Cebranopadol, a Mixed Opioid Agonist, Reduces Cocaine Self-administration through Nociceptin Opioid and Mu Opioid Receptors

Qianwei Shen¹, Yulin Deng², Roberto Ciccocioppo¹* and Nazzareno Cannella¹

¹ School of Pharmacy, Pharmacology Unit, University of Camerino, Camerino, Italy, ² School of Life Sciences, Beijing Institute of Technology, Beijing, China

Cocaine addiction is a widespread psychiatric condition still waiting for approved efficacious medications. Previous studies suggested that simultaneous activation of nociceptin opioid (NOP) and mu opioid (MOP) receptors could be a successful strategy to treat cocaine addiction, but the paucity of molecules co-activating both receptors with comparable potency has hampered this line of research. Cebranopadol is a non-selective opioid agonist that at nanomolar concentration activates both NOP and MOP receptors and that recently reached phase-III clinical trials for cancer pain treatment. Here, we tested the effect of cebranopadol on cocaine self-administration (SA) in the rat. We found that under a fixed-ratio-5 schedule of reinforcement, cebranopadol (25 and 50 µg/kg) decreased cocaine but not saccharin SA, indicating a specific inhibition of psychostimulant consumption. In addition, cebranopadol (50 µg/kg) decreased the motivation for cocaine as detected by reduction of the break point measured in a progressive-ratio paradigm. Next, we found that cebranopadol retains its effect on cocaine consumption throughout a 7-day chronic treatment, suggesting a lack of tolerance development toward its effect. Finally, we found that only simultaneous blockade of NOP and MOP receptors by concomitant administration of the NOP antagonist SB-612111 (30 mg/kg) and naltrexone (2.5 mg/kg) reversed cebranopadol-induced decrease of cocaine SA, demonstrating that cebranopadol activates both NOP and classical opioid receptors to exert its effect. Our data, together with the fairly advanced clinical development of cebranopadol and its good tolerability profile in humans, indicate that cebranopadol is an appealing candidate for cocaine addiction treatment.

Keywords: cocaine, nociceptin opioid, mu opioid, opioid system, buprenorphine, addiction

INTRODUCTION

Cocaine is a widely abused illicit drug with a worldwide prevalence of 0.4%, accounting for 14–20 million individuals who have used cocaine in their lifetime (1). Cocaine addiction, for which approved medications are still needed, causes physical, psychiatric, socioeconomic, and judicial problems (2), representing therefore a major social burden. The fourth member of the opioid receptor superfamily, the nociceptin opioid (NOP) receptor, is considered a novel potential target...
for medication of cocaine addiction due to its anti-rewarding properties and limited side effects of NOP agonists (3–7).

Nociceptin is the endogenous ligand of the NOP that has been proven to inhibit cocaine-induced conditioned place preference (CPP) in the rat (8) and NOP knockout (KO) mice showed higher cocaine-induced CPP than their wild-type counterparts (9). In addition, nociceptin abolished cocaine-elicited psychomotor sensitization in wild type but not NOP KO mice (10). Noteworthy, intracerebroventricular injection of nociceptin failed to induce either place preference or aversion even at a dose as high as 1,000 ng/rat (11). Consistently, NOP agonists have limited abuse liability (12), thus representing promising tools for the treatment of cocaine addiction. Yet, direct evidence of the effect of NOP agonists on cocaine self-administration (SA) is still lacking.

A growing body of evidence indicates that buprenorphine—an NOP receptor and mu opioid (MOP) receptor agonist and kappa (KOP) and delta (DOP) opioid receptors antagonist—reduces cocaine intake partially via NOP agonism (13). Buprenorphine combined with low, but not high, doses of naltrexone, maintained its potential to prevent cocaine SA in non-dependent rats (14), which could be explained by residual activation of MOP and NOP (13). Clinical trials demonstrated that only a high dose of buprenorphine (≥16 mg/day) was effective in reducing cocaine use, possibly due to simultaneous activation of NOP as MOP was fully occupied at this dose (3, 6). Similar results were obtained by buprenorphine treatment in alcohol-prefering rats in which high doses decreased alcohol intake through NOP activation, but low doses increased intake by activating MOP (15) However, the limitation of buprenorphine as potential cocaine addiction treatment is owed to its too low affinity to NOP compared with MOP (about 50 times lower) (16), and its related abuse liability (17). Therefore, we hypothesize that compounds with high affinities and strong potency to both NOP and classical opioid receptors may have promising potential for treatment of cocaine addiction with reduced side effects.

