Varenicline decreases nicotine self-administration and cue-induced reinstatement of nicotine-seeking behaviour in rats when a long pretreatment time is used

Bernard Le Foll1,2,3, Munmun Chakraborty-Chatterjee1, Shaul Lev-Ran1,3, Chanel Barnes1, Abhiram Pushparaj1, Islam Gamaleddin1, Yijin Yan1, Maram Khaled1 and Steven R. Goldberg1

1 Translational Addiction Research Laboratory, Centre for Addiction and Mental Health, Toronto, Canada
2 Departments of Family and Community Medicine, Pharmacology, Psychiatry, Institute of Medical Sciences, University of Toronto, Toronto, Canada
3 Addiction Program, Centre for Addiction and Mental Health, Toronto, Canada
4 Preclinical Pharmacology Section, Behavioral Neuroscience Research Branch, Intramural Research Program, National Institute on Drug Abuse, National Institutes of Health, Department of Health and Human Services, Baltimore, MD, USA

Abstract

Effects of varenicline (Champix), a nicotinic partial agonist, were evaluated on subjective effects of nicotine (drug discrimination), motivation for nicotine taking (progressive-ratio schedule of intravenous nicotine self-administration) and reinstatement (cue-induced reinstatement of previously extinguished nicotine-seeking behaviour). Effects on motor performance were assessed in rats trained to discriminate nicotine (0.4 mg/kg) from saline under a fixed-ratio (FR 10) schedule of food delivery and in rats trained to respond for food under a progressive-ratio schedule. At short pretreatment times (5–40 min), varenicline produced full or high levels of partial generalization to nicotine’s discriminative-stimulus effects and disrupted responding for food, while there were low levels of partial generalization and no disruption of responding for food at 2- or 4-h pretreatment times. Varenicline (1 and 3 mg/kg, 2-h pretreatment time) enhanced discrimination of low doses of nicotine and to a small extent decreased discrimination of the training dose of nicotine. It also dose-dependently decreased nicotine-taking behaviour, but had no effect on food-taking behaviour under progressive-ratio schedules. Finally, varenicline significantly reduced the ability of a nicotine-associated cue to reinstate extinguished nicotine-seeking behaviour. The ability of varenicline to reduce both nicotine-taking and nicotine-seeking behaviour can contribute to its relatively high efficacy in treating human smokers.

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Introduction

Smoking is currently responsible for the death of one in ten adults worldwide (about 5 million deaths each year). Half the people that smoke today – that is about 650 million people – will eventually be killed by tobacco (Tobacco Advisory Group RCP, 2000) and this morbidity and mortality can be mostly avoided if subjects stop smoking (Department of Health and Human Services, 1990). Several pharmacological treatments (nicotine replacement, bupropion, varenicline) have demonstrated their efficacy as treatments for smoking cessation (Le Foll & George, 2007). However, despite treatment a majority of subjects will relapse and there is the need to develop novel and more effective medications. The availability of drugs that have demonstrated their utility in humans allows testing the predictive value of the animal models that
are used (Lerman et al. 2007). There is a strong need to test those drugs in the various animals used in this field to validate the preclinical approaches (Lerman et al. 2007).

Varenicline is a novel medication that is available for the treatment of smokers (Cahill et al. 2007). It is a \(\alpha_2\beta_2\) subtype nicotinic receptor partial agonist that has actions on other nicotinic receptor subunits (Rollema et al. 2007). Varenicline appears more effective than nicotine-replacement therapy and bupropion in helping smokers quit smoking (Cahill et al. 2007). The effects of varenicline have been evaluated in various animal models. First, it has been reported that varenicline administered 15 min prior to the session significantly decreased nicotine self-administration under a 5-response fixed-ratio (FR 5) schedule of reinforcement in rats (Rollema et al. 2007). In rats trained to self-administer nicotine under a progressive-ratio (PR) schedule, varenicline maintained lower break-points than nicotine when it was tested by substitution (Rollema et al. 2007). In rats trained to discriminate nicotine from saline, varenicline produced full generalization in one study when administered 5 min before the session (Rollema et al. 2007) and partial generalization in another study when administered 25 min before the session (LeSage et al. 2009). Varenicline has been reported to be effective in blocking nicotine-induced reinstatement of drug-seeking behaviour in rats when administered 15 min prior to the session (O’Connor et al. 2010). Although varenicline also reduced cue-reactivity in human smokers (Franklin et al. 2011), it was not effective in reducing cue-induced reinstatement of drug-seeking behaviour in rats (O’Connor et al. 2010). However, since this study used a short pretreatment time (15 min), the agonist properties of varenicline may have interfered with its ability to reduce cue-induced reinstatement. In a more recent study, varenicline tended to decrease cue-induced reinstatement of nicotine-seeking behaviour when administered at a 30 min pretreatment time, although effects did not reach statistical significance (Wouda et al. 2011).

