

R-Modafinil Attenuates Nicotine-Taking and Nicotine-Seeking Behavior in Alcohol-Preferring Rats

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(±)-Modafinil (MOD) is used clinically for the treatment of sleep disorders and has been investigated as a potential medication for the treatment of psychostimulant addiction. However, the therapeutic efficacy of (±)-MOD for addiction is inconclusive. Herein we used animal models of self-administration and *in vivo* microdialysis to study the pharmacological actions of R-modafinil (R-MOD) and S-modafinil (S-MOD) on nicotine-taking and nicotine-seeking behavior, and mechanisms underlying such actions. We found that R-MOD is more potent and effective than S-MOD in attenuating nicotine self-administration in Long–Evans rats. As Long–Evans rats did not show a robust reinstatement response to nicotine, we used alcohol-preferring rats (P-rats) that display much higher reinstatement responses to nicotine than Long–Evans rats. We found that R-MOD significantly inhibited intravenous nicotine self-administration, nicotine-induced reinstatement, and nicotine-associated cue-induced drug-seeking behavior in P-rats. R-MOD alone neither sustained self-administration in P-rats previously self-administering nicotine nor reinstated extinguished nicotine-seeking behavior. The *in vivo* brain microdialysis assays demonstrated that R-MOD alone produced a slow-onset moderate increase in extracellular DA. Pretreatment with R-MOD dose-dependently blocked nicotine-induced dopamine (DA) release in the nucleus accumbens (NAc) in both naive and nicotine self-administering rats, suggesting a DA-dependent mechanism underlying mitigation of nicotine's effects. In conclusion, the present findings support further investigation of R-MOD for treatment of nicotine dependence in humans.

Neuropsychopharmacology (2015) **40**, 1762–1771; doi:10.1038/npp.2015.24; published online 11 February 2015

INTRODUCTION

Tobacco smoking remains a serious global health threat and the single most preventable cause of hospitalizations and deaths in developed countries (Carson *et al*, 2013). Although many smokers attempt to quit, a majority relapse repeatedly. The United States Food and Drug Administration (FDA) has approved the use of nicotine replacement, bupropion, and varenicline for smoking cessation (Carson *et al*, 2013). Yet, only 20–30% of smokers treated with these medications achieve long-term abstinence (Cahill *et al*, 2007; Carson *et al*, 2013; Hughes *et al*, 2007). Therefore, the challenge of developing more effective and/or alternative

medications to achieve smoking cessation and prevent relapse remains critical. One strategy is the preclinical study of medications already clinically available that act on neurobiological mechanisms related to nicotine reward and addiction.

(±)-Modafinil ((±)-MOD) and its R-enantiomer (R-MOD) are FDA approved for the treatment of narcolepsy and other sleep disorders (Garnock-Jones *et al*, 2009). (±)-MOD is also used, off label, as a cognitive enhancer and for treatment of attention-deficit/hyperactivity disorder (Schmaal *et al*, 2013). Recently, (±)-MOD has been clinically investigated for the treatment of cocaine and methamphetamine addiction. As (±)-MOD is a mild psychostimulant and a dopamine transporter (DAT) inhibitor, but with no reported abuse liability (Jasinski and Kovacevic-Ristanovic, 2000; Myrick *et al*, 2004), it has been suggested as useful for the treatment of psychostimulant dependence. Camacho and Stein (2002) first reported that (±)-MOD was effective in treatment of amphetamine dependence in a patient with comorbid social phobia. This finding was later supported by a number of studies demonstrating that (±)-MOD was effective in reducing cocaine and methamphetamine use, subjective euphoric effects, craving, and withdrawal symptoms (Dackis *et al*, 2005; Dackis *et al*, 2003; Goudriaan *et al*, 2013; Hart *et al*, 2008; Malcolm *et al*, 2006; McGregor *et al*,

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³The first two authors contributed equally to this work. Received 9 September 2014; revised 1 January 2015; accepted 12 January 2015; accepted article preview online 23 January 2015

2008; Shearer *et al*, 2009). In experimental animals, (\pm)-MOD has been reported to block reinstatement of drug-seeking caused by cocaine, heroin, methamphetamine, or drug-associated environmental cues (Mahler *et al*, 2014; Reichel and See, 2010; Tahsili-Fahadan *et al*, 2010).

In contrast, several recent clinical trials failed to confirm the therapeutic efficacy of (\pm)-MOD in treatment of cocaine (Dackis *et al*, 2012) or methamphetamine dependence (Anderson *et al*, 2012; De La Garza *et al*, 2010; Lee *et al*, 2013). Similarly, conflicting results are reported for (\pm)-MOD as an adjunctive pharmacotherapeutic aid for smoking cessation. Although (\pm)-MOD was reported to attenuate self-reported nicotine 'drug strength,' it had no effect on tobacco withdrawal symptoms (Sofuoglu *et al*, 2008). In two other studies, (\pm)-MOD failed to increase tobacco abstinence duration or reduce craving and withdrawal (Perkins *et al*, 2013; Schnoll *et al*, 2008).

Given that (\pm)-MOD is a mixture of enantiomers and that R-MOD has higher affinity and potency at blocking the DAT (Cao *et al*, 2010; Loland *et al*, 2012) and an improved pharmacokinetic (PK) profile (Robertson and Hellriegel, 2003; Wong *et al*, 1999), we hypothesized that R-MOD may be more potent and effective than S-MOD or (\pm)-MOD in attenuating drug-taking and drug-seeking behavior. We tested this hypothesis in this study. We first investigated the effects of R- and S-MOD on nicotine self-administration in Long-Evans rats. As Long-Evans rats failed to display robust reinstatement to nicotine-seeking behavior, we used alcohol-preferring rats (P-rats), which display higher vulnerability to nicotine (Le *et al*, 2006), to study the effects of R-MOD on nicotine self-administration and relapse to nicotine-seeking behavior. We then evaluated the abuse liability of R-MOD itself in both self-administration and reinstatement tests. Finally, we used *in vivo* microdialysis to study the effects of R-MOD on nicotine-enhanced extracellular DA to explore the mechanisms underlying the above behavioral effects.

