GPR83 engages endogenous peptides from two distinct precursors to elicit differential signaling

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Running Title: Peptide-mediated differential signaling at GPR83

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List of abbreviations: CCK, cholecystokinin; CHO, chinese hamster ovary; CPE, carboxypeptidase E; GPe, globus pallidus external segment; PLC, phospholipase C; PPN, pedunculopontine nucleus; proCCK, procholecystokinin; SN, substantia nigra; SNpc, substantia nigra pars compacta; VTA, ventral tegmental area
ABSTRACT

PEN is an abundant neuropeptide that activates GPR83, a G protein-coupled receptor that is considered a novel therapeutic target due to its roles in regulation of feeding, reward, and anxiety-related behaviors. The major form of PEN in the brain is 22 residues in length. Previous studies have identified shorter forms of PEN in mouse brain and neuroendocrine cells; these shorter forms were named PEN18, PEN19 and PEN20, with the number reflecting the length of the peptide. The C-terminal five residues of PEN20 are identical to the C-terminus of a procholecystokinin (proCCK)-derived peptide, named proCCK56-62, that is present in mouse brain. ProCCK56-62 is highly conserved across species although it has no homology to the bioactive cholecystokinin domain. ProCCK56-62 and a longer form, proCCK56-63 were tested for their ability to engage GPR83. Both peptides bind GPR83 with high affinity, activate second messenger pathways, and induce ligand-mediated receptor endocytosis. Interestingly, the shorter PEN peptides, ProCCK56-62, and ProCCK56-63 differentially activate signal transduction pathways. Whereas PEN22 and PEN20 facilitate receptor coupling to Gαi, PEN18, PEN19 and ProCCK peptides facilitate coupling to Gαs. Furthermore, the ProCCK peptides exhibit dose dependent Gα subtype selectivity in that they facilitate coupling to Gαs at low concentrations and Gαi at high concentrations. These data demonstrate that peptides derived from two distinct peptide precursors can differentially activate GPR83, and that GPR83 exhibits Gα subtype preference depending on the nature and concentration of the peptide. These results are consistent with the emerging idea that endogenous neuropeptides function as biased ligands.

Key Words: G protein-coupled receptors, functional selectivity, signaling bias, post-translational peptide processing, feeding, reward.
Significance Statement (1-2 sentences; 80 words):

We found that peptides derived from proCCK bind and activate GPR83, a G protein-coupled receptor that is known to bind peptides derived from proSAAS. Different forms of the proCCK- and proSAAS-derived peptides show biased agonism, activating $G_{\alpha s}$ or $G_{\alpha i}$ depending on the length of the peptide and/or its concentration.
INTRODUCTION

GPR83 first identified in murine thymoma cells as a receptor upregulated after treatment with the glucocorticoid dexamethasone (Harrigan et al., 1989; Harrigan et al., 1991; Wang et al., 2001), was later found to exhibit expression in the brain (Brezillon et al., 2001; Pesini et al., 1998). Studies reported robust GPR83 expression in regions involved in modulation of food intake (Lueptow et al., 2018; Mack et al., 2019; Muller et al., 2013b) and reward behaviors (Fakira et al., 2019; Lueptow et al., 2018; Pesini et al., 1998). Moreover, GPR83 knockout mice demonstrate altered food intake and anxiety related behaviors (Fakira et al., 2021; Muller et al., 2013b; Vollmer et al., 2013). Recent studies show that knockdown of GPR83 in the ventral striatum attenuates morphine-induced reward (Fakira et al., 2019). Together, this provides support for involvement of GPR83 in regulation of feeding, stress and reward behaviors, and indicate that it is a potential therapeutic target (Lueptow et al., 2018).

In 2016, the neuropeptide PEN was identified as a potent and selective ligand for GPR83 (Gomes et al., 2016), an observation that has been independently confirmed (Foster et al., 2019; Nataliya Parobchak; et al., 2020). PEN was first identified in 2000 in a search for substrates of carboxypeptidase E (CPE), an important enzyme in the biosynthesis of most peptide hormones and neuropeptides (Fricker et al., 2000). The PEN precursor, named proSAAS, gives rise to several distinct peptides including big and little forms of SAAS, GAV, and LEN. Previous studies have implicated big LEN and PEN in the regulation of feeding (Wardman et al., 2011). Big LEN is a ligand for GPR171, which like GPR83 has been implicated in feeding and reward behaviors (Gomes et al., 2013; Wardman et al., 2016). Little LEN does not bind to GPR171, illustrating the importance of peptide processing on the resulting biological activity.
Most studies on PEN focused on the full-length peptide, which is 22 amino acids in length (Fricker et al., 2000; Gomes et al., 2016; Wardman et al., 2011). Smaller forms of PEN lacking 2 or more C-terminal residues were identified in mouse brain and in AtT20 cells overexpressing proSAAS (Fricker, 2010; Mzhavia et al., 2002). These forms were named PEN20, PEN19, and PEN18, with the number indicating the length (Mzhavia et al., 2002) (Fig. 1). Interestingly, PEN20 has a similar C-terminal sequence as a peptide derived from procholecystokinin (proCCK) that has been identified by peptidomic analyses of mouse brain (Fricker, 2010) (Fig. 1). This proCCK peptide corresponds to residues 56-62 of proCCK (numbering from the initiation Met). ProCCK 56-62 and PEN20 have the same five residues on their C-termini: LGALL. Shorter and longer forms of this proCCK peptide were also found in mouse brain. The ‘LGALL’ sequence in proCCK is highly conserved across species, but there are no studies reporting the biological activity of this peptide. Most of the previous studies on proCCK-derived peptides have focused on the 8-residue cholecystokinin (CCK) peptide, or N-terminally-extended forms CCK-12, CCK-33 and others (Beinfeld, 2003; Goetze and Rehfeld, 2019). CCK plays an important physiological role in digestion and functions as a neuropeptide in the central nervous system to modulate satiety and suppress hunger (Okonkwo et al., 2020; Rehfeld, 2017).