To further investigate this issue, we first assessed the influence of pretreatment time on FR responding for food, the ability of varenicline to generalize to a nicotine cue and its effects of nicotine dose-response curve in a drug-discrimination paradigm in rats. We then evaluated the effects of varenicline on nicotine self-administration maintained under a PR schedule and the effects of varenicline on cue-induced reinstatement of nicotine-seeking behaviour. Control experiments were performed in rats trained to respond for food.

### Materials and methods

#### Animals

Male Long–Evans rats (Charles River, Canada) for the food and nicotine self-administration and PR experiments were experimentally naive at the start of the study. They initially weighed 250–275 g. Rats were individually housed in a temperature- and humidity-controlled room on a 12-h reversed light/dark cycle (lights off 07:00 hours). Experiments were conducted during the dark phase. Prior to any experimental manipulation, animals were given a minimum of 7 days to habituate to the colony room, during which they were weighed and handled.

For the nicotine discrimination study, male Sprague–Dawley rats (Charles River, USA) initially weighing 250–275 g were used. They were housed individually in a temperature- and humidity-controlled room on a regular 12-h light/dark cycle (lights on 07:00 hours). Experiments were conducted during the light phase.

For all experiments, water was available ad libitum and a diet restriction was maintained throughout the studies (~20 g/d).

All the experimental procedures described in this report were performed in compliance with the guidelines of the Canadian Council on Animal Care and corresponding NIH guidelines, and were reviewed and approved by the institutional Animal Care Committee. All efforts were made to minimize animals’ suffering, and to reduce the number of animals required. Use of a repeated-measures design contributed to the latter.

Techniques for initial training and surgery were similar to those previously reported (Corrigall & Coen, 1989; Corrigall et al. 2001). Animals were trained to press a lever on a schedule in which each press resulted in the delivery of a 45-mg food pellet (continuous reinforcement, CRF, no associated cues). Once trained, each animal was surgically prepared with a chronic intravenous catheter implanted in the jugular vein; the catheters exited between the scapulae. Surgery was performed under anaesthesia induced by xylazine (10 mg/kg), given intraperitoneally (i.p.) and ketamine hydrochloride (75 mg/kg i.p.). Incision sites were infiltrated with the local anaesthetic bupivacaine (0.125%). Buprenorphine was given for post-operative analgesia (0.01 mg/kg), given subcutaneously (s.c.), and a single dose of penicillin (30000 units, i.m.) was administered before surgical procedures. Animals were allowed to recover for a 1-wk period before drug self-administration sessions were begun.
Drugs

Nicotine hydrogen tartrate (Sigma-Aldrich, USA) was dissolved in saline, the pH was adjusted to 7.0 (±0.2), and the solution was filtered through a 0.22-mm syringe filter (Fisher Scientific, USA) for sterilization purposes. All nicotine doses are reported as free base concentrations. Nicotine was administered intravenously (i.v.) in a volume of 100 μl/kg/injection or s.c. in a volume of 1 ml/kg.

Varenicline (a gift from Pfizer, USA) was diluted in saline, the pH was adjusted to 7.0 (±0.2) and was administered i.p. in a volume of 1 ml/kg body weight.

Acquisition of the nicotine or food self-administration behaviour

Self-administration sessions were conducted in experimental chambers equipped with two levers (Med Associates, USA). Session start was signalled by the illumination of a house-light; extinction of this light indicated the time-out (TO) period. Rapid delivery of the self-administered drug (approximately 1-s delivery time) was achieved with Med Associates Model PHM-104 pumps. Unit doses were 100 μl/kg; volume adjustments were used to accommodate inter-animal or between-session differences in body weight. Responding on one of the levers resulted in drug delivery when schedule requirements were met, while responding on the other lever was recorded but did not produce any change of lights or drug infusion (active levers are counterbalanced). Self-administration sessions occurred mostly 5 d per week.

