CAMP-Mediated Mechanisms for Pain Sensitization during Opioid Withdrawal

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Chronic opioid-induced drug dependence and withdrawal syndrome after opioid cessation remain a severe obstacle in clinical treatment of chronic pain and opioid drug addiction. One of the key symptoms during opioid withdrawal is a state of sensitized pain. The most significant molecular adaptation induced by chronic opioids in the brain is upregulation of the CAMP-signaling pathway. Although the CAMP system is known to have multiple effects on central neuron functions, how its upregulation mediates behavioral opioid dependence and withdrawal-induced pain in vivo remains unclear. In this study, we demonstrate that withdrawal from chronic morphine significantly upregulates the mRNA level of adenylyl cyclase (AC) VI and VIII isoforms and immunoreactivity of ACV/VI in the nucleus raphe magnus (NRM), a brainstem site critically involved in opioid modulation of pain. In cellular studies of NRM neurons containing μ-opioid receptors, we show that morphine withdrawal significantly increases glutamate synaptic transmission via a presynaptic mechanism mediated by an upregulated CAMP pathway. Morphine withdrawal also enhances the hyperpolarization-activated current in these neurons by increased intracellular CAMP. Both of the withdrawal-induced CAMP actions increase the excitability of these μ-receptor-containing neurons, which are thought to facilitate spinal pain transmission. Furthermore, in morphine-dependent rats in vivo, blocking the CAMP pathway significantly reduces withdrawal-induced pain sensitization. These results illustrate neurobiological mechanisms for the CAMP-mediated withdrawal pain and provide potential therapeutic targets for the treatment of opioid dependence and withdrawal-related problems.

Key words: glutamate EPSCs; Ih; adenylyl cyclase; nucleus raphe magnus; hyperalgesia; opioid dependence

Introduction
Repeated use of opioid drugs, either for clinical treatment of chronic pain or as a result of drug abuse, induces neuroadaptive changes in the brain that form the neurobiological basis for drug dependence and addiction (Nestler, 2001; Williams et al., 2001; Chao and Nestler, 2004). Abrupt abstinence or withdrawal from opioid drugs causes a series of severe adverse symptoms, which keep drug-dependent individuals craving continued opioids. One of the core withdrawal symptoms is an increase in pain sensitivity (pain sensitization or hyperalgesia), which has been well documented both in animal studies (Mao et al., 1995; Nestler and Aghajanian, 1997) and in clinical reports, including those of opioid addicts (Doverty et al., 2001; Angst et al., 2003; Compton et al., 2003). Despite decades of research the underlying neurobiological mechanisms for opioid dependence and withdrawal remain unclear, leaving current treatment for opioid dependence and withdrawal mostly empirically with limited success. Early studies, using in vitro models of cell lines, showed that chronic morphine induces a compensatory increase in the adenylyl cyclase (AC) activity and intracellular CAMP concentration, and with morphine withdrawal the CAMP concentration overshoots to above premorphine levels (Sharma et al., 1975; Brandt et al., 1976). Subsequently, this chronic morphine-induced molecular adaptation of the CAMP cascade has been confirmed in many opioid addiction-related brain areas, and such a CAMP adaptation has been widely related to opioid dependence and withdrawal (for review, see Nestler and Aghajanian, 1997; Nestler, 2001; Williams et al., 2001; Watts, 2002; Chao and Nestler, 2004). Previous in vitro studies that used CAMP analogs have identified several direct CAMP-mediated actions on central neurons, including increasing synaptic transmission and augmenting the hyperpolarization-activated current (Ih) (for review, see Williams et al., 2001). It has been demonstrated recently that GABA synaptic transmission is augmented via an upregulated CAMP cascade in several brain areas of morphine-withdrawn rats (Bonci and Williams, 1997; Ingram et al., 1998; Jolas et al., 2000). The behavioral significance of this GABA synaptic adaptation remains to be demonstrated. Although the CAMP pathway has been implicated in the expression of many somatic symptoms of opioid withdrawal (Punch et al., 1997), the mechanisms by which this adaptive upregulation of the CAMP pathway mediates the behavioral syndrome of opioid dependence and withdrawal remain unclear. Especially lacking are mechanisms linking chronic opioid-induced CAMP upregulation in vivo to the behavioral...
symptom of pain sensitization during opioid withdrawal. In this study, we used a rat model of opioid dependence and withdrawal to investigate effects of chronic morphine on AC expression levels and, consequently, on CAMP-mediated synaptic actions and pain behaviors, focusing on neurons in the nucleus raphe magnus (NRM), a brainstem site critically involved in opioid modulation of pain transmission in the spinal cord (Scholz and Woolf, 2002; Fields, 2004).

