Nasal Oxytocin Administration Reduces Food Intake without Affecting Locomotor Activity and Glycemia with c-Fos Induction in Limited Brain Areas

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Abstract
Recent studies have considered oxytocin (Oxt) as a possible medicine to treat obesity and hyperphagia. To find the effective and safe route for Oxt treatment, we compared the effects of its nasal and intraperitoneal (IP) administration on food intake, locomotor activity, and glucose tolerance in mice. Nasal Oxt administration decreased food intake without altering locomotor activity and increased the number of c-Fos-immunoreactive (ir) neurons in the paraventricular nucleus (PVN) of the hypothalamus, the area postrema (AP), and the dorsal motor nucleus of vagus (DMNV) of the medulla. IP Oxt administration decreased food intake and locomotor activity and increased the number of c-Fos-ir neurons not only in the PVN, AP, and DMNV but also in the nucleus of solitary tract of the medulla and in the arcuate nucleus of the hypothalamus. In IP glucose tolerance tests, IP Oxt injection attenuated the rise of blood glucose, whereas neither nasal nor intracerebroventricular Oxt affected blood glucose. In isolated islets, Oxt administration potentiated glucose-induced insulin secretion. These results indicate that both nasal and IP Oxt injections reduce food intake to a similar extent and increase the number of c-Fos-ir neurons in common brain regions. IP Oxt administration, in addition, activates broader brain regions, reduces locomotor activity, and affects glucose tolerance possibly by promoting insulin secretion from pancreatic islets. In comparison with IP administration, the nasal route of Oxt administration could exert a similar anorexigenic effect with a lesser effect on peripheral organs.

Key Words
Oxytocin · Nasal treatment · Paraventricular nucleus · Dorsal motor nucleus of vagus · Glucose tolerance · Obesity · Insulin release

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Introduction

Oxytocin (Oxt) is a neurohypophysial hormone that regulates uterine contraction during labor and milk ejection [1]. Oxt is synthesized in the paraventricular (PVN) and supraoptic nucleus of the hypothalamus. It is released peripherally after being shuttled to the pituitary [2], or centrally in the brain to regulate neuronal process [3]. Recent studies have clarified new functions of Oxt in the central nervous system, including increased trust [4], regulation of social recognition [5] and development of mother-infant bonding [6]. In addition, our previous study employed Oxt in the PVN-driven anorexigenic circuit in rats [7].

The anorexigenic effect of Oxt has been reported since the early 1990s. Olson et al. [8] showed that Oxt and its agonist decrease food intake when administered centrally in rats, and Arletti et al. [9] showed that intracerebroventricular (icv) injection of Oxt inhibits food intake together with water intake. More recent studies have shown that mice with deficient Oxt or Oxt receptor (Oxt-R) develop late-onset obesity [10, 11].

Obesity is linked to numerous diseases including type 2 diabetes, cardiovascular events, and certain forms of cancer [12, 13]. Several antiobesity drugs, including diethylpropion, fenproporex, mazindol, fluoxetine, and sibutramine, have been developed [14], but there is little evidence for effective and safe treatments.

We have previously shown that subchronic peripheral Oxt treatment through a subcutaneously implanted osmotic minipump reduces hyperphagia and obesity in high-fat diet-fed obese mice, and that intraperitoneal (IP) Oxt treatment similarly reduces hyperphagia [15]. Zhang et al. [16] have recently reported that the nasal administration of Oxt decreases body weight (BW) in obese subjects. Ott et al. [17] showed that nasal Oxt treatment reduces reward-driven food intake. These reports indicate that subcutaneous, IP, and nasal routes of Oxt treatment have common actions of reducing obesity and hyperphagia. However, a comparison of these actions and underlying mechanisms following different administration routes of Oxt has not been previously undertaken.

We performed a comparative study on the effects of nasal and IP administration of Oxt on feeding, locomotor activity, and glucose tolerance in mice, and found that both nasal and IP Oxt injections reduce food intake to a similar extent with increased c-Fos-immunoreactive (ir) neurons in common brain regions. IP Oxt administration also increased c-Fos-ir neurons in additional brain regions, reduced locomotor activity, and affected glucose tolerance possibly by promoting insulin secretion from pancreatic islets.

