

Effects of Chronic Buspirone Treatment on Cocaine Self-Administration

Nancy K Mello^{*1}, Peter A Fivel¹, Stephen J Kohut¹ and Jack Bergman¹

¹Alcohol and Drug Abuse Research Center, Harvard Medical School-McLean Hospital, Belmont, MA, USA

Cocaine abuse and dependence is a major public health problem that continues to challenge medication-based treatment. Buspirone (Buspar) is a clinically available, non-benzodiazepine anxiolytic medication that acts on both serotonin and dopamine systems. In recent preclinical studies, acute buspirone treatment reduced cocaine self-administration at doses that did not also decrease food-reinforced behavior in rhesus monkeys (Bergman *et al*, 2012). The present study evaluated the effectiveness of chronic buspirone treatment on self-administration of cocaine and food. Five adult rhesus monkeys (*Macaca mulatta*) were trained to self-administer cocaine and food during four 1-h daily sessions under a second-order schedule of reinforcement (FR2 [VR 16;S]). Buspirone (0.32 and 0.56 mg/kg/h) was administered intravenously through one lumen of a double-lumen catheter every 20 min for 23 h each day for 7–10 consecutive days. Each buspirone treatment period was followed by saline control treatment until drug- and food-maintained responding returned to baseline levels. Buspirone significantly reduced responding maintained by cocaine, and shifted the dose–effect curve downwards. Buspirone had minimal effects on food-maintained responding. In cocaine discrimination studies, buspirone (0.1–0.32 mg/kg, IM) did not antagonize the discriminative stimulus and rate-altering effects of cocaine in four of six monkeys. These findings indicate that buspirone selectively attenuates the reinforcing effects of cocaine in a nonhuman primate model of cocaine self-administration, and has variable effects on cocaine discrimination.

Neuropsychopharmacology (2013) **38**, 455–467; doi:10.1038/npp.2012.202; published online 17 October 2012

Keywords: buspirone; cocaine; drug self-administration; dopamine D₃ receptor; dopamine D₄ receptor; 5-HT_{1A}

INTRODUCTION

Buspirone (Buspar) is a non-benzodiazepine anxiolytic approved by the FDA for the treatment of anxiety. Buspirone has a complex pharmacology and its mechanisms of anxiolytic action are poorly understood. Both dopamine and serotonin receptors appear to contribute to buspirone's behavioral effects. Buspirone acts as a 5-HT_{1A} partial agonist (Newman-Tancredi *et al*, 1998; Wong *et al*, 2007), and also binds to dopamine D₂, D₃, and D₄ receptors (Bergman *et al*, 2012; Kula *et al*, 1994; Tallman *et al*, 1997). In addition to amelioration of anxiety, buspirone has also been used to reduce cigarette smoking (Cinciripini *et al*, 1995; Henningfield *et al*, 2005; Hilleman *et al*, 1992; West *et al*, 1991) and is being considered for treatment of marijuana dependence and withdrawal (Elkashef *et al*, 2008; McRae-Clark *et al*, 2009; Weinstein and Gorelick, 2011). Buspirone has been well tolerated in clinical studies, and there is little evidence of abuse liability in humans (Griffith

et al, 1986). Recent interest in buspirone for the treatment of cocaine abuse and dependence is based in part on evidence that buspirone has dopamine D₃ and D₄ selective actions, which may mediate its ability to decrease cocaine self-administration (Bergman *et al*, 2012).

Many preclinical studies have evaluated the effects of buspirone on the abuse-related characteristics of cocaine, and most have concluded that buspirone's interactions with cocaine reflect its dopamine D₂ antagonist properties. Neuropharmacological studies have demonstrated that buspirone selectively blocks the D₃ and D₄ subtypes of the D₂ receptor and that this selectivity extends to its dopamine antagonist effects in the functional β -arrestin assay (Bergman *et al*, 2012; Kula *et al*, 1994). This is of interest because of the suggested importance of dopamine D₃ and D₄ receptors in the abuse-related effects of cocaine, and therefore, in the design of candidate treatment medications. The potential role of dopamine D₃ and D₄ receptor antagonists and partial agonists in drug abuse treatment medications has stimulated considerable research (see for review Heidbreder *et al*, 2005; Heidbreder and Newman, 2010; Le Foll *et al*, 2005; Le Foll *et al*, 2000; Newman *et al*, 2005; Newman *et al*, 2012). The recent discovery of buspirone's dopamine D₃ and D₄ antagonist activity, combined with our observation that an acute dose of buspirone reduced cocaine self-administration by rhesus monkeys (Bergman *et al*, 2012), led us to examine the effects

*Correspondence: Dr NK Mello, Alcohol and Drug Abuse Research Center, Harvard Medical School-McLean Hospital, 115 Mill Street, Belmont, MA 02478, USA, Tel: +1 617 855 2746, Fax: +1 617 855 2519, E-mail: nmello@mclean.harvard.edu

Preliminary data were reported at the 2011 meeting of the American College of Neuropsychopharmacology.

Received 20 June 2012; revised 28 August 2012; accepted 29 August 2012

of chronic treatment with buspirone on cocaine self-administration, as well as the effects of buspirone pretreatment on cocaine discrimination.

The effects of buspirone on the abuse-related effects of cocaine have been inconsistent across species and test procedures. In nonhuman primates, buspirone did not share discriminative stimulus properties with cocaine (Koetzner, 1996) and did not maintain self-administration (Balster, 1990; Balster and Woolverton, 1982; Griffiths *et al*, 1991), suggesting that buspirone may have relatively low abuse potential. In drug discrimination studies in rodents, buspirone also did not substitute for cocaine, but either dose-dependently reduced cocaine-appropriate responding and shifted the cocaine dose-effect function to the right (Callahan and Cunningham, 1997), or had no effect (Rapoza, 1993). The effects of buspirone on the reinforcing effects of cocaine in nonhuman primates have not been widely studied. Early studies in rhesus monkeys showed that acute treatment with buspirone (0.1–0.56 mg/kg) significantly increased cocaine self-administration, whereas a high dose, 1.0 mg/kg, significantly decreased rates of cocaine self-administration under a fixed ratio (FR) 10 schedule of reinforcement (Gold and Balster, 1992). Rate-increasing effects of buspirone also were observed under a second-order schedule of IV cocaine self-administration in squirrel monkeys (Nader and Barrett, 1990). The effects of chronic buspirone treatment on cocaine self-administration also have not been widely studied. In contrast to buspirone's acute effects, daily treatment with the relatively low single doses (0.1 and 0.3 mg/kg, IV) had little effect on cocaine self-administration over a 10-day period (Gold and Balster, 1992). It was suggested that multiple daily injections would be necessary to maintain a constant buspirone blood level (Gold and Balster, 1992).

We recently reported that acute administration of relatively low doses of buspirone (0.1 or 0.3 mg/kg, IM) selectively reduced cocaine self-administration maintained under a FR 30 schedule in rhesus monkeys (Bergman *et al*, 2012). Buspirone produced a downward shift in the cocaine dose-effect curve (0.003–0.1 mg/kg/injection) with inconsistent effects on food-maintained responding (Bergman *et al*, 2012). The current study extends these findings to examine the effects of chronic buspirone treatment on cocaine- and food-maintained responding. Evaluation of the chronic effects of candidate treatment medications is important to determine whether tolerance develops to acute effects during repeated administration, and whether medication effects on the abused drug are sustained over time (Mello, 2005; Mello and Negus, 1996).

The present report examines the effects of 7–10 days of buspirone treatment on IV cocaine- and food-maintained responding using a procedure in which IV saline or buspirone injections were delivered every 20 min for 23 h each day. This procedure was designed to ensure that steady-state levels of the treatment medication and its metabolites were present during the four drug and food sessions each day (Negus and Mello, 2003). This procedure is especially valuable for studies of a relatively short-acting medication such as buspirone with active metabolites that may have differing durations of action (Bergman *et al*, 2012; Dockens *et al*, 2006; Gammans *et al*, 1986). The effects of buspirone were also studied in a conventional two-key, drug

discrimination procedure to determine whether it attenuated or enhanced the cocaine discriminative stimulus.

MATERIALS AND METHODS

Subjects

Eleven male rhesus monkeys (*Macaca mulatta*) that weighed between 6 and 10 kg were studied. Each monkey was studied as his own control across saline and buspirone treatment conditions. All monkeys had a history of cocaine self-administration. Each day, monkeys received multiple vitamins, fresh fruit and vegetables, and Lab Diet Jumbo Monkey Biscuits (PMI Feeds, St Louis, MO) to supplement a banana-flavored pellet diet, fortified with vitamin C (Formula 4TUR banana flavor, grain-based pellet, Purina Mills Test Diet, Richmond, IN). Food supplements were given twice a day between 0900 and 0930 hours, and 1300 and 1330 hours. Monkeys were maintained at 90% free-feeding weight. Water was continuously available from an automatic watering system. A 12-h light-dark cycle was in effect (lights on 0800–2000 hours), and the experimental chamber was dark during food and drug self-administration sessions.

Animal maintenance and research were conducted in accordance with the guidelines provided by the Institute of Laboratory Animal Resources (ILAR-NRC, 1996) and the NIH Office of Laboratory Animal Welfare. The facility is licensed by the US Department of Agriculture, and protocols were approved by the Institutional Animal Care and Use Committee. Monkeys were observed at least twice every day, and any changes in general activity were noted. In addition, the health of the monkeys was regularly monitored by consultant veterinarians trained in primate medicine. Operant food and drug acquisition procedures provided an opportunity for enrichment and for monkeys to manipulate their environment (Line, 1987). Monkeys had visual, auditory, and olfactory contact with other monkeys throughout the study.

Cocaine Self-Administration

Surgical procedures. Double lumen Silicone rubber catheters (I.D. 0.028 in, O.D. 0.088 in) (Saint Gobain Performance Plastics, Beaverton, MI) were surgically implanted in the internal jugular, external jugular, or femoral vein to permit IV drug self-administration and IV buspirone or saline administration. All surgical procedures were performed under aseptic conditions. Monkeys were initially sedated with ketamine (5–10 mg/kg, IM), and atropine (0.05 mg/kg) SC or IM was administered to reduce salivation. Following insertion of an endotracheal tube, anesthesia was maintained with isoflurane (1–2% mixed with oxygen). After surgery, monkeys were given procaine penicillin G at 20 000 units/kg, IM twice daily for 5 days, or cephalexin 20 mg/kg, PO twice daily for 5 days. An analgesic dose of buprenorphine (0.032 mg/kg, IM) and Metacam (meloxicam) (0.1 mg/kg, SC) was administered twice daily for 3 days.

