These results indicate that conditional Oxtr deletion may delay maturation of newly generated DGCs.

We next evaluated whether Oxtr deletion impacted mature DGCs. To characterize the morphological and physiological properties, we performed electrophysiological analysis of newly generated DGCs in WT and Oxtr⁻/⁻ mice at 14 and 28 days after the last BrdU injection.

**Fig. 3** Conditional deletion of Oxtr impairs the survival of newly generated DGCs. **a** Schematic representation of the experimental designs for comparing the proliferation of newly generated DGCs in WT and Oxtr⁻/⁻ mice. Mice were given a single injection of BrdU (50 mg/kg) and were killed 2 h after BrdU injection. **b** Schematic representation of the experimental designs for comparing the survival of newly generated DGCs in WT and Oxtr⁻/⁻ mice. Mice were injected six times intraperitoneally with BrdU (50 mg/kg) at 12 h intervals and were killed 14 and 28 days after the last BrdU injection. **c** Representative immunofluorescence images of hippocampal sections from WT and Oxtr⁻/⁻ mice double or triple stained for BrdU (green), Ki67 (red), DCX (red), and NeuN (blue) after BrdU injection. Scale bar, 100 μm. **d** Quantification of the total number of BrdU⁺ cells in the DG of WT and Oxtr⁻/⁻ mice at 2 h after BrdU injection (n = 4 mice per genotype, unpaired two-tailed Student’s t test). **e** Quantification of the total number of Ki67⁺ cells in the DG of WT and Oxtr⁻/⁻ mice at 2 h after BrdU injection (n = 5 mice per genotype, unpaired two-tailed Student’s t test). **f** Quantification of the total number of BrdU⁺ cells in the DG of WT and Oxtr⁻/⁻ mice at 14 and 28 days after BrdU injection (n = 5 mice for each group; **P < 0.001, unpaired two-tailed Student’s t test). **g** Quantification of the total number of DCX⁺ cells in the DG of WT and Oxtr⁻/⁻ mice at 14 days after the last BrdU injection (n = 5 mice per genotype; ***P < 0.001, unpaired two-tailed Student’s t test). Data represent the mean ± s.e.m.
electrophysiological properties of DGCs, we recorded miniature excitatory postsynaptic currents (mEPSCs) from mature DGCs with the use of whole-cell patch-clamp recordings and biocytin was routinely included in the intracellular solution to allow post hoc identification of the recorded neurons (Fig. 6a). We observed no significant differences between Oxtr−/− and WT mice in total dendritic length ($P = 0.19$; Fig. 6b) and branch number of DGCs ($P = 0.22$; Fig. 6c). Sholl analysis revealed no significant difference in the dendritic complexity of DGCs between Oxtr−/− and WT mice ($F_{(1,225)} = 0.14, P = 0.71$; Fig. 6d). mEPSCs were recorded in the presence of tetrodotoxin (TTX, 0.5 μM) and gabazine (10 μM; Fig. 6e). A significant difference in cumulative interevent interval...
distribution was observed between mature DGCs from Oxtr−/− and WT mice (Fig. 6). However, no significant difference in cumulative amplitude distribution was observed between two groups (Fig. 6g). The mean frequency of mEPSCs in DGCs from Oxtr−/− mice was significantly less than those from WT mice (P < 0.05; Fig. 6h). No significant difference was observed between Oxtr−/− and WT mice in the mean amplitude of mEPSCs (P = 0.11; Fig. 6i).

OXT controls adult neurogenesis through CA3 neurons. Having established that conditional deletion of Oxtr results in impaired survival and maturation of newly generated DGCs, we next investigated the possible mechanisms by which OXT controls adult hippocampal neurogenesis. Based on the foregoing observations in conjunction with the suggested role of CA3 pyramidal neurons in regulating adult neurogenesis, we tested whether OXT neurons of the PVN send projections to the CA3 region of the hippocampus. To identify PVN-CA3 projection neurons, we injected a monosynaptic retrograde tracer cholera toxin B subunit (CTB) into the CA3 and examined the distribution of retrogradely CTB-labeled cells appeared in the PVN (Fig. 7a, b). We also labeled neurosecretory neurons with intraperitoneal injection of the retrograde tracer, fluoro-gold (4% w/v in 100 µl saline)34. Immunoreactivity for CTB was clearly detected in fluoro-gold and OXT + parvocellular neurons of the PVN (Fig. 7b), but not the SON (Supplementary Fig. 4a). We also observed that PVN neurons send their projecting fibers directly to the CA3 (Supplementary Fig. 4b). These results indicate that the CA3 contains a dense population of OXT-expressing neurons and receives direct inputs from OXT neurons in the PVN.

