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The effects of two highly selective dopamine D₃ receptor antagonists (SB-277011A and NGB-2904) on food self-administration in a rodent model of obesity

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Abstract

In the current study, we examined the effect of the selective D₃ receptor antagonists SB-277011A and NGB 2904 on operant food self-administration (FSA) in Zucker obese and lean rats. Obese (Ob) and lean (Le) Zucker rats were maintained under a restricted feeding regimen (70% of ad-libitum rat chow) and were trained to lever press for food during daily, 2 hour fixed-ratio 4 (FR4) schedules. Once rats reached a stable baseline for FSA, they were injected with vehicle until a stable FSA criterion was achieved. Animals then received daily injections of different random doses of SB-277011A (3, 10, and 30 mg/kg i.p.), and NGB-2904 (0.3, 1 and 3 mg/kg i.p.). SB-277011A produced a significant decrease in both food intake and active lever responses in both Ob and Le rats. In contrast, NGB-2904 did not decrease food intake levels or lever presses for food in Ob and Le rats. These results suggest that along with its involvement in seeking behavior for drugs of abuse, the D₃ dopamine receptor may also be involved in seeking behavior for natural reinforcers such as food.

Keywords

Hunger; Ingestive behavior; Operant conditioning; Anorexia; Anorexigenic

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1. Introduction

Food consumption is one of the most widely studied reward survival mechanisms (Fetissov et al., 2002). The overconsumption of food is one of the major factors that has contributed to a significant increase in the incidence of obesity (Erlanson-Albertsson and Zetterstrom, 2005; Isasi et al., 2006; Hedley et al., 2004; Ogden et al., 2002). Although the pathophysiology of obesity remains to be elucidated, it is generally thought to result from a dysregulation in energy homeostasis which is mediated in part through food intake regulation (Cecil et al., 2006). It has been shown that various peptides, hormones and brain neurotransmitters are involved in food intake regulation (Erlanson-Albertsson and Zetterstrom, 2005). Dopamine (DA) is known to regulate a broad range of biological functions such as locomotor activity, cognition, food intake, and hormone secretion (Palmiter, 2007; Pijl, 2003) as well as reinforcement of addictive substances and behaviors (Di Chiara, 2002; Di Chiara and Bassareo, 2007). The dopaminergic perikarya from the ventral tegmental area (VTA) send projections that innervate a number of limbic and telencephalic structures, including the olfactory tubercle, amygdala, frontal and limbic cortices, medial prefrontal cortex (mPFC), and the nucleus accumbens (NAc), a brain area that plays a significant role in reward-related behavior and reward learning (Day and Carelli, 2007; Di Chiara and Bassareo, 2007; Fenu and Di Chiara, 2003). DA plays an essential role in food-seeking behavior (Wise, 2006a,b), reward prediction (Phillips et al., 2007; Schultz et al., 2000), reward motivation (to seek and obtain reward) (McClure et al., 2003; Phillips et al., 2007) and facilitation of conditioned learning (Fenu and Di Chiara, 2003). Since DA has been shown to mediate food intake through brain reward mechanisms (McQuade et al., 2004), DA modulation may be responsible for regulating the reward or reinforcement necessary to maintain, enhance or attenuate feeding behavior (Berridge, 1996).

The hormone leptin is secreted by adipocytes (Hagan et al., 1999) and is an important signal in the regulation of energy balance (Hagan et al., 1999). Rodents and humans with homozygous mutations in the leptin or leptin receptor genes manifest hyperphagia and severe obesity (Hagan et al., 1999). Zucker obese (fa/fa) rats are widely used as animal models in obesity research due to their phenotypic similarities with obese humans such as metabolic and cardiopulmonary deficits (Brooks-Asplund et al., 2002); type 2 diabetes mellitus (Hunt et al., 1976), hypertension (Alonso-Galicia et al., 1996), upper airway narrowing and poor exercise capacity (Dockstader et al., 2001; Lee et al., 2005). These rats also exhibit hyperphagia, hypertriglycemia, and hyperinsulinemia (Boulange et al., 1979). Their genotype (fa/fa) denotes that they are homozygous for defective leptin receptors, which prevents leptin signaling in these animals (Malcher-Lopes et al., 2006).

It has previously been shown that anorectic concentrations of leptin in the brain reduces the firing of mesolimbic DAergic neurons *in vivo* and suppresses DA-related motivational aspects of feeding (Krugel et al., 2003) which may indicate decreased DA levels in leptin receptor deficient obese Zucker (fa/fa) rats as one of the causes for obesity (Brunetti et al., 1999). Furthermore, in mice deficient in both DA and leptin, DA was shown to be required for the initiation of food intake in the absence of leptin (Szczyepka et al., 2000). Leptin receptors have been found to be extensively co-expressed within DA neurons of the ventral tegmental area (VTA) and substantia nigra pars compacta (SNc) (Figlewicz et al., 2003) suggesting that DA release in relation to food intake is modulated by leptin. We recently reported that leptin receptor deficient obese Zucker rats display lower D2 receptor (D₂) levels than lean rats and that these levels were modulated by food restriction (Michaelides et al., 2006; Thanos et al., 2007). In addition, microPET assessment of D₂ receptor density in these rats suggest that D₂ receptor availability is differentially influenced by food restriction in obese as compared to lean rats, (Michaelides et al., 2006; Thanos et al., 2007).