The intravenous catheter exited in the mid-scapular region and was protected by a tether system consisting of a custom-fitted nylon vest connected to a flexible stainless-steel

cable and fluid swivel (Lomir Biomedical, Malone, NY). This flexible tether system permits monkeys to move freely. Catheter patency was evaluated periodically by administration of a short-acting barbiturate, methohexital sodium (4 mg/kg) through the catheter lumen. If muscle tone decreased within 10 s after drug administration, the catheter was considered patent.

Behavioral procedures and apparatus for drug self-administration. Monkeys lived in stainless-steel chambers (64 × 64 × 79 cm) equipped with a custom-designed operant response panel (28 × 28 cm), a pellet dispenser (Gerbrands Model G5210, Arlington, MA) and two syringe pumps (Model 981210, Harvard Apparatus, South Natick, MA), one for each lumen of the double-lumen catheter. During food self-administration sessions, the response key (6.4 × 6.4 cm) on the operant panel was illuminated with a red light. Completion of the response requirement under a FR 2, variable ratio 16 (FR 2, [VR 16:S]) schedule resulted in presentation of a 1 s-red light beneath the response key. Completion of a second VR16 resulted in delivery of a 1-g banana-flavored pellet (Purina, Estherville, IA). During drug self-administration sessions, the response key was illuminated with a green light, and completion of the response requirement under a FR 2, [VR 16:S] schedule resulted in delivery of saline or a cocaine solution over 1 s through one lumen of the double-lumen catheter. A 10-s time-out followed delivery of each drug injection or food pellet, during which stimulus lights remained off and responding had no scheduled consequences. If 25 food pellets or 20 injections were delivered before the end of the 1-h session, then all stimulus lights were turned off, and responding had no scheduled consequences for the remainder of that session. Thus, a monkey could earn a maximum of 100-food pellets/day and 80 cocaine or saline injections/day.

The four daily food self-administration sessions began at 1100, 1500, 1900, and 0600 hours the next morning, and the four daily drug self-administration sessions began at 1200, 1600, 2000, and 0700 hours the next morning. The daily sequence of sessions is shown schematically in Figure 1. Room lights were off during all experimental sessions. Schedules of reinforcement were programmed with custom-designed software and IBM-compatible computers and interface systems (Med Associates, St Albans, VT). Additional details of this apparatus have been described previously (Mello *et al*, 1995). Drug doses were varied by computer-controlled changes in pump infusion duration (Fivel, 2011).

Training procedure. Monkeys were initially trained to self-administer 1-g banana-flavored food pellets until food-maintained performance was stable; subjects then were trained to respond for cocaine (0.1 mg/kg/injection). Stability was defined as at least 3 days when food- or drug-maintained responding varied by no more than 20 percent and there were no upwards or downwards trends. Once consistent cocaine-maintained responding occurred, the unit dose was reduced to 0.01 or 0.032 mg/kg to limit the disruptive effects of each IV cocaine injection, and facilitate higher levels of IV self-administration behavior throughout the session. Extinction training consisted of sessions in

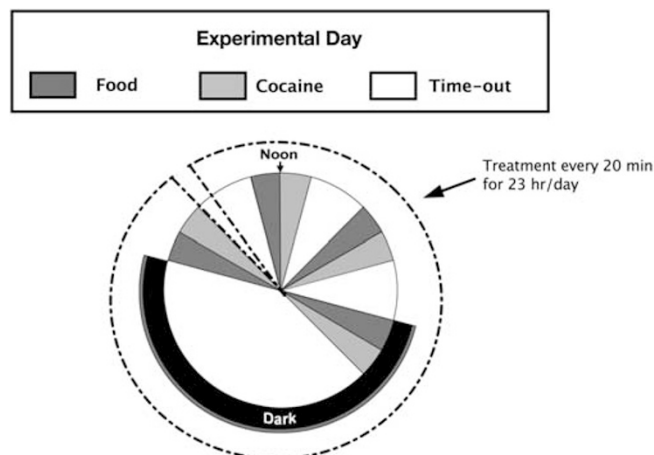


Figure 1 Schematic representation of daily experimental procedures. Daily food self-administration sessions are shown as black wedges. Daily cocaine self-administration sessions are shown as gray wedges. Time out periods when responding had no scheduled consequences are shown as open areas. The broken circle indicates the 23-h time period when IV buspirone or saline treatment was administered every 20 min. The 1-hour during the first time out period when daily health checks and general maintenance occurred are indicated by a break in the circle during the first morning time out. Experiments ran 7 days per week throughout the year.

which saline was substituted for the cocaine training dose. Once stable low levels of saline-maintained responding were observed during extinction, drug dose-effect curves were determined over a dose range of 0.0032–0.10 mg/kg/injection IV cocaine. Saline and different doses of cocaine were presented in an irregular order.

Cocaine dose-effect curve determinations. Training continued until monkeys met the following criteria for stable food and cocaine self-administration under the FR2 [VR16:S] schedule of reinforcement: (1) 3 consecutive days during which the number of drug injections/day varied by no more than 20% of the 3-day mean with no upward or downward trend, and (2) the mean number of food pellets and injections delivered per day was equal to or greater than 60. Once responding was stable, saline and each dose of cocaine (0.0032–0.10 mg/kg/injection) was substituted for a minimum of 7 days and until responding was stable according to the above criteria, or for a maximum of 10 days. Following substitution of each dose of cocaine, monkeys were returned to the maintenance dose of cocaine for at least 3 days, and until responding was stable to ensure reliable baseline responding prior to substitution of the next dose of cocaine. Cocaine doses were presented in an irregular order that differed across monkeys.

Testing procedures. Buspirone doses (0.32 and 0.56 mg/kg/h) were selected on the basis of our previous report of the acute effects of buspirone (Bergman *et al*, 2012) and previous studies in rhesus monkeys (Gold and Balster, 1992). The effects of buspirone on the reinforcing effects of cocaine were evaluated using a procedure similar to that used in our studies of the effects of d-amphetamine on cocaine self-administration (Negus and Mello, 2003). Saline or a dose of buspirone was administered through one lumen of a double-lumen catheter every 20 minutes for 23 h each

day. Buspirone treatment was suspended during 0900 and 1000 hours each morning to permit experimental changes and husbandry. This is shown schematically in Figure 1. The total injection volume delivered was 6.9 ml in 69 injections. This medication administration procedure was developed to ensure that relatively short-acting drugs would be continuously present during the test sessions (Negus and Mello, 2003). Each treatment dose was studied for 7–10 days until responding was stable. Successive buspirone doses were separated by an interval of saline treatment until drug- and food-maintained responding returned to baseline levels. The saline treatment interval was used to prevent any carryover effects from the preceding treatment.

Sedation ratings. Immediately following the noon cocaine self-administration session, sedative effects were rated on a simple scale adapted from a scale designed to measure sedation and prolactin levels following administration of a dynorphin analog (Butelman *et al*, 1999). These ratings were made daily by a trained observer to determine if sedation might have contributed to any changes in food or cocaine self-administration observed during buspirone treatment.

Data analysis. The primary dependent variables were the total number of cocaine or saline injections and food pellets earned per day. The number of cocaine injections self-administered during the last 3 days of each substitution condition were averaged. A linear regression analysis was used to test for main effects of treatment and cocaine dose and a treatment X cocaine dose interaction for total cocaine injections and food pellets earned. Where significant main effects were found, one-way analysis of variance (ANOVA) with Dunnett's *post-hoc* tests were used to determine which cocaine self-administration points differed between saline and buspirone treatment doses (Sheskin, 2000). To determine which doses of cocaine produced significantly more cocaine injections compared to saline self-administration, paired *t*-tests with Bonferroni step-down error corrections were performed. All statistical procedures were accomplished using SPSS v.20 and figures were drawn using GraphPad Prism v.5.

Two of the five monkeys lost their catheters before studies of buspirone's effects on 0.0032 mg/kg/injection cocaine could be completed. Thus, each data point for saline and 0.01–0.10 mg/kg/injection cocaine (with saline and buspirone treatment) reflects 4–5 monkeys. Data for 0.0032 mg/kg/injection cocaine during buspirone treatment reflect three monkeys.

Measurement of buspirone plasma levels. Blood samples were collected from the saphenous vein in five monkeys for analysis of buspirone levels in plasma. Each monkey had 48 h of exposure to 0.32 mg/kg/h IV buspirone before sample collection. No cocaine self-administration sessions were conducted at that time. Blood samples were collected between 0900 and 0920 hours when buspirone was not being administered (see Figure 1). Monkeys were lightly sedated with ketamine (3–5 mg/kg, IM) to permit blood collection.

Plasma concentrations of buspirone were analyzed in duplicate by gas chromatography by NMS Labs, Willow Grove, PA. The assay sensitivity was 5 ng/ml.

Drug source and preparation. Cocaine HCl was provided by the National Institute on Drug Abuse (Rockville, MD, USA) and prepared in sterile saline (0.9%). Buspirone HCl was prepared in sterile water (Sigma-Aldrich, St Louis, MO, USA). Self-administered drugs were sterile-filtered with a 0.22- μ syringe-driven filter.

Cocaine Discrimination

Apparatus and behavioral procedures for cocaine discrimination. Each monkey lived in a stainless-steel chamber (56 × 71 × 69 cm) equipped with a computer-controlled operant panel and food pellet dispenser. Discrimination training and testing procedures were identical to those described previously (Mello and Negus, 2007). Training sessions consisted of 1–5 cycles, and each cycle consisted of a 15-min time-out period followed by a 5-min response period. During the time-out, all stimulus lights were off and responding had no scheduled consequences. During the response period, the right and left response keys were illuminated red or green, and monkeys could earn up to 10 food pellets under a FR 30 schedule. On training days, monkeys received saline or 0.40 mg/kg cocaine IM at the beginning of each cycle. Following saline administration, responding on only the green, saline-appropriate key produced food. Following 0.40 mg/kg cocaine administration, only responding on the red, drug-appropriate key produced food. Responses on the inappropriate key reset the FR requirement on the appropriate key. If the training dose of cocaine was administered, it was administered only during the last cycle. The principal dependent variables were (a) percent injection-appropriate responses for the entire cycle, and (b) response rate in responses per second. Training was complete when the following criteria were met during seven of eight consecutive training sessions: (1) $\geq 80\%$ injection-appropriate responding before the first reinforcer; (2) $\geq 90\%$ injection-appropriate responding for the entire cycle; (3) ≥ 10 responses per second during saline training cycles.