Materials and Methods

Chronic morphine treatment and morphine withdrawal. Rats were treated with chronic morphine for 6 d by subcutaneous implantation of morphine pellets as described previously (Pan, 2003a). Placebo pellets were used as controls. For real-time PCR and Western blotting experiments, naloxone (1 mg/kg, i.p.) was injected on day 7 in both morphine- and placebo-treated rats 5–7 h before tissue samples were taken. For recordings, the morphine withdrawal was induced either by the application of naloxone (1 μM) to morphine-dependent brain slices maintained in 5 μM morphine in vitro or by the incubation of morphine-dependent slices in a morphine-free solution for at least 3 h (spontaneous withdrawal).

Relative quantitative real-time PCR and Western blotting. Tissue samples containing the NRM were taken from morphine-withdrawn or placebo control rats (n = 8). Total RNA was prepared by using an RNAqueous-4 PCR kit (Ambion, Austin, TX) and reverse transcribed by using a SYBR Green reverse transcription (RT)–PCR Reagents kit (Applied Biosystems, Foster City, CA). The sequences of primers designed with the Primer Express software (Applied Biosystems) included the following: for ACI, CCAGGGAAGATTCGATGC and GCGCTGTCACAGCAGCAGTGTCA and CTCTGGGAAAAGTTCAGGTTGA; for ACV, CTGAGCTAGCTCTGCTATGTG and GGAGTCCCTGGGGGAGGC; for ACVIII, TTGCTTCTCAGCCTGCTG and CTCTGGGAAGTTGCAGTTGGA; for ACVI, CTGCCCTCAGCCTGCTG and CTGAGGTCAAGGATGCGTGA; and for β-actin, CACCCACACTGTGCCCATCTA and AGGCTCGGTCGGGCCAGCAC. Primer concentrations were optimized. Real-time PCRs were performed by using SYBR Green I Master Mix and the ABI 7000 Sequence Detector. Amplification efficiencies were comparable, as indicated by the lines in plots of the following (in mM): 126 NaCl, 2.5 KCl, 1.2 NaH2PO4, 1.2 MgCl2, 2.4 CaCl2, 11 glucose, and 25 NaHCO3, saturated with 95% O2/5% CO2, pH 7.2–7.4. Slices were maintained at ~35°C throughout the recording experiment. Neonatal rats were used for better visualization of neurons in brain slices with an infrared Nomarski microscope. It has been demonstrated that the physiological and pharmacological properties of neurons from these young rats are indistinguishable from those of adult rats (Pan et al., 1997). Visualized whole-cell voltage-clamp recordings were made from identified neurons with a glass pipette (resistance, 3–5 MΩ) filled with a solution containing the following (in mM): 126 potassium gluconate, 10 NaCl, 1 MgCl2, 11 EGTA, 10 HEPES, 2 ATP, and 0.25 GTP, pH adjusted to 7.3 with KOH; osmolality, 280–290 mOsm/L. An AxoPatch 1-D amplifier and AxoGraph software 4.7 (Axon Instruments, Union City, CA) were used for data acquisition and on-line/off-line data analyses. A seal resistance of ≥2 GΩ and an access resistance of ≤15 MΩ were considered acceptable. Series resistance was compensated optionally. The access resistance was monitored throughout the experiment. All NRM cells that were recorded were classified into either a cell type that contains the μ-opioid receptor (termed “secondary cell”) or the other cell type that lacks the μ-receptor (termed “primary cell”), according to the criteria described in our previous study (Pan et al., 1990). Electrical stimuli of constant current (0.25 ms, 0.2–0.4 mA) were used to evoke EPSCs with bipolar-stimulating electrodes placed lateral (200–400 μm) to the recording electrode within the NRM. A pair of EPSCs was evoked by two stimuli with an interval of 40 ms. Paired pulse ratio (PPR) was calculated by dividing the amplitude of the second EPSC by the first one. The PPR has been used widely to determine synaptic action site (Manabe et al., 1993; Perkel and Nicoll, 1993; Dobrunz and Stevens, 1997; Ungless et al., 2001; Bie and Pan, 2003). It has an inverse relationship with the probability of presynaptic neurotransmitter release. Thus a decrease in the PPR indicates an increase in presynaptic transmitter release and vice versa. Miniature EPSCs (mEPSCs) were obtained in 60 s epochs in the presence of tetrodotoxin (TTX; 1 μM). The AxoGraph software was used to detect and measure the amplitude and intervals of the synaptic events and to analyze their distribution data. The amplitude of the 1st was determined by the difference in the current values between the beginning and the end of a 2 s hyperpolarizing voltage step from a holding potential of –50 mV. Ioff data were fit with the Boltzmann equation, and then the voltage value (V1/2) at which one-half of the maximum conductance is activated was determined from the Ioff activation curve (Pan, 2003b). In some experiments, NRM slices from chronic morphine-treated rats were incubated in the AC inhibitor cis-N-(2-phenylcyclopentyl)azacyclotridec-1-en-2-amine (MDL12330A; 100 μM) or in the protein kinase A (PKA) inhibitor N-[2-(p-bromocinnamylamino)ethyl]-5-isouquinolinesulfonamide (H89; 10 μM) for at least 1 h. Firing of action potentials was evoked by a 3 s depolarizing current step in control, in morphine-withdrawn slices, and in withdrawn slices treated with the Ioff blocker 4-(N-ethyl-N-phenylamino)-1,2-dimethyl-6-(methylamino)pyridinium chloride (ZD7288; 10 μM for >1 h). ZD7288 at 25–100 μM has been shown to selectively block the Ioff in many types of central neurons, including μ-receptor-lacking neurons in the NRM (Harris and Constanti, 1995; Khakh and Henderson, 1998; Takigawa et al., 1998; Larkman and Kelly, 2001; Mellor et al., 2002; Pan, 2003b). Statistical analyses of the mEPSCs were performed by using Statview software (SAS Institute, Cary, NC) with the Kolmogorov–Smirnov test. Other numerical data were analyzed statistically with Student’s t tests and presented as mean ± SEM. Generally, only one neuron was recorded in one slice, and one to two slices from each rat were used for recording.
Drugs generally were applied via the bath solution unless otherwise specified.