There are many examples where two or more distinct neuropeptides bind to a common receptor. A classic example is the opioid peptides, with >20 different enkephalin-containing peptides produced from proenkephalin, prodynorphin, and proopio-melanocortin (Fricker et al., 2020). All of these peptides bind to the three opioid receptors (mu, delta, and kappa), albeit with different affinities and efficacies. Recently, different opioid peptides were found to produce distinct profiles of second-messenger activation at all three opioid receptors (Gomes et al., 2020). Therefore, the first question addressed in this study was whether PEN20 and shorter peptides
bound to GPR83 and if so, whether their activities differed from those of PEN. The second question was whether the proCCK peptides can bind and activate GPR83. Our finding that GPR83 binds the truncated PEN peptides and also the conserved proCCK-derived peptides suggests that it can integrate signaling from distinct neuropeptides. Furthermore, our finding that these peptides exhibit differences in the type and extent of intracellular signaling serves as a foundation for understanding GPR83 signaling both in normal cell function and in pathological disease states, and for the development of small molecule GPR83 ligands as novel therapeutics for the treatment of anxiety, depression and addiction.
Materials and Methods

Materials

Mouse PEN (mPEN), was obtained from Phoenix Pharmaceuticals, Inc (Burlingame, CA). CHO cells were from ATCC (Manassas, VA). F12 media was from Gibco/Thermo Fisher (Waltham, MA). Fetal bovine serum (FBS) was from BioWest (Riverside, MO). PEN18, PEN19, PEN20, PEN22, CCK56-63, CCK56-62, and DYKK-tagged mouse GPR83 (mGPR83) cDNA were from Genscript (Piscataway, NJ). Protease inhibitor cocktail (cat. No. P2714), p-nitrophenylphosphorylcholine, were from Sigma-Aldrich (St. Louis, MO). \(^{125}\)I (cat No. NEZ033L001MC) and \(^{35}\)S]GTP\(_{\gamma}\)S (cat. No. NEG030H250UC) were from PerkinElmer (Shelton, CT). Pierce Iodination Reagent was from Thermo Scientific (Waltham, MA). GF/B filters were from Brandel, Inc. (Gaitherburg, MD). Forskolin, and IBMX were from Tocris Bioscience (Minneapolis, MN). HitHunter cAMP detection kit was from DiscoveRx (Fremont, CA). Anti-DYKK monoclonal antibodies (cat. No. 14793) were from Cell Signaling Technology (Danvers, MA). Anti-mouse antibody coupled to horseradish peroxidase was from Vector Laboratories (Burlingam, CA). GPR83 knockout (genotyped by quantitative polymerase chain reaction analysis) and control wild-type mouse brains were generated as described (Muller et al., 2013b). Calcium 6 assay kit was from Molecular Devices (San Jose, CA).

Cell culture

CHO cells were grown in F12 media containing 10 % FBS, 1X penicillin-streptomycin solution at 37°C and 10% CO\(_2\)/O\(_2\). Cells stably expressing DYKK-tagged mGPR83 (CHO-GPR83) were generated using lipofectamine as per manufacturer’s protocol, and maintained in growth media containing 500 \(\mu\)g/ml G418. Cells free from mycoplasma contamination were used in our studies. Receptor expression levels were routinely checked by ELISA, binding assays and qRT-PCR.
(Gomes et al., 2016). Cells were used up to 6 passages, after which a fresh vial was thawed from liquid nitrogen stocks.

**Membrane preparation**

Membranes from CHO and CHO-GPR83 cells, and from the striatum of wild-type (WT striatum) or GPR83 knockout (GPR83 KO striatum) mice were prepared as described previously (Gomes et al., 2016). Briefly, cells/striatum were homogenized in 25 volumes (1 g wet weight/25 ml) of ice-cold 20 mM Tris-Cl buffer containing 250 mM sucrose, 2 mM EGTA, and 1 mM MgCl₂ (pH 7.4) followed by centrifugation at 27,000 g for 15 min at 4°C. The pellet was resuspended in 25 ml of the same buffer, and the centrifugation step was repeated. The resulting membrane pellet was resuspended in 40 volumes (of original wet weight) of 2 mM Tris-Cl buffer containing 2 mM EGTA and 10% glycerol (pH 7.4). The protein content of the homogenates was determined using the Pierce BCA Protein Assay Reagent, after which homogenates were stored in aliquots at −80°C until use.

**[125I]Tyr-rPEN Displacement Assay**

Tyr-rPEN (200 μg) was radioiodinated using [125I] and Pierce Iodination Reagent as described in the manufacturer’s protocol (Thermo Scientific). The specific activity of the iodinated peptide was 53.5 Ci/mmol at the time of iodination (the radiolabeled peptide was used within 60 days of iodination).

Displacement binding assays were carried out for 1 hour at 37°C using 50 mM Tris-Cl (pH 7.4) containing protease inhibitor cocktail, 3 nM of [125I]Tyr-rPEN, different concentrations (0 to 10 μM) of either PEN18, PEN19, PEN20, PEN22, PEN11-20, PEN11-22, CCK56-63, or CCK56-62 and membranes (20 μg) from either CHO-GPR83 cells, parental CHO cells, or from the striatum of wild-type mice. At the end of the incubation period, samples were filtered using a Brandel
filtration system and GF/B filters. Filters were washed three times with 3 ml of ice-cold 50 mM Tris-Cl (pH 7.4) and bound radioactivity was measured using a scintillation counter. Membranes from CHO cells alone or GPR83 KO mice striatum were used as controls showing that $[^{125}\text{I}]$Tyr-rPEN did not exhibit binding that could be displaced by PEN22 (Supplemental Fig. 1A and 1I).