The effects of acute treatment with buspirone on cocaine discrimination were studied in six monkeys. Buspirone (0.1–0.32 mg/kg, IM) was administered 15 min before determination of a cumulative cocaine discrimination dose–effect curve (0.013–1.3 mg/kg, IM). A control cocaine dose–effect curve was determined before each test with buspirone or saline. If responding did not meet criterion levels of discrimination performance, training was continued until criterion levels of performance occurred for at least 2 consecutive days.

Data analysis. Individual subject data are shown as cocaine-appropriate responding (over the entire response period) calculated as a percentage of responses on the cocaine lever out of total responses on the saline and cocaine levers and response rate in responses per second (excluding responses during time out) as a function of cocaine dose. Cocaine ED₅₀ values were calculated for

individual monkeys by log-linear interpolation as the dose of cocaine that produced 50% cocaine-appropriate responding. Data points for percentage of cocaine-appropriate responding were excluded from analysis if response rate was less than 20% of control cocaine levels for individual subjects, however, response rate data for those points are included. In pretreatment studies, buspirone was considered to alter the cocaine discrimination dose-effect curve if the test ED_{50} value did not fall within the 95% confidence limits of two baseline ED_{50} values.

RESULTS

Buspirone's Effects on Cocaine Self-Administration

When only saline was available for self-administration during saline treatment, monkeys took an average of 15.87 (± 6.61) injections per day (Figure 2, top panel). During saline treatment, a cocaine unit dose of 0.01 mg/kg/injection was at the peak of the dose-effect curve, and monkeys self-administered an average of 79.8 (± 0.20) of a possible 80 injections per day. Figure 2 shows that treatment with buspirone (0.32 mg/kg/h) for 7–10 days shifted the ascending

limb (0.0032 mg/kg/injection) and the descending limb (0.10 mg/kg/injection) of the cocaine dose-effect curve downwards. Buspirone significantly reduced cocaine self-administration of 0.01 mg/kg/injection, at the peak of the dose-effect curve during saline control treatment ($P < 0.05$), and shifted the peak 1/2 log unit to the right. Treatment with a higher dose of buspirone (0.56 mg/kg/h) also shifted the ascending limb of the cocaine dose-effect curve downwards. However, there were no significant differences between the effects of 0.32 and 0.56 mg/kg/h, buspirone on cocaine-maintained responding.

Buspirone's Effects on Food-Maintained Responding

During saline treatment and saline self-administration, monkeys took the maximum number of food pellets available (Figure 2, bottom panel). Food-maintained responding was not affected by increasing doses of cocaine during saline treatment. During treatment with 0.32 and 0.56 mg/kg/h buspirone, food-maintained responding was not significantly altered on the ascending limb of the cocaine dose-effect curve when buspirone's effects on cocaine self-administration were greatest. During sessions in which cocaine unit doses of 0.01 and 0.32 mg/kg/injection were studied, treatment with 0.32 mg/kg/h buspirone decreased food-maintained responding by 14 and 23%, whereas treatment with 0.56 mg/kg/h buspirone decreased food-maintained responding by 26 and 33%. At the highest unit dose of cocaine (0.1 mg/kg/injection), food-maintained responding decreased by 33% and 34% during treatment with 0.32 and 0.56 mg/kg/h buspirone, respectively.

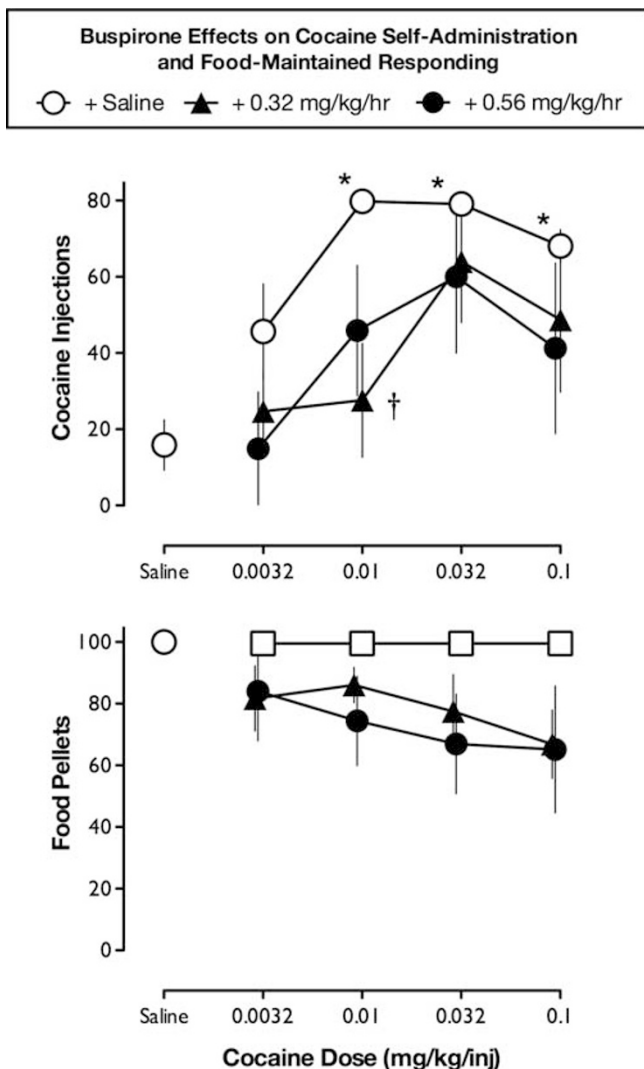


Figure 2 The effects of chronic treatment with buspirone or saline on cocaine- and food-maintained responding. Dose-effect curves for cocaine self-administration (0.0032–0.10 mg/kg/injection) are shown in the top panel for the group of monkeys. The unit doses of cocaine are shown on the abscissae, and the number of injections per day is shown on the left ordinate. Points above 'saline' show data from saline treatment sessions when saline was available for self-administration. Cocaine self-administration during chronic saline treatment is shown as open circles. Cocaine self-administration during chronic treatment with buspirone is shown as filled triangles (0.32 mg/kg/h) or filled circles (0.56 mg/kg/h). Food-maintained responding during saline treatment and saline or cocaine self-administration is shown as open squares in the bottom panel. Food-maintained responding during treatment with buspirone is shown as filled triangles (0.32 mg/kg/h) or filled circles (0.56 mg/kg/h). Each data point for saline and 0.01–0.10 mg/kg/injection cocaine reflects 4–5 monkeys. Data for 0.0032 mg/kg/injection cocaine during buspirone treatment reflect three monkeys. Each data point is based on the last 3 days of a 7- to 10-day treatment (mean \pm SEM). The regression analysis indicated a significant main effect of treatment ($P < 0.05$) on cocaine self-administration. One-way ANOVA with Dunnett's post-test indicated that treatment with 0.32 mg/kg/h buspirone reduced cocaine self-administration at 0.01 mg/kg/injection ($F_{2,13} = 4.83$; $P < 0.05$) and a trend for significance with 0.56 mg/kg/h buspirone treatment ($P = 0.076$). Paired t -tests found that cocaine injections were higher than saline self-administration at doses of 0.01, 0.032, and 0.1 ($t = -9.797, -9.372, \text{ and } -6.778$; $P < 0.05$) mg/kg/injection with saline treatment. The regression analysis on food pellets earned indicated a significant main effect of buspirone treatment ($P < 0.05$), but post-tests indicated that food-maintained responding during buspirone treatment were not significantly different from baseline at any cocaine dose. Asterisks denote data points that were significantly different from saline self-administration. Daggers indicate data points that were significantly lower than during saline control treatment.

To further evaluate the effects of buspirone on food-maintained responding in the absence of cocaine, data from the first food session of the experimental day were analyzed. This food session began at 1100 hours, 3 h after the previous

cocaine self-administration session ended at 0800 hours. Figure 3 shows that monkeys self-administered the maximum number of pellets available in the session during the last 3 days of saline control and chronic buspirone treatment.

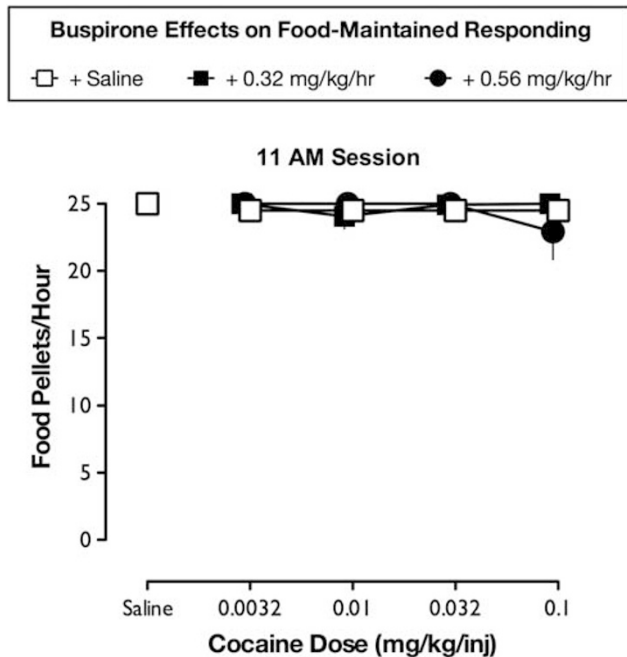


Figure 3 Effects of Buspirone on food-maintained responding in the absence of cocaine. Food-maintained responding during the 1100 hours session, 4 h after the last cocaine session are shown for the last 3 days of each cocaine dose condition. Saline treatment is shown as open squares, buspirone treatment is shown as black squares and black circles for doses of 0.32 and 0.56 mg/kg/h, respectively. The number of food pellets taken in the first food session of the experimental day is shown on the ordinate. The dose of cocaine available during that 7- to 10-day period is shown on the abscissa.