**Microinjection and behavioral experiments.** Male Wistar rats (250–300 g) were maintained lightly anesthetized in a stereotaxic apparatus with a constant intravenous infusion of methohexital (10 mg/ml at 0.8 ml/h). A 26-gauge double-guide cannula was aimed at the NRM (anteroposterior, –10.0 from the bregma; lateral, 0; ventral, –10.5 from dura with the skull leveled) (Paxinos and Watson, 1986). Control placement of the cannula was 1.5 mm dorsal to the NRM placement. Tail-flick latency to a radiant heat stimulus was measured every 2 min. The heat intensity was set to 4.6 mV (ZD7288 or AP-5 and CNQX was microinjected immediately before naloxone). Drug effects were analyzed statistically by an ANOVA for repeated measures and the Tukey–Kramer test of post hoc analysis. For histological verification of cannula placements, 0.5 μl of methylene blue was injected through the double-injector cannula. The rat brain was fixed in 4% paraformaldehyde for 30 min, and then NRM slices (200 μm thick) were cut and examined for cannula placement under a microscope. Data from rats in which cannula placements were outside of the NRM area were excluded from the results. Morphine and placebo pellets were kindly supplied by the National Institute on Drug Abuse. All other drugs were purchased either from Sigma–Aldrich or from Tocris Cookson (Ellisville, MO).

**Results**

**Morphine withdrawal increases mRNA level and immunoreactivity of AC**

We next determined whether cAMP analogs could modulate glutamate synaptic transmission in normal NRM slices from morphine-naïve rats, focusing on a class of NRM cells (termed secondary cells) that express μ-opioid receptors (Pan et al., 1997). In these cells methionine-enkephalin (10 μM), an endogenous opioid peptide acting on the μ-opioid receptor, produced a membrane hyperpolarization of 24 ± 4.5 mV (n = 12), an effect similar to what we have reported previously (Pan et al., 1997, 2000). Forskolin (10 μM), an AC activator, significantly increased the amplitude of glutamate-mediated evoked EPSCs in these cells sensitive to μ-receptor agonists (control, 95.1 ± 4.4 pA; forskolin, 144.1 ± 14.2 pA; p < 0.01; n = 12) (Fig. 2A). The membrane-permeable cAMP analog 8-bromo-cAMP (8-br-cAMP; 1 mM) also increased the EPSC amplitude (control, 110.9 ± 14.8 pA; 8-br-cAMP, 166.6 ± 19.2 pA; p > 0.05; n = 4). To determine the synaptic site of this cAMP action, we first used the paradigm of PPR. Forskolin significantly decreased the PPR of glutamate EPSCs (control, 1.73 ± 0.21; forskolin, 1.03 ± 0.14; p < 0.01; n = 5) (Fig. 2B) and so did 8-br-cAMP (control, 1.70 ± 0.08; 8-br-cAMP, 1.10 ± 0.07; p < 0.01; n = 4) (Fig. 2C). This indicates that a presynaptic site is involved in the cAMP action on glutamate release. Forskolin also produced a significant increase in the frequency, but not the amplitude, of the mEPSCs (for frequency: control, 5.70 ± 1.03 Hz; forskolin, 8.46 ± 1.07 Hz; p < 0.05; for amplitude: control, 20.76 ± 2.04 pA; forskolin, 20.64 ± 2.40 pA; p > 0.05; n = 5) (Fig. 2D). The opioid antagonist naltxone (1 μM) had no effect on the frequency or amplitude of the mEPSCs in these control cells (Fig. 2D).