Cebranopadol is a compound already tested in both completed and ongoing phase-II and phase-III clinical trials for pain treatment (11, 18). Radioligand binding data revealed sub-nanomolar affinity to both rat and human NOP and MOP 20 times higher than human DOP and 3–4 times higher than human KOP receptors. Cebranopadol is a full agonist for human NOP, MOP, and DOP and partial agonist for KOP receptors (19). Here, we employed cebranopadol to test our hypothesis. In a work concomitantly conducted by others, cebranopadol decreased cocaine fixed ratio 1 (FR1) SA in escalated rats and reduced cue-induced reinstatement of cocaine seeking (20). Here we expanded this investigation by looking at the effect of the drug on cocaine reinforcement and motivation by FR5 and progressive-ratio (PR) SA. Moreover, to validate cebranopadol selectivity for cocaine reinforcement, we tested this compound on saccharin SA. Lastly, the NOP antagonist SB-612111 and the non-selective opioid antagonist naltrexone were employed to determine the pharmacological mechanism by which cebranopadol abolishes cocaine consumption. Rats with stable cocaine SA baseline were also treated chronically with cebranopadol or buprenorphine and their vehicle for seven consecutive days to evaluate if the effect of cebranopadol is maintained after repeated treatments.

**METHODS AND MATERIALS**

**Animals**

Male Wistar rats (Charler River, Italy) weighing 270–320 g at the beginning of the experiments were used. Animals’ body weight at the time of tests was very similar with groups with an individual difference not larger than 462.4 ± 8.4. Pairs of rats were housed in a room with artificial 12/12 h light/dark cycle (lights off at 8.00 a.m.), at constant temperature (20–22°C) and humidity (45–55%). Food (4RF18, Mucedola, Settimo Milanese, Italy) and water were provided ad libitum except during session time. All experiments were conducted during the dark phase of the light/dark cycle. Rats were allowed to acclimate to the housing room for 1 week and were handled three times before any experimental manipulation. All procedures were carried out in accordance with the recommendations of the European Community Council Directive and the National Institutes of Health for the Care and Use of Laboratory Animals and were approved by the University of Camerino Internal Ethical Committee for the Laboratory Animal Protection and Use (CEAPA) (Protocol no. 3796/12).

**Drugs**

Cocaine hydrochloride and morphine hydrochloride (Sigma, USA) were dissolved in sterile saline. Saccharin (Sigma, Italy) was dissolved in tap water. Cebranopadol (Biochempartner Co., Ltd., China) for operant tests was diluted to a fine suspension with 5% DMSO and 95% glucose (5%) and administered per os (p.o.) by gavage. We chose p.o. administration to mimic the most common administration route in human. Cebranopadol for place conditioning was dissolved in 10% DMSO + 5% Cremophor EL + 85% saline and administered intraperitoneally (i.p.). For CPP, we used the i.p. route to facilitate the rapid absorption of the drug that is important to achieve the expression of place conditioning. Cebranopadol doses were selected based on previously reported p.o. ED30 value. Previous work demonstrated that pharmacological efficacy of cebranopadol is maintained for least 9 h (19). The selective NOP antagonist SB-612111 was obtained from Tocris (USA) and was dissolved in 1-M H3PO4 in distilled water (1:1), the opioid receptors antagonist naltrexone (Sigma, USA) was dissolved in distilled water. Evidence demonstrates that the doses chosen for the two antagonists result in an appropriate brain bioavailability and significant central nervous system effects (21, 22). Pharmacokinetic data on naltrexone and SB-612111 interactions are not available.

**Catheter Implantation**

Animals were anesthetized by intramuscular injection of 100–150 µL of a solution containing tiletamine cloridrate (58.17 mg/mL) and zolazepam chlorohydrate (57.5 mg/mL). For IV surgery, incisions were made to expose the right jugular vein. A catheter made from micro-renathane tubing (ID = 0.020”, OD = 0.037”) was subcutaneously positioned between the vein and the back. After insertion into the vein, the proximal end of the catheter was anchored to the muscles underlying the vein with surgical silk. The distal end of the catheter was attached to a stainless steel cannula bent at an angle of 90°. The cannula was inserted...
into a support made by dental cement on the back of the animals and was covered with a plastic cap. Immediately after surgery, rats were treated intramuscularly with 200 µL of enrofloxacin (50 mg/mL, Baytril, Germany).

Rats were allowed to recover 1 week before SA training. Catheters were flushed with 100 µL/rat of heparinized saline (20 U/mL) prior to each SA session, when appropriate 0.5 mg/mL of enrofloxacin was added to the flushing solution. Catheter patency was confirmed by intravenous injection of 150 µL/rat of pentothal sodium (25 mg/mL, Intervet, Italy) at the end of experimental procedures.

SA Apparatus
The SA stations consisted of operant conditioning chambers (Med Associate Inc.) enclosed in sound attenuating and ventilated environmental cubicles. Each chamber was equipped with two retractable levers located in the front panel of the chamber. Before the beginning of the session, an infusion pump delivering cocaine was connected by a polyethylene tube to the catheter. The infusion pump was activated by responses on the right (active) lever, while responses on the left (inactive) lever were recorded but did not result in any programmed consequences. Activation of the pump resulted in delivery of 0.1 mL of fluid. An IBM compatible computer controlled the delivery of cocaine solution and recording of the behavioral data.

