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Investigation of endocannabinoid modulation of conditioned responding evoked by a nicotine CS and the Pavlovian stimulus effects of CP 55,940 in adult male rats

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Abstract

Rationale — The cannabinoid CB1 receptor antagonist/inverse agonist rimonabant (SR 141716) has been shown to block reinforcing and rewarding effects of nicotine. Research has not investigated whether the cannabinoid system is involved in the interoceptive stimulus effects of nicotine functioning as a conditional stimulus (CS).

Objective — We examined the effects of rimonabant and the CB1/2 receptor agonist, CP 55,940, on responding evoked by a nicotine CS in rats. Additionally, we determined whether CP 55,940 functioned as a CS or a Pavlovian positive drug feature

Materials and methods — Pavlovian discrimination training involved intermixed nicotine (0.2 mg base/kg) and saline sessions with intermittent access to water only on nicotine. Antagonism tests with rimonabant (0.1–3 mg/kg) and substitution tests with CP 55,940 (0.003–0.1 mg/kg) followed. An effective dose of CP 55,940 was tested against the nicotine generalization curve. A separate group received CS training with CP 55,940 (0.01 mg/kg). Two other groups were trained using CP 55,940 (0.01 or 0.03 mg/kg) as a positive drug feature in which a brief light CS signaled access to water only on CP 55,940 sessions.

Results — Rimonabant blocked nicotine-evoked responding. CP 55,940 partially substituted for nicotine and enhanced responding to lower nicotine doses. Overall, CP 55,940 did not acquire control of conditioned responding in either Pavlovian drug discrimination task.

Conclusions — The cannabinoid system was involved in the CS effects of nicotine. This finding is counter to the operant drug discrimination research with nicotine as a discriminative stimulus, warranting further research into this possible dissociation.

Keywords: appetitive conditioning, associative learning, cannabinoid receptor, nicotinic acetylcholine, occasion setting, smoking cessation

Introduction

Recent attention has been given to cannabinoid compounds for use in smoking cessation. Of particular interest have been compounds related to the cannabinoid CB1 receptor antagonist/inverse agonist rimonabant, or SR 141716. Cannabinoid antagonists have appeared promising for use as smoking pharmacotherapies, and several pharmaceutical companies have worked to develop a marketable compound. For example, another CB1 antagonist, surinabant (SR 147778; Lamota et al. 2008; Rinaldi-Carmona et al. 2004) completed Phase II clinical trials in Europe in 2008 before development was discontinued because of adverse side effects.

Preclinical findings in rodents describing the role of cannabinoid activation on behavioral effects of nicotine have been reviewed elsewhere (e.g., Beardsley and Thomas 2005; Castañé et al. 2005). Briefly, rimonabant has been found to decrease nicotine self-administration (Cohen et al. 2002; Kodas et al. 2007), cue-induced reinstatement of nicotine seeking (Cohen et al. 2004; De Vries et al. 2005), and nicotine-conditioned place preferences (Forget et al. 2005; Le Foll and Goldberg 2004). The operant discriminative stimulus (SD) effects of nicotine do not appear to be affected by pretreatment with rimonabant (Cohen et al. 2002; Le Foll and Goldberg 2004; Zaniewska et al. 2006).
These cannabinoid effects on nicotine led us to examine the potential role for cannabinoid activation in the expression of the conditional stimulus (CS) effects of nicotine in an appetitive Pavlovian conditioning task (cf. Besheer et al. 2004). In this task, rats receive a subcutaneous (SC) injection of nicotine or saline before placement in a conditioning chamber. On nicotine sessions, liquid sucrose (i.e., the unconditioned stimulus; US) is delivered intermittently. Sucrose is not available on intermixed saline sessions. Using head entries into the sucrose receptacle before the first sucro delivery as a measure of the conditioned response (i.e., CR; goal tracking; Farwell and Ayres 1979), nicotine serves as an interoceptive CS as evidenced by increased dipper entries on nicotine compared to saline sessions. The most notable procedural distinction between this task and the operant drug discrimination models is that there is no explicit response requirement for reinforcement to be delivered in the discriminated goal-tracking task. Although a rat must insert its head into the dipper to access the sucrose, the delivery of the sucrose is determined by the experimenter—not the rat. Recent research suggests that the CS effects of nicotine might involve somewhat different neuropharmacological processes than that of a S\textsuperscript{D}. Specifically, N-methyl-D-aspartate receptor blockade attenuated conditioned responding evoked by a nicotine CS (Murray and Bevins 2007a), but has not been shown to affect operant responding controlled by S\textsuperscript{D} effects of nicotine (Kim and Brioni 1995; Zakharova et al. 2005). Therefore, we examined the effects of the cannabinoid CB\textsubscript{1} receptor antagonist/inverse agonist, rimonabant (Pan et al. 1998; Rinaldi-Carmona et al. 1994) and the non-selective cannabinoid CB\textsubscript{1/2} receptor agonist, CP 55,940 (Little et al. 1988; Thomas et al. 1998) on nicotine-evoked conditioned responding. We also examined whether CP 55,940 served as a CS or as a Pavlovian drug feature.

