RESEARCH PAPER

Pancreatic polypeptide and its central Y4 receptors are essential for cued fear extinction and permanent suppression of fear

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BACKGROUND AND PURPOSE
Avoiding danger and finding food are closely related behaviours that are essential for surviving in a natural environment. Growing evidence supports an important role of gut-brain peptides in modulating energy homeostasis and emotional-affective behaviour. For instance, postprandial release of pancreatic polypeptide (PP) reduced food intake and altered stress-induced motor activity and anxiety by activating central Y4 receptors.

EXPERIMENTAL APPROACH
We characterized [K30(PEG2)]hPP2-36 as long-acting Y4 receptor agonist and injected it peripherally into wildtype and Y4 receptor knockout (Y4KO) CS7Bl/6NCrl mice to investigate the role of Y4 receptors in fear conditioning. Extinction and relapse after extinction was measured by spontaneous recovery and renewal.

KEY RESULTS
The Y4KO mice showed impaired cued and context fear extinction without affecting acquisition, consolidation or recall of fear. Correspondingly, peripheral injection of [K30(PEG2)]hPP2-36 facilitated extinction learning upon fasting, an effect that was long-lasting and generalized. Furthermore, peripherally applied [K30(PEG2)]hPP2-36 before extinction inhibited the activation of orexin-expressing neurons in the lateral hypothalamus in WT, but not in Y4KO mice.

CONCLUSIONS AND IMPLICATIONS
Our findings suggests suppression of excessive arousal as a possible mechanism for the extinction-promoting effect of central Y4 receptors and provide strong evidence that fear extinction requires integration of vegetative stimuli with cortical and subcortical information, a process crucially depending on Y4 receptors. Importantly, in the lateral hypothalamus two peptide systems, PP and orexin, interact to generate an emotional response adapted to the current homeostatic state. Detailed investigations of feeding-relevant genes may thus deliver multiple intervention points for treating anxiety-related disorders.

Abbreviations
CS, conditioned stimulus; CS+, conditioned stimulus that was paired with an unconditioned stimulus; CS−, conditioned stimulus that was not paired with an unconditioned stimulus; NPY, neuropeptide Y; NTS, nucleus tractus solitarii; PP, pancreatic polypeptide; US, unconditioned stimulus
**Introduction**

Avoiding danger and finding food are two intimately associated, life-sustaining behaviours that are strongly modulated by emotions (Sakurai and Mieda, 2011). Maladaptation within such survival circuits can induce dysregulated, pathological behaviour, resulting in the development of feeding or anxiety disorders (Myers and Davis, 2007; Finger et al., 2010; Steransson, 2013; Steransson et al., 2013). In the USA and Europe, an estimated number of 100 million people are suffering from anxiety-related diseases (Wittchen et al., 2011; Kessler et al., 2012). Thus, identifying critical components of the underlying circuitry may provide new treatment strategies. Interestingly, neuropeptides are essential modulators of both energy homeostasis and anxiety-related behaviours (Hokfelt et al., 2003; Coll et al., 2007; Bowers et al., 2012; Schellekens et al., 2012). For instance, gut-brain peptides, such as neuropeptide Y (NPY), are released during states of hunger or acute danger (Bowers et al., 2012; Holzer et al., 2012). While the anxiolytic and fear-reducing properties of NPY are increasingly evident (Heilig, 1995; Karlsson et al., 2008; Tasan et al., 2011; Verma et al., 2012), the role of pancreatic polypeptide (PP) and its Y$_4$ receptor in fear-related behaviour is still poorly understood.

PP, which preferentially activates Y$_4$ receptors, is synthesized mainly in endocrine cells of the pancreas and released into the blood circulation by a cholinergic, vagus-dependent mechanism predominantly regulating gastrointestinal processes and appetite (Lin et al., 2009; Holzer et al., 2012; Shi et al., 2013). In the CNS, Y$_4$ receptors have a very distinct distribution, with particular high expression in the hypothalamus and the nucleus tractus solitarii (NTS), two brain areas that are accessible to peripherally released PP and important for mediating emotionally driven autonomic responses (Dumont et al., 2007; Tasan et al., 2009). Thus, PP may be considered as a signalling peptide communicating the internal metabolic state to respective CNS centres that generate an adaptive behavioural response. This may include satiety signals but also accompanying emotional behaviour, such as altered anxiety and fear processing.

Recently, we demonstrated that genetic deletion of Y$_4$ receptors (Y4KO) results in increased novelty-induced motor activity, reduced anxiety and improved stress coping (Painsipp et al., 2008; Tasan et al., 2009; Painsipp et al., 2010). Overexpression of the endogenous Y$_4$ receptor agonist PP, on the other hand, resulted in reduced anxiety-like behaviours. Interestingly, chronic peripheral application of PP reduced anxiety-like behaviours, while i.c.v. injection of PP had no effect (Asakawa et al., 1999; Asakawa et al., 2003), suggesting that only peripherally released PP may properly interact with central Y$_4$ receptors.

Fear and anxiety-related behaviours are predominantly controlled by the amygdala complex and can be tested by Pavlovian fear conditioning, a simple form of associative learning (LeDoux, 2000). In the fear conditioning paradigm, an unconditioned stimulus (US), usually a mild electric foot shock, is repetitively paired with a conditioned stimulus (CS), typically represented by an auditory stimulus. After a few pairings, the CS alone can elicit a typical fear reaction, termed conditioned response (CR). Subsequently, repetitive presentations of the CS in the absence of the US result in a gradual decrease of the learned fear response, a process called fear extinction. Recent evidence suggests that central NPY inhibits the expression and promotes the extinction of learned fear (Gutman et al., 2008; Fendt et al., 2009; Verma et al., 2012). Furthermore, we demonstrated that short-term fasting impairs fear consolidation but facilitates fear extinction by increasing feedforward inhibition in an amygdala microcircuit (Verma et al., 2016). This feedforward inhibition was reduced in Y4KO mice resulting in impaired fear extinction. These data indicate an important role of gut-derived peptides in emotional control, thus linking life-sustaining circuitries for fear and hunger.