To demonstrate that CA3 pyramidal neurons are functionally modulated by OXT, we analyzed the effects of OXT on electrical membrane properties of CA3 neurons (Fig. 7c). Under whole-cell current-clamp condition, bath application of OXT (1 µM) consistently caused a membrane depolarization by 3.98 ± 1.44 mV (n = 8) in CA3 pyramidal neurons from WT mice. The OXT-induced membrane depolarization was not observed in CA3 pyramidal neurons from Oxtr−/− mice (0.98 ± 0.21 mV, n = 10; Fig. 7d, e) and was prevented by the selective OXTR antagonist injection (100 pA, 500 ms) (Fig. 7d, e). The enhancing effect of OXT on action potential firing in CA3 pyramidal neurons was also prevented by the pretreatment of the hippocampal slices with L-371257 (1 µM; Supplementary Fig. 5c, d). To further validate whether OXT may act indirectly to enhance intrinsic excitability of hippocampal interneurons through activation of CA3 OXTR + pyramidal neurons, CA3 OXTR + interneurons in slices from Oxtr−/− mice were targeted for recording, filled with biocytin to allow post hoc reconstruction (Supplementary Fig. 6a). In contrast to what has been observed in CA3 OXTR + pyramidal neurons, bath application of OXT (1 µM) did not significantly affect membrane potential in the absence of L-371257 (1 µM) in CA3 OXTR − interneurons (Supplementary Fig. 6b–d).

We employed optogenetic approach to further validate that DGCs receive a direct back projection from CA3 pyramidal neurons. We unilaterally injected the CA3 region with a recombinant adeno-associated virus capsid DJ (AAV-DJ) serotype expressing the light-sensitive channelrhodopsin 2 (ChR2) targeted with enhanced yellow fluorescent protein (EYFP), under control of the CaMKIIα promoter [AAV-CaMKIIα-hChR2 (H134R)-EYFP], favoring expression within excitatory neurons25. Acute brain slices were prepared 3 weeks after virus injection. Our immunohistochemical analysis revealed robust and unilateral expression of EYFP in the subgranular zone of the DG and the stratum radiatum (Fig. 7h). We used whole-cell recording to examine postsynaptic currents in DGCs while optically stimulating ChR2–EYFP-positive projections in the DG. Blue light pulses reliably induced postsynaptic currents in DGCs. These optically evoked postsynaptic currents were not significantly affected by gabazine (10 µM) but were blocked by coapplication of 6-cyano-7-nitroquinolinoxide-2, 3-dione (CNQX; 20 µM) and D-2-amino-5-phosphonopentanoic acid (APV; 50 µM), indicating that they were excitatory postsynaptic responses (Fig. 7i). We confirmed that DGCs indeed receive a direct back projection from CA3 pyramidal neurons.

Given that chemogenetics permits bidirectional manipulation of neuronal activity with anatomical, genetic, and temporal precision, we also used in vivo chemogenetic approaches to study the role of CA3 pyramidal neuronal activity in controlling adult hippocampal neurogenesis. The experimental procedure is depicted in Fig. 8a. We bilaterally injected the CA3 region with an AAV-DJ viral vector expressing an engineered Gq-coupled human M3 (hM3Dq) tagged with mChirite (hM3Dq-mChirite) or G418-coupled human M4 (hM4Di) tagged with mCherry (hM4Di-mCherry), under control of the CaMKIIα promoter. One week later, WT and Oxtr−/− mice were subjected to multiple BrdU injections and analyzed BrdU incorporation 2 weeks later.

Fig. 4 Conditional deletion of Oxtr delays the morphological maturation of newly generated DGCs. a Confocal three-dimensional reconstruction of dendrites of EGFP + DGCs from WT and Oxtr−/− mice at 14 dpi. Scale bar, 50 µm. b, c Summary bar graphs depicting b the total dendritic length and c branch number of EGFP + DGCs from WT and Oxtr−/− mice at 14 dpi (n = 11 neurons from 4 mice per genotype; **P < 0.01, ***P < 0.001, unpaired two-tailed Student’s t test). d Sholl analysis of dendritic complexity of EGFP + DGCs from WT and Oxtr−/− mice at 14 dpi (n = 11 neurons from 4 mice per genotype; *P < 0.05, **P < 0.01, two-way ANOVA with Bonferroni’s post hoc test). e Representative confocal images showing EGFP + DGCs from WT and Oxtr−/− mice at 28 dpi. Scale bar, 50 µm. f, g Summary bar graphs depicting f the total dendritic length and g branch number of EGFP + DGCs from WT and Oxtr−/− mice at 28 dpi (n = 11 neurons from 5 mice per genotype; unpaired two-tailed Student’s t test). h Sholl analysis of dendritic complexity of EGFP + DGCs from WT and Oxtr−/− mice at 28 dpi (n = 11 neurons from 5 mice per genotype; two-way ANOVA). i Top panel, schematic representations of the retroviral vectors used for in vivo genetic manipulation under the control of the Ubi-1 promoter. Bottom panels, representative confocal images showing Retro-EGFP + and Retro-Cre-2A-EGFP + DGCs in Oxtr−/− mice at 14 dpi. Scale bar, 50 µm. j Summary bar graphs depicting the total dendritic length of Retro-EGFP + and Retro-Cre-2A-EGFP + DGCs in Oxtr−/− mice at 14 dpi (n = 5 neurons from 3 mice for each group, unpaired two-tailed Student’s t test). k Sholl analysis of dendritic complexity of Retro-EGFP + and Retro-Cre-2A-EGFP + DGCs in Oxtr−/− mice at 14 dpi (n = 5 neurons from 3 mice for each group, two-way ANOVA). Data represent the mean ± s.e.m.
Mice were injected intraperitoneally daily with clozapine-N-oxide (CNO, 10 mg/kg), a synthetic ligand of hM3Dq and hM4Di, for continuous 2 weeks until killing. Post hoc histological examination of brain sections revealed robust and bilateral expression of hM3Dq or hM4Di in CA3 pyramidal neurons (Fig. 8b). In a subset of mice, we performed ex vivo electrophysiological