MATERIALS AND METHODS

Subjects

Male Long-Evans rats (Charles River Laboratories, Raleigh, NC) and P-rats (Indiana University Medical Center, Indianapolis, IN) were used. All animals were housed individually in a climate-controlled room under a 12 h light/dark cycle. Food and water were available *ad libitum* throughout the experiments. All experimental procedures were conducted in accordance with the Guide for the Care and Use of Laboratory Animals and were approved by the Animal Care and Use Committee of the National Institute on Drug Abuse of the US National Institutes of Health.

Drugs

R-MOD and S-MOD were prepared in the Medicinal Chemistry Section, National Institute on Drug Abuse-Intramural Research Program, from intermediates provided by Dr Thomas Prisinzano (University of Kansas) according to literature procedures (Cao *et al*, 2010). R-MOD and S-MOD were dissolved in sterile water containing 10% DMSO and 15% Tween-80 for intraperitoneal (i.p.) injection.

In the intravenous (i.v.) R-MOD replacement experiment, R-MOD was dissolved in saline.

Experiment 1: Nicotine Self-Administration

Initial oral sucrose self-administration. The procedures for oral sucrose self-administration were as previously reported (Xi *et al*, 2006b). Rats were deprived of food and water 2 h before self-administration and then allowed access to 5% oral sucrose under fixed ratio-1 (FR-1) reinforcement conditions for 5–7 days (3 h/day). After stable self-administration was achieved, one group of animals was used to study the effects of R-MOD on sucrose self-administration, whereas other groups of rats were used for i.v. nicotine self-administration below.

Intravenous nicotine self-administration. The procedures for jugular catheter surgery and nicotine self-administration were as previously reported (Le *et al*, 2006; Xi *et al*, 2006b). Intravenous self-administration experiments were conducted in operant response test chambers from MED Associates (Georgia, VT). Each test chamber had an active lever and an inactive lever. Depression of the active lever activated the infusion pump; depression of the inactive lever was counted but had no consequence. After 7 days of recovery from surgery, rats were initially trained to self-administer nicotine (30 μ g/kg/infusion) under FR-1 reinforcement. Each nicotine infusion delivered a volume of 0.08 ml/infusion over 5 s and was paired with presentation of a stimulus light and tone. After stable nicotine self-administration was achieved (≥ 10 nicotine infusions per 3 h session; $< 20\%$ variability in daily nicotine infusions across two consecutive sessions; an active/inactive lever press ratio exceeding 2:1), some animals were switched to other unit doses of nicotine (7.5, 15, 30, or 60 μ g/kg/infusion) under different reinforcement schedules (FR-1, FR-2, or FR-5) to more fully characterize nicotine self-administration behavior. To confirm whether the operant lever response was reinforced by nicotine, an active/inactive lever switching test was conducted in a subset of rats, in which the previous nicotine-paired active lever became inactive, whereas the previous inactive lever became active. The effects of R- or S-MOD on nicotine self-administration were evaluated in Long-Evans rats and P-rats. After each test, animals then received an additional 3–5 days of self-administration of nicotine alone until stable self-administration was reestablished. The order of testing for the various doses of R-MOD or S-MOD was counterbalanced.

R-MOD substitution test. After stable nicotine self-administration was established, an additional group of P-rats was divided into two subgroups ($n = 6$ each): (1) nicotine was replaced by R-MOD (0.5 mg/kg/infusion); and (2) nicotine was replaced by saline (0.08 ml/infusion). As animals might take several days to support self-administration for a novel reinforcer, each replacement test was repeated for 3 days. The doses of R-MOD were chosen on two grounds. First, the maximal solubility of R-MOD in saline (vehicle) is ~ 2 mg/ml, making 0.5 mg/kg/infusion the maximum feasible unit dose. Second, the cumulative i.v. R-MOD dose within the initial 30 min was ~ 2.5 –5 mg/kg

(5–10 infusions \times 0.5 mg/kg/infusion, i.v.), and this is close to or higher than the i.p. dose of 10 mg/kg of R-MOD that was found to be effective in attenuation of nicotine self-administration.

Experiment 2: Relapse to Nicotine-Seeking Behavior in P-Rats

To maximize active lever response during self-administration and reinstatement testing, animals were allowed to self-administer nicotine (30 or 60 μ g/kg/infusion) under progressively increased reinforcement schedules (FR-1–FR-5). After stable nicotine self-administration was established, animals were divided into three groups to evaluate the effects of R-MOD in the following experiments.

Nicotine-induced reinstatement. The extinction procedure was the same as described previously (Xi et al, 2006b). During extinction, nicotine was replaced by saline, and the nicotine-associated cue light and tone were turned off. Active lever pressing led only to saline infusion. After the animals met the extinction criteria (≤ 10 lever presses for 3 consecutive days), the rats were divided into three subgroups to study the effects of R-MOD on reinstatement of drug-seeking behavior induced by nicotine priming (0.15 mg/kg, s.c.).

Cue-induced nicotine-seeking behavior. Following establishment of stable nicotine self-administration as described above, the second group of rats underwent 3 weeks of withdrawal in the housing facility. On test day, the rats were placed into the same self-administration chambers to observe cue-induced drug-seeking behavior (i.g., lever pressing) under extinction conditions during which lever pressing resulted in presentation of nicotine-associated cue light and cue tone, but not nicotine. Each rat was tested three times under different R-MOD doses in a counterbalanced manner. The interval between drug tests was 2–3 days.

R-MOD-induced reinstatement. After stable nicotine self-administration was achieved, the third group of rats underwent extinction, as described above, until drug-seeking behavior was extinguished. Then, animals were divided into three subgroups and tested for reinstatement of drug-seeking behavior triggered by R-MOD (0, 30, and 100 mg/kg, i.p.) in P-rats.

