

Augmented Accumbal Serotonin Levels Decrease the Preference for a Morphine Associated Environment During Withdrawal

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Recent studies have found that acute morphine administration increases serotonin (5-HT) transmission within the nucleus accumbens and other forebrain regions. In contrast, 5-HT transmission is depressed during withdrawal from chronic morphine. We show that pharmacological agents that increase brain 5-HT levels (fluoxetine or 5-hydoxytryptophan, 5-HTP) abolish the preference of chronically morphine-treated, withdrawn rats for a morphine-associated environment. Similar results were seen when fluoxetine was microinjected into the

nucleus accumbens. Conversely, rats given morphine acutely showed an enhanced preference for a morphine-associated environment when pretreated with these agents. Fluoxetine also decreased the heightened anxiety found in morphine withdrawn rats. The results of our study indicate that drugs that augment 5-HT levels may reduce the desire for morphine during withdrawal.

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In addition to gross neurological and somatic signs the abstinence syndrome associated with opiate dependence involves important subjective responses including anxiety, anhedonia, depression and drug craving (Jaffe 1990). The control and elimination of the subjective symptoms, which linger long after the physical symptoms of opiate withdrawal have dissipated, has been a major obstacle to the successful treatment of opiate dependence (Childress et al. 1994). Understanding the alterations in brain chemistry that occur during abstinence from opiate abuse may lead to improved treatments for these symptoms, and greater success in treating opiate addiction.

Acute systemic administration of morphine enhances brain 5-HT synthesis (Garcia-Sevilla et al. 1980) and tryptophan hydroxylase activity (Boadle-Biber et al. 1987). Acute morphine also enhances the turnover and release of 5-HT and dopamine (DA) in widespread areas of the forebrain (Spampinato et al. 1985; Tao and Auerbach 1994). The effect of morphine on 5-HT release appears to be mediated by activation of dorsal raphe neurons (Spampinato et al. 1985; Tao and Auerbach 1994) and may reflect disinhibition caused by direct inhibition of GABA transmission. A similar mechanism has also been proposed for the excitatory effects of opiates on DA neurons in the ventral tegmental area (Bloom et al. 1972; Bonci and Williams 1997).

Neurons in the nucleus accumbens have been shown to be critically involved in the reinforcing properties of opiates (Dworkin et al. 1988; Pettit et al. 1984; Zito et al. 1985). Although morphine administration enhances 5-HT and DA levels within the accumbens (Spampinato et al. 1985; Tao and Auerbach 1995; Wise et al. 1995), the transmitter systems involved in opiate reward are not decisively known. It has been shown that chemical le-

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sions of DA systems or blockade with dopamine antagonists do not affect opiate self-administration (Ettenberg et al. 1982; Gerrits and Van Ree 1996; Pettit et al. 1984) or morphine place preference in animals (Mackey and van der Kooy 1985; however, also see Bozarth and Wise 1981; Smith et al. 1985; Spyraki et al. 1983). On the other hand, chemical lesions of 5-HT terminals in the accumbens impair morphine self-administration without affecting responding for food or water (Smith et al. 1987), and also block the acquisition of morphine conditioned place preference (CPP) (Spyraki et al. 1988).

Withdrawal from opiates decreases both dopamine and 5-HT transmission in the brain. A single dose of morphine induces measurable reductions in both 5-HT and DA levels at 24 hours post-injection (Antkiewicz-Michaluk et al. 1995). Withdrawal from longer chronic opiate treatments produces profound decreases in both dopamine (Acquas et al. 1991; Pothos et al. 1991; Rossetti et al. 1992) and 5-HT transmission (Ahtee 1980; Tao et al. 1998). Reduced 5-HT levels have been linked to both depression (Maes and Meltzer 1995) and compulsive behavior (Dolberg et al. 1996), two disorders that have recently been proposed to play a role in addictive behaviors (Childress et al. 1994; Koob et al. 1998; Markou et al. 1998; O'Brien et al. 1998). Thus, withdrawal-induced reductions in monoamine levels could be responsible for somatic and well as subjective symptoms of opiate withdrawal. Previously, we found that dopamine transmission in the nucleus accumbens regulated somatic signs of opiate withdrawal (Harris and Aston-Jones 1994).