In rats, D₃ receptors and D₃ receptor mRNA is predominantly located in limbic brain regions such as the NAc, the islands of Calleja, and the VTA and SNc (Booze and Wallace, 1995; Bouthenet et al., 1991; Curran and Watson, 1995; Diaz et al., 1995, 2000; Levant, 1997; Shafer and Levant, 1998; Sokoloff et al., 1990). In humans D₃ receptors have been localized in the NAc, internal globus pallidus, ventral pallidum, septum, islands of Calleja, amygdala, and VTA (Landwehrmeyer et al., 1993; Meador-Woodruff et al., 1994; Suzuki et al., 1998). Until recently, it has been difficult to characterize the functional role of D₃ receptors in the CNS due to the lack of selective D₃ receptor agents. This obstacle has been surmounted by the synthesis of selective D₃ receptor antagonists. SB-277011A, a selective D₃ receptor antagonist, has an 80- to 100-fold selectivity for the D₃ receptor compared to other DA receptors and has a high affinity for human (pKi 7.95) and rat (pKi 7.97) cloned D₃ receptors compared to D₂ receptors (pKi=5.98). In addition, a number of studies indicated that SB-277011A significantly decreases the rewarding/reinforcing actions of various drugs of abuse as well as drug-seeking behavior. In rats, SB-277011A has been reported to 1) attenuate cue-induced cocaine self-administration behavior (Di Ciano and Everitt, 2003; Vorel et al., 2002); 2) block the expression of cocaine and heroin-induced conditioned place preference (CPP) (Ashby et al., 2003; Vorel et al., 2002); 3) attenuate alcohol intake in ethanol preferring rats (Heidbreder et al., 2007; Thanos et al., 2005) and 4) suppress alcohol self-administration, and prevent reinstatement of alcohol-seeking in mice (Heidbreder et al., 2007; Thanos et al., 2005). In addition, SB-277011A significantly attenuated nicotine-triggered relapse to nicotine seeking (Andreoli et al., 2003; Pak et al., 2006) as well as sucrose-seeking behavior induced by sucrose-associated cue reintroduction (Cervo et al., 2007).

Another selective D₃ antagonist is NGB-2904, which has 155-fold selectivity for the primate D₃ over the D₂ and over 800-fold selectivity for the rat D₃ (Newman et al., 2003). In rats, NGB-2904 dose-dependently decreased cocaine self-administration behavior (Xi et al., 2006; Gilbert et al., 2005) and significantly attenuated the cocaine-induced leftward shift in brain stimulation reward paradigm (BSR). Finally, while NGB-2904 pretreatment was shown to decrease cocaine triggered reinstatement of cocaine-seeking behavior, it had no effect on sucrose-plus-sucrose-cue-triggered reinstatement of sucrose-seeking behavior (Xi et al., 2006).

To our knowledge, there are no published reports regarding the effects of highly selective D₃ antagonists (such as SB-277011A and NGB-2904) in an operant FSA paradigm in obese rats. The present study consisted of two experiments that aimed at examining the effects of these two selective D₃ antagonists SB-277011A (Experiment 1) and NGB-2904 (Experiment 2) in Zucker obese and lean rats in the FSA task. We chose to utilize the obese (fa/fa) Zucker rat based on its severe hyperphagia and weight gain. These characteristics, coupled with the leptin receptor mutation provide an original examination of changes in food-seeking and FSA behavior.

2. Materials and methods

2.1. Experiment 1

2.1.1. Animals—Eight-week old Zucker obese (fa/fa) ($n=6$; 372 ± 15.4 g) and lean (Fa/?) rats ($n=6$; 257 ± 12.3 g) were purchased from Charles River Laboratories (Wilmington, MA). Each animal was allowed to acclimate for 7 days after arriving at the Brookhaven National Laboratory Animal Facility. Animals were restricted to 15 g of Purina Rodent Chow per day with ad-libitum water access. Animals were individually housed in standard 9 in. \times 24 in. \times 9 in. plastic cages with wire covers. The home cage environment was set at 22 ± 1 °F with approximately 60% humidity and reverse 12 h/12 h light/dark cycle with lights off at 0700 h and on at 1900 h. Ad-libitum access to water was maintained but access to food was restricted to 2 h daily during operant sessions throughout the entire experiment. All animals received

food supplementation following each operant session to maintain them on a limited food access diet of 70% of the food consumed by two similarly aged and ad-libitum fed groups of lean and obese rats. On average, lean rats consumed 13.2 g/day and obese rats consumed 22 g/day. Experiments were conducted in conformity with the National Academy of Sciences Guide for the Care and Use of Laboratory Animals (NAS and NRC, 1996) and Brookhaven National Laboratory Institutional Animal Care and Use Committee protocols.

2.1.2. Drugs—SB-277011A (Reavill et al., 2000; Bull et al., 2000) (trans-N-[4-[2-(6-cyano-1,2,3,4-tetrahydroisoquinolin-2-yl)ethyl] cyclohexyl]-4-quinolinecarboxamide) was obtained from GlaxoSmithKline (Harlow, Essex, U.K.). A 3% methylcellulose w/v (M-7140 Sigma) and 0.9% saline solution was used as the vehicle. All rats were injected with 3, 10 and 30 mg/kg of SB-277011A which was prepared in concentrations of 3, 10, and 30 mg/ml for physiological use in Zucker obese (fa/fa) and Zucker lean (Fa/?) rats.