**GTPγS Assay**

$[^{35}\text{S}]$GTPγS binding assays were carried out as described previously (Gomes et al., 2013; Gomes et al., 2016). Briefly, membranes (20 μg) from CHO-GPR83 cells, parental CHO cells, or striatum from wild-type mice were incubated for 1 hour at 30°C with either full-length mPEN (PEN22), PEN18, PEN19, PEN20, PEN11-22, PEN11-20, CCK56-63, or CCK56-62 (0 to 10 μM final concentration) in the presence of 2 mM GDP and 0.5 nM $[^{35}\text{S}]$GTPγS in 50 mM HEPES buffer containing 5 mM MgCl$_2$, 100 mM NaCl, 1 mM EDTA (pH 7.4), and protease inhibitor cocktail. Nonspecific binding was determined in the presence of 10 μM cold $[^{35}\text{S}]$GTPγS. Basal values represent values obtained in the presence of GDP and in the absence of ligand. At the end of the incubation period, samples were filtered using a Brandel filtration system and GF/B filters. Filters were washed three times with 3 ml of ice-cold 50 mM Tris-Cl (pH 7.4), and bound radioactivity was measured using a scintillation counter. In a separate set of experiments, membranes (20 μg) from CHO, CHO-GPR83 cells, WT or GPR83 KO striatum were incubated in the absence or presence of 10 μM (final concentration) of either PEN18, PEN19, PEN20, PEN22, PEN11-22, PEN11-20, CCK56-63, or CCK56-62.

**Adenylate cyclase activity assay**

Membranes (2 μg per well) from CHO-GPR83 cells or WT striatum were incubated for 30 min at 37°C with either full-length mPEN (PEN22), PEN18, PEN19, PEN20, PEN11-22, PEN11-
20, CCK56-63, or CCK56-62 (0 to 10 μM final concentration) in assay buffer (50 mM HEPES, 10 mM MgCl₂, 100 mM NaCl, 200 μM ATP, 10 μM GTP, pH 7.4) containing 40 μM forskolin, 1X protease inhibitor cocktail and 100 μM IBMX (3-isobutyl-1-methylxanthine), and cAMP levels were measured using the HitHunter cAMP detection kit for membranes from DiscoveRx as described in the manufacturer’s protocol. In a separate set of experiments, membranes (2 μg per well) from CHO, CHO-GPR83 cells, WT or GPR83 KO striatum were incubated in the absence or presence of 10 μM (final concentration) of either PEN18, PEN19, PEN20, PEN22, PEN11-22, PEN11-20, CCK56-63, or CCK56-62. For experiments involving toxins, membranes were pre-incubated with either pertussis toxin (100 ng/ml) or cholera toxin (100 μg/ml) for 15 min at RT prior to carrying out the assay.

**Phospholipase C Assay**

PLC activity was measured by incubating membranes from CHO-GPR83 cells or striatum from wild-type mice (10 μg) with either full-length mPEN (PEN22), PEN18, PEN19, PEN20, (0 to 10 μM final concentration) for 1 hour at 37°C in 0.1 M Tris-Cl buffer (pH 7.4) containing 10 mM CaCl₂, 80 mM p-nitrophenylphosphorylcholine, and protease inhibitor cocktail (De Silva and Quinn, 1987). The amount of p-nitrophenol released was measured at 410 nm. In a separate set of experiments, membranes (10 μg) from CHO, CHO-GPR83 cells, WT or GPR83 KO striatum were incubated in the absence or presence of 10 μM (final concentration) of either PEN18, PEN19, PEN20, or PEN22.

**Intracellular Ca²⁺ Release Assay**

These assays were carried out essentially as described previously (Gomes et al., 2013; Gomes et al., 2016) except that Calcium 6 dye was used instead of Fluo-4 NW. Briefly, CHO-GPR83 cells expressing a chimeric hGα₁₅/i₃ protein were plated into black poly-l-lysine–coated 384-well clear-
bottom plates (40,000 cells/25μ growth media/well). On the next day cells were incubated with 25 μl Calcium 6 dye in HBSS buffer containing protease inhibitor cocktail for 1 hour at 37°C. Cells were then treated with buffer or either full-length mPEN (PEN22), PEN18, PEN19, PEN20, (0 to 10 μM final concentration). Increases in intracellular Ca^{2+} levels were measured for ~210 s at excitation 485 nm and emission 525 nm using the FLIPR system. In a separate set of experiments, CHO or CHO-GPR83 cells were incubated in the absence or presence of 10 μM (final concentration) of either PEN18, PEN19, PEN20, or PEN22.

**Endocytosis Assay**

Receptor endocytosis assays were carried out using previously described protocols (Gupta et al., 2015; Gupta et al., 2014). Briefly, CHO-GPR83 cells were seeded into 24-well plates (2 × 10^5 cells per well). Cells were labeled with anti-DYKK monoclonal antibodies (1:1000 in PBS containing 1% FBS) for 1 hour at 4°C, washed three times with growth media, followed by treatment with either full-length mPEN (PEN22), PEN18, PEN19, PEN20, (0 to 10 μM final concentration) in media containing protease inhibitor cocktail at 37°C for 1 h. Cells were briefly fixed (3 min) with 4% paraformaldehyde followed by three washes (5 min each) with PBS followed by incubation with anti-mouse antibody coupled to horseradish peroxidase (1:2000 in PBS containing 1% FBS) for 1 hour at 37°C. Cells were washed three times with 1% FBS in PBS (5 min each wash), and color was developed by the addition of the substrate o-phenylenediamine [5 mg/10 ml in 0.15-m citrate buffer (pH 5) containing 15 μl of H_2O_2]. Absorbance at 490 nm was measured with a Bio-Rad ELISA reader. For time course experiments, cells were treated with 100 nM of different peptides for different time periods (0-120 min).