Experiment 3: Locomotor Activity in P-Rats

This experiment was designed to evaluate the psychomotor stimulating effects of R-MOD. Drug-naive rats were placed in locomotor detection chambers (Accuscan, Columbus, OH) and habituated for 1 h. Each rat randomly received vehicle or one dose of R-MOD (0, 30, and 100 mg/kg, i.p.). Following the injection, locomotor activity was recorded for 2 h in 10 min intervals. Each animal was tested 3 times under different R-MOD doses in a counterbalanced manner. The time interval was 2–3 days between tests. The distance counts (cm) were used to evaluate the effects of R-MOD on locomotion.

Experiment 4: *In Vivo* Microdialysis in P-Rats

Intracranial guide cannula surgery and *in vivo* microdialysis procedures were as reported previously (Xi et al, 2006b). Two guide cannulae were surgically implanted into the NAc (AP + 1.6, ML \pm 2.0, DV $-$ 4.0 mm, 6° from vertical) to collect extracellular fluid samples. After 7 days of recovery from surgery, microdialysis probes were inserted into the NAc 12 h before the onset of microdialysis. After 2 h of baseline sample collection, each animal received one dose of R-MOD (30 and 100 mg/kg, i.p.) and/or nicotine (0.4 mg/kg, s.c.). R-MOD was given 40 min before nicotine to observe the effects of R-MOD on nicotine-enhanced extracellular NAc DA. After collection, samples were frozen at -80°C . Dialysate DA was measured with the ESA electrochemical detection system (ESA, Chelmsford, MA) (Xi et al, 2006b).

After completion of the microdialysis experiments, rats were deeply anesthetized with a high dose of pentobarbital and perfused transcardially with 0.9% saline followed by 10% formalin. Brains were removed and placed in 10% formalin for histological verification of microdialysis probe locations in rat brain.

Data Analysis

All data are presented as means \pm SEM. One-way or two-way analysis of variance (ANOVA) for repeated measures over time or drug dose was used to analyze the effects of R-MOD or S-MOD in various experiments. Whenever a significant main effect was found, individual group comparisons were carried out using the Student–Newman–Keuls method.

RESULTS

R-MOD Inhibits Nicotine Self-Administration in Long-Evans Rats

Supplementary Figure S1A shows the mean numbers of active and inactive lever responses during daily 3 h sessions of oral sucrose and i.v. nicotine self-administration in 22 rats displaying stable nicotine self-administration. Thirty four rats were not included as they were either sick, catheter clogged, or did not meet stable nicotine self-administration criteria. Supplementary Figure S1B shows representative nicotine self-administration records from six different animals, illustrating regular patterns of nicotine self-administration during daily 3 h sessions. Supplementary Figure S1C shows that R-MOD significantly inhibited nicotine (30 μ g/kg/infusion) self-administration in Long-Evans rats in a dose-dependent manner. One-way ANOVA for repeated measures over drug dose demonstrated a statistically significant R-MOD treatment main effect ($F_{4,28} = 5.46$, $p < 0.01$). Supplementary Figure S1D shows representative nicotine self-administration records before and after 30 mg/kg R-MOD administration, illustrating a typical extinction pattern—initial high-rate lever pressing followed by cessation of lever pressing, suggesting a reduction in nicotine rewarding effects. Supplementary Figure S1E shows that 30 mg/kg R-MOD significantly shifted the nicotine dose–response self-administration curve downward. Two-way ANOVA for repeated measures over nicotine dose revealed a significant R-MOD treatment main

effect ($F_{1,7} = 30.99$, $p < 0.001$). Supplementary Figure S1F shows that S-MOD, at the same doses (10–30 mg/kg), had no effect on nicotine self-administration. However, when the dose was increased to 100 mg/kg, S-MOD significantly inhibited nicotine self-administration ($F_{4,24} = 5.33$, $p < 0.05$).

Long-Evans Rats Display Low Reinstatement Response to Nicotine

We then studied whether R-MOD inhibited nicotine-induced reinstatement of drug-seeking behavior. Supplementary Figure S2A shows the active and inactive lever presses during nicotine (30 μ g/kg/infusion) self-administration, extinction, and reinstatement testing, illustrating that nicotine priming (0.15 mg/kg, s.c.) did not reinstate nicotine-seeking behavior in Long-Evans rats. We then increased the nicotine dose from 30 to 60 μ g/kg/infusion during self-administration in an additional group of rats (Supplementary Figure S2B). We again found that nicotine priming failed to reinstate a robust lever response to nicotine in Long-Evans rats.

Alcohol-Preferring Rats Display High Reinstatement Response to Nicotine

Therefore, we turned to the use of P-rats in this study based on evidence that they display increased vulnerability to nicotine self-administration and relapse to drug seeking (Le *et al*, 2006). Supplementary Figure S2C shows that P-rats displayed significantly higher active lever responding during self-administration in a reinforcement schedule-dependent manner and significantly higher reinstatement responding to nicotine compared to Long-Evans rats, under the same experimental conditions (Supplementary Figure S2B).

R-MOD Inhibits Nicotine Self-Administration in P-Rats

Nicotine-reinforced self-administration in P-rats was then characterized. Figure 1a shows a typical dose-response-relationship for self-administration a compensatory decrease in lever presses with increased nicotine dose. To determine whether such lever responses were truly reinforced by nicotine, and not by carryover from initial sucrose self-administration, we performed an active and inactive lever switch test in a group of rats. Figure 1b shows that P-rats rapidly learnt to press the new active lever for nicotine self-administration while also displaying increased lever responding on the new inactive lever (previous active lever).

Figure 1c shows that R-MOD pretreatment significantly and dose-dependently inhibited nicotine (60 μ g/kg/infusion) self-administration in P-rats (infusions, $F_{2,14} = 15.19$, $p < 0.001$; active lever responses, $F_{2,14} = 10.33$, $p < 0.001$). This reduction in nicotine self-administration is not due to nonspecific locomotor impairment, as the same doses of R-MOD neither altered sucrose self-administration (Figure 1d) nor inhibited locomotor behavior (Supplementary Figure S3). In fact, R-MOD (at 100 mg/kg), produced a modest increase in locomotor behavior.