2.2. Experiment 2

2.2.1. Animals—Eight-week old Zucker obese (Ob) (fa/fa) ($n=6$; 365 ± 17 g) and lean (Le) (Fa/?) rats ($n=6$; 243 ± 10 g) (Charles River Laboratories) were used in this study. All other details were the same as stated above for Experiment 1.

2.2.2. Drugs—NGB 2904 *N*-(4-[4-{2,3-dichlorophenyl}-1-piperazinyl] butyl)-2-fluorenylcarboxamide was synthesized in the Medicinal Chemistry Section, NIDA-IRP. All rats were injected with 0.3, 1 and 3 mg/kg of NGB-2904 (0.3, 1 and 3 mg/ml solutions) which was dissolved in 25% 2-hydroxy-propyl- β -cyclodextrin (w/v); (vehicle solution (Sigma/RBI, Saint Louis, MO)).

2.3. Experiments 1 and 2

2.3.1. Apparatus—Both experiments utilized the 45 mg dustless precision food pellets (Prod#: F0021; Bio-Serv Inc.; Frenchtown, NJ) throughout the operant conditioning task. Clear acrylic operant test chambers measuring 32 \times 25 \times 33 cm were used (Coulbourn Instruments, Allentown, PA). Each test cage was enclosed in an environment isolation chamber to keep out any outside environmental stimuli. Cage floors were constructed of stainless steel horizontal bars spaced 2 cm apart. The test cages were equipped with two randomly assigned and counterbalanced response levers; a reinforced (R) and a non-reinforced (NR) lever, with cue lights located above each. R lever presses led to a delivery of a food pellet. A lever press on the NR lever had no consequences. All data from the test chamber was recorded using Graphic State software version 3.0.

2.3.2. Procedures—Sessions were run daily for 2-h per day. Animals were started on a fixed ratio 1 (FR1, 1 lever press/1 pellet dispensed) schedule of reinforcement and were increased by one FR level until reaching FR4. Upon completion of each session, animals were promptly returned to their home cage. This schedule was used to train (the rats to press the lever) and feed the rats until baseline criteria ($\leq 20\%$ change in daily food intake for five consecutive sessions) were met. After completion of the food training protocol, each rat was injected intraperitoneally (i.p.) with approximately 0.3 ml of the vehicle 15 min prior to the operant sessions under the FR4 schedule that included a 30 second time-out period until the vehicle baseline criteria (daily food intake $\leq 20\%$ for three consecutive sessions) was met. Upon meeting the vehicle baseline criteria, all the rats in Experiment 1 continued on the same protocol using a randomized Latin square design of SB-277011A administration (3 mg/kg; 10 mg/kg and 30 mg/kg for three consecutive sessions at each dose). The doses of SB-277011A were selected based on previous studies (Cervo et al., 2007; Heidbreder et al., 2007; Thanos et al., 2005). For Experiment 2, the similar randomized design was used for NGB-2904 after vehicle criteria was achieved (0.3 mg/kg; 1 mg/kg and 3 mg/kg). The doses of NGB-2904 were based

on data indicating that they produced significant effects in other behavioral paradigms (0.1, 1 and 5 mg/kg dose-dependently inhibited cocaine SA) (Xi et al., 2004, 2006). One day after the last drug treatment of SB-277011A for Experiment 1 or NGB-2904 for Experiment 2, rats received vehicle for three days.

2.3.3. Data analysis—Food intake (number of pellets), lever presses, were collected during each experimental session. All data was analyzed using a Two-way Repeated Measures Analysis of Variance (RM ANOVA), followed by pairwise multiple comparison using the Holm–Sidak method. The vehicle data represent the average of the first (baseline), intermittent (between treatments) and last (after the last drug treatment) vehicle sessions.

3. Results

3.1. Experiment 1

3.1.1. Food Intake—SB-277011A effects on food consumed are shown in Fig. 1. A two-way repeated measures ANOVA revealed significant differences for strain [$F(1,143)=27.619$; $p<0.001$] and treatment [$F(3, 143)=34.501$; $p<0.001$] but no significant interaction effect. Subsequent multiple pairwise comparisons (Holm–Sidak; $p<0.05$) revealed that in Le rats food intake was significantly lower at 30 mg/kg SB-277011A treatment compared to all other treatments [vehicle ($t=7.191$), 3 mg/kg ($t=7.010$), 10 mg/kg ($t=6.497$)]. In Ob rats food intake was also significantly lower at 30 mg/kg SB-277011A treatment compared to all other treatments [vehicle ($t=5.136$), 3 mg/kg ($t=7.007$), 10 mg/kg ($t=2.913$)]. Also, 10 mg/kg SB-277011A treatment in Ob rats decreased food intake compared to 3 mg/kg ($t=4.094$). Finally, Ob rats consumed more food than Le rats at the 3 mg/kg ($t=4.697$) and 30 mg/kg ($t=4.700$) SB-277011A doses.