Materials and methods

Subjects For all experiments, we used male Sprague-Dawley rats (314 ± 10 g at start of study) from Harlan (Indianapolis, IN, USA) that were previously used in brief cocaine/novelty place conditioning experiments (Reichel and Bevins 2008; unpublished research). Rats were taken from control and low-dose cocaine groups; care was taken to match histories as much as possible across the different conditions of the present report. Before the start of the present studies, rats were handled for at least 3 min/day for 3 days. They were housed individually in clear 48.3 × 26.7 × 20.3 cm (l × w × h) polycarbonate cages lined with wood shavings in a temperature- and humidity-controlled colony. Food was continuously available in the home cage. Rats were 23-h water-restricted; access to water for the hour occurred after each daily session. All sessions were conducted during the light portion of a 12-h light:dark cycle with lights on at 0600 h. Protocols were approved by the University of Nebraska-Lincoln Animal Care and Use Committee and followed the “Guidelines for the Care and Use of Mammals in Neuroscience and Behavioral Research” (National Research Council 2003).

Apparatus Eight conditioning chambers (ENV-008CT; Med Associates, Inc., St. Albans, VT, USA) measuring 30.5 × 24.1 × 21.0 cm (l × w × h) were used. Sidewalls were aluminum; the ceiling and front and back walls were clear polycarbonate. Each chamber was equipped with a recessed receptacle (5.2 × 5.2 × 3.8 cm; l × w × d) on one sidewall. A dipper arm raised a 0.1-ml cup of distilled water into the receptacle. Water, rather than sucrose, served as the US because cannabinoid compounds can affect caloric consumption (e.g., Glick and Milloy 1972; Xie et al. 2007). An infrared emitter/detector unit, 1.2 cm into the receptacle and 3 cm above the chamber floor, monitored head entries into the dipper. A second infrared emitter/detector unit, 4 cm above the rod floor, bisected the chamber 14.5 cm from the sidewall containing the receptacle. This unit provided a measure of locomotor activity in the chamber. Two white stimulus lights (2.54 cm diameter; 28 V, 100 mA) were each mounted on the sidewall on either side of the dipper receptacle, 14.6 cm above the metal rod floor and 3.5 cm from either the front or the back wall. Illumination of the lights served as the discrete CS in the CP 55,940 positive feature experiments. Each chamber was enclosed in a light- and sound-attenuating cubic fitting with a fan to provide airflow and mask noise. A personal computer with Med Associates interface and software (Med-PC for Windows, version IV) controlled water deliveries and recorded dipper entries and locomotor activity.

Drugs (−)-Nicotine hydrogen tartrate (Sigma, St. Louis, MO, USA), rimonabant (RTI International, Research Triangle Park, NC, USA), and CP 55,940 (RTI International) were used. Nicotine was mixed in saline then adjusted to a pH of 7.0 ± 0.2. Rimonabant and CP 55,940 were mixed in a 1:1:18 Tween-80: 100% ethanol: distilled water solution (IPI). Rimonabant and CP 55,940 were injected intraperitoneally (IP). Rimonabant was given at 2 mg/kg with a 40 min IPI (cf. Wiley et al. 1995a); CP 55,940 was given at a volume of 1 ml/kg with a 30 min IPI (cf. De Vry and Jentzschi 2003; Mauler et al. 2002). Nicotine doses are reported in base form.