Fig. 5 Effects of Oxtr deletion on excitatory and inhibitory synaptic transmission of newly generated DGCs. a Representative traces of sEPSCs recorded from newly generated DGCs in slices from WT and Oxtr−/− mice at 14 dpi. b, c Cumulative probability plots of sEPSC b interevent intervals and c amplitude (n = 6 neurons from 4 mice per genotype, Kolmogorov-Smirnov test) in newly generated DGCs from WT and Oxtr−/− mice at 14 dpi. d, e Summary bar graphs depicting the averaged d frequency and e amplitude of sEPSCs in newly generated DGCs from WT and Oxtr−/− mice at 14 dpi (n = 6 neurons from 4 mice per genotype; *P < 0.05, unpaired two-tailed Student’s t test). f Left panel, Sample traces of evoked IPSCs recorded at different holding potentials ranging from −70 to −20 mV (10 mV step) from newly generated DGCs in slices from WT and Oxtr−/− mice at 10 dpi. Right panel, current-voltage plot of the peak current of evoked IPSCs obtained from the sample traces. g Summary data comparing the reversal potential of GABA-mediated IPSCs (E_GABA) of newly generated DGCs from WT and Oxtr−/− mice at 10, 14, and 28 dpi (n = 4–5 neurons from 4 mice for each group; *P < 0.05, **P < 0.01, two-way ANOVA with Bonferroni’s post hoc test). h Summary data comparing the calculated intracellular chloride concentrations ([Cl]i) in newly generated DGCs from WT and Oxtr−/− mice at 10, 14, and 28 dpi (n = 4–5 neurons from 4 mice for each group; *P < 0.05, ***P < 0.001, two-way ANOVA with Bonferroni’s post hoc test). Data represent the mean ± s.e.m.
recordings to confirm the effects of CNO in AAV-infected neurons. Application of CNO (1 μM) significantly decreased spiking responses to +150 pA square current pulses in hM4Di-expressing (hM4Di+) neurons but increased spiking responses in hM3Dq-expressing (hM3Dq+) neurons (Fig. 8c). In Oxtr−/− mice, we found that the total number of BrdU+ cells was significantly increased by hM3Dq/CNO-based approach (Fig. 8d, e). Conversely, hM4Di/CNO-based approach led to a decrease in the total number of BrdU+ cells in WT mice (Fig. 8d, e). To evaluate whether chronic CNO administration may cause cell death of CA3 pyramidal neurons, we used terminal deoxynucleotidyl transferase-mediated dUTP nick-end labeling (TUNEL) assay to assess the extent of apoptotic death. No TUNEL-positive cells were found in the CA3 of WT mice with or without hM3Dq/CNO treatment compared with DNase-treated positive control (Supplementary Fig. 7), confirming that our chronic CNO treatment protocol does not induce apoptotic cell death.

Since Oxtr deletion in Oxtr−/− mice was not restricted to the CA3, the loss of OXTR function in other hippocampal or forebrain regions may also contribute to impair the survival and maturation of newly generated DGCs. To circumvent this limitation, in a subset of experiments, we used Oxtrf/f mice in combination with Cre recombinase-mediated gene deletion in a localized fashion through bilateral stereotaxic injections of AAV-
Cre-GFP or control AAV-GFP vectors targeting the CA3, under control of the CaMKIIα promoter, favoring expression to excitatory neurons. One day before stereotaxic injection of AAV-GFP or AAV-Cre-GFP, mice were subjected to multiple BrdU injections, and the number of BrdU+ cells was measured 2 weeks after the last BrdU injection (Supplementary Fig. 8a). The successful transduction of AAV-GFP or AAV-Cre-GFP was confirmed by immunohistochemistry (Supplementary Fig. 8b). Quantitative real-time PCR analysis confirmed the loss of Oxtr mRNA expression in the CA3 2 weeks following the stereotactic injection of AAV-Cre-GFP (Supplementary Fig. 8c). Consistent with the results from Oxtr−/− mice, we found a significant reduction of the total of BrdU+ cells in Oxtrf/f mice with AAV-Cre-GFP injections compared with those injected with of...
Discussion