3.1.2. Reinforced (R) and non-reinforced (NR) lever responses—We examined the mean number of R lever responses using a two-way repeated measures ANOVA (Fig. 2) which revealed a significant main effect of strain [$F(1, 143)=10.917$; $p=0.004$] and SB-277011A treatment [$F(3, 143)=30.288$; $p<0.001$] but no interaction effect. Subsequent multiple pairwise comparisons (Holm–Sidak; $p<0.05$) revealed that in Le rats R lever responses were significantly lower at 30 mg/kg SB-277011A treatment compared to all other treatments [vehicle ($t=7.021$), 3 mg/kg ($t=7.296$), 10 mg/kg ($t=4.980$)]. In Ob rats R lever responses were also significantly lower at 30 mg/kg SB-277011A treatment compared to all other treatments [vehicle ($t=4.479$), 3 mg/kg ($t=5.898$), 10 mg/kg ($t=3.639$)]. Finally, Ob rats elicited a greater number of R lever responses than Le rats only at the 30 mg/kg dose of SB-277011A ($t=3.008$).

A two-way repeated measures ANOVA did not reveal any significant main effects (strain [$F(1, 143)=2.732$; $p=0.117$], treatment [$F(3, 143)=3.556$; $p=0.021$], interaction [$F(3, 143)=0.142$; $p=0.935$]) on the number of NR lever responses in response to SB-277011A treatment (Fig. 2).

3.2. Experiment 2

3.2.1. Food intake—NGB-2904 effects on food intake are shown in Fig. 3. A two-way repeated measures ANOVA revealed significant differences for treatment [$F(3, 143)=4.369$; $p=0.008$], interaction [$F(3, 143)=5.094$; $p=0.004$] but no significant strain effect. Subsequent multiple pairwise comparisons (Holm–Sidak; $p<0.05$) revealed that in Ob rats food intake was significantly greater at 3 mg/kg NGB-2904 treatment compared to all other treatments [vehicle ($t=3.936$), 0.3 mg/kg (3.845), 1 mg/kg ($t=3.802$)]. NGB-2904 did not affect food intake in Le rats since we did not detect any significant differences as a function of NGB-2904 treatment. Finally, at 3 mg/kg Ob rats consumed significantly more food compared to Le rats ($t=3.443$).

3.2.2. R and NR lever responses—We examined the mean number of R lever responses using a two-way repeated measures ANOVA (Fig. 4) which revealed a significant main effect of strain [$F(1, 143)=8.662$; $p=0.009$], treatment [$F(3, 143)=4.221$; $p=0.01$] as well as interaction [$F(3, 143)=3.865$; $p=0.01$] effects. Subsequent multiple pairwise comparisons (Holm–Sidak; $p<0.05$) revealed that in Ob rats R lever responses were significantly greater at 3 mg/kg NGB-2904 treatment compared to all other treatments [vehicle ($t=3.313$), 0.3 mg/kg ($t=2.940$), 1 mg/kg ($t=3.406$)]. NGB-2904 did not affect R lever responses in Le rats since we did not detect any significant differences. Finally, at 1 mg/kg ($t=4.190$) Le rats elicited significantly more R lever responses compared to Ob rats.

A two-way repeated measures ANOVA did not reveal any significant main effects (strain [$F(1, 143)=0.391$; $p=0.540$], treatment [$F(3, 143)=2.256$; $p=0.093$], interaction [$F(3, 143)=1.997$; $p=0.126$]) on the number of NR lever responses in response to NGB-2904 treatment (Fig. 4).

4. Discussion

4.1. SB-277011A

The role of the D₃ receptor in obesity and feeding-related behaviors has not been elucidated. One study found that male D₃ receptor deficient mice become obese when fed a high fat diet for a period of three months but did not gain weight when fed regular chow for the same amount of time (McQuade et al., 2004). Quite surprisingly, the increases in weight gain and adiposity were reported independently of increased food intake. The same study also reported that male D₃ knockout mice showed increased plasma concentrations of leptin, but not insulin in response to the high fat diet and not the chow. Subsequently, the authors concluded that the D₃ receptor is involved in the regulation of body weight and body fat when consuming diets differing in palatability and fat content. Although a gene-knockout model of D₃ inactivation does not necessarily parallel pharmacological studies of D₃ inactivation, the above finding is very important in providing insight into the potential involvement of the D₃ in reward and reward-seeking.

The effect of SB-277011A on food intake in obese animals has not been previously reported. One study in non-obese animals examined the effects of SB-277011A on drinking sucrose solution and showed no effect on R lever responses (Di Ciano et al., 2003). Similarly, a recent experiment (again in non-obese rats) showed that SB-277011A (10 mg/kg) did not modify conditioned reinstatement triggered by sucrose pellet-associated cues (Cervo et al., 2007). SB-277011A was also shown to have no effect on the number of pellets earned or the response rate in a food self-administration experiment conducted under a progressive-ratio (PR) schedule in non-obese rats even though it was reported that two out of eight rats tested showed a greater than 50% reduction in food intake at the 30 mg/kg dose (Ross et al., 2007).

Contrary to these previous studies, the present study is the first to examine the effects of SB-277011A on obese Zucker rats in an operant FSA task. The current results demonstrated that SB-277011A attenuated the amount of food consumed and R lever presses in both Ob and Le Zucker rats. Specifically, both Ob and Le rats exhibited a decrease in the amount of food consumed during the high (30 mg/kg) doses as compared to vehicle (Fig. 1). Both groups also showed significantly lower R lever responses at the highest dose (30 mg/kg) than at any other treatment. At this same dose, Ob rats showed greater R lever responses than their Le counterparts. SB-277011A administration did not interfere with R vs. NR lever response discrimination since NR lever responses comprised only a small percentage of total lever responses.

