Central neuropeptide Y receptors are involved in 3rd ventricular ghrelin induced alteration of colonic transit time in conscious fed rats
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Abstract

Background: Feeding related peptides have been shown to be additionally involved in the central autonomic control of gastrointestinal functions. Recent studies have shown that ghrelin, a stomach-derived orexigenic peptide, is involved in the autonomic regulation of GI function besides feeding behavior. Pharmacological evidence indicates that ghrelin effects on food intake are mediated by neuropeptide Y in the central nervous system.

Methods: In the present study we examine the role of ghrelin in the central autonomic control of GI motility using intracerebroventricular and IP microinjections in a freely moving conscious rat model. Further the hypothesis that a functional relationship between NPY and ghrelin within the CNS exists was addressed.

Results: ICV injections of ghrelin (0.03 nmol, 0.3 nmol and 3.0 nmol/5 µl and saline controls) decreased the colonic transit time up to 43%. IP injections of ghrelin (0.3 nmol – 3.0 nmol kg⁻¹ BW and saline controls) decreased colonic transit time dose related. Central administration of the NPY₁ receptor antagonist, BIBP-3226, prior to centrally or peripherally administration of ghrelin antagonized the ghrelin induced stimulation of colonic transit. On the contrary ICV-pretreatment with the NPY₂ receptor antagonist, BIIE-0246, failed to modulate the ghrelin induced stimulation of colonic motility.

Conclusion: The results suggest that ghrelin acts in the central nervous system to modulate gastrointestinal motor function utilizing NPY₁ receptor dependent mechanisms.

Background

The presence or absence of food in the gut stimulates the release of several regulatory peptides. These orexigenic (NPY, AGRP, ghrelin, MCH, Orexin-A, ...) and anorexi-
functions like secretion and motility. For example fasted motor activity of the GI tract, e.g. the colon, is observed after intracerebroventricular (ICV) injection of neuropeptide Y whereas CRF ICV-treatment cause the disruption of fasted colonic motor activity [1]. Stomach-derived ghrelin is the first peripheral orexigenic peptide identified [2-6]. There is convincing evidence from several groups of investigators that ghrelin acts in the CNS and the periphery to simulate not only feeding but also GI function such as gastric acid secretion and gastric motility in rodents [7,9-11]. However, it is still unknown whether ghrelin is involved in the CNS control of other digestive functions besides gastric acid secretion and motility. Recent studies suggest that CNS-signaling by circulating ghrelin is mediated downstream by neurons of arcuate nucleus and the paraventricular nucleus of the hypothalamus, in particular, neurons expressing neuropeptide Y and agouti-related protein (AGRP) [12-14]. Furthermore it has been demonstrated that there is an anatomical interaction and functional relationship between ghrelin and neuropeptide Y. Using electrophysiological recordings Cowley et al have found that ghrelin stimulated the activity of arcuate NPY-ergic neurons and mimicked the effect of NPY in the paraventricular nucleus of the hypothalamus [15]. In addition ghrelin simulates food intake through hypothalamic NPY_1 receptors [1,16,17]. Thus, the question came up "are NPY receptors involved in the ghrelin effect on GI function?" Among others, neuropeptide Y plays a role in the CNS control of gastrointestinal function [1,18]. NPY activates at least six receptor subtypes, NPY_1 to NPY_6. NPY binds preferentially with high affinity to Y_1 and Y_2 receptors, and there is evidence suggesting that these two receptor subtypes are involved in CNS regulation of digestive function by NPY action in arcuate nucleus and the paraventricular nucleus of the hypothalamus [18].

Taken together there is overwhelming evidence that ghrelin, beside its satiety modulatory capacity, is involved in the CNS control of digestive function of the upper gastrointestinal tract. In the CNS ghrelin and NPY, the most potent orexigenic neuropeptides known, are anatomical associated and functionally related. Moreover hypothalamic NPYergic neurons are downstream mediators of feeding related ghrelin action.