Here we examined the role of 5-HT transmission in subjective signs of opiate withdrawal. In the current series of experiments, we used a conditioned place preference (CPP) paradigm to test the possibility that decreased 5-HT levels during morphine withdrawal may be involved in the withdrawal-enhanced preference for a morphine environment. We hypothesize that after the physical signs of opiate withdrawal have dissipated decreased 5-HT may be responsible for some of the subjective signs of opiate withdrawal that persist. We also propose that the subjective signs of opiate withdrawal in rats could be expressed as an enhanced place preference for morphine and/or increased anxiety. We tested whether 5-HT enhancing agents might decrease the preference for morphine in withdrawing animals using CPP. In the CPP paradigm, drug preference is measured by the amount of time an animal spends in an environment that has been associated with morphine. Environmental drug cues have been shown to be potent stimulators of opiate craving in humans (Ehrman et al. 1992) and CPP models in animals the cue-elicited conditioning that motivates drug-taking behavior (Bardo et al. 1995; Tzschentke 1998). Because CPP training elicits an operant approach response even in the absence of drug, animals can be tested in a morphine-free state without the problems associated with extinction as might occur in a self-administration paradigm when morphine is not present at testing. Furthermore, CPP allows us to address the role of environmental cues associated with morphine withdrawal.

We also tested whether enhancing 5-HT levels in morphine-withdrawn animals would reduce the anxiety associated with morphine withdrawal (Harris and Aston-Jones 1993). For this, we used the conditioned defensive burying paradigm to measure the anxiety associated with drug withdrawal because of its speed, simplicity and reliability (Treit 1985) . This task measures species-specific defensive behavior and, unlike drug discrimination or conflict paradigms, does not require extensive behavioral training of the animals. Furthermore, in contrast to tasks such as the elevated plus maze or the light-dark box, this task contains both active instinctual and passive learning components that can be separately measured.

MATERIALS AND METHODS

Subjects

Male Sprague-Dawley rats (200-250 g) from Harlan (Indianapolis, IN) were used in all experiments. Rats were group housed in accordance with NIH guidelines on a 12-h light/dark cycle with food and water available ad libitum. All animal procedures were approved by the Institutional Animal Care and Use Committees of the Philadelphia Veterans Administration Medical Center and the University of Pennsylvania.

Drugs

Morphine pellets and morphine sulfate powder were provided by the National Institute on Drug Abuse. Fluoxetine was purchased from Tocris Cookson Inc (St Louis, MO) and 5-HTP was purchased from RBI (Natick, MA). Both drugs were dissolved in sterile distilled water. Morphine sulfate was dissolved in sterile saline. All vehicle injections consisted of sterile distilled water, and all drugs were administered via intraperitonal injection.

Chronic Drug Treatment

Rats under halothane anesthesia were implanted subcutaneusly with two 75 mg morphine tablets. Previous studies have shown that morphine pellets are a reliable way to induce physical dependence (Gold et al. 1994; Yoburn et al. 1985). The signs of physical dependence begin to wane around day 14 as the morphine pellets dissolve (Gold et al. 1994). Deprivation withdrawal (a model of abstinence) was induced by removing the pellets after 14 days.