R-MOD Itself Failed to Sustain Self-Administration in P-Rats

Figure 1e shows that R-MOD substitution for nicotine did not sustain stable self-administration in rats already displaying stable nicotine self-administration. Although there is no significant change observed on the first day of substitution, continuation of substitution for 2 more days caused a significant reduction in the total number of active lever responses. Two-way ANOVA for repeated measures over time revealed a significant time main effect ($F_{10,90} = 7.88$, $p < 0.001$), but did not reveal a significant treatment (saline vs R-MOD) main effect ($F_{1,9} = 0.82$, $p > 0.05$). After cessation of saline or R-MOD replacement testing, nicotine self-administration gradually recovered to the basal levels before the substitution test. Figure 1f shows representative patterns of self-administration during R-MOD substitution testing, illustrating that the self-administration behavior underwent gradual extinction over the 3 h test period. This pattern of extinction was essentially identical to that seen when saline was substituted for nicotine.

R-MOD Inhibits Nicotine- or Cue-Induced Nicotine-Seeking Behavior in P-Rats

Figure 2a shows active lever responses during the last session of nicotine self-administration, last session of extinction, and reinstatement testing, illustrating that pretreatment with R-MOD significantly attenuated nicotine-induced reinstatement of drug-seeking behavior ($F_{2,14} = 18.09$, $p < 0.001$). R-MOD also significantly inhibited inactive lever responding (Figure 2b).

We also used an animal model of incubation craving (Marchant *et al*, 2013), in which rats were tested in a single extinction session after 3 weeks of withdrawal to study the effects of R-MOD on nicotine cue-induced drug-seeking behavior. We found that R-MOD pretreatment significantly attenuated nicotine cue-induced drug seeking in P-rats (Figure 2c: $F_{2,16} = 15.21$, $p < 0.001$). R-MOD had no effect on inactive lever responses in this test (Figure 2d). Figure 2e and 2f shows that R-MOD alone failed to induce reinstatement of nicotine-seeking behavior under the same experimental conditions in additional groups of rats.

R-MOD Blocked Nicotine-Enhanced NAc DA Release

Figure 3a shows that systemic injections of R-MOD produced a slow-onset, dose-dependent increase in extracellular NAc DA in naive rats. This effect lasted for 2–3 h with a peak effect at ~ 1 h after R-MOD. Two-way ANOVA for repeated measures over time revealed a statistically significant R-MOD treatment main effect ($F_{2,18} = 6.01$, $p = 0.01$) and a significant time main effect ($F_{9,18} = 4.17$, $p < 0.01$). *Post hoc* individual group comparisons revealed a statistically significant increase in extracellular DA after 100 mg/kg, but not 30 mg/kg, R-MOD (Figure 3a).

Figure 3b shows the time courses of the changes in extracellular DA before and after each drug administration in drug-naive rats, illustrating that both R-MOD and nicotine elevated extracellular DA levels immediately after injection. We then used two methods to quantitatively