Here, we have characterized the role of peripheral PP in acquisition, recall/expression and extinction of conditioned fear and investigated the particular contribution of Y$_4$ receptors in these processes. We decided to stimulate Y$_4$ receptors by peripheral injection of a novel derivative of PP, the PEGylated peptide [K$_{30}$(PEG2)]hPP$_{2-36}$, which we have characterised as a selective agonist of Y$_4$ receptors. While both lipidation and PEGylation of PP prolong plasma half-life, PEGylation also inhibits arrestin recruitment and receptor internalization, two effects that are independent of the PEGylated amino acid residue (Mäde et al., 2014) and may significantly prolong the activation of Y$_4$ receptors. Our results showed that [K$_{30}$(PEG2)]hPP$_{2-36}$ acting on central Y$_4$ receptors promoted the cued fear extinction probably by inhibiting orexin-expressing neurons in the lateral hypothalamus, resulting in a stable, long-term suppression of fear.
Methods

Synthesis and characterization of [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36}

Peptide synthesis. The peptide was synthesized as described previously (Bellmann-Sickert et al., 2011) using fluorenlymethoxycarbonyl-/tert-butyl (Fmoc/tBu) protecting group strategy on a Rink amide resin (Iris Biotech, Marktriedt, Germany; loading capacity: 0.7 mmol·g\textsuperscript{-1}, mesh size 100–200). The peptide sequence was elongated automatically by a fully automated parallel peptide synthesizer (Syrto II, Biotage, Uppsala, Sweden). Amino acids, N,N-diisopropylcarbodiimide (DIC) and oxyma were dissolved in N,N-dimethylformamide (DMF) to a final concentration of 0.5 M and applied with an 8-fold excess. Double couplings of 2 × 30 min were performed per cycle. Fmoc deprotection was carried out with 40% piperidine in DMF for 3 min followed by 20% piperidine for 10 min. Lysine at position 30 was introduced with (4-methoxyphenyl)diphenylmethyl (Mmt) side chain protection. N-terminal proline was either introduced as tert-butoxycarbonyl (Boc) protected or was labelled with 5-carboxytetramethylrhodamine (TAMRA) using 2 equivalents TAMRA, 2 equivalents O-(7-azabenzotriazol-1-yl)-N,N,N',N’tetramethyluronium-hexafluoroarsenate (HATU) and 1.9 equivalents N,N-diisopropylethylamine dissolved to 0.5 M in DMF and incubation for 2 h at room temperature. After complete synthesis of the peptide backbone, the Mmt group was cleaved off by repeatedly (15 times) applying 1 mL of a solution of 2% (v/v) trifluoracetic acid (TFA) and 5% (v/v) triisopropsilanes (TIS) in dichloromethane for 2 min and washing with dichloromethane. For FEGylation, 2 equivalents of α-methoxy-o-carboxylic acid succinimidyl ester poly(ethylene glycol) (MeO-PEG-NHS, 2000 Da; Iris Biotech) were dissolved in DMF/dichloromethane (2:1 (v/v)) to a concentration of 0.06 M. 4-Dimethylpiperidine and DIC were added to a concentration of 0.1 M. The mixture was added to the resin and shaken for 16 h at room temperature and washed subsequently with DMF and dichloromethane. Peptide was cleaved from the resin with a mixture of TFA, 1,2-ethanedithiol and para-thioninolise (18:1:1, v/v/v) for 3 h at room temperature and precipitated from ice-cold diethylther. Methionines were reduced by applying a mixture of TFA, trimethylsilylformic acid and 1,2-ethanedithiol (100:1:6:1.2, v/v/v) for 20 min at room temperature and precipitation from ice-cold diethylther. Precipitates were washed with ice-cold diethylther, dried under vacuum and dissolved in a mixture of water and tert-butanol (3:1, v/v). Products were purified and analysed by reversed phase high performance liquid chromatography using a gradient consisting of 0.1% TFA in water (eluent A) and 0.08% TFA in acetonitrile (eluent B) ranging from 20% to 60% B in A for 40 min. Purification was done by using a Jupiter Proteo 90 Å with 10 μm pore size and dimensions of 21.2 × 250 mm (Phenomenex, Aschaffenburg, Germany). Peptide purity was confirmed to be >95%. Peptide identity was proven by matrix-assisted laser desorption/ionization mass spectrometry using an Ultraflex III matrix-assisted laser desorption/ionization time-of-flight/time-of-flight (MALDI-TOF/TOF) mass spectrometer (Bruker, Bremen, Germany). The MS results are shown in Supporting Information Figure S1.

Characterisation of activity at Y receptors. Agonist activity at Y receptors was assessed by measuring inositol phosphate accumulation in COS-7 cells, as described previously (Hofmann et al., 2013). We used COS-7 cells that stably expressed one of the four human Y receptors (hYR) and a chimeric G\textsubscript{i}o protein (Kostenis et al., 1997) to channel cAMP inhibition to phospholipase C stimulation and hence inositol 1,4,5-triphosphate (IP\textsubscript{3}) production. Stably transfected COS-7-hYR-GluA6q4myr cells were cultivated in DMEM with 4.5 g·L\textsuperscript{-1} glucose and L-glutamine supplemented with 10% (v/v) heat-inactivated FCS, 100 units mL\textsuperscript{-1} penicillin, 100 μg mL\textsuperscript{-1} streptomycin, 1.5 mg mL\textsuperscript{-1} G418-sulfate and 146 μg mL\textsuperscript{-1} hygromycin B in a humidified atmosphere at 37°C and 5% CO\textsubscript{2}. COS 7 cells were a gift from Prof. Dr. Torsten Schönberg (Leipzig University). Stable cell lines were prepared as described in Mäde et al., 2014. Cells were grown in 48-well plates (90,000 cells in 500 μL per well) for 24 h. Cells were labelled with 2 μCi mL\textsuperscript{-1} myo-[2-3H]-inositol (PerkinElmer, Rodgau, Germany) in culture medium without penicillin and streptomycin for at least 16 h followed by stimulation for 1 h with peptide solutions (10⁻⁵–10⁻¹¹ M, depending on ligand potency) in DMEM with 4.5 g·L\textsuperscript{-1} glucose and L-glutamine containing 10 mM Lici (Sigma-Aldrich). Cells were lyes applying 0.1 M sodium hydroxide for 5 min at room temperature followed by neutralization with 0.2 M formic acid. Samples were diluted in dilution buffer (5 mM Na-borate and 0.5 mM Na-EDTA). Cell debris was removed. Samples were loaded on columns packed with the anion exchange resin AG 1-X8 formate (BIO-RAD, Munich, Germany), washed with glycerolphosphate elution buffer (5 mM Na-borate, 60 mM Na-formate) and water and eluted with elution buffer (1.0 M ammonium formate, 0.1 M formic acid). Eluates were diluted with scintillation cocktail Ultima Gold (Perkin Elmer) and measured by a scintillation counter (Tri-Carb 2910 TR from PerkinElmer). The values for decay per minute obtained at the highest used wild type concentration (corresponds to 100% activity) were normalized to a mean value to which all other agonists were related. For each compound, global regression using GraphPad Prism 5.0. All compounds were examined in duplicate, in at least two independent experiments.