In the present study, we examined the role of endogenous OXT signaling in adult hippocampal neurogenesis. There are three major findings. First, OXTR is not expressed in neural progenitor cells reside within the subgranular zone or mature granule cells of adult DG, whereas it is enriched in the CA2 and CA3 of the hippocampus. Second, conditional deletion of Oxtr from hippocampal excitatory neurons leads to impaired survival and maturation of newly generated DGCs. Third, OXT enhances the excitability of hippocampal CA3 pyramidal neurons and CA3 neuronal activity regulates adult hippocampal neurogenesis under basal conditions. Together, these results highlight a non-cell autonomous role for OXT in promoting adult hippocampal neurogenesis via OXTR expressed in CA3 pyramidal neurons.

OXTR is widely expressed in the brain. Early studies using radiolabeled ligands or transgenic approaches have demonstrated abundant expression of OXTR in the hippocampus, however, no studies have yet reported whether OXTR is expressed in developing DGCs in adult mouse brain. Using Oxtr<enu5-Neow/+ mice in conjunction with immunofluorescent staining for proteins expressed at distinct stages of cell differentiation, we demonstrate here for the first time that OXTR is not expressed in neural progenitor cells or mature granule cells of adult mouse DG. Instead, we confirmed in the previous findings that OXTR is highly expressed in hippocampal CA2 and CA3 pyramidal neurons. We also found that OXTR is present in a subset of calretinin-expressing cells in the hilus of the DG. Given that hilar calretinin-expressing cells in mouse DG could belong to either GABAergic interneurons or mossy cells, additional studies are needed to clarify the neuronal identity of hilar OXTR-expressing cells.

A recent study using pharmacological approaches has proposed that OXT could protect the hippocampus from the detrimental effects of elevated glucocorticoids by promoting adult neurogenesis. It has been demonstrated that both acute peripheral and intra-hippocampal OXT administration enhances cell proliferation, and repeated OXT administration increases the number of newly generated DGGs in adult rat hippocampus. We have extended these findings by underscoring the importance of OXTR signaling in the regulation of adult hippocampal neurogenesis under basal conditions. This view is supported by the observation that conditional deletion of Oxtr from hippocampal excitatory neurons resulted in impaired survival and maturation of newly generated DGCs. Our data suggest, however, that the progenitor cell proliferation and early differentiation into immature neurons were unaltered by conditional deletion of Oxtr. Because we observed no significant changes in cell proliferation in the DG following conditional deletion of Oxtr by an in vivo BrdU incorporation assay, it might be possible that the level of endogenous OXT in the hippocampus is not enough to exert a tonic stimulation of progenitor cell proliferation. Indeed, peripheral OXT administration was previously found to enhance cell proliferation in the adult rat DG in a dose-dependent manner.

Newly generated DGCs exhibit features of dendrites as early as 2 weeks after birth and display overall morphological and functional properties of fully mature GCs at 6 weeks of cell age. In addition, the rate of newly generated DGC maturation highly correlates with the pattern of neuronal activity. We demonstrated that conditional deletion of Oxtr decreased dendritic complexity of newly generated DGCs, as shown by decreased dendritic length and branching, at 14 dpi. However, the suppressive effect of Oxtr deletion on dendrite maturation of newly generated DGCs is short lived and dendritic complexity returns to normal levels at 28 dpi. In accordance with morphological observations, our electrophysiological data indicate that the maturation of excitatory synaptic transmission and the developmentally regulated shift in the actions of GABA from excitation to inhibition on newly generated DGCs were delayed by conditional deletion of Oxtr. These data suggest that, although endogenous OXT signaling has effects on morphological properties of newly generated DGCs during their early development, some compensatory mechanisms may exist during late developmental stages to normalize the morphological and physiological changes caused by Oxtr deletion.