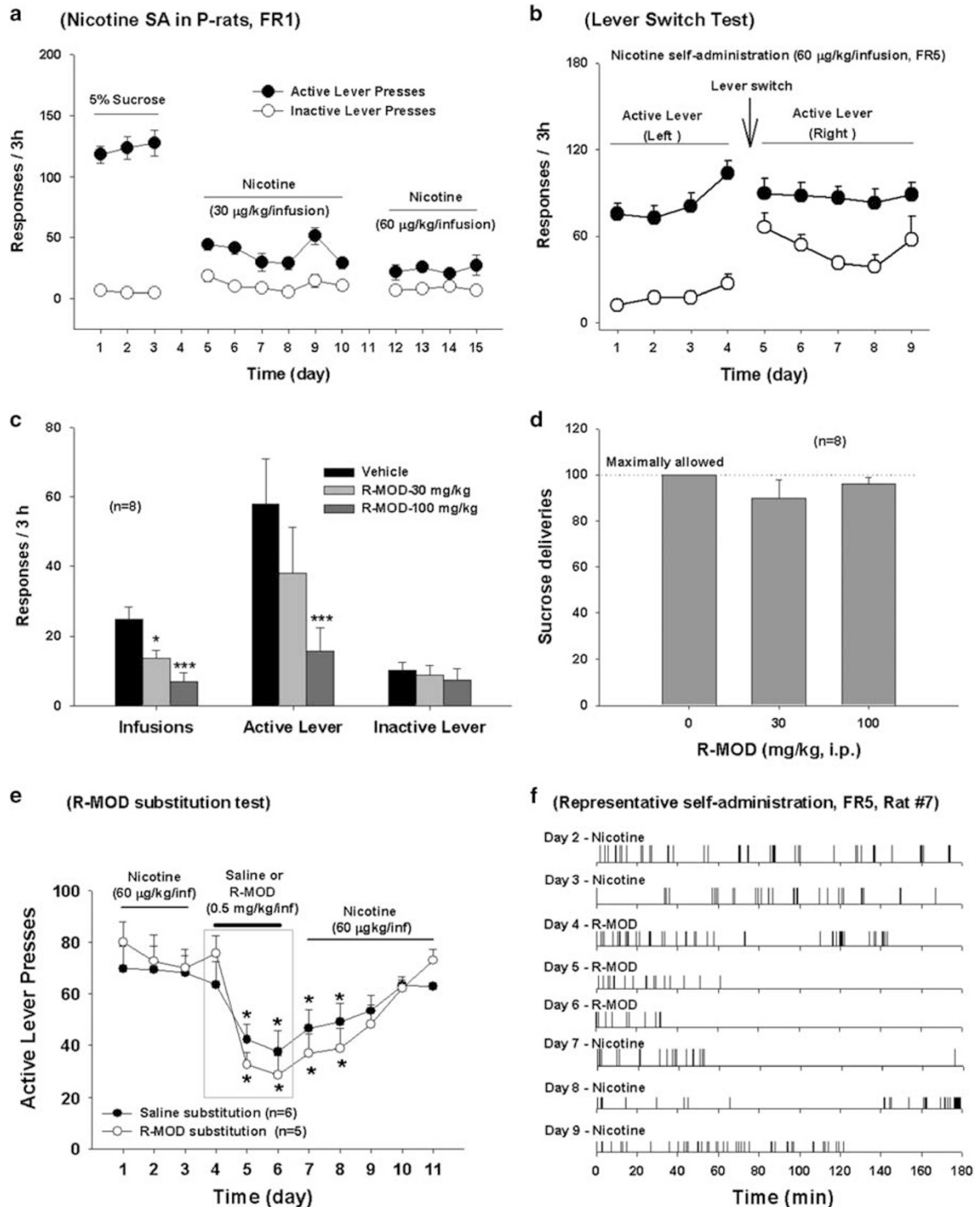


Figure 1 Effects of R-MOD on nicotine self-administration in P-rats. (a) Stable nicotine self-administration maintained by two different doses (30 or 60 µg/kg/infusion, FR-1) of nicotine in rats pretrained on oral sucrose self-administration; (b) active and inactive lever switch results, illustrating that the lever responses were reinforced by nicotine (60 µg/kg/infusion, FR-5), not by previous oral sucrose self-administration; (c) R-MOD (30 and 100 mg/kg) dose-dependently inhibited nicotine self-administration; (d) R-MOD, at the same doses that inhibited nicotine self-administration, had no effect on oral sucrose self-administration. (e) Active lever presses during nicotine self-administration and 3 days of R-MOD replacement for nicotine, illustrating that R-MOD cannot sustain self-administration in P-rats previously self-administering nicotine. (f) Representative records of nicotine or R-MOD self-administration, illustrating a typical pattern of extinction of drug seeking, ie, initial burst-like lever response followed by cessation of lever responding, during 3 h test period after R-MOD substitution for nicotine. * $p < 0.05$; *** $p < 0.001$, compared with predrug baseline in each dose group.

evaluate the effects of R-MOD pretreatment on nicotine-induced DA release. First, we compared the absolute amounts of DA (fmol/sample) produced by nicotine in the

presence and absence of R-MOD pretreatment (Figure 3c), and found that R-MOD pretreatment significantly attenuated nicotine-enhanced DA in a dose-dependent manner.

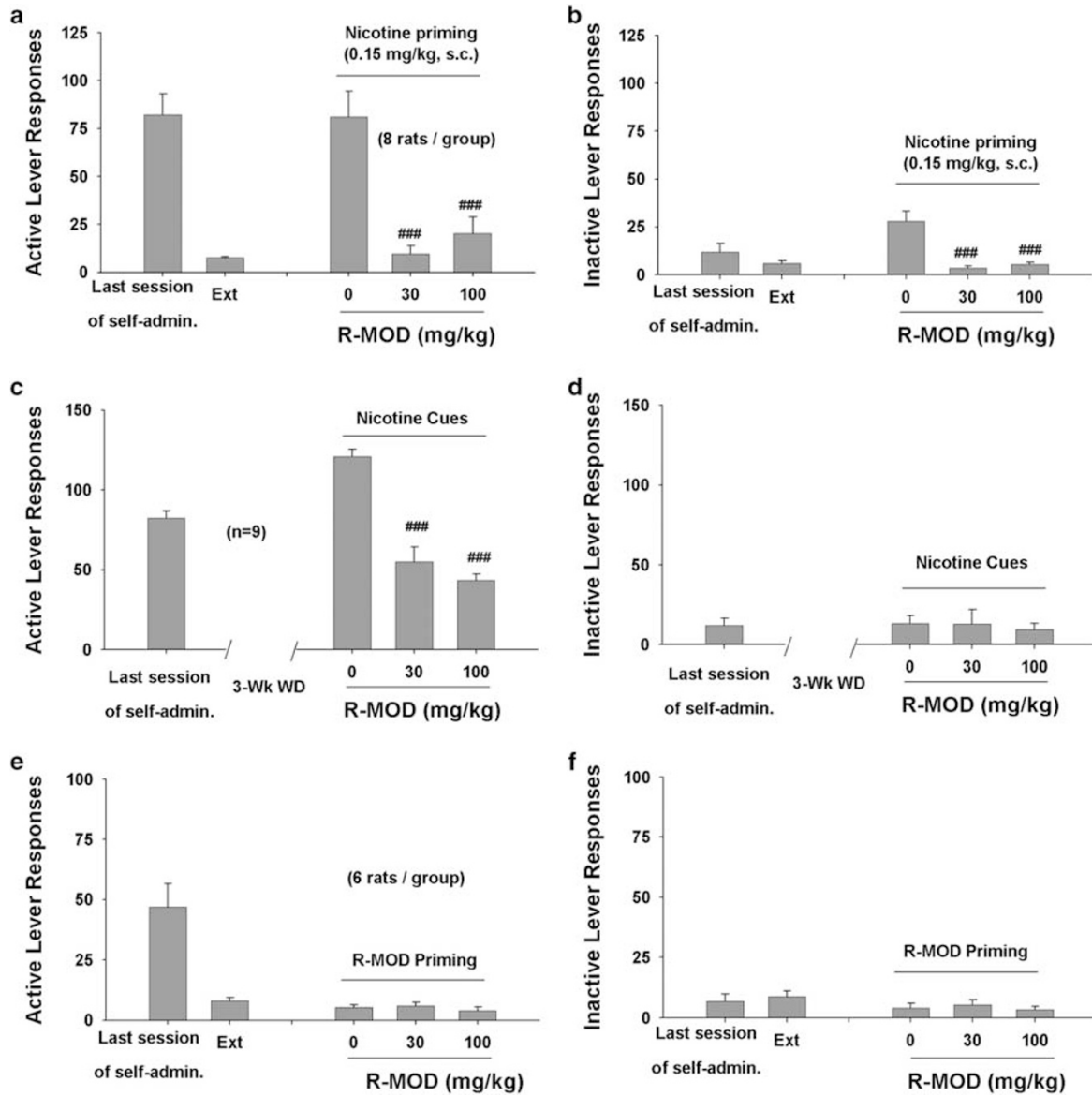


Figure 2 Effects of R-MOD on nicotine- or nicotine-associated cue-induced reinstatement of nicotine seeking in P-rats. (a, b) Active and inactive lever responses during the last session of nicotine self-administration, last session of extinction, and nicotine-triggered reinstatement testing, illustrating that R-MOD significantly inhibited nicotine priming-induced reinstatement. (c, d) Active and inactive lever responses during the last session of nicotine self-administration and cue-triggered reinstatement testing, illustrating that R-MOD significantly inhibited nicotine-associated cue-induced nicotine-seeking behavior in rats after 3 weeks of withdrawal from the last nicotine self-administration. (e, f) Active and inactive lever responses during the last session of nicotine self-administration, extinction, and reinstatement testing, illustrating that R-MOD (30 or 100 mg/kg) failed to reinstate nicotine-seeking behavior. ### $p < 0.001$, compared with vehicle control.