**Data Analysis and Statistics**
Data was analyzed using Prism 9.0 software. Data throughout the manuscript is presented as Mean±SD. Displacement binding assay data (Fig. 2B, 3A, 4B & C, Supplemental Fig. 1A & I, Supplemental Table 1 & 3 was analyzed by comparing both One site-Fit logIC₅₀ and Two-sites-Fit logIC₅₀ curves to determine for each data set which of the two equations fits best. In Fig. 2B, assays with PEN18, PEN19 and PEN20 were carried out 3 independent times, and with PEN22 6 independent times each with triplicate determinations. In Fig. 2C, assays were carried out 6 independent times each with triplicate determinations. In Fig. 2D, assays were carried out 3 independent times each with quadruplicate determinations. In Fig. 2E, assays were carried out 3 independent times each with quadruplicate determinations; data was analyzed using one-way ANOVA with Tukey’s multiple comparison. In Fig. 2F, assays were carried out 6 independent times each with triplicate determinations. In Fig. 2G-I, Fig. 3A, 3B, assays were carried out 3 independent times each with triplicate determinations. In Fig. 3C, assays were carried out 3 independent times each with triplicate determinations, and data was analyzed using unpaired t-test with Welch’s correction. In Fig. 3D assays were carried out 3 independent times each with quadruplicate determinations. In Fig. 3E, assays were carried out 3 independent times each with quadruplicate determinations, and data was analyzed using unpaired t-test with Welch’s correction. In Fig. 3F, assays were carried out 3 independent times each triplicate determinations. In Fig. 4B, 4C and 4D, assays were carried out 6 independent times each triplicate determinations. In Fig. 4E, assays were carried out 5 independent times each triplicate determinations for PEN11-20 and PEN11-22 and 3 independent times each triplicate determinations for PEN22; data was analyzed using one-way ANOVA with Tukey’s multiple comparison. In Fig. 4F, assays were carried out 6 independent times each triplicate determinations. In Fig. 4G, assays were carried out 6 independent times each with quadruplicate determinations. In Fig. 4H and 4I, assays were carried out 4
independent times each with quadriplicate determinations. In Supplemental Fig. 1A, assays were carried out 6 independent times each triplicate determinations. In Supplemental Fig. 1B and 1C, assays were carried out 6 independent times each triplicate determinations, data was analyzed using two-way ANOVA with Tukey’s multiple comparison. In Supplemental Fig. 1D-F, assays were carried out 3 independent times each with quadriplicate determinations. In Supplemental Fig. 1G, assays were carried out 6 independent times each with triplicate determinations; data was analyzed using two-way ANOVA with Tukey’s multiple comparison. In Supplemental Fig. 1H, assays were carried out 3 independent times each with triplicate determinations; data was analyzed using two-way ANOVA with Tukey’s multiple comparison. In Supplemental Fig. 1I, assays were carried out 3 independent times, each with triplicate determinations. In Supplemental Fig. 1J, assays were carried out 3 independent times each with triplicate determinations; data was analyzed using unpaired t-test with Welch’s correction. In Supplemental Fig. 2A, assays were carried out 6 independent times each with triplicate determinations; data was analyzed using unpaired t-test with Welch’s correction. In Supplemental Fig. 2B, assays were carried out 5 independent times each with quadruplicate determinations; data was analyzed using two-way ANOVA with Tukey’s multiple comparison. In Supplemental Fig. 2C and 2D, assays were carried out 5 independent times each with quadruplicate determinations. In Supplemental Fig. 2B-2I, assays were carried out 4 independent times each with quadriplicate determinations.
RESULTS

Binding of C-terminally truncated PEN peptides to heterologously expressed GPR83 and characterization of the signaling pathways.

We first investigated whether the C-terminally truncated PEN peptides, PEN18, PEN19, and PEN20 (Fig. 2A) can displace [125I]Tyr-rPEN binding to membranes from CHO cells stably expressing GPR83 (CHO-GPR83). [125I]Tyr-rPEN exhibits robust binding to CHO-GPR83 membranes and not to CHO membranes alone (Supplemental Fig. 1A). Full-length PEN (referred to as ‘PEN22’) as well as PEN18, PEN19, and PEN20 exhibit biphasic displacement binding curves with high-affinity in the sub-nM range and low-affinity in the high nM - μM range in CHO-GPR83 membranes (Fig. 2B, Supplemental Fig.1A, Supplemental Table 1). Moreover, while PEN22 (10 μM) completely displaces [125I]Tyr-PEN binding to CHO-GPR83, this is not the case with PEN18, PEN19 and PEN20 which show only partial displacement (Fig. 2B, Supplemental Table 1).

The pharmacological properties of the C-terminally truncated PEN peptides were characterized using signaling assays previously used to characterize PEN22-GPR83 signaling (Gomes et al., 2016). C-terminally truncated PEN peptides activate G proteins as measured using the [35S]GTPγS binding assay in CHO-GPR83 but not CHO membranes (Supplemental Fig. 1B, Fig. 2C). PEN22 and PEN20 exhibit similar potencies, although PEN20 exhibits a lower but not statistically significant difference in efficacy (118±3%) compared to PEN22 (126±3%) (Fig. 2C, Supplemental Table 2). PEN19 was less potent (777±2 nM) and less efficacious (111±4%) than PEN22 while PEN18 could not induce increases in [35S]GTPγS binding (Fig. 2C, Supplemental Table 2). Since the GTPγS binding assay primarily detects Gαi proteins (Harrison and Traynor, 2003), these results do not rule out the possibility that PEN18 and PEN19 could signal via other Gα proteins. To
investigate whether C-terminally truncated PEN peptides signal through $G_{\alpha s}$ or $G_{\alpha i}$, we used the adenylyl cyclase activity assay. C-terminally truncated PEN peptides modulated adenylyl cyclase activity in CHO-GPR83 but not CHO membranes (Supplemental Fig. 1C). Treatment of CHO-GPR83 cell membranes with PEN22 or PEN20 led to a dose-dependent decrease in adenylyl cyclase activity and this could be blocked by pertussis toxin (Fig. 2D, E, Supplemental Fig. 1, Supplemental Table 2). Interestingly, PEN18 or PEN19 treatment led to a dose-dependent increase in adenylyl cyclase activity that could be substantially attenuated by cholera toxin treatment (Fig. 2D, E, Supplemental Fig. 1, Supplemental Table 2). Taken together, these results are consistent with the notion that PEN22 and PEN20 facilitate $G_{\alpha i}$-mediated signaling while PEN18 and PEN19 facilitate $G_{\alpha s}$-mediated signaling by engaging GPR83, suggesting biased signaling by endogenous C-terminally truncated PEN peptides.