Conditioned Place Preference Procedure

Testing occurred in a Plexiglas apparatus consisting of two distinct compartments separated by a tunnel. One compartment had a grid floor with black walls, and the second compartment had a mesh floor with black and white stripes on the walls. Each compartment was equipped with photocells to automatically record time in each compartment (MED Associates). On the first day (Day 14 of chronic morphine treatment), the rats were allowed to freely explore all of the apparatus for 15 min, and the amount of time spent in each compartment was recorded. None of the animals had an initial bias for either compartment and were randomly assigned to one compartment for morphine conditioning in a balanced design. Conditioning began five days later after all somatic signs of opiate withdrawal in the chronic morphine group had dissipated. On each of the next three days, rats were injected with either saline or morphine (20 mg/kg or 10 mg/kg ip in chronic morphine animals, and 10 mg/kg ip for the acute morphine group) in the morning and afternoon. Rats were confined to one side of the box after each injection by means of an opaque Plexiglas divider for 40 min. Morphine and saline treatments were alternated in morning and afternoon sessions for each animal, so that rats given morphine in the morning were given saline in the opposite compartment in the afternoon, and on subsequent days received saline in the morning, and morphine in the afternoon. One week following conditioning, acute and chronic morphine animals were pretreated with either vehicle, fluoxetine (10 mg/kg or 3.5 mg/kg) or 5-HTP (20 mg/kg), and 20 min later given free access to the apparatus for 15 min. The amount of time spent in each compartment was recorded. As an acute conditioning control, an additional group of six animals was given injections of saline in both compartments and morphine injections (10 mg/ kg) in the home cage.

In a separate set of experiments, chronically morphine-treated withdrawn rats were given one-sided conditioning. Rats in these experiments were treated the same as above, except that one compartment was paired with morphine and the other compartment was only experienced on the preconditioning and test days. In this case the non-paired compartment was relatively novel to the animals and not associated with opiate withdrawal. Fifty-three animals were in the acute morphine groups and fifty-two animals were in the chronic morphine groups.

Intracerebral Injections

Under sodium pentobarbital anesthesia rats were implanted with bilateral chronic indwelling guide cannulae (26 gauge) aimed 2 mm above the nucleus accum-

bens (AP +1.6, ML 3.0, DV 6.3). Cannulae were angled 15° (dorsolateral to ventromedial) to avoid penetrating and allowing injectate to diffuse into the ventricle. The tips of inner injection cannulae (30 gauge) extended 2 mm below the guide cannulae into the injection sites. Control injections were made into the caudate nucleus (AP +1.6, ML 3.0, DV 6.1) in separate animals, 2 mm above accumbens injections. Animals were given one week to recover from surgery before receiving chronic drug treatments. Microinjections (0.5 µl volume on each side over 5 min) of fluoxetine (30 μM) or vehicle (artificial cerebral spinal fluid) were made 20 min prior to the final preference test. Animals were habituated to the procedure by mock injections (injection cannulae did not extend into the injection sites and no fluid was injected) given prior to each conditioning day. Twentythree animals were included in the intra-accumbal groups, with group n = 5-6. Eleven animals had dorsal striatal injections.

Conditioned Defensive Burying Paradigm

As has been described previously for this task (Harris and Aston-Jones 1993; Treit et al. 1981), individual rats were placed in a chamber with an electrified prod inserted through a hole in the chamber wall. Electric current was supplied by a constant current source (MED Associates, 8 mA in amplitude) through two uninsulated silver wires wrapped around the prod. While exploring the prod, rats received one shock, and subsequently buried the prod using bedding material from the floor of the chamber. Burying time was measured for 15 min. Enhanced anxiety was evidenced by significantly shorter latencies to begin burying, as well as by an increase in burying duration, relative to salinetreated animals. Previous studies have validated this procedure to measure anxiety as evidenced by response to classical anxiolytic drugs (Treit 1985; Treit et al. 1981). Chronic and acute morphine animals pretreated with vehicle or fluoxetine were tested in the burying paradigm within 15 min of finishing the preference test. A third group consisted of chronic saline animals treated with vehicle.

5-HTP-Induced Head-Shake Behavior

Six animals from the vehicle-treated chronic morphine group and six from the vehicle-treated acute morphine group were administered 5-HTP (50 mg/kg) 30 min after the preference test. Head shake behavior as described previously (Lucki et al. 1984) consisted of a rapid rhythmic shaking of the head in a radial motion. The total number of head shakes during a 70-min period after drug administration were recorded. One-half of the animals were scored by a blind independent ob-

server. There were no differences between the scores from the two observers.