In the present study we scrutinize the hypothesis that central neuropeptide Y receptor activation is involved in the ghrelin induced modulation of gastrointestinal motility using a microinjection-model in conscious fed and freely moving rats.

Methods

Animals

All experimental components described were performed in accordance with the requirements of German legisla-

tion for the protection of animals and were licensed and supervised by the appropriate government body.

Male Sprague-Dawley rats with a mean body weight of 350 ± 50 g were maintained on a 12 : 12 h photoperiod. They were housed in colony cages under conditions of controlled humidity and temperature (22 ± 2°C) for at least 7 days prior to the surgical procedure. The animals were fed a standard rat diet (Altromin®, Lage, Germany) an tap water ad libitum. After surgical procedures, rats were housed individually. During experimental procedure the animals had continuous access to food and water.

Drugs

Ghrelin (Bachem, Heidelberg, Germany) doses of 0.03 nmol (100 ng), 0.3 nmol (1 µg) and 3 nmol (10 µg)/5 µl were dissolved in 0.15 M sterile saline (B. Braun, Melsungen, Germany). The NPY_1 receptor antagonist, BIBP-3226 (200 nmol/5 µl; Sigma-RBI, Natrix, MA, USA) [see Ref.: [19]] and the NPY_2 receptor antagonist, BIIE-0246 (120 nmol/5 µl; Boehringer-Ingelheim, Biberach, Germany) [see Ref.: [21]] were dissolved in sterile 0.15 M saline. The NPY receptor antagonists were used in similar equipotent nanomolar concentrations. The used intracerebroventricular concentrations of the receptor antagonists were comparable with the ICV-dosages used by other groups in rodents [18,20]. Probes were aliquoted and frozen (-80°C). Fresh aliquots were thawed on each experimental day before injections. Any excess was discarded. In our hands nanomolar concentrations of BIBP-3226 and BIIE-0246 were effective in antagonization of NPY receptor subtypes without any side effects. In particular no central depressive effects or conspicuous behavior was observed after BIBP-3226 treatment [18].

Cerebral cannula

For surgical procedures, rats were anesthetized with a mixture of ketamine (75 mg kg⁻¹ i.p., Parke-Davis, Freiburg, Germany) and xylazine (5 mg kg⁻¹ i.p., Bayer AG, Leverkusen, Germany). Animals were positioned in a stereotactic apparatus (David Kopf Instruments, Tujunga, CA). The head was fixed in a nose-down-position (-3 mm) and the skull exposed. Then trepanation of the skullcap was performed according to coordinates obtained from Paxinos and Watson [22] (mm from bregma: anterior-posterior = -3.30; lateral = ± 0.0; dorsoventral = -3.8). According to these coordinates a 22-gauge guide cannula (Bilarney / Plastic one, Düsseldorf, Germany) was implanted into the third ventricle. The cannula was anchored by dental cement and stainless steel screws affixed to the skull. Dummy cannulas (28 G), extending 2 mm beyond the guide cannula tips, were inserted to prevent blockage. After cerebral surgery, animals were individually housed. The animals were allowed 4 days
recovery after guide cannula surgeries before the abdominal surgical procedures were performed.

**Colonic catheter**
This method was performed as described elsewhere [1]. Prior to all abdominal surgeries, the animals were food deprived overnight. Four days after cerebral surgery, rats were anaesthetized with a mixture of ketamine (75 mg kg\(^{-1}\)) and xylazine (5 mg kg\(^{-1}\)). After laparotomy a polyethylene microcatheter (inside diameter, 1.2 mm; outside diameter 1.7 mm; Becton Dickinson, New Jersey, USA) was chronically implanted into the proximal colon 1 cm distal from the caecocolonic junction. The catheter was fixed at the colonic wall by a purse-string suture and routed subcutaneously to the interscapular region, where it was exteriorized through the skin and secured. The animals were allowed 7 days recovery after abdominal surgeries before the beginning of habituation training sessions. Experiments were performed in fed, conscious rats.