Nicotine CS training and testing Rats (n = 16) were given an injection of 0.2 mg/kg nicotine each day for 3 days in the home cage to attenuate the initial locomotor suppressant effects of nicotine (cf. Murray and Bevins 2007a).
Daily training sessions began the day after the last nicotine injection. Rats received either 0.2 mg/kg nicotine or saline before the start of each 20-min session. During nicotine sessions, there were 36 deliveries of 4 s access to water. Four different programs that varied when water was delivered were created to discourage timing of water deliveries. The average time before the first water delivery across programs was 137 s with a range of 124–152 s. No water was delivered during saline sessions, but there were 4-s “empty” intervals to maintain consistency between nicotine and saline sessions. Session types and programs were randomly assigned with the restriction that no more than two nicotine or two saline sessions occurred in a row. Training continued for 28 sessions. Following acquisition of the discrimination, rats entered testing. On the first four consecutive days of each 5-day cycle, rats received two nicotine and two saline training sessions as described previously. If the discrimination criterion was met (see later), a 4-min test session occurred in place of a training session on day 5; water was withheld in testing. If the criterion was not met, the rat remained in its home cage on that day. Nicotine generalization was tested first, followed by tests of rimonabant antagonism then CP 55,940 substitution. Finally, nicotine generalization was conducted again concurrently with CP 55,940 pretreatment of the nicotine dose–effect curve. Within each phase, the test doses were randomly intermixed for each rat. All ligands and doses in a phase were completed before beginning the next phase.

**CP 55,940 CS training**  A separate set of rats (n = 16) received either 0.01 mg/kg CP 55,940 or vehicle before each 20-min session. Water was delivered only in the CP 55,940 sessions. The programs and their order were the same as the nicotine CS experiment. Following 56 sessions of training with the CP 55,940 CS, rats were randomly assigned to one of two groups. One group remained with CP 55,940 training as described for another eight CP 55,940 and eight vehicle sessions. The other group was switched from CP 55,940 training to 0.2 mg/kg nicotine training for eight nicotine and eight saline sessions.

**CP 55,940 feature-positive training**  Rats (0.01/0.03 mg/kg group, n = 14; 0.03 mg/kg group, n = 6) received dipper training consisting of three 50-min sessions. Each daily session was initiated with the rat’s first head entry into the receptacle. The probability of receiving 4 s access to water decreased from 0.267 to 0.05 per 60 s over the three sessions. Acquisition training began the day after the last dipper training session. Rats were injected with CP 55,940 or vehicle 30-min before chamber placement. During each 20-min session, there were eight 15-s light CS presentations. On CP 55,940 sessions, each offset of the light was followed immediately by 4-s access to water. Four different programs were used to vary the timing of light presentations and water delivery. The average time to the first light onset was 135 s (range of 90–180 s) with a mean intertrial interval of 120 s (range of 74–165 s). On vehicle sessions, light presentations were matched with those of CP 55,940 sessions and 4-s “empty” intervals occurred in place of water deliveries to ensure identical session length. CP 55,940 and vehicle sessions were intermixed randomly with the restrictions that no more than two of a session type occurred in a row. The 0.01/0.03 mg/kg group received 60 training sessions at 0.01 mg/kg followed by 60 training sessions at 0.03 mg/kg CP 55,940; the 0.03 mg/kg group received 120 training sessions.

**Dependent measures**  In the CS experiments, the primary dependent measure was rate of dipper entries per second before the first water delivery. To allow for comparable measurement between drug (i.e., water) and vehicle (i.e., no water) sessions, the program types were matched for timing of the intervals from which the dipper entry rate was taken. The dependent measure for test sessions was the dipper entry rate in the first 2 min of the test. To qualify to test, dipper entry rate on each nicotine session was a minimum of 0.01 entries per second higher than each saline session within that testing cycle. Rate of locomotor activity in the chamber (beam breaks per second during the same interval as used for dipper entries) was also analyzed. In feature-positive experiments, the primary dependent measure was the initial elevation score of a session. This score was calculated as the number of dipper entries recorded during the first 15-s presentation of the light CS minus the number of dipper entries recorded during the 15-s interval before the light (CS period–pre-CS period). A positive value indicates more dipper entries during the CS. Locomotor activity was also recorded throughout the session.