In this study, rats acquired nicotine self-administration under a FR schedule of reinforcement and the unit dose was 30 μg/kg per infusion of nicotine base. Session duration was 60 min, and the TO period (turning off of the house-light and illumination of a cue light above the active lever) following each infusion was 1 min. During the first 5 d of acquisition, each lever press during the time-in period resulted in the delivery of an infusion (FR 1), then the response requirement was increased to FR 2 for 3 d and then to the final value of FR 5 (i.e. animals were required to make five lever presses for each drug infusion) for 5 d.

For food studies the apparatus, the stimuli associated with food delivery and the schedule of the acquisition were exactly the same as described above. The rats received a food pellet (45 mg precision pellets, BioServ) instead of a nicotine injection.

Testing under the PR schedule

After training under FR 5, the animals were switched to a PR schedule where the response requirement increased with each successive injection or food pellet delivery. The response requirement progression was based on the formula $5e^{(0.25\text{[inj.number + 3]})} - 5$, with the first two values replaced by 5 and 10 (modified from Roberts, 1992). Thus, the response requirements for successive injections were 5, 10, 17, 24, 32, 42, 56, 73, 95, 124, 161, 208, etc. The break-point was defined as the highest ratio completed prior to the first 30-min period without a response on the active lever in both nicotine and food self-administration.

Sessions under the PR schedule lasted a maximum of 4 h. The animals were allowed 10–15 d depending upon time of stabilization of nicotine or food self-administration on the PR schedule before testing with an i.p. injection of varenicline or vehicle. A counterbalanced within-subject design was employed for the testing. The baseline (BL) condition represented a vehicle (saline) injection given 2 h prior to session start, as performed for varenicline injections. Rats in the nicotine self-administration experiments were also tested in a session where saline was substituted for nicotine (a saline-substitution test conducted after all doses of varenicline were tested on the PR schedule). Testing for catheter patency was conducted at multiple time-points throughout the PR schedule portion of the experiment and subjects analysed in the Results section exclude rats with non-patent catheters.

Cue-induced reinstatement of nicotine seeking

After the rats were tested for the effects of varenicline on nicotine self-administration (PR), an additional few nicotine self-administration sessions were conducted without any treatment and the self-administration behaviour was then extinguished. During the extinction phase rats were placed into the self-administration chamber with the house-light illuminated and responses on the active or inactive levers were recorded, but had no consequences. The criterion for extinction was <20 active lever presses per 1-h session over two consecutive days. After stable extinction, these rats were tested for the effects of varenicline (0.3–3 mg/kg) on cue-induced reinstatement of nicotine-seeking behaviour in a counterbalanced within-subject design. Reinstatement tests were conducted under conditions identical to those of self-administration sessions, except that (1) a single presentation of the cues (light above the active lever on and house-light off for 1 min) was delivered response-independently immediately at the start of the session and (2) responses on the active lever (on a FR 5 schedule) resulted in contingent presentation of the cues without nicotine availability (no injections). Responses on the inactive lever were
recorded but had no programmed consequence. The testing sessions lasted 1 h.

**Nicotine discrimination procedure**

Rats acquired food-maintained behaviour as described previously (Justinova et al. 2009; Le Foll et al. 2005, 2008). Under a discrete-trial schedule of food-pellet delivery, rats learned to respond on one lever after an injection of a training dose of 0.4 mg/kg nicotine and on the other lever after an injection of 1 ml/kg of saline vehicle (n=10). Injections of nicotine or saline were given s.c. 10 min before the start of the session. At the start of the session, a white house-light was turned on and in its presence the rats were required to make ten consecutive responses (FR 10 schedule of food delivery) on the lever appropriate to the pre-session treatment. The completion of ten consecutive responses on the correct lever produced the delivery of a 45-mg food pellet and initiated a 45-s TO period during which lever-press responses had no programmed consequences and the chamber was dark. Responses on the incorrect lever had no programmed consequences other than to reset the FR requirement on the correct lever. After each TO period, the house-light was again turned on and the next trial began. Each session ended after the completion of 20 FR trials or after 30 min elapsed, whichever occurred first. Discrimination-training sessions were conducted 5 d per week under a double alternation schedule (i.e. DDSSDDSS etc., D = drug, S = saline). Training continued until there were eight consecutive sessions during which rats completed at least 90% of their responses during the session on the correct lever and no more than four responses occurred on the incorrect lever during the first trial. Test sessions with other doses and other drugs were then initiated.