Metabolic stability in human blood plasma. Proteolytic stability in human blood plasma was assessed as described previously (Mäde et al., 2014) with minor adaptations. 15 nmol of the N-terminally TAMRA-labelled compound was dissolved in 10 μL of distilled water and 1490 μL of human blood plasma (citrate stabilized). The mixture was incubated for 96 h at 37°C. At the indicated time points, samples (150 μL) were withdrawn and equal volumes of ethanol and acetonitrile were added in order to precipitate plasma proteins. Samples were incubated at −20°C overnight and centrifuged at 12,000 g for 30 s. Supernatants were again incubated at −20°C for 20 min and afterwards filtered using SpinX tubes (0.22 μm, Costar). Filtrates were directly subjected to reverse phase high performance liquid chromatography (RP-HPLC) using a Varitide RPC column (200 Å, 6 μm pore size, 4.6 × 250 mm) and a gradient of 20 to 60% B in A for 30 min. Intact peptide was followed by fluorescence detection at an excitation wavelength of 525 nm and an emission wavelength of 572 nm. Degradation of each compound was independently performed twice (n = 2) and is presented as mean ± SEM (Supporting Information Figure S3).
Animals

All animal care and experimental procedures were conducted in accordance with international laws and policies (Directive 2010/63/EU of the European parliament and of the council of 22 September 2010 on the protection of animals used for scientific purposes; Guide for the Care and Use of Laboratory Animals, US National Research Council, 2011) and were approved by the Austrian Ministry of Science. Animal studies are reported in compliance with the ARRIVE guidelines (Kilkenny et al., 2010; McGrath & Lilley, 2015). All effort was taken to minimize the number of animals used and their suffering.

Experiments were performed in adult male mice (10–16 weeks old, weighing 25–30 g). All mice were kept and bred in the animal facility of the Medical University of Innsbruck. Y4KO mice were backcrossed to a C57Bl/6NCrl background for at least 10 generations and compared with respective WT controls before the experiments. For both mouse lines, experimental animals were derived from homozygote breeding pairs. Generation of Y4KO mice has been described in detail previously (Sainsbury et al., 2002). Deletion of Y4 receptor genes was confirmed in all mice used for the experiments by PCR and agarose gel electrophoresis. Further characterization of receptor deletion was performed in randomly selected mice by in situ hybridization and receptor autoradiography (using human [125I]-PP as ligands for Y4 receptors), as described in detail previously (Gobbi et al., 1998; Tasan et al., 2009).

Mice were housed in groups of three to five animals per cage under standard laboratory conditions (12 h/12 h light/dark cycle, lights being on at 07:00 h, food and water ad libitum).

The mouse was chosen as an appropriate animal species, because we wanted to study the underlying cause and potential treatment options for human fear-related disorders. Pavlovian fear conditioning was selected as a well-established model to study the development and extinction of fear. Because human anxiety disorders can be only studied adequately in mammals, our results have no implications for replacement, refinement or reduction. However, the parameters of the behavioural experiments, including fear conditioning (Verma et al., 2012) and fasting (Verma et al., 2016), were refined according to our previous experiments and were selected as the least harmful possible while still allowing the study to be conducted.

Experimental conditions. The number of animals per experiment was based on previous data (Verma et al., 2012) and a corresponding power analysis. The exact group size for the individual experiments is shown in the corresponding figure legends. In general, a group size of 7–10 animals was used. An unequal group size in Figures 1 and S2 was due to a limited availability of male Y4KO mice and respective, age-matched controls.

For randomization, half the litter of wildtype (WT) mice was in the control group, while the other half was in the treatment group. Blinding was implemented as follows. The operator was blinded to the group identity, but not to animals of the same group. Thus, for i.p. injection of a drug, two different solutions were prepared: solution A (e.g. saline) and solution B (e.g. [K30(PEG2)]hPP2-36). Importantly, the data analyst was blinded (histology), or analyses were performed by automated computer software (behavioural experiments).

Mice were killed by an i.p. injection of a lethal dose of thiopental (500 mg·kg⁻¹; Sandoz, Austria) to remove tissue for histological analyses or by CO2 inhalation after the final behavioural experiments.

Behavioural experiments. For behavioural experiments, [K30(PEG2)]hPP2-36 was dissolved in 0.9% saline to yield a final drug concentration of 1 mg·kg⁻¹.

Fear conditioning paradigm. Naïve WT (C57Bl/6NCrl) or Y4KO mice were used for fear conditioning experiments. Fear conditioning experiments were conducted during the light phase of the light/dark cycle in sound proof chambers (TSE Systems, Bad Homburg, Germany) by repetitive exposure of the mice to an auditory stimulus co-terminating with a foot shock, as described in detail previously (Verma et al., 2012).
Fear conditioning and context testing were performed in context A consisting of a transparent acrylic mouse conditioning chamber with a metal grid floor. The whole set-up was enclosed by a sound attenuating chamber. Illumination was 80 lux, and chambers were cleaned with 70% ethanol. Fear recall as well as fear extinction and extinction recall was performed in a different context consisting of a dimly illuminated (10 lux) chamber with black, smooth walls and the floor cleaned with 1% acetic acid (context B).

Fear conditioning procedure. On day 1 (context A), mice were subjected to a differential fear conditioning paradigm in which one auditory stimulus served as a CS because it was explicitly paired (CS+, 30 s white noise, 80 dB) with the US, whereas the second auditory stimulus was not paired (CS−, 30 s, 3.5 kHz, 80 dB). All animals received five CS− and five CS+ in an alternating order, starting with a CS+. The US co-terminating with each CS+ consisted of a foot shock. The shock intensity was set to 0.5 mA (2 s), a minimum threshold at which both WT and Y4KO mice displayed respective behavioural reactions in the sensitivity analysis (Verma et al., 2016). On days 2 and 3, mice were tested for their context (context A for 15 min) and CS-related fear memory (context B) respectively.