Discrepancies between this study and previous experiments that have assessed the effect of SB-277011A on food intake include: a) animal strain (Zucker obese rats used here versus non-obese rats in other studies); b) task (operant FR4 FSA task versus other FR and variable-interval tasks); c) age of the animals (adolescent animals used here versus adult animals in other studies); d) solid versus liquid food (solid food pellet used here versus liquid sucrose solution used in other studies); e) palatability of food (regular chow pellets versus sucrose or high fat pellets) and f) caloric intake value. It may be possible that the Zucker Ob rat is more sensitive to D₃ receptor antagonism than non-obese rats and that leptin (leptin receptor is deficient in Zucker Ob rats) may regulate this effect. Thus, while the present results support that SB-277011A may be valuable in reducing food intake in obese rats; future experiments assessing D₃ levels in these obese rats as well as the role of the D₃ should also examine among other things the interaction with leptin on food intake and weight gain.

4.2. NGB-2904

NGB-2904 showed no attenuation of R lever presses or food intake in the Ob or Le rats. Previous studies have shown that NGB-2904 did not affect the reinstatement response to sucrose in rats (Xi et al., 2006) or food-maintained operant responding in rhesus monkeys (Martelle et al., 2007). In our study, NGB-2904 had a minimal effect on operant FSA. Specifically, the only significant differences found was at the high dose (3 mg/kg) where Ob rats consumed more food than the Le rats. This was a similar effect to that observed with 3 mg/kg SB-277011A. While higher doses (>3 mg/kg) of NGB-2904 were not tested in this study future investigation on this is warranted. NGB-2904 did not have an effect on R lever presses except that at the highest dose tested (3 mg/kg) where there was no longer a difference between Ob and Le. No significant differences for NR responses were observed with NGB-2904.

Although both SB-277011A and NGB-2904 are highly selective for the D₃, SB-277011A may not be as selective as NGB-2904 in rats (Newman et al., 2003). Nevertheless, such a comparison has not been thoroughly evaluated as the binding profile of SB-277011A has been extensively examined (specifically, it has been shown that SB-277011A has selectivity for the D₃ receptor compared to at least 150 other targets) while NGB-2904 has not undergone this degree of testing. Since SB-277011A's D₃/D₂ ratio (100) is not as high as that of NGB-2904 (800), it is possible that SB-277011A may be binding to some D₂ as well as D₃ receptors. This may serve as an explanation for the SB-277011A effect on decreasing operant FSA in Zucker rats as observed in this study even though it has been shown not affect progressive-ratio for food reinforcement and to not modify sucrose-seeking behavior induced by sucrose-associated cues in rats (Ross et al., 2007; Cervo et al., 2007) as well as decreasing drug reward as observed in other studies (for review see Heidbreder et al., 2005), particularly since the D₂ is significantly involved in both drug and food consumption (Volkow and Wise, 2005). In contrast to this explanation, SB-277011A (at doses as high as 90 mg/kg) was shown not to exhibit any of the behavioral properties attributed to D₂ receptor antagonists (Pak et al., 2006). Therefore, it is not likely that SB-277011A would be interacting significantly with D₂ receptors at the doses used in this study.

4.3. Zucker Ob rats and FSA

One important finding of this study was that Ob and Le rats did not show significant differences in operant FSA during the vehicle sessions. Taking the severe hyperphagia that Ob rats exhibit into account, one would expect that these rats would show increased R lever responses and consequently consume a lot more food than their Le counterparts. Although our FSA results indicated that Ob rats had a greater food intake than Le rats following vehicle treatment, it was not statistically significant. There are several possible factors that may explain this finding. Previous studies have examined FSA in both Ob Zucker and diet-induced obese (DIO) rats under an FR schedule and experimental procedures similar to ours (Glass et al., 1999; la Fleur

et al., 2007). In these studies, operant lever responses for food did not differ between obese and lean animals (Glass et al., 1999; la Fleur et al., 2007). Nevertheless, operant responding behavior in obese rats seems to be subject to the specific schedule of reinforcement. Indeed, experiments that examined FSA and utilized variable-interval schedules of reinforcement [i.e. progressive-ratio (PR)] demonstrated the opposite: Ob Zucker and DIO rats show increased lever responses compared to lean controls (la Fleur et al., 2007; Vasselli et al., 1980). The increased operant responding in obese rats as demonstrated using PR schedules has been interpreted to reflect the increased incentive value and motivational properties of food in obese rats (la Fleur et al., 2007). However, it has been postulated that obese rats do not show increased lever responses for food under FR schedules because of the influence of satiety signals in responding for food. Under FR schedules, it is believed that rats achieve satiety after a certain number of lever responses, which in turn decreases the motivation for food and eventually the lever response (la Fleur et al., 2007). An exception to this was the Greenwood et al. study (Greenwood et al., 1974) which showed increased lever responses for Ob Zucker rats compared to Le and which assessed operant responding under an FR schedule but over a period of 24 h. A possible explanation for this is that Ob and Le Zucker rats differ in their circadian feeding patterns and therefore variability in circadian rhythms between the two strains may contribute to differences observed in the Greenwood et al. study. Indeed, since then it has been shown that Ob Zucker rats differ from Le in circadian rhythms for temperature, locomotor activity and feeding (Fukagawa et al., 1992; Mistlberger et al., 1998; Murakami et al., 1995).