**Data analyses**  For acquisition, two-way repeated measures analyses of variance (ANOVs) were conducted with Drug (nicotine versus saline or CP 55,940 versus vehicle) as one factor and Session as the other factor for the dipper entry and activity measures. Generalization, antagonism, and substitution tests were analyzed with one-way repeated measures ANOVAs with Drug Dose as the factor. Finally, the regeneration of the nicotine generalization curves with CP 55,940 pretreatment were examined with a two-way repeated measures ANOVA using Drug (CP 55,940 or none) as one factor and Nicotine Dose as the other factor. Significant effects in substitution and antagonism ANOVAs were followed with Tukey’s honestly significant difference (HSD) tests. Significant interactions in the two-way acquisition ANOVAs prompted selective use of paired t tests with Bonferroni’s correction to compare drug conditions within
each session. Median effective doses [ED$_{50}$s (95% confidence intervals)] were calculated using the least squares linear regressions on nicotine generalization curves and testing that resulted in full blockade of the CS effects of nicotine. Statistical significance was declared when $p < 0.05$ for all tests. Only significant values are reported.

**Results**

**Nicotine CS training and testing**  As shown in Figure 1a, the discrimination was acquired by the eighth day of training. This impression was supported by a significant Drug × Session interaction, $F(13, 195) = 13.03, p < .001$, followed by paired $t$ tests with Bonferroni’s correction indicating that for sessions 4–14, dipper entries were higher on nicotine than saline sessions, $t_{s}(15) ≥ 4.38, ps < 0.001$. Overall higher response levels on nicotine compared to saline was also supported by a main effect of Drug, $F(1, 15) = 100.49, p < 0.001$. Although inspection of the inset graph does not suggest a systematic effect of nicotine on locomotor activity, there was a Drug × Session interaction, $F(13, 195) = 1.84, p < .05$. However, none of the post-hoc comparisons were significant.

As shown in Figure 1b, conditioned responding diminished as the test dose of nicotine decreased from the training dose (0.2 mg/kg) to saline [ED$_{50}$ = 0.052 (0.03–0.074) mg/kg]. There was a main effect of Drug Dose, $F(5, 75) = 17.93, p < 0.001$. Follow-up Tukey’s HSD tests found that 0.05, 0.1, and 0.2 mg/kg nicotine evoked greater responding than saline, and that 0.025, and 0.4 mg/kg evoked lower responding than the 0.2 mg/kg training dose of nicotine, HSD$_{mmd}$ = 0.046. There was no significant change in locomotor activity (inset graph) as a function of nicotine dose.

Results from antagonism and substitution testing are shown in Figure 2. During these phases, two rats were removed from the study for failure to maintain the discrimination ($n = 14$). Pretreatment with increasing doses of rimonabant decreased the nicotine-evoked CR (Figure 2a). The main effect of Drug Dose, $F(4, 52) = 17.87, p < 0.001$, showed 1 and 3 mg/kg rimonabant decreased nicotine-evoked responding compared to saline pretreatment, HSD$_{mmd}$ = 0.045 [ED$_{50}$ = 1.09 (0.49–1.69) mg/kg]. Of those two doses, 1 mg/kg was also different from the vehicle baseline. Relative to vehicle pretreatment, the highest dose of rimonabant reduced nicotine-induced locomotor activity (inset graph), $F(4, 52) = 4.15, p < 0.01$, to a level comparable to vehicle only, HSD$_{mmd}$ = 0.078 (i.e., no nicotine).

There was an inverted U-shaped pattern of CP 55,940 substitution for the nicotine CS (Figure 2b). There was a main effect of Drug Dose, $F(4, 52) = 6.77, p < 0.001$. Conditioned responding on 0.01 mg/kg CP 55,940 was significantly higher than vehicle alone, HSD$_{mmd}$ = 0.028. This dose also differed from the 0.2 mg/kg nicotine baseline. The 0.1 mg/kg dose of CP 55,940 reduced locomotor activity relative to vehicle alone and nicotine (inset graph), $F(4, 52) = 6.61, p < 0.001$, HSD$_{mmd}$ = 0.06.

