Orphan GPR110 (ADGRF1) targeted by N-docosahexaenoylethanolamine in development of neurons and cognitive function

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Docosahexaenoic acid (DHA, 22:6n-3) is an omega-3 fatty acid essential for proper brain development. N-docosahexaenoylethanolamine (synaptamide), an endogenous metabolite of DHA, potently promotes neurogenesis, neuritogenesis and synaptogenesis; however, the underlying molecular mechanism is not known. Here, we demonstrate orphan G-protein coupled receptor 110 (GPR110, ADGRF1) as the synaptamide receptor, mediating synaptamide-induced bioactivity in a cAMP-dependent manner. Mass spectrometry-based proteomic characterization and cellular fluorescence tracing with chemical analogues of synaptamide reveal specific binding of GPR110 to synaptamide, which triggers cAMP production with low nM potency. Disruption of this binding or GPR110 gene knockout abolishes while GPR110 overexpression enhances synaptamide-induced bioactivity. GPR110 is highly expressed in fetal brains but rapidly decreases after birth. GPR110 knockout mice show significant deficits in object recognition and spatial memory. GPR110 deorphanized as a functional synaptamide receptor provides a novel target for neurodevelopmental control and new insight into mechanisms by which DHA promotes brain development and function.
Omega-3 fatty acids are essential nutrients. Docosahexaenoic acid (DHA, 22:6n-3), an omega-3 fatty acid highly enriched in the brain, plays a significant role in brain development, as well as cognitive and memory function, but specific molecular mechanisms responsible for this beneficial effect are not clearly understood. DHA promotes neurite outgrowth, synaptogenesis and synaptic function in cultured hippocampal neurons and induces neuronal differentiation of neural stem cells (NSCs) in vivo and in vitro. Several pre- and postsynaptic proteins involved in neurotransmission and synapse formation are reduced in the DHA-depleted brains, indicating an important role of DHA in maintaining synapse integrity in vivo.

N-docosahexaenoyl ethanolamine, an endogenous metabolite of DHA, significantly increases neuritogenesis, synaptogenesis and glutamatergic synaptic activity and potently induces neurogenic differentiation. The similarity of its structure to a well-known endocannabinoid N-arachidonylethanolamine (anandamide), suggests that synaptamide may use the biosynthetic mechanism involving hydrolysis of N-acylphosphatidylethanolamine (NAPE) that has been established for anandamide. The significant decreases in synaptamide recently reported in the brain regions of NAPE-phospholipase D (NAPE-PLD) knockout (KO) mice support such a mechanism via N-docosahexaenoylphosphatidylethanolamine (NDPE). On the basis of the synaptic property and its endocannabinoid-like amide structure, the term ‘synaptamide’ was coined for N-docosahexaenoyl ethanolamine. We used ‘synaptamide’ to represent N-docosahexaenoyl ethanolamine since the abbreviation ‘DHEA’ is a widely used and accepted term for the steroid, dehydroepiandrosterone. The neurogenic, neuritogenic and synaptic bioactivities of DHA and synaptamide are qualitatively similar, but the latter is significantly more potent. The endogenous level of synaptamide in brain is affected by the dietary omega-3 fatty acid intake, which alters brain DHA content. Furthermore, DHA-derived neurogenic, neuritogenic and synaptic bioactivity is considerably enhanced when fatty acid amide hydrolase is inhibited. These findings suggest that synaptamide is a principal mediator for the observed bioactivity of DHA, most likely involving a specific target receptor.

G-protein coupled receptors (GPCR) are the major regulators of intracellular signalling and are important drug targets, but many including most of the adhesion GPCRs (aGPCRs) still remain orphan without identified natural ligands. GPR40 and GPR120 of class A GPCRs were deorphanized as fatty acid receptors that preferentially bind to long chain free fatty acids including DHA. In this paper, using a chemical and proteomic strategy altogether with molecular and cellular characterization, we identified orphan GPR110 (ADGRF1), a member of the less-known aGPCRs, as the functional synaptamide receptor mediating synaptamide-induced neurite growth and synaptogenesis in cortical neurons and neurogenic differentiation of NSCs. We suggest that synaptamide-activated GPR110 signalling is an important molecular mechanism for the neurodevelopmental actions of omega-3 fatty acids.

**Results**

**Synaptamide bioactivity is cAMP-dependent.** Previously, we observed that synaptamide promotes neurite growth and synaptogenesis in hippocampal neurons and induces neurogenic differentiation and cyclic adenosine monophosphate (cAMP)/protein kinase A signalling at a low nM range in NSCs. Therefore, we first examined whether the bioactivity of synaptamide depends on cAMP production using two model systems, cortical neurons and NSCs in culture (Fig. 1).

**GPR110 is the synaptamide-binding protein.** To identify the GPCR that binds to synaptamide, we devised a strategy using a pull down approach coupled to mass spectrometry (MS). A biotinylated synaptamide analogue was prepared for affinity-purification (AP); G1, (4Z,7Z,10Z,13Z,16Z,19Z)-N-(2-(5-(2-(5-thieno(3,4-d)imidazol-4-yl))-2-oxohexahydro-1H-thieno(3,4-d)imidazol-4-yl)pentanamide)ethyl)docosa-4,7,10,13,16,19-hexaenamide (Fig. 2a). This analogue exhibited bioactivity similar to synaptamide, inducing cAMP production (Fig. 2b) and neurite outgrowth (Supplementary Fig. 5). Using G1, we searched for GPCRs according to the scheme illustrated in Fig. 2c. Mouse fetal brains or NSCs were lysed under a mild condition, treated with G1, and incubated with streptavidin beads for affinity-purification. The proteins thus detected were considered as background. This approach allowed us to identify two peptides, G(432–442)R (GTPVTVQIQSTR) and G(460–471)K (GHVFIEPDQFK) (Fig. 2d), that uniquely belong to GPR110 in G1-treated fetal brain and NSC samples (Supplementary Table 1). No other GPCRs were detected using this approach. GPR110 is an orphan receptor that belongs to the adhesion GPCR family and has seven transmembrane (7TM) helices with multiple N-linked glycosylation sites. The molecular mass of GPR110 is about 100 kD, but lower molecular weight isoforms as well as differentially glycosylated forms have been reported.

To confirm that GPR110 is indeed the G1-binding protein, C-terminal HA-tagged GPR110 (GPR110-HA) was expressed in HEK cells and subjected to the AP/MS procedure described above. The mouse GPR110-HA (mGPR110-HA) expressed in HEK cells was detected mainly at the apparent molecular weight of 100 kD along with a minor band at ~130 kD using both HA and N-terminal targeting mGPR110 antibodies (Supplementary Fig. 6a). After immunoprecipitation (IP) using HA antibody, the IP product was applied to SDS PAGE, and the gel bands at
~100 and ~130 kD were cut out and analysed by mass spectrometry. Both N- and C-terminal, as well as transmembrane region tryptic peptides were detected (Supplementary Fig. 6b, Supplementary Tables 2 and 3), confirming that these protein bands were derived from the full length mGPR110-HA, and therefore, possibly represent different states of posttranslational modification. When the lysate from mGPR110-HA-expressing cells was treated with G1 followed by the streptavidin affinity purification, HA antibody detected the G1-binding protein at ~100 kD (Fig. 2e,f), confirming binding of G1 to the expressed mGPR110-HA. The competitive binding assayed according to the strategy described above (Fig. 2c) revealed dose-dependent displacement of G1 by synaptamide (Fig. 2e), verifying that both G1 and synaptamide bind to the same receptor, GPR110. Among other fatty acid ethanolamides including anandamide (AEA), n-3 and n-6 docosapentaenoylethanolamide (DPEAn-3, DPEAn-6) or oleoylethanolamide (OEA), only synaptamide significantly displaced G1 binding to GPR110 (Fig. 2f), indicating that synaptamide is a specific ligand to GPR110.