PEN22 was previously shown to exhibit both $G_{\alpha q}$ and $G_{\alpha i}$-mediated signaling at GPR83 (Gomes et al., 2016). We therefore examined the ability of C-terminally truncated PEN peptides to signal via $G_{\alpha q}$ proteins using the phospholipase C (PLC) assay, and also by measuring their ability to increase intracellular Ca$^{2+}$ levels. C-terminally truncated PEN peptides increase PLC activity in CHO-GPR83 but not CHO membranes (Supplemental Fig. 1G) with sub-nM to nM potency (Fig. 2F, Supplemental Table 2). However they differ in their efficacy in CHO-GPR83 membranes with PEN22 being the most efficacious and PEN18 and PEN19 the least efficacious (Fig. 2F, Supplemental Table 2). C-terminally truncated PEN peptides increase intracellular Ca$^{2+}$ levels in CHO-GPR83 but not CHO cells (Supplemental Fig. 1H). In CHO-GPR83 cells, treatment with PEN19 or PEN20 increased intracellular Ca$^{2+}$ levels although with much lesser efficacy than PEN22, while PEN18 treatment caused an increase only at the highest concentration tested (10 $\mu$M) (Fig. 2G, Supplemental Table 2). These results are consistent with the notion that C-
terminally truncated PEN peptides can also signal via $G_{\alpha q}$ proteins thereby suggesting the promiscuity of $G_{\alpha}$ protein subtype coupling by GPR83.

Exposure to agonists often leads to endocytosis of G protein-coupled receptors (GPCRs) which can modulate cellular responses (Lobingier and von Zastrow, 2019). Therefore, we looked at the ability of C-terminal truncated PEN peptides to induce GPR83 endocytosis. These peptides cause a dose and time-dependent endocytosis of GPR83 although there are differences in the extent of receptor endocytosed as compared to PEN22 (Fig. 2H, I, Supplemental Table 2). At a highest concentration (10 μM) PEN18 induces ~32%, PEN19 and PEN20 ~51% while PEN22 induces ~70% receptor endocytosis (Fig. 2H, Supplemental Table 2). Even after prolonged exposure to these peptides (2 h) we find that PEN18 is the least effective at inducing GPR83 endocytosis (Fig. 2I; Supplemental Table 2). Taken together, these results show that C-terminally truncated PEN peptides exhibit differential signaling and endocytosis of GPR83.

Binding of C-terminally truncated PEN peptides to striatal GPR83 and characterization of the signaling pathways.

First we examined $[^{125\text{I}}]$Tyr-rPEN binding WT and GPR83 KO striatal membranes and found robust binding WT but not GPR83 KO membranes (Supplemental Fig. 1I). Next, we examined if C-terminally truncated PEN peptides could bind to and signal through native GPR83 in the striatum. As found with the cell line studies described above, all of the PEN peptides exhibit biphasic displacement binding curves with high-affinity binding in the sub-nM range and low-affinity binding in the high nM-μM range (Fig. 3A, Supplemental Table 1). C-terminally truncated PEN peptides in striatal membranes induce dose-dependent increases in $[^{35\text{S}}]$GTPγS binding (Fig. 3B, Supplemental Table 2). Like PEN22, the peptides exhibit nM potencies but differ in their efficacies with PEN19 and PEN20 being least efficacious (Fig. 3B, Supplemental Table 2). This
signaling appears to be mediated through GPR83 based on the finding that peptide-mediated increases in $[^{35}\text{S}]\text{GTP}\gamma\text{S}$ binding are not seen in GPR83 KO striatal membranes (Fig. 3C). In the adenylyl cyclase activity assays, PEN18 and PEN19 cause modest increases in activity while treatment with PEN20 and PEN22 cause robust increases (Fig. 3D, Supplemental Table 2). These increases in activity were abrogated in GPR83 KO striatal membranes (Fig. 3E). In the phospholipase C activity assay the peptides cause dose-dependent increases in activity (Fig. 3F, Supplemental Table 2) with more robust effects being observed with PEN20 and PEN22 compared to PEN18 and PEN19 (Fig. 3F, Supplemental Table 2); these increases in activity were not seen in GPR83 KO striatal membranes (Supplemental Fig. 1J). Together these results show that C-terminal truncated PEN peptides can bind to and signal through GPR83 in the striatum. Moreover the peptides can elicit $G_\alpha_i$, $G_\alpha_s$, and $G_\alpha_q$-mediated signaling in this brain region.

**Binding of procholecystokinin-derived peptides to GPR83 and characterization of the signaling pathways**

Based on the finding that PEN20 and other related peptides bound to and signaled through GPR83, it was of interest to test the proCCK peptide that shares the C-terminal sequence with PEN20 (i.e. CCK56-62) (Fig. 1). In addition, another frequently-detected peptide that contains one additional C-terminal residue was also examined (i.e. CCK56-63) (Fig.1). Synthetic peptides representing the C-terminal region of two PEN peptides (i.e. PEN11-20 and PEN11-22) were included in the analysis for comparison (Fig. 4A). Both of the proCCK peptides as well as the shorter PEN peptides displaced $[^{125}\text{I}]\text{Tyr-PEN}$ binding to CHO-GPR83 cell membranes (Fig. 4B, Supplemental Table 3) and striatal membranes (Fig. 4C, Supplemental Table 3), showing biphasic displacement profiles with high affinity binding in the sub-nM range and low affinity binding in the high nM range. All of these peptides also induce robust increases in $[^{35}\text{S}]\text{GTP}\gamma\text{S}$ binding in
WT striatal membranes, although they are less potent and less efficacious than PEN22 (Fig. 4D, Supplemental Table 4). The increases in [35S]GTPγS binding caused by treatment with CCK56-62 and CCK56-63 are substantially reduced in GPR83 KO striatal membranes (Supplemental Fig. 2A). Together, these results support the idea that the ‘LGALL’ motif common to all of the peptides is responsible for binding to GPR83.