Fear extinction procedure. Because male Y4KO mice on C57BL/6NCrl background did not show fear extinction, we performed an extensive extinction protocol. In these experiments, we performed a total of six extinction sessions (context B), each consisting of 15 CS+ (30 s, inter stimulus interval 5 s), and extinction recall was tested the following day (five CS+ presentations, context B). Reinstatement was induced by an unsignalled foot shock in context A (day 20) and tested on day 21 by exposing the mice to five CS+ in context B. Behaviour was recorded by a video camera and scored offline by a pixel-based analysis software (http://topowatch.sourceforge.net/, TopoWatch v0.3). The parameters of the programme were validated by comparison with the manual analysis of two experienced observers. A detailed validation procedure of the analysis software for detecting freezing behaviour including baseline activity and reactive motor activity in an unfamiliar environment relevant for fear conditioning (e.g. the fear conditioning box) has been published previously (Verma et al., 2012).

Peripheral injection of Y4 receptor agonist on fear-related behaviour. Fear conditioning was performed in WT (C57Bl6/NCrl) and Y4KO mice on day 1 (context A) as described in the previous paragraph. To avoid interference with postprandial released PP, mice were fasted overnight (16 h) before injection of the drug followed by fear extinction training. Water was freely available during the fasting period. Based on our previous results, a 16 h fasting period was required to reach our experimental aim (Verma et al., 2016). A control experiment was performed without food restriction (non-fasting). [K30(PEG2)]hPP2-36 (1 mg·kg−1) or saline was injected i.p. 60 min before behavioural testing, as indicated in the timeline of the respective experiments. For extinction experiments, mice were divided into two groups, receiving an i.p. injection of either the long-acting Y4 receptor agonist [K30(PEG2)]hPP2-36 (1 mg·kg−1) or saline in controls followed after 60 min by extinction training (extinction 1; 5 CS− and 25 CS+, context B). Mice were then tested on day 3 for extinction recall under drug-free conditions by exposure to five CS− (context B). To exclude any bias of group selection, another extinction session was performed on day 4 (extinction 2; 25 CS+, context B) followed by extinction recall on day 5 (five CS+; context B), both under drug-free conditions. Spontaneous recovery (day 12) and renewal (day 13) were tested by exposing the mice to five CS+ in contexts B and A respectively.

Determination of motor activity after [K30(PEG2)]hPP2-36 injection. Male WT mice were injected i.p. with either the long-acting Y4 receptor agonist [K30(PEG2)]hPP2-36 (1 mg·kg−1) or saline. Activity measurements were performed for 24 h in a novel cage. Mice were single-housed in standard cages with food and water ad libitum. Movements of the mice were detected by an infrared sensor throughout light and dark phase (TSE LabMaster InfraMot, Bad Homburg, Germany).

Histological analysis

A separate group of WT and Y4KO mice were fasted overnight (16 h), injected i.p. with [K30(PEG2)]hPP2-36 or saline and subjected to extinction training 60 min later. Using a cryostat 40 μm coronal, free-floating sections were cut for subsequent immunohistochemistry.

Immunohistochemistry procedures. Immunohistochemical analysis was performed on free-floating, PFA-fixed, 40 μm thick coronal sections using indirect peroxidase labelling, as described previously (Tasan et al., 2011). The following antisera were used: polyclonal rabbit anti-c-Fos (1:20 000 PC38, Calbiochem) and polyclonal rabbit anti-pan-orexin (1:1000, Abcam ab6214, Lot:GR63996-1). In brief, coronal sections were incubated free-floating in 10% normal goat serum (Biomedica, Vienna, Austria) in Tris–HCl buffered saline (TBS; 50 mM, pH 7.2) for 90 min, followed by incubation with primary antisera. The resulting complex was visualized by incubation with HRP-coupled secondary antibody (1:250 P0448; Dako, Vienna, Austria) at room temperature for 150 min, followed by tyramide amplification solution (1:100, TSA Fluorescein) for 6 min. After staining, sections were exposed to 0.01 M of HCl for 20 min at room temperature to denature HRP and first primary antibodies, and incubation with the secondary primary antibody was performed as described for c-Fos except that TSA CY3 was used for staining. Sections were mounted on slides and covered using Vectashield mounting medium (Vector laboratories, Inc., Burlingame, CA, USA).

The number of c-Fos positive cells was obtained bilaterally from six matched sections per animal showing the lateral hypothalamus, at a magnification of 400×, in multiple separate fields and mean values were calculated for each mouse. Results are presented as number of immunoreactivity positive cells per section and expressed as mean ± SEM. Analysis of dual labelling immunofluorescence was carried out as described elsewhere (Tasan et al., 2011). In brief, for each brain area containing a region of interest, four matched sections per mouse (seven mice per group) were processed for either c-Fos/orexin for dual localization. Identification of dual-labelled cells was performed at 400 times magnification within the respective brain area in each section.
Results

Pharmacological characterization of [K^{30}(PEG2)]hPP_{2-36}

Agonist activity of the peptide at human Y receptors. Activity at the human receptors, hY1, hY2, hY4 and hY5, was measured by accumulation of IP₃, after channeling cAMP inhibition to phospholipase C activity, in COS7 cells. Results from the concentration-response studies are shown in Supporting Information Figure S2 and summarized in Table 1. This modified PP peptide, [K^{30}(PEG2)]hPP_{2-36} was found to selectively activate hY₄ receptors with an EC\(_{50}\) of 9.1 nM, whereas with all other Y receptors, no full curves could be obtained over the concentration range used (\(10^{-11} - 10^{-5}\)M). For comparison, the native human PP has an EC\(_{50}\) value of 1.3 nM at the hY₄ receptor (Bellmann-Sickert et al., 2011; Mäde et al., 2014), indicating that PEGylation had only a minor effect on agonist potency.

Proteolytic stability assay in human blood plasma. Peptide stability was evaluated by incubating a fluorescently labelled derivative in human blood plasma and subsequent chromatographic analysis of fluorescent degradation products. TAMRA-[K^{30}(PEG2)]hPP_{2-36} exhibited a high degree of stability towards proteolysis with 92% intact peptide after an incubation period of 96 h (Supporting Information Figure S3). This is in accordance with previous results for different hPP analogues that all showed a substantial elevation of proteolytic stability in different media. The native hPP showed about 60% intact compound after incubation for 96 h, under the same experimental conditions (Bellmann-Sickert et al., 2011; Mäde et al., 2014).

Role of peripheral PP in the modulation of motor activity

PP is the main ligand for the Y₄ receptor. It is produced in the pancreas and released upon ingestion of food. Thus, to investigate the effect of peripherally released PP and its relation to Y₄ receptors, we injected the long-acting Y₄ receptor agonist [K^{30}(PEG2)]hPP_{2-36} i.p. into WT and Y₄KO mice, followed by behavioural testing.