Effects of Buspirone on Daily Cocaine- and Food-Maintained Responding in Representative Individual Monkeys

During saline treatment, each monkey earned all of the cocaine injections (0.01 mg/kg/injection) and food pellets that were available each day (top row; Figure 4a and b). During baseline control determinations, immediately before buspirone treatment, each monkey continued to self-administer high levels of cocaine and food.

Figure 4a (row 2) shows that initiation of buspirone treatment (0.32 mg/kg/h) immediately reduced cocaine self-administration to 34% percent of control levels and this decrease was sustained throughout 8 days of treatment. Food-maintained responding was reduced to 58% percent of control levels on day 2 of treatment, but returned to control levels on day 5. After cessation of buspirone treatment, monkey MM 30 increased cocaine self-administration to 70% percent of control levels within 4 days, but cocaine self-administration did not meet stability criteria (3 days of >60 injections) for 20 days.

The effects of treatment with a higher dose of buspirone (0.56 mg/kg/h; row 3) initially decreased both cocaine self-administration and food-maintained responding to 26% and 56%, respectively, on day 1. However, cocaine self-administration returned to control levels by day 3 and food-maintained responding returned to control levels by day 5. Cocaine self-administration continued to be variable for 3 days after cessation of buspirone (0.56 mg/kg/h) treatment, whereas food-maintained responding remained stable.

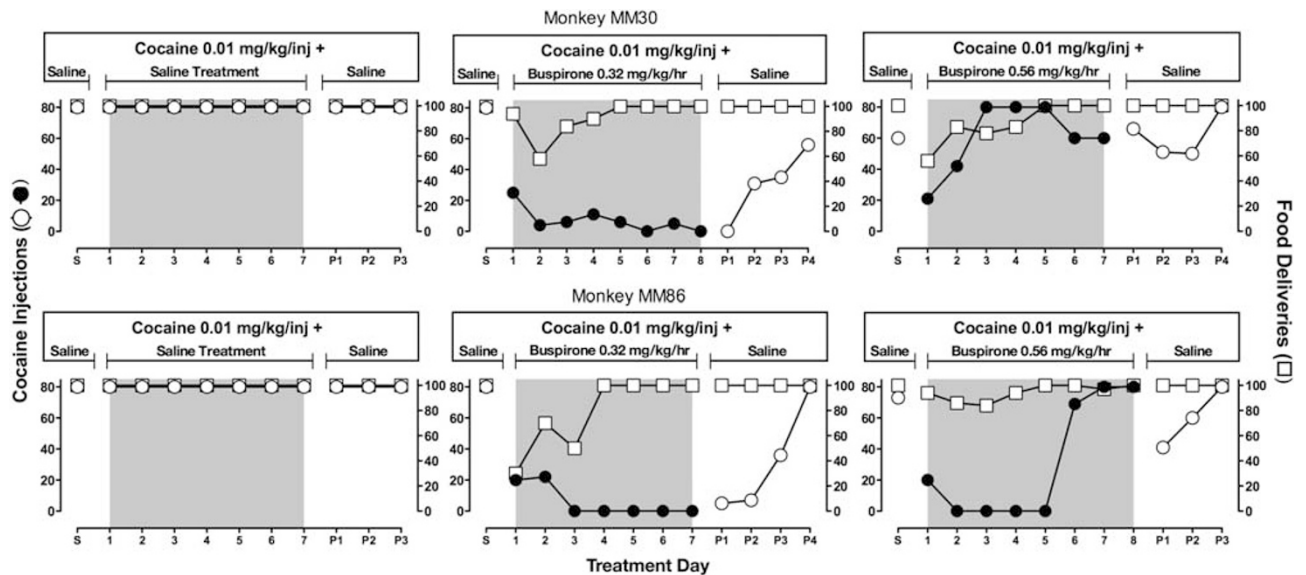


Figure 4 The effects of buspirone on cocaine- and food-maintained responding in individual monkeys (MM 30 and MM 86). Abscissae: consecutive days during and following saline (top panel) or buspirone treatment (0.32 mg/kg/h; middle panel) or (0.56 mg/kg/h; bottom panel). Left ordinates: number of cocaine (0.01 mg/kg/injection) injections per day is shown as open circles during saline treatment and closed circles during 0.32 mg/kg/h and 0.56 mg/kg/h buspirone treatment. Right ordinates: number of food pellets (1 g) earned per day is shown as open squares. Points above saline (S) in each panel show the mean number of injections/day and food pellets/day during 3 days of saline treatment baseline control conditions (mean + SEM).

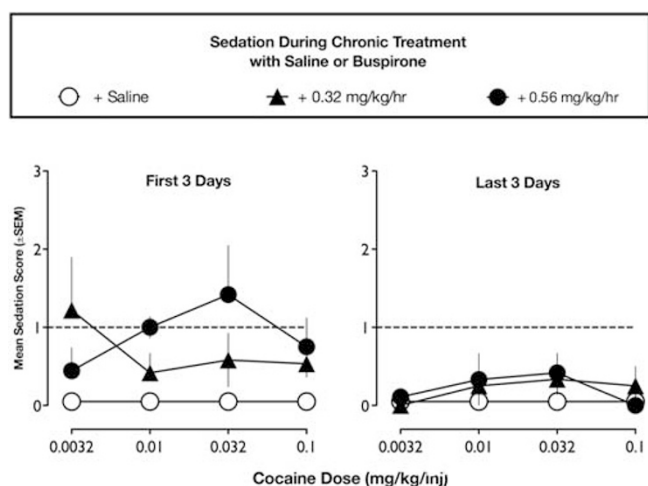


Figure 5 Sedation ratings during chronic buspirone treatment and cocaine self-administration. Cocaine unit doses are shown on the abscissae, and sedation scale ratings immediately following the noon cocaine self-administration session are shown on the left ordinates. Sedation was scored as: 0 = no observable sedation; alert to environment; 1 = mildly sedated, quieter than usual, but reaches for food treats; 2 = moderately sedated, does not reach for food treats but does respond to noise in the room; 3 = heavily sedated, lying on floor of cage, no response to experimenter. Sedation ratings during chronic treatment with saline (open circles), 0.32 (filled triangles), and 0.56 (filled circles) mg/kg/h buspirone are shown during the first 3 days of treatment (left panel) and during the last 3 days of treatment (right panel). Each data point for saline and 0.01–0.10 mg/kg/injection cocaine reflects 4–5 monkeys. Data for 0.0032 mg/kg/injection cocaine during buspirone treatment reflect three monkeys.

Figure 4b (row 2) shows that cocaine self-administration decreased to about 25% of control levels when treatment with 0.32 mg/kg/h buspirone was initiated. Subsequently, cocaine self-administration was completely abolished on treatment days 3–7. Food-maintained responding was also initially reduced to about 30% of control levels but returned to baseline within 4 days. When treatment with a higher dose of buspirone (0.56 mg/kg/h) began, cocaine self-administration immediately fell to 25% of control levels and remained low for 5 days (Figure 2b, row 3). On days 6, 7, and 8, cocaine self-administration returned to control levels. Upon cessation of buspirone treatment, cocaine self-administration fell to about 50% of control levels then returned to baseline by days 3 and 4. Food-maintained responding was minimally affected during or after buspirone treatment.

Recovery of Cocaine Self-Administration and Food-Maintained Responding After Buspirone Treatment

After saline treatment, monkeys usually met stability criteria within 3 days. However, at the lowest dose of cocaine (0.0032 mg/kg/injection), monkeys took an average of $8.0 (\pm 2.88)$ days to reach stability after saline treatment (Figure 4 a and b). At the highest cocaine dose studied (0.1 mg/kg/injection), recovery was more rapid and took an average of 5.6 ± 1.78 days. When 0.32 mg/kg/h buspirone was discontinued, recovery of cocaine self-administration increased by an average of $4.4 (\pm 2.42)$ days and ranged from 2 to 26 days. The increase in days to reach stability was

consistent across all cocaine doses. When 0.56 mg/kg/h buspirone was discontinued, recovery of cocaine self-administration increased by an average of 2.38 (± 2.55) days. Food-maintained responding was not disrupted at any cocaine dose during saline treatment and all monkeys met stability criteria within 3 days post-treatment. Although a slight decrease in food-maintained responding was evident during treatment with 0.32 and 0.56 mg/kg/h buspirone treatment (see Figure 3), food-maintained responding recovered almost immediately after treatment cessation.

Buspirone Plasma Levels

After 48 h of exposure to 0.32 mg/kg/h, IV buspirone, in the absence of concurrent cocaine self-administration, buspirone plasma levels averaged 60.2 ± 6.85 ng/ml. The buspirone plasma values ranged from 39 to 74 ng/ml.

Buspirone's Side Effects and Safety During Chronic Treatment

Ratings of buspirone's sedative effects, recorded immediately following the noon cocaine self-administration sessions, are shown in Figure 5. Data are shown for the first 3 days after buspirone treatment began (left panel), and for the last 3 days before the treatment was terminated (right panel). There were minimal sedative effects immediately following each noon cocaine self-administration session. Monkeys were alert, responsive to the investigator, and would take food treats. During buspirone treatment (0.32 mg/kg/h), sedation scores indicated either mild sedation or no sedation. At the higher dose of buspirone, only one monkey was heavily sedated, and all other scores indicated either mild (1) or no sedation (0) at all doses of cocaine. In general, sedative effects with the higher dose of buspirone treatment were lower during the last 3 days than during the first 3 days of treatment.