**Morphine withdrawal increases glutamate synaptic transmission through upregulated cAMP cascade**

We first used the real-time RT-PCR technique to examine whether withdrawal from chronic morphine changed the expression of AC at the mRNA level in the NRM. We focused on ACI, ACV, ACVII, and ACVIII, the four AC isoforms that are inhibited by acute opioids (Aviaron-Reiss et al., 1997). Amplification plots demonstrated that the cycle thresholds for ACVII and ACVIRA mRNA levels were upregulated in the NRM area during morphine withdrawal (Fig. 1A, B). When normalized to internal reference β-actin, the mRNA level of both ACVII and ACVIII was increased significantly in withdrawn rats (Fig. 1C). There was a small increase in ACV mRNA, but the difference did not reach statistical significance. ACI mRNA was not changed. To determine changes in these AC isoforms at the protein level, we used Western blotting for AC immunoreactivity. After being normalized to β-actin, the immunoreactivity of ACV/VI was enhanced significantly in the NRM from morphine-withdrawn rats, whereas ACI and ACVIII immunoreactivity was unchanged (Fig. 1D, E). Thus it appears that morphine withdrawal increases ACVII and ACVIII mRNA levels and ACV/VI protein levels in the NRM.
synaptic transmission in NRM cells, morphine withdrawal should increase glutamate synaptic transmission in these cells. We then tested this hypothesis by using naloxone-precipitated and morphine-abstinent withdrawn slices from chronic morphine-treated rats. In morphine-dependent slices, naloxone (1 μM) significantly increased the amplitude of glutamate EPSCs in the μ-receptor-containing cells (dependent, 138.3 ± 20.1 pA; plus naloxone, 191.5 ± 29.6 pA; p < 0.01; n = 11) (Fig. 3A). It also reduced the PPR in these cells (dependent, 1.37 ± 0.13; plus naloxone, 1.08 ± 0.09; p < 0.01; n = 11) (Fig. 3C). In control slices from placebo-treated rats, naloxone was ineffective on the EPSC or the PPR in these cells (control, 1.94 ± 0.29; plus naloxone, 2.02 ± 0.36; p > 0.05; n = 8). In contrast to the effect of naloxone in these μ-receptor-containing cells, naloxone had no effect in the other type of cells lacking the μ-receptor from morphine-treated rats (treated, 128.6 ± 16.7 pA; plus naloxone, 123.4 ± 17.1 pA; p > 0.05; n = 8) (Fig. 3B). Furthermore, naloxone significantly increased the frequency, but not the amplitude, of the mEPSCs in the μ-receptor-containing cells (for frequency: dependent, 5.13 ± 1.3 Hz; plus naloxone, 9.47 ± 1.56 Hz; p < 0.05; for amplitude: dependent, 20.51 ± 0.7 pA; plus naloxone, 21.52 ± 1.01 pA; p > 0.05; n = 8) (Fig. 3 E,F). Naloxone was without effect either on the PPR or on the mEPSCs in the NRM cells lacking μ-receptors. These results suggest that naloxone-precipitated withdrawal enhances glutamate synaptic transmission presynaptically by increasing glutamate release only in μ-receptor-containing NRM cells.

Next we examined whether this naloxone withdrawal-induced enhancement of glutamate neurotransmission was mediated by an upregulated cAMP cascade. We found that the EPSC-enhancing effect of forskolin in the μ-receptor-expressing cells was significantly larger in withdrawn slices than that in control slices (Fig. 3A,D), likely indicating a sensitized or upregulated AC system (Bonci and Williams, 1997; Ingram et al., 1998). Forskolin also increased the mEPSC frequency, but not the amplitude, in the withdrawn cells (Fig. 3E,F). In morphine-dependent slices treated with the AC inhibitor MDL12330a (100 μM), naloxone was no longer effective on the evoked EPSC amplitude (dependent/MDL12330a, 129.4 ± 17.3 pA; plus naloxone, 126.9 ± 20.2 pA; p > 0.05; n = 6) (Fig. 3G) or on the PPR (dependent/MDL12330a, 1.53 ± 0.07; plus naloxone, 1.50 ± 0.11; p > 0.05; n = 6) (Fig. 3H). Similar treatment with the PKA inhibitor H89 (10 μM) also abrogated the effect of naloxone (for EPSC: dependent/H89, 123.2 ± 13.5 pA; plus naloxone, 128.4 ± 9.9 pA; p > 0.05; n = 8; for PPR: dependent/H89, 1.46 ± 0.11; plus naloxone, 1.39 ± 0.13; p > 0.05; n = 7) (Fig. 3I). In addition, consistent results were obtained in NRM spontaneous withdrawal slices. In these withdrawn slices, μ-receptor-containing cells displayed a significantly larger EPSC (control, 111.5 ± 7.8 pA; n = 76; withdrawn, 140.5 ± 8.2 pA; n = 27; p < 0.01) (Fig. 4A,B). They also exhibited significantly smaller PPRs than those in control slices (control, 1.89 ± 0.13, n = 33; withdrawn, 1.12 ± 0.04, n = 15; p < 0.01), indicating enhanced synaptic release of glutamate in these spontaneously withdrawn cells. Treatment of these withdrawn slices with MDL12330a (100 μM) significantly reduced the withdrawal effect on the PPR (plus MDL12330a, 1.59 ± 0.08; p < 0.01; n = 16) (Fig. 4C,D). These results obtained from both naloxone-induced and spontaneous withdrawal slices indicate the involvement of the cAMP pathway in the increased glutamate neurotransmission.