Synaptamide binding activity to human GPR110 (hGPR110) was also examined (Supplementary Fig. 7). The hGPR110-HA expressed in HEK cells showed apparent molecular weight of ~130 and ~110 kD (Supplementary Fig. 7a). After deglycosylation using PNGase F, these bands shifted to ~100 kD. All three bands detected at ~130, ~110 and ~100 kD contained N and C-terminal as well as transmembrane region amino acid sequence.

Figure 1 | cAMP-dependent induction of neurite outgrowth, synaptogenesis and neurogenesis by synaptamide. In cortical neurons, 10 nM synaptamide (Syn) increased cAMP production, neurite outgrowth (a) and synaptogenesis (b) while pretreatment with SQ22,536 for 30 min prevented these effects. Neurite outgrowth and synaptogenesis were evaluated after treating neurons from day 1 in vitro (DIV1) for 2 or 6 days, respectively, while cAMP production and CREB phosphorylation were determined after stimulating DIV3 neurons with synaptamide for 10 min. Neurons and nuclei were visualized using MAP2 antibody (green) and DAPI (blue). Co-localization analysis of presynaptic marker synapsin1 (red) and postsynaptic marker PSD95 (green) revealed increased synapses (overlapping synapsin1/PSD95 puncta indicated by arrows) by synaptamide treated cortical neurons but not in the presence of SQ22,536 (b). For quantification of synaptic protein puncta, 20 cells were scored per well for 3 wells per each group. Synaptamide at 10 nM induced G-protein activation (c). The $\gamma$ GTTP binding was measured after treating NSC membranes with 10 nM synaptamide for 15 min with or without overnight pretreatment with pertussis toxin (PTX, 0.1 μg ml$^{-1}$). While expected increases in cAMP production were observed after PTX pretreatment of cortical cells or NSCs, synaptamide remained capable of increasing cAMP (d). Values are expressed as means ± s.e.m. of biological triplicates ($n = 3$), representing three independent experiments. Statistical analysis was performed using unpaired Student t-test. $^* P < 0.05$, $^*^* P < 0.01$, $^*^*^* P < 0.001$, $^P < 0.05$, $^*^P < 0.01$, $^*^*^P < 0.001$ in comparison with the corresponding control. $^\dagger$, relative to control. Scale bars, 20 μm (a) and 5 μm (b).
Figure 2 | Identification of synaptamide receptor and associated G-protein. Biotinylated synaptamide analogue G1 (a) increased cAMP production in DIV3 neurons after 10 min stimulation (b). After mouse fetal brains or NSCs were lysed in PBS containing 0.5% Triton X-100, treated with G1, affinity-purified using streptavidin beads and subjected to SDS/PAGE, the entire gel was cut into several bands for tryptic digestion and protein identification (ID) by mass spectrometric analysis (c). The MS/MS spectra of GPR110 peptides G(432–442)R and G(460–471)K detected from the gel band in the 100–130 kD molecular weight region are shown (d). Binding of G1 to GPR110 was confirmed by the western blot detection of mGPR110-HA expressed in HEK cells at ~100 kD after streptavidin pull-down while synaptamide dose-dependently decreased the mGPR110-HA recovered after G1-streptavidin pull-down (e). Among fatty acid ethanolamides, only synaptamide displaced the G1 binding to mGPR110-HA (f). GPR110 expression in the plasma membrane was detected (g). The cAMP response induced by 10 nM synaptamide increased in a GPR110 gene-dose-dependent manner (h). GPR110 interaction with Ga was identified by co-immunoprecipitation of overexpressed mGPR110-HA with endogenous Ga (i). Syn, synaptamide; n3, omega-3; n6, omega-6; DPEA, N-docosapentaenoylethanolamine; OEA, N-oleoylethanolamine; AEA, N-arachidonylethanolamine. Results in (d) and (i) represent two independent experiments. Data in (b, e, f, h) are means ± s.e.m. of biological triplicates (n = 3), representing three independent experiments. **P < 0.01, ***P < 0.001 versus control, and #P < 0.05, ##P < 0.01, ###P < 0.001 versus G1 binding by unpaired Student t-test. *, relative to control. Scale bar, 5 μm (g).
we have examined endocytic receptors as fluorescent puncta to microscopically determine specific binding of bodipy-neurons (Fig. 3f). As the fatty acyl ethanolamides are lipophilic, this analogue also retained bioactivity comparable to synaptamide (Fig. 3e). ((4-

permanently expressing CRE-luc2P as a cAMP sensor 25, synaptamide binds to hGPR110 as was observed with mGPR110. mGPR110, hGPR110 showed G1 binding which was displaced by G1* ((4-

GPR110 was overexpressed in HEK293 cells permanently expressing CRE-luc2P as a cAMP sensor 25. GPR110 expression was detected in the plasma membrane and synaptamide-induced cAMP production increased in a gene-dose dependent manner (Fig. 2g,h). Co-immunoprecipitation (IP) assay indicated that GPR110-HA specifically associates with Gzs (Fig. 2i, Supplementary Fig. 8). When HEK cells expressing GPR110-HA were lysed and subjected to co-IP using HA antibody, only Gzs but not Gzi and Gzq was detected from the pull-down precipitate (Fig. 2i). Reversed IP using Gzs antibody detected GPR110-HA, confirming the association of Gzs with GPR110 (Supplementary Fig. 8a). The interaction between GPR110-HA and GPF-Gzs was confirmed in HEK cells expressing both GPR110-HA and GPF-Gzs (Supplementary Fig. 8b). These data are consistent with the pertussis toxin-sensitive increase in $[\gamma,^{35}S]$ GTP binding and cAMP increases observed after synaptamide treatment (Fig. 1c,d).

Synaptamide binds to GPR110 in living cells. To capture the interaction between GPR110 and synaptamide in living cells, in-cell cross-linking 26 was performed using a cross-linkable analogue G1* ((4Z,7Z,10Z,13Z,16Z,19Z)-N-(7-amino-4-5-(3aS,4S,6aR)-2-oxohexahydro-1H-thieno[3,4-dimidazol-4-yl]pentanamido)heptyl) docosa-4,7,10,13,16,19-hexaenamide) that carries a primary amine group (Fig. 3a). This analogue also retained synaptamide-like bioactivity, increasing cAMP (Fig. 3b) and neurite outgrowth (Supplementary Fig. 5). Live HEK cells expressing mGPR110-HA were treated with G1*, cross-linked with a lysine-specific cross-linker disuccinimidylsuberate (DSS), washed extensively and lysed before AP using streptavidin beads as illustrated in Fig. 3c. Subsequent WB analyses using HA-antibody detected GPR110-HA in the G1*-treated, DSS-cross-linked sample, confirming G1* binding to GPR110-HA in live HEK cells (Fig. 3d). The same in-cell cross-linking approach confirmed the binding of endogenous GPR110 to G1* in mouse NSCs (Fig. 3d). In this case, the G1*-bound GPR110 at ~100 kD was detected by peroxidase conjugated streptavidin, as the GPR110 antibody currently available was not sufficiently sensitive to detect a minute level of cross-linked GPR110. Of note, endogenous expression of mGPR110 in NSCs and cortical neurons was clearly indicated by the protein band at ~100 kD, particularly after IP enrichment using this antibody (Supplementary Fig. 9a).

