A bifunctional nociceptin and mu opioid receptor agonist is analgesic without opioid side effects in nonhuman primates

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Misuse of prescription opioids, opioid addiction, and overdose underscore the urgent need for developing addiction-free effective medications for treating severe pain. Mu opioid peptide (MOP) receptor agonists provide very effective pain relief. However, severe side effects limit their use in the clinical setting. Agonists of the nociceptin/orphanin FQ peptide (NOP) receptor have been shown to modulate the antinociceptive and reinforcing effects of MOP agonists. We report the discovery and development of a bifunctional NOP/MOP receptor agonist, AT-121, which has partial agonist activity at both NOP and MOP receptors. AT-121 suppressed oxycodone’s reinforcing effects and exerted morphine-like analgesic effects in nonhuman primates. AT-121 treatment did not induce side effects commonly associated with opioids, such as respiratory depression, abuse potential, opioid-induced hyperalgesia, and physical dependence. Our results in nonhuman primates suggest that bifunctional NOP/MOP agonists with the appropriate balance of NOP and MOP agonist activity may provide a dual therapeutic action for safe and effective pain relief and treating prescription opioid abuse.

INTRODUCTION

The recent marked increase in misuse and abuse of prescription opioids and the epidemic of opioid overdose mortality have greatly affected our society (1–3). Although mu opioid peptide (MOP) receptor agonists remain the most effective and widely used analgesics, their abuse liability and possibility of causing respiratory arrest have contributed to escalating medical and economic burdens and fueled the opioid crisis in the global community (4, 5).

Multiple scientific strategies have been tested to develop safe and effective medications for pain and opioid addiction (6, 7). As finely articulated by Volkow and Collins (7), the crisis surrounding this dynamic epidemic of opioid use disorders falls into three critical needs that can be addressed with novel scientific approaches for (i) reversing opioid overdose, (ii) treating opioid addiction, and (iii) finding safe, new approaches for effective pain relief without the side effects of opioids. The lack of alternative analogs that offer opioid-like pain relief without associated adverse effects is a major contributor to the current opioid crisis (2, 7, 8).

Kappa opioid peptide (KOP) and delta opioid peptide (DOP) receptor agonist–based analgesics, although effective, have different adverse effects, such as dysphoria and convulsions, and have a narrow therapeutic window (6, 9). Current efforts to modulate MOP agonist–induced adverse effects have centered on MOP agonists that favor intracellular G protein (guanine nucleotide–binding protein) over arrestin signaling, based on the rationale that arrestin signaling is associated with undesirable effects of opioids (10, 11). TRV-130 (oliceridine) was developed as a G protein–biased MOP agonist with potency and efficacy similar to morphine, but with fewer side effects (12). However, repeated administration of TRV-130 retained abuse liability in a rodent model (13). New approaches are still needed to obtain effective opioid-like analgesic efficacy without MOP agonist–associated side effects.

Agonists targeted to the fourth opioid receptor subtype, the nociceptin/orphanin FQ (N/OFQ) peptide (NOP) receptor, have been shown to modulate the pain-relieving and addictive properties of MOP agonists (14–17). The NOP receptor and its endogenous ligand N/OFQ are expressed in the same neuronal circuitry as the other opioid receptors and generally inhibit neuronal transmission (16, 18). In nonhuman primates, selective NOP receptor agonists produce antinociception and antihypersensitivity at the spinal, supraspinal, and systemic levels (19–21). NOP agonists do not produce respiratory depression and reinforcing effects (abuse potential) in primates, and they enhance MOP agonist–induced analgesia synergistically at spinal and systemic levels (21–23). Given the inhibitory effects on dopamine release and neurotransmission by the N/OFQ-NOP receptor system (14, 24, 25), we hypothesized that bifunctional NOP/MOP agonists may be a viable approach to develop nonaddictive analogs (9, 26–28).

Several bifunctional NOP/MOP agonists have been discovered in the past few years (27–30). Although these compounds, such as AT-201 and AT-212, had antinociceptive effects in rodent models of acute and chronic pain, they exerted rewarding effects (27, 28, 31). In addition, some aspects of the antinociceptive functional profiles of NOP/MOP agonists in rodents did not generalize to rhesus monkeys (for instance, the opposing effects of NOP antagonists) (22, 30, 32, 33). The balance of NOP versus MOP agonist efficacy appears to be a key factor for developing a suitable nonaddictive analgesic devoid of MOP agonist liabilities.

Differences between rodents and primates in the antinociceptive and pharmacological profiles of NOP- and MOP-targeted compounds have been extensively documented (22, 34, 35), and there is ample evidence of neuroanatomic, neurochemical, and neuropharmacological similarities between primates and humans (9, 36, 37). Thus, we conducted in vivo pharmacological studies in monkeys as a translational platform to determine the effects of a dual NOP/MOP agonist on a variety of behavioral and physiological responses.
Using receptor structure–guided drug design and rational structure–activity relationship (SAR) exploration, we developed a bifunctional NOP/MOP agonist, AT-121, that produced potent analgesic effects without inducing opioid-associated hyperalgesia or physical dependence in nonhuman primates. In addition, AT-121 attenuated the reinforcing effects of oxycodone as measured by drug self-administration in primates. Our results suggest that AT-121 might be an effective safer alternative to opioids for pain relief.