During the test sessions, a range of doses of varenicline were substituted for the training dose of nicotine. The influence of different pretreatment time was evaluated at the following pretreatment times: 5, 10, 20, 40, 120, and 240 min before the session. Varenicline, given 120 min before the session, was also administered together with nicotine (given 10 min before the session). Test sessions were identical to training sessions, with the exception that both levers were active and ten consecutive responses on either one of the two levers resulted in the delivery of a food pellet. Switching responses from one lever to the other lever reset the ratio requirement. In a test phase, a single alternation schedule was introduced and test sessions were usually conducted on Tuesdays and Fridays. Thus, a 2-wk sequence starting on Monday was: DTDSTDST (T = test). In this way, test sessions occurred with equal probability after saline and drug sessions. Test sessions were conducted only if the criterion of 90% accuracy and not more than four incorrect responses during the first trial was maintained in two preceding training sessions.

For the drug-discrimination studies, two independent measures of behaviour were collected: a measure of discrimination performance expressed as the percentage of nicotine-associated responses and a measure of motor performance expressed as response rate. The percentage of nicotine-associated responses during each session (training or test) reflected the percentage of the number of responses elicited on the nicotine-associated lever relative to the total number of responses elicited on both levers during a session. The percentage of nicotine-associated responses was individually calculated for each rat and then expressed as a group mean (±S.E.M.). Nicotine-associated lever selection data were excluded from analysis if a rat had fewer than ten responses during the test session. Full generalization to the nicotine cue was defined as a percentage of responding on the nicotine-associated lever of ≥80%. No generalization to the nicotine cue was defined as a percentage of responding on the nicotine-associated lever of 20% or lower. Partial generalization to the nicotine cue was defined as a percentage of responding on the nicotine-associated lever ranging from >20% to <80%.

Response rates (responses/s) during each session were calculated by dividing the total number of responses on both levers during a session by the total session length. Response rates were individually calculated for each rat and then expressed as group means (±S.E.M.)

**Data analysis**

For drug-discrimination studies, results were subjected to two-way ANOVAs followed by LSD post-hoc tests. For drug self-administration studies, repeated-measures ANOVAs, followed when appropriate by post-hoc Dunnett’s tests for comparisons with the BL condition (the BL value was the mean of the values the day before each test session with an injection with the appropriate vehicle) for self-administration studies under the PR schedule of reinforcement; and by post-hoc Newman–Keuls tests for multiple comparisons for studies on reinstatement of nicotine-seeking behaviour. Changes were considered significant when p < 0.05.
Results

Influence of pretreatment time on response rates and ability of varenicline to generalize to the nicotine cue

Two-way ANOVA performed on the response rates of the rats indicated that there was a significant effect of varenicline dose \((F_{1,88} = 68.5, p < 0.0001)\), a significant effect of pretreatment time \((F_{5,88} = 7.5, p < 0.0001)\) and a significant interaction between varenicline dose and pretreatment time \((F_{5,88} = 7.3, p < 0.0001); \text{ see Fig. 1b}.\)

Post-hoc analysis indicated that there was no disruption of response rates in rats receiving 1 mg/kg varenicline (all \(p > 0.38\)). In contrast, 3 mg/kg had marked effects on the ability of the rats to respond. Rats receiving varenicline 5 and 10 min \((p < 0.0001\) for both) and 20 min \((p < 0.01)\) before the session had a significant disruption of responding compared to rats receiving varenicline 120 or 240 min before the sessions.

Two-way ANOVA performed on the percentage of lever presses performed on the nicotine-associated lever indicated no significant effect of varenicline dose \((F_{1,88} = 0.003, p = 0.99)\), a significant effect of pretreatment time \((F_{5,88} = 3.7, p < 0.01)\) and no significant interaction between varenicline dose and pretreatment time \((F_{5,88} = 0.8, p = 0.6)\). Rats receiving 1 mg/kg varenicline 5 min prior to the session had full generalization, whereas rats administered at 10, 20, 40, 120 and 240 min before the test session displayed partial generalization (see Fig. 1a). The ability of 1 mg/kg varenicline to generalize for nicotine cue was
significantly lowest in rats receiving varenicline 120 min (p=0.007) and 240 min (p=0.003) before the session compared to 5-min pretreatment time. The ability of 3 mg/kg varenicline to generalize for nicotine cue was significantly lower in rats receiving varenicline 240 min (p=0.02) before the session compared to 5-min pretreatment time.

**Effect of varenicline administered 2 h before the session on nicotine discrimination**

Varenicline administered in combination with various doses of nicotine or alone did not significantly modify the rate of responding by the rats when administered 120 min before the session. Two-way ANOVAs performed on response rates indicated that there was no significant effect of varenicline (F_{1,152}=1.7, p=0.18), no significant effect of nicotine dose (F_{3,152}=0.6, p=0.7) and no significant interaction between varenicline and nicotine (F_{9,152}=0.7, p=0.7; Fig. 1d).