**Figure 2.** Activation of cAMP pathway presynaptically increases glutamate synaptic transmission in μ-receptor-containing NRM cells from control rats. A–C, Superimposed glutamatergic EPSC (A) and EPSC pairs normalized to the first EPSC (B, C) in control and in forskolin (10 μM) or forskolin plus forskolin. cAMP, 8-Br-cAMP; Con, control; Forsk, forskolin; Nal, naloxone.

**Morphine withdrawal enhances hyperpolarization-activated current through upregulated cAMP cascade**

cAMP also modulates the voltage dependence of the I_h or hyperpolarization-activated, cyclic nucleotide-gated (HCN) channels (Ingram and Williams, 1994; Luthi and McCormick, 1999; Wainger et al., 2001; Mellor et al., 2002). These μ-receptor-expressing NRM cells, which display a large I_h (Pan, 2003b), had typical I_h properties, including blockade by cesium and by the I_h channel blocker ZD7288, but not by barium (data not shown). In control cells, the inclusion of 8-br-cAMP (1 mM) in the recording pipette increased the average amplitude of the I_h and I_h tails at submaximal levels, but not the maximum (Fig. 5 A,B). Analysis with the Boltzmann equation revealed that the cAMP analog produced a positive shift of 8.7 mV, on average, in the I_h voltage dependence measured by V_1/2 at the half-maximal current without affecting its maximum conductance (control, −76.1 ± 0.9 mV, n = 12; 8-br-cAMP, −67.4 ± 1.7 mV, n = 15; p < 0.001) (Fig. 5C). This is consistent with the I_h-modulating mechanism by cAMP described in previous studies (Ingram and Williams, 1994; Luthi and McCormick, 1999; Wainger et al., 2001; Pan, 2003b). Forskolin produced a smaller effect with a positive shift of 2.1 mV, which did not reach statistical significance (control, −75.1 ± 1.1 mV; forskolin, −73.0 ± 2.2 mV; p > 0.05; n = 9). Because we have shown above that the cAMP pathway in the NRM likely is upregulated during morphine withdrawal, one would expect that morphine withdrawal would modulate the I_h similarly in these μ-receptor-containing cells. Indeed, when compared with control slices, μ-receptor-containing cells in spontaneously withdrawn slices showed a positive shift of 3.5 mV in V_1/2 (control, −76.1 ± 0.9 mV, n = 12; withdrawn, −72.6 ± 0.6 mV, n = 14; p < 0.01) (Fig. 5D). Treatment with ML121330a (100 μM) mostly blocked this withdrawal-induced V_1/2 shift (withdrawn/ML121330a, −75.8 ± 0.8 mV; n = 18; p < 0.01 when compared with V_1/2 in withdrawal and p > 0.05 when compared with V_1/2 in control) (Fig. 5E). These data indicate that morphine withdrawal-induced I_h enhancement likely is mediated by the upregulated cAMP cascade.
Inhibition of cAMP cascade attenuates withdrawal pain

Morphine withdrawal-induced increase in both glutamate synaptic transmission and the membrane-depolarizing \( I_h \) would increase the activity of these \( \mu \)-receptor-expressing NRM cells. Consistently, these cells in spontaneously withdrawn slices displayed more-positive resting membrane potentials (control, \(-52.6 \pm 0.8\) mV, \( n = 22 \); withdrawn, \(-50.8 \pm 0.7\) mV, \( n = 23 \); \( p < 0.05 \)) and a higher evoked firing rate (control, \( 17.6 \pm 2.5\) Hz, \( n = 21 \); withdrawn, \( 27.7 \pm 1.7\) Hz, \( n = 31 \); \( p < 0.01 \)) (Fig. 6). Treatment of the withdrawn slices with a low concentration of the \( I_h \) blocker ZD7288 (10 \( \mu \)M) significantly reduced the firing rate (withdrawn/ZD7288, \( 18.2 \pm 1.5\) Hz; \( p < 0.01 \); \( n = 22 \)) (Fig. 6), indicating an important \( I_h \) role in control of the cell excitability.