Materials and Methods

Animals

Male C57BL/6J mice (aged 6 weeks) were obtained from Japan SLC (Hamamatsu, Japan). Animals were maintained on a 12-hour light/dark cycle. The dark and light phases started at 19:30 h and 7:30 h, respectively. Mice were allowed ad libitum access to water and a standard diet (CE-2; Clea, Osaka, Japan). Experimental procedures and care of animals were carried out according to the Jichi Medical University Institute of Animal Care and Use Committee. The mice used in the experiments were well habituated to minimize stress.

Measurements of Food Intake and Locomotor Activity after Oxt Administration

Following deprivation of food for 2 h before the dark phase, the animals received 0.1–10 μg/10 μl nasal Oxt or 40 and 400 μg/kg IP Oxt (Peptide Institute, Osaka, Japan) 30 min before the dark phase. For the nasal administration, 10 μl of Oxt or vehicle was dropped into the nasal cavity. Cumulative food intake was measured for the following 0.5, 1, 2, 6, and 24 h. For the measurements during the dark phase, red light with minimum intensity was used to prevent mice from being stimulated by bright light. Locomotor activity was measured by an activity monitoring system (ACTIMO-100; Shinfactory, Fukuoka, Japan). The doses of nasal Oxt were 0.1, 1 and 10 μg, based on the minimum effective dose of Oxt (1 μg) on food intake determined by pilot experiments. The dose of IP Oxt was based on a previous report [15] showing that 400 μg/kg Oxt markedly and long-lastingly suppressed food intake.

Immunostaining of c-Fos after Oxt Administration

Food was deprived for 2 h before the dark phase until perfusion. The animals received nasal (1 μg/10 μl) or IP Oxt (400 μg/kg) administration 30 min before the dark phase. After 90 min, the mice were transcardially perfused as described previously [7]. Coronal sections of 40-μm thickness were cut with a freezing microtome. Sections at 120-μm intervals between −0.58 and −8.0 mm from the bregma were used for c-Fos immunohistochemistry. Sections were rinsed in phosphate-buffered saline (PBS; 0.01 M, pH 7.4), incubated in PBS containing 2% normal goat serum and 2% bovine serum albumin (BSA), then incubated with rabbit anti-c-Fos antiserum (sc-52, 1:5,000; Santa Cruz, Calif., USA). Subsequently, the sections were incubated with biotinylated goat anti-rabbit IgG (1:500; Vector Laboratories Inc., Calif., USA), and an avidin-biotin complex (ABC kit; Vector Laboratories Inc.). Immunoreactions were visualized by incubating in diaminobenzidine (DAB) solution containing nickel ammonium.

Bilateral sections of the hypothalamus and brainstem between −0.58 and −8.0 mm from the bregma were used for c-Fos immunohistochemistry. Sections were rinsed in phosphate-buffered saline (PBS; 0.01 M, pH 7.4), incubated in PBS containing 2% normal goat serum and 2% bovine serum albumin (BSA), then incubated with rabbit anti-c-Fos antiserum (sc-52, 1:5,000; Santa Cruz, Calif., USA). Subsequently, the sections were incubated with biotinylated goat anti-rabbit IgG (1:500; Vector Laboratories Inc., Calif., USA), and an avidin-biotin complex (ABC kit; Vector Laboratories Inc.). Immunoreactions were visualized by incubating in diaminobenzidine (DAB) solution containing nickel ammonium.

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Glucose Tolerance Test and Measurement of Plasma Insulin

Mice aged 10 weeks (or with a BW of 22–25 g) were cannulated in the lateral ventricle (0.5 mm caudal to bregma, 1.0 mm lateral from the midline, and 2.2 mm below the skull surface; ICM-23G09 Inter Medical, Osaka, Japan) and housed individually. BW and food intake were not significantly different before the operation and for 10 days thereafter. On the day of the experiment, food was deprived at 09:00 h and an IP glucose tolerance test (IPGTT; 2 g/kg) was started at 13:00 h. IPGTT was performed 20 min after nasal (0.1, 1, 10 μg), icv (0.4, 4 μg) or IP (40, 400 μg/kg) Oxt injection. Oxt doses for icv injection were determined based on our previous report [7]. Blood was sampled by cutting the surface of the tail skin under local anesthesia with EMLA cream (AstraZeneca K.K., London, UK) without restraint. Blood glucose levels were measured using Glucocard (Arkray, Kyoto, Japan).