A pressing question that follows these observations is how conditional deletion of Oxtr from hippocampal excitatory neurons impairs neurogenesis in the DG. Our results indicate that CA3 pyramidal neurons exert the stimulatory effect of OXT on neurogenesis in the DG of adult mice. By using optogenetic stimulation and immunofluorescence imaging, we found that CA3 pyramidal neurons functionally project to DGCs. In support of our observations, a recent viral-genetic tracing study has provided evidence that newly generated DGCs receives excitatory afferents from CA3 pyramidal neurons. In addition, lesions to the CA3 region of the hippocampus have been found to decrease the survival of newly generated DGCs. Thus, excitatory inputs from
CA3 to newly generated DGCs are crucial in maintaining the survival of newly generated DGCs. How do CA3 pyramidal neurons influence the survival of newly generated DGCs? Considering that network activity plays a critical role in the survival of newly generated neurons in the adult brain\textsuperscript{33–35}, it is therefore possible that OXT may increase the excitability of CA3 pyramidal neurons, thereby promoting network activity and newly generated DGC survival. In support of this view, we observed that bath application OXT induced a membrane depolarization and increased the number of action potentials in response to postsynaptic depolarizing current injection, suggesting a role for OXT in regulating the excitability of CA3 pyramidal neurons. Our findings that chemogenetic activation of CA3 pyramidal neurons enhanced the survival of newly generated DGCs in Oxtr\textsuperscript{−/−} mice, whereas chemogenetic inhibition of CA3 pyramidal neurons decreased the number of newly generated DGCs in WT mice, clearly confirm that CA3 pyramidal neuron activity can influence the survival of newly generated DGCs.

OXT may exert its biological effects through the activation of OXTR-expressing interneurons. It was previously demonstrated...
that OXT activates a subpopulation of GABAergic interneurons within the lateral subdivision of the central nucleus of the amygdala (CeA) that inhibits activity in medial CeA neurons, thereby attenuating conditioned fear responses. Furthermore, OXT has been shown to enhance hippocampal spike transmission by modulating fast-spiking interneurons. Very recent work also reports that OXT can activate a small subset of fast-spiking hilar GABAergic interneurons, which provide local inhibition to mossy cells in the DG. This in context, our immunofluorescent staining study demonstrates that a small fraction of hilar OXTR-expressing cells are also immunoreactive for GAD67, calretinin, or parvalbumin. It is worth noting that, although parvalbumin interneuron activation has been shown to effectively promote newborn neuronal progeny survival and development via GABA signaling, this mechanism cannot account for the observed impaired survival of newly generated DGCs in our Oxt conditional knockout mice. Indeed, we conditionally deleted Oxt under the control of the CaMKIIα promoter, which was not expressed in parvalbumin-expressing interneuron. A recent study has demonstrated that CaMKIIα was expressed in ~40% calretinin-immunoreactive neurons in the DG of 4- to 5-week-old mice; however, we rarely detected double-labeled neurons immunoreactive for CaMKIIα and calretinin in 10-week-old mice (Lin et al., unpublished observation). This may exclude a role of OXTR-expressing calretinin interneurons in mediating the effects of conditional deletion of Oxt on adult DG neurogenesis. Since Oxt deletion in conditional Oxt knockout mice was not restricted to the hippocampus, we could not exclude the possibility that the observed impairment of adult neurogenesis in Oxt–/– mice may be partially mediated through OXTR-expressing excitatory neurons within extra-hippocampal regions that also provide afferent inputs to the DG, such as the amygdala. Although we have conducted site-specific manipulation experiments to confirm the role of hippocampal CA3 OXT signaling in regulating DG neurogenesis, our study cannot definitively rule out the possibility that CA3 pyramidal neurons may, in part, control DG neurogenesis indirectly through the regulation of subsets of hilar GABAergic interneuron activity. Further research is warranted to test this possibility.

In conclusion, we provide compelling evidence that conditional deletion of Oxt from hippocampal excitatory neurons results in reduced survival and impaired maturation of newly generated DGCs, suggesting a potential role for endogenous OXT signaling in regulating adult hippocampal neurogenesis. Our results also reveal that CA3 pyramidal neurons play an important role in the regulation of adult hippocampal neurogenesis and OXT controls adult hippocampal neurogenesis through an indirect non-cell autonomous mechanism by OXTR expressed in CA3 pyramidal neurons. Given the prominent role of endogenous OXT signaling in enhancing adult hippocampal neurogenesis, we therefore speculate that it may serve as a potential therapeutic target for neurodegenerative disorders.

**Methods**

**Animals.** Adult male C57BL/6 (8-12 weeks old), homozygous Oxt–/– (Oxtfl/fl), and CaMKIIα-Cre transgenic mice were originally obtained from The Jackson Laboratory and bred within our animal facility. Oxt–/– mice were crossed to CaMKIIα-Cre mice to generate Oxt conditional knockout (Oxt–/–) mice in the C57BL/6 genetic background. The heterozygous Oxtfl/+ mice were generated as described previously. Mice were genotyped by a PCR-based method using genomic DNA isolated from tail samples. The primers used were as follows: Oxtforward (5′-GGCTCAGGTTGCAGCT-3′) and reverse (5′-GGCAGGAAACAGGGTTGATTA-3′); Cre, forward (5′-GGGCTCTGGGAAATGATCACTTAC-3′) and reverse (5′-GTGGGGACACAGGGTTGATTA-3′); and Ver, forward (5′-GGGCTCTGGGAAATGATCACTTAC-3′) and reverse (5′-GTGGGGACACAGGGTTGATTA-3′). Mice were housed in groups of three in a humidity- and temperature-controlled (25 ± 1 °C) vivarium on a 12 h-light/dark cycle with access to food and water ad libitum. All experimental procedures were conducted in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and were approved by the Institutional Animal Care and Use Committee of National Cheng Kung University. All efforts were made to minimize the number of animals used and their suffering.