Previous studies, including ours, have shown that these \( \mu \)-receptor-containing NRM cells may facilitate spinal pain transmission (Urban and Gebhart, 1999; Pan et al., 2000; Porreca et al., 2002; Bie and Pan, 2003; Fields, 2004). Therefore, we hypothesized that the cAMP-mediated mechanisms described above in these cells contributed to the symptom of pain sensitization (hyperalgesia) during opioid withdrawal. In behavioral experiments in rats in vivo, naloxone (3 \( \mu \)g/kg, i.v.) had no effect on pain behaviors in placebo-treated rats, but it significantly decreased the pain threshold measured by the tail-flick test in rats treated with chronic morphine, indicating morphine withdrawal-induced pain sensitization (Fig. 7A). Microinjection of the glutamate receptor antagonists AP-5 (197 ng in 1 \( \mu \)l) and CNQX (276 ng in 1 \( \mu \)l) into the NRM significantly attenuated the hyperalgesia (\( n = 4 \) rats) (Fig. 7A). This is consistent with our previous studies of acute morphine withdrawal, which suggest that glutamate-mediated activation of the \( \mu \)-receptor-expressing cells, and not the other \( \mu \)-receptor-lacking NRM cells, primarily mediates the withdrawal hyperalgesia (Pan et al., 1997, 2000; Bie and Pan, 2003). Supporting a role for \( I_h \), in the control of the firing of these cells shown above in vitro, microinjection of a low dose of ZD7288 (293 ng in 1 \( \mu \)l) also significantly reduced the withdrawal pain (\( n = 5 \) rats) (Fig. 7B). Pretreatment by microinjection of the AC inhibitor MDL12330a (377 ng in 1 \( \mu \)l) into the NRM significantly antagonized this withdrawal-induced pain sensitization (\( n = 6 \) rats) (Fig. 7C). Furthermore, NMR microinjection of the PKA inhibitor H89 (519 ng in 1 \( \mu \)l; \( n = 6 \) rats) or Rp-cAMP (8.9 \( \mu \)g in 1 \( \mu \)l; \( n = 5 \) rats) before the naloxone injection also suppressed the withdrawal pain (Fig. 7D, E). However, microinjection of the same dose of Rp-cAMP into a site directly dorsal to the NRM failed to alter the withdrawal-induced hyperalgesia (\( n = 3 \) rats) (Fig. 7F), indicating a local NRM-mediated effect for the reduction in withdrawal pain. Figure 8 shows representative cannula placements within the NRM and dorsal to the NRM.

Discussion

The current study has identified two neuroadaptations induced by in vivo administration of chronic morphine in rats: an increase in glutamate synaptic transmission and an enhancement of the \( I_h \) in the same population of \( \mu \)-receptor-containing NRM neurons. Both adaptations appear to be mediated by chronic morphine-induced upregulation of the cAMP pathway in vivo. The current

![Figure 3. Naloxone-precipitated morphine withdrawal presynaptically increases glutamate synaptic transmission by cAMP upregulation. A, B, Glutamate EPSCs in control, in naloxone, and in naloxone plus forskolin from a \( \mu \)-receptor-containing cell (A) and from a \( \mu \)-receptor-lacking cell (B) of a morphine-dependent rat. Abbreviations are as in Figure 2. The following results were obtained from \( \mu \)-receptor-containing cells. C, Normalized EPSC pairs in control and in naloxone in a morphine-dependent cell. D, Percentage increase by forskolin of the EPSC amplitudes in control (open circles; \( n = 12 \)) and in withdrawn cells (filled circles; \( n = 6 \)), \( * p < 0.05 \); \( ** p < 0.01 \). E, Frequency (\( F \)) and amplitude (\( A \)) distributions of the mEPSCs in control, in naloxone, and in naloxone plus forskolin from the same morphine-dependent cell. G–I, Glutamate EPSCs (G, superimposed) and normalized EPSC pairs (H, I, staggered) in control and in naloxone from morphine-dependent cells treated with MDL12330a (MDL; 100 \( \mu \)M) or H89 (10 \( \mu \)M).](https://example.com/image3)