**Intraperitoneal (IP) and intracerebroventricular (ICV) microinjection**
The doses of ghrelin were calculated according to the lowest effective doses to stimulate food intake [see Ref.: [23]]. For IP injection a 1 ml syringe (Hamilton, Reno, NV, USA) was used. Ghrelin and vehicle were injected IP after central administration of vehicle or NPY receptor antagonist. For IP injections the low dose of ghrelin administered peripherally was 0.3 nmol kg\(^{-1}\)/ml 0.15 M saline and the high dose of ghrelin was 3 nmol kg\(^{-1}\)/ml saline. NPY receptor antagonists were injected ICV 15 min before ghrelin was given peripherally at doses of 200 nmol/rat (BIBP-3226) and 120 nmol/rat (BIIE-0246) respectively.

For ICV injections a 1 µl micro syringe (Hamilton, Reno, NV, USA), attached to a 32 G injection needle via a PE-50 tube-catheter was used. The stainless steel injection cannulas (32 G) were cut to protrude 2 mm beyond the tips of the guide cannulas. The conscious animals were gently restrained by hand, the injection needle was inserted through the guide cannula, and vehicle (5 µl 0.15 M saline) or NPY receptor antagonists (BIBP-3226 200 nmol/5 µl; BIIE-0246 120 nmol/5 µl) and ghrelin (0.03 nmol, 0.3 nmol or 3 nmol/5 µl), were consecutively injected ICV slowly over a 60 s period. We used a 15 min time interval between ICV injection of receptor antagonist or vehicle and ICV ghrelin administration. The injection needle was left in place for 2 min after injection to allow diffusion of the solution and to prevent back flow. Then dummy cannulas were reinserted into the guide cannulas. After the last experimental testing session, the rats were anesthetized and 5 µl of alcian blue 8GX were injected ICV.

**Colonic transit time measurement**
Colonic transit time was calculated by using an enteral dye-marker. Trypan blue, a non-absorbable dye, was injected in 0.2 ml volume through the catheter positioned in the proximal colon and followed by a 0.2 ml saline flush immediately after the ICV or IP microinjection. Colonic transit time was evaluated as the time interval between dye injection and the discharge of the first blue pellet. Faecal pellet output was monitored continuously by a self-developed, automated observation system that mechanically registers the time of all bowel movements for 24 h. The device consists of a conveyor belt placed under the mesh bottom cage, which transports faecal pellets with defined velocity to a collector.

**Brain histology**
The methods were performed as described in previous studies [17]. When experiments were completed, rats were anaesthetized with ketamine (75 mg kg\(^{-1}\)i.p.) and xylazine (5 mg kg\(^{-1}\)i.p.), and 0.05% alcian blue 8GX was microinjected intracerebroventricular under the same conditions as vehicle or peptide. The anaesthetized animals were transcardially perfused with phosphate buffered saline (PBS) buffer (0.1 M, pH 7.4) followed by Zamboni’s fixative (2% formaldehyde and 2% picric acid in 0.1 M PBS buffer, pH 7.4). The brains were removed and cryoprotected in 25% sucrose. The site of injection was confirmed by inspection of intracerebroventricular dye distribution. Animals that received injections outside of the 3rd ventricle were excluded from data analysis.

**Experimental design**
*Experiment I: Effect on colonic motor function of peripheral (IP) and central (ICV) ghrelin administration*
The aim of the first experiment was to determine whether exogenous ghrelin would alter colonic motor function. Thus in experiment I, dose response effects of ghrelin in the cerebrospinal fluid (ICV) and the periphery (IP) on colonic transit time were examined. Ghrelin or saline as a vehicle was administered ICV or IP in conscious lightly restrained rats as previously described. For IP injection the low dose of ghrelin administered peripherally was 0.3 nmol kg\(^{-1}\) BW and the high dose of ghrelin was 3.0 nmol kg\(^{-1}\) BW. After injections, rats were subsequently returned to their home cages and maintained in a non-stressful environment to monitor colonic transit time. In order to minimize interindividual variation, and to reduce the number of animals needed to perform this study, animals were tested twice in this study. In randomized order, each rat received vehicle and a single dose of ghrelin or vehicle ICV or IP. The time interval between the experiments performed on the same animal was at least 4 days.
Experiment II: Effect of BIBP-3226 and BIIE-0246 pretreatment on centrally and peripherally injected ghrelin induced modulation of colonic transit