In parallel, we microscopically examined the binding of synaptamide to GPR110 in living cells by using a fluorescent analogue of synaptamide, bodipy-synaptamide 10-(5-(2-(4Z,7Z,10Z,13Z,16Z,19Z)-docosa-4,7,10,13,16,19-hexaenamido)ethylamino)-5-oxopentyl)-5,5-difluoro-1,3,7,9-tetramethyl-5H-dipyrrrolo(1,2-c,1′-d)(1,3,2)diazaborinin-4-ium-5-uide, (Supplementary Fig. 12a). This analogue also retained bioactivity comparable to synaptamide as indicated by the cAMP production in cultured cortical neurons (Fig. 3f). As the fatty acyl ethanolamides are lipophilic, non-specific binding to cell membranes made it difficult to microscopically determine specific binding of bodipy-synaptamide to the receptors on the cell surface. For this reason, we have examined endocytic receptors as fluorescent puncta based on the fact that most GPCRs are internalized after ligand binding 27. Indeed, the cortical neurons treated with bodipy-synaptamide for 30 min showed punctate fluorescence signals in the cell body and neurites (Fig. 3g). We observed that an N-terminal targeting mGPR110 antibody used to detect full length GPR110 in Supplementary Figs S6a and S9a dose-dependently decreased synaptamide-induced cAMP production in both cortical neurons and NSCs (Supplementary Fig. 9b), suggesting that this antibody successfully interfered with the binding of synaptamide to GPR110. When GPR110 is blocked by the pretreatment with this N-terminal targeting anti-GPR110 antibody under a non-permeable condition, the fluorescent puncta were no longer visible, indicating that those puncta signals were derived from the binding of bodipy-synaptamide to GPR110 on the surface of cortical neurons. The pretreatment of the neurons with synaptamide also abolished the puncta, indicating the binding of synaptamide to GPR110. Both microscopic and in-cell cross-linking data consistently demonstrated that synaptamide binds to GPR110 in living cells. Synaptamide potently induced cAMP production with EC50 in the low 2–5 nM range in cultured cortical neurons and NSCs (Fig. 3h). The same N-terminal targeting GPR110 antibody completely blocked the synaptamide-induced cAMP production, indicating the importance of synaptamide binding to GPR110 to increase cAMP. A physical binding assay based on microscale thermophoresis 28,29 using immunopurified hGPR110 and bodipy-synaptamide, a fluorescent synaptamide analogue, indicated apparent Kd in the low nM range (Supplementary Fig. 10), further supporting synaptamide as a potent GPR110 ligand.

Neither GPR110 binding nor synaptamide-like bioactivity was observed with other endogenous N-acyl ethanolamides. Only synaptamide blocked cellular binding of bodipy-synaptamide detected as the fluorescent endocytic receptor puncta (Supplementary Fig. 11), and significantly displaced G1 bound to GPR110 (Fig. 2e), indicating that synaptamide is a specific ligand to GPR110. Accordingly, other fatty acid ethanolamides did not significantly induce cAMP production (Supplementary Fig. 12a) or cortical neurite outgrowth (Supplementary Fig. 12b).

N-terminal part of GPR110 is important for synaptamide binding. The observation that the N-terminal targeting antibody blocked the synaptamide binding to GPR110 (Fig. 3g) and synaptamide-induced cAMP production (Fig. 3h, Supplementary Fig. 9b) led us to test the significance of GPR110 N-terminus in synaptamide-induced GPR110 activation. GPR110 has a long N-terminus with an autoproteolysis-inducing (GAIN) domain containing a GPCR proteolysis site (GPS). As a cleaved form derived from GPS hydrolysis is considered important for GPR110 activation 30, we prepared the HA-tagged mutant forms shown in Fig. 4a. These hGPR110 mutants overexpressed in HEK-CRE-luc2P cells were tested for synaptamide analogue G1 binding and cAMP production (Fig. 4b,c). Double mutation at GPS (DM, H565A/T567A) to minimize the autocleavage at GPS altered neither ligand binding ability nor synaptamide-induced cAMP production observed with wild type GPR110 (WT). When the N-terminus was truncated at GPS (CTF), both G1-binding and cAMP production induced by synaptamide were either no longer apparent or greatly diminished at best. In contrast, the N-terminal side of the cleavage product (NTF) showed considerable binding to G1 although cAMP production was not induced as expected from the lack of the C-terminus for G-protein coupling. Furthermore, synaptamide did not affect GPR110 autolysis (Fig. 4d). These data clearly indicate the importance of the N-terminal sequence for synaptamide-induced GPR110 activation.

GPR110 is a functional receptor for synaptamide. The functional significance of GPR110 was tested for synaptamide-induced cAMP production and neurite outgrowth after disrupting synaptamide-GPR110 interaction through pretreatment with...
the N-terminal targeting GPR110 antibody described above or by GPR110 gene silencing or overexpression (Fig. 5). Blocking GPR110 by pretreatment of cortical neurons with the GPR110 antibody for 30 min prevented a synaptamide-induced increase in cAMP (Fig. 5a), CRE transcriptional activity (Fig. 5b) and neurite outgrowth (Fig. 5c,d). Similarly, synaptamide-induced cAMP production and neurogenic differentiation of NSCs was diminished after pretreatment with the same GPR110 antibody while the...
Data represent three independent experiments (HEK293 cells permanently expressing CRE-luc2P as a cAMP sensor and treated with 10 nM synaptamide for 16 h. M45: empty vector expressing cells. Synaptamide did not affect hGPR110 autocleavage (GPS (DM) to prevent the autocleavage did not alter the ligand binding ability or synaptamide-induced cAMP production. The treatment with 10 nM (C)-type full size hGPR110 (1–910, WT), GPS cleavage sequence (HL^T) double mutant (H565A/T567A, DM), functional receptor for synaptamide. Synaptamide to GPR110 clearly indicate that GPR110 is a control cells. These data altogether with physical binding of synaptamide effect was observed in empty vector-transfected overexpressed in HEK293-CRE-luc2P cells, 10 nM synaptamide outgrowth (Fig. 5f,g) and cAMP production (Fig. 5h). In contrast almost completely abolished synaptamide-induced neurite scramblified with GFP-expressing shRNAs compared with the scrambled control RNA (Sc) (Fig. 5e). The GPR110 knockdown was specific to GPR110 KO animals, indicating that the probe used for this assay was specific to gpr110. Similarly, the gpr110 signal detected after in situ hybridization also indicated expression of gpr110 in the P0 brain and adult hippocampal DG, confirming the qPCR results (Fig. 6b). The gpr110 was strongly detected in the glomerular structure of kidney but was not detected in the kidney from GPR110 expression in brain. The temporal and regional specificity of GPR110 expression in the brain was apparent. The GPR110 mRNA expression was notably high in NSCs and fetal brains (E14) and decreased significantly after birth (Fig. 6a). In 4 month old adult brains, cortical expression of GPR110 is minimal as reported earlier (Lum et al.24); however, hippocampus showed a considerable signal at a level comparable to that of newborn brains (P0). The dentate gyrus (DG) in particular, where neurogenesis occurs in adulthood, showed higher GPR110 expression in comparison to other hippocampal regions. Neurogenesis occurs in adulthood, showed higher GPR110 expression in comparison to other hippocampal regions. Neurogenesis occurs in adulthood, showed higher GPR110 expression in comparison to other hippocampal regions.
GPR110 KO abolishes synaptamide bioactivity. The essential role of GPR110 in synaptamide-induced bioactivity was further confirmed using GPR110 KO mice recently produced in our laboratory. The endogenous gpr110 was deleted in GPR110 KO mice using both in situ hybridization and PCR for P0 kidney and brain cortex (Figs 6b and 7a). The endogenous synaptamide level in the P0 cortex from WT and KO animals was similar (Fig. 7b). While WT cortical neurons significantly increased cAMP production and neurite growth following synaptamide treatment, no synaptamide effect was observed in neurons prepared from KO mice (Fig. 7c,d, Supplementary Fig. 16). Deletion of gpr110 also prevented synaptamide-induced synaptogenesis seen in WT neurons (Fig. 1b). The expression of synapsin1 and PSD95, as well as the synapse number evaluated by synapsin1/PSD95 overlapping...
puncta were significantly lower in KO neurons compared with WT (Fig. 7e). More importantly, synaptamide increased neither synapsin1 and PSD95 expression nor synaptogenesis, unlike the case with WT neurons. These data indicate the importance of GPR110 in mediating synaptamide bioactivity. To determine the in vivo significance of GPR110, the synapse number was evaluated using synapsin1/PSD95 and synapsin1/Homer1 co-localizing puncta in the cortical synaptic zone (Fig. 7f). Although the endogenous synaptamide level was similar (Fig. 7b), the lack of synaptamide receptor GPR110 resulted in a severe deficit not only in individual synaptic protein puncta but also in synapse number. Western blot analysis indicated decreases in the synapsin1, PSD95 and Homer1 protein level in KO animals (Supplementary Fig. 17), further confirming the synaptic protein loss microscopically evaluated in Fig. 7f. These in vitro and in vivo data from GPR110 KO mice provide clear evidence that synaptamide promotes synapse development by acting through GPR110.