![Figure 4. Morphine abstinence-induced spontaneous withdrawal increases glutamate synaptic transmission by cAMP upregulation in \( \mu \)-receptor-containing cells. A, Representative EPSCs in neurons from a control slice and from a spontaneously withdrawn slice. B, Group data of PPRs from the same three groups as in A. Numbers in the columns indicate the cell numbers. C, Normalized EPSC pairs in neurons from a control slice, a withdrawn slice, and a withdrawn slice treated with MDL12330a (MDL). D, Group data of PPRs from the same three groups as in C. \( * p < 0.05 \); \( ** p < 0.01 \). Withdr., Withdrawal. Error bars represent SEM.](https://example.com/image4)
Morphine withdrawal enhances the $I_h$ by cAMP upregulation in µ-receptor-containing cells. A. Current traces (aligned on the instantaneous current level) recorded with a control and an 8-br-cAMP-filled (1 mM) pipette during a voltage step (~50 to ~80 mV). B. Averaged amplitudes of $I_h$ tails recorded with control ($n = 12$) and 8-br-cAMP-filled pipettes ($n = 15$). C–E, $I_h$ activation curves in control ($V_{1/2} = –76.1$ mV; $n = 12$) and in 8-br-cAMP ($V_{1/2} = –67.4$ mV; $n = 15$; $p < 0.001$) (C), in control (same as in C), in morphine-withdrawn cells ($V_{1/2} = –72.6$ mV; $n = 14$; $p < 0.01$) (D), in withdrawn cells (same as in D), and in another group of withdrawn cells treated with MDL12330a (MDL) ($V_{1/2} = –75.8$ mV; $n = 18$; $p < 0.01$) (E). Con, Control. Error bars represent SEM.

Figure 6. Inhibition of $I_h$ channels reduces evoked firing of µ-receptor-containing cells in withdrawn slices. Evoked firing of action potentials in cells from a control slice, a withdrawn slice, and a withdrawn slice (Withdr.) treated with ZD7288 (10 μM) is shown.

Figure 7. Inhibition of the cAMP pathway attenuates morphine withdrawal-induced pain sensitization. A–F, Plots of tail-flick latencies in placebo-treated rats (open circles) and in morphine-treated rats (filled symbols). Naloxone-induced withdrawal hyperalgesia (filled circles) was reduced by NRM microinjection of AP-5 and CNQX (A; filled squares; $n = 4$ rats), by ZD7288 (B; filled squares; $n = 5$), by MDL12330a (MDL) (C; filled triangles; $n = 6$), by H89 (D; filled diamonds; $n = 6$), and by Rp-cAMP (E; filled diamonds; $n = 5$). *$p < 0.05$; **$p < 0.01$ (ANOVA for repeated measures and the Tukey–Kramer test of post hoc analysis). F. Same microinjections of Rp-cAMP as in E, with cannula placement dorsal to the NRM (n = 3 rats). Error bars represent SEM.

serotonergic dorsal raphe nucleus (Jolias et al., 2000; Chao and Nestler, 2004). The complete molecular mechanisms for the chronic opioid-induced cAMP upregulation remain to be demonstrated (Watts, 2002; Chao and Nestler, 2004). Recent research shows that AC has nine different isoforms for which the brain distributions and cellular locations are highly specific, indicating complex and multiple presynaptic and postsynaptic actions of the enzyme (Williams et al., 2001; Watts, 2002). Although ACI, ACIII, and ACVIII display no synaptic plasticity of long-term potentiation in the hippocampus and amygdala of the human alcoholic brain (Yamamoto et al., 2001). Mice lacking both ACI and ACVIII display no synaptic plasticity of long-term potentiation in the hippocampus (Wong et al., 1999). Our current results study also shows that both of the cAMP-mediated actions, which increase the activity of these presumably pain-facilitating neurons, contribute to the pain sensitization during morphine withdrawal.