Effect of Acute Cebranopadol on Cocaine SA
Fixed Ratio
Rats (n = 8) were initially trained for 2-h daily cocaine SA sessions under FR1 schedule of reinforcement for 10 days, then reinforcement schedule was increased to FR5 until stable baseline of responding (<10% variation for three consecutive days) was reached. Following each cocaine infusion (0.25 mg/0.1 mL intravenous), a 20-s time out (TO) period was presented during which responses at the active lever had no programmed consequences. Cebranopadol (0, 25, 50 µg/kg p.o. 1 h before test) was tested in a Latin-square counterbalanced design. At least, a 3-day interval during which cocaine SA baseline was reestablished was allowed between drug tests.

Progressive Ratio
For PR experiments, a new cohort of rats (n = 8) was trained to cocaine SA as described above. After training, the effect of cebranopadol on cocaine SA under PR schedule of reinforcement was tested. During PR sessions, the response requirements necessary to receive a single cocaine dose increased according to the following scale: 5, 11, 18, 26, 35, 45, 56, 68, 82, 98, 116, 136, 158, 182, 208, 236, 268, and 304. PR session stopped after 6 h or if the required ratio was not achieved within 1 h, whichever came first. The breakpoint (BP) corresponding to the last ratio completed was used as a measure of motivation. One hour before test, rats (n = 8) were treated with cebranopadol (0, 25, 50 µg/kg p.o.) in a Latin-square counterbalanced design. Drug treatment was performed every fourth day. Baseline FR5 cocaine SA was reestablished between PR tests.

Effect of Acute Cebranopadol on Saccharin SA
Another group of rats (n = 8) was used to test the effect of cebranopadol on oral saccharin (0.2% w/v) SA. Rats were trained to 30-min daily sessions under an FR1 schedule of reinforcement until stable lever pressing baseline was reached. At this point, the effect of cebranopadol (0, 25, 50 µg/kg p.o.) was tested in a Latin-square counterbalanced design. Drug treatment was performed every fourth day, 1 h before test. Baseline FR1 saccharin SA was reestablished between tests. We used FR1 because we expected, based on our previous experience, that this schedule of reinforcement would produce in 30 min of saccharin SA—a range of lever responses comparable to the initial phase (30–60 min) of cocaine SA in FR5.

Effect of Chronic Treatment with Cebranopadol on Cocaine SA
The effect of repeated cebranopadol administrations was tested to evaluate if the effect of the drug on cocaine intake was maintained following chronic treatment. For seven consecutive days, two groups of rats (n = 7–8/group) with similar cocaine SA baseline were treated with cebranopadol (25 µg/kg p.o.) or its vehicle given 1 h before initiating the 2-h SA sessions under FR5.

Effect of SB-612111 and Naltrexone on Cebranopadol-Induced Inhibition of Cocaine SA
To explore the mechanisms of action of cebranopadol, we tested the effect of the selective NOP antagonist SB-612111 (30 mg/kg i.p.), the MOP/KOP/DOP antagonist naltrexone (Nal, 2.5 mg/kg i.p.) or their combination on cebranopadol-induced inhibition of cocaine SA (23, 24). One hour before session, rats (n = 10) received cebranopadol (50 µg/kg p.o.) or its vehicle alone or in combination with SB-612111, naltrexone or both antagonists together. Rats were subjected to all treatment conditions in a Latin-square design. An interval of at least 3 days during which cocaine SA baseline was reestablished was allowed between drug tests.

Cebranopadol-Induced Place Conditioning
To test whether cebranopadol has rewarding effects per se, we measured the place conditioning induced by the drug in a two compartment apparatus using an unbiased schedule of conditioning. The first day rats (n = 8/group) were allowed to explore the entire apparatus for 15 min (preconditioning) to confirm lack of innate preference for one of the two compartments. From the second day, animals underwent one 25-min conditioning session per day for 6 days. Cebranopadol groups (10 and 50 µg/kg i.p.) received three cebranopadol treatments in one compartment or three vehicle treatments in the opposite compartment every other day. The control group received vehicle injections in both compartments. Treatments were balanced across compartments and days. On the seventh day, drug free animals were allowed to explore the entire apparatus for
15 min. An operator unaware of treatment conditions recorded time spent in each compartment scoring test video clips. A rat was considered to have entered a compartment when all its four paws stepped inside. Rats received treatments 15 min before session.

We treated a group of rats (n = 7) with morphine (5 mg/kg i.p.) with the same conditioning schedule as a positive control to check whether our protocol is suitable to assess opioid-induced place conditioning. Place-conditioning protocol was designed to maximize drug effect based on meta-analysis data (25).