GPR110 KO animals exhibit memory deficit. To further determine the in vivo significance of GPR110, the memory function was evaluated by novel object recognition and Morris water maze tests. Both WT and GPR110 KO mice at 2–4 months (2.7 ± 0.6 versus 3.0 ± 0.8 month old, WT versus KO; P = 0.15; t-test) explored the novel object for a longer time than the familiar one; however, GPR110 KO mice spent significantly less time exploring the novel object than WT mice (P < 0.05; t-test; Fig. 7g), even though the total exploration time for both objects is similar between the genotypes (17.8 ± 0.78 versus 18.04 ± 0.89 s). In the Morris water maze test, WT and GPR110 KO mice showed similar learning curves from day 1 to day 4, with the escape latencies and distance to platform for GPR110 KO mice being similar to the WT mice (P > 0.05; t-test; Supplementary Fig. 18), suggesting that both WT and GPR110 KO mice had comparable visual acuity, swimming skills and ability to learn the task of reaching the platform. As escape latencies are not a sensitive measure of reference memory31, the probe trials were performed. During the
Figure 7 | Synaptamide bioactivity abolished by GPR110 KO. No expression of gpr110 in GPR110 KO kidney and brain cortex indicated by PCR and in situ hybridization (a). The endogenous synaptamide level (mean ± s.d., biological triplicates, n = 3) in WT and KO PO brains determined by mass spectrometry was comparable (b). In cortical neurons from KO mice, synaptamide up to 100 nM showed no effect on cAMP production (c) or neurite outgrowth (d). Co-localization analysis of presynaptic marker synapsin1 (red) and postsynaptic marker PSD95 (green) revealed increased synapses (overlapping puncta indicated by arrows) by synaptamide treatment in WT but not in KO cortical neurons (e). For quantification of synaptic protein puncta in primary neuron culture, 20 cells were microscopically scored per condition, and the data represent three independent experiments. In vivo synaptogenesis evaluated in cortical synaptic zone of P10 mouse brain showing severe loss of synapsin1, Homer1 and PSD95 puncta as well as synapsin1/Homer1 or synapsin1/PSD95 colocalizing puncta in the KO brain, indicating significant loss of synapses (f). KO animals exhibited significantly reduced recognition memory (g) and spatial memory compared with WT (h). Schematic representation of synaptamide signalling (i). DHA-derived synaptamide binds to GPR110 which in turn triggers Gqα activation, cAMP increase and CREB phosphorylation to promote neurite growth, synaptic protein expression and NSC neurogenic differentiation. Data represent three (a–e) and two (f) independent experiments. Values are mean ± s.e.m. of biological triplicates (n = 3) unless specified otherwise. *P < 0.05, **P < 0.01, ***P < 0.001, #P < 0.05, ##P < 0.01, ###P < 0.001 versus corresponding control (c–e) or WT platform quadrant time (h) by unpaired Student's t-test. For the comparison of the time in four quadrants within the WT or KO group (h), one-way ANOVA followed by Tukey post hoc test was used with the statistical significance at P < 0.05. Means designated with the same letter (a or b) are not significantly different. NT, N-terminus. N.S., Not significant. * relative to control. Scale bars, 100 μm (a) and 5 μm (e,f).
Discussion

Synaptamide is an endogenous metabolite of DHA with potent bioactivity on neurite growth and neurogenesis. Taking advantage of chemical analogues of synaptamide, molecular interaction of synaptamide with the target receptor was probed using cell- and mass spectrometry-based approaches in this study. We discovered that synaptamide is an endogenous ligand to the orphan GPR110, triggering Gz-coupled cAMP production. Synaptamide-induced neurogenic differentiation and neurite outgrowth were dependent on synaptamide binding to GPR110, indicating the importance of the GPR110 signalling for the observed bioactivity of synaptamide. Synaptamide signalling through GPR110 activation leading to neurogenesis, neurogenesis and synaptogenesis is schematically depicted in Fig. 7i.

The signalling mechanisms as well as biological function are still elusive for many adhesion GPCRs largely due to lack of known agonists, although some activation mechanisms and functional involvement have recently emerged34. The GAIN domain/GPS has been shown to mediate receptor activity either in a GPS-cleaved or GPS-intact fashion35,36. It has been suggested that autoproteolysis at the GPS is critical for the trafficking and function of adhesion GPCR37,38, and the resulting short N-terminal stalk region of the 7TM domain was recently shown to be a tethered agonist for the activation of GPR56 and GPR110 (ref. 39). In contrast, the proteolytic cleavage is not required for the function of some adhesion GPCRs such as BA13 (ref. 39), GPR133 (ref. 40) and latrophilin-1 (LAT-1) receptors41, suggesting the importance of the highly conserved GAIN/GPS motif itself for receptor signalling. We observed that synaptamide remained capable of binding to and activating the cleavage resistant mutant form (H565A/T567A) of GPR110 (Fig. 4), indicating that GPS cleavage is not necessary for synaptamide-induced GPR110 activation. In fact, synaptamide was not able to bind to or activate GPR110 when the N-terminus was truncated, suggesting that this endogenous ligand signals in a manner distinctively different from the GPS cleavage-dependent mechanism. The requirement of the N-terminus for synaptamide-induced GPR110 activation enabled us to successfully employ an N-terminal targeting antibody as a tool to block the synaptamide action, providing additional evidence that GPR110 is the synaptamide target receptor mediating neurogenic differentiation of NSCs and neurite growth. With synaptamide identified as an endogenous ligand for GPR110 in this study, molecular details of GPR110 activation may now be further investigated under a physiologic context.

Although aGPCRs have seven TM like other GPCR families, intracellular G protein coupling has not been well-established. Enhancement of constitutive GPR110 activity was observed when GPR110 and Gzq were co-overexpressed42. In addition, basal G protein activity was shown to increase when isolated GPR110-expressing membranes were reconstituted with Gzq but not with any other G proteins, suggesting Gzq as an exclusive GPR110 partner at least in vitro43. Nevertheless, it is not clear whether Gzq coupling is an endogenous G protein signalling mechanism for ligand-activated GPR110. In this work, using a co-IP approach, we demonstrated the association of GPR110 with endogenous Gzs but not with Gzi and Gzq, and this coupling was further verified using overexpressed Gzs (Fig. 2i, Supplementary Fig. 8). The fact that synaptamide induces a cAMP increase that is insensitive to pertussis toxin (Fig. 1) is consistent with GPR110 coupling with Gzs.