In experiment II the hypothesis that ghrelin acts at the CNS to modulate colonic motor function via a NPY receptor dependent pathway was addressed. Therefore, we determined if a pretreatment with selective NPY$_1$- (BIBP-3226) and NPY$_2$ (BIIE-0246) receptor antagonists administered into the cerebrospinal fluid would block the alterations of colonic motor activity induced by centrally and peripherally administered ghrelin. The animals were pretreated with the NPY-Y$_1$ and -Y$_2$ receptor antagonists, injected ICV or vehicle (0.15 M saline), 15 min prior to ICV or IP ghrelin injections. Thereafter colonic transit time was assessed as described above.

Data analysis

The criterion used to include results in the data analysis of the ICV-injected group was the correct placement of the ICV cannulas.

Results

Effect of peripherally (IP) and centrally (ICV) administered ghrelin on colonic motor function

In experiment I, dose response effect of peripherally and centrally administered ghrelin on colon transit time in fed and freely moving rats were examined. As demonstrated in Fig. 1, ghrelin injected into the cerebrospinal fluid (CSF) stimulates colonic transit time dose dependently. In rats microinjected with vehicle into the CSF or IP, the average colonic transit time was 322 ± 8 min. As demonstrated in Fig. 1, 0.03 nmol, 0.3 nmol and 3 nmol/5 µl ghrelin injected ICV dose-dependently decreased transit time by 24%, 34% and 43% respectively in conscious fed rats. Peripherally administered ghrelin accelerated transit time up to 36% (Fig. 1).

Effect of ICV NPY receptor antagonist pretreatment on 3rd ventricular and peripherally ghrelin induced stimulation of colonic transit

The hypothesis that ghrelin acts in the brain to stimulate colonic transit via NPY receptor dependent mechanisms was addressed. As shown in Fig. 2, pre-treatment with BIBP-3226 (200 nmol) which is a selective NPY$_1$ receptor antagonist 15 min prior to ICV application of 0.3 nmol ghrelin totally blocked the ghrelin induced effect on colonic transit. Application of BIBP-3226 into the CSF of the control group that was microinjected with vehicle, had no effect. Changes in colonic transit time induced by IP injection of ghrelin (3.0 nmol kg$^{-1}$ BW) were canceled by ICV injection of NPY$_1$ receptor antagonist, BIBP-3226. (Fig.: 2). Pretreatment with the selective NPY$_2$ receptor antagonist, BIIE-0246, failed to affect the ghrelin induced alteration of colonic transit time. (Fig.: 2)

Discussion

The present experiments using a freely moving conscious rat model permit the measurement of colonic motility in rats in the physiological fed status. The results demonstrated that ghrelin given ICV and IP stimulates gastrointestinal motility indicated by shortened colonic transit time. In addition we found that the NPY type 1 receptor is primarily involved in the ghrelin induced modulation of fasted motor activity of the colon.