RESULTS
Bifunctional NOP/MOP receptor agonists were developed by structure-guided drug design and optimization starting from a NOP receptor–selective chemical scaffold

To design a bifunctional ligand with the desired spectrum of binding affinity and functional efficacy at the NOP and MOP receptors within a single chemical scaffold, we began with structure-activity optimization of a new chemical lead from our NOP ligand library, 1, which showed modest binding affinity for the NOP receptor but about 10-fold lower affinity for the MOP receptor (Fig. 1A and Table 1). This early lead had low partial agonist efficacy at the NOP receptor in the guanosine 5′-O-[γ-thio]triphosphate (GTPγS) functional assay and no agonist efficacy at MOP (Table 1). To improve binding affinity and bifunctional NOP/MOP efficacy, we used structure-guided SAR optimization, starting by docking 1 in the active-state NOP receptor structure we previously developed (38). We hypothesized that substituting cationic groups for the nonionic cyanomethyl group on the isoquinolinone nitrogen of 1 would improve NOP binding affinity by enabling interactions with the anionic amino acids in the extracellular loop 2 (EL2) of the NOP receptor toward which the ligand 1 was oriented. We synthesized analogs 2 to 6 (Fig. 1) to explore this SAR hypothesis. All analogs were synthesized from

Fig. 1. Structure-based design and optimization of bifunctional NOP-MOP ligands. (A) Top: Docking and binding interactions of compound 2 (shown in green stick representation) into the active-state NOP homology model (38), shown in cartoon representation (pink). Amino acids in a 4 Å radius around the ligand are shown as pink sticks and are labeled. Polar interactions of the 2-aminoethyl moiety of 2 with Glu194 and Glu199 in the EL2 of the receptor are shown as green dotted lines. Bottom: Structures of the isoquinolinolone-based bifunctional ligands and their binding affinities at NOP and MOP. (B) Docking and binding interactions of compound 5 (AT-121) (shown in cyan) in the active-state NOP structure (pink). Note the interactions of the nitrogens of the sulfamide group with the Glu194 and Glu199 of the EL2 loop of NOP. Gln280, the nonconserved amino acid between NOP and the opioid receptors [His297 in MOP, shown as green sticks in (C)], does not interact with the ligand. (C) Docking and binding interactions of bifunctional compound 5 in the MOP active-state crystal structure (73), shown in cyan cartoon representation. The interacting amino acids are shown as cyan stick representations. The lipophilic isopropylcyclohexyl group binds to a nonpolar pocket lined with Met131, His297 (green sticks), and Trp293 in the MOP binding pocket.
interacted with conserved amino acids in the NOP and also at NOP receptors (Table 1). Docking the bifunctional NOP/MOP ligand (AT-121) (Fig. 1A and fig. S1 and described in Supplementary Materials and Methods. Values are means ± SEM of three independent experiments run in triplicate. NT, not tested (compounds with binding affinity K_i >100 nM were not tested in the functional assay); DPDPE, (d-penicillamine d-penicillamine)-enkephalin.

Further ligand optimization by introducing the larger polar ethylsulfamide group onto the isoquinolinone nitrogen significantly increased MOP receptor affinity (K_i = 16.49 nM) compared to analogs 2 and 3, maintaining high affinity at the NOP receptor (K_i = 3.67 nM), resulting in the bifunctional NOP/MOP ligand, 5 (AT-121) (Fig. 1A and Table 1). The SAR showed that the ethylsulfamide group was important for high affinity at both receptors, because the closely related N-ethylmethanesulfonamide analog 6 had much lower affinity at MOP and also at NOP receptors (Table 1). Docking the bifunctional NOP/MOP ligand 5 into the active-state NOP receptor (Fig. 1B) and MOP receptor (Fig. 1C) showed that 5 interacted with conserved amino acids in both receptors (NOP Asp130 3,32 and MOP Asp147 3,32, NOP Met132 3,36 and MOP Met151 3,36, and NOP Trp276 4,39 and MOP Trp293 4,39).