Zucker Ob rats exhibit differences in food regulatory messengers such as leptin, ghrelin and insulin among others (Michaelides et al., 2006; Thanos et al., 2007). It has been shown that intra-cerebroventricular administration of both leptin and insulin decreased FSA in normal rats (Figuelewicz et al., 2006). Differences in concentrations of these messengers may also contribute to the lack of difference observed between Ob and Le rats during vehicle sessions in the operant FSA task.

Recently it was shown that Ob Zucker rats have lower D_2 receptor levels than Le rats and that chronic food restriction leads to increases in D_2 receptors in both strains (Michaelides et al., 2006; Thanos et al., 2007). The negative correlation between weight and D_2 receptor levels was consistent with what was observed clinically between the Body Mass Index (BMI) and D_2 receptors (Wang et al., 2001, 2004, 2003). Furthermore, Ob rats show a distinct DA profile that responds uniquely to fasting and food restriction (Michaelides et al., 2006; Thanos et al., 2007). Therefore, differences in DA and its receptors between Ob and Le rats may also contribute to the operant FSA behavior observed, especially since pharmacological manipulation of DA and D_2 receptors has been shown to modulate food operant responding (Barrett et al., 2004; Ishiwari et al., 2004). Future experiments measuring D_3 receptor levels in Ob and Le Zucker rats may shed light on the involvement of D_3 receptors in weight gain and food intake and how these responses may be modulated by leptin.

4.4. Limitations

D_3 receptor levels have not been characterized in obese clinical or preclinical studies partly because of the lack of availability of highly selective D_3 receptor radioligands. It is possible that there exists a difference in D_3 receptor levels with obesity and this of course has not been determined between obese and non-obese rats (this would produce a differential sensitivity to D_3 receptor antagonism). This remains to be evaluated in future experiments. Different D_3 profiles between Ob and Le Zucker rats may serve as a potential contribution to the Ob Zucker phenotype. This is possible since, it has been recently shown that they have been characterized with different D_2 receptors and DA profiles (Hamdi et al., 1992; Michaelides et al., 2006; Thanos et al., 2007) and therefore differences in other receptors and transmitters may exist.

4.5. Conclusion

The findings of this experiment, taken together with evidence that lack of D₃ receptors contributes to increased adiposity and disruption of leptin and insulin levels in response to a high fat diet but not regular chow (McQuade et al., 2004) supports a modulatory role for the D₃ receptor with respect to obesity. At the doses examined, SB-277011A significantly inhibited operant FSA behavior while NGB-2904 did not in Zucker Ob rats. While both SB-277011A and NGB-2904 have been previously shown to decrease drug intake in rodents, and may be valuable pharmacotherapeutic agents for drug addiction; SB-277011A displayed properties in the Zucker obese rat that warrant further investigation with respect to food-seeking and weight loss in obesity. Furthermore, higher doses of NGB-2904 need to be examined. Our findings thus as well as other reports support the further investigation of the contribution and interaction of the D₃ receptors with several key signaling messengers (like leptin, insulin and ghrelin) in modulating reward and satiety for natural reinforcers such as food. Furthermore, these results may indicate that the interaction and involvement of these signals may function differently for drug reward versus natural rewards like food.

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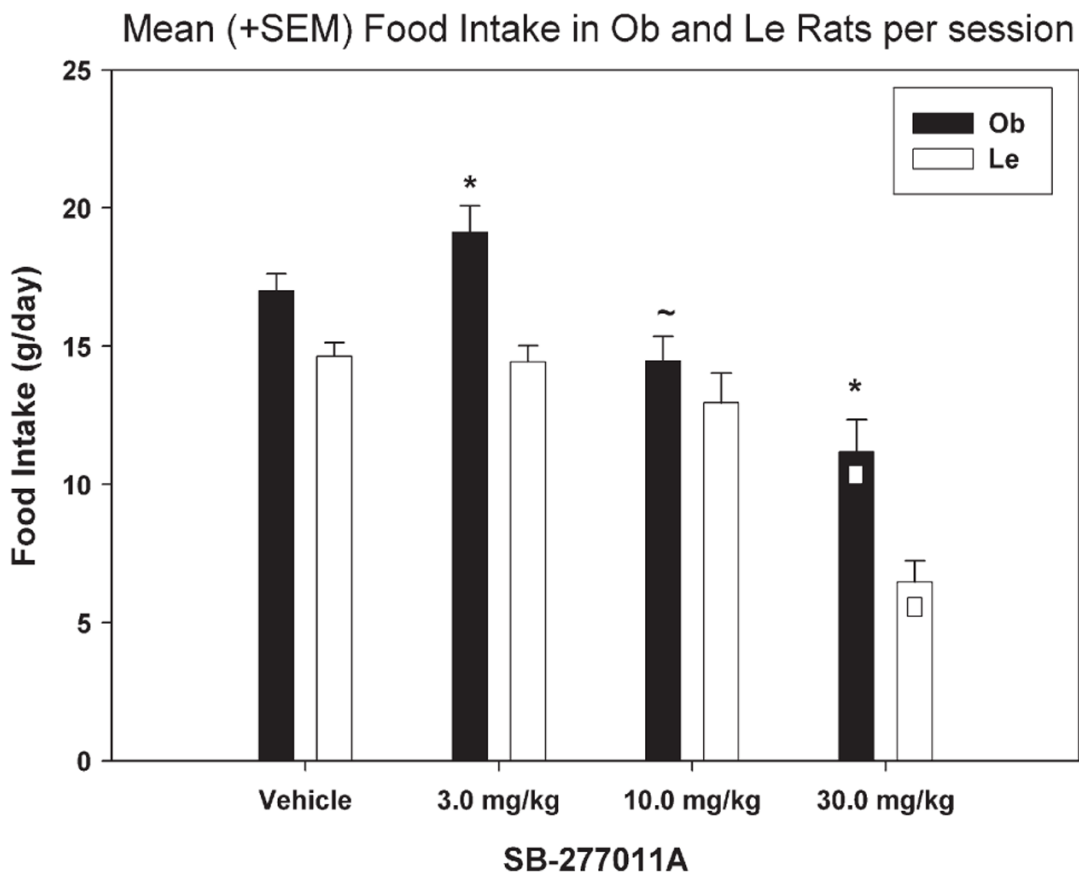