To exclude non-specific effects of [K^{30}(PEG2)]hPP_{2-36} on motor activity that may compromise freezing detection during extinction training, we first injected male WT mice i.p. with [K^{30}(PEG2)]hPP_{2-36} or saline and subsequently observed the activity of the mice for 24 h in a novel cage (Supporting Information Figure S4). Repeated two-way ANOVA revealed no difference between treatment groups (\(F_{(1/13)} = 0.03, P > 0.05\)), but

Table 1

| hY₁ | hY₂ | hY₄ | hY₅ |
|-----|-----|-----|-----|
| EC\(_{50}\) | pEC\(_{50}\) | EC\(_{50}\) | pEC\(_{50}\) | EC\(_{50}\) | pEC\(_{50}\) | EC\(_{50}\) | pEC\(_{50}\) |
| > 1000 | n. d. | > 1000 | n. d. | 9.1 | 8.0 ± 0.2 | 402 | 6.4 ± 0.1 |

Peptide activity determined from concentration-response curves from the inositol phosphate turnover assay. Inositol phosphate accumulation was measured by stimulating COS7 cells stably transfected with the respective Y receptor subtype and a chimeric G\(_{i/o}\) protein, over a range of peptide concentrations (Supporting Information Figure S2). Experiments were performed in duplicate, at least twice. Data in the Table are EC\(_{50}\) (shown as nM) and pEC\(_{50}\) (± SEM) values, generated by nonlinear regression using GraphPad Prism 5.0. n.d., not determined.
different activity over time ($F_{23/209} = 8.93, P < 0.05$), indicating that i.p. injection of $[K^{30}(PEG2)]hPP_{2-36}$ did not influence general motor activity and preserved physiological activity changes during light/dark cycles.

**Effect of peripherally injected $[K^{30}(PEG2)]hPP_{2-36}$ on cued fear extinction**

Specific impairment of cued fear extinction in Y4KO mice suggests that activation of Y4 receptors may facilitate fear extinction. Thus, to investigate if peripherally released PP is the main actor mediating the effect of Y4 receptors on cued fear extinction learning and to differentiate acute effects of Y4 receptor stimulation from permanent Y4 receptor deletion, we injected $[K^{30}(PEG2)]hPP_{2-36}$ i.p. into male WT mice 60 min before fear testing and consecutive extinction training (Figure 2A). Following fear acquisition, male WT mice were divided into two equal groups (Figure 2B). All mice were fasted overnight for 16 h to avoid interference with endogenous PP release that naturally occurs after feeding. As shown in Figure 2C, i.p. injection of $[K^{30}(PEG2)]hPP_{2-36}$ significantly facilitated extinction learning in WT mice ($F_{1/22} = 5.80, P < 0.05$). The effect of facilitated fear extinction was consolidated into an improved extinction memory as reflected by a reduction of freezing during extinction recall performed 24 h later under drug-free and non-fasting conditions (Figure 2D, $t_{14} = 2.20, P < 0.05$). A consecutive extinction trial (Figure 2E, extinction 2) was performed on day 4 followed by extinction recall 2 on day 5, both in the absence of drug and with food and water available *ad libitum*, demonstrating that extinction under drug-free, non-fasting conditions yields similar levels of CS-induced freezing in both groups (Figure 2F, $t_{14} = 0.42, P > 0.05$).

Central Y4 receptors may be targeted by peripherally released PP; however, also, NPY released in the CNS displays high affinity for Y4 receptors (Bard et al., 1995; Gehlert et al., 1996), while PP may act also on Y6 receptors (Yulyaningsih et al., 2014). Thus, to investigate if the extinction-promoting effect is due to peripherally applied PP acting specifically on Y4 receptors, we injected $[K^{30}(PEG2)]hPP_{2-36}$ 60 min before extinction in fasted (16 h) Y4KO mice (Figure 3A). In contrast to concomitantly tested WT mice (Figure 3B and E, $F_{1/16} = 6.18, P < 0.05$), Y4KO mice did not respond to $[K^{30}(PEG2)]hPP_{2-36}$ injection (Figure 3C–G, $F_{1/16} = 0.02, P > 0.05$), indicating the specific activity of $[K^{30}(PEG2)]hPP_{2-36}$ on Y4 receptors. Further analysis revealed that WT mice injected with $[K^{30}(PEG2)]hPP_{2-36}$ already displayed reduced freezing at CS13, while no such effect was apparent in Y4KO mice (Figure 3E, $F_{1/16} = 8.38, P < 0.05$). Importantly, there was a difference in freezing behaviour at CS20 between WT and Y4KO mice only in the $[K^{30}(PEG2)]hPP_{2-36}$-treated group (Figure 3F, $F_{1/16} = 12.02, P < 0.05$). Although an apparent difference in freezing levels may be indicated during the end of the extinction session (e.g. CS25) also in Y4KO mice, neither the overall course of the extinction session nor a specific testing of CS25 was significantly different. These additional analyses support a drug effect in WT mice, but the absence of an effect in Y4KO mice. Interestingly, and as shown previously (Verma et al., 2016), impaired fear extinction was rescued in Y4KO mice upon fasting, further demonstrating the close relationship of energy homeostasis and emotional processing (Figure 3C and G).

Taken together, we conclude that peripherally injected $[K^{30}(PEG2)]hPP_{2-36}$ specifically promoted cued fear extinction learning by activating Y4 receptors in fasted mice.

**Role of PP and Y4 receptors in the permanence of extinction memory**

Fear memories are strong and often persist lifelong, whereas extinction memories are labile and transient, resulting in relapse of fear in particular upon stressful situations. To investigate the permanence of extinction memory, we employed