AT-121 is a bifunctional NOP/MOP partial agonist

Functional efficacy at both receptors was determined for all compounds using a guanosine-5′-O-(3-[35S]thio) triphosphate ([35S]GTPγS) binding assay in cell membranes from Chinese hamster ovary cells (CHO-hNOP and CHO-hMOP) stably expressing the human NOP and MOP receptors as we have reported previously (40–42). Concentration-response curves were run in parallel with the prototypical full agonists N/OFQ for the NOP receptor and DAMGO ([d-Ala^2, N-MePhe^4, Gly-ol]-enkephalin) for the MOP receptor as reference agonists for the assay. Functional efficacies of the test compounds are reported as % stimulation of [35S]GTPγS binding compared to a full response (100% stimulation) by the reference full agonists (Table 1). The potencies of the functional response are reported as EC_50 (half maximal effective concentration) values determined by nonlinear regression analysis and are presented in Table 1.
The initial lead compound 1 showed low efficacy and potency at the NOP receptor and no agonist efficacy at the MOP receptor in the GTPγS assay. Analogs 2 to 4 had nanomolar binding affinity at NOP but poor to low agonist potencies and efficacies at NOP and MOP in the GTPγS assay, and did not have a desirable bifunctional NOP/MOP agonist profile. Analog 5 (AT-121), on the other hand, showed high potency (EC₅₀ = 35 nM) and partial agonist efficacy at the NOP receptor (Table 1) relative to N/OFQ, and high potency (EC₅₀ = 20 nM) and partial agonist activity at the MOP receptor relative to DAMGO. AT-121 showed significantly lower binding affinity at the DOP or KOP receptors (Kᵢ > 100 nM) (Table 1). Overall, these findings indicate that AT-121 has bifunctional NOP/MOP agonist activity, showing high potency and partial agonist efficacy at both NOP and MOP receptors. We selected AT-121 as a prototype to explore our hypothesis that bifunctional NOP/MOP agonists having an appropriate balance of NOP and MOP agonist efficacies may be a viable approach to develop analgesics with reduced side effects. Characterization of the pharmacokinetics and brain permeability of AT-121 in rats and assessment of stability of AT-121 in monkey plasma (shown in figs. S2 and S3 and tables S1 to S3, respectively) showed that AT-121 had suitable pharmacokinetic properties for further in vivo studies.

AT-121 produces potent antinociceptive and antiallodynic effects

MOP receptor agonists change nociceptive thresholds and produce antinociception in nonhuman primates and humans (43–45). We used the warm water tail-withdrawal assay in primates to determine the functional efficacy of AT-121 for changing the nociceptive threshold. Subjects’ baselines values are presented in table S4. After subcutaneous administration (0.003 to 0.03 mg/kg), AT-121 produced antinociceptive effects against an acute noxious stimulus, 50°C water, in a dose-dependent (F₃,₉ = 163.9; P < 0.05) and time-dependent (F₄,₁₂ = 114; P < 0.05) manner (Fig. 2A). The minimum effective dose of AT-121 to produce full antinociception was 0.03 mg/kg. This dose had a 3-hour duration of action, which subsided by 6 hours (Fig. 2A). To determine the effects of AT-121 on pain sensitivity, we used a clinically relevant model, capsaicin-induced allodynia, which is widely applied to evaluate analgesics in humans (46, 47). Systemic AT-121 exerted a dose-dependent inhibitory effect on capsaicin-induced thermal allodynia in 46°C water (F = 22.5; P < 0.05) (Fig. 2B).

Next, we conducted antagonist studies using the NOP receptor antagonist J-113397 and the MOP receptor–selective dose of the opioid receptor antagonist naltrexone (19, 23). Pretreatment with

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**Fig. 2. Effects of systemic administration of AT-121 on modulating sensory processing in nonhuman primates.** (A) Effect of AT-121 administration on acute noxious stimulus, 50°C water. (B) Bar graph showing the effect of AT-121 on antihypersensitivity against capsaicin-induced allodynia in 46°C water. (C) Effects of NOP receptor antagonist J-113397 (0.1 mg/kg) and MOP receptor antagonist naltrexone (0.03 mg/kg) on AT-121–induced antinociception. (D) Comparison of antinociceptive potency of AT-121 and morphine. (E) Comparison of itch-scratching responses elicited by AT-121 (0.03 mg/kg) and morphine (1 mg/kg) at antinociceptive doses. Each data point represents mean ± SEM (n = 4). All compounds were delivered subcutaneously. Data were analyzed by two-way analysis of variance (ANOVA) with repeated measures (A) or one-way ANOVA with repeated measures (B and E), followed by Bonferroni’s multiple comparisons test. *P < 0.05, significantly different from vehicle condition from the first time point to the corresponding time point.
J-113397 (0.1 mg/kg) or naltrexone (0.03 mg/kg) produced small degrees (two- to threefold dose ratio) of rightward shift of the dose-response curve for AT-121–induced antinociception. Furthermore, combined pretreatment with both J-113397 and naltrexone provided a larger rightward shift (30-fold) of AT-121’s dose-response curve (Fig. 2C). These findings indicate that both NOP and MOP receptors contributed to antinoceceptive effects of AT-121. Analysis of the dose-response curves showed that systemic AT-121 was more potent than morphine [median effective dose (ED₅₀) = 0.01 mg/kg versus 1 mg/kg] (Fig. 2D). To examine whether AT-121 elicited an itch sensation, we compared its effects with morphine, which has been shown to elicit scratching responses in monkeys (48). AT-121 (0.03 mg/kg) did not significantly increase scratching responses (P > 0.9) (Fig. 2E). In contrast, morphine (1 mg/kg) increased the number of scratches in the same subjects (P < 0.05). Together, the data show that systemic AT-121 had a promising analgesic profile in primates distinct from typical mu opioid analgesics such as morphine.