Buspirone's Effects on Cocaine Discrimination

During the training days preceding test days, monkeys responded exclusively on the saline key during saline cycles and almost exclusively on the cocaine key ($99.95\% \pm 0.05$) during cocaine cycles. Mean response rates (\pm SEM) were $2.40 (\pm 0.44)$ and $2.18 (\pm 0.50)$ responses per second during saline and cocaine training cycles, respectively. Control data for each individual monkey are shown in the shaded areas in Figure 6 (for discriminative stimulus effects) and Figure 7 (for response rates). Figures 6 and 7 show the discriminative stimulus and rate-altering effects of cocaine administered alone or 15 min after pretreatment with buspirone (0.1–0.32 mg/kg). All monkeys showed dose-dependent increases in cocaine-like responding during baseline cumulative cocaine tests. ED_{50} values for cocaine discrimination in individual monkeys are shown in Table 1. In three monkeys (89B013, A056, and CH599), buspirone produced a significant rightward shift in the cocaine discrimination dose–effect curve. All monkeys responded at or near 100% at the training dose of cocaine (0.4 mg/kg) during baseline conditions. Buspirone pretreatment reduced cocaine-like responding in four monkeys (89B013, A056, RQ1499, and CH599). The highest dose of buspirone (0.32 mg/kg) also

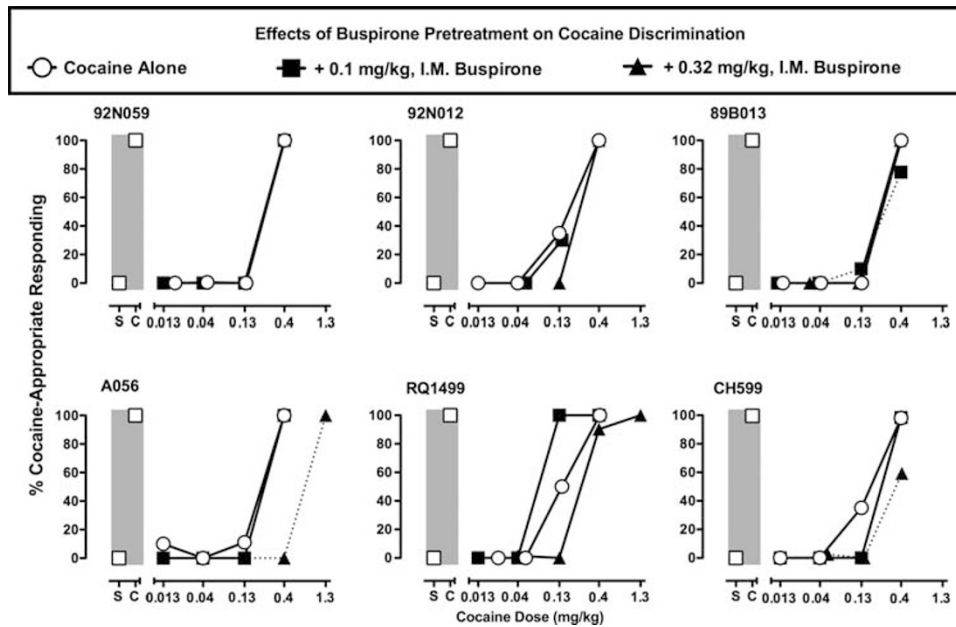


Figure 6 Discriminative stimulus effects of cocaine administered alone or 15 min after pretreatment with buspirone (0.1 or 0.32 mg/kg, IM). Abscissae: dose of cocaine in mg/kg (log scale). Ordinates: % cocaine-appropriate responding for the entire session. Points above 'S' and 'C' show control data obtained during saline and cocaine training cycles, respectively. Dose-effect curves for 'cocaine alone' show mean data from two determinations in each monkey. Dose-effect curves for cocaine + buspirone show data from one determination in each monkey. Dashed connecting lines indicate dose-effect curves with significantly greater ED₅₀ values compared with 'cocaine alone'.

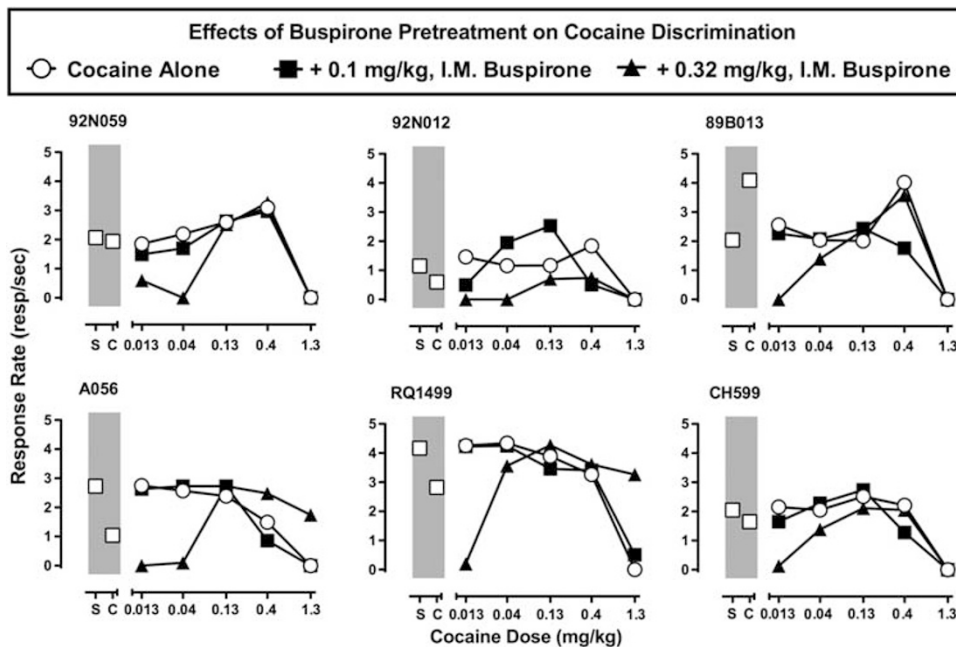


Figure 7 Rate-altering effects of cocaine administered alone or after pretreatment with buspirone: abscissae: dose cocaine in mg/kg (log scale). Ordinates: response rate in responses per second. Points above 'S' and 'C' show control data obtained during saline and cocaine training cycles, respectively. Dose-effect curves for 'cocaine alone' show mean data from two determinations in each monkey. Dose-effect curves for cocaine + buspirone show data from one determination in each monkey.

attenuated the rate-decreasing effects of the highest dose of 1.3 mg/kg cocaine in two of those monkeys.

In contrast to drug self-administration studies where buspirone was administered every 20 min as mg/kg/h, a single bolus IM dose in drug discrimination produced significant sedative effects. During the first two cycles of

cumulative cocaine tests, buspirone (0.32 mg/kg, IM) produced dose-dependent decreases in response rates, overt catalepsy-like behavior, and monkeys were unresponsive to approach of investigator. These rate-decreasing and catalepsy-like effects dissipated during later cycles (about 45–65 min after buspirone administration). Buspirone's transient

Table 1 ED₅₀ Values (mg/kg) of Cocaine Alone (95% confidence intervals) and in Combination with Buspirone for Substituting for Cocaine Drug Discrimination

ID	Cocaine alone	+ 0.1 mg/kg Buspirone	+ 0.32 mg/kg Buspirone
RQ1499	0.13 (0.04–0.40)	0.07	0.24
92N059	0.23 (0.23–0.23)	0.23	0.23
A056	0.19 (0.13–0.27)	0.23	0.72 ^a
92N012	0.16 (0.08–0.32)	0.29	0.23
CH599	0.16 (0.08–0.33)	0.23	0.34 ^a
89B013	0.23 (0.23–0.23)	0.25 ^a	0.23

^aDenotes ED₅₀ values after buspirone fell outside of the 95% confidence intervals of baseline ED₅₀ values.

disruption of behavior may indicate that these effects were attenuated by increasing cumulative doses of cocaine, or may reflect buspirone's relatively short duration of action.

DISCUSSION

This report describes the effects of chronic buspirone treatment on the reinforcing effects of cocaine. These findings confirm and extend our previous findings that acute doses of buspirone dose-dependently reduced cocaine self-administration by nonhuman primates (Bergman *et al*, 2012). Buspirone was selected for study, in part, because it is FDA approved for the treatment of anxiety. Our finding that chronic buspirone can selectively reduce cocaine self-administration by rhesus monkeys may have important clinical implications. The preclinical IV drug self-administration model used in this study has good predictive validity for drug abuse treatment medications that are clinically available (Mello, 2005).

The relation of our data to previous reports of the interactions between buspirone and cocaine in drug self-administration and drug discrimination studies are described below. Possible receptor mechanisms contributing to buspirone's effects on cocaine-maintained responding are discussed. Buspirone's dopamine D₂, D₃, and D₄ receptor activity are examined in the context of recent discoveries of the potential role of D₃ antagonists and partial agonists in modulating cocaine's abuse-related effects. Some strengths, limitations, and translational implications of this study are also described.

Buspirone's Effects on Cocaine Self-Administration

Our results using a chronic buspirone treatment paradigm are consistent with our previous study of the acute effects of buspirone on cocaine self-administration (Bergman *et al*, 2012), and overlap with data from previous studies in rhesus monkeys (Gold and Balster, 1992). A single high dose of buspirone (1.0 mg/kg, IV) administered 15 min before a 1-h cocaine self-administration session, decreased cocaine-maintained responding, whereas acute or repeated administration of lower doses (0.1–0.56 mg/kg, IV) had no effect or increased responding (Gold and Balster, 1992). It was suggested that buspirone's relatively short duration of

action may have contributed to the observed lack of effect on cocaine self-administration (Gold and Balster, 1992). In the present study, IV buspirone was administered every 20 min for 23 h each day for 7–10 days to insure that significant levels of buspirone and its metabolites were present throughout each food and cocaine self-administration session conducted between 1100 and 2000 hours each day (see Figure 1). Our current findings suggest that 0.32 mg/kg/h buspirone is sufficient to selectively reduce cocaine self-administration by rhesus monkeys, and a slightly higher dose of buspirone is not more effective.

In the present study, buspirone (0.32 and 0.56 mg/kg/h) did not enhance the stimulant effects of cocaine over a dose range of 0.0032–0.10 mg/kg/injection. However, in squirrel monkeys, buspirone- (0.001–0.03 mg/kg) related increases in cocaine-maintained responding were observed under a second-order schedule (Nader and Barrett, 1990). It is likely that this reflected disruptions in responding rather than direct stimulant effects of buspirone because IV cocaine was not given until the end of the session (Nader and Barrett, 1990). In rats, buspirone's effects on cocaine self-administration varied as a function of the schedule of reinforcement (Homberg *et al*, 2004). Buspirone (1.25 and 2.5 mg/kg) significantly reduced cocaine self-administration maintained on a progressive ratio schedule, but increased cocaine self-administration when the response requirement was low (FR1) (Homberg *et al*, 2004). Similar effects have been reported for D₃ selective antagonist and partial agonist treatment of cocaine self-administration in rats (Gal and Gyertyan, 2003; Gyertyan *et al*, 2007; Heidbreder *et al*, 2005).