For the measurement of plasma insulin, mice aged 10 weeks (22–26 g BW) were used. IP Oxt injection (400 μg/kg) and IPGTT procedures followed the same protocol and time course. The mice were decapitated, and their blood was collected 10 min after IPGTT. After centrifugation, plasma insulin concentration was measured by ultrasensitive mouse insulin ELISA kit (Morinaga Institute of Biological Science, Inc., Yokohama, Japan).

Isolation of Islets and Measurement of Insulin Secretion

Mice with similar age and BW to those in the IPGTT were used. Islets were isolated as previously described [18] and cultured overnight in DMEM solution (Sigma, St. Louis, Mo., USA). After 30 min of preincubation in 2 mM glucose solution and 30 min under static incubation conditions in Krebs-Ringer Buffer (mM): 120 NaCl, 4.7 KCl, 2.5 CaCl₂, 1 KH₂PO₄, 1.2 MgSO₄, 10 HEPES, 25 NaHCO₃, pH 7.4 (with NaOH), plus 0.1% BSA, experimental glucose (2, 5, 20 mM) with or without Oxt (10 nM), insulin secretion was measured. Islets were then incubated overnight at –20°C with acidified ethanol solution (95% ethanol, 5% acetic acid) to extract all insulin. Insulin was measured using a Morinaga mouse ELISA kit (Morinaga Institute of Biological Science, Inc.).

Statistical Analysis

All data are presented as means ± SEM. The statistical analysis of experimental data of c-Fos-ir neurons, insulin secretion, and plasma insulin was carried out using Student’s t test. Data for food intake, locomotor activity, and glucose tolerance test were analyzed by repeated measures of two-way ANOVA with treatment (saline vs. Oxt including dose responses) and time as factors. Post hoc multiple comparisons were made using Tukey’s test. Significance was set at p < 0.05 for all analyses.

Results

Effect of Oxt Administration on Food Intake and Locomotor Activity

In order to compare the effects of nasal and IP Oxt administration, we administered an approximately equally effective amount of Oxt for nasal and IP administration (nasal: 0.1–10 μg/25 g BW, i.e. approx. 4–400 μg/kg; IP: 40–400 μg/kg). Nasal administration of Oxt at doses of 0.1–10 μg/10 μl was performed in mice weighing 20–25 g. These doses correspond to approximately 4–400 μg/kg.

Nasal administration of 0.1 μg Oxt did not affect the food intake significantly (F₁,₆₀ = 2.53, p > 0.05; fig. 1a). Administration of 1 μg Oxt significantly decreased the food intake (F₁,₅₆ = 27.4, p < 0.01; fig. 1b), and 10 μg Oxt (F₁,₉₂ = 27.16, p < 0.01; fig. 1c). Tukey’s test indicated that nasal administration of 1 μg Oxt reduced the cumulative food intake 6 and 24 h after administration (fig. 1b), and 10 μg Oxt reduced the food intake 0.5, 1, 2, 6, and 24 h after administration (fig. 1c) without changing locomotor activity (F₁,₂₉₉ = 0.10, p > 0.05; fig. 2a, c).

IP administration of 40 μg/kg Oxt failed to significantly affect the food intake (F₁,₂₄ = 2.27, p > 0.05; fig. 1d) and 400 μg/kg Oxt markedly decreased the food intake at 0.5, 1, 2, and 6 h (F₁,₂₄ = 38.9, p < 0.01; fig. 1e) but not at 24 h. IP injection of 400 μg/kg Oxt showed a tendency to decrease locomotor activity (F₁,₁₁₈ = 2.6, p > 0.05) for the first 7 h and significantly decreased cumulative locomotor activity during the dark phase (fig. 2b, d).

The tendency and time course of reduction in the food intake were similar between nasal and IP administration (fig. 1b, d). However, at a higher dose (approx. 400 μg/kg), nasal administration of Oxt (10 μg/μl; fig. 1c) reduced the cumulative food intake for up to 24 h, but the extent of food intake reduction during 0.5–6 h was smaller compared to IP administration (fig. 1e).