**AT-121 lacks reinforcing effects and attenuates reinforcing effects of oxycodone**

To examine and compare the reinforcing strengths of compounds, we used a progressive-ratio schedule of reinforcement, which is commonly used for evaluating abuse potential (49, 50). Monkeys were allowed to self-administer single doses of cocaine and remifentanil and various doses of AT-121 and oxycodone (0.3 to 10 μg/kg per injection). Substitution of saline between test compounds resulted in a low number of reinforcers (three or fewer injections) (Fig. 3A). The reinforcing strength of remifentanil (0.3 μg/kg per injection) was similar to that of cocaine (0.03 mg/kg per injection) (Fig. 3A). There was no significant difference between the reinforcing strengths of saline and AT-121 (F = 1.2; P > 0.1) (Fig. 3A). In contrast, oxycodone produced reinforcing effects dose-dependently (F = 9.7; P < 0.05) (Fig. 3B). In addition, similar to buprenorphine (0.1 mg/kg) and naltrexone (0.01 mg/kg), pretreatment with AT-121 (0.03 mg/kg) effectively attenuated the reinforcing effects of oxycodone (F = 25.5; P < 0.05) (Fig. 3C). This attenuation by AT-121 was selective against oxycodone, because AT-121 did not attenuate the reinforcing effects of food pellets (Fig. 3D). Collectively, AT-121 was devoid of reinforcing effects (abuse potential) and attenuated reinforcing effects of oxycodone, suggesting its therapeutic potential for opioid addiction.

**Higher doses of AT-121 do not compromise physiologic functions**

To characterize the appropriate dosage of AT-121, we measured physiologic parameters in monkeys implanted with radiotelemetric transmitters able to monitor real-time respiratory and cardiovascular activities (32). A systemic dose (0.03 mg/kg) of AT-121 that produced full antinociception did not affect respiratory function (respiration rate and minute volume), cardiovascular activity (heart rate and blood pressure), or body temperature of monkeys (Fig. 4). At a dose (0.3 mg/kg) about 10 to 30 times higher than its antinoceceptive doses (0.01 to 0.03 mg/kg), AT-121 also did not significantly change any physiologic parameters (all F values = 1 to 4, P > 0.1) during the first 40 min and the 6-hour period (Fig. 4 and fig. S4). In contrast, heroin (1 mg/kg intramuscularly) rapidly (<20 min) caused respiratory depression in a single monkey, which required naltrexone (0.01 mg/kg) to reverse it (Fig. 4B). For safety reasons, this particular experiment was only conducted in one monkey. These findings illustrate that unlike heroin, AT-121 did not cause respiratory and cardiovascular concerns in primates. Moreover, animals did not show sedation or motor impairment when they were tested with these doses of AT-121. Animals were alert and did not display eye closure while being tested with these doses of AT-121.

**Repeated exposure to AT-121 does not cause physical dependence**

After repeated exposure to mu opioid analgesics, primates and humans quickly develop physical dependence (35, 51, 52). In such conditions, antagonist-precipitated withdrawal is indicated as changes in respiratory and cardiovascular parameters in primates (32, 35). Compared to the vehicle-treated condition (0.1 ml/kg), naltrexone (0.01 mg/kg) precipitated withdrawal signs on day 4 in morphine-treated monkeys (two injections of 1.8 mg/kg daily for 3 days). These withdrawal signs included increases in respiratory rate (F₁,₃ = 49;
After short-term exposure to morphine (two injections of 1.8 mg/kg–m), development of opioid-induced hyperalgesia and tolerance (53–55). After repeated exposure to mu opioid analgesics, animals and humans develop opioid-induced hyperalgesia and tolerance (53–55).

Repeated exposure to AT-121 does not cause opioid-induced hyperalgesia

After repeated exposure to mu opioid analgesics, animals and humans develop opioid-induced hyperalgesia and tolerance (53–55). After short-term exposure to morphine (two injections of 1.8 mg/kg for 1 day), monkeys had a lower threshold to capsaicin-induced hypersensitivity on day 2 ($F_{1,5} = 10.2; P < 0.05$), shown as decreased tail-withdrawal latencies in 46°C water (Fig. 6A). In contrast, AT-121–treated monkeys (two injections of 0.03 mg/kg for 1 day) displayed similar hypersensitivity to capsaicin on day 2 ($F_{1,5} = 2.4; P > 0.1$) (Fig. 6B). After long-term exposure to morphine (two injections of 1.8 mg/kg daily for 4 weeks), morphine-treated monkeys showed a decrease in antinociceptive effects produced by morphine (1 and 1.8 mg/kg; $P < 0.05$) (Fig. 6C), indicative of tolerance development. In contrast, after the same duration of repeated administration, AT-121–treated monkeys (0.03 mg/kg) did not show tolerance to antinociceptive effects of AT-121 at 0.01 or 0.03 mg/kg (Fig. 6D). These results demonstrate that unlike morphine, repeated administration of AT-121 did not cause opioid-induced hyperalgesia and may have a slower development of analgesic tolerance compared to morphine.