**Statistical Analysis**

The effect of acute cebranopadol on cocaine and saccharin SA, as well as the effect of SB-612111 and naltrexone on cebranopadol-induced reduction of cocaine infusions, was analyzed by one-way analysis of variance (ANOVA) with treatment as a repeated measure. The effect of chronic cebranopadol on cocaine SA was analyzed by two-way ANOVA with one factor between (treatment) and one factor within (time). Inactive-level responses were separately analyzed and used as an additional measure to monitor the specificity of drug effects. The Newman–Keuls test was used for *post hoc* analysis when appropriate.

In the CPP experiment, to verify that our protocol was unbiased, we used the *t*-test for dependent samples to compare the time rats spent in each compartment during the preconditioning test. For the control group, absence of side preference was tested also on the test day.

Preference score was defined as the time spent in the drug-paired compartment minus the time spent in the vehicle-paired compartment. For the control group, which received vehicle in both compartments, the “drug-paired compartment” was randomly assigned before test. We analyzed cebranopadol place conditioning by one-way ANOVA with groups as independent factors (cebranopadol: 0, 10, and 50 µg/kg).

Results were expressed as mean ± SEM and statistical significance was set at *p* < 0.05.

**RESULTS**

**Acute Cebranopadol Selectively Reduced Cocaine SA**

Rats rapidly acquired a stable baseline of cocaine responding under FR5 contingency. ANOVA on the effect of cebranopadol on cocaine SA revealed an overall effect of doses [$F(2, 7) = 43.1069, *p* < 0.00001] (Figure 1A). *Post hoc* analysis revealed a significant decrease in the number of cocaine reinforced responding at both doses of cebranopadol tested (*p* < 0.001). Inactive lever pressing was very low and was not affected by drug treatment [$F(2, 7) = 0.9143, *p* > 0.05].

When the effect of cebranopadol on motivation for cocaine was evaluated under PR contingency, overall ANOVA of the break points found an overall effect of doses [$F(2, 7) = 6.915, *p* < 0.01]. As shown in Figure 1B, *post hoc* tests indicated a significant reduction (*p* < 0.05) of break point following administration of the highest dose (50 µg/kg) of the drug.

**Acute Cebranopadol Selectively Increased Saccharin SA**

Rats rapidly acquired saccharin SA under FR1 contingency. ANOVA on the effect of cebranopadol on saccharin SA revealed an overall effect of doses [$F(2, 7) = 4.859, *p* < 0.05]. As shown in Figure 2, *post hoc* analysis showed a significant increase (*p* < 0.05) in saccharin reinforcement earned following administration of 25 µg/kg of cebranopadol. At a higher dose (50 µg/kg), the drug did not modify saccharin SA. Inactive lever pressing was

![Figure 1](image-url)
Figure 2 | Effect of cebranopadol on saccharin SA. The dose of 25 µg/kg, but not 50 µg/kg, increased the number of saccharin rewards self-administered (top panel). Inactive lever response was not affected by cebranopadol treatments (bottom panel). Data are presented as mean ± SEM. Statistical differences: *p < 0.05 vs. vehicle; #p < 0.05 vs. 25 µg/kg. SA, self-administration.

Figure 3 | Chronic cebranopadol treatment on cocaine self-administration. Two groups of rats with similar baseline of cocaine intake received cebranopadol (25 µg/kg) or its vehicle for seven consecutive sessions. The number of cocaine infusions earned by the cebranopadol-treated group decreased during treatment in respect to baseline, and returned to pretreatment level from the first day posttreatment. In the vehicle-treated group, cocaine self-administration remained stable throughout the experiment. Data are presented as mean ± SEM and show the last 2 days of pretreatment, 7 days of treatment, and 2 days of posttreatment. Statistical differences: *p < 0.05, **p < 0.01, and ****p < 0.0001 vs. the last day pretreatment.

Reduction of Cocaine SA Maintained Following Chronic Administration of Cebranopadol

Analysis of variance of cocaine infusion earned during chronic cebranopadol treatment revealed an overall effect of days [F(10, 13) = 4.74, p < 0.001], while overall effect of groups was not significant [F(1, 13) = 1.00, p = 0.334]. However, there was a significant treatment-by-group interaction [F(10, 130) = 4.51, p < 0.001]. Post hoc tests indicated that there were no significant differences between the first three treatment days and pretreatment SA level, whereas a significant inhibition in the number of cocaine reinforced responding was detected from days 4 to 7 (Figure 3). When treatment was discontinued, cocaine SA returned to pretreatment levels. Analysis of the inactive lever responding found no effect of group [F(1, 13) = 0.48, p > 0.05], days [F(10, 13) = 0.96, p > 0.05], or group-by-day interaction [F(10, 130) = 1.42, p > 0.05]. This result seems in part to contrast with the effect of acute administration of cebranopadol, where the dose of 25 µg/kg reduced cocaine SA. This discrepancy is probably due to the different treatment design, Latin-square rotation of each dose in the acute experiment and between subjects in the chronic one. To confirm that this apparent discrepancy was due to the experimental design, using the t-test analysis for dependent samples, we compared the first drug test session with the last cocaine-training day. As expected, results showed a significant [t(7) = 4.05; p < 0.01] reduction of cocaine SA following cebranopadol.