**Quantitative real-time PCR (qPCR).** Total RNA was isolated from hippocampal CA2, CA3, and hypothalamus tissue lysates using a Tri Reagent kit (Molecular Research Center) and treated with RNase-free DNase (RQ1; Promega) to remove potential contamination by genomic DNA. Total RNA (2 μg) from samples was reverse transcribed using a SuperScript CDNA synthesis kit (Invitrogen). qPCR was performed using the Roche LightCycler instrument (Roche Diagnostics) using the FastStart DNA Master SYBR Green I kit (Roche Applied Science) according to the manufacturer’s instructions. The primers used in this experiment for Oxt were as follows: forward (5′-TTTCTTCTGACGATTTGAGG-3′) and reverse (5′-CCCTTGAGGACCAAGCAG-3′). The PCR reactions were run for 40 cycles. Each experiment was run in triplicate with expression results normalized to β-actin rRNA.

**Immunofluorescence.** Oxtfl/cre-Venus-Neo1 mice were deeply anesthetized with sodium pentobarbital (50 mg/kg, intraperitoneally) and perfused transcardially with PBS and 4% paraformaldehyde. After the perfusion, brains were removed and immersed in 4% paraformaldehyde for 4 h. Frozen sections (8 μm) were incubated in solution containing 30% sucrose that immersed in 4°C for at least 48 h before slicing. Coronal sections were sectioned to a 40 μm thickness, washed with 0.3% Triton X-100, and then incubated for blocking with solution containing 3% goat serum in PBS. After blocking, the sections were incubated in the primary antibodies against CaMKIIα (1:500, Millipore, AB1778), β-actin (1:500; Millipore, AB5808), calretinin (1:500; Millipore, MAB1568), parvalbumin (1:150; Swant, PV235), calbindin (1:500; Millipore, AB1778), CaMKIIα (1:500; Novus Biologicals, NB100-81830), and GAD67 (1:500, Millipore, MAB5406). Finally, sections were washed with PBS-T (10 mM Tris-HCL, 150 mM NaCl, and 0.025% Tween 20; pH 7.4) and then incubated with the secondary Alexa Fluor 568 antibody (Life Technologies) for 2 h at room temperature in blocking buffer. The immunostained sections were collected on separate gelatin-subbed glass slides, rinsed extensively in PBS, and mounted with ProLong Gold Antifade Reagent (Invitrogen). Fluorescence images of neurons were captured using an Olympus Fluorescence microscope at ×100 magnification with sequential acquisition setting at a resolution of 1024 × 1024 pixels, z-stack with 15–20 optical sections. All images were imported into NIH Image J software (National Institutes of Health) for analysis, and all the parameters used were kept consistent during capturing.

**BrdU injection and quantification of newly generated DGCs.** For proliferation assay, mice were received single pulse of BrdU (50 mg/kg; Sigma–Aldrich) injection. One hour after BrdU was injected, mice were perfused with 4% paraformaldehyde, and brain tissue was collected. For quantification of the survival neurons, mice were injected six times intraperitoneally with BrdU at 12 h intervals and killed by transcardial perfusion after 2 and 4 weeks after the last BrdU injection. For the study of the CA3 neuronal activity on newborn neurons survival, AAV-hmD2q or AAV- hM4Di was bilaterally injected into the CA3 region. One week later, mice were injected six times intraperitoneally with BrdU at 12 h intervals and daily intraperitoneally injected with CNO (10 mg/kg; Sigma–Aldrich) for continuous 2 weeks until killing. The entire dentate gyrus (DGC), ~0.9 to ~4.2 mm from bregma) was sectioned coronally at a thickness of 40 μm using a sliding microtome (Leica SM2000R).