Histology and Data Analysis

Animals from the intracerebral injection groups were killed with an overdose of sodium pentobarbital. Injections of pontine sky blue dye (0.5 µl) were made at injection sites to allow subsequent histological confirmation. Brains were removed and frozen with methylbutane and sectioned (40 µm). Sections were stained with neutral red to determine the exact location of the injection site. Only animals with injection sites located in the shell region of the nucleus accumbens or dorsal striatum (for the control injection group) were included in the data analysis. Place conditioning data were analyzed by calculating the time spent in the morphinepaired chamber minus the time spent in the salinepaired (or novel) chamber. The resulting difference score was compared between groups. In addition, a within-group measurement of conditioned place preference was assessed by comparing the difference in time spent in the morphine- and saline-paired side (or novel) preconditioning vs. postconditioning. Behavioral data were analyzed using a one-way or two-way analysis of variance. Where necessary post hoc analysis was performed with a Newman-Keuls test.

RESULTS

Place Conditioning

Our paradigm produced a significant preference for the morphine-paired environment in both acutely and chronically treated rats (p < .01; $F_{(1,18)} = 26.65$ for acute, and $F_{(1,15)} = 27.51$ for chronic animals). The acute control group given saline injections paired with both compartments and morphine in the home cage did not show a significant place preference (difference score = $25.8 \pm$ 5.4 seconds, p > .53). A significantly enhanced preference for the morphine-paired environment was seen in rats given prior chronic morphine treatment (relative to the acute morphine group) (Figure 1A; $F_{(1,13)} = 13.14$, p < .01). This enhanced preference was seen in the chronic morphine rats regardless of whether they were conditioned using the 10 mg/kg or 20 mg/kg dose of morphine ($F_{(2,18)} = 6.3$, p < .01, see Table 1). The acute group was found to be significantly different from both chronic groups (p < .05), but the two chronic groups did not significantly differ from each other (p < .16).

When animals from the chronic morphine group with two-sided conditioning were given systemic injections of fluoxetine on the test day, the preference for the morphine environment was reduced (3.5 mg/kg dose, p <.05) or eliminated (10 mg/kg dose, p < .01) ($F_{(2,15)} = 10.13$, p < .01). Similar results were seen following the systemic administration of the 5-HT precursor 5-HTP on the test

day (Figure 1B, $F_{(1,10)} = 93.05$, p < .01). To tell whether fluoxetine decreased an aversion to the saline-paired environment in which morphine dependent animals might have experienced withdrawal, we also used a one-sided conditioning procedure. Fluoxetine (10 mg/kg) was also found to abolished the preference for the morphinepaired environment (Figure 2A) when administered to chronic morphine rats given one-sided conditioning $(F_{(1,14)} = 16.22, p < .01)$. These experiments also revealed that both the one-sided (Figure 1A) and two-sided (Figure 2A) conditioning procedures yielded similar levels of preference for the morphine-paired environment.

Similar experiments were performed with intra-accumbens microinjections of fluoxetine or vehicle to test whether the nucleus accumbens was involved in the ability of 5-HT enhancing agents to decrease morphine preference in the chronic morphine group. Fluoxetine (30 μM) microinjected into the accumbal shell on the test day blocked the preference for the morphine-associated environment in the chronic morphine rats during both two-sided and one-sided conditioning ($F_{(1.9)}$ 13.3, p < .01 for two-sided conditioning, data shown in Figure 2B and $F_{(1,10)} = 12.9$, p < .01 for one-sided conditioning, the means were: Veh = 249 ± 69 and Flu = -4.0 ± 18). These results are similar to the findings obtained above with systemic fluoxetine or 5-HTP treatment. In contrast, fluoxetine had no effect on preference in such animals when injected dorsal to the accumbens along the cannula tract, in the caudate nucleus ($F_{(1.8)}$ = 0.10, p < .75, means Veh = 242 ± 44 Flu = 214 ± 50).