Effect of SB-612111 and Naltrexone on Cebranopadol-Induced Inhibition of Cocaine SA

To investigate the pharmacological mechanism by which cebranopadol inhibits cocaine intake, we tested the effect of SB-612111, a selective NOP antagonist, and of the MOP/KOP/DOP antagonist naltrexone, on cebranopadol-induced inhibition of cocaine SA. When number of infusions was analyzed, ANOVA revealed an overall effect of treatment [F(4, 8) = 8.147, p < 0.001]. As shown in Figure 4, Newman–Keuls post hoc test indicated, as expected, that 50 µg/kg cebranopadol significantly reduced the number of cocaine reinforced responding (p < 0.001). Neither SB-612111 nor naltrexone alone were able to antagonize cebranopadol effect. In contrast, when SB-612111 and naltrexone were coadministered they completely reversed cebranopadol-induced reduction of cocaine SA (p < 0.01).

Inactive lever responding was very low, and as shown by overall ANOVA, was not affected by treatments [F(4, 8) = 1.152, p > 0.05].

Cebranopadol Not Inducing Place Conditioning

Control rats did not show preference for either of the compartments, demonstrating that place conditioning was performed in unbiased conditions [t(8) = 0.44; p = 0.67]. Morphine-treated rats
showed marked development of CPP with a significant preference score \( t(13) = -2.33; p < 0.05 \), demonstrating that our protocol was suitable to test place preference induced by drugs targeting the opioid system (Figure 5 inset). For cebranopadol, ANOVA revealed no overall effect of groups \( F(2.21) = 1.83; p > 0.05 \), indicating that cebranopadol did not induce significant place-conditioning effects (Figure 5).

**DISCUSSION**

Results showed that cebranopadol significantly reduced cocaine SA under fixed and progressive ratio schedule of reinforcements, indicating reduced motivation for cocaine following drug treatment. Our data are in good agreement with work concomitantly conducted by de Guglielmo et al. (20), showing that cebranopadol reduces cocaine SA in rats after escalation of cocaine intake in 6-h sessions. The effect of cebranopadol was substance-specific because when we tested cebranopadol on saccharin SA, we found a slight increase in lever pressing at the lowest dose and no changes at the highest dose. A similar finding was described by de Guglielmo et al. (20) who attributed that using sweet condensed milk as reinforcer demonstrated the same behavioral pattern. The tendency of cebranopadol to increase the consumption of sweet solutions is consistent with its ability to activate MOP receptors. In fact, sweet intake is markedly enhanced following activation of this opioid receptor by morphine, DAMGO or other selective agonists (26–29). In addition, the fact that cebranopadol does not decrease, or even increases, responses for natural reinforcers demonstrates that the effect on cocaine was not secondary to sedation or to general disruption of motor activity. In the chronic study, cebranopadol slightly decreased cocaine SA at the beginning of the treatment; the effect progressively increased following repeated administrations and became significant after 4 days. This finding supports the feasibility of chronic drug administration as a therapeutic strategy for cocaine addiction. We also demonstrated that the inhibitory effect of cebranopadol on cocaine SA is blocked by coadministration of SB-612111 and naltrexone, while compounds alone did not affect the efficacy of cebranopadol. This indicates that the effect of cebranopadol on cocaine SA is mediated by two possibly independent mechanisms involving NOP and classical opioid receptors. Only simultaneous blockade of these two pathways warrants full inhibition of cebranopadol effects on cocaine. Finally, in a place-conditioning test, we found that in contrast with morphine, cebranopadol did not elicit significant expression of place preference, although a trend to an effect was detected (30, 31). Additional studies (i.e., drug discrimination, operant intravenous SA) will have to be carried out to thoroughly evaluate the abuse potential of this compound.