Fluorescent immunolabelling was used for counting the number of newly generated DGCs. For BrdU/Ki67, BrdU/DCX, or BrdU/NeuN double labeling, float-free sections were denatured in 10 mM saline-sodium citrate buffer at 85 °C for 20 min and then incubated at 72 °C for 30 min in 2 H CCL. Sections were rinsed twice for 5 min at 25 °C in PBS-T. Sections were incubated in the primary antibodies against BrdU (1:500; Millipore, AB4072), Ki67 (1:500; Abcam, ab158360), DCX (1:500; Cell Signaling Technology, #4604), NeuN (1:1000; Millipore, ABN78), calretinin (1:500; Millipore, MAB1568), parvalbumin (1:150; Swant, PV235), calbindin (1:500; Millipore, AB1778), CaMKIIα (1:500; Novus Biologicals, NB100-81830), and GAD67 (1:500, Millipore, MAB5406). Finally, sections were washed with PBS-T (10 mM Tris-HCL, 150 mM NaCl, and 0.025% Tween 20; pH 7.4) and then incubated with the secondary Alexa Fluor 568 antibody (Life Technologies) for 2 h at room temperature. The nuclei were visualized using 4′,6-diamidino-2-phenylindole (DAPI, 1:5000; Sigma–Aldrich) staining. The immunostained sections were collected on separate gelatin-subbed glass slides, rinsed extensively in PBS, and mounted with ProLong Gold Antifade Reagent (Invitrogen). Quantification of BrdU-labeled cells was performed using a modified unbiased stereology protocol as described previously. Every sixth section covering the entire DG was processed for BrdU immunohistochemistry. All BrdU-labeled cells in the granule cell layer, subgranular zone, and hilus were counted under fluorescent illumination at ×600 using an Olympus BX51 microscope coupled to an Olympus DP70 digital camera. Fluorescent stereology analysis was applied to count BrdU-labeled cells. Total cell numbers were estimated by multiplying the number of cells counted in every sixth section by six.
Fluorescent in situ hybridization. Fluorescent in situ hybridization was performed using RNAscope® Multiplex Fluorescent Reagent Kit 2.0 according to the manufacturer’s instructions (Advanced Cell Diagnostics). Briefly, brain sections (16 μm) were fixed in 4% paraformaldehyde for 15 min and dehydrated through graded ethanol solutions (50, 70, and 100%) for 5 min each. Sections were subjected to reagent pretreat 3 at 25 °C for 30 min and then hybridized with probes at 40 °C for 2 h in a humidified oven. The Ovx-O1 probe (Cat# 450011) and the CaMKII probe (Cat# 450012) were designed to target 200 bp of the artificial cerebellar gene and Oxtr mRNA. After hybridization, brain sections were sequentially applied with a series of probe signal amplification steps, rinsed with ACD wash buffer twice for 2 min between each step and finally counterstained with DAPI and mounted with Vectashield antifade mounting medium (Vector Laboratories) containing DAPI (1:5000; Sigma-Aldrich).

Retrovirus production, stereotaxic injection, and analysis. Engineered self-inactivating murine retroviruses expressing EGFP were used to label proliferating cells and their progeny in the DG of adult mice as described previously. High titers of engineered retroviruses (1 × 10^8 unit/ml) were produced by co-transfection of retroviral vectors and VSV-G into HEK293GP cells followed by ultracentrifugation of viral supernatant. The Cre recombinase (Addgene, plasmid #20781) was cloned into a retroviral expression vector (Ubi-X-2A-EGFP).

Adult male WT or homozgyous Ovx+/− mice were anesthetized and the purified retroviruses were stereotactically injected into the DG at 4 sites (0.5 μl/site at 0.25 μl/min) with the following coordinates (anterior-posterior = −2 mm from bregma, lateral = ±1.6 mm, ventral = 2.5 mm; anterior-posterior 3 mm from bregma, lateral = ±2.6 mm, ventral = 3.2 mm) in accordance with the description by Franklin et al. (2013) and the progress of perforation was continuously monitored until the series resistance had stabilized to ≤30 MΩ. The intracellular chloride concentration was calculated according to the Nernst equation: [Cl]i = [Cl]o × (EClABAF/RT), where the extracellular Cl− concentration [Cl]o, is 124 mM.

Electrophysiological recordings. Hippocampal slices were prepared using standard procedures as described previously. Briefly, were anesthetized with isoflurane (1.5%–2.5%) and were rapidly decapitated, and the brain was rapidly bathed in ice-cold oxygenated sucrose cutting solution (containing (in mM): sucrose 234, KCl 2.5, CaCl2 0.5, MgCl2 7, NaHCO3 25, NaH2PO4 1.2, and glucose 11 at pH 7.3–7.4 and equilibrated with 95% O2–5% CO2). Hippocampal slices (250 μm) were prepared using a vibrating microtome (VT1200; Leica) and transferred to a holding chamber (1 ml) containing artificial cerebellar medium (ACSF) (containing (in mM): NaCl 117, KCl 4.7, CaCl2 0.5, MgCl2 1.2, NaHCO3 25, NaH2PO4 1.2, and glucose 11 at pH 7.3–7.4 and equilibrated with 95% O2–5% CO2) and maintained at room temperature for at least 1 h before use.