Unlike the effects seen in the chronic morphine animals, similar pretreatments with fluoxetine on the test day in rats acutely treated with morphine caused a significant enhancement in the preference for the morphine-paired environment at both the 3.5 (p < .01) and 10 mg/kg (p < .05) doses (Figure 1A $F_{(2.20)} = 11.2$, p < .05.01). A similar enhancement was seen with 5-HTP administration (Figure 1B, $F_{(1,10)} = 47.47$, p < .01). For the acute morphine-treated rats the increase in brain 5-HT levels caused by fluoxetine or 5-HTP may mimic some of the stimulus cueing properties of morphine, leading the animals to spend more time in the morphine-associated environment. To test this hypothesis, we pretreated acute morphine animals with morphine on the test day (10 mg/kg). These rats also showed an enhanced preference for the morphine environment (Figure. 2C, $F_{(1,10)} = 68.4$, p < .01), resembling the results found for such animals pretreated with fluoxetine or 5-HTP. The exact number of seconds each group in each of the drug conditions spent in the paired chambers can be seen in Table 1.

Head Shake Response

To test the idea that brain 5-HT systems were altered in the chronically morphine treated rats relative to the acute morphine rats, we injected animals with 5-HTP

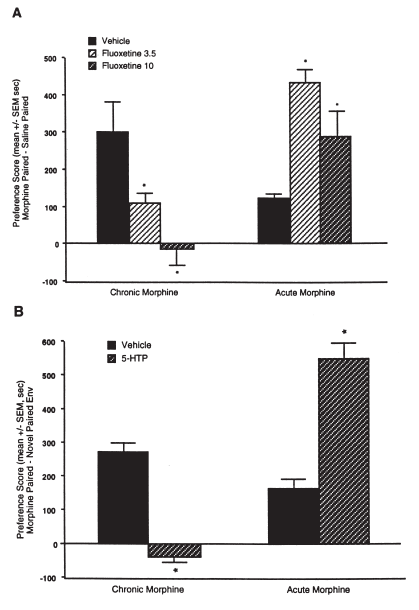


Figure 1. Preference scores for the morphine-paired environment expressed as the time spent in the morphine-paired side minus the time spent in the saline-paired side on the test day. Groups were pretreated on the test day with vehicle, systemic fluoxetine (A, 3.5 and 10 mg/kg, ip), or systemic 5-HTP (B, 20 mg/kg, ip). Fluoxetine given systemically (A) significantly reduced the preference for the morphine-paired environment in the chronic morphine animals at the 3.5 mg/kg dose and abolished preference at the 10 mg/ kg dose. Fluoxetine at both doses significantly enhanced preference in the acute morphine groups (A). Similarly, 5-HTP (B) significantly reduced the preference for the morphine-paired environment in the chronic morphine treated animals but enhanced preference in acute morphine animals. The symbol * indicates a significant difference between vehicle and drug groups (p < .05).

(50 mg/kg) and measured head shake behavior, an indirect measure of brain 5-HT function (Bedard and Pycock 1977; Darmani et al. 1997), for 1 hr. Acutely treated morphine rats showed significantly more head shakes than chronically treated morphine rats (19.2 \pm 2.7 vs. 5.1 \pm 0.36, F_(1,10) = 25.7, p < .01). The number of head-shake responses seen in the acute group was similar to what was seen when the same dose of 5-HTP was given to drug-naive rats (18.5 \pm 1.7, n = 4). Together, these results indicate that 5-HT transmission was depressed in the chronic morphine rats at the time of testing.

Conditioned Burying Response

Chronic morphine treatment significantly decreased the latency to begin burying and significantly increased the duration of burying in vehicle-pretreated rats even when tested one week after the last morphine injection $(F_{(4,31)} = 26.59, p < .01;$ the chronic morphine group was different from all other groups, p < .01). Fluoxetine pretreatment in the chronic morphine group increased the latency to begin burying and produced burying behavior similar to that seen in drug naive animals (Figure 3A). Fluoxetine also significantly reduced the enhanced amount of time that the morphine withdrawn rats buried (Figure 3b). No changes were seen with burying behavior in the acute morphine animals given fluoxetine.