Next, we examined if PEN11-20, PEN11-22, CCK56-62 & CCK56-63 exhibit differences in downstream signaling. In the adenylyl cyclase activity assay, PEN11-20 and PEN11-22 exhibit increases in activity in CHO-GPR83 but not CHO membranes (Supplemental Fig. 2B-D, Supplemental Table 4); these increases are blocked by pre-treatment with cholera toxin but not pertussis toxin (Fig. 4E, Supplemental Fig. 2C & D), indicating that these peptides activate Gαs-mediated signaling. In contrast, CCK56-62, and CCK56-63 exhibit a bell-shaped profile of adenylyl cyclase activity with a peak in the sub-nM to nM range (Fig. 4G, Supplemental Fig. 2, Supplemental Table 4); the peptides have no effects on adenylyl cyclase activity in CHO cell membranes (Fig. 4G). Pretreatment with cholera toxin attenuates CCK56-62 or CCK56-63-mediated adenylyl cyclase activity seen at low doses (0.1 to 10 nM), while pertussis toxin treatment attenuates the activity seen at higher doses of the peptides, and a combination of both toxins completely abrogates signaling by both peptides (Fig. 4H, I, Supplemental Fig. 2). Moreover, PEN11-20, PEN11-22, CCK56-62, and CCK56-63 induce GPR83 endocytosis with a potency in the nM range (Fig. 4F, Supplemental Table 4). Together, these results indicate that the proCCK peptides that share sequence identity with PEN can bind to and differentially signal at GPR83 in a dose-dependent manner. Furthermore, PEN binding and signaling through GPR83 is largely mediated by the C-terminal portion of the peptide and that GPR83 is activated by peptides from
proCCK (in addition to proSAAS) and this leads to differential signaling allowing for fine-tuning of GPR83 signaling.
DISCUSSION

Our findings expand the repertoire of endogenous GPR83 ligands and highlight the complexities of this neuropeptide-receptor signaling system. One major finding is that C-terminal truncated PEN-derived peptides can bind and signal through GPR83. These truncated peptides appear to be major forms of PEN in brain and other tissues (Fricker, 2010; Mzhavia et al., 2001; Mzhavia et al., 2002). Rather than representing inactive degradation fragments, we find that the truncated peptides exhibit differences in their abilities to activate downstream signaling pathways, a phenomenon commonly referred to as “biased agonism”. There are several other examples of biased GPCR signaling arising from small differences in a peptide’s length. For example, binding of angiotensin II to angiotensin receptors triggers G protein-mediated signaling while the binding of a peptide that is one amino acid shorter (Ang1-7) leads to β-arrestin mediated signaling (Teixeira et al., 2017). Other examples of this phenomenon are found with opioid peptides, where pentapeptide enkephalins and C-terminally-extended enkephalins show differences in G protein versus beta-arrestin signaling at mu, delta, and kappa opioid receptors (Gomes et al., 2020). The present findings that various PEN peptides show biased signaling through GPR83 extends this phenomenon, and implies that the peptidases which convert PEN22 into shorter peptides produce a subtle change in the signaling profiles.

Another major finding of the present study is that proCCK derived peptides bind and signal via GPR83 although this does not rule out that the peptides could also signal via an as yet unidentified receptor since residual signaling is detected in GPR83 KO striatal membranes (Supplemental Fig. 2A). At GPR83 these peptides activate $G_{\alpha_s}$-mediated signaling at low concentrations ($\leq 1$ nM) and $G_{\alpha_i}$-mediated signaling at higher concentrations. This supports our observations that GPR83 can signal via $G_{\alpha_s}$, $G_{\alpha_i}$ and $G_{\alpha_q}$ proteins (Gomes et al., 2016), and extends
the dynamics of signaling by endogenous peptides. It has been reported that neuropeptide Y and other agonists of the type 2 neuropeptide Y receptor bind to GPR83 (Sah et al., 2007). However, another study found that high concentrations of neuropeptide Y caused only a small displacement in $[^{125}\text{I}]$Tyr-PEN binding to human GPR83, suggesting that the binding site for the two peptides may be distinct (Gomes et al., 2016). A recent study reported that FAM237A and to a lesser extent FAM237B peptides induce $\beta$-arrestin recruitment and inositol phosphate recruitment via GPR83 although the authors were not able to detect consistent changes in cAMP levels (Sallee et al., 2020). Finally, the basal activity of GPR83 has been reported to be modulated by the N-terminal domain of the receptor and by Zn$^{+2}$ (Muller et al., 2016; Muller et al., 2013a). Together these findings suggest that the activity of GPR83 can be regulated by different modalities. However, only the activation of GPR83 by PEN has been replicated by independent investigators (Foster et al., 2019; Nataliya Parobchak; et al., 2020), which is a criterion for ‘de-orphanizing’ a receptor. The other reported ligands (including the proCCK peptides in the present study) await confirmation.

If CCK56-62 and CCK56-63 serve as endogenous GPR83 ligands, these peptides should be secreted in the vicinity of cells expressing GPR83. High levels of proCCK mRNA are expressed in cortex, hippocampus, thalamus, and several other regions. In contrast, GPR83 mRNA is highly expressed in nucleus accumbens, caudoputamen, and olfactory tubercle as well as specific nuclei in other regions (Fig. 5). Because neuropeptides can be secreted from nerve terminals located far from the cell bodies where their mRNA is expressed, it is important to consider neuronal projections. GPR83 is expressed on interneurons in the striatum that receive direct inputs from various regions where proCCK is expressed (Enterria-Morales et al., 2020; Fakira et al., 2019)(Klug et al., 2018). Cortico-striatal projections directly release proCCK peptides into the striatum (Morino et al., 1994; Schiffmann and Vanderhaeghen, 1991; You et al., 1994). ProCCK
mRNA is also expressed in thalamostriatal projections (Burgunder and Young, 1988) providing an additional source of proCCK peptides in the striatum. Neurons in the ventral tegmental area (VTA) and substantia nigra pars compacta (SNpc) have also been shown to release CCK into the striatum and olfactory tubercle (Hokfelt et al., 1980b; Zaborszky et al., 1985). In addition, it is possible that GPR83 expressed in the cortex is activated by proCCK peptides released from cortical interneurons (Adams et al., 2003; Brezillon et al., 2001; Lueptow et al., 2018; Nguyen et al., 2020; Pesini et al., 1998; Wang et al., 2001; Whissell et al., 2015). Taken together, it is likely that proCCK56-62 and related peptides are secreted in the proximity of GPR83.