Fig. 1. Mean (+SEM) number of grams of food intake per 2-h session, following injection of: vehicle, 3, 10 or 30 mg/kg of SB-277011A. □ Denotes significance between specific treatment relative to all others, ~denotes significance between 10 and 3 mg/kg in Ob rats and * denotes significance between Ob and Le rats at specific dose. The vehicle column data represent the average of the first (baseline), intermittent (between treatments) and last (after the last drug treatment) vehicle sessions.

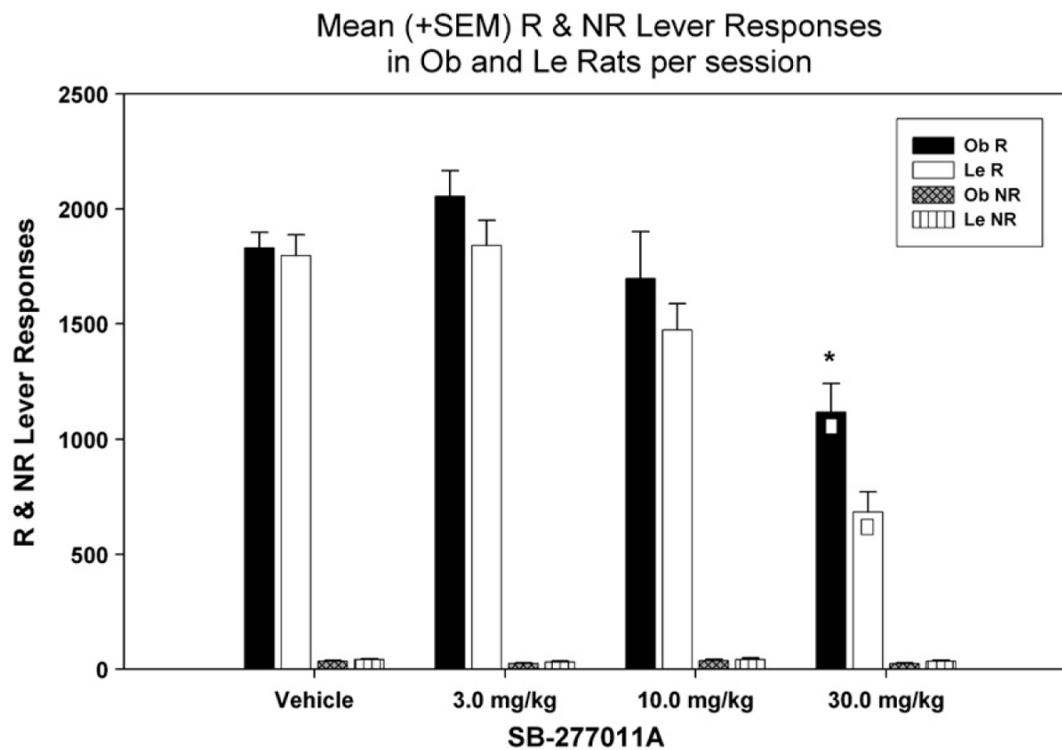


Fig. 2. Mean (+SEM) number of R (total and during time-out) and NR lever responses per 2-h session, following injection of: vehicle, 3, 10, or 30 mg/kg of SB-277011A. □ Denotes significance between specific treatment relative to all others, and * denotes significance between Ob and Le rats at specific dose. The vehicle column data represent the average of the first (baseline), intermittent (between treatments) and last (after the last drug treatment) vehicle sessions.