**DISCUSSION**

This study provides four significant findings in nonhuman primates indicating the translational potential of AT-121, a bifunctional NOP/MOP agonist, as a safe, nonaddictive analgesic and possible treatment for prescription opioid abuse (dual therapeutic action). First, AT-121 produced morphine-comparable antinociceptive and antihypersensitive effects mediated by NOP and MOP receptors. Second, unlike oxycodone, AT-121 lacked reinforcing effects, suggesting little or no abuse liability or apparently much less abuse liability than oxycodone. AT-121 selectively attenuated reinforcing effects of oxycodone, suggesting its treatment potential for prescription opioid abuse. Third, unlike heroin, AT-121 was safe and did not compromise respiratory and cardiovascular activities at 10 times the analgesic doses. Fourth, AT-121 produced less opioid-induced hyperalgesia, physical dependence, or tolerance than morphine after repeated administration.

AT-121 is about 100-fold more potent than morphine in producing antinociceptive effects. Because NOP agonists increased nociceptive threshold and inhibited capsaicin-induced allodynia in humans (43, 46, 47), the antinociceptive and antiallodynic effects of AT-121 suggest that its functional efficacy as an analgesic may be similar to MOP agonists. Although few pain models are established in primates, several rodent studies support the rationale for developing bifunctional NOP/MOP agonists as effective analgesics across diverse pain modalities (16, 18, 21). Previous studies in nonhuman primates showed that co-administration of NOP and MOP agonists produced synergistic antinociception without eliciting itch and respiratory depression (22). In addition, pretreatment with both NOP and MOP antagonists produced a much larger rightward shift of the AT-121 dose-response curve. Together, these findings with AT-121 support the enhanced potency and broad therapeutic window displayed by ligands with dual NOP/MOP agonist activity (9, 22).

Compared to highly abused drugs like oxycodone and cocaine, in this study, AT-121 did not produce reinforcing effects in monkeys. In our intravenous drug self-administration procedure in primates, considered a gold standard to evaluate the abuse potential of drugs (56, 57), AT-121 showed little to no abuse liability. The NOP partial agonist buprenorphine, on the other hand, produced mild-to-moderate reinforcing effects under the same schedule (32). Although AT-121 had comparable agonist efficacy as buprenorphine at MOP receptors (58), its higher agonist efficacy at NOP receptors (compared to that of buprenorphine) likely counteracts its MOP
receptor-mediated reinforcing effects. AT-121 attenuated reinforcing effects of oxycodone without affecting reinforcing effects of food pellets, indicating its selective attenuation of reinforcing effects of a prescription opioid. Because AT-121 was devoid of abuse liability compared to current medications such as buprenorphine, our findings suggest that bifunctional NOP/MOP agonists have potential as “replacement opioid therapy” for patients prescribed opioids and at risk for opioid dependence. Given the medical burden associated with misuse and abuse of prescription opioids (1, 2, 59), these findings shed light on the future potential of bifunctional NOP/MOP agonists.

Fig. 5. Development of physical dependence on morphine or AT-121 in nonhuman primates after short-term repeated administration. (A to D) Effects of naltrexone on respiration rate (A), minute volume (B) heart rate (C), and mean arterial pressure (D) in morphine- or vehicle-treated monkeys. (E to H) Effects of J-113397 (0.03 mg/kg) and naltrexone (0.01 mg/kg) on respiration rate (E), minute volume (F), heart rate (G), and mean arterial pressure (H) in AT-121- or vehicle-treated monkeys. Data are shown as changes from the baselines (before antagonist treatment). Each data point represents mean ± SEM (n = 4) from each individual data averaged from a 15-min time block. All compounds were delivered intramuscularly. Data were analyzed by two-way ANOVA with repeated measures, followed by Bonferroni’s multiple comparisons test. *P < 0.05, significantly different from vehicle from 15 to 30 min to the corresponding time point.

Fig. 6. Development of opioid-induced hyperalgesia and tolerance in nonhuman primates after repeated administration of morphine or AT-121. (A and B) Tail-withdrawal latencies in 46°C water after topical capsaicin (0.4 mg/ml × 0.3 ml) in vehicle-treated (A and B), morphine (1.8 mg/kg)–treated (A), and AT-121 (0.03 mg/kg)–treated (B) animals (n = 6). (C and D) Tail-withdrawal latencies in 50°C before (BL) and after (day 30) repeated administration of morphine (1.8 mg/kg, n = 4) (C) or AT-121 (0.03 mg/kg, n = 5) (D). Each data point represents mean ± SEM. Data were analyzed by two-way ANOVA with repeated measures, followed by Bonferroni’s multiple comparisons test. *P < 0.05, significantly different from the vehicle at 15 and 30 min. #P < 0.05, significantly different from BL values.
agonists as having a dual therapeutic action as nonaddictive analgesics and as medications to treat opioid use disorders (9, 16, 26). However, the bifunctional NOP/MOP partial agonist profile of AT-121 is distinguishable from that of cebranopadol (GRT6005), a NOP and opioid receptor agonist currently in clinical trials for acute and chronic pain (60). In contrast to AT-121, cebranopadol is a full agonist at the NOP and MOP receptors (29). Although cebranopadol had a wider therapeutic window than morphine and classical MOP agonists, it generalized to a morphine discriminative stimulus and showed a conditioned place preference in rodents (61, 62), suggestive of intrinsic rewarding properties and the potential for abuse liability.