DISCUSSION

The present results indicate that administration of either the 5-HT reuptake inhibitor, fluoxetine, or the 5-HT precursor, 5-HTP, strongly attenuated the preference of morphine-withdrawn subjects for a morphine-paired environment. This effect was specific for rats

Table 1. Place Conditioning Data Expressed as Group Means ± SEM in Seconds

	Chronic Morphine Vehicle	Acute Morphine Vehicle	Chronic Morphine Flu (10)	Acute Morphine Flu (10)	Chronic Morphine Flu (3.5)	Acute Morphine Flu (3.5)	Chronic Morphine Flu (ace)	Chronic Morphine 5-HTP	Acute Morphine 5-HTP
Two-Sided Conditioning									
Morphine 20	489 ± 51		274 ± 14		337 ± 25		289 ± 24	289 ± 18	
Saline 20	186 ± 37		286 ± 18		225 ± 12		234 ± 50	334 ± 10	
Morphine 10	413 ± 23	386 ± 12		487 ± 48		545 ± 27			663 ± 34
Saline 10	229 ± 37	261 ± 10		195 ± 26		107 ± 17			111 ± 15
One-sided									
Conditioning									
Morphine 20	492 ± 42		295 ± 29				245 ± 18		
Novel	209 ± 23		308 ± 40				249 ± 27		

Morphine and Saline 10 and 20 refer to the conditioning dose of morphine in mg/kg and the corresponding saline injections. Numbers listed refer to the number of seconds spent in the morphine, saline or novel paired chamber on the test day. Flu = fluoxetine, 10 and 3.5 = dose in mg/kg, ace = intra-accumbens injections of fluoxetine

given chronic morphine, and did not occur in acutely treated morphine rats. In contrast to the effects seen in the chronic morphine treated rats, the acute morphine rats showed an augmented preference for the morphine environment when pretreated with the 5-HT enhancing drugs. Direct application of fluoxetine into the nucleus accumbens also abolished the preference for the morphine-associated environment in the chronically treated withdrawn animals, indicating that the accumbens is the likely site of action for the effect of fluoxetine. The decrease in preference seen in the chronic morphine rats following the administration of the 5-HT enhancing drugs occurred at a time when 5-HT transmission was reduced, as measured by the 5-HTP-induced head shake response. Fluoxetine was also shown to be effective in reducing anxiety in the chronically morphinetreated, withdrawn rats as measured in a defensive burying paradigm.

The fact that fluoxetine and 5-HTP were both effective in abolishing the preference for the morphine-associated environment in morphine-withdrawn rats indicates that a 5-HT mechanism is involved. Both drugs have been shown to substantially enhance forebrain 5-HT levels for many hours when given in the doses used in this experiment (Guan and McBride 1988; Gudelsky and Nash 1996; Perry and Fuller 1993; Rutter and Auerbach 1993). Fluoxetine blocks 5-HT reuptake to enhance 5-HT levels (Fuller et al. 1991) and 5-HTP is a 5-HT precursor and is decarboxylated in the brain to form 5-HT. As these two compounds enhance 5-HT levels by different mechanisms, it is unlikely that their similarities in effects result from nonspecific actions.

Previous studies have shown that 5-HT enhancing agents reduce the self-administration of opiates (Rockman et al. 1980; Wang et al. 1995). In addition, two research groups have reported that fluoxetine substantially reduces heroin craving in human addicts (Gerra et al. 1995; Maremmani et al. 1992). These studies, to-

gether with the current findings, indicate that enhancing 5-HT transmission in opiate experienced subjects may decrease the desire for morphine.