For recording, sections were transferred to a submergence-type recording chamber and fixed at the glass bottom of the chamber with a nylon grid on a platinum frame. The chamber was constantly perfused with ACSF at 32.0 ± 0.5 °C with a rate of 2–3 ml/min. Conventional whole-cell and g-aminobutyric acid-perfused patch-clamp recordings were made from newly generated (EGFP)+ DGCs, mature (EGFP)+ DGCs or CA3 pyramidal neurons by using a patch-clamp amplifier (Axopatch 200B, Molecular Devices) under infrared differential interference contrast microscope. Mature DGCs were recorded from the outer portion of the granule cell layer. Data acquisition and analysis were performed using a digitizer (Digidata 1440 A, Molecular Devices) and pCLAMP 9 software (Molecular Devices). Synthetic responses were evoked using a bipolar stainless steel stimulating electrode placed in the middle molecular layer ~100 μm away from the recorded cell. For measurement of E_CABA, in newly generated DGCs, pharmacologically isolated GABA_A receptor-mediated IPSCs were recorded using g-aminobutyric acid-perfused patch under voltage-clamp at different holding potentials in the presence of CNQX (20 μM: Tocris Bioscience) and APV (50 μM: Tocris Bioscience) to eliminate glutamate and g-aminobutyric acid stock solution (50 μg/ml) was prepared in dimethyl sulfoxide and was then diluted in the pipette solution (containing (in mM): CsCl 135, MgCl2 2, EGTA 0.5, HEPES 10, pH 7.3, pH 7.3, 290–295 mOsM) to a final concentration of 25 μg/ml. Following gigaseal formation, the progress of perforation was continuously monitored until the series resistance had stabilized to ≤30 MΩ. The intracellular chloride concentration was calculated according to the Nernst equation: [Cl]i = [Cl]o × (EClABAF/RT), where the extracellular Cl− concentration [Cl]o, is 124 mM.

IPSeps and mIPSeps were recorded from newly generated or mature DGCs held under voltage-clamp at −70 mV in the presence of gabazine (10 μM: Tocris Bioscience) and analyzed offline using commercially available software (Mini Analysis 4.3; Synaptosoft). Recordings of mIPSeps were conducted in the presence of TTX (0.5 μM: Tocris Bioscience). The composition of intracellular solution was (in mM): K-glucuronate 120, KCl 15, HEPES 10, MgCl2 4, EGTA 0.1, Na2ATP 4, Na,GTP 0.3, phosphocreatine 7, and 0.5% w/v bicucullin, 280–290 mMos without bicucullin, pH 7.3 with KOH. The detection threshold for eIPSeps and mIPSeps was set at 3 pA.

To examine the effects of OXT on intrinsic membrane properties of OXTR-expressing CA3 pyramidal neurons, hippocampal slices were prepared from Ovx-Venus knock-in mice. After a stable record of membrane potential under current-clamp condition, OXT (1 μM) was recorded to the recorded cell by a 5-min bath application. In some experiments, hyperpolarizing (−50 pA) and depolarizing current pulses (100 pA) of 500 ms were injected through the recording pipette in current-clamp mode to measure neuronal excitability. The number of action potentials triggered was counted. Bicucullin was routinely included in the intracellular solution to allow post hoc staining of the recorded neurons. OXT and L-371257 were purchased from Tocris Bioscience.

To confirm the expression of engineered hM3Dq or hM4Di receptors in CA3 neurons, a depolarizing current pulse (150 pA, 500 ms) was injected into the control, hM3Dq or hM4Di (hM3Dq or hM4Di expression) neurons to induce spiking. Following 10 min continuous recordings, CNO (1 μM; Sigma-Aldrich) was applied into the ACSF and a second depolarizing current pulse (+150 pA, 500 ms) or hyperpolarizing current pulse (−50 pA, 500 ms) was injected into the hM3Dq−, hM3Dq+, hM4Di−, and hM4Di+ neurons to induce spiking in order to compare spiking before and after CNO application.

For in vitro optical stimulation experiments, a multi-mode optical fiber with core diameter of 200 μm (Thorlabs), coupled to a diode-pumped solid-state laser of specific wavelength (473 nm blue laser; LaserRig Technologies), was used. The ending power on brain slices was ~5 mW/mm² and synaptic responses were recorded at 70 mV in voltage-clamp mode.

Statistical analysis. No statistical tests were used to predetermine sample sizes, but our sample sizes were consistent with the previous publication using similar approaches. Mice were randomly assigned to viral injection experiments, and investigators were blinded to the group allocation while performing cell number counting, morphological analysis, and electrophysiological recordings. The results presented are mean ± s.e.m. Statistical analysis was performed using the Prism 6 software package (GraphPad software). The significance of any difference between two groups was calculated using the paired or unpaired two-tailed Student’s t test. One-way ANOVA or repeated measure two-way ANOVA tests were used for multiple groups’ comparison and Bonferroni’s post hoc analyses were used to assess the significance between isolated groups. Kolmogorov-Smirnov test were found to be not significant. The actual statistical test used is described in the figure legends.
used in comparison of cumulative frequency distribution. N represents the number of animals used. Values of P < 0.05 were considered significant.

Data availability. The authors declare that all data supporting the findings of this study are available within the article and its Supplementary Information File.

Received: 16 October 2016 Accepted: 19 July 2017
Published online: 14 September 2017

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