AT-121 at antinociceptive doses and at 10-fold higher dose did not cause significant respiratory depression or affect cardiovascular activities immediately after administration or during and after its analgesic action. In contrast, a high dose of heroin quickly suppressed respiration rate. In contrast to the rapid respiratory depression/arrest by heroin or fentanyl (32) at analgesic doses, AT-121 demonstrates a wide safety window in primates. Overall, these in vivo findings in primates demonstrate that after acute administration, bifunctional NOP/MOP partial agonists such as AT-121 are safe, nonaddictive analgesics with a wider therapeutic window than traditional opioids.

Development of such bifunctional NOP/MOP agonists with a suitable profile of partial agonist efficacy at NOP and MOP receptors to treat pain and opioid addiction might help to reduce the burden related to opioid abuse (4, 5, 7).

After repeated administration, opioid analgesics often cause adverse events, such as opioid-induced hyperalgesia, physical dependence, and tolerance (52, 53, 55). After short-term exposure (1 or 3 days), morphine-treated primates developed opioid-induced hyperalgesia (enhanced sensitivity to pain) and physical dependence (precipitated withdrawal signs). In a side-by-side comparison with morphine, AT-121–treated primates did not develop opioid-induced hyperalgesia or physical dependence. Moreover, after long-term exposure (4 weeks), morphine-treated primates developed tolerance to its antinociceptive effects. In contrast, AT-121–treated primates showed a slower development of tolerance to antinociceptive effects. Although more frequent dosing and longer durations of treatment could result in tolerance, our findings indicate that bifunctional NOP/MOP agonists like AT-121 may have advantages over morphine in repeated or chronic dosing regimens. Central NOP receptors have an intriguing plasticity under chronic pain states (18, 21). Unlike down-regulation of MOP receptors, mRNA expression of NOP receptors remained unchanged in the spinal cord of diabetic primates with neuropathy (63). Future studies are warranted to investigate whether bifunctional NOP/MOP agonists cause tolerance to develop more slowly compared to clinically used MOP agonists in patients with chronic pain.

There were some limitations of this study. The effect of potential sex differences could not be evaluated with the limited number of primates. It is important to characterize the behavioral responses to bifunctional NOP/MOP receptor agonists in male and female subjects. Although primate studies are labor-intensive, studies with larger sample sizes can address these limitations and are warranted to further validate our findings.

In conclusion, the rational design and modulation of the bifunctional profile of a nonmorphinan chemical structure (42, 64, 65) resulted in the identification of a potent NOP/MOP bifunctional agonist AT-121. The functional utility of AT-121 extends beyond that of the recently reported orvinol analog BU08028, a universal opioid ligand with NOP/MOP agonist activity (32), or the NOP/MOP full agonist cebranopadol (29). In vivo findings with AT-121 in primates support bifunctional NOP/MOP partial agonists as safe, nonaddictive analgesics and suggest their potential as treatments for opioid use disorders. Generally, AT-121 was devoid of several adverse effects associated with clinically used MOP agonists after acute and repeated/chronic administration. It is pivotal to further investigate the interactions between NOP and MOP receptors with pharmacologic tools with varying efficacies at NOP versus MOP receptors (26, 66–68). Primate models will continue to be a translational bridge to facilitate research and development of bifunctional NOP/MOP agonists as treatment for pain and substance abuse.

MATERIALS AND METHODS

Study design

The main objective of this study was to design and optimize bifunctional NOP/MOP receptor agonists and establish the functional profile of a newly discovered agonist as a safe, nonaddictive analgesic and potential treatment for opioid abuse (dual therapeutic action) in primate models. For behavioral and physiological measurements, primate subjects were trained by using the pole and collar procedure to be transported between their home cages and procedure rooms and habituated to different experimental conditions. These healthy subjects were randomly selected for each study component listed below. On the basis of our previous experience using power and statistical analysis for quantifying primate behavioral and physiological responses (34, 44, 69–71), the sample size of four subjects is sufficient to determine the magnitude, duration of action, and dose dependency of ligand-induced effects. All behavioral measurements were conducted during the day time (9:00 a.m. to 3:00 p.m.) by laboratory staff who were blinded to dosing conditions.

Subjects

All animal care and experimental procedures were conducted in accordance with the Guide for the Care and Use of Laboratory Animals as adopted and promulgated by the U.S. National Institutes of Health and approved by the Institutional Animal Care and Use Committee of Wake Forest University. Fifteen adult male and female rhesus monkeys (Macaca mulatta), 10 to 18 years, 6.5 to 12.4 kg, were kept at an indoor facility accredited by the Association for Assessment and Accreditation of Laboratory Animal Care International. Animals were individually housed in species-specific rooms with environmental controls set to maintain 21 to 25°C, 40 to 60% relative humidity, and a 12-hour light-dark cycle (light: 6:30 a.m. to 6:30 p.m.). The daily diet consisted of about 20 to 28 biscuits (Purina Monkey Chow; Ralston Purina Co.), fresh fruit, and water ad libitum. Small amounts of primate treats and various cage enrichment devices were supplied as forms of environmental enrichment. Animals were not exposed to any opioid compound for 1 month before experiments. This study is reported in accordance with the Animal research: Reporting in vivo experiments (ARRIVE) guidelines for reporting experiments involving animals (72).