Effect of Oxt Administration on the Number of c-Fos-ir Neurons

After nasal administration of 1 μg Oxt, the number of c-Fos-ir neurons was significantly increased in the PVN of the hypothalamus. The number was 137.1 ± 25.6/sec- tion in controls versus 284.6 ± 37.4/sec in Oxt-injected groups (p < 0.05; fig. 3a, b, e). Nasal Oxt administration also increased the number of c-Fos-ir neurons in the medulla, the area postrema (AP; 10.6 ± 3.5/sec in controls vs. 27.3 ± 4.7/sec in Oxt-injected groups), and the dorsal motor nucleus of vagus (DMNV; 32.3 ± 6.9/sec in controls vs. 69.2 ± 4.7/sec in Oxt-injected groups; fig. 3c–e). IP administration of Oxt also increased the number of c-Fos-ir neurons in the PVN (71.7 ± 18.1/sec in controls vs. 188.0 ± 35.2/sec in Oxt-injected groups), DMNV (14.7 ± 3.7/sec in controls vs. 43.4 ± 4.8/sec in Oxt-injected groups; fig. 3f, g, j), nucleus of the solitary tract (NTS; 39.5 ± 9.8/sec in controls vs. 153.3 ± 6.0/sec in Oxt-injected groups), and the arcuate nucleus (ARC; 68.9 ± 12.2/sec in controls vs. 128.6 ± 7.0 in Oxt groups), consistent with our previous...
Thus, IP but not nasal administration of Oxt increased c-Fos-ir neurons in the NTS and ARC.

**Effect of Oxt Administration on Glucose Metabolism**

The effect of Oxt on the elevated levels of blood glucose during IPGTT was examined. Nasal and icv Oxt administration had no effect on blood glucose levels in IPGTT (F_{3,84} = 0.28, p > 0.05, and F_{2,54} = 1.02, p > 0.05, respectively; fig. 4a, c). In contrast, IP administration of 40 and 400/μg/kg Oxt (F_{2,36} = 13.10, p < 0.01) dose-dependently attenuated the elevation of blood glucose at 30 and 60 min of IPGTT (fig. 4b). The alteration of glycemia prompted us to examine whether Oxt affects insulin release from pancreatic islets. Ten minutes after IPGTT, plasma insulin concentration was significantly increased (2.19 ± 0.26 ng/ml), compared with the saline-injected group (0.93 ± 0.12 ng/ml; fig. 4d). Incubation of isolated islets with 10 nM Oxt-enhanced insulin secretion in the stimulatory (20 mM), but not in the low (2 mM) and basal (5 mM) glucose conditions (fig. 4e).

**Discussion**

It has recently been reported that nasal administration of Oxt reduces BW in obese humans [17]. To the best of our knowledge, the present study is the first to show the effect of nasal Oxt in reducing food intake in rodents without affecting locomotor activity or blood glucose level. This may suggest a selective anorexigenic action of nasal treatment of Oxt.

Neumann et al. [19] reported the increase of brain and plasma Oxt concentrations after nasal administration in rats and mice. The dose (12 μg/10 μl) and application method used in their study are almost identical to those used here (10 μg/10 μl). Hence, we speculate that Oxt levels in the brain and plasma may have been increased after nasal administration of Oxt in the present study. Nasal administration of Oxt may lead to its inhalation into the lungs [20], thereby increasing the plasma Oxt level. If so, pharmacokinetics of Oxt may be similar between nasal and IP applications. Hence, the similar anorexigenic effects of...
nasal and IP Oxt treatments shown in the present study may have been mediated largely by the increase of peripheral Oxt concentration. In addition, Oxt is reported to cross the blood-brain barrier (BBB) from the blood to the brain by 0.05% [21]. The permeation of Oxt through the BBB depends on the concentration of Oxt in vessels. Since our study used a high dose of Oxt via nasal and IP routes, a significant amount of the peptide may have reached the central nervous system by crossing the BBB [3, 22].

On the other hand, this study has shown that both nasal and IP administration of Oxt induces c-Fos in PVN, AP, and DMNV. In addition, only IP administration of Oxt induces c-Fos in ARC and NTS.