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**Figure 2**

Cued fear extinction is facilitated by peripheral injection of the long-acting Y4 receptor agonist $[K^{30}(PEG2)]hPP_{2-36}$. (A) Experimental procedure of fear conditioning and extinction experiments. (B) Following fear acquisition, male WT mice were divided into two equal groups; (C) peripheral injection of $[K^{30}(PEG2)]hPP_{2-36}$ in fasted mice (16 h, before and during fear extinction) resulted in facilitated extinction of CS-induced fear compared with saline-injected controls. (D) Freezing to the CS was reduced in the $[K^{30}(PEG2)]hPP_{2-36}$ group during extinction recall tested under drug-free conditions with food available *ad libitum*. (E) An additional fear extinction session of the saline and $[K^{30}(PEG2)]hPP_{2-36}$ groups under drug-free and fed conditions resulted in (F) equal freezing levels during extinction recall 2, demonstrating similar composition of the groups (saline: $n = 8$, $[K^{30}(PEG2)]hPP_{2-36}$: $n = 8$) Data are expressed as mean ± SEM, *$P < 0.05$, significantly different from saline.
Figure 3
Peripheral injection of \([K^{30}(PEG2)]hPP_2.36\) facilitates fear extinction by activation of Y4 receptors. (A) Experimental timeline of fear conditioning experiments. (B) Facilitated fear extinction in male WT mice but not in (C) Y4KO mice after peripheral injections of \([K^{30}(PEG2)]hPP_2.36\) 60 min before fear extinction. (D) Equal freezing levels of all groups during the first CS, demonstrating equal fear expression. (E) Reduced freezing level of \([K^{30}(PEG2)]hPP_2.36\)-injected WT mice compared with those of all other experimental groups (CS13). (F) While both WT groups displayed extinction of conditioned fear, Y4KO mice exhibited still high freezing levels (CS20) and (G) extinction of conditioned fear in all four treatment groups at CS 25 (WT: saline: \(n=10\), \([K^{30}(PEG2)]hPP_2.36\): \(n=10\); Y4KO: saline: \(n=10\), \([K^{30}(PEG2)]hPP_2.36\): \(n=10\); data are expressed as mean ± SEM. *\(P<0.05\), PP significantly different from saline. #\(P<0.05\), WT significantly different from Y4KO.

Figure 4
Treatment with \([K^{30}(PEG2)]hPP_2.36\) before fear extinction results in long-term suppression of fear. (A) Experimental timeline of fear conditioning experiments. (B) Male WT mice were divided into two groups with equal fear acquisition. (C) Facilitated extinction during three consecutive extinction sessions (E1–E3) in \([K^{30}(PEG2)]hPP_2.36\)-treated mice compared with saline-injected controls and equal freezing levels during extinction recall (ER), but decreased freezing of \([K^{30}(PEG2)]hPP_2.36\) group during (D) spontaneous recovery and (E) renewal on days 12 and 13 respectively (saline: \(n=7\), \([K^{30}(PEG2)]hPP_2.36\): \(n=7\); data are expressed as mean ± SEM. *\(P<0.05\), significantly different from saline.

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two well-established paradigms, spontaneous recovery and renewal testing. Male WT mice were fasted for 16 h, injected with [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} or saline and subjected to an extinction training 60 min later on day 2 (Figure 4A). Two additional extinction sessions under drug-free, non-fasting conditions were added to yield equal levels of fear expression (Figure 4A). In mice that had received [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} before extinction training 1, reduced freezing was evident during spontaneous recovery (Figure 4D, \( t_{(12)} = 2.24, P < 0.05 \)) and renewal (Figure 4E, \( t_{(12)} = 2.28, P < 0.05 \)) tests that were performed on days 12 and 13 respectively. These experiments demonstrate that [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} injection before extinction training creates a long-lasting extinction memory with permanent suppression of fear.

**Influence of homeostatic balance on the effect of PP on extinction learning**

In order to avoid possible interference with endogenously released PP, all previous experiments were performed after a 16 h period of fasting. However, we have previously shown that fasting alone is sufficient to promote extinction learning (Verma et al., 2016). To investigate if short-term fasting (16 h) controls the responsiveness to PP, we repeated the experiments in non-fasted mice by injecting [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} i.p. 60 min before extinction training (Supporting Information Figure S5A). Interestingly, when we performed extinction training in non-fasted WT mice, peripheral injection of [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} did not facilitate extinction learning anymore (Figure SSC and D, \( F_{(1/10)} = 0.30, P > 0.05 \)), suggesting that food deprivation is essential for the action of [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} on fear extinction.

**Effect of peripheral injection of [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} on the acquisition and consolidation of fear memories**

Although male Y4KO mice did not display any alterations in acquisition, consolidation or recall/express expression of fear, a possible Y4 receptor-mediated effect may have been masked here by developmental adaptations. Thus, it is important to investigate the acute effects of Y4 receptor stimulation during fear acquisition and consolidation in WT mice. To characterize the effect of PP and Y4

![Figure 5](image)

**Figure 5**

Peripheral injection of [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} before fear extinction specifically reduces the activation of orexin neurons in the lateral hypothalamus. Male WT and Y4KO mice were fasted before and during fear extinction (16 h) and injected with [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36} 60 min before fear extinction; all mice were perfused 90 min after the end of extinction training. Representative images of immunohistochemistry for c-Fos, orexin and overlay for (A–C) saline-injected and (D–F) [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36}-injected WT mice. Arrows point to examples of dual-labelled neurons, higher magnification in A and F with Hoechst staining for labelling nuclei; scale bar 100 \( \mu \)m. Histograms depicting quantification of (G) percentage of cFos/orexin double-labelled neurons and (H) number of orexin neurons in the lateral hypothalamus of WT and Y4KO mice (WT: saline: \( n = 7 \); [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36}: \( n = 7 \); Y4KO: saline: \( n = 7 \); [K\textsuperscript{30}(PEG2)]hPP\textsubscript{2-36}: \( n = 7 \); data are expressed as mean ± SEM. *\( P < 0.05 \), significantly different from saline. #\( P < 0.05 \), WT significantly different from Y4KO.
receptors in fear acquisition and consolidation we injected i.p. [K30(PEG2)]hPP2-36 before or after fear acquisition respectively (Supporting Information Figures S6 and S7). Injection of the long-acting Y4 receptor agonist 60 min before fear acquisition did not alter context fear (Figure S6C and D, F(1,19) = 0.70, P > 0.05) or CS-induced fear (t(8) = 0.51, P > 0.05) tested 24 and 48 h later respectively. Furthermore, injection of [K30(PEG2)]hPP2-36 immediately after fear acquisition, a schedule to test fear consolidation, did also not change context fear (Figure S7C and D, F(1,19) = 1.35, P > 0.05) or CS-induced freezing (t(8) = 0.45, P > 0.05). Collectively, these data suggest that PP does not interfere with the acquisition and consolidation of conditioned fear but rather displays a specific, feeding-dependent effect on the acquisition of fear extinction.

Activation of CNS neurons by peripherally injected [K30(PEG2)]hPP2-36 during fear extinction

PP released from the pancreas acts on Y4 receptors in the periphery but may also target the CNS (Dumont et al., 2007; Tasan et al., 2009). To identify brain areas that are differentially activated by PP during fear extinction learning, WT and Y4KO mice were fasted for 16 h, injected i.p. with [K30(PEG2)]hPP2-36 and subjected to fear extinction training. For immediate early gene mapping, brains were removed 90 min after behavioural testing. As shown in Figure 5, peripheral injection of [K30(PEG2)]hPP2-36 significantly reduced the percentage of cFos-positive, orexin neurons in the lateral hypothalamus (Figure 5G) in WT but not in Y4KO mice. These data indicate that fasting and fear exposure result in high arousal during extinction learning and that peripherally injected PP may promote fear extinction in part by reducing stress levels through inhibition of orexin neurons.