Sensory assays

Acute nociception

The warm water tail-withdrawal assay was used to evaluate thermal antinociceptive effects of AT-121 and morphine. Monkeys were seated in primate restraint chairs in a designated procedure.
room, and the lower parts of their shaved tails (~15 cm) were immersed in a thermal flask containing water maintained at 42°, 46°, or 50°C. Water at 42° and 46°C was used as nonnoxious stimulus, and water at 50°C was used as an acute noxious stimulus. All tail-withdrawal latencies were measured at each temperature using a computerized timer by individuals who were blinded to the experimental conditions. If a monkey did not remove its tail within 20 s (cutoff), the flask was removed and a maximum time of 20 s was recorded. Test sessions began with baseline measurements at each temperature. Subsequent tail-withdrawal latencies were measured at multiple time points after subcutaneous administration of the test compound. For dose-response curves, the test compound was administered by a cumulative dosing procedure with a 30-min inter-injection interval. Tail-withdrawal latencies were measured at 20 min after each injection. A single dose of the NOP receptor–selective antagonist J-113397 (0.1 mg/kg) or the MOP receptor–selective antagonist naltrexone (0.03 mg/kg) was administered subcutaneously 15 min before determination of dose-response curves to determine the NOP and MOP receptor components mediating AT-121–induced antinoception. The doses and pretreatment time for J-113397 and naltrexone were chosen based on previous studies (19, 22).

Capsaicin-induced thermal allodynia

Antiallodynic effects of AT-121 were evaluated by using a 1-hour pretreatment time regimen (1 hour before capsaicin administration). Capsaicin (1.2 mg/ml × 0.3 ml) was administered topically via a bandage attached on the terminal 3 to 5 cm of the tail for 15 min (70). The allodynic response was manifested as reduced tail-withdrawal latency from a maximum value of 20 s to ~2 to 3 s in 46°C water. This allodynic effect peaked at 15 min after removal of the capsaicin bandage, which is when tail-withdrawal latency was measured in 46°C water to determine the antiallodynic effects of the test compound (23, 70). For evaluating the potential development of opioid-induced hyperalgesia, a low dose (0.4 mg/ml × 0.3 ml) of capsaicin was used; this dosing condition elicited moderate, reduced tail-withdrawal latency (10 to 12 s in 46°C water), which allowed detection of enhanced hypersensitivity. Monkeys received AT-121 (0.03 mg/kg) or morphine (1.8 mg/kg) on day 1 (two injections: ~9:00 a.m. and 16:00 p.m.). Their responses to capsaicin were measured on the morning of day 2.

Itch-scratching responses

Scratching activity as a behavioral response to itch sensation was recorded on videotapes when monkeys were in their home cages (48). A 15-min recording session was conducted at 1 hour after subcutaneous administration of AT-121 (0.03 mg/kg) or morphine (1 mg/kg). A scratch was defined as one brief (<1 s) episode of scraping contact of the forepaw or hind paw on the skin surface of other body parts. Total scratches were counted by individuals who were unaware of the experimental conditions.

Drug self-administration

Monkeys with indwelling intravenous catheters and subcutaneous vascular access ports were used to evaluate the reinforcing effects of the test compound under a progressive-ratio schedule as described previously (32). Briefly, lever-pressing responses were maintained by injections of remifentanil (0.3 μg/kg) until responding was stable (mean ± three injections for three consecutive sessions with no trend). Dose-response curves were determined in each monkey by substituting saline, cocaine (0.03 mg/kg per injection), or a range of doses of AT-121 (0.3 to 10 μg/kg per injection) or oxycodone (0.3 to 10 μg/kg per injection) for the maintenance dose in a randomized order. Doses were available for at least five consecutive sessions and until responding was deemed stable. AT-121 (0.03 mg/kg), buprenorphine (0.1 mg/kg), or naltrexone (0.01 mg/kg) was administered intramuscularly 30 min before starting the progressive-ratio schedule of oxycodone (3 μg/kg per injection) to evaluate each compound’s potential attenuation of reinforcing effects by oxycodone. In addition, vehicle (0.1 ml/kg) or AT-121 (0.03 mg/kg) was administered intramuscularly 30 min before starting the fixed ratio (FR10) schedule of banana-flavored food pellets to determine whether AT-121 alters the value of other reinforcers or affects general motor function.