Buspirone's Effects on Cocaine Discrimination

There has been relatively little attention to the effects of buspirone on cocaine discrimination. In the present study, buspirone (0.1 and 0.32 mg/kg, IM) did not consistently alter the discriminative stimulus and rate-altering effects of cocaine, and similar findings have been reported in rats (Rapoza, 1993). We interpret these data to suggest that monkeys were able to recognize cocaine, even after relatively high acute doses of buspirone. However, these data are not concordant with previous findings in rats trained to discriminate cocaine (5 mg/kg, IM) from saline (Callahan and Cunningham, 1997). In that study, buspirone (2.5–20 mg/kg) dose-dependently reduced cocaine discrimination and shifted the cocaine dose-effect function to the right (Callahan and Cunningham, 1997). A combination of procedural and species differences may have contributed to these discrepant findings. In rhesus monkeys trained to discriminate cocaine (0.1 mg/kg, IV) from saline, buspirone (0.03–0.56 mg/kg, IV) did not substitute for cocaine, but the effects of buspirone pretreatment were not tested (Koetner, 1996). The finding that buspirone did not substitute for cocaine (Koetner, 1996) is consistent with reports that buspirone does not maintain self-administration in rhesus monkeys and baboons, and has low abuse potential (Balster, 1990; Balster and Woolverton, 1982; Griffiths *et al*, 1991).

Buspirone's Receptor Mechanisms

The mechanisms by which buspirone selectively reduces cocaine self-administration by rhesus monkeys are unclear,

and probably reflect an interaction between dopamine and 5-HT_{1A} receptors. Recent radioligand-binding studies indicated that bupirone has high affinity for dopamine D₃ and D₄ receptors, as well as for D₂ receptors (Bergman *et al*, 2012). The functional activity of bupirone also was examined in live cell β -Arrestin assays. It was found that bupirone had no agonist activity, and antagonized dopamine activation of D₃ and D₄ receptors with similar potencies. We concluded that bupirone was a functional antagonist at D₂-like dopamine receptors with the highest affinity/potency for the D₃ and D₄ receptor subtypes (Bergman *et al*, 2012). These findings were consistent with earlier radioligand-binding studies of bupirone (Kula *et al*, 1994; Tallman *et al*, 1997).

Although the present study does not permit a clear distinction between the role of D₂, D₃, and D₄ receptors in bupirone's effects on cocaine self-administration, it may be useful to review recent evidence concerning bupirone's D₂, D₃, and D₄ activity. Many investigators have concluded that bupirone's behavioral effects are primarily mediated through D₂ antagonist actions (Callahan and Cunningham, 1997; Rijnders and Slangen, 1993). This conclusion is usually based on the similarity between the effects of bupirone and other known D₂ antagonists. For example, bupirone's effects were similar to those of haloperidol and supiride in drug discrimination studies in rats (Rijnders and Slangen, 1993). In addition, the finding that gepirone, a selective 5-HT_{1A} partial agonist, did not affect cocaine's discriminative stimulus effects in rats was interpreted to suggest that bupirone's reduction of cocaine's discriminative stimulus effects primarily reflected its D₂ antagonist properties and not its 5-HT_{1A} agonist effects (Callahan and Cunningham, 1997). Gepirone also failed to alter cocaine self-administration by rhesus monkeys (Gold and Balster, 1992). Nonetheless, an influence of 5-HT_{1A} receptors on stimulant abuse is possible, and may involve complex pre- and post-synaptic actions (Muller *et al*, 2007).

An emerging literature attests to the potential importance of dopamine D₃ and D₄ receptors activity in the modulation of cocaine's abuse-related effects (see for review Heidbreder *et al*, 2005; Heidbreder and Newman, 2010; Le Foll *et al*, 2005; Le Foll *et al*, 2000; Newman *et al*, 2005; Newman *et al*, 2012). Dopamine D₃ receptor agonists, antagonists, and partial agonists each can influence the behavioral effects of cocaine under some conditions. However, the relative effectiveness of D₃ receptor partial agonists and antagonists for reducing the abuse-related effects of cocaine remains unresolved. D₃ selective agonists produced cocaine-like effects in rhesus and squirrel monkeys, and in rats trained to discriminate cocaine from saline (Achat-Mendes *et al*, 2010; Acri *et al*, 1995; Lamas *et al*, 1996; Spealman, 1996). D₃ selective agonists also maintained self-administration in rats (Caine and Koob, 1993) and rhesus monkeys (Nader and Mach, 1996). Consistent with clinical reports that agonist therapies are effective in reducing cocaine abuse (Herin *et al*, 2010), the D₃ preferring agonists, such as 7-OH-DPAT, also have been shown to reduce cocaine self-administration by rats (Caine and Koob, 1995).

Like D₃ receptor agonists, D₃ receptor antagonists also decrease cocaine self-administration under several behavioral procedures in rhesus monkeys, but do not appear to have abuse potential. Novel D₃ antagonists and partial

agonists each attenuated the reinforcing and discriminative stimulus effects of cocaine under some conditions in rhesus monkeys (Beardsley *et al*, 2001; Claytor *et al*, 2006; Martelle *et al*, 2007; Nader *et al*, 1999). In contrast, in squirrel monkeys, a D₃ partial agonist and a D₃ antagonist did not alter cocaine self-administration (Achat-Mendes *et al*, 2010; Achat-Mendes *et al*, 2009). In comparison, a D₂ antagonist, studied under the same conditions, decreased cocaine self-administration and cocaine discrimination, and cocaine-induced reinstatement (Achat-Mendes *et al*, 2010). In a reinstatement paradigm, D₃ and D₂ agonists dose-dependently reinstated cocaine seeking and D₃ and D₂ antagonists attenuated cocaine reinstatement (Achat-Mendes *et al*, 2010).

In rats, selective dopamine D₃ receptor antagonists (SB-27701_{1A} and NGB 2904) reduced cocaine self-administration maintained on a progressive ratio schedule, and significantly lowered progressive ratio breakpoints (Xi *et al*, 2005; Xi *et al*, 2006); see for review Xi and Gardner, 2007). Direct infusion of SB-27701_{1A} into the basolateral amygdala (but not the nucleus accumbens shell or dorsal striatum) also reduced cocaine self-administration maintained on a second-order schedule (Di Ciano, 2008). Selective D₃ receptor partial agonists (RGH-237) blocked cue-induced reinstatement of cocaine seeking, but had no effect on sucrose or water seeking (Gyertyan *et al* 2007). A number of investigators have reported that D₃ selective compounds have no effect on cocaine self-administration maintained on a low-response requirement, such as a FR 1 schedule (Gyertyan *et al*, 2007; Pilla *et al*, 1999; Xi and Gardner, 2007; see for review Heidbreder *et al*, 2005), but are effective at higher response requirements (eg, FR 10 and progressive ratio schedules) (Xi and Gardner, 2007; Xi *et al*, 2005). Selective D₃ receptor antagonists also inhibited cocaine-induced reinstatement (Gyertyan *et al*, 2007), biphasically increased and decreased cocaine self-administration maintained on a FR schedule (Gyertyan *et al*, 2007), and reduced cocaine-enhanced brain stimulation (Peng *et al*, 2009). A D₃ receptor antagonist reduced cocaine self-administration maintained on both fixed and progressive ratio schedule in rats and mice (Song *et al*, 2012). Interestingly, a D₃ antagonist did not inhibit cocaine self-administration in D₃ receptor knockout mice (Song *et al*, 2012). Taken together, these data illustrate that the response requirement is a critical determinant of dopamine D₃ receptor effects on the abuse-related effects of cocaine.

Strengths and Limitations of the Study

This evaluation of chronic daily bupirone treatment was designed to model clinical treatment trials. An important strength of this study is that treatment effects on both food- and cocaine-maintained responding were studied. This procedure facilitates evaluation of the extent to which the effects of a treatment drug are selective for cocaine or reduces both food- and cocaine-maintained responding, indicating a general disruption of operant responding. We conclude that the sustained decreases in cocaine self-administration reflected the effects of bupirone treatment and not other confounding factors for the following reasons: (1) there was no evidence of catheter malfunction because all monkeys resumed cocaine self-administration at saline

treatment baseline levels after buspirone treatment; (2) decreases in cocaine self-administration were not associated with sedative effects that interfered with operant responding; (3) buspirone, across the dose range studied on cocaine self-administration, did not produce catatonia or other adverse behavioral effects; (4) buspirone's effects were selective for cocaine and had modest initial effects on food-maintained responding that usually dissipated during the course of treatment. Moreover, this is the first study in which buspirone was administered intermittently (every 20 min for 23 h) throughout each day to insure that doses were stable during food and cocaine sessions. Finally, the same monkeys were studied as their own control across successive treatment conditions to minimize any influence of idiosyncratic individual reactions.

One possible limitation of this study is that buspirone doses used clinically for the treatment of anxiety are usually lower than the dose necessary to reduce cocaine self-administration by rhesus monkeys (0.32 mg/kg/h). After oral administration, the usual clinical dose of buspirone (20–40 mg in divided doses) reached maximum concentrations within 1 h and the elimination half-life was about 2.5 h (Gammans *et al*, 1986; Goldberg, 1984; Mahmood and Sahajwalla, 1999; Mandrioli and Mercolini, 2010). In a clinical trial to evaluate the effectiveness of buspirone for prevention of relapse to cocaine, oral doses of 60 mg/day are planned (Winhusen *et al*, 2012).

There has been relatively little attention to the pharmacokinetics of buspirone in rhesus monkeys. Two reports conclude that the pharmacokinetics of buspirone appear to be similar in humans and in rhesus monkeys (Gammans *et al*, 1986; Marathe *et al*, 1999). For example, plasma concentrations of C14 labeled buspirone were similar over 1 h (Gammans *et al*, 1986). When rhesus monkeys were given repeated oral doses of buspirone (12.5, 25 and 50 mg/kg/day), the peak levels (C_{max}) were dose dependent, but there was considerable variability between animals (Marathe *et al*, 1999). We also observed considerable variability between animals in plasma buspirone levels after repeated IV doses for 48 h (0.32 mg/kg/h). Although the plasma levels in rhesus monkeys ranged from 39 to 74 ng/ml, it is important to note that there were no adverse behavioral or medical consequences of chronic buspirone treatment. There were no significant changes in blood chemistry assessments at 3, 4, 10, or 14 days after discontinuation of buspirone. Differences in the route (oral *vs* IV) and rate (bolus *vs* three injections per hour) of buspirone administration in humans and rhesus monkeys as well as differences in analytic procedures (gas chromatography, radioimmunoassay) make it difficult to compare effective doses.