Cebranopadol is a high-affinity agonist at both MOP and NOP receptors and is almost equipotent in activating both receptors. At higher concentrations, it also activates KOP and DOP receptors (19). The pharmacological profile of cebranopadol is partially mimicked by buprenorphine, a well-characterized compound that acts as a high-affinity partial agonist at MOP and low-affinity NOP agonist, but is an antagonist at KOP.
and DOP receptors (32). Like cebranopadol, buprenorphine reduced cocaine intake in the rat (13, 33, 34). This effect was confirmed also in rhesus monkeys (35) and in heroin-dependent patients co-abusing cocaine (36). Importantly, clinical data demonstrated that reduction of cocaine consumption by buprenorphine is independent of the effect on heroin intake and occurs only at doses higher than 24 mg/kg (36). This dose is substantially higher (>16 mg/day) compared with those needed to fully occupy MOP, and most likely KOP receptors, as the drug shows equivalent affinity for both receptors (32, 36–39). Based on these receptor affinity profiles, we argued that the inhibitory effect of buprenorphine on cocaine consumption was mediated by activation of NOP, which occurs at substantially higher drug doses (15, 32, 40, 41). Contrary to our expectation, we found that selective blockade of NOP did not prevent the effect of buprenorphine. However, the effect was fully prevented if NOP and MOP receptors were simultaneously blocked (13). Based on this observation, we concluded that buprenorphine reduces cocaine consumption with mechanisms involving both NOP and MOP receptors. Here we demonstrated that, cebranopadol despite acting as a KOP and DOP agonist, whereas buprenorphine is an antagonist, also attenuated cocaine intake. Based on these data, it is tempting to speculate that KOP-dependent mechanisms might not be relevant in mediating the effects of these drugs on cocaine consumption.

In addition to NOP, MOP, and KOP receptors, although at higher concentrations, cebranopadol binds also to DOP (19). Based on present results, we cannot fully exclude the possible involvement of this opioid receptor subtype in mediating the effect of cebranopadol on cocaine. However, this hypothesis is discouraged by the lower affinity of this compound for DOP and by the fact that psychostimulant consumption is blocked by DOP antagonism (42). Cebranopadol is at phase-III stage of development for the treatment of pain, and toxicological data demonstrated that it is well tolerated in humans (11). Its safety profile was confirmed by studies in rodents showing that, contrary to classic opioid analgesics, cebranopadol did not produce respiratory depression and impairment of motor coordination at pharmacological effective doses (19). The opportunity to test this compound in cocaine addicts is therefore relatively close.

Theoretically, the ability of cebranopadol to act as KOP agonist could rise concerns on the use of this compound in drug addicts. In fact, previous studies have shown that activation of KOP receptors is associated with dysphoria, exacerbation of anxiety, and enhanced motivation to take drugs of abuse (43–50). For instance, it has been shown that activation of KOP recruiting stress response mechanisms, is able to reinstate extinguished CPP for cocaine in the rat (51, 52). Moreover, in monkeys trained to self-administer cocaine, the administration of selective KOP agonists facilitates reinstatement of drug seeking and enhances preference for cocaine if animals are trained to a concurrent-choice schedule—choice between food or cocaine (45, 53). Here we showed that in the place-conditioning test cebranopadol did not evoke aversion and produced a trend to a preference probably mediated by its ability to activate MOP receptors. Hence, unlike selective KOP agonist, cebranopadol appeared to be devoid of negative affective properties in the rat (54–58).

Attenuation of drug effect following repeated administration is always a concern when chronic treatment is required in the clinic setting. Interestingly, an earlier study exploring the analgesic effect of cebranopadol revealed that the anti-allodynic effect of the drug is maintained for substantially longer time compared with chronic morphine, which may reflect reduced propensity to produce pharmacological tolerance (19). Here, we further explored this phenomenon by investigating the effect of 1-week treatment with cebranopadol on cocaine SA. Results demonstrated that the effect improved along the 7 days of drug administration, with an important trend toward an increase at the end of treatment. Classically, repeated administrations of opioid agonists lead to rapid development of tolerance; using cebranopadol here this effect did not occur (59, 60). In contrast with other opioidergic agents, buprenorphine shows reduced propensity to evoke tolerance. For instance, reduction of cocaine SA in monkeys was fully retained for a period as long as 120 days (35). Based on these observations, it is tempting to speculate that the pharmacological profile of cebranopadol—mimicking buprenorphine properties, but showing even higher safety and possibly lower abuse liability profile—may be suitable for a chronic therapy in cocaine addicts.

The reasons why molecules with mixed MOP/NOP agonist profile have low tendency to produce tolerance remain unclear at present and, to some extent, contrast with older data, suggesting that activation of central nociceptin system facilitates analgesic tolerance elicited by chronic morphine (61). To explain this phenomenon, further studies should analyze the molecular mechanisms and, possibly, the intracellular signaling pathways that are engaged when MOP and NOP receptors, which are largely co-expressed in the brain, are co-activated (62). Interestingly, cebranopadol differently from other agonists shows low ability to stimulate the opioid receptors-β-arrestin pathway that is typically involved in receptor internalization or desensitization (63, 64).