Discussion and conclusions

The important function of neuropeptides in regulating fear-related and extinction-related processes is becoming increasingly evident (Bowers et al., 2012; Singewald et al., 2015). However, while most data concern the activity of central neuropeptides, much less is known about the interaction of the gut–brain axis in emotional-affective behaviour (Cryan and Dinan, 2012; Holzer and Farzli, 2014). Here, we demonstrated that peripherally released PP is crucially and specifically involved in the extinction of cued fear probably by acting on central Y4 receptors. In particular, we demonstrated that peripheral injection of the Y4 receptor agonist [K30(PEG2)]hPP2-36 significantly promoted extinction learning in fasted, but not in fed, mice and results in a lasting suppression of fear. An absence of an effect in Y4KO mice suggests that it is mediated by activation of Y4 receptors. Furthermore, peripheral application of [K30(PEG2)]hPP2-36 before fear extinction training resulted in reduced activation of orexin neurons in the lateral hypothalamus, providing a possible interaction point of central and peripheral peptidergic systems during fear processing. While the percentage of activated cFos/orexin neurons in Y4KO mice did not respond to peripheral injection of [K30(PEG2)]hPP2-36, their activation was generally reduced compared with that in WT mice. Multiple changes may occur during embryonic and also postnatal development that may account for this reduced activation upon genetic deletion of Y4 receptors. Such a reduced activity of orexin neurons may be even more pronounced under non-fasted conditions and could in part explain the impaired fear extinction of Y4KO mice.

While Y4 receptors are expressed in discrete regions of the brain, such as specific nuclei of the hypothalamus and brain stem, their main ligand, PP, is not expressed in the CNS (Pieribone et al., 1992) but synthesized in endocrine F cells of pancreatic islets (Holzer et al., 2012). PP is released postprandially via the vagus nerve to act locally in the gut, but it is also released into the circulation activating Y4 receptors in brain regions that are open to the blood–brain barrier (Dumont et al., 2007).

We have demonstrated recently that peripherally applied hPP specifically activates different brain areas in a time-dependent manner, suggesting direct and indirect activation of limbic circuits (Tasan et al., 2009). In particular, activation of the NTS may be crucial for the central effects of peripherally released PP. The NTS is important for sensing glucose levels and it responds to hypoglycemia by stimulating food intake, inducing release of stress hormones and increasing sympathetic activity (Renner et al., 2010; Rinaman, 2010; Wu et al., 2012). Interestingly, extinction-promoting effects are known for glucocorticoids and yohimbine, both compounds that increase stress and sympathetic activity (Barrett and Gonzalez-Lima, 2004; Yang et al., 2006; Holmes and Quirk, 2010; Blundell et al., 2011). Activation of the NTS may thus promote fear extinction by linking the internal homeostatic situation to emotional arousal. Recent evidence has demonstrated that i.p. injection of radioactive-labelled [125I]hPP specifically accumulates in the area postrema and the NTS, suggesting that peripherally released PP directly acts on Y4 receptors in these brain areas (Dumont et al., 2007). NTS neurons have also prominent reciprocal projections to limbic brain areas, in particular to hypothalamic nuclei. Importantly, there is a mutual connection between the NTS and the paraventricular nucleus of the hypothalamus, an area responsible for modulation of sympathetic outflow (Affleck et al., 2012). The important role of the hypothalamus in PP and Y4 receptor-mediated effects is further emphasized by a recent study using MRI scans in fasted rats demonstrating the activation of specific hypothalamic nuclei upon i.p. injection of hPP (Hankir et al., 2011). Furthermore, in the arcuate nucleus of the hypothalamus, PP inhibits GABA neurons by a presynaptic attenuation of glutamate release (Acuna-Goycolea et al., 2005), presenting thus another potential node of interaction with the central stress system.

We have demonstrated that [K30(PEG2)]hPP2-36 acting on central Y4 receptors, specifically facilitates cued fear extinction while not affecting fear acquisition and fear consolidation. Extinction of conditioned fear is a learning process that leads to a gradual decrease in fear expression and involves plasticity in a distributed neural network, including prefrontal cortex, amygdala and hippocampus (Orsini and Maren, 2012). Interestingly, the NTS, a brain stem nucleus with high abundance of Y4 receptors, has also reciprocal connections with different amygdala nuclei and infralimbic cortex (Ricardo and Koh, 1978; Danielsen et al., 1989; Affleck et al., 2012; Garcia-Medina and Miranda, 2013). Thus, Y4 receptors in the NTS may modify signalling in the amygdala and in the infralimbic cortex, thereby specifically facilitating fear extinction.

Furthermore, we found that the extinction-promoting effect of PP depended on two important prerequisites: first, the presence of Y4 receptors and, second, a defined period of food
restriction. In Y4KO mice, impaired extinction acquisition was rescued by a 16 h fasting period but was, however, not further enhanced by peripheral application of [K30(PEG2)]hPP2-36, indicating that the extinction-facilitating effect of PP was mediated by Y4 receptors. In WT mice, fasting also accelerated extinction learning, and this effect was further enhanced by injection of [K30(PEG2)]hPP2-36 in the present study. In contrast, injection of [K30(PEG2)]hPP2-36 did not alter fear extinction in fed WT mice. Thus, [K30(PEG2)]hPP2-36 facilitates cued fear extinction only in fasted WT mice, but not in fed WT mice or Y4KO mice. Compared with our previous study (Verma et al., 2016), the slightly higher freezing levels in the current experiments were likely to be due to the stress of i.p. injections shortly before extinction training.