The brain regions expressing high levels of proCCK mRNA are distinct from those expressing high levels of proSAAS (Fig. 5). Little is known regarding projections of proSAAS-expressing neurons, but due to the broad expression of low levels of proSAAS throughout the brain, it is likely that GPR83 is exposed to secreted PEN and related peptides. In brain, GPR83 is likely to integrate signals from a broad range of brain areas, with cortical-striatal projections releasing proCCK peptides, and projections from other regions releasing PEN. Recently, GPR83 was reported to be expressed on neurons that project from spinal cord to lateral parabrachial nucleus of the pons, a region that represents a major ascending spinal pathway for processing of tactile and noxious stimuli (Choi et al., 2020). Both proSAAS and proCCK are abundantly expressed in spinal cord, and thus both PEN and proCCK peptides are potential ligands for GPR83 in this tissue (Chakraborty et al., 2006; Feng et al., 2001; Larsson and Rehfeld, 1979; Royds et al., 2020).

The vast majority of previous reports on proCCK-derived peptides have focused on CCK-containing peptides, which includes a number of N-terminally-extended peptides (Fig. 1). The peptide CCK56-63 is cleaved from proCCK at single basic cleavage sites Arg55 and Arg64.
Cleavage of proCCK at Arg 64 generates a long form of CCK designated CCK39 (Rehfeld et al., 2007). While the Arg55 cleavage site on the N-terminus of CCK56-63 does not appear to have been previously reported in the CCK field, another site further upstream (Arg45) produces the long CCK form named CCK58 (Rehfeld et al., 2007). Interestingly, intraperitoneal injections of synthetic CCK58 or CCK8 produced comparable reductions of meal size, but CCK58 increased the satiety ratio and this effect was not seen with CCK8 (Overduin et al., 2014; Sayegh et al., 2014). If CCK58 activates GPR83, this could potentially account for the additional effects seen with the longer peptide. Although this was not directly tested in the present study, our finding that CCK56-62 and CCK56-63 produce similar effects indicate that the LGALL motif does not need to be at the extreme C-terminus of the peptide in order to activate GPR83, raising the possibility that CCK58 also activates this receptor. If this prediction is confirmed, it could potentially explain the different biological effects of CCK8 and CCK58 observed by Reeve and colleagues (Overduin et al., 2014; Sayegh et al., 2014).

Some of the peptidases that generate the various peptides described in the present study can be predicted based on previous peptidomic studies on mice lacking specific peptide-processing enzymes. Cleavage at Arg45 and Arg64 produces the peptide CCK46-63, which was detected many times in mouse brain peptidomics studies (Fricker, 2010). This peptide, as well as the shorter CCK56-63 used in the present study are both produced in the late secretory pathway by CPE, based on peptidomics analyses of Cpe null mice (Lim et al., 2006; Zhang et al., 2008). CCK56-63 is produced from CCK46-63 by prohormone convertase 2, based on the changes observed in Pcks2 null mice (Pan et al., 2006; Zhang et al., 2010). C-terminally truncated forms were detected many times in mouse brain, including CCK46-62 (ending with LGALL), as well as shorter C-terminally truncated forms that were not tested in the present study. Similarly, PEN22 and shorter forms have
been frequently detected in mouse brain and in proSAAS-expressing cell lines (Fricker, 2010; Mzhavia et al., 2001; Mzhavia et al., 2002). PEN22 is produced from proSAAS mainly via furin acting at the consensus sites RxRR on either side of PEN22, followed by removal of the C-terminal R residues by carboxypeptidase D. In contrast, PEN20 is produced later in the secretory pathway and requires CPE for its formation, based on the absence of this peptide in Cpe null mice (Lim et al., 2006; Zhang et al., 2008). It is not clear if the C-terminal trimming of PEN20 and CCK56-63 into shorter peptides occurs within secretory granules or by extracellular enzymes following secretion of the peptides.

Taken together, this study sheds light on the complexity of the GPR83 signaling system, and raises the exciting and intriguing possibility of exploiting the potential biased signaling at GPR83 to develop therapeutics targeting this receptor for the treatment of obesity, addiction and reward-related disorders.
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Footnotes:

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Legends for Figures

Fig. 1. Schematic representation of peptides generated from the processing of proSAAS and proCCK.

Processing of the proSAAS precursor at monobasic and dibasic sites yields Big SAAS, Big GAV, KEP, Little SAAS, Little GAV, PEN, Big LEN and Little LEN. Mass spectrometric analysis detects the presence of PEN (i.e. PEN22) as well as shorter PEN peptides (PEN18, PEN19, PEN20) (Fricker, 2010; Mzhavia et al., 2002). Processing of proCCK yields CCK8, CCK12, CCK33, CCK39, CCK58. In addition, mass spectrometric analyses detects the presence of CCK peptides containing the highly-conserved LGALL motif that is present in PEN (Fricker, 2010; Mzhavia et al., 2002).

Fig. 2. Binding and signaling by C-terminally truncated PEN peptides at CHO-GPR83.

(A) Sequence of full-length PEN (PEN22) and of the C-terminally truncated peptides, PEN18, PEN19 and PEN20. (B) C-terminally truncated PEN peptides displace $[^{125}\text{I}]-\text{Tyr-rPEN}$ binding (3 nM) to membranes (20 μg) from CHO cells expressing GPR83 (CHO-GPR83). Data represent Mean±SD (n=3 for PEN18, PEN19 and PEN20 and n=6 for PEN22) (C) C-terminally truncated PEN peptide-mediated increases in $[^{35}\text{S}]\text{GTP}_S$ binding to CHO-GPR83 membranes (20 μg). Data represent Mean±SD (n=6). (D) Modulation of adenylyl cyclase activity by C-terminally truncated PEN peptides in CHO-GPR83 membranes (2 μg). Data represent Mean±SD (n=3). (E) Effect of pertussis toxin (PTX) and cholera toxin (CTX) treatment on PEN18-, PEN19- , PEN20- and PEN22-mediated inhibition of adenylyl cyclase activity. Data represent Mean±SD (n=3). *p<0.05; **p<0.01; ***p<0.001; n.s. = not significant (one-way ANOVA with Tukey’s multiple comparison). (F) C-terminally truncated PEN peptide-mediated changes in phospholipase C activity in CHO-GPR83 cell membranes. Data represent Mean±SD (n=6). (G) C-terminally
truncated PEN peptide-mediated increases in intracellular Ca^{2+} levels in CHO-GPR83 cells. Data represent Mean±SD (n=3). (H) Dose-dependent modulation of GPR83 endocytosis by C-terminally truncated PEN peptides. (I) Time course of GPR83 internalization by C-terminally truncated PEN peptides. Data (H-I) represent mean±SD (n=3).