Since its discovery in cell lines in vitro (Sharma et al., 1975; Brandt et al., 1976), chronic opioid-induced upregulation of the cAMP signaling pathway has been demonstrated in many brain areas after chronic morphine treatment (Nestler and Aghajanian, 1997; Jolias et al., 2000; Nestler, 2001; Williams et al., 2001; Chao and Nestler, 2004). These brain areas are critical either for the reinforcing effects of drugs of abuse including opioids in drug addiction, such as the ventral tegmental area (VTA) and the nucleus accumbens, or for the expression of somatic signs of opioid withdrawal, such as the noradrenergic locus ceruleus and the
important in the control of resting membrane potential, cell ex-
memory (Salin et al., 1996). Second, they augment the
mission, a well studied synaptic plasticity implicated in learning and
et al., 2001), including the enhancement of long-term potentia-
motor and postsynaptic mechanisms (Greengard et al., 1991; Ra-
central glutamatergic and GABAergic synapses via both presyn-
cellular cAMP concentration after opioid withdrawal are a widely
activity of these NRM cells during morphine withdrawal.
In addition, the present study provides evidence for another
ular effect of an upregulated CAMP cascade also induced by in vitro
administration of morphine. It shows that morphine with-
drawing augments the $I_h$, likely via an upregulated CAMP pathway
The superactivated CAMP cascade and an overshoot in intra-
cellular CAMP concentration after opioid withdrawal are a widely
distributed molecular adaptation to chronic opioids (Maldonado
however, how such an adaptation in the cAMP pathway leads to
behavioral dependence on opioids and withdrawal syndrome is still unclear, as are the underlying cellular mechanisms for the
chronic opioid-induced behaviors. In cell preparations in vitro,
the CAMP analogs or AC activators generally have two actions on
central neurons. First, they increase neurotransmitter release in
central glutamatergic and GABAergic synapses via both presyn-
aptic and postsynaptic mechanisms (Greengard et al., 1991; Ra-
motor and postsynaptic mechanisms (Greengard et al., 1991; Ra-
mitability, and firing patterns (Ingram and Williams, 1994; Luthi
and McCormick, 1999; Williams et al., 2001; Mellor et al., 2002).
The molecular and cellular mechanisms for CAMP modulation of the
$I_h$ channel (HCN channels) have been characterized in central
neurons (Wainger et al., 2001; Williams et al., 2001).
However, in contrast to these CAMP analog-mediated mecha-
nisms found in vitro, demonstration of the cellular effects of an
upregulated endogenous CAMP cascade induced by in vivo ad-
ministration of opioids in intact animals currently is limited only to
GABAergic synapses. It has been shown recently that opioid
withdrawal increases GABA synaptic transmission through an
upregulated CAMP cascade in the VTA, the nucleus accumbens,
and the periaqueductal gray (PAG) of chronic morphine-treated
rats (Bonci and Williams, 1997; Ingram et al., 1998; Williams et al.,
Interestingly, this chronic opioid effect in all of these
previous reports was observed exclusively on the inhibitory
GABA synapses, with no effect on glutamate synapses. In fact,
chronic morphine treatment has been reported to decrease
NMDA EPSCs in the nucleus accumbens (Martin et al., 1999). In
contrast, our results in the present study suggest that morphine
withdrawal presynaptically increases glutamate synaptic trans-
mission, likely via an upregulated CAMP cascade in $\mu$-receptor-
expressing NRM neurons. The sources of these glutamatergic
terminals presently are unknown. Based on the current results, it
appears that both the removal of the inhibitory effect of
$\mu$-receptors by naloxone or morphine abstinence and the CAMP-
dependent activation of glutamate synaptic inputs increase the
activity of these NRM cells during morphine withdrawal.

The mechanisms for their activation in those pain conditions
mostly are unknown at present. Our previous studies have shown
that these cells also play a critical role in the hyperalgesia induced
by withdrawal from a single dose of acute morphine (Pan et al.,

indicating that withdrawal from chronic morphine increases the
mRNA levels of ACVI and ACVIII and immunoreactivity of
ACV/VI in the NRM are generally in agreement with previous
reports indicating a role for NRM ACV/VI and ACVIII in chronic
morphine-induced synaptic and behavioral adaptations.

Figure 8. Placements of NRM microinjections. Photomicrographs show representative mi-
croinjection sites within the NRM area (A) and dorsal to the NRM (B). Scale bar, 1 mm.
2000; Bie and Pan, 2003). The present study suggests that withdrawal from chronic morphine activates these cells by increasing their glutamate synaptic inputs and by augmenting the excitatory $I_{\text{ex}}$, both likely via an upregulated cAMP pathway after chronic morphine treatment. Thus evidence from this study indicates that activation of these pain-facilitating cells contributes to the pain sensitization during morphine withdrawal. This is in line with previous substantial evidence demonstrating the involvement of glutamate receptors in the development of opioid dependence and the expression of withdrawal behaviors (Trujillo and Akl, 1991; Mao et al., 1995; Christie et al., 1997; Vekovischeva et al., 2001; Varney and Gereau, 2002; Inoue et al., 2003) as well as in withdrawal-induced hyperalgesia (Mao et al., 1995; Urban and Gebhart, 1999).

In summary, the present study shows two adaptive changes in pain-modulating brainstem neurons as a consequence of chronic morphine treatment. It provides possible cellular mechanisms for the behavioral pain sensitization during opioid withdrawal. Understanding the roles of these brainstem neurons and their activation mechanisms in abnormal pain conditions is beneficial to the development of improved therapies for opioid dependence and withdrawal.

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