However, short-term fasting may have contributed to improved extinction learning by increasing arousal. This is also reflected by the activation of orexin neurons in the lateral hypothalamus. Orexin neurons are involved in the modulation of arousal, energy balance and reward processing (Sakurai, 2014). Interestingly, peripherally injected [K30(PEG2)]hPP2-36 inhibited the activation of orexin neurons but promoted extinction in fasted mice. Experimental evidence indicates that orexin-expressing neurons express functional Y4 receptors (Campbell et al., 2003) and injection of hPP was suggested to inhibit fasting-induced increases in orexin mRNA (Sainsbury et al., 2010). However, whether peripherally released PP is able to pass the blood–brain barrier and directly activate Y4 receptors on orexin neurons or the inhibition of orexin neurons is rather indirect remains to be demonstrated (Dumont et al., 2009). Orexin and in particular the orexin-1 receptor have been suggested to increase stress and fear-related behaviours (Furlong et al., 2007). Thus, the extinction-promoting effect of the peripherally applied Y4 receptor agonist [K30(PEG2)]hPP2-36 may be explained by reducing the orexin-induced high fear/stress levels evident during extinction learning. More direct, recent data, favouring the hypothesis that PP and Y4 receptors mediate the facilitation of extinction by inhibiting orexin neurons, come from the work of Flores et al. (2014). The authors demonstrated that blockade of orexin-1 receptors specifically promotes the consolidation of extinction memory (Flores et al., 2014).

We demonstrated previously that fasting promotes fear extinction in WT mice and can even rescue impaired fear extinction in Y4KO mice (Verma et al., 2016). Many different brain circuits are activated upon acute fasting, in particular during concomitant fear exposure; however, only some of these activity changes may promote fear extinction. For instance, increased attention is necessary for successful extinction learning; however, an overly active arousal system may inhibit extinction consolidation. Thus, in fasted mice, PP may suppress an overstimulated arousal system bringing it back to an efficient level, but this mechanism may be ineffective in fed mice.

Importantly, also, NPY is crucially involved in both food intake (Loh et al., 2015) and modulation of emotional affective behaviour, including fear conditioning and extinction (Tasan et al., 2016). Thus, the NPY system may be at a central node for adapting emotional responses to the present homeostatic situation. It is further conceivable that PP released in the periphery will interact with the central NPY system to achieve this integration of feeding behaviour and emotional processing in the CNS.

A crucial finding is that the suppression of fear behaviour by [K30(PEG2)]hPP2-36 treatment before fear extinction is long-lasting and independent of context as demonstrated by the reduced freezing levels during spontaneous recovery and renewal. Interestingly, freezing behaviour in [K30(PEG2)]hPP2-36-treated mice was even lower than during extinction recall. This suggests that application of [K30(PEG2)]hPP2-36 not only promoted extinction learning but also strengthened consolidation and permanence of an acquired extinction memory, an effect that may persist even after washout of the drug. Further experiments are needed to elucidate this effect in more detail.

Fear extinction is considered as neurobiological basis for exposure therapy in humans. However, a major drawback is currently that extinction memory is rather labile resulting in frequent re-emergence of fear. Thus, a supportive pharmacological intervention that can stabilize an acquired extinction memory during psychotherapy is of high relevance.

Importantly, we highlight here a close interaction of two life-sustaining circuits, hunger and fear, and we present a peptide that is involved in both feeding and fear. Thus, a thorough analysis of critical mediators of the gut–brain axis may yield further promising drug targets for treating fear and anxiety-related disorders.

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Author contributions

D.V. and B.H. performed the research; D.V. and R.O.T. analysed the data; H.H. contributed the knockout mice for the study; A.B.S., V.T. and K.B.S. synthesized the Y4 receptor agonist [K30(PEG2)]hPP2-36 and performed the in vitro characterization; R.O.T. designed the research study and R.O.T. and G.S. wrote the paper.

Conflict of interest

The authors declare no conflicts of interest.

Declaration of transparency and scientific rigour

This Declaration acknowledges that this paper adheres to the principles for transparent reporting and scientific rigour of preclinical research recommended by funding agencies, publishers and other organizations engaged with supporting research.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:
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**Figure S1** RP-HPL chromatogram (A) and MALDI-TOF mass spectrum (B) of [K30(PEG2)]hPP. RP-HPLC was performed on a Jupiter 4u Proteo 90 Å (Phenomenex) applying a gradient of 20 to 60% of eluent B (0.08% TFA in acetonitrile) in eluent A (0.1% TFA in water) over 40 min.

**Figure S2** Inositol phosphate turnover assay of the activity of [K30(PEG2)]hPP2-36 at all human Y receptors. Inositol phosphate accumulation was measured by stimulating COS7 cells stably transfected with the respective Y receptor subtype and a chimeric Gi/o-protein with different peptide concentrations. Experiments were performed in duplicate, at least twice. Data points shown are means ± SEM. Curves were generated by nonlinear regression using GraphPad Prism 5.0.

**Figure S3** Proteolytic stability of [K30(PEG2)]hPP2-36 in human blood plasma. Peptide was incubated in human blood plasma at 37°C for the indicated periods. The degradation assay was carried out in two independent experiments (n = 2) and results are presented as means ± SEM.

**Figure S4** Activity measurements over 24 h after i.p. injection of a Y4 receptor agonist or saline. Motor activity in a novel cage was unchanged after i.p. injection of the Y4 receptor agonist [K30(PEG2)]hPP2-36 (saline: n = 7, [K30(PEG2)]hPP2-36: n = 8; data are expressed as mean ± SEM).

**Figure S5** Peripheral [K30(PEG2)]hPP2-36 injection does not affect fear extinction in non-fasted mice. (A) Experimental timeline of fear conditioning and extinction experiments, (B) after fear acquisition mice were divided into two equal groups, (C) fear extinction and (D) extinction recall, is similar between non-fasted [K30(PEG2)]hPP2-36-injected WT mice and saline-injected controls (saline: n = 6, [K30(PEG2)]hPP2-36: n = 6; data are expressed as mean ± SEM).

**Figure S6** Peripheral injection of the Y4 receptor agonist [K30(PEG2)]hPP2-36 does not influence acquisition of conditioned fear. (A) Experimental set-up with 16 h of fasting before and during fear acquisition, (B) no change in fear acquisition, (C) context fear and (D) CS-induced fear expression after peripheral injection of [K30(PEG2)]hPP2-36 60 min before fear acquisition compared with saline-injected controls (saline: n = 6, [K30(PEG2)]hPP2-36: n = 6; data are expressed as mean ± SEM).

**Figure S7** The Y4 receptor agonist [K30(PEG2)]hPP2-36 did not affect the consolidation of conditioned fear. (A) Experimental procedure of fear conditioning experiments. (B) Following fear acquisition, mice were divided into two groups that were injected with [K30(PEG2)]hPP2-36 or saline, (C) no change in context fear extinction and (D) CS-induced freezing of [K30(PEG2)]hPP2-36 injected mice compared to saline injected controls (saline: n = 6, [K30(PEG2)]hPP2-36 n = 6; data are expressed as mean ± SEM).