There is convincing evidence that the most effective appetite stimulating peptides, NPY and ghrelin, act in the CNS and the periphery to simulate not only feeding but also GI function such as gastric acid secretion and gastric motility [1,8-10]. Stomach derived ghrelin, first described in 1999 by Kojima et al., is the first peripheral orexigenic peptide identified [4]. Ghrelin was identified as endogenous ligand for the GH secretagogue receptor (GHS R) and a peripheral metabolic signal informing the brain about stomach nutrient load [17,24]. Physiological studies suggest a functional relationship of ghrelin and NPY within the brain. It has been demonstrated that peripherally (i.v.) and central (ICV) administered ghrelin increases the expression of the immediate early gene c-fos, a marker of neuronal activity, in the arcuate nucleus and the PVN in awake fed rats [25]. Furthermore, exogenous ghrelin increases mRNA levels for NPY into the arcuate nucleus and simulates food intake through hypothalamic NPY$_1$ receptors [14,16,26]. Further Fujino et al. have demonstrated that ICV pretreatment with neuropeptide Y antiserum completely blocked ghrelin induced gastric and duodenal motoractivity [9]. Taken together these data suggest that there is an anatomical interaction and functional relationship between ghrelin and NPY within the brain. Six recognized subtypes of neuropeptide Y receptors have been described (NPY$_1$ to NPY$_6$). Two of these, NPY$_1$ and NPY$_2$ receptors, are found in high density in the hypothalamus. There is compelling evidence that, in particular, NPY$_1$ and NPY$_2$ receptors are involved in the CNS regulation of gastrointestinal function. [1,8,27] For this reason, we focused on neuropeptide Y$_1$ and Y$_2$ receptor pathways in the present study and did not investigate the role of neuropeptide Y receptor subtypes Y$_3$ and Y$_5$ which are also expressed in the hypothalamus and are also involved in the autonomic control of feeding behavior and GI function. In the present study pretreatment with the NPY$_1$ receptor antagonist, BIBP-3226, blocked stimulation of colonic motility induced by systemic microinjection of exogenous ghrelin (ICV and IP). In our hands BIIE-2046, which is a selective antagonist of the NPY Y$_2$ recep-
tor, failed to affect the ghrelin induced induction of fasted motor activity. It was previously described that knocking out NPY significantly decreases ghrelin stimulated feeding [17,28]. In this context Fujino et al. have recently demonstrated that the ghrelin induced fasted gastroduodenal motor activity in rats is blocked by ICV injection of GHS-R antagonist as well as NPY antiserum [9]. The results presented by Fujino et al. also suggest that the vagal pathway may mediate the action of centrally administered ghrelin on gastroduodenal motility [9]. Thus we can speculate that central NPY pathways, e.g. centrally NPY receptor activation, are the primary downstream mediator of circulating ghrelin. This interpretation is consistent with neuroanatomical and physiological facts: Neuropeptide Y works at two sites, locally within the arcuate nucleus to inhibit POMC neuronal activity and at afferent-terminal sites, in particular the paraventricular nucleus of the hypothalamus. Guan et al. have shown that neuropeptide Y- and ghrelin like immunoreactive (LI) neurons within the arcuate nucleus could influence each other by complex synaptic transmissions [29]. Furthermore Cowley et al. have demonstrated that ghrelin stimulated the activity of arcuate neuropeptide Y-LI neurons and mimicked the effect of neuropeptide Y in the PVN [15]. Compelling evidence showed that NPY projections from the arcuate nucleus (ARC) to the PVN are involved in the CNS regulation of food intake and other physiological functions of the organism, e.g. digestive function, by neuroendocrine and autonomic pathways [17,18]. For example NPY released from ARC neurons activates NPY-Y1 receptors in the hypothalamus, e.g. the PVN, and results in the stimulation of GI motor function [18]. Furthermore arcuate