In apparent discrepancy with our results, nasal administration of Oxt (1 μg) has been reported to have no effect on c-Fos-ir neurons in rats [23]. However, this discrepancy may be explained by the dose of Oxt. In the study by Ludwig et al. [23], the dose of Oxt used was 1 μg for rats (BW: 250–300 g). In the present study, it was 1 μg for mice (BW: 25–30 g), a factor 10 times higher when corrected for BW. Oxt doses that induced c-Fos in the present study correspond to 1–100 times of the total Oxt content of ap-
proximately 150 ng in the mouse pituitary [24]. Hence, it is suggested that a relatively high dose of nasal Oxt is required to induce c-Fos expression. In addition, the present study used a higher dose of Oxt (approx. 10 μg nasal, approx. 400 μg/kg IP) than that used in a human study (approx. 48 μg nasal) [17]. The different dose requirement between humans and rodents is possibly due to different anatomical structures and/or to the different approach for nasal Oxt administration between humans (vaporizer or nasal spray) and mice (dropping the Oxt-containing solution on the nose by a pipette).

In this study, Oxt administration via the nasal versus the IP route increased the number of c-Fos-ir neurons in both common and distinct brain areas. Both nasal and IP administration of Oxt increased c-Fos-ir neurons in the PVN, AP, and DMNV. The increase in the PVN may underlie the common action of nasal and IP Oxt administration to suppress food intake, since PVN is recognized as
Fig. 4. Effect of Oxt on blood glucose levels in glucose tolerance test and insulin levels. a–c Blood glucose profiles in IPGTT after Oxt administration via nasal (0.1, 1, and 10 μg; n = 6–11; a), IP (40 and 400 μg/kg; n = 5; b) and icv administration (0.4 and 4 μg; n = 5–8; c). Differences were assessed by repeated measures two-way ANOVA with Oxt treatment and time as factors. For each time point, post hoc multiple comparisons were made using Tukey’s test. * p < 0.05, ** p < 0.01 versus control. d Plasma insulin levels after IP administration of Oxt. ** p < 0.01. e Insulin secretion from isolated pancreatic islets under static incubation with low (2 mM), basal (5 mM), and stimulatory (20 mM) glucose concentrations. Open bars indicate the control group (n = 11) and solid bars indicate the Oxt-treated group (n = 11). Differences were assessed for significance by Student’s t test. * p < 0.05.
an integrative center for feeding and energy metabolism [25, 26]. The neurons in the ARC of the hypothalamus sense various nutrients and hormones and convey neural information to the neurons in PVN, where peripheral and central information is integrated [27]. In contrast, only IP Oxt increased c-Fos-ir neurons in the ARC and NTS. Several pathways may be considered for the common and distinct brain signalling by nasal and IP Oxt. First, given that the cribriform plate is the only structure separating the nasal cavity from the olfactory bulb in rodents, nasal Oxt administration may activate the Oxt-R present in the nasal cavity from the olfactory bulb [28, 29]. Second is the route mediated by the AP. c-Fos-ir neurons were increased in the AP after both IP and nasal application of Oxt. The AP is known to have a leaky BBB, and the neurons in the AP project to NTS neurons, which in turn project to the ARC and PVN [30]. Hence, it is possible that nasal and IP injections of Oxt exert their effects on feeding via activating neurons in AP. However, in this study Oxt increased c-Fos-ir neurons in the NTS only after IP injection and not after nasal administration. This may indicate that neural signal transduction through the AP to NTS projection after nasal application of Oxt was not intensive enough to induce c-Fos-ir neurons in the NTS. Third is the route via vagal afferents. In this study, only IP administration of Oxt increased the number of c-Fos-ir neurons in the NTS, where the vagal afferent neurons terminate [31]. Oxt-R is widely expressed in the peripheral tissues including the nodose ganglion [32]. Hence, it is possible that IP-administered Oxt interacts with Oxt-R on the vagal afferent nodose ganglion neurons to signal to the NTS.

Regarding the reduced locomotor activity found only with IP administration of Oxt, the underlying mechanisms could be due to the passage of Oxt from the blood to the brain, thereby activating responsible brain regions, or to the consequences of the peripheral actions of very high levels of Oxt following IP administration. It is important to note that a high peripheral Oxt concentration can also activate vasopressin receptors that have been reported to regulate locomotor activity [33]. Further studies are required to elucidate the mechanism underlying the effect of IP Oxt administration on locomotor activities.