Physiological responses

Freely moving monkeys implanted with the D70-PCTR telemetry transmitter were used to evaluate the effects of AT-121 and MOP receptor agonists on physiologic functions as described previously (32). Respiration, heart rate, blood pressure, and temperature were measured and analyzed with Ponemah software version 5.2 (Data Sciences International). For acute drug effects, data from the 30-min interval before drug administration were collected as baseline and then at each time point (1, 3, and 6 hours) after intramuscular administration of AT-121 (0, 0.03, and 0.3 mg/kg) compared to a high dose of heroin (1 mg/kg). For detecting precipitated withdrawal signs after 3 days (two injections per day; first injection at ~09:00 a.m. and second injection at ~16:00 p.m.) of intramuscular administration of AT-121 (0.03 mg/kg) or morphine (1.8 mg/kg), data from the 30-min interval before administration of antagonist were collected and then continuously for 1 hour after antagonist administration on day 4. The mean value of each 15-min time block was generated from each subject to represent the measured outcome for each single data point.

Drugs

(i) The chemical synthesis procedures (fig. S1), (ii) in vitro pharmacological characterization of compounds 1 to 6, (iii) molecular docking and computational modeling, (iv) in vivo rodent pharmacokinetic data (fig. S2 and tables S1 and S2), and (v) in vitro primate plasma stability data (fig. S3 and table S3) are described and presented in detail in the Supplementary Materials. AT-121 was dissolved in a solution of dimethyl sulfoxide/1% (w/v) hydroxypropyl cellulose in a ratio of 3:97. Morphine sulfate, buprenorphine HCl, heroin (diacromorphine HCl), remifentanil HCl, and naltrexone HCl [National Institute on Drug Abuse (NIDA)] were dissolved in sterile water. (~)Cocaine HCl and oxycodone HCl (NIDA) were dissolved in sterile 0.9% saline. J-113397 (R&D Systems) was dissolved in a solution of dimethyl sulfoxide/Tween 80/sterile water in a ratio of 1:1:8. Capsaicin (Sigma-Aldrich) was dissolved in 70% (v/v) dimethyl sulfoxide/sterile water. Heroin (diacromorphine HCl), remifentanil HCl, and naltrexone HCl [National Institute on Drug Abuse (NIDA)] were dissolved in sterile water. Oxycodone and remifentanil were dissolved in sterile 0.9% saline. J-113397 (R&D Systems) was dissolved in a solution of dimethyl sulfoxide/Tween 80/sterile water in a ratio of 1:1:8. Capsaicin (Sigma-Aldrich) was dissolved in 70% (v/v) ethanol. For systemic administration, drugs were administered at a volume of 0.1 ml/kg. There was a minimum 1-week interval between drug administrations.

Statistical analysis

Mean values ± SEM were calculated from individual data for all study endpoints. Comparisons were made for the same monkeys across all test sessions in the same experiment. Data were analyzed by either two-way ANOVA with repeated measures, followed by Bonferroni’s multiple comparisons test, or one-way ANOVA with repeated measures, followed by Bonferroni’s multiple comparisons test. The criterion for significance for all tests was set at P < 0.05. To
analyze nociceptive responses, individual tail-withdrawal latencies were converted to the percentage of maximum possible effect by using the formula defined as [(test latency – control latency)/(cutoff latency, 20 s – control latency)] × 100. Raw data for all experiments are presented in Table S5.

**SUPPLEMENTARY MATERIALS**

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Materials and Methods

Fig. S1. Chemical synthesis scheme for compounds 1 to 6.

Fig. S2. AT-121 plasma and brain concentrations after a single subcutaneous administration of 3 mg/kg to male Sprague-Dawley rats.

Fig. S3. In vitro plasma stability assessment of AT-121 in nonhuman primate plasma.

Fig. S4. Effects of systemic administration of AT-121 on physiologic functions of freely moving nonhuman primates implanted with telemetry probes.

Table S1. Brain sample records for AT-121 brain penetration after subcutaneous administration of 3 mg/kg to male Sprague-Dawley rats.

Table S2. Pharmacokinetic parameters (plasma and brain) for AT-121 after subcutaneous administration in male Sprague-Dawley rats.

Table S3. Plasma stability assessment of AT-121 in nonhuman primate plasma.

Table S4. Baseline values for the nonhuman primate tail-withdrawal assay that are normalized across different dosing conditions.

Table S5. Raw data.

References (74, 75)

**REFERENCES AND NOTES**


T. M. Tschentschke, K. Rutten, Mu-opioid peptide (MOP) and nociceptin/orphanin FQ peptide (NOP) receptor activation both contribute to the discriminative stimulus properties of cebranopadol in the rat. Neuropharmacology 129, 100–108 (2018).


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A bifunctional nociceptin and mu opioid receptor agonist is analgesic without opioid side effects in nonhuman primates

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A dual-targeting painkiller

Opioids are among the most effective treatments for severe pain. Their pain-relieving effects are mediated by activation of the mu opioid peptide (MOP) receptor. Unfortunately, selective MOP agonists induce diverse side effects, including respiratory depression, tolerance, hyperalgesia, and dependence. Recently, activation of the nociceptin/orphanin FQ peptide (NOP) receptor has been reported to enhance MOP agonist-induced analgesia without producing side effects. Now, Ding et al. have developed a bifunctional MOP/NOP agonist, called AT-121, that showed potent analgesic effects in nonhuman primates without inducing hyperalgesia, respiratory depression, or dependence. The results suggest that bifunctional MOP/NOP agonists might represent a safe and effective pharmacological tool for treating severe pain.