Altogether these results demonstrate that cebranopadol, in addition to its already known analgesic profile, shows efficacy in attenuating cocaine consumption. This effect is specific and does not generalize toward other ingesta (i.e., saccharin). Moreover, present findings demonstrate that NOP and classical opioid receptor pathways are both involved in mediating these actions of cebranopadol on cocaine. One limitation of the study is that cebranopadol was tested against a fixed dose of cocaine. This does not allow to draw conclusions on whether the effect of this opioid panagonist depends upon its ability to shift the classically U-shaped dose response curve of cocaine toward the right (reflecting attenuation of reward) or toward the left (reflecting potentiation of reward). In recent years, despite significant efforts made to develop new medications for the treatment of cocaine addiction, very little is available for clinical use. The fairly advanced clinical development of cebranopadol makes this compound an ideal candidate for an immediate clinical investigation in cocaine-dependent patients.
ETHICS STATEMENT

All procedures were carried out in accordance with the European Community Council directive and the National Institutes of Health guidelines for the care and use of laboratory animals.

AUTHOR CONTRIBUTIONS

QS performed self-administration experiments, analyzed data, and wrote the manuscript. YD contributed to design of the experiment and provided a critical reading. RC wrote the manuscript. NC run place conditioning, analyzed data, and wrote the manuscript.

ACKNOWLEDGMENTS

Many thanks to Alfredo Fiorelli, Rina Righi, and all the staff of the animal facility for their technical assistance.

FUNDING

This study was supported by the grant of the University of Camerino and NIH/NIAAA (grant number AA014351 to RC).
Cebranopadol Reduces Cocaine Intake

30. Martell BA, O'Connor PG, Kerns RD, Becker WC, Morales KH, Kosten TR, et al. Systematic review: opioid treatment for chronic back pain: prevalence, efficacy, utilization and research needs. *Ann Intern Med* (2007) 146:116–27. doi:10.7326/0003-4814-146-2-20070116-00006

31. Sehgal N, Manchikanti L, Smith HS. Prescription opioid abuse in chronic pain: a review of opioid abuse predictors and strategies to curb opioid abuse. *Pain Physician* (2012) 15:ES67–92.

32. Huang P, Kehner GB, Cowan A, Liu-Chen LY. Comparison of pharmacological activities of buprenorphine and norbuprenorphine: norbuprenorphine is a potent opioid agonist. *J Pharmacol Exp Ther* (2001) 297:688–95.

33. Carroll ME, Lac ST. Effects of buprenorphine on self-administration of cocaine and a nondrug reinforcer in rats. *Psychopharmacology* (1992) 106:439–46. doi:10.1007/BF02448122

34. Sorge RE, Stewart J. The effects of chronic buprenorphine on intake of heroin and cocaine in rats and its effects on nucleus accumbens dopamine levels during self-administration. *Psychopharmacology* (2006) 186:28–41. doi:10.1007/s00213-006-0485-1

35. Mello NK, Lukas SE, Kamien JB, Mendelson JH, Drieje J, Cone EJ. The effects of chronic buprenorphine treatment on cocaine and food self-administration by rhesus-monkeys. *J Pharmacol Exp Ther* (1992) 260:1185–93.

36. Montoya ID, Gorelick DA, Preston KL, Schroeder JR, Umbricht A, Cheskin IJ, et al. Randomized trial of buprenorphine for treatment of concurrent opiate and cocaine dependence. *Clin Pharmacol Ther* (2004) 75:43–48. doi:10.1016/j.clpt.2003.08.034

37. Kosten TR, Rosen MI, Schottenfeld R, Ziedonis D. Buprenorphine for cocaine and opiate dependence. *Psychopharmacology* (2012) 215:ES67–92. doi:10.1007/s00213-011-2502-0

38. Kosten TR, Rosen MI, Schottenfeld R, Ziedonis D. Buprenorphine and cocaine self-administration and naltrindole, a delta-selective opioid antagonist. *J Pharmacol Exp Ther* (2004) 309:381–4.

39. Redila VA, Chavkin C. Stress-induced reinstatement of cocaine seeking is mediated by the kappa opioid system. *Psychopharmacology* (2007) 232:525–33. doi:10.1007/s00213-007-12548

40. Redila VA, Chavkin C. Stress-induced reinstatement of cocaine seeking is mediated by the kappa opioid system. *Psychopharmacology* (2008) 200:118–26. doi:10.1007/s00213-008-1122-y

41. Lahiri S, Srivastava A, Carlezon WA. Role of kappa-opioid receptors in stress and anxiety-related behavior. *Psychopharmacology* (2013) 229:435–52. doi:10.1007/s00213-013-3195-5

42. Reid LD, Glick SD, Menkens KA, French ED, Bilsky EJ, Porreca F. Cocaine and antidepressant-like effects in rats. *J Pharmacol Exp Ther* (1995) 273:1409–12. doi:10.1097/00001756-199507100-00012

43. Reiersen PM, Howard JL, Shelton KL, Carroll FI. Differential effects of the novel kappa opioid receptor antagonist, JDTic, on reinstatement of cocaine-seeking induced by footshock stressors vs cocaine primes and its antidepressant-like effects in rats. *Psychopharmacology* (2005) 183:118–26. doi:10.1007/s00213-005-0167-4