The dose of CP 55,940 that partially substituted for the nicotine CS (0.01 mg/kg) was assessed to determine if it could shift the nicotine dose–effect curve. During this phase, one more rat was removed for poor discrimination performance ($n = 13$). As shown in Figure 2c, CP 55,940 appeared to enhance nicotine-appropriate responding. There was a main effect of Drug, $F(1, 12) = 7.07, p < .001$, and a Drug × Nicotine Dose interaction, $F(3, 36) = 2.88, p < .05$. CP 55,940 enhanced re-
sponding to saline and to 0.1 mg/kg nicotine, HSD\textsubscript{mmmd} = 0.038 [ED\textsubscript{50} = 0.043 (0.025–0.06) mg/kg for nicotine alone; 0.049 (0.022–0.075) mg/kg for CP 55,940 pretreatment]. Although CP 55,940 showed partial substitution for the training dose of nicotine twice and enhanced responding to a low dose of nicotine, these effects did not translate into a shift in the ED\textsubscript{50} for nicotine. A Type I error seems unlikely given the replication of partial substitution. The outcome may instead be a product of not having enough test doses along the linear portion of the dose–effect curves. Locomotor activity appeared relatively stable across nicotine doses (inset graph). However, there was a significant Drug × Nicotine Dose interaction, \(F(3, 36) = 5.39, p < 0.01\); post-hoc comparisons found no differences HSD\textsubscript{mmmd} = 0.069.

**CP 55,940 CS training** When CP 55,940 was trained as the CS, the discrimination was not acquired even when training continued for twice as long as the nicotine CS (Figure 3a). There was a significant Drug × Session interaction, \(F(27, 405) = 1.90, p < 0.01\). However, none of the comparisons were significant following Bonferroni’s correction tests. Activity (Figure 3c) was similar regardless of drug treatment.

For rats that remained on CP 55,940 CS training, CP 55,940 evoked slightly higher levels of dipper entries than vehicle (Figure 3b, top panel) as shown by a main effect of Drug, \(F(1, 7) = 5.87, p < 0.05\). Although this result suggests potential CS effects of CP 55,940, the discrimination was weak and not maintained throughout training. There were no differences in activity for this subset of rats (Figure 3d, top panel). In contrast, rats switched to nicotine quickly acquired the discrimination (Figure 3b, bottom panel). There was a significant Drug × Session interaction, \(F(7, 49) = 14.74, p < 0.001\). There was more responding on nicotine than saline for sessions 31 and 33–36, \(t(7) ≥ 3.88, ps < 0.0063\). For activity (Figure 3d, bottom panel), there was a significant Drug × Session interaction, \(F(7, 49) = 2.25, p < 0.05\). Nicotine decreased activity on session 30, \(t(7) = 4.20, p < 0.0063\).

**CP 55,940 feature positive training:** 0.01/0.03 mg/kg group The light CS came to evoke a goal-tracking CR (Figure 4a). The main effect of Session supports the observation that dipper entries during the light presentations steadily increased, \(F(29, 377) = 4.48, p < 0.001\). However, rats did not discriminate between 0.01 mg/kg CP 55,940 sessions (water reinforced) and the vehicle sessions (non-reinforced). There was no main effect of Drug or Drug × Session interaction. After switching from 0.01 mg/kg to 0.03 mg/kg CP 55,940, weak discrimination performance emerged [main effect of Drug, \(F(1, 13) = 18.13, p < .001\)]. Response rates were stable across trials resulting in no main effect of Session or Drug × Session interaction. CP 55,940 evoked higher locomotor activity than vehicle at both training doses (Figure 4b). For

**Figure 2.** Panels a and b show dipper entries per second during rimonabant and CP 55,940 tests, respectively. *Inset graphs* for each panel show locomotor activity per second during the tests. *denotes significant difference from vehicle condition of the given drug test. * denotes a further significant difference from the control baseline (no nicotine for rimonabant and 0.2 mg/kg nicotine for CP 55,940). Panel c shows dipper entries per second for pretreatment of a range of nicotine doses with 0.01 mg/kg CP 55,940. *denotes significant difference from the same dose of nicotine with no pretreatment. The * inset graph shows locomotor activity per second
each dose, there were main effects of Drug, $F_s \geq 15.65$, $p_s \leq .002$, but no main effect of Session or Drug × Session interaction.

CP 55940 feature positive training: 0.03 mg/kg group  Similar to the 0.01/0.03 mg/kg group, when rats were trained with 0.03 mg/kg CP 55,940 from the outset a very weak discrimination developed (data not shown). There was a main effect of Drug, $F(1, 295) = 6.97$, $p < 0.05$, with the light CS evoking slightly higher initial elevation scores after CP 55,940 than vehicle administration. There was no effect of Session or Drug × Session interaction. Locomotor activity was slightly higher on CP 55,940 sessions than vehicle sessions, with an overall decrease in activity across sessions (data not shown).

These observations were supported by main effects of Drug, $F(1, 5) = 7.41$, $p < 0.05$, and Session, $F(59, 295) = 2.41$, $p < 0.001$, and a Drug × Session interaction, $F(59, 295) = 1.39$, $p < 0.05$. However, after Bonferroni’s correction, none of the comparisons were significant.

Discussion

Rats readily acquired the Pavlovian drug discrimination when 0.2 mg/kg nicotine was paired with a water US. This finding extends previous research from our laboratory to a new US. Past research used food restriction and a sucrose US. The studies that used 0.2 mg/kg nicotine as the training dose yielded similar to slightly higher $ED_{50s}$ [from 0.049 in Murray and Bevins (2007a)
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Reichel et al. (2007a) found that the nicotine dose of 0.075 mg/kg was used in the present study, which is higher than the 0.052 mg/kg nicotine found in the present study. We also found a role for endocannabinoid activation in the expression of the interoceptive CS effects of nicotine. Rimonabant partially blocked the CS effects of nicotine without affecting locomotor activity at 1 mg/kg. At 3 mg/kg, there was a concurrent reduction in locomotion and dipper entries, a finding that is potentially consistent with a response competition account of reduced locomotion after administration of rimonabant (Tallett et al. 2007). In that study, 1.5 and 3 mg/kg rimonabant reduced locomotion while enhancing grooming and scratching behaviors. Future research involving video recording of behavior during test sessions would be needed to assess this speculation. Our finding with the 1 mg/kg dose of rimonabant complements the well-documented role of endocannabinoids in the US or reinforcing effects of nicotine (e.g., Castañé et al. 2002; Kodas et al. 2007). However, the results are in contrast to the operant drug discrimination literature. In those studies, rimonabant at a range of 0.3 to 3 mg/kg (Cohen et al. 2002; Le Foll and Goldberg 2004) or even 5 to 10 mg/kg (Zaniewska et al. 2006) did not affect the distribution of nicotine-appropriate responding. In those studies, the effect of rimonabant on rates of responding was mixed, with two of the three studies showing no change in response rate (Le Foll and Goldberg 2004; Zaniewska et al. 2006). Furthermore, in the current study, 0.01 mg/kg CP 55,940 partially substituted for the training dose and enhanced responding to a low dose of nicotine. However, in the operant-conditioning study by Zaniewska et al. (2006), pretreatment with a higher dose of CP 55,940 (0.1 mg/kg) had no effect on nicotine discrimination performance or rate of responding.

There are several possible reasons for the differences between the present studies and those just described.
For instance, the dose of nicotine used as an $S^D$ was 0.4 mg/kg; the present study used 0.2 mg/kg nicotine as the CS training dose. The 0.4 mg/kg dose during the nicotine generalization test did not evoke a CR (i.e., dipper entries were at saline levels), a finding consistent with the suggestion that the interoceptive stimulus effects of 0.2 mg/kg differ from 0.4 mg/kg. In Murray and Bevins (2007a), 0.4 mg/kg also did not generalize to 0.2 mg/kg nicotine. Additionally, extinction of the 0.4 mg/kg nicotine CS proceeded more slowly than the 0.2 mg/kg CS (Murray and Bevins 2007b). Finally, in the operant drug discrimination task, rats trained on 0.4 mg/kg learned the discrimination faster than rats trained on 0.2 mg/kg (cf. Chance et al. 1977). Another reason may be that previous research with CP 55,940 or rimonabant and the stimulus effects of nicotine used the two-lever operant-discrimination task with separate schedules of reinforcement in force depending on drug state. The present research used a Pavlovian discriminated goal-tracking task in which the reinforcer was made available independent of responding. If this distinction eventuates to be important, then present results are the second neuropharmacological distinction found between the stimulus effects of nicotine as a CS versus an $S^D$. The first one was described by Murray and Bevins (2007a). In that study, the $N$-methyl-$d$-aspartate receptor channel blocker MK-801 attenuated conditioned responding evoked by a nicotine CS. In contrast, nicotine-appropriate operant responding was unchanged after MK-801 pretreatment in Zakharova et al. (2005) and Kim and Brioni (1995).

Because the interoceptive effects of CP 55,940 enhanced conditioned responding to the nicotine CS, we hypothesized that this cannabinoid agonist would also function as a CS. However, after twice the training as that for nicotine, there was still no suggestion that conditioned responding was coming under control of the CP 55,940 drug state. The subset of rats subsequently switched to nicotine quickly acquired the Pavlovian discrimination. Thus, CP 55,940 shares some stimulus effects with nicotine as measured by partial substitution for the nicotine CS and its enhancement of conditioned responding to a low non-training dose of nicotine. The stimulus effects of 0.01 mg/kg CP 55,940 itself, however, were not sufficient to function as a CS for access to water. Notably, switching to nicotine training revealed the locomotor-inhibiting effects of early nicotine exposure and suggests no cross-tolerance developed with CP 55,940 exposure.

Research from our laboratory has shown that thus far, drugs other than nicotine (i.e., amphetamine, caffeine, and chlor Diazepoxide) do not readily come to control a CR when trained as a CS (Murray et al. 2007; Palmatier and Bevins 2007). In contrast, up to this point, every drug state tested by us has functioned as a positive drug feature occasioning when a discrete CS will be reinforced. That list includes amphetamine, bupro-
mation in the present studies is due to the memory-deficit effects of cannabinoid activation. This seems unlikely given our findings that CP 55,940 increased nicotine-appropriate responding, whereas rimonabant decreased nicotine-appropriate responding. This pattern is the opposite of what would be expected had memory function been the primary factor involved in the cannabinoid modification of the nicotine CS. In addition, CP 55,940 effectively served as a S\(^D\) indicating that it can acquire control over behavior (e.g., Wiley et al. 1995a). Overall, we are led to the conclusion that at the dose used, and within the training conditions of the present research, CP 55,940 is not effective as a Pavlovian stimulus (i.e., a CS or positive feature).

Of course, acceptance of this conclusion without the above caveat would be premature given that there are numerous experimental parameters that could be manipulated. One example is the training dose. We started with 0.01 mg/kg CP 55,940 because this dose significantly modified conditioned responding to the nicotine CS. The 0.01 mg/kg CP 55,940 is a relatively low dose in the cannabinoid literature (cf. De Vry and Jentzsch 2003; Wiley et al. 1995a); a higher dose may be a more effective drug feature. Indeed, a weak discrimination developed when the training dose was increased to 0.03 mg/kg CP 55,940. However, when the higher dose of CP 55,940 served as the drug feature at the start of training (i.e., 0.03 mg/kg group) discrimination performance was still weak and inconsistent. Although operant drug discrimination research has shown that 0.03 mg/kg CP 55,940 was sufficient to function as an S\(^D\) (De Vry and Jentzsch 2003; Mauler et al. 2002), perhaps an even higher dose was needed in the Pavlovian positive feature task. Another potential parametric change is to increase the length of training. We trained the CP 55,940 CS for twice the number of sessions as the nicotine CS, and the CP 55,940 positive features were trained for more than twice that of the CP 55,940 CS. Further, De Vry and Jentzsch (2003), as well as Mauler et al. (2002), showed that rats acquired the 0.03 mg/kg CP 55,940 operant discrimination in a median of 38 training sessions (cf. to the 120 training sessions in the present CP 55,940 experiments). It seems unlikely that more training would have conditioned a stronger discrimination in the present studies.

Overall, the results of the present research indicate a mediating role of endocannabinoids in the nicotine CS. However, the cannabinoid agonist, CP 55,940, does not have Pavlovian stimulus effects using the current parameters. Because rimonabant does not alter schedule-controlled responding by a nicotine S\(^D\) (Cohen et al. 2002; Le Foll and Goldberg 2004; Zaniewska et al. 2006), and because CP 55,940 functions as an effective operant S\(^D\) at the doses used here (De Vry and Jentzsch 2003; Mauler et al. 2002; Wiley et al. 1995a, b), the current research suggests a potential dissociation between the mechanisms mediating operant and Pavlovian stimulus effects of nicotinic and cannabinoidergic compounds that warrant further investigation (cf. Murray and Bevins 2007a).

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