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**Figure 1**

Effect of ghrelin injected into the 3rd ventricle (ICV) and intraperitoneally (IP) on colonic transit time. Ghrelin injected ICV as well as IP induced a dosed-related stimulation of propulsive colonic motor activity. MI = microinjection The bars represent the mean ± SEM. * P < 0.05 vs. vehicle. ■ = vehicle-group; A = 0.03 nmol ghrelin /5 µl 0.15 M saline; B = 0.3 nmol ghrelin /5 µl 0.15 M saline; C = 3 nmol ghrelin /5 µl 0.15 M saline; D = 0.3 nmol ghrelin kg⁻¹ BW; E = 3.0 nmol ghrelin kg⁻¹ BW
NPYergic neurons have been thought to regulate feeding behavior by NPY receptor subtypes Y1 and Y5 in the PVN and adjacent areas [17]. Pretreatment with a Y1, but not other receptor antagonist markedly inhibited ghrelin-induced feeding, pointing to NPY receptor Y1 as one of the downstream pathways [9,17]. With regard to the characteristic physiological feature that peripheral ghrelin does not cross the blood-brain barrier in rodents it is important to note that the arcuate nucleus is the only hypothalamic structure located outside the brain-blood barrier [30]. Thus we can speculate that circulating ghrelin modulates gastrointestinal motility via activation of NPY-Y1 receptors, in the arcuate nucleus. This hypothesis is in good agreement with our observation that the effect of peripherally (IP) administered ghrelin on colonic motility is blocked by ICV pretreatment with the specific NPY Y1 receptor antagonist, BIBP-3226. The NPY Y2 receptor antagonist BIIE-2046 injected ICV fails to affect the stimulated colonic motoractivity induced by ICV or IP injection of ghrelin. MI = microinjection

**Figure 2**
Effect of pretreatment with NPY receptor antagonists on fasted motor activity of the colon induced by centrally (ICV) and peripherally (IP) administered ghrelin. BIBP-3226, which is a selective NPY Y1 receptor antagonist, injected ICV antagonizes the stimulation of colonic transit induced by ghrelin injected in the same route and IP. The NPY Y1 receptor antagonist BIIE-2046 injected ICV fails to affect the stimulated colonic motoractivity induced by ICV or IP injection of ghrelin. MI = microinjection

The bars represent the mean ± SEM. * P < 0.05 vs. vehicle-group; #P < 0.05 vs. ghrelin-group
synaptically [31,32] The question of whether neuropeptide Y4 or Y5 receptors in the CNS are involved in the CNS control of gastrointestinal function should be examined in future studies.

Conclusion
We hypothesize that circulating ghrelin exhibits its effect by activating hypothalamic neurons, in particular neurons in the arcuate nucleus bearing GHS- and NPY-Y1 receptors. Further this ghrelin induced neuronal activation leads to stimulation of GI motor function by activation of higher hypothalamic brain sites, e.g. activation of neuronal projections within the paraventricular nucleus of the hypothalamus. On the other hand it is possible that the site of action of circulating ghrelin is not the hypothalamus but other brain sites. In our model using 3rd ventricular injection of ghrelin this could simply mean that the peptide gained access to the 4th ventricle and reached further caudal brain sites, e.g. NTS, DVC and medulla oblongata. With respect to the distribution of GHS- and NPY receptors in the CNS this hypothesis is possible, but how ghrelin action on any of these brain sites would modulate digestive function is not known. This question should be examined in future studies.

In summary, we presented evidence that ghrelin is involved in the CNS control of GI function. Apart from humoral pathways ghrelin acts into the CNS to control GI function by a mechanism of action involving neuropeptide Y pathways. Further this study support the hypothesis giving by Chen et al. that ghrelin has an absolute requirement for neuropeptide Y pathways to unfold its physiological effects [28].

List of abbreviations
AGRP agouti-related peptide
ARC arcuate nucleus
CART cocain- and amphetamine-regulated transcript
CCK cholecystokinin
CNS central nervous system
CSF cerebrospinal fluid
CRF corticotropin releasing factor
DVC dorsal motor nucleus of vagus
GHS-R growth hormone secretagogue receptor
GLP-1 glucagon like peptide-1
ICV intracerebroventricular
MCH melanocortin hormone
MI microinjection
NPY neuropeptide Y
NTS nucleus of the solitary tract
POMC proopiomelanocortin
PVN paraventricular nucleus of the hypothalamus

Competing interests
The author(s) declare that they have no competing interests.

Authors’ contributions
J JT participated in the design and coordination of the study, performed the microinjection studies and drafted the manuscript. CGT was the surgeon in charge and participated in the animal experiments. M-KHS and SM were involved in the design and coordination of the study. MR participated in the analysis and interpretation of data and revised the manuscript critically.

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