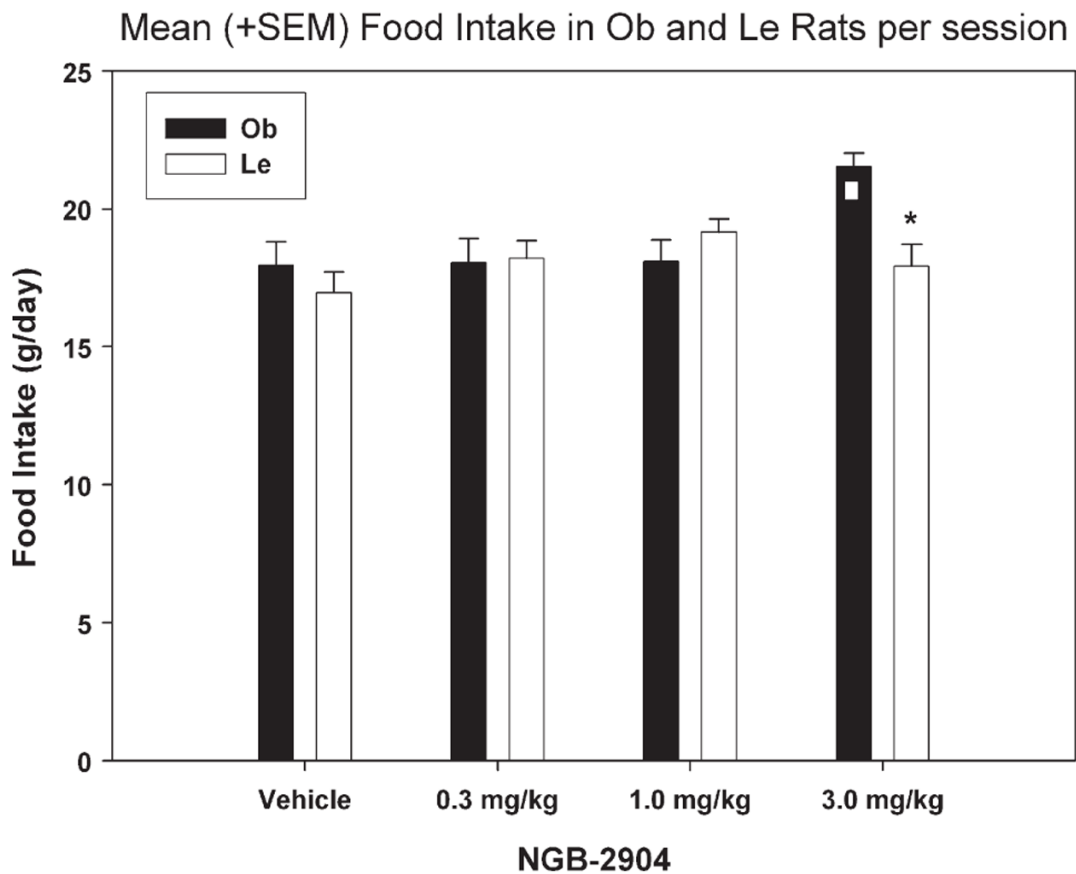


Fig. 3. Mean (+SEM) number of grams of food intake per 2-h session, following injection of: vehicle, 0.3, 1, or 3 mg/kg of NGB-2904. □ Denotes significance between specific treatment relative to all others, and * denotes significance between Ob and Le rats at specific dose. The vehicle column data represent the average of the first (baseline), intermittent (between treatments) and last (after the last drug treatment) vehicle sessions.

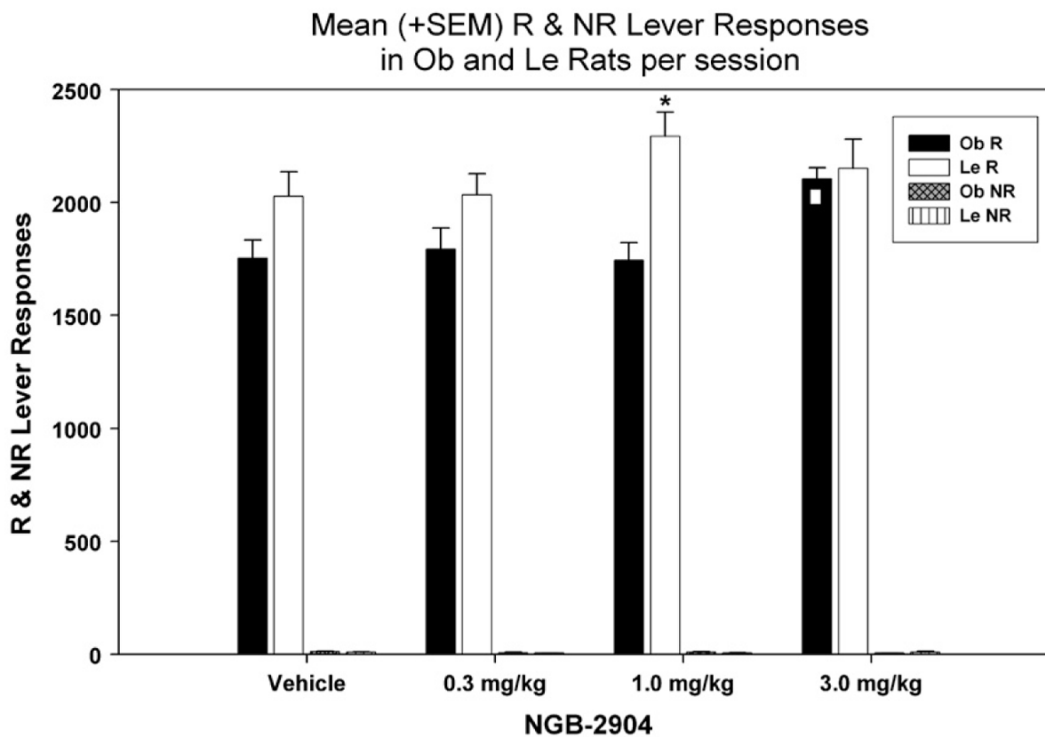


Fig. 4. Mean (+SEM) number of R (total and during time-out) and NR lever responses per 2-h session, following injection of: vehicle, 0.3, 1, or 3 mg/kg of NGB-2904. □ Denotes significance between specific treatment relative to all others, and * denotes significance between Ob and Le rats at specific dose. The vehicle column data represent the average of the first (baseline), intermittent (between treatments) and last (after the last drug treatment) vehicle sessions.