**Fig. 3. Binding and signaling by C-terminally truncated PEN peptides at striatal membranes.**

(A) C-terminally truncated PEN peptides displace [^{125}I]-Tyr-PEN binding (3 nM) to striatal membranes (20 μg). Data represent Mean±SD (n=3). (B-C) C-terminally truncated PEN peptide-mediated increases in [^{35}S]GTPγS binding to striatal membranes (20 μg) from wild-type (WT) (B,C) and GPR83 knockout (GPR83 KO) (C) mice. Data represent Mean±SD (n=3). In (C) *p<0.05; **p<0.01; WT vs GPR83 KO; Unpaired t-test with Welch’s correction. (D-E) Modulation of adenylyl cyclase activity by C-terminally truncated PEN peptides in striatal membranes (2 μg) from WT (D,E) and GPR83 k/o (E) mice. Data represent Mean±SD (n=3). In (E) *p<0.05; **p<0.01 WT vs GPR83 k/o; Unpaired t-test with Welch’s correction. (F) C-terminally truncated PEN peptide-mediated changes in phospholipase C activity in striatal membranes (10 μg). Data represent mean±SD (n=3).

**Fig. 4. Binding and signaling by proCCK-derived peptides at CHO-GPR83.**

(A) Sequence of proCCK derived peptides that exhibit similarity to peptides derived from PEN 22 (PEN11-20 and PEN11-22). (B-C) ProCCK-derived peptides and related PEN peptides displace [^{125}I]-Tyr-PEN binding (3 nM) to membranes (20 μg) from CHO-GPR83 (B) and WT striatum (C). Data represent mean ±SD (n=6). (D) ProCCK-derived peptide- and related PEN peptide-mediated increases in [^{35}S]GTP γS binding to striatal membranes (20 μg). Data represent mean ±SD (n=6). (E) PEN11-20 and PEN11-22-mediated increase in adenylyl cyclase activity in
CHO-GPR83 cell membranes is blocked by CTX but not by PTX. Data represent mean ±SD (n=5 for PEN11-20 and PEN11-22 and n=3 for PEN22). **p< 0.01; ***p< 0.001 (One-Way ANOVA with Tukey’s multiple comparison test. (F) GPR83 endocytosis by proCCK-derived peptides and related PEN peptides in CHO-GPR83 cells. Data represent mean ±SD (n=6) (G) Modulation of adenylyl cyclase activity by proCCK-derived peptides in CHO and in CHO-GPR83 membranes (2 μg). Data represent mean ±SD (n=6). (H-I) A combination of CTX and PTX is needed to block CCK56-63-mediated changes in adenylyl cyclase activity in CHO-GPR83 (H) and striatal (I) membranes. Data represent mean±SD (n=4).

**Fig. 5.** Characterization of GPR83, proSAAS, and proCCK brain distribution in mice. The data for the mRNA expression was obtained from Allen Brain Mouse Atlas (https://mouse.brain-map.org/experiment), which shows in situ hybridization of sagittal mouse brain sections probed with GPR83 (A), proCCK (B), and proSAAS (gene name Pcsk1n) (C). Bar graphs represent the average raw expression value for different brain regions; each dot represents a biological replicate. OLF, olfactory area; HPF, hippocampal formation; CTXsp, cortical subplate; STR, striatum; PAL, pallidum; TH, thalamus; HY, hypothalamus; MB, midbrain; P, pons; MY, medulla; CB, cerebellum. D. Diagram showing potential projections of proCCK and proSAAS derived-neuropeptide release into brain regions with high levels of GPR83 expression. Projections of proCCK neurons and the expression of proSAAS-derived peptides was obtained from previous studies (Adams et al., 2003; Brezillon et al., 2001; Burgunder and Young, 1988; Lueptow et al., 2018; Morino et al., 1994; Nguyen et al., 2020; Pesini et al., 1998; Schiffmann and Vanderhaeghen, 1991; Wang et al., 2001; Whissell et al., 2015; You et al., 1994). Solid pink arrows represent the proCCK neuronal projections to striatum and olfactory tubercle (orange) both of which express
GPR83 mRNA. Dashed green lines represent projections to the striatum and olfactory tubercle that express proSAAS mRNA and these are potential sources of PEN in the GPR83-rich brain regions.
Peptide Sequence
PEN22  SVDQDLGPEVPPENVLGALLRV
PEN20  SVDQDLGPEVPPENVLGALL
PEN19  SVDQDLGPEVPPENVLGAL
PEN18  SVDQDLGPEVPPENVLGA

AC activity (% basal)

Ligand [M]
PEN22 PEN20 PEN19

AC activity (% basal)

Ligand [M]
PEN18 PEN19 PEN20

AC activity (% basal)

Ligand [M]
PEN18 PEN19 PEN20

AC activity (% basal)

Ligand [M]
PEN18 PEN19 PEN20

PLC levels (% basal)

Ligand [M]
PEN18 PEN19 PEN20

Calcium levels (% basal)

Ligand [M]
PEN18 PEN19 PEN20

% Cell surface receptors

Ligand [M]
PEN18 PEN20 PEN22

% Cell surface receptors

Ligand [M]
PEN18 PEN19 PEN20

% Cell surface receptors

Ligand [M]
PEN18 PEN20 PEN22

Fig. 2

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Fig. 3
Fig. 5