Translational Implications of Buspirone's Reduction of Cocaine Self-Administration by Rhesus Monkeys

Evaluations of candidate medications in this preclinical drug self-administration model have shown good concordance with clinical treatment trials (Mello, 2005). This model is valuable for medication development, because often candidate treatment drugs are not FDA approved for evaluation in humans. The reliability and predictive validity of this model has been repeatedly demonstrated (see for review Haney and

Spealman, 2008; Mello, 2005; Mello and Negus, 1996). An optimal treatment medication would have low abuse liability and minimal side effects in addition to selective reduction of the abuse-related effects of the target drug of abuse. Buspirone appears to have no abuse liability in rhesus monkeys (Balster, 1990) and, despite its putative anxiolytic effects, increased tension, nervousness, and dysphoria were reported in clinical laboratory studies (Cole, 1982; Griffith *et al*, 1986; Rush *et al*, 1995). Chronic buspirone treatment produced only mild and transient sedation in rhesus monkeys in the present study, and did not increase ratings of drowsy or sleepy in humans (Rush *et al*, 1995). It is possible that the combination of dopamine antagonism and 5-HT_{1A} effects may be valuable in the development of medications for cocaine abuse. Some side effects usually observed during treatment with dopamine D₂ receptor antagonists (catatonia, catalepsy) were not observed during buspirone treatment and cocaine self-administration. This observation is consistent with an extensive literature showing that 5-HT_{1A} agonists, including buspirone, attenuate dopamine D₂-induced catalepsy (Kleven *et al*, 2005; Prinssen *et al*, 2002; Prinssen *et al*, 2000).

In summary, the present study indicates that chronic buspirone treatment reduced IV cocaine self-administration by nonhuman primates. Although it is unlikely that a single drug will be effective in all individuals, if buspirone proves to be as effective in clinical settings as in the preclinical laboratory, it will be a useful addition to the current treatment options for cocaine abuse. Although the relative contribution of buspirone's 5-HT_{1A} and dopamine D₂, D₃, and D₄ activity remains to be determined, buspirone has the important advantage of FDA approval for clinical use for another indication. Basic research to clarify the effects of 5-HT_{1A} and dopamine receptor subtypes on the abuse-related effects of cocaine are urgently needed to optimize such strategies for medication development.

ACKNOWLEDGEMENTS

This work was supported in part by grant number R01-DA002519 and R01-DA026892 (NK Mello, PI) from the National Institute on Drug Abuse, NIH. We thank Meredith Mahnke, Sherilyn Pattel, and Olga Smirnova for excellent technical assistance.

DISCLOSURE

The authors declare no conflict of interest.

REFERENCES

- Achat-Mendes C, Grundt P, Cao J, Platt DM, Newman AH, Spealman RD (2010). Dopamine D3 and D2 receptor mechanisms in the abuse-related behavioral effects of cocaine: studies with preferential antagonists in squirrel monkeys. *J Pharmacol Exp Ther* 334: 556–565.
- Achat-Mendes C, Platt DM, Newman AH, Spealman RD (2009). The dopamine D3 receptor partial agonist CJB 090 inhibits the discriminative stimulus but not the reinforcing or priming effects of cocaine in squirrel monkeys. *Psychopharmacology (Berl)* 206: 73–84.

- Acri JB, Carter SR, Alling K, Geter-Douglass B, Dijkstra D, Wikstrom H *et al* (1995). Assessment of cocaine-like discriminative stimulus effects of dopamine D3 receptor ligands. *Eur J Pharmacol* **281**: R7–R9.
- Balster RL (1990). Abuse potential of buspirone and related drugs. *J Clin Psychopharmacol* **10**: 31S–37S.
- Balster RL, Woolverton WL (1982). Intravenous buspirone self-administration in rhesus monkeys. *J Clin Psychiatry* **43**: 34–37.
- Beardsley PM, Sokoloff P, Balster RL, Schwartz JC (2001). The D3R partial agonist, BP 897, attenuates the discriminative stimulus effects of cocaine and D-amphetamine and is not self-administered. *Behav Pharmacol* **12**: 1–11.
- Bergman J, Roof RA, Furman CA, Conroy JL, Mello NK, Sibley DR *et al* (2012). Modification of cocaine self-administration by buspirone (Buspar®): potential involvement of D3 and D4 dopamine receptors. *Int J Neuropsychopharmacol* **25**: 1–14.
- Butelman ER, Harris TJ, Kreek MJ (1999). Effects of E-2078, a stable dynorphin A(1–8) analog, on sedation and serum prolactin levels in rhesus monkeys. *Psychopharmacology (Berl)* **147**: 73–80.
- Caine SB, Koob GF (1993). Modulation of cocaine self-administration in the rat through D-3 dopamine receptors. *Science* **260**: 1814–1816.
- Caine SB, Koob GF (1995). Pretreatment with the dopamine agonist 7-OH-DPAT shifts the cocaine self-administration dose-effect function to the left under different schedules in the rat. *Behav Pharmacol* **6**: 333–347.
- Callahan PM, Cunningham KA (1997). Modulation of the discriminative stimulus properties of cocaine: comparison of the effects of fluoxetine with 5-HT_{1A} and 5-HT_{1B} receptor agonists. *Neuropharmacology* **36**: 373–381.
- Cinciripini PM, Lapitsky L, Seay S, Wallfisch A, Meyer WJI, Van Vunakis H (1995). A placebo-controlled evaluation of the effects of buspirone on smoking cessation: differences between high- and low-anxiety smokers. *J Clin Pharmacol* **15**: 182–191.
- Clayton R, Lile JA, Nader MA (2006). The effects of eticlopride and the selective D3-antagonist PNU 99194-A on food- and cocaine-maintained responding in rhesus monkeys. *Pharmacol Biochem Behav* **83**: 456–464.
- Cole JO (1982). New anxiety agents. *Psychopharmacol Bull* **18**: 45–46.
- Di Ciano P (2008). Drug seeking under a second-order schedule of reinforcement depends on dopamine D3 receptors in the basolateral amygdala. *Behav Neurosci* **122**: 129–139.
- Dockens RC, Salazar DE, Fulmore IE, Wehling M, Arnold ME, Croop R (2006). Pharmacokinetics of a newly identified active metabolite of buspirone after administration of buspirone over its therapeutic dose range. *J Clin Pharmacol* **46**: 1308–1312.
- Elkashef A, Vocci FJ, Huestis M, Haney M, Budney A, Gruber A *et al* (2008). Marijuana neurobiology and treatment. *Subst Abuse* **29**: 17–29.
- Fivel PA (2011). Computer-controlled drug doses for IV drug self-administration. *Exp Clin Psychopharmacol* **19**: 131–133.
- Gal K, Gyertyan I (2003). Targeting the dopamine D3 receptor cannot influence continuous reinforcement cocaine self-administration in rats. *Brain Res Bull* **61**: 595–601.
- Gammans RE, Mayol RF, LaBudde JA (1986). Metabolism and disposition of buspirone. *Am J Med* **80**: 41–51.
- Gold LH, Balster RL (1992). Effects of buspirone and gepirone on i.v. cocaine self-administration in rhesus monkeys. *Psychopharmacology (Berl)* **108**: 289–294.
- Goldberg HL (1984). Buspirone hydrochloride: a unique new anxiolytic agent. Pharmacokinetics, clinical pharmacology, abuse potential and clinical efficacy. *Pharmacotherapy* **4**: 315–324.
- Griffith JD, Jasinski DR, Casten GP, McKinney GR (1986). Investigation of the abuse liability of buspirone in alcohol-dependent patients. *Am J Med* **80**: 30–35.
- Griffiths RR, Lamb RJ, Sannerud CA, Ator NA, Brady JV (1991). Self-injection of barbiturates, benzodiazepines and other sedative-anxiolytics in baboons. *Psychopharmacology (Berl)* **103**: 154–161.
- Gyertyan I, Kiss B, Gal K, Laszlovszky I, Horvath A, Gemesi LI *et al* (2007). Effects of RGH-237 [N-{4-[4-(3-aminocarbonyl-phenyl)-piperazin-1-yl]-butyl}-4-bromo-benzamide], an orally active, selective dopamine D(3) receptor partial agonist in animal models of cocaine abuse. *J Pharmacol Exp Ther* **320**: 1268–1278.
- Haney M, Spealman R (2008). Controversies in translational research: drug self-administration. *Psychopharmacology (Berl)* **199**: 403–419.
- Heidbreder CA, Gardner EL, Xi Z-X, Thanos PK, Mugnaini M, Hagan JJ *et al* (2005). The role of central dopamine D3 receptors in drug addiction: a review of pharmacological evidence. *Brain Res Rev* **49**: 77–105.
- Heidbreder CA, Newman AH (2010). Current perspectives on selective dopamine D₃ receptor antagonists as pharmacotherapeutics for addictions and related disorders. *Ann NY Acad Sci* **1187**: 4–34.
- Henningfield JE, Fant RV, Buchhalter AR, Stitzer ML (2005). Pharmacotherapy for nicotine dependence. *CA Cancer J Clin* **55**: 281–299, quiz 322–283, 325.
- Herin DV, Rush CR, Grabowski J (2010). Agonist-like pharmacotherapy for stimulant dependence: preclinical, human laboratory, and clinical studies. *Ann NY Acad Sci*.
- Hilleman DE, Mohiuddin SM, Del Core MG, Sketch MH Sr. (1992). Effect of buspirone on withdrawal symptoms associated with smoking cessation. *Arch Intern Med* **152**: 350–352.
- Homberg JR, Arends B, Wardeh G, Raaso HS, Schoffelmeer AN, de Vries TJ (2004). Individual differences in the effects of serotonergic anxiolytic drugs on the motivation to self-administer cocaine. *Neuroscience* **128**: 121–130.
- ILAR-NRC (1996). *Guide for the Care and Use of Laboratory Animals*. National Academy Press: Washington, DC, pp 125.
- Kleven MS, Barret-Grevoz C, Bruins-Slot L, Newman-Tancredi A (2005). Novel antipsychotic agents with 5-HT_{1A} agonist properties: role of 5-HT_{1A} receptor activation in attenuation of catalepsy induction in rats. *Neuropharmacology* **49**: 135–143.
- Koetznner L, Riley AL, Glowa JR (1996). Discriminative stimulus effects of dopaminergic agents in rhesus monkeys. *Pharmacol Biochem Behav* **54**: 517–523.
- Kula NS, Baldessarini RJ, Keabian JW, Neumeyer JL (1994). S-(+)-aporphines are not selective for human D3 dopamine receptors. *Cell Mol Neurobiol* **14**: 185–191.
- Lamas X, Negus SS, Nader MA, Mello NK (1996). Effects of the putative dopamine D3 receptor agonist 7-OH-DPAT in rhesus monkeys trained to discriminate cocaine from saline. *Psychopharmacology (Berl)* **124**: 306–314.
- Le Foll B, Goldberg SR, Sokoloff P (2005). D3 receptor and drug dependence: effects on reward or beyond? *Neuropharmacology* **49**: 525–541.
- Le Foll B, Schwartz J-C, Sokoloff P (2000). Dopamine D3 receptor agents as potential new medications for drug addiction. *Eur Psychiatry* **15**: 140–146.
- Line SW (1987). Environmental enrichment for laboratory primates. *J Am Vet Med Assoc* **190**: 854–859.
- Mahmood I, Sahajwalla C (1999). Clinical Pharmacokinetics and pharmacodynamics of buspirone, an anxiolytic drug. *Clin Pharmacokinet* **36**: 277–287.
- Mandrioli R, Mercolini L, Raggi MA (2010). Metabolism of benzodiazepine and non-benzodiazepine anxiolytic-hypnotic drugs: an analytical point of view. *Curr Drug Metab* **11**: 815–829.
- Marathe PH, Shen F, Markham P, Greene DS (1999). Pharmacokinetics of buspirone following oral administration to rhesus monkeys. *J Pharm Pharmacol* **51**: 601–607.
- Martelle JL, Clayton R, Ross JT, Reboussin BA, Newman AH, Nader MA (2007). Effects of two novel D3-selective compounds, NGB