Two-way ANOVA for repeated measures over time did not reveal a statistically significant R-MOD main effect ($F_{2,24} = 1.72, p > 0.05$), but revealed a statistically significant time main effect ($F_{10,240} = 4.90, p < 0.001$) and treatment \times time interaction ($F_{10,240} = 3.98, p < 0.05$). Individual group comparisons revealed a significant reduction in peak DA at 60 min compared with vehicle control group (Figure 3c). Second, we compared nicotine-induced % changes in DA over baseline in the presence or absence of R-MOD pretreatment (Figure 3d). We found that R-MOD pretreatment significantly blunted nicotine-induced increases in extracellular DA (Figure 3d). Two-way ANOVA for repeated measures over time did not reveal a statistically significant

R-MOD main effect ($F_{2,24} = 1.12, p > 0.05$), but revealed a statistically significant time main effect ($F_{10,240} = 5.89, p < 0.001$) and treatment \times time interaction ($F_{10,240} = 3.23, p < 0.05$). Individual group comparisons revealed a significant reduction in DA at several time points compared with vehicle control group (Figure 3d).

Finally, we investigated the effects of R-MOD pretreatment on nicotine-induced DA enhancement in nicotine self-administration rats after 2 weeks of withdrawal (Figure 4a). Figure 4b shows extracellular DA levels before and after R-MOD and/or nicotine administration. As R-MOD pretreatment elevated extracellular DA before the nicotine injection, we used the same methods, as described above, to

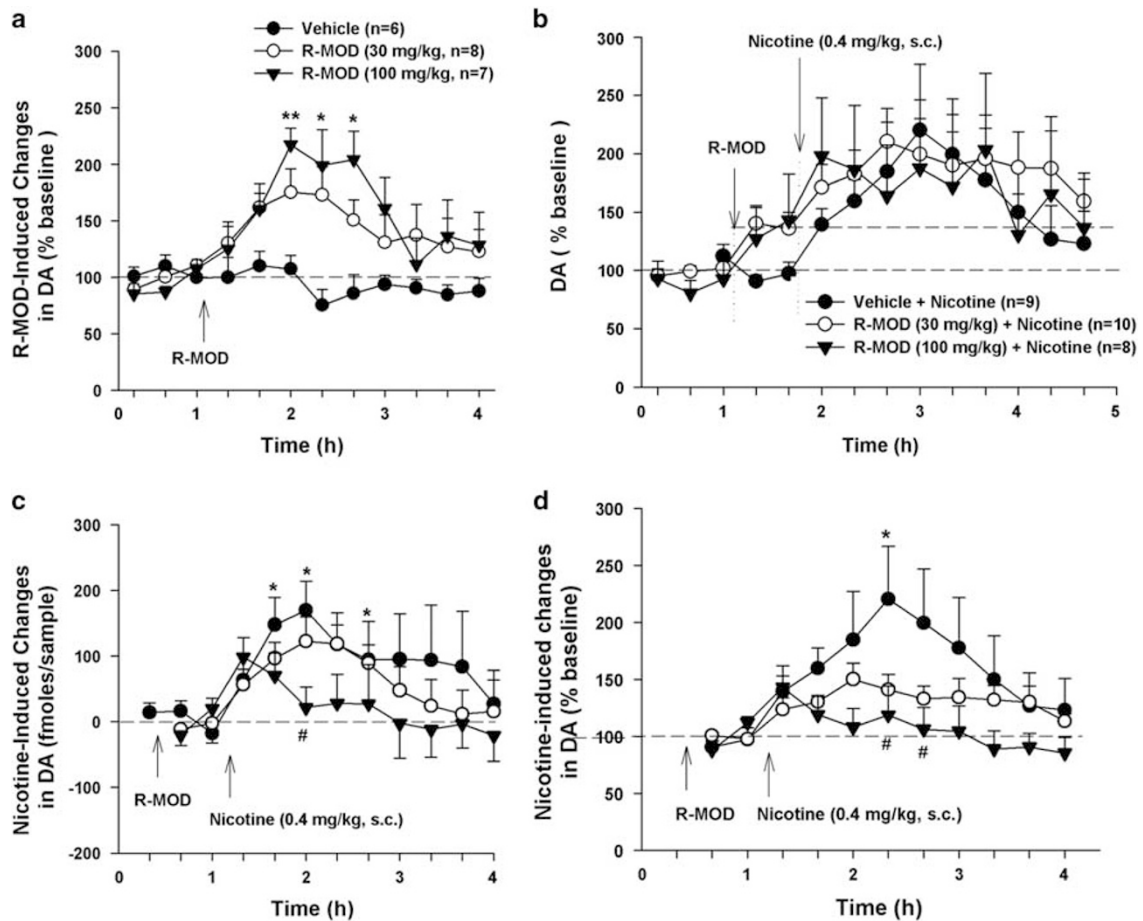


Figure 3 Effects of R-MOD on basal and nicotine-enhanced DA in the nucleus accumbens in naive rats without nicotine self-administration history. (a) R-MOD alone produced a slow-onset, dose-dependent increase in extracellular DA; (b) extracellular DA levels before and after R-MOD and/or nicotine administration, illustrating that each drug elevated extracellular DA levels. (c) Nicotine-induced changes in extracellular NAc DA (fmol/sample) in the presence or absence of R-MOD pretreatment, by subtracting basal level (mean value of two samples before nicotine) of extracellular DA from DA level after nicotine administration; (d) nicotine-induced % changes in DA over baseline (mean value of two samples before nicotine injection) in the presence or absence of R-MOD pretreatment, illustrating a significant reduction in nicotine-enhanced extracellular DA in a dose-dependent manner after R-MOD administration. * $p < 0.05$; ** $p < 0.01$, compared with baseline before R-MOD (a) or nicotine (b, c) administration; # $p < 0.05$, compared with vehicle control group.