44. McLaughlin JP, Land BB, Li S, Pintar JE, Chavkin C. Prior activation of kappa opioid receptors by US0, 488 mimics repeated forced swim stress to potentiate cocaine place preference conditioning. *Neuropsychopharmacology* (2006) 31:787–94. doi:10.1038/npj.2006.00550.x

45. Valdez GR, Platt DM, Bowlett JK, Riedi-Betschen D, Speelman RD. \(\kappa\) Agonist-induced reinstatement of cocaine seeking in squirrel monkeys: evidence for a role in opioid and stress-related mechanisms. *J Pharmacol Exp Ther* (2007) 323:252–33. doi:10.1124/jpet.107.12548

46. Funk D, Coen K, Lé A. The role of kappa opioid receptors in stress-induced reinstatement of alcohol seeking in rats. *Brain Behav* (2014) 4:366–76. doi:10.1002/brb3.222

47. Schindler AG, Li S, Chavkin C. Behavioral stress may increase the rewarding valence of cocaine-associated cues through a dynorphin/\(\kappa\)/opioid receptor-mediated mechanism without affecting associative learning or memory retrieval mechanisms. *Neuropsychopharmacology* (2010) 35:1932–42. doi:10.1038/npp.2010.67

48. Al-Hanawi R, Mccall JG, Foshage AM, Bruchas MR. Locus coeruleus kappa-opioid receptors modulate reinstatement of cocaine place preference through a noradrenergic mechanism. *Neuropsychopharmacology* (2013) 38:2484–97. doi:10.1038/npp.2013.151

49. Negus SS. Effects of the kappa opioid agonist U50, 488 and the kappa opioid antagonist nornalbinaltorphimine on choice between cocaine and food in rhesus monkeys. *Psychopharmacology* (2004) 176:204–13. doi:10.1007/s00213-004-1878-7

50. Al-Hanawi R, Mccall JG, Foshage AM, Bruchas MR. Locus coeruleus kappa-opioid receptors modulate reinstatement of cocaine place preference through a noradrenergic mechanism. *Neuropsychopharmacology* (2013) 38:2484–97. doi:10.1038/npp.2013.151

51. Schindler AG, Li S, Chavkin C. Behavioral stress may increase the rewarding valence of cocaine-associated cues through a dynorphin/\(\kappa\)/opioid receptor-mediated mechanism without affecting associative learning or memory retrieval mechanisms. *Neuropsychopharmacology* (2010) 35:1932–42. doi:10.1038/npp.2010.67

52. Ehrich JM, Messinger DI, Knakal CR, Kuhr JR, Schattauer SS, Bruchas MR, et al. Kappa opioid receptor-induced aversion requires p38 MAPK activation in VTA dopamine neurons. *J Neurosci* (2015) 35:12917–31. doi:10.1523/JNEUROSCI.00523-15.2015

53. Freye E, Latala L. Development of opioid tolerance—molecular mechanisms and clinical consequences. *Anesthesiol Intensivmed Notfallmed Schmerzther* (2003) 38:14–26. doi:10.1002/smi.2003.36558

54. Jamison RN, Mao JR. Opioid analgesics. *Mayo Clin Proc* (2015) 90:957–68. doi:10.1016/j.mayocp.2015.04.010

55. Tian JH, Zhang W, Fang Y, Xu W, Grandy DK, Han JS. Endogenous orphanin FQ: evidence for a role in the modulation of electroacupuncture analgesia and the development of tolerance to analgesia produced by morphine and electroacupuncture. *Br J Pharmacol* (1998) 124:21–6. doi:10.1038/sj.bjp.0701788

56. Otawa A, Brunori G, Mercatelli D, Wu J, Cippitelli A, Zou B, et al. Knock-in mice with NOP-eGFP receptors identify receptor cellular and regional localization. *J Neurosci* (2015) 35:11682–93. doi:10.1523/JNEUROSCI.5122-14.2015

57. Rizzi A, Cerles MC, Ruzza C, Malafacini D, Ferrari F, Bianco S, et al. Pharmacological characterization of cebranopadol a novel analgesic acting as mixed Nociceptin/Orphanin FQ and opioid receptor agonist. *Pharmacol Res Perspect* (2016) 4:e00247. doi:10.1016/j.prrep.2016.03.005

58. Rajagopal S, Shenoy SK. GPCR desensitization: acute and prolonged phases. *Cell Signal* (2017). doi:10.1016/j.cellsig.2017.01.024

**Conflict of Interest Statement:** The authors declare that they have no competing or conflicting interest and no financial disclosures to include.

The reviewers LR-K and MS and handling editor declared their shared affiliation.

Copyright © 2017 Shen, Deng, Ciccocioppo and Cannella. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) or licensor are credited and that the original publication is not modified. No use, distribution or reproduction is permitted which does not comply with these terms.