- 2904 [N-(4-(4-(2, 3-dichlorophenyl) piperazin-1-yl) butyl)-9H-fluorene-2-carboxamide] and CJB 090 [N-(4-(4-(2, 3-dichlorophenyl) piperazin-1-yl) butyl)-4-(pyridin-2-yl) benzamide], on the reinforcing and discriminative stimulus effects of cocaine in rhesus monkeys. *J Pharmacol Exp Ther* **321**: 573–582.
- McRae-Clark AL, Carter RE, Killeen TK, Carpenter MJ, Wahlquist AE, Simpson SA et al (2009). A placebo-controlled trial of buspirone for the treatment of marijuana dependence. *Drug Alc Depend* **105**: 132–138.
- Mello NK (2005). Marian W Fischman memorial lecture (2004). Evaluation of drug abuse treatment medications: concordance between clinical and preclinical studies. In: Dewey WL (ed). *Problems of Drug Dependence 2004: Proceedings of the 66th Annual Scientific Meeting, The College on Problems of Drug Dependence, Inc.* US Department of Health and Human Services, National Institutes of Health: Bethesda, MD, pp 82–104.
- Mello NK, Negus SS (1996). Preclinical evaluation of pharmacotherapies for treatment of cocaine and opiate abuse using drug self-administration procedures. *Neuropsychopharmacology* **14**: 375–424.
- Mello NK, Negus SS (2007). Effects of d-amphetamine and buprenorphine combinations on speedball (cocaine + heroin) self-administration by rhesus monkeys. *Neuropsychopharmacology* **32**: 1985–1994.
- Mello NK, Negus SS, Lukas SE, Mendelson JH, Sholar JW, Drieze JM (1995). A primate model of polydrug abuse: cocaine and heroin combinations. *J Pharmacol Exp Ther* **274**: 1325–1337.
- Muller CP, Carey RJ, Huston JP, De Souza Silva MA (2007). Serotonin and psychostimulant addiction: focus on 5-HT_{1A}-receptors. *Prog Neurobiol* **81**: 133–178.
- Nader MA, Barrett JE (1990). Effects of chlordiazepoxide, buspirone, and serotonin receptor agonists and antagonists on responses of squirrel monkeys maintained under second-order schedules of intramuscular cocaine injection or food presentation. *Drug Dev Res* **20**: 5–17.
- Nader MA, Green KL, Luedtke RR, Mach RH (1999). The effects of benzamide analogues on cocaine self-administration in rhesus monkeys. *Psychopharmacology (Berl)* **147**: 143–152.
- Nader MA, Mach RH (1996). Self-administration of the dopamine D3 agonist 7-OH-DPAT in rhesus monkeys is modified by previous cocaine exposure. *Psychopharmacology (Berl)* **125**: 13–22.
- Negus SS, Mello NK (2003). Effects of chronic d-amphetamine treatment on cocaine- and food-maintained responding under a second-order schedule in rhesus monkeys. *Drug Alc Depend* **70**: 39–52.
- Newman AH, Blaylock BL, Nader MA, Bergman J, Sibley DR, Skolnick P (2012). Medication discovery for addiction: translating the dopamine D3 receptor hypothesis. *Biochem Pharmacol* **84**: 882–890.
- Newman AH, Grundt P, Nader MA (2005). Dopamine D3 receptor partial agonists and antagonists as potential drug abuse therapeutic agents. *J Med Chem* **48**: 3663–3679.
- Newman-Tancredi A, Gavaudan S, Conte C, Chaput C, Touzard M, Verrielle L et al (1998). Agonist and antagonist actions of antipsychotic agents at 5-HT_{1A} receptors: a [35S]GTPgammaS binding study. *Eur J Pharmacol* **355**: 245–256.
- Peng X-Q, Ashby CR, Spiller K, Li X, Li J, Thomasson N et al (2009). The preferential dopamine D3 receptor antagonist S33138 inhibits cocaine reward and cocaine-triggered relapse to drug-seeking behavior in rats. *Neuropharmacology* **56**: 752–760.
- Pilla M, Perachon S, Sautel F, Garrido F, Mann A, Wermuth CG et al (1999). Selective inhibition of cocaine-seeking behaviour by a partial dopamine D3 receptor agonist. *Nature* **400**: 371–375.
- Prinssen EPM, Colpaert FC, Koek W (2002). 5-HT_{1A} receptor activation and anti-cataleptic effects: high-efficacy agonists maximally inhibit haloperidol-induced catalepsy. *Eur J Pharmacol* **453**: 217–221.
- Prinssen EPM, Koek W, Colpaert FC, Kleven MS (2000). Repeated treatment with 8-OH-DPAT induces tolerance to its ability to produce the 5-HT_{1A} behavioural syndrome, but not to its ability to attenuate haloperidol-induced catalepsy. *Behav Pharmacol* **11**: 299–305.
- Rapoza D (1993). Buspirone fails to affect the discriminative stimulus effects of cocaine. *Pharmacol Biochem Behav* **45**: 179–183.
- Rijnders HJ, Slangen JL (1993). The discriminative stimulus properties of buspirone involve dopamine-2 receptor antagonist activity. *Psychopharmacology (Berl)* **111**: 55–61.
- Rush CR, Critchfield TS, Troisi JR, Griffiths RR (1995). Discriminative stimulus effects of diazepam and buspirone in normal volunteers. *J Exp Anal Behav* **63**: 277.
- Sheskin DJ (2000). *Handbook of Parametric and Nonparametric Statistical Procedures*. 2nd edn. Chapman and Hall: Boca Raton, FL.
- Song R, Yang R-F, Wu N, Su R-B, Li J, Peng X-Q et al (2012). YQ14: a novel dopamine D3 receptor antagonist that inhibits cocaine self-administration in rats and mice, but not in D3 receptor-knockout mice. *Addict Biol* **17**: 259–273.
- Spealman RD (1996). Dopamine D3 receptor agonists partially reproduce the discriminative stimulus effects of cocaine in squirrel monkeys. *J Pharmacol Exp Ther* **278**: 1128–1137.
- Tallman F, Primus RJ, Brodbeck R, Cornfield L, Meade R, Woodruff K et al (1997). I. NGD 94-1: identification of a novel, high-affinity antagonist at the human dopamine D4 receptor. *J Pharmacol Exp Ther* **282**: 1011–1019.
- Weinstein AM, Gorelick DA (2011). Pharmacological treatment of cannabis dependence. *Curr Pharm Des* **17**: 1351–1358.
- West R, Hajek P, McNeil A (1991). Effect of buspirone on cigarette withdrawal symptoms and short-term abstinence rates in a smokers clinic. *Psychopharmacology (Berl)* **104**: 91–96.
- Winhusen T, Brady KT, Stitzer M, Woody G, Lindblad R, Kropp F et al (2012). Evaluation of buspirone for relapse-prevention in adults with cocaine dependence: an efficacy trial conducted in the real world. *Contemp Clin Trials*.
- Wong H, Dockens RC, Pajor L, Yeola S, Grace JEJ, Stark AD et al (2007). 6-Hydroxybuspirone is a major active metabolite of buspirone: assessment of pharmacokinetics and 5-hydroxytryptamine_{1A} receptor occupancy in rats. *Drug Metab Dispos* **35**: 1387–1392.
- Xi Z-X, Gardner EL (2007). Pharmacological actions of NGB 2904, a selective dopamine D3 receptor antagonist, in animal models of drug addiction. *CNS Drug Rev* **13**: 240–259.
- Xi Z-X, Gilbert JG, Pak AC, Ashby CR Jr, Heidbreder CA, Gardner EL (2005). Selective dopamine D3 receptor antagonism by SB-277011A attenuates cocaine reinforcement as assessed by progressive-ratio and variable-cost-vrifiable-payoff fixed-ratio cocaine self-administration in rats. *Eur J Neurosci* **21**: 3427–3438.
- Xi Z-X, Newman AH, Gilbert JG, Pak AC, Peng X-Q, Ashby CR Jr et al (2006). The novel dopamine D3 receptor antagonist NGB 2904 inhibits cocaine's rewarding effects and cocaine-induced reinstatement of drug-seeking behavior in rats. *Neuropsychopharmacology* **31**: 1393–1405.