quantitatively evaluate nicotine-induced DA release in the absence or presence of R-MOD pretreatment. We found that R-MOD pretreatment significantly and dose-dependently blocked nicotine-enhanced extracellular DA, as assessed by absolute amounts of DA release (Figure 4c, $F_{2,16} = 7.11$, $p < 0.01$, two-way repeated measures ANOVA over time) or percentage changes over the new baseline immediately before nicotine administration (Figure 4d, $F_{2,16} = 9.27$, $p < 0.01$). Histological examination indicated that active microdialysis membranes spanned the length of the core and shell compartments of the NAc (Supplementary Figure S4).

DISCUSSION

The major findings in this study include: (1) R-MOD is more potent and effective than S-MOD at attenuating nicotine self-administration in Long-Evans rats; (2) P-rats display higher lever responses for nicotine self-administration and higher reinstatement responses to nicotine priming than Long-

Evans rats; (3) R-MOD effectively attenuates nicotine self-administration, nicotine-induced reinstatement, and cue-induced nicotine-seeking in P-rats; (4) R-MOD substitution for nicotine fails to sustain self-administration and R-MOD alone also fails to reinstate nicotine-seeking behavior; and (5) pretreatment with R-MOD attenuates nicotine-enhanced DA release in drug-naive rats and nicotine self-administering rats. Together, these findings demonstrate that R-MOD inhibits nicotine-taking and nicotine-seeking behavior in experimental animals, an effect likely mediated by a DA-dependent mechanism.

R-MOD and S-MOD display different pharmacokinetic and pharmacodynamic profiles and one might affect the other in the racemic mixture. Specifically, R-MOD has a ~3-fold higher affinity for the DAT than S-MOD (Cao *et al*, 2010; Loland *et al*, 2012) and is more metabolically stable and longer acting (Dinges *et al*, 2006; Garnock-Jones *et al*, 2009). In addition, R-MOD appears to bind to the DAT more like atypical DA uptake inhibitors, exemplified by JHW 007, than like cocaine (Loland *et al*, 2012; Okunola-Bakare *et al*, 2014). In this study, we found that R-MOD is