Varenicline given 120 min before the session, followed by various doses of nicotine given immediately prior to the session, significantly modified nicotine discrimination performance of the rats. Two-way ANOVAs performed on percentage of responses on the nicotine lever indicated a significant effect of varenicline (F_{1,152}=12.9, p<0.0001), a significant effect of nicotine dose (F_{3,152}=8.8, p<0.0001) and a significant interaction between varenicline and nicotine (F_{9,152}=4.4, p<0.0001) (Fig. 1c). Post-hoc analysis indicated that in rats pretreated with 1 mg/kg or 3 mg/kg varenicline, followed by administration of 0.1–0.4 mg/kg doses of nicotine there was no significant change in discrimination performance compared to pretreatment with vehicle (all p>0.55 and p>0.26 for 1 and 3 mg/kg varenicline, respectively). Although, the discrimination of the training dose of nicotine (0.4 mg/kg) appeared to be antagonized to a small extent by varenicline pretreatment, the effect did not reach statistical significance. Discrimination performance with lower nicotine dose 0.03 mg/kg was enhanced after pretreatment with 1 and 3 mg/kg varenicline, compared to vehicle-treated animals (all p<0.01), but this was not the case with the lowest dose of nicotine (0.01 mg/kg), where the effect of varenicline pretreatment was only additive.

**Effects of varenicline (0.3, 1 and 3 mg/kg i.p., 2-h pretreatment time) on motivation for food assessed by the PR schedule**

The repeated-measures ANOVA performed on the number of food pellets that the rats (n=10) received following varenicline or vehicle administration (2 h pretreatment time) showed no main effect of treatment (F_{4,56}=1.7, p>0.05) (Fig. 2a).

The repeated-measures ANOVA performed on inactive lever responding showed no main effect of treatment (F_{4,56}=9.1, p<0.0001). The means ± S.E.M. for BL 0.3, 1.0, 3.0 mg/kg and saline substitution were 22.3±4.3, 22.4±5.1, 26.4±7.9, 16.6±3.9 and 61.8±7.4, respectively. Multiple comparisons showed that the saline substitution resulted in significantly higher inactive lever responding compared to all other treatments (p<0.0001). However, no dose of varenicline produced any significant difference in inactive lever responding compared with BL.

**Effects of varenicline (0.3, 1 and 3 mg/kg i.p., 120 min pretreatment time) on cue-induced reinstatement of nicotine-seeking behaviour**

The ANOVAs performed on active and inactive lever presses of rats (n=8) indicated a main effect of lever type (F_{1,56}=770, p<0.0001), of varenicline (F_{4,56}=94.3, p<0.0001) and a significant interaction between lever type and varenicline treatment (F_{4,56}=79.8, p<0.0001; Fig. 3). The post-hoc analysis showed that cue presentation elicited a significant increase of the active (p=0.0002), but not the inactive (p=0.6) lever presses, under the vehicle condition compared to extinction responding. Pretreatment with 0.3 mg/kg varenicline significantly increased the cue-induced reinstatement (p=0.008) compared to vehicle, whereas pretreatment...
with 1 and 3 mg/kg varenicline significantly decreased the reinstatement (both $p < 0.001$). In addition, no significant changes in responding on the inactive lever were observed under all conditions (all $p > 0.6$; mean number of inactive lever presses ± S.E.M. under the various testing conditions fluctuated between 6.3 ± 1.3 and 9.4 ± 2.7).

**Discussion**

This is the first study to evaluate the impact of pretreatment time on discriminative-stimulus effects induced by varenicline; the first to evaluate the effect of varenicline on motivation to respond for nicotine under a PR schedule of reinforcement; and the first to report a significant decrease in the ability of nicotine-associated cues to induce nicotine-seeking behaviour following varenicline administration.

In the present study, varenicline produced full or high levels of partial generalization to the nicotine cue when administered at very short pretreatment times (5, 10, 20 min). At later time-points, varenicline produced lower levels of partial generalization. These data are consistent with previous findings by Rollema et al. (2007) where a pretreatment time of 5 min was used. Interestingly, another study reporting a lesser ability of varenicline to generalize to nicotine was performed using a longer pretreatment time of 25 min (LeSage et al. 2009). However, since those two studies did not directly evaluate the impact of pretreatment time, our study is the first to reveal that this aspect is important.