Synaptamide biosynthesis was widely observed in brain cells and homogenate preparations30,31,33, although the exact biosynthetic mechanism has yet to be established. In addition, endogenous expression of GPR110 was detected in cultured cortical neurons and NSCs (Supplementary Fig. 9), as well as in fetal and adult brain regions (Fig. 6). The persistent presence of both synaptamide biosynthetic activity and GPR110 expression in brain cells from embryonic to adult stages, despite varying extent, suggests sustained operation of synaptamide/GPR110 signalling in the brain. Previously, expression of this receptor was shown to be low except in kidney33, however, we found that in the fetal brain and NSCs, GPR110 mRNA expression is notably higher than in postnatal brains, and is even comparable to that in kidney (Fig. 6a). Furthermore, hippocampal DG, which retains neurogenic capacity throughout life43, showed sustained expression of GPR110, suggesting a role of GPR110 in promoting neurogenesis even after embryonic development. GPR110 KO mice recently produced in our laboratory showed no appreciable phenotype in terms of viability or fertility as has been previously described44; however, detailed phenotypic characterization needs to be performed when a sufficient number of KO animals become available. Nevertheless, GPR110 KO abolished the synaptamide bioactivity (Fig. 7c–e) and significantly reduced synaptic protein expression and synapse number in developing brains (Fig. 7f, Supplementary Fig. 17) despite no alteration in endogenous synaptamide levels (Fig. 7b), indicating the importance of synaptamide action through GPR110 for synapse development. While GPR110 may not be critical for survival, the synaptamide/GPR110 signalling may be an essential mechanism to meet the developmental need for active neurogenesis, neurogenesis and synaptogenesis which are vital for proper development of brain function. Indeed, the recognition memory and spatial memory were significantly compromised in adult GPR110 KO mice, supporting the notion that developmental deficits in synaptamide/GPR110 signalling can lead to suboptimal brain function at a later stage. Considering the close link between DHA and synaptamide levels in brain11,14, inadequate synaptamide-activated GPR110 signalling, particularly during development, may provide an explanation for learning disability and memory deficits associated with brain DHA reduction caused by insufficient dietary omega-3 fatty acid intake45. Likewise, genetic or environmental modification of GPR110 may also have significant ramifications in development of brain function.

In conclusion, synaptamide, an endogenous metabolite of DHA, is a specific GPR110 ligand which triggers cAMP-dependent signal transduction leading to potent induction of neurogenesis, neuritogenesis and synaptogenesis in developing neurons. GPR110 deorphanized as the functional receptor for synaptamide provides a mechanism that explains the critical importance of omega-3 fatty acid nutrition in brain development. Furthermore, GPR110 emerges as a potential novel target for controlling physiological and pathophysiological processes of neurodevelopment and function.

Methods

Animals. Time pregnant female C57BL/6 mice or Wistar rats were obtained from NIH-NCI animal production program or Charles River Laboratories (Portage, MI).
respectively, and GPR110 (adhesion G protein-coupled receptor F1: Adgrf1) heterozygous mice on C57BL/6 background were generated by Knockout Mouse Project (KOMP) repository. Animals were housed in SPF facility and acclimated for a day before brains were collected for the preparation of primary cortical neurons or neural stem cells (NSCs), in situ hybridization or immunohistochemistry. GPR110 KO mice were generated by heterozygote mating in our laboratory. All experiments in this study were carried out in accordance with the guidelines for the care and use of animals approved by the National Institute on Alcohol Abuse and Alcoholism (LMS-HK13 or 41).

Primary cell culture and transfection. Primary cortical neurons were prepared from C57BL/6 mice brains according to the established protocol46–48. Briefly, cortices were isolated from P0 pups and digested with 100 U of papain for 30 min at 37 °C and mechanically disrupted by pipetting several times in neurobasal medium (In VitroGen, Carlsbad, CA) supplemented with 2% B27 (In VitroGen, Carlsbad, CA) and 0.1% GlutaMax (In VitroGen, Carlsbad, CA). The dissociated cortical neurons were seeded in poly-D-lysine-coated 24-well plates (0.5 × 10^6 cells per well) or 60-mm dishes (2.5 × 10^5 cells per dish) for neurite outgrowth analysis or western blotting, respectively. Electroporation of primary cortical cells was performed before plating using an AMAXA V4XC-2024 4D-Nucleofector X kit (Lonza, Basel, Switzerland). In these studies, 5 × 10^5 cells were transfected into a new tube, centrifuged at 800 g for 10 min at room temperature and the pellet was suspended in 100 µl of transfection solution. After addition of 3 µg of plasmid DNA, cells were transfected into the electroporation cuvette and pulsed. Subsequently, the pulsed cells were plated in 1,000 µl of medium (20 mM HEPES, 100 mM sodium butyrate, bicarbonate CO2 incubator (5% CO2, 37 °C) for 2 h before the medium change. HEK293 and A549 cells that were authenticated and certified with non-detectable mycoplasma contamination were obtained from American Type Culture Collection (ATCC, Manassas, VA). HEK293 cells were cultured in DMEM/F12 (1:1) (Thermo-Millipore), total 1% (v/v) fetal bovine serum (FBS, Invitrogen, Carlsbad, CA) in the humidified CO2 incubator. A549 cells were cultured in F12K (Invitrogen, Carlsbad, CA) with 10% FBS. GPR110-HA transfection in HEK293 and A549 cells were performed using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s instructions.

NSCs were cultured by the neurosphere method described earlier11,49. Briefly, rat or mouse cortical NSCs were dissociated into single cells by repeated pipetting in a serum-free condition (N2 medium) containing DMEM/F12 (1:1), 0.6% (wt/vol) glucose, 0.1125% (wt/vol) sodium bicarbonate, 2 mM l-glutamine, 5 mM HEPES, 100 µg/ml human apo-transferrin, 20 nM progesterone, 30 nM sodium selenite, 60 µM putrescine and 25 µg/ml insulin. The dissociated cells were cultured in 60-mm dishes at 5 × 10^4 cells per dish in N2 medium with 20 ng/ml basic fibroblast growth factor (bFGF) and 2.5 µM vitamin E and 2.5 µM vitamin E. After 16 h, the CRE activity was measured with dual luciferase reporter assay kit (Promega) and the renilla luciferase activity was normalized to firefly luciferase activity. The effect of GPR110 overexpression on synaptamide activity was also evaluated using CRE-luc2P HEK293 cells (Promega). CRE-luc2P HEK293 cells were plated in 24-well plates at 0.25 × 10^5 cells per well and transfected with full length GPR110-HA (WT), MgPr110-DM-HA (HG56/5767A GPS double mutant), MgPr110-N-terminal truncation GPR110-HA C-terminal fragment (CCT), MgPr110-N-terminal fragment (NTF) or empty vector M45 (GeneCopoeia) for 24 h, and stimulated with 10 nM synaptamide, in DMEM containing 0.01% BSA and 40 µM vitamin E. After 16 h, the CRE activity was determined using the Dual-Glo luciferase assay kit (Promega), following the manufacturer’s instructions. All cell lines were tested negative for mycoplasma contamination.

Luciferase assay for CRE activity. Cortical cells were plated in 24-well plates at 2.5 × 10^5 cells per well and cultured in neurobasal medium supplemented with 2% B27, 1% glucose/hydrocortisone mixture (1:3). Transient transfection of inducible CRE-responsive firefly luciferase reporter and Renilla luciferase construct (4010) (Qiagen, Valencia, CA) was performed on DIV 2 using Lipofectamine 2000 (Invitrogen). After transfection for 24 h, cells were stimulated with simpatamide, DHA or OA in DMEM containing 0.01% BSA and 40 µM vitamin E. After 18 h, the CRE activity was assayed using the Dual Luciferase Reporter Assay kit (Qiagen) and the renilla luciferase activity was normalized to firefly luciferase activity. The effect of GPR110 overexpression on synaptamide activity was also evaluated using CRE-luc2P HEK293 cells (Promega). CRE-luc2P HEK293 cells were plated in 24-well plates at 0.25 × 10^5 cells per well and transfected with full length GPR110-HA (WT), MgPr110-DM-HA (HG56/5767A GPS double mutant), MgPr110-N-terminal truncation GPR110-HA C-terminal fragment (CCT), MgPr110-N-terminal fragment (NTF) or empty vector M45 (GeneCopoeia) for 24 h, and stimulated with 10 nM synaptamide, in DMEM containing 0.01% BSA and 40 µM vitamin E. After 16 h, the CRE activity was determined using the Dual-Glo luciferase assay kit (Promega), following the manufacturer’s instructions. All cell lines were tested negative for mycoplasma contamination.