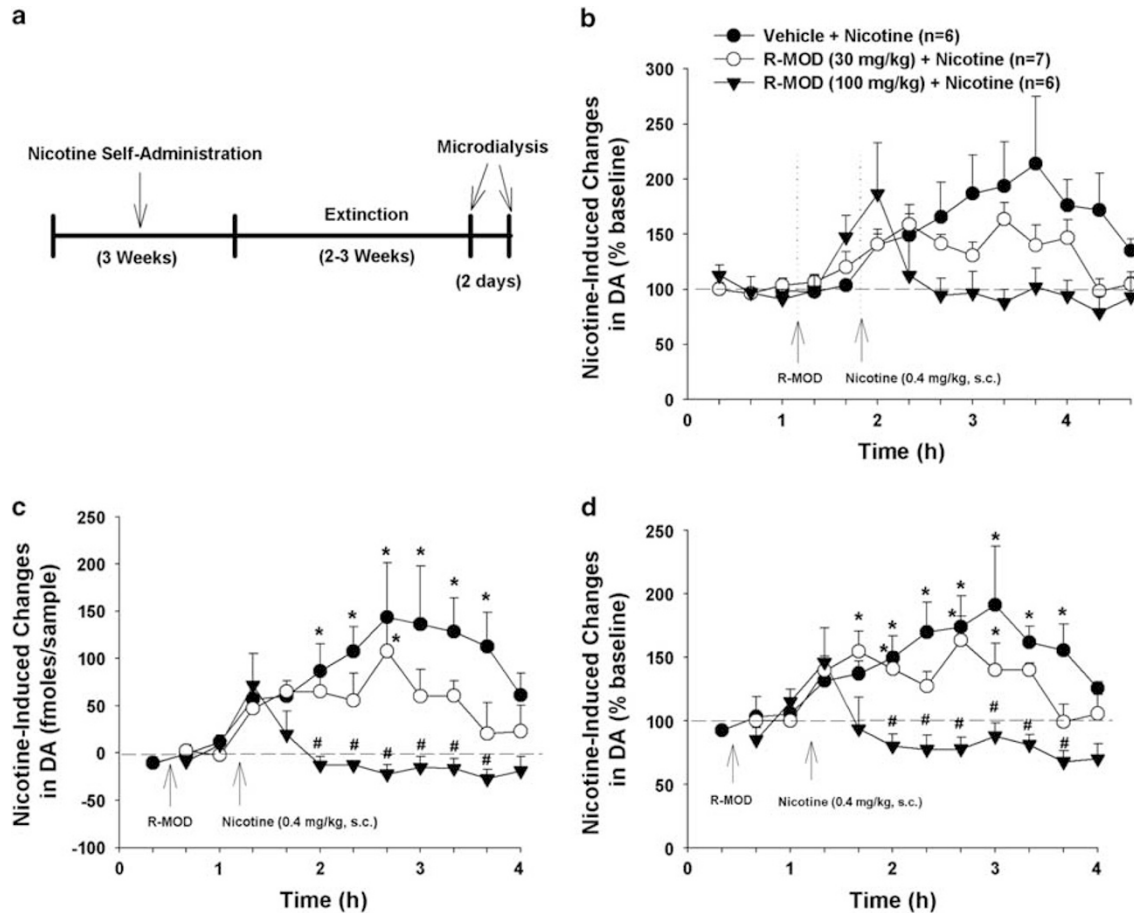


Figure 4 Effects of R-MOD on basal and nicotine-enhanced DA in the nucleus accumbens in rats with nicotine self-administration history. (a) The general experimental procedures; (b) extracellular DA levels before and after R-MOD and/or nicotine administration, illustrating that both drugs elevated extracellular DA levels. (c) Nicotine-induced changes in extracellular NAc DA (fmol/sample) in the presence or absence of R-MOD pretreatment, by subtracting basal level (mean value of two samples before nicotine) of extracellular DA from DA level after nicotine administration; (d) nicotine-induced % changes in DA over baseline (mean value of two samples before nicotine injection) in the presence or absence of R-MOD pretreatment, illustrating a significant reduction in nicotine-enhanced extracellular DA in a dose-dependent manner after R-MOD administration. * $p < 0.05$; compared with baseline before R-MOD (a) or nicotine (b, c) administration; # $p < 0.05$, compared with vehicle control group.

more effective than S-MOD in inhibiting nicotine self-administration in Long-Evans rats. The minimum effective doses (10–30 mg/kg) of R-MOD in attenuating nicotine self-administration and relapse to nicotine seeking are also significantly lower than that of (\pm)-MOD (300 mg/kg) in attenuating cocaine-, methamphetamine-, or cue-induced reinstatement of drug-seeking behavior (Deroche-Gamonet *et al*, 2002; Holtz *et al*, 2012; Mahler *et al*, 2014; Reichel and See, 2010). We note that the effective doses of R-MOD in inhibiting nicotine self-administration are higher in P-rats (30–100 mg/kg) than in Long-Evans rats (10–30 mg/kg). This may be related to the higher nicotine dose used in the P-rats (60 μ g/kg/infusion) than that used in the Long-Evans rats (30 μ g/kg/infusion). We also note that (\pm)-MOD produces conditioned place preference (Nguyen *et al*, 2011; Shuman *et al*, 2012). However, in this study, R-MOD substitution failed to sustain self-administration in rats previously self-administering nicotine. R-MOD itself also failed to reinstate drug-seeking behavior, suggesting that R-MOD has significantly lower abuse potential than nicotine. This may be related to the lower (10- to 20-fold)

affinity of R-MOD for the DAT than cocaine, and different pharmacokinetic and pharmacodynamic profiles (Gold and Balster, 1996; Loland *et al*, 2012; Newman *et al*, 2010; Paterson *et al*, 2010). Taken together, the present findings suggest that R-MOD may be more useful than (\pm)-MOD in treatment of substance abuse and addiction, and should be investigated for smoking cessation.

Although the present findings are observed mainly in selectively bred P-rats, they may generalize to other populations of rats or humans as R-MOD is also effective in attenuating nicotine self-administration in Long-Evans rats, and only a small population of human subjects develop addiction to nicotine, similar to P-rats that display higher vulnerability to nicotine (Le *et al*, 2006). We acknowledge that only a 3h short access to nicotine was used in the present nicotine self-administration, and this may not reflect certain key aspects of the nicotine addiction state in a manner seen after more prolonged daily nicotine consumption.

The mechanisms by which R-MOD inhibits nicotine-taking and nicotine-seeking behaviors are unclear. It is

unlikely due to nonspecific sedation or locomotor impairment as R-MOD neither alters oral sucrose self-administration nor inhibits locomotor behavior. In contrast, R-MOD, at the high dose of 100 mg/kg, moderately increased open-field locomotor activity. This is consistent with previous findings that (\pm)-MOD facilitates expression of locomotor sensitization and increased locomotor activity in rodents (Edgar and Seidel, 1997; Paterson et al, 2010). Although (\pm)-MOD may modulate multiple neurotransmitter systems, the DAT is the only target that has been directly implicated in its therapeutic efficacy (Loland et al, 2012; Mereu et al, 2013 for review). An important finding in this study is that R-MOD pretreatment significantly blocks nicotine-enhanced DA in the NAc in either drug-naive rats or nicotine self-administering rats, suggesting a DA-dependent mechanism underlying the behavioral effects observed in this study. In addition, systemic administration of R-MOD alone also elevated NAc DA. This effect has been attributed to its DAT inhibition (Loland et al, 2012). As virtually all drug-dependent humans and experimental animals display reduced basal levels of extracellular DA or DA receptor (particularly D2/3) densities after abstinence (Diana, 2011; Volkow et al, 2009), this hypofunctional NAc DA transmission has been postulated to be correlated with increased drug craving and increased susceptibility to relapse (Diana, 2011; Volkow et al, 2009). Accordingly, the R-MOD-induced increase in extracellular DA would be expected to renormalize extracellular DA, and therefore contribute to relief of nicotine craving and relapse to nicotine-seeking behavior. The mechanisms by which R-MOD inhibits nicotine-enhanced DA are unclear. One possibility is that R-MOD-induced increases in basal levels of DA may activate presynaptic DA (D2 and D3) autoreceptors, therefore inhibiting subsequent nicotine-enhanced DA. Thus, reduced DA response to nicotine, combined with normalized extracellular DA produced by R-MOD pretreatment, may explain the antagonism of nicotine self-administration and relapse to nicotine-seeking behavior after R-MOD administration.

In conclusion, this study demonstrates that low doses of R-MOD significantly inhibit nicotine-taking and nicotine-seeking behavior in rats. As R-MOD itself has lower reinforcing effects than nicotine, the present preclinical findings support clinical investigation of R-MOD as a promising medication for treatment of nicotine dependence in humans.

FUNDING AND DISCLOSURE

The authors declare no conflict of interest.

ACKNOWLEDGEMENTS

This research was supported by the Intramural Research Program (IRP) of the National Institute on Drug Abuse (NIDA), National Institutes of Health (NIH). We acknowledge Ms J Cao for synthesizing S-MOD, Dr Tom Prisinzano for supplying intermediates and methods for the syntheses of R- and S-modafinil, and Drs Yavin Shaham and Gianluigi Tanda for their helpful suggestions on experimental design and comments on an earlier draft of this manuscript.

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