In the present study, nasal and IP administration of Oxt increased the number of c-Fos-ir neurons in the PVN of male mice, suggesting a possible peripheral-central Oxt axis. The result is in agreement with a recent report that the IP administration of Oxt activates PVN Oxt neurons in male rats [34]. Negative feedback effects of Oxt on Oxt neurons were reported under particular conditions in virgin and pregnant female rats, in contrast to positive feedback effects in male and lactating female rats [35–37]. These reports suggest that Oxt activates Oxt neurons in male rodents. Hence, our results could reflect the direct effects of exogenously administered Oxt on PVN Oxt neurons or indirect effects mediated by the nasal/IP Oxt-activated neurons that project to the PVN Oxt neurons.

We have previously reported that chronic Oxt treatment ameliorates the impaired glucose tolerance in high-fat diet-induced obese mice [15]. Since chronic Oxt treatment also ameliorates obesity, the improvement of glucose tolerance by Oxt could be due to its direct effect on glucose metabolism or secondary to the amelioration of obesity. In the present study, IP administration of 40 μg/kg Oxt induced a small effect on food intake but markedly decreased blood glucose levels at 30 min of IPGTT. Furthermore, Oxt elevated plasma insulin level in vivo and enhanced glucose-induced insulin secretion in isolated islets in vitro. These results taken together suggest that IP-administered Oxt substantially reaches the pancreas and directly interacts with islet β-cells to enhance glucose-induced insulin release, thereby improving glucose tolerance. Insulin, released postprandially [38], might enter the brain and inhibit food intake [27]. It was also reported that insulin activates vagal afferent neurons, which could be linked to the regulation of food intake [39]. Therefore, it is possible that insulin secretion induced by IP Oxt may partly contribute to the anorexigenic effect of IP Oxt.

In our study, nasal Oxt administration failed to affect glucose tolerance in mice. In dogs, in contrast, nasal Oxt at a similar dose (3–4 μg/kg) was reported to increase plasma insulin and glucagon levels [40]. Also, in humans, nasal administration of Oxt (24 IU) attenuated the elevated blood glucose following the test buffet [17]. The apparent discrepancy in the effect of nasal Oxt on glucose homeostasis could be due to species differences or methodology of treatment as stated above. The nasally administered Oxt might be more easily transferred to the periphery in dogs and humans than in mice.

In order to assess the direct cerebral effect of Oxt on glucose metabolism, we performed the IPGTT after icv injection of Oxt. icv injection of Oxt (0.4, 4 μg/mice) had no effect on glucose metabolism. Although regulation of glucose metabolism through the brain is established, whether this process may be modulated by Oxt is not clear [41, 42]. Our results suggest that Oxt in the brain has little effect on glucose metabolism under the experimental conditions studied here.
When considering the physiological aspect of Oxt on females, it shows different effects on food intake under different conditions. In the preparturition period, Oxt secretion is very low in order to reduce the risk of preterm delivery and to prevent the loss of the accumulating neurophysial Oxt store [43], which is accompanied by an increase in food intake in both mice and rats [42, 44]. During the parturition and lactation periods, in spite of the extensive activation of Oxt neurons, food intake is dramatically elevated [44]. It is suggested that the involvement of Oxt in feeding behavior is unique in these periods when the energy intake and expenditure are in different states from those in male or virgin animals.

Recent reports have shown that Oxt treatment reduces reward-driven food intake [17]. It is well known that the reward process, including feeding, sexual [45] and social [46] rewards, is related to the mesocorticolimbic dopamine system in the brain [47, 48]. Oxt appears to impact this system [47], in which dopamine is one of the neurotransmitters playing a major role in addiction. It has recently been reported that Oxt modulates the negative aspect of addiction [49] and that Oxt treatment can attenuate the addictive cocaine-seeking behavior in humans [50]. Hyperphagia is considered an addictive behavior for food. The common mechanisms for hyperphagia and addiction have been reported [51]. Therefore, Oxt as a neuropeptide may serve to keep the reward system at a moderate level.

In conclusion, nasal and IP administration of Oxt have similar anorexigenic effects. However, nasal administration of Oxt may have a lesser effect on the peripheral tissue, including pancreatic β-cells, compared to IP administration. Hence, nasal administration may have the advantage of activating specific brain regions without affecting glucose tolerance and locomotion.

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Disclosure Statement

The authors have no conflicts of interest to disclose.

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