In the present experiments, nicotine (30 μg/kg per infusion) supported self-administration under a PR schedule in Long–Evans rats at levels comparable to previously reported studies (Donny et al. 1999; Forget et al. 2010a,b). Consistent with previous findings obtained using FR schedules (George et al. 2011; O’Connor et al. 2010), varenicline in the present study dose-dependently reduced the ability of nicotine to maintain self-administration behaviour. Interestingly, the effect of varenicline was of the same magnitude as the effect of substituting saline for nicotine, indicating that the reinforcing efficacy of nicotine was fully reversed by varenicline. Moreover, in our study, varenicline did not affect responding for food under the same schedule of reinforcement. Some investigators have reported that varenicline, administered 15 min prior to the session, can disrupt responding for food (O’Connor et al. 2010), while others have reported that varenicline increases responding for food or sucrose when administered either 15 or 30 min prior to the session (Rollema et al. 2007; Wouda et al. 2011). Our study indicates that pretreatment time is a critical factor that can influence the effects of varenicline on rates of responding (see Fig. 1). Other factors could have some influence under specific conditions, for example use of naive animals, different route of administration, etc. Moreover, it should be noted that Sprague–Dawley rats were used for the
drug-discrimination experiments and Long–Evans rats for the food and nicotine self-administration experiments, respectively. Although using those two strains allowed us to be consistent with our previously published work, we cannot exclude the possibility that the pretreatment–time effect may not be similar in the two rodent species. In addition, further parametric studies varying the unit dose of varenicline and nicotine on different addictive models could be of value in clarifying the interactions between nicotine and varenicline. Those parametric studies could explain why low doses of varenicline increased cue-induced nicotine seeking, while higher doses of varenicline decreased cue-induced nicotine seeking.

The major finding of the present experiments is that varenicline significantly reduces cue-induced reinstatement of nicotine seeking (Fig. 3). These findings are consistent with previous studies in humans indicating that varenicline can reduce the effects of cue presentation on brain activity in human subjects (Franklin et al. 2011). In contrast, no significant effects of varenicline on cue-induced reinstatement have previously been reported in rats, although a trend towards significant attenuation was recently reported when varenicline was administered 30 min prior to cue-induced reinstatement (Wouda et al. 2011). We believe that the long 2-h pretreatment time used here could explain the different effects found in our present study and previous studies.

One limitation of the current study is the use of only one drug. Since varenicline has potentially other targets than α4β2 nicotinic receptors, we cannot directly implicate a particular nicotinic subunit in these effects (Rollema et al. 2007). However, this was not the goal of the present experiments, which were performed to explore the effects of an established treatment compound, varenicline, in animal models that will be used to screen for future medications for treatment of nicotine dependence. It will be of value to expand the present findings to other models such as intracranial self-stimulation and withdrawal, which could reflect other aspects of the addiction cycle (Lerman et al. 2007). Some of the biochemical results reported with varenicline indicate that varenicline can produce both agonist and antagonistic effects depending on the manner it is administered (Rollema et al. 2009). Here, varenicline administered without nicotine produced partial generalization to the nicotine cue in the drug-discrimination paradigm, suggesting some agonistic effects. In contrast, when administered in the drug-reinstatement paradigm (in the absence of nicotine), varenicline reduced nicotine-seeking. This effect could be mediated through the agonistic properties of varenicline, although we did not perform experiments to explore this hypothesis directly. When administered in combination with nicotine, varenicline decreased the motivation to self-administer nicotine under the PR nicotine self-administration paradigm and displayed a partial agonist profile in the drug-discrimination paradigm, where it enhanced discrimination of low doses of nicotine and, to a small extent, attenuated discrimination of the training dose of nicotine (although this was not statistically significant). These findings suggest that the antagonistic properties of varenicline can be demonstrated under certain conditions.

In conclusion, the present study explored time-course of effects of varenicline pretreatment and showed that pretreatment time significantly affected
generalization of varenicline to a nicotine cue and also affected rates of responding for food in a drug-discrimination setting. Thus, varenicline pretreatment time should be considered an important variable under different experimental conditions. Moreover, the present findings support and extend previous findings that varenicline is able to act on several aspects of nicotine dependence. Animals pretreated with varenicline displayed decreased motivation for nicotine as assessed by a PR drug self-administration procedure. In addition, varenicline caused the presentation of nicotine-associated stimuli to be less effective at reinstating nicotine-seeking behaviour. These results support the use of nicotine-taking and nicotine-seeking paradigms as screens for evaluating future medications for nicotine dependence treatment.

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