RT-PCR analysis. Brain tissues were homogenized using a glass Teflon homogenizer. RNA was extracted with an RNeasy mini kit (Qiagen) and measured by a NanoDrop 1000 spectrophotometer (Thermo Scientific). cDNA synthesis was carried out using a cDNA reverse transcription kit (Applied Biosystems) according to manufacturer’s protocol. Briefly, 1 µg of isolated RNA, 50 nM random RT primer, 0.25 µM dNTPs, 0.25 µM M-200 U-1 reverse transcriptase and 0.25 µL 10 nM RNAase inhibitor were used for each RT reaction. The forward/ reverse primers used in polymerization chain reactions include 5'-GACCAAGGAGGCACAAACCCTC-3'/5'-TGTGAAAGGCCACCCAGGATGG-3' and 5'-AAAAAGGGTTCTCGAGTGCTTCG-3'/5'-AAGGATGCGCCCAATCTTGTGTC-3' for GPR110 and 5'-CAACACGTCGATCATCCT-3'/5'-CAGCCCCCTTTGCCG-3' for glyceraldehydes-3-phosphate dehydrogenase (GAPDH), respectively. Quantitative real-time PCR was carried out using a Quantitect SYBR Green PCR kit (Qiagen) and an ABI Prism 7900 HT system. Briefly, 1 µl cDNA, 6.5 µl of Quantitect SYBR Green master mix, 1 µl of each of GPR110 primers and the control GAPDH primers was used in each reaction (96-well plate). The thermal cycler program was composed of 2 min at 50 °C, 15 min at 95 °C followed by 40 cycles of denaturation at 95 °C for 15 s, annealing at 55 or 57 °C for 10 s and extension at 72 °C for 30 s. The relative quantification of mRNA expression was performed using the comparative threshold method.

cAMP assay. Approximately, 1.25 × 10^5 NSCs or 0.25 × 10^5 cortical cells were cultured in 0.5 ml medium unless otherwise specified. Cultured NSCs (DIV 4) or cortical cells (DIV 3) were transfected with plasmids that were added to date the cAMP levels were determined using cAMP AXP assay kit (Cell Signaling Technology, Danvers, MA) according to the manufacturer’s protocol. Briefly, cAMP from the cell lysate was added to the kit to displace HRP-linked cAMP which was bound to an anti-cAMP XP Rabbit mAb immobilized on a 96-well plate. After removing excess HRP-substrate TMB solution, the colorimetrically determined at 450 nm. In some cases, cells were pretreated with 100 µM adenosyl cyclase inhibitor SQ22,536 (Sigma) or 0.4 µg ml−1 IgG (Cell signaling Technology, Thermo Scientific). Raw data was analyzed using Graphpad Prism 5.0 software.
Danvers, MA) or N-terminal targeting GPR110 antibody (Abmart) for 30 min before the stimulation with synaptamide. For evaluating synaptamide-induced cAMP production in GPR110 overexpressing cells, GPR110 residues were transfected with HA-tagged mGPR110-HA or empty vector (M45) for 24h, and treated with 10–100 nM synaptamide for 10 min. In some cases, the transfected cells were pretreated with 0.4 µg ml⁻¹ GPR110 antibody or control IgG for 30 min before the stimulation with synaptamide.

Affinity purification and MS analysis. Fetal mouse brain or HEK293 cells overexpressing mouse or human full length GPR110-HA (WT), hGPR110-DM-HA (H565A/T567A GPS double mutant), N-terminal truncated hGPR110-HA C-terminal fragment (CTF), C-terminal truncated hGPR110-HA N-terminal fragment (NTF) or empty vector M45 (GeneCoppoeia) were mildly lysed in phosphate buffered saline (PBS) containing 0.5% Triton X-100 and protease inhibitors. The lysate was treated with 1 µM G1 or biotin control for 20 min, followed by incubation with streptavidin beads at 25°C for 30 min. After washing, the affinity-purified products were eluted with 2 x lithium dodecyl sulphate buffer (LDS) for 30 min at 37°C and subjected to SDS-PAGE for western blot (WB) or MS analysis. MS analysis and protein identification were carried out using the methods described in Supplementary Fig. 22. Briefly, the entire gel was cut into several bands and each band diced into small pieces. For identification of G1 binding GPR110-HA, affinity purified GPR110-HA was run on the gel and the bands of interest were cut off in the apparent molecular weight around 90 and 110 kD where positive detection by HA antibody (Santa Cruz Biotechnology, 1:200) was indicated. Following reduction by DTT and alkylation, the affinity-purified GPR110-HA was subjected to SDS-PAGE. For identification of G1 binding GPR110-HA, the Affinity-purified hGPR110-HA in PBS containing 0.5% Triton X-100 and protease/ phosphatase inhibitors followed by incubation with streptavidin beads at 25°C for 30 min and the beads were washed three times with the lysis buffer. The affinity-purified products were eluted with 2 x LDS at 37°C for 30 min and subjected to SDS-PAGE analysis.

Immunopurification and deglycosylation. HEK cells expressing mouse or human HA-tagged GPR110 were lysed with PBS containing 0.5% Triton X-100 and protease/phosphatase inhibitors followed by immunoprecipitation with anti-HA antibody (Santa Cruz) for co-IP western blot analysis. For mass spectrometric analysis, the immunoprecipitated GPR110-HA were washed five times with lysis buffer, and GPR110-HA was eluted with HA-peptide (Thermo Fisher, 1 mg ml⁻¹ in lysis buffer). For deglycosylation, the immunopurified GPR110-HA was subjected to N-linked and O-linked deglycosylation using PNGase F or a deglycosylation enzyme mix from New England BioLabs in accordance with the manufacturer’s instruction. GPR110-HA samples with or without deglycosylation were subjected to SDS-PAGE, Coomassie staining, in-gel tryptic digestion and MS analysis.

Competition binding assay. HEK293 cells overexpressing human or mouse GPR110-HA was mildly lysed in PBS containing 0.5% Triton X-100 and protease inhibitors. The lysate was treated with 1 µM G1 for 30 min in the presence of 0–10 µM synaptamide. The treated lysate was incubated with streptavidin beads at 25°C for 30 min and the beads were washed three times with the lysis buffer. The affinity-purified products were eluted by incubating with 2 x LDS at 37°C for 30 min and subjected to SDS-PAGE for western blot analysis.

In-cell cross-linking and in-cell binding assay procedures. GPR110-HA overexpressed HEK293 cells or NSCs were washed with PBS, treated with 1 µM G1* (the cross-linkable analogue of G1) or biotin for 20 min and cross-linked by incubating with 1 µM DSS for 30 min at 25°C. Cross-linking was quenched with 30 mM Tris–HCl for 10 min. Cells were washed thoroughly using 0.1% BSA-containing buffer, and lysed with PBS containing 1% Triton X-100 and protease inhibitors. The lysate was incubated with streptavidin beads at 25°C for 30 min and washed three times. The affinity-purified products were recovered by incubating the streptavidin beads with 2 x LDS at 37°C for 30 min, and subjected to SDS-PAGE for western blot analysis. To monitor fluorescent endocytic receptor puncta formation in living cells, cortical cells were treated with 100 nM bodipy-synaptamide for 30 min with or without pretreatment with various fatty acyl ethanolamides (1 µM) or 0.4 µg ml⁻¹ N-terminal targeting GPR110 antibody for 30 min. After washing with PBS-T, the cells were fixed with 0.4% paraformaldehyde (Sigma) in PBS (pH 7.4) for 30 min for microscopic analysis.