The enhancement of deficient 5-HT function in the withdrawn animals by fluoxetine and 5-HTP apparently reduced their motivation to spend time in the morphine-paired environment. The 10 mg/kg dose of fluoxetine and the 20 mg/kg dose of 5-HTP completely abolished the preference for the morphine environment in the chronically treated animals. The lower 3.5 mg/kg dose of fluoxetine had a smaller, although significant effect, and did not reduce preference below that of the acute group. The complete suppression of preference by the higher doses of the 5-HT enhancing agents does not appear to be due to a non-specific effect, e.g. a memory deficit, as similar treatment of acute morphine animals did not reduce preference for the morphine environment. The difference in effectiveness of the different doses of fluoxetine may reflect the fact that fluoxetine dose-dependently increases forebrain 5-HT (Guan and McBride 1988; Rutter and Auerbach 1993) and therefore the lower dose may not completely reverse 5-HT changes in the chronic morphine animals. This indicates that the motivation of withdrawn animals to seek out morphine-associated stimuli may be due in part to neurochemical changes involving the 5-HT system. The fact that the morphine withdrawn rats showed significantly reduced head shaking behavior following 5-HTP administration establishes that 5-HT transmission was deficient at the time of preference testing. These findings are consistent with other reports showing evidence for reduced 5-HT transmission during morphine withdrawal (Ahtee 1980; Tao et al. 1998).

The chronic morphine rats showed a much larger preference for the morphine-paired environment than the acutely morphine-treated rats. This effect was seen whether the chronic animals were conditioned with the 10 or 20 mg/kg dose of morphine. This result indicates

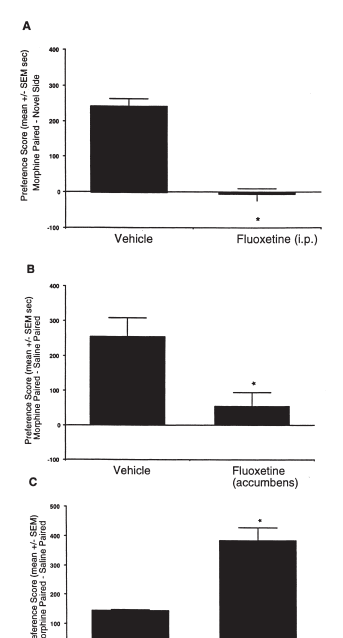


Figure 2. Preference scores for the morphine-paired environment expressed as the time spent in the morphine-paired side minus the time spent in the novel (**A**) or saline-paired side on the test day (**B**, **C**). Groups were pretreated on the test day with vehicle or systemic fluoxetine (**A**, 10 mg/kg, ip in chronic morphine animals given one-sided conditioning), intracranial fluoxetine (**B**, 30 μ M microinjected into the accumbens of chronic morphine animals) or morphine (**C**, 10 mg/kg in acute morphine animals). Fluoxetine given systemically (**A**) significantly reduced the preference for the morphine-paired environment in the chronic morphine animals given one-sided conditioning. Micro-injections of fluoxetine (**B**) into the shell region of the nucleus accumbens of

Morphine (i.p.)

Saline

that the motivation driving preference for the morphine-related stimuli may be different in the chronic and acute morphine conditions. The negative reinforcement associated with alleviation of morphine withdrawal could summate with the positive reinforcing aspects of morphine in the dependent animals. Similar "negative reinforcement theories" of morphine addiction have been proposed previously (Koob and Bloom 1988; Nader and van der Kooy 1996). According to this theory, abstinence following chronic drug treatment produces a negative state (i.e., dysphoria, anhedonia, anxiety, etc.) and this negative state is mentally contrasted to the memories of the euphoric properties of morphine, increasing the likelihood that subjects will be drawn to morphine and morphine-associated stimuli.

Alternatively, other experiments have shown sensitization to the rewarding properties of morphine using the place preference paradigm (Lett 1989; Shippenberg et al. 1996, 1998; Contarino et al. 1997). In those studies as in ours, rats treated chronically with morphine showed enhanced place preference for a morphine-associated environment relative to rats conditioned acutely with morphine. Notably, unlike our experiment, rats were not made physically dependent on morphine. Therefore, the enhanced preference seen in the chronically treated animals in our study may be due to a combination of sensitization to the rewarding effects of morphine as well as the alleviation of the negative symptoms associated with morphine withdrawal.

It has been proposed that the motivational forces underlying morphine addiction may at least in part result from subjective withdrawal symptomatology (Koob and Bloom 1988). Furthermore, it has been hypothesized that the neural substrates that mediate the reinforcing actions of drugs are modified by chronic drug use and subsequently become the mechanