Discriminative Stimulus Effects of Tramadol in Humans

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ABSTRACT

Tramadol is an unscheduled atypical analgesic that acts as an agonist at μ-opioid receptors and inhibits monoamine re-uptake. Tramadol can suppress opioid withdrawal, and chronic administration can produce opioid physical dependence; however, diversion and abuse of tramadol is low. The present study further characterized tramadol in a three-choice discrimination procedure. Nondependent volunteers with active stimulant and opioid use (n = 8) participated in this residential laboratory study. Subjects were trained to discriminate between placebo, hydromorphone (8 mg), and methylphenidate (60 mg), and tests of acquisition confirmed that all volunteers could discriminate between the training drugs. The following drug conditions were then tested during discrimination test sessions: placebo, hydromorphone (4 and 8 mg), methylphenidate (30 and 60 mg), and tramadol (50, 100, 200, and 400 mg). In addition to discrimination measures, which included discrete choice, point distribution, and operant responding, subjective and physiological effects were measured for each test condition. Both doses of hydromorphone and methylphenidate were identified as hydromorphone- and methylphenidate-like, respectively. Lower doses of tramadol were generally identified as placebo, with higher doses (200 and 400 mg) identified as hydromorphone, or opioid-like. The highest dose of tramadol increased ratings on the stimulant scale, but was not significantly identified as methylphenidate-like. Tramadol did not significantly increase subjective ratings associated with reinforcement. Taken together, these results extend previous work with tramadol as a potential medication for the treatment of opioid dependence and withdrawal, showing acute doses of tramadol exhibit a profile of effects similar to opioid agonists and may have abuse liability in certain populations.

Introduction

Tramadol is an unscheduled atypical analgesic marketed as Ultram (Ortho-McNeil-Janssen Pharmaceutical, Titusville, NJ) and as generic. Tramadol exerts its analgesic effects in humans and animals through activation of two different systems; it is an agonist at μ-opioid receptors and inhibits monoamine re-uptake, specifically serotonin and nor-epinephrine (Raffa et al., 1992; Desmeules et al., 1996; for review see Grond and Sablotzki, 2004; Ide et al., 2006). The racemic form of tramadol has affinity for μ-opioid receptors but is less potent than morphine (Raffa et al., 1992). A metabolite formed after first-pass metabolism, O-desmethyl-tramadol (M1), possesses a higher affinity for μ-opioid recep-

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ABBR EVIATIONS: HM, hydromorphone; MPH, methylphenidate; VAS, visual analog scale.

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References

Cicero, F. T., Moye, A. L., Blakely, D. J., Nettles, C. A., and Atkinson, B. F. (1999). Monitoring tramadol use in the United States in 1994 (Cicero et al., 2005, Woody et al., 2003). Consistent with this unique pharmacological profile, tramadol exhibits some opioid agonist-like effects, but has lower abuse potential than typical opioid analgesics (Zacny, 2005, Epstein et al., 2006). For example, tramadol maintained lower rates of self-administration compared with lefetamine, morphine, and remifentanil in monkeys and rats (Yanagita, 1978, O’Connor and Mead, 2010), but retains analgesic effects (Raffa et al., 1992; Ide et al., 2006). These preclinical findings are consistent with results from clinical laboratory studies suggesting that tramadol does not produce significant morphine-like effects (Preston et al., 1991; Cami et al., 1994; Adams et al., 2006; Lofwall et al., 2007). Although chronic tramadol administration has the ability to produce physical dependence in the laboratory (Yanagita, 1978, Lanier et al., 2010), epidemiological and postmarketing surveillance of tramadol reports low abuse and diversion since its introduction in the United States in 1994 (Cicero et al., 1999, 2005, Woody et al., 2003; Inciardi et al., 2006).
Modest opioid agonist activity in an oral product is desirable for opioid withdrawal treatment. Ideally, a medication would exhibit enough efficacy to relieve opioid withdrawal symptoms, but not enough to support significant abuse or regulatory scheduling that would limit clinical availability. Because it exhibits some opioid agonist characteristics but with lower abuse liability compared with full \( \mu \)-opioid receptor agonists, tramadol may be a useful therapeutic agent for opioid dependence. In opioid-dependent volunteers maintained on morphine, tramadol suppressed spontaneous opioid withdrawal induced by placebo substitution and did not significantly increase subject-rated effects of feeling high, drug liking, or drug effect (Carroll et al., 2006; Lofwall et al., 2007). In methadone-maintained volunteers, acute tramadol challenges failed to elicit significant morphine-like effects or precipitate withdrawal symptoms (Cami et al., 1994). Taken together, these data suggest tramadol may be useful in treating opioid withdrawal.

The current study used a drug discrimination paradigm to expand on prior human laboratory studies examining the effects of tramadol in experienced drug users. Drug discrimination is a behavioral tool that is useful for distinguishing a test drug from other drug classes, as well as for distinguishing activity at different opioid receptor systems (Herling and Woods, 1981, Young et al., 1984; Kamien et al., 1993; Dykstra et al., 1997). In prior studies acute tramadol elicited only modest opioid agonist-like subjective effects, suggesting possible nonopioidergic mechanisms. In animal drug discrimination procedures, tramadol fully substituted for morphine in morphine-trained rats, and this effect was attenuated with the opioid receptor antagonist naloxone (Ren and Zheng, 2000). Filip et al. (2004) reported an enhancement of tramadol discrimination after reboxetine, a norepinephrine reuptake inhibitor, and milnacipram, a serotonin and norepinephrine reuptake inhibitor, but not the selective serotonin reuptake inhibitors fluoxetine or venlafaxine. Although these compounds did not substitute for tramadol in tramadol-trained rats, these data suggest norepinephrine, and possibly serotonin, may play a role in the discriminative stimulus effects of tramadol.

To investigate further the pharmacological profile of tramadol the discriminative and subjective effects of tramadol were examined in humans. Because tramadol exerts activity at both opioid and monoamine systems, nondependent volunteers with recent sporadic opioid and stimulant use were trained to discriminate placebo, hydromorphone (HM), and methylphenidate (MPH) in a three-choice discrimination procedure (e.g., Preston et al., 1987; Jones et al., 1999). Doses of hydromorphone, methylphenidate, and tramadol were then tested. It was hypothesized that volunteers would successfully acquire the discrimination and that higher doses of tramadol would be identified primarily as an opioid agonist, but engender less opioid agonist-like subjective effects compared with hydromorphone.

### Materials and Methods

#### Subjects

Participants were volunteers with current sporadic opioid and stimulant use (including cocaine in all subjects), but they were not physically dependent on opioids or stimulants (Table 1). Eight male volunteers completed the study. Females were enrolled; however, none completed the protocol. Participants underwent routine medical screening that included a medical history, physical examination, EKG, chemistry, hematology, urine drug testing, and routine medical urinalysis testing (e.g., specific gravity, pH, etc.). Medical staff not involved in the study as investigators reviewed all results, and all subjects were found to be without significant medical problems. The Structured Clinical Interview for the DSM-IV was completed to ensure volunteers were not physically dependent on substances (except caffeine and nicotine). In addition, participants were monitored drug-free for 48 h after residential unit admission to ensure there was no evidence of physical dependence on drugs other than caffeine and nicotine.

Pregnancy and significant medical or psychiatric illness (e.g., insulin-dependent diabetes, schizophrenia) were exclusionary. Individuals seeking treatment were not enrolled in the study and were assisted in referral to community-based treatment programs. The Institutional Review Board approved the study, and all volunteers gave written informed consent and were paid for their participation.

#### Study Setting

Participants lived on a closed, 14-bed residential unit for the duration of the study. Breathalyzer testing for alcohol was completed on the day of admission and randomly at least twice weekly. In addition, urine samples were collected at admission and daily throughout the study and tested intermittently for the presence of illicit drugs using an EMIT system (Olympus AU400; Syva Co., San Jose, CA). No evidence of unauthorized alcohol or drug abuse was detected during the study. Participants did not have access to caffeinated beverages and were allowed to smoke cigarettes ad libitum, except 90 min before and during experimental sessions.

#### Drugs

Drugs were encapsulated in red/white capsules and filled with lactose. Each volunteer received four red/white capsules on each session day. Lactose-filled capsules served as placebo. During training sessions and tests of acquisition, volunteers received placebo, hydromorphone (8 mg), and methylphenidate (60 mg). During discrimination sessions, volunteers received placebo, hydromorphone (4 and 8 mg), methylphenidate (30 and 60 mg), and tramadol (50, 100, 200, and 400 mg). Compounds were obtained from commercial sources: hydromorphone (Abbott Laboratories, Abbott Park, IL), methylphenidate (Novartis Consumer Health, East Hanover, NJ), and tramadol (PriCara, Raritan, NJ).

### Table 1

#### Demographics

<table>
<thead>
<tr>
<th>Variable</th>
<th>Total (n = 8)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>40.3 (±2.3)</td>
</tr>
<tr>
<td>Sex (male)</td>
<td>8</td>
</tr>
<tr>
<td>Race (white)</td>
<td>5</td>
</tr>
<tr>
<td>Education (years)</td>
<td>12.4 (±0.4)</td>
</tr>
<tr>
<td>Opioid use</td>
<td></td>
</tr>
<tr>
<td>Years since first opioid use</td>
<td>15.8 (±2.3)</td>
</tr>
<tr>
<td>Days of opioid use in last 30 days</td>
<td>12.8 (±2.7)</td>
</tr>
<tr>
<td>Intravenous users</td>
<td>4</td>
</tr>
<tr>
<td>Lifetime use (years)</td>
<td>4.9 (±1.3)</td>
</tr>
<tr>
<td>Cocaine use</td>
<td></td>
</tr>
<tr>
<td>Years since first cocaine use</td>
<td>17.5 (±4.3)</td>
</tr>
<tr>
<td>Days of cocaine use in last 30 days</td>
<td>8 (±1.8)</td>
</tr>
<tr>
<td>Intravenous users</td>
<td>0</td>
</tr>
<tr>
<td>Lifetime use (years)</td>
<td>6.8 (±2.1)</td>
</tr>
<tr>
<td>Other drugs used in last 30 days*</td>
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</tr>
<tr>
<td>Alcohol</td>
<td>7</td>
</tr>
<tr>
<td>Cannabis</td>
<td>2</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>2</td>
</tr>
</tbody>
</table>

* Number of volunteers that used each drug.
All drug administration was double-blind. Each participant was assigned three arbitrary letters that corresponded to each training drug condition (placebo, 8 mg of hydromorphone, 60 mg of methylphenidate). Letters varied across volunteers, but remained unchanged for each volunteer throughout participation. Capsule administration occurred at 9:00 AM on each session day, which was 30 min before the start of postdrug assessments and 90 min before the start of discrimination assessments.

General Methods

After volunteers completed informed consent, they were admitted and oriented to the residential unit. Volunteers were informed that the purpose of the study was to examine the effects of tramadol and that they would be required to discriminate between placebo, an opioid, and a stimulant. Examples of each of these were given and participants were told during test sessions that they might experience no effects, opioid agonist effects, stimulant effects, or other effects. Volunteers were instructed to attend closely to the effects of each letter-coded drug. They were informed that correct identification of the administered drug by letter code would result in a monetary bonus. All volunteers had a practice session for familiarization with study procedures and measures; these data were not included in the analyses. Volunteers were permitted to eat a light breakfast (e.g., toast and juice) 45 min before sessions, but then they were allowed only water until session end.

There were three phases for each volunteer, although staff and participants were aware of only two phases (the second and third phases were indistinguishable). Daily sessions were conducted weekdays (Monday–Friday).

Discrimination Training (Phase 1)

The purpose of this phase was to train subjects to identify each condition by letter code. In random order each participant received at least two exposures to each training drug condition: placebo, hydromorphone (8 mg), and methylphenidate (60 mg). During training exposures, volunteers were informed which letter they were receiving immediately before drug administration and were again informed of the letter code at session end.

Test of Acquisition (Phase 2)

After training sessions, acquisition of discrimination was tested. The purpose of this phase was to test whether volunteers could identify each training drug condition by the correct letter code. Each volunteer received at least two exposures of each training drug condition in randomized order. Subjects were not informed of the letter code of the drug before drug administration. At each session end, volunteers were informed of the letter code of the administered drug condition and whether they had earned a monetary bonus for correctly identifying the drug by letter code. The criterion for acquisition of the discrimination was at least 67% correct responses for the combined drug conditions and at least one correct response for each drug condition.

Discrimination Test Sessions (Phase 3)

During this phase, doses of hydromorphone (4 and 8 mg), methylphenidate (30 and 60 mg), tramadol (50, 100, 200, and 400 mg), and placebo were tested in a random order. These sessions were conducted in the same manner as the test of acquisition sessions (phase 2), except that no feedback on letter code was provided on discrimination test days. Test of acquisition sessions (i.e., feedback about the letter code given after the session was completed) were interspersed with discrimination test sessions.

Experimental Sessions

Subject-Rated and Physiological Effects. Subject-rated effects and pupil diameter were collected 15 min before capsule administration, which was used for baseline, and at 90, 60, 90, 120, and 150 min after capsule administration. Pupil diameter was measured using a Neuroptics Pupilometer (Neuroptics Inc., Irvine, CA).

At each time point, volunteers completed three computer questionnaires rating the subjective effects of the drug condition administered: 1) visual analog scales (VAS), 2) an adjective rating scale, and 3) a pharmacological class questionnaire. On VAS items, volunteers placed an arrow along a 100-point line anchored with “not at all” and “extremely” to indicate the degree of effect produced by the drug condition. Participants rated drug effects as high, like, good effects, bad effects, sick, desire for cocaine now, similar to opioid, and similar to stimulant. In addition, participants rated the degree to which each drug condition was similar to each of the training drugs, as identified by letter code (e.g., similar to drug X; similar to drug Y). Volunteers rated adjectives on a five-point scale from 0 (no effect) to 4 (extremely). The adjective list constituted a 16-item opioid agonist scale (carefree, coating, drive, drugged, dry mouth, energetic, friendly, good mood, heavy or sluggish feeling, nervous, nodding, pleasant sick, relaxed, skin itchy, talkative/soapboxing, turning of stomach) and a 27-item stimulant scale (confused, craving for cocaine, difficulty concentrating, dizzy/lightheaded, drug effect, excited, fearful, feel a thrill, feeling of power, fidgety, headache, hungry, irritable, jittery, nausea, numbness, restless, seeing/hearing things, shaky (hands), sleepy, stimulated, suspicious, sweating, thirsty, tingling, tired, tremor). On the pharmacological class questionnaire, volunteers indicated which drug class was most similar to the drug condition they received that day. Ten drug classes were listed with descriptive labels and examples of each: placebo, opiates, phenothiazines, barbiturates, antidepressants, opiate antagonists, hallucinogens, benzodiazepines, stimulants, and other.

Discrimination Procedures. Discrimination was assessed at 90 and 120 min after capsule administration using three procedures: 1) discrete choice, 2) point distribution, and 3) operant responding. During discrete choice, volunteers chose the letter of the training drug that they thought they received. In point distribution, volunteers distributed 50 points among the three training drug letters depending on how certain they were of the identity of the administered drug. Lastly, volunteers emitted operant responses on computer keys that corresponded to the training letters, on a fixed interval 1-s schedule for 8.5 min. Points were earned for responses on each training drug. Payments during phases 2 and 3 for test of acquisition sessions were based on the accuracy of responses. The maximum possible payment for discrimination tasks was $10/session. On discrimination test days, payments were based on an average of the payments received for test of acquisition sessions.

Data Analysis

Data from the eight subjects who completed the protocol were included in the final analysis. To preserve testing in a random order, some volunteers received more than one exposure to a test drug during discrimination testing (phase 3). For data analysis purposes only the first exposure to each test drug was included, with the exception of sessions repeated because of malfunctions. To encompass the peak effects for each training drug condition, data from only the 120-min time point were used for analysis of the three discrimination measures. Peak effects for each session were determined for subjective and physiological measures. For most measures, the reported value was a peak increase; however, the peak increase and peak decrease were analyzed for pupil diameter. Means for the discrimination measures and peak effects for subjective and physiological measures were both analyzed using a repeated-measures regression model with an exchangeable covariance structure and an effect of drug condition. Pairwise comparisons were examined using a conservative one-step procedure, Tukey’s honestly significant difference. Peak placebo effects were compared with each drug condition. In addition, all tramadol conditions were compared with hydromorphone (4 and 8 mg) and methylphenidate (30 and 60 mg).
TABLE 2
Summary of discrimination measures
Numbers shown for operant responses and point distribution (maximum total = 50) are mean (S.E.M.). F ratio indicates a statistically significant overall regression (P < 0.0001, df = 8, 56). Lack of variability for discrete choice precluded statistical analysis. Numbers for discrete choice are mean percentage for each drug condition. All data are from the 120-min time point.

<table>
<thead>
<tr>
<th>HM (mg)</th>
<th>MPH (mg)</th>
<th>Tramadol (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Operant responses</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F Placebo</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>Placebo</td>
<td>16.4</td>
<td>455.5 (9.6)</td>
</tr>
<tr>
<td>HM</td>
<td>14.3</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>MPH</td>
<td>102.7</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td><strong>Point distribution</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>17.2</td>
<td>50.0 (0.0)</td>
</tr>
<tr>
<td>HM</td>
<td>15.0</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td>MPH</td>
<td>109.0</td>
<td>0.0 (0.0)</td>
</tr>
<tr>
<td><strong>Discrete choice</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Placebo</td>
<td>100</td>
<td>25</td>
</tr>
<tr>
<td>HM</td>
<td>0</td>
<td>75</td>
</tr>
<tr>
<td>MPH</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

<sup>a</sup>p < 0.05 vs. placebo.
<sup>b</sup>p < 0.05 vs. 4 mg HM.
<sup>c</sup>p < 0.05 vs. 8 mg HM.
<sup>d</sup>p < 0.05 vs. 30 mg MPH.
<sup>e</sup>p < 0.05 vs. 60 mg MPH.

Fig. 1. Reinforced operant responses for placebo (circles), hydromorphone (squares), and methylphenidate (triangles) during discrimination test sessions. Results are shown for the 120-min time point (Table 2). Placebo was associated with significantly higher methylphenidate-appropriate responding, whereas hydromorphone was associated with significantly higher hydromorphone-appropriate responding. Hydromorphone was identified as hydromorphone 75 to 100% of the time, whereas methylphenidate was identified as methylphenidate 100% of the time.
increased, volunteers identified it as hydromorphone on the majority of occasions (63–75%). The highest dose of tramadol was identified as methylphenidate or placebo 25 and 12% of the time, respectively (Table 2).

**Subjective and Physiological Effects.** On the pharmacological class questionnaire, volunteers identified placebo as placebo on 100% of occasions (Table 3). Doses of hydromorphone were predominantly identified as an opioid agonist on 75% (4 mg) and 100% (8 mg) of occasions, whereas both doses of methylphenidate were identified as a stimulant on 100% of occasions. Lower doses of tramadol were generally identified as placebo on 100% (50 mg) and 75% (100 mg) of occasions. As the dose or tramadol increased, identifications as placebo decreased. Higher doses of tramadol (200 and 400 mg) were primarily identified as an opioid agonist (63%). The remaining identifications for these doses were split between placebo and stimulant (Table 3).

**VAS Ratings of Similarity.** Volunteers rated how similar test doses were to each of the training drugs, as identified by letter code, and to a drug class (i.e., opioid, stimulant) on a 100-point visual analog scale (Table 4). Generally, volunteers rated placebo as most similar to placebo (Fig. 2). Compared with placebo, both doses of hydromorphone (4 and 8 mg) and one dose of tramadol (200 mg), but not methylphenidate, were rated significantly similar to hydromorphone (Fig. 2) and opioid (Table 4). Methylphenidate (30 and 60 mg), but not hydromorphone or tramadol, was rated significantly similar to methylphenidate and stimulant compared with placebo (Fig. 2; Table 4). Subject ratings of similarity of each drug test condition were consistent between training drugs identified by letter code and drug class.

**VAS Ratings of Effects.** Hydromorphone (8 mg), but not methylphenidate or tramadol, significantly increased ratings of like and good effects. Compared with placebo, hydromorphone (8 mg) and methylphenidate (60 mg) increased ratings of high and drug effect; however, the highest dose of methylphenidate significantly increased ratings of bad effects (Fig. 3; Table 4). Tramadol did not significantly increase ratings of like or good effects compared with placebo.

**Adjectives.** Volunteers rated a series of adjectives after each test drug, which constituted an opioid agonist scale and a stimulant scale. Compared with placebo, 8 mg of hydromorphone increased ratings on the opioid agonist scale, whereas both doses of methylphenidate and 400 mg of tramadol significantly increased ratings on the stimulant scale (Fig. 3).

**Pupil Diameter.** Hydromorphone (8 mg) significantly decreased pupil diameter compared with placebo and all doses of tramadol (Table 4). A statistically significant increase in pupil diameter was not observed (data not shown).

### Discussion

The purpose of the present study was to extend earlier work with tramadol by investigating the discriminative stimulus, subjective, and physiological effects of tramadol in non-dependent drug-experienced humans. Given that tramadol is an atypical analgesic that exerts agonist activity at μ-opioid receptors and inhibits monoamine reuptake, subjects were trained to discriminate placebo, an opioid receptor agonist, hydromorphone (8 mg), and a monoamine uptake-inhibiting stimulant, methylphenidate (60 mg; study phase 1). In subsequent discrimination testing, doses of hydromorphone occasioned hydromorphone-appropriate responding during the operant response discrimination task (Fig. 1), whereas doses of methylphenidate occasioned methylphenidate-appropriate responding. Higher doses of tramadol (200 and 400 mg) were associated with hydromorphone-, but not methylphenidate-, appropriate responding. A similar pattern of results was observed across all three discrimination tasks (Table 2).

Examination of subject-rated measures revealed additional effects of the three test drugs. Consistent with previous reports that drug discrimination and drug self-report measures are sensitive to detect the effects of stimulants and opioids (Kelly et al., 2003), effects obtained with these measures were similar across drug test conditions (Bickel et al., 1989). The pattern of results indicates that the 200-mg dose of tramadol engenders effects similar to an opioid agonist, whereas a higher dose of tramadol (400 mg) exerts mixed behavioral effects characteristic of an opioid agonist and stimulant.

To our knowledge, the current study is the first to report the acquisition of a three-choice discrimination using placebo, hydromorphone, and methylphenidate in nondependent drug-experienced humans. The use of methylphenidate and hydromorphone as training drugs in human discrimination procedures can detect stimulant- and opioid-like discriminative effects, respectively. Methylphenidate has been shown to share discriminative stimulus effects with other stimulants such as methamphetamine and d-amphetamine, suggesting that this drug is a useful pharmacological tool to detect stimulant-like effects for novel compounds (Stoops et al., 2005; Sevak et al., 2009). Likewise, hydromorphone has been used as a training drug to detect opioid agonist versus antagonist effects, as well as partial versus full μ-opioid receptor agonist effects (Preston et al., 1987; Jones et al., 1999; Preston and Bigelow, 2000). In the present study, the three-choice discrimination procedure was sensitive to detect opioid agonist effects and stimulant effects, as shown by differential responses to doses of hydromorphone and methylphenidate in discrimination tasks and subject-rated effects.

Doses of tramadol revealed a unique behavioral profile in the present study. Higher doses of tramadol resulted in hy-
dromorphone-appropriate responding, a pattern that was preserved across discrimination tasks (Table 2). Tramadol did not significantly increase ratings of drug liking, good effects, high, or drug effects, but did significantly increase scores on the stimulant scale at the highest dose tested. This unique pattern of results (i.e., hydromorphone-appropriate responding and stimulant scale scores) probably reflects tramadol’s activity at both \( /H_9262\)-opioid receptors and the monoamine system. However, the lack of subjective effects that are generally associated with increased abuse liability (e.g., significantly increased ratings of drug liking or good effects) is consistent with tramadol’s reportedly lower reinforcement efficacy and abuse (Cicero et al., 2005; Adams et al., 2006; Raffa, 2006; O’Connor and Mead, 2010). Although subjects

### Table 4

<table>
<thead>
<tr>
<th>F</th>
<th>Placebo</th>
<th>HM (mg) 4</th>
<th>HM (mg) 8</th>
<th>MPH (mg) 30</th>
<th>MPH (mg) 60</th>
<th>Tramadol (mg) 50</th>
<th>Tramadol (mg) 100</th>
<th>Tramadol (mg) 200</th>
<th>Tramadol (mg) 400</th>
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<tr>
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<td></td>
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<td></td>
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<tr>
<td>Similar to placebo</td>
<td>2.1</td>
<td>100.0 (0.0)</td>
<td>50.0 (18.9)</td>
<td>40.1 (16.7)</td>
<td>56.3 (17.5)</td>
<td>62.5 (18.3)</td>
<td>87.5 (12.5)</td>
<td>75.0 (16.4)</td>
<td>83.9 (12.5)</td>
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<td>Similar to HM</td>
<td>9.4**</td>
<td>0.0 (0.0)</td>
<td>86.9 (12.4)</td>
<td>100.0 (0.0)</td>
<td>12.5 (12.5)</td>
<td>13.4 (12.4)</td>
<td>12.5 (12.5)</td>
<td>25.0 (16.4)</td>
<td>72.0 (14.5)</td>
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<tr>
<td>Similar to MPH</td>
<td>47.3**</td>
<td>0.4 (0.4)</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>100.0 (0.0)</td>
<td>100.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>4.5 (4.5)</td>
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<tr>
<td>Similar to opioid</td>
<td>8.9**</td>
<td>0.0 (0.0)</td>
<td>62.9 (15.9)</td>
<td>85.3 (9.9)</td>
<td>12.5 (12.5)</td>
<td>7.4 (6.4)</td>
<td>12.5 (12.5)</td>
<td>7.4 (5.3)</td>
<td>50.0 (14.9)</td>
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<tr>
<td>Similar to stimulant</td>
<td>21.1**</td>
<td>0.4 (0.4)</td>
<td>0.0 (0.0)</td>
<td>0.9 (0.9)</td>
<td>79.9 (13.4)</td>
<td>85.5 (9.5)</td>
<td>12.5 (12.5)</td>
<td>7.4 (5.3)</td>
<td>50.0 (14.9)</td>
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<tr>
<td>Like</td>
<td>4.4**</td>
<td>0.0 (0.0)</td>
<td>29.0 (11.1)</td>
<td>44.1 (11.8)</td>
<td>11.3 (4.2)</td>
<td>23.3 (7.0)</td>
<td>13.1 (13.1)</td>
<td>6.1 (4.9)</td>
<td>14.9 (8.3)</td>
</tr>
<tr>
<td>Bad effects</td>
<td>3.6*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.1)</td>
<td>20.6 (12.0)</td>
<td>26.0 (11.7)</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>2.3 (1.2)</td>
</tr>
<tr>
<td>Good effects</td>
<td>5.3**</td>
<td>0.0 (0.0)</td>
<td>24.3 (8.3)</td>
<td>47.5 (11.9)</td>
<td>13.0 (4.6)</td>
<td>24.5 (6.2)</td>
<td>0.0 (0.0)</td>
<td>7.1 (5.4)</td>
<td>13.6 (8.2)</td>
</tr>
<tr>
<td>High</td>
<td>4.7**</td>
<td>0.0 (0.0)</td>
<td>19.8 (5.8)</td>
<td>29.0 (6.5)</td>
<td>22.1 (9.1)</td>
<td>37.1 (9.7)</td>
<td>0.4 (0.4)</td>
<td>5.8 (3.5)</td>
<td>10.5 (4.6)</td>
</tr>
<tr>
<td>Drug effect</td>
<td>5.0**</td>
<td>0.0 (0.0)</td>
<td>18.8 (5.2)</td>
<td>31.6 (6.4)</td>
<td>25.8 (9.5)</td>
<td>42.9 (10.0)</td>
<td>0.4 (0.4)</td>
<td>4.4 (3.0)</td>
<td>18.6 (7.5)</td>
</tr>
<tr>
<td>Subject Adjectives</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Opioid agonist scale</td>
<td>2.3*</td>
<td>1.4 (0.7)</td>
<td>2.9 (1.0)</td>
<td>4.8 (2.1)</td>
<td>2.1 (0.5)</td>
<td>3.4 (1.2)</td>
<td>0.4 (0.4)</td>
<td>1.6 (0.7)</td>
<td>2.1 (1.0)</td>
</tr>
<tr>
<td>Stimulant scale</td>
<td>8.6**</td>
<td>1.4 (0.8)</td>
<td>3.6 (1.0)</td>
<td>3.6 (1.2)</td>
<td>8.0 (1.6)</td>
<td>10.6 (2.3)</td>
<td>0.6 (0.4)</td>
<td>0.9 (0.5)</td>
<td>4.9 (2.0)</td>
</tr>
<tr>
<td>Pupil diameter</td>
<td>2.2*</td>
<td>-0.5 (0.2)</td>
<td>-1.2 (0.2)</td>
<td>-1.9 (0.2)</td>
<td>-0.6 (0.3)</td>
<td>-0.3 (0.2)</td>
<td>-0.5 (0.1)</td>
<td>-0.2 (0.1)</td>
<td>-0.8 (0.3)</td>
</tr>
</tbody>
</table>

*p < 0.05, df (8, 56).

**p < 0.001, df (8, 56).

*p < 0.05 vs. placebo.

*p < 0.05 vs. 4 mg HM.

*p < 0.05 vs. 8 mg HM.

*p < 0.05 vs. 30 mg MPH.

*p < 0.05 vs. 60 mg MPH.

*p < 0.05 vs. placebo.  
*p < 0.05 vs. 4 mg HM.  
*p < 0.05 vs. 8 mg HM.  
*p < 0.05 vs. 30 mg MPH.  
*p < 0.05 vs. 60 mg MPH.

**p < 0.001 vs. 4 mg HM.  
**p < 0.001 vs. 8 mg HM.  
**p < 0.001 vs. 30 mg MPH.  
**p < 0.001 vs. 60 mg MPH.

### Figure 2

Volunteers were asked to rate how similar each test drug condition was to each drug, as identified by letter code. Data represent mean (S.E.M.) peak change from baseline for visual analog scale scores for

**p < 0.001 versus hydromorphone (4 and 8 mg); # p < 0.05 versus methylenidate (30 and 60 mg).  
# p < 0.05 versus methylphenidate (4 and 8 mg); \# p < 0.05 versus methylenidate (30 and 60 mg).  
* p < 0.05 versus hydromorphone (4 and 8 mg); \* p < 0.05 versus methylenidate (30 and 60 mg).  
\* p < 0.05 versus methylphenidate (4 and 8 mg); \* p < 0.05 versus methylenidate (30 and 60 mg).
reported stimulant-like effects for tramadol, they may have generally disliked such effects despite their history of stimulant abuse, given the ratings of bad effects associated with methylphenidate.

One explanation for this profile of effects (i.e., mild opioid-like effects with reduced abuse liability) is tramadol’s slow onset and lower efficacy at \(\mu\)-opioid receptors compared with a full \(\mu\)-opioid receptor agonist. Tramadol’s parent compound is approximately 6000 times weaker than morphine; however, the M1 metabolite formed after first-pass metabolism has higher affinity for \(\mu\)-opioid receptors compared with the parent form and possesses analgesic activity (Hennies et al., 1988). This profile may increase the potential of tramadol to serve as medication for opioid dependence. In addition, the absence of stimulant-like effects is a further advantage if tramadol is used in a drug-abusing population. These results are in line with several laboratory studies that have supported tramadol as a potential treatment for opioid dependence (Carroll et al., 2006; Lofwall et al., 2007; Lanier et al., 2010) and are also consistent with retrospective studies examining tramadol as a treatment for opioid withdrawal (Tamaskar et al., 2003; Threlkeld et al., 2006). Taken together, these converging lines of work suggest tramadol may be a useful medication for the treatment of patients with low levels of opioid dependence or for the treatment of mild to moderate opioid withdrawal.

This mixed profile of discriminative and subjective effects is consistent with previous human studies. Preston et al. (1991) reported no significant ratings of drug liking or decreased pupil size after doses of tramadol in nondependent opioid-using volunteers. In other studies, tramadol has been shown to engender \(\mu\)-opioid receptor-like effects (Zacny, 2005, Epstein et al., 2006). Preclinical studies have confirmed that effects of tramadol such as analgesia are mediated via both opioid and nonopioid mechanisms (Raffa et al., 1992; Ide et al., 2006). More specifically, Filip et al. (2004) reported that tramadol discrimination in rats was probably mediated by \(\mu\)-opioid receptors, norepinephrine, and possibly serotonergic activity.

Doses of hydromorphone significantly decreased pupil diameter. In the present study, tramadol failed to change pupil diameter significantly. Examination of the time-dependent changes in pupil dilation and constriction after doses of tramadol revealed a delayed, but not significant, pupillary effect compared with hydromorphone (data not shown). Previous reports of tramadol’s pupillary effects have been mixed. Tramadol has been shown to both significantly decrease pupil size (Zacny, 2005; Epstein et al., 2006) and have no effect (Preston et al., 1991) in nondependent opioid volunteers given similar doses as the present study. One possible explanation for the present lack of pupillary effects with tramadol may be differences in metabolism rates related to the polymorphic isoenzyme cytochrome P450 2D6. Individuals who are poor metabolizers of tramadol and express this polymorphism do not show significant miosis after tramadol administration (Fliegert et al., 2005). An alternative explanation may involve the lower efficacy of tramadol compared with full \(\mu\)-opioid receptor agonists.

The present study expanded knowledge on the effects of tramadol in several ways. First, this study included a large dose range of oral tramadol (for example, compared with Preston et al., 1991; but see Epstein et al., 2006 for a report...
on oral doses as high as 700 mg). In addition, whereas previous studies have focused on tramadol’s opioid effects, the present work examined tramadol’s stimulant-like effects along with opioid-like effects. This study also tested tramadol using a human laboratory drug discrimination three-choice procedure. Along with previous reports of tramadol’s utility as a potential treatment medication for opioid withdrawal, the present work brings together in one study assessments of the discriminative stimulus and subjective effects for both opioids and stimulants and the abuse liability of oral tramadol in a drug-experienced nondependent population.

To our knowledge, this study is the first to evaluate the discriminative stimulus effects of tramadol in opioid nondependent humans. We used a novel three-choice discrimination procedure with placebo, hydromorphone, and methylphenidate as training drugs. High doses of tramadol shared discriminative stimulus effects with hydromorphone, but not methylphenidate, suggesting a role for μ-opioid receptors in the acute discriminative stimulus effects of tramadol. Consistent with its lower abuse potential compared with full mu-opioid receptor agonists, tramadol did not increase positive subject ratings associated with reinforcement efficacy, such as drug liking and good effects. However, the highest dose of tramadol increased subject rated scores on a stimulant scale. Taken together these data suggest that μ-opioid receptors are involved in the discriminative stimulus effects of tramadol and stimulant-like effects emerge with higher doses of tramadol, but that this profile of effects is still consistent with a modest abuse liability for tramadol.

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Authorship Contributions
Participated in research design: Bigelow, Lanier, and Strain.
Conducted experiments: Duke, Bigelow, Lanier, and Strain.
Performed data analysis: Duke and Strain.
Wrote or contributed to the writing of the manuscript: Duke, Bigelow, and Strain.
Other: Strain acquired funding for the research.

References