Plasmid constructs and shRNA. CMV expression vectors for C-terminal HA-tagged GPR110 constructs were prepared by GeneCoppoeia (Rockville, MD). The construct contains the full-length open reading frame sequence of human GPR110 (NM_153840.2) and mouse GPR110 (NM_133776.1). The hGPR110-DM-HA have H565A and T567A GPS double mutations in full-length human GPR110. N-terminal fragment (NTF)-HA is a N-terminal domain of human GPR110 residues 567–910. The control empty vector is M45. The pCDNA expression vector for GFP-tagged Gzs was kindly provided by Dr Mark Rasenick (University of Illinois College of Medicine, Chicago, IL). Knockdown of GPR110 by shRNA was performed using GPR110 OmicsLiu short hairpin RNA clones with target sequences of 5′gagaagacgctagagccgctatccgctccgcgcacggtagcagagcaagaattg and 5′ccgctgttggtttttagtttagatattgacac a for sh2 both of which co-express GFP protein to facilitate visualization of transfected cells. The shRNA and non-targeting scrambled shRNA (Sc) vectors were obtained from GeneCoppoea (Rockville, MD).

Co-immunoprecipitation. HEK293 cells were co-transfected with empty vector (CMV), HA-tagged GPR110, and/or GFP-tagged Gzs by using Lipofectamin 2000 (Invitrogen). Cells were lysed in ice-cold lysis buffer (Thermo Scientific, Danvers, MA). Cell lysates were pre-cleared with protein A-garose beads to minimize non-specific binding. As an additional control for non-specific binding, samples were also immunoprecipitated with mouse IgG. The GPR110-HA protein was immunoprecipitated from cell lysates (1 µg of total protein for each sample) using anti-HA antibody (Santa Cruz Biotechnology, Inc., Dallas, TX). The co-immunoprecipitated products were subjected to SDS-PAGE and immunoblotted using antibodies against Gzs/GQ or GFP (Millipore, Temecula, CA, 1:1,000). Separately, endogenous Gzs was immunoprecipitated from cell lysates (1 µg of total protein for each sample) using anti-Gzs (Millipore). The same conditions were used for Gzs and anti-HA antibody (Santa Cruz Biotechnology, 1:200). Alternatively, GPR110 from cortical cells was immunoprecipitated using a mouse GPR110 antibody (Abmart, Shanghai, China).

Western blotting. Proteins in cell lysates (generally 30 µg protein) were electrophoresed in 4–12% Bis-Tris gels at 100 V using MOPS SDS running buffer. Proteins were electrophoretically transferred to a PVDF or nitrocellulose membrane at 100 V for 1.5 h. The membrane was blocked with 5% milk or BSA in TBS containing 0.1% Tween 20 (TBS-T) at room temperature for 1 h. Blots were incubated with primary antibodies at 4°C overnight. After washing with TBS-T buffer, the membranes were incubated for 60 min with peroxidase-conjugated secondary antibodies in TBS-T, and the labelled proteins were detected with chemiluminescence reagents (Thermo Fisher Scientific, Rockford, IL). Unless specified otherwise, α-tubulin, or β-actin (Abcam, 1:1,000) were used as the loading control for all western blotting analyses. Western blot bands were quantified using a Kodak Gel Logic 440 imaging system with ImageQuant 5.1 software (Molecular Dynamics, Sunnyvale, CA). All uncropped western blots can be found in Supplementary Fig. 22.

Binding of bodipy-synaptamide to immunopurified hGPR110-HA. Direct binding of the bodipy-synaptamide to hGPR110-HA was measured by microscale thermophoresis (MST) with a Monolith NT.115 instrument (NanoTemper Technologies, Munich, Germany). Bodipy-synaptamide at a fixed concentration (20 nm) was mixed with increasing concentrations of immunopurified hGPR110-HA in PBS containing 0.5% Triton X-100 and protease inhibitors. After incubation for 10 min at 4°C, the samples were centrifuged at 14,000 g for 4 min and loaded into MST premium capillaries (NanoTemper). Measurements were performed with various LED light. Data analyses were performed using the NanoTemper analysis software.

RNA in situ hybridization. Tissue sections cut at 20 µm thickness were processed for RNA in situ detection using the RNAScope Detection Kit (Chromogenic) according to the manufacturer’s instructions (Advanced Cell Diagnostics, Hayward, CA). Briefly, frozen tissue samples were incubated with Pretreat 1 buffer for 10 min at Room temperature (RT). Slides were boiled in Pretreat 2 buffer for 15 min, followed by incubation with Pretreat 3 buffer for 30 min at 40°C. RNAScope probes used were gpr110 (NM_133776.2, cat#300031) and daph (negative control probe, EF191515, region: 414–862, cat# 310043).

Synaptic protein puncta analysis. For synaptic protein puncta analysis, 10 day old (P10) GPR110 KO and littermate WT mice from GPR110 heterozygote mating were used. The brains were fixed overnight with 4% paraformaldehyde in PBS at 4°C, cryoprotected with 30% sucrose in PBS and then embedded in 30% sucrose in PBS/OCT (Tissue-Tek, Sakura, Japan), and cryosectioned at 15 µm using Leica CM3050S (Leica, Germany). The coronal sections were washed and permeabilized in PBS with 0.2% Triton-X 100 (TBST; Roche, Switzerland) and blocked in 5% Normal Goat Serum (NGS) in PBS-T for 1 h at room temperature. The primary antibodies were incubated in 1% NGS including Anti-PDSS1 (1:500, Thermo, MA), anti-Homer1 (1:500, Abcam, MA) and anti-Synapsin1 (1:250, Cell signaling, MA) were diluted in 5% NGS containing PBST. The brain sections were incubated overnight at 4°C with...
these primary antibodies followed by Alexa-fluorophore conjugated secondary antibodies (1:200 in PBST with 5% NgLS, Life technology, MA) for 2 h at room temperature. Slides were mounted and images were acquired on a Zeiss LSM700 confocal laser-scanning microscope. Brain sections were stained with pre- 
(synapsin-1) and post-synaptic (PSD95, Homer1) marker pairs. Three random fields in the cortex synaptic zone were examined per brain section, three separate sections per mouse and three mice were used for analyses. Five μm thick confocal z-stacks (optical section depth 0.33 μm, 15 optical sections/z-stack) of the synaptic zone in cortex area were imaged at ×100 magnification. Maximum projections of three consecutive optical sections (corresponding to 1 μm total depth) were generated from the original z-stack. Analyses were performed blinded to generator. A plug-in Puncta Analyzer (available upon request from Duke University at https://sites.duke.edu/eroglulab/publications/) under the Image J analysis software platform was used to count the number of pre-, post-, and co-localized synaptic protein puncta. This quantification method is based on the fact that pre- and post-synaptic proteins are not within the same cellular compartments and would appear co-localized only at synapses due to their close proximity. Detecting two identical objects were placed in the chamber with two objects, one object was from the first trial while the other object trial was performed 24 h later to assess object memory. Mice were placed in the chamber with two objects, one object was from the first trial while the other object. Mice were allowed to freely explore the two objects for 10 min or till 20 s of exploration. A mouse was considered to explore the objects when it was looking toward the object with its nose within 2 cm of the object. The second trial was performed 24 h later to assess object memory. Mice were placed in the chamber with two objects, one object was from the first trial while the other object was a new object that the mice had not encountered before. The mice were allowed to explore the objects for 10 min or till the total exploration time was 20 s. The positions of the objects were reversed for about half of the mice tested. The floor of the chamber was divided into four equal sized quadrants. A 10 cm² transparent platform was placed 10 cm away from the walls. A seamless pool with 120 cm diameter was filled with water. A total of 34 mice were used. Four mice were excluded from the study due to technical reasons.

**Behavioural testing.** Animals were housed in SPF facility with a 12 h light dark cycle. Male mice from 2 to 4 months of age were used for behavioural experiments, which were performed during the light cycle with the experimenter blinded to the experimental groups.

**Novel object recognition test.** A modified procedure for novel object recognition test was followed35 using the ANY-maze video tracking and analysis software (Stoeiling Co., Illinois). A total of 35 mice were used. Mice were introduced in a 40 x 40 cm² opaque acrylic chamber for 5 min each for habituation. On the next day, two objects were placed in the chamber and the mice were allowed to explore for ~ 10 min away from the walls and from each other. Mice were allowed to freely explore the two objects for 10 min or till 20 s of exploration. A mouse was considered to explore the objects when it was looking toward the object with its nose within 2 cm of the object. The second trial was performed 24 h later to assess object memory. Mice were placed in the chamber with two objects, one object was from the first trial while the other object was a new object that the mice had not encountered before. The mice were allowed to explore the objects for 10 min or till the total exploration time was 20 s. The positions of the objects were reversed for about half of the mice tested. The floor of the chamber was divided into four equal sized quadrants. A 10 cm² transparent platform was placed in the centre of one of the quadrants such that it was about 5 mm below the surface of the water. A total of 34 mice were used. Four mice were excluded from the study due to technical reasons. One mouse had difficulty in swimming; one bee mice were by Ipoleh and Eroglu34. Briefly, 1 μm thick maximum projections are separated into red and green channels, background subtracted (rolling ball radius = 50) to detect discrete puncta without introducing noise. The Puncta Analyzer plug-in uses an algorithm to detect the number of puncta that are in close alignment across the two channels, yielding quantified co-localized puncta. At least five z-stack sections per brain section were taken per session and three brain sections per animal were analysed. Considering three randomly selected, two objects were placed in the chamber, one object was from the first trial while the other object. Mice were allowed to freely explore the two objects for 10 min or till 20 s of exploration. A mouse was considered to explore the objects when it was looking toward the object with its nose within 2 cm of the object. The second trial was performed 24 h later to assess object memory. Mice were placed in the chamber with two objects, one object was from the first trial while the other object was a new object that the mice had not encountered before. The mice were allowed to explore the objects for 10 min or till the total exploration time was 20 s. The positions of the objects were reversed for about half of the mice tested. The floor of the chamber was divided into four equal sized quadrants. A 10 cm² transparent platform was placed in the centre of one of the quadrants such that it was about 5 mm below the surface of the water. A total of 34 mice were used. Four mice were excluded from the study due to technical reasons. One mouse had difficulty in swimming; one bee mice were by Ipoleh and Eroglu34.

**Preparation of biotinylated synaptotagmin analogue G1.** The synthetic preparation scheme for G1, (4Z,7Z,10Z,13Z,16Z,19Z)-N-(2-[(3aS,4aS,6R)-2-oxo-octahydro-1H-thieno(3,4-d)-imidazol-4-yl]pentamido)ethyl)docosa-4,7,10,13,16,19-hexaenamide, is shown in Supplemental Fig. 19. A vial charged with DHA (75 mg, 0.23 mmol) was treated with ethane-1,2-diamine (137 mg, 2.28 mmol) followed by HATU (87 mg, 0.23 mmol) and diisopropylamine (0.060 ml, 0.342 mmol). The reaction was performed for 16 h, then concentrated to yield a product. A modified procedure for novel object recognition test was followed35 using the ANY-maze video tracking and analysis software (Stoeiling Co., Illinois). A total of 35 mice were used. Mice were introduced in a 40 x 40 cm² opaque acrylic chamber for 5 min each for habituation. On the next day, two objects were placed in the chamber and the mice were allowed to explore for ~ 10 min away from the walls and from each other. Mice were allowed to freely explore the two objects for 10 min or till 20 s of exploration. A mouse was considered to explore the objects when it was looking toward the object with its nose within 2 cm of the object. The second trial was performed 24 h later to assess object memory. Mice were placed in the chamber with two objects, one object was from the first trial while the other object was a new object that the mice had not encountered before. The mice were allowed to explore the objects for 10 min or till the total exploration time was 20 s. The positions of the objects were reversed for about half of the mice tested. The floor of the chamber was divided into four equal sized quadrants. A 10 cm² transparent platform was placed in the centre of one of the quadrants such that it was about 5 mm below the surface of the water. A total of 34 mice were used. Four mice were excluded from the study due to technical reasons. One mouse had difficulty in swimming; one bee mice were by Ipoleh and Eroglu34.
3.20 mmol), HATU (1.22 g, 3.20 mmol) and diisopropylethylamine (1.12 ml, 6.40 mmol) in DMF (10 ml). Both reactions were concentrated in vacuo before half of the DMF. The rest was purified via preparative reverse phase column chromatography (5 to 90% MeCN in water, both solvents have 1% TFA, Combi Flash Companion, 100 RedSep RF Gold column). TLC analysis on fractions was done in normal phase with silica gel plates, elution with ~10% MeOH/DCM to yield 4 (1.22 g, 66% yield). LC/MS gradient 4 to 100% acetonitrile (0.05% TFA) over 3.0 min; RT 3.23 min, ESI (M+H)3+

with PAA) to provide (4-(3-(4-((3,4-dimethyl-4-yl)-pentan-1-yl)thio)hexa-2,5-dienamido)-2-oxoethoxy)1H-thieno[3,4-d]imidazol-4(3H)-yl)pentanamide (5).

The crude diamine was then purified with DCM/MeOH, passed through a bicarbonate cartridge, concentrated and added a solution in DMF (0.5 ml) to a mixture of DHA (35 µl, 0.17 mmol), HATU (64 mg, 0.17 mmol) and diisopropylethylamine (78 µl, 0.45 mmol) in DMF (1 ml). The reaction mixture was stirred 2 h and purified by reverse phase chromatography on a Waters semi-preparative HPLC with a Luna C18 column (5 micron, 30 × 75 mm) at a flow rate of 45 ml min−1. A gradient of 10 to 50% acetonitrile in water, each with 0.1% TFA over 8 min was used during the purification. Fractions (collection triggered by UV detection at 220 nm) containing compound were passed through PL HICO MP Resin and lyophilized. Pure product 6 was isolated in low yield (7.0 mg, 10.3 µmol, 4.6% yield over two steps). LC/MS gradient 4 to 100% acetonitrile (0.05% TFA) over 3.0 min; RT 3.63 min; ESI (M+H)+

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The crude diamine was then purified with DCM/MeOH, passed through a bicarbonate cartridge, concentrated and added a solution in DMF (0.5 ml) to a mixture of DHA (35 µl, 0.17 mmol), HATU (64 mg, 0.17 mmol) and diisopropylethylamine (78 µl, 0.45 mmol) in DMF (1 ml). The reaction mixture was stirred 2 h and purified by reverse phase chromatography on a Waters semi-preparative HPLC with a Luna C18 column (5 micron, 30 × 75 mm) at a flow rate of 45 ml min−1. A gradient of 10 to 50% acetonitrile in water, each with 0.1% TFA over 8 min was used during the purification. Fractions (collection triggered by UV detection at 220 nm) containing compound were passed through PL HICO MP Resin and lyophilized. Pure product 6 was isolated in low yield (7.0 mg, 10.3 µmol, 4.6% yield over two steps). LC/MS gradient 4 to 100% acetonitrile (0.05% TFA) over 3.0 min; RT 3.63 min; ESI (M+H)+

with PAA) to provide (1H-thieno[3,4-d]imidazol-4(3H)-yl)pentanamide (heptyl)-docos-4,7,10,13,16,19-hexaenamide (6, G1).

Preparation of bodipy-snapyrimidine. The synthetic preparation scheme for bodipy-snapyrimidine, 10 is used. The reaction was stirred for 16 h and washed with saturated aqueous NaHCO3, extracted with EtOAc. The organic layer was dried (MgSO4), concentrated and purified by flash silica gel chromatography (10–30% MeOH/DCM, TLC stained with PAA) to provide 10. (4×7,10,13,16,19Z)-N-(7-amino-4-(3,4-dimethyl-4-yl)-pentan-1-yl)thieno[3,4-d]imidazol-4(3H)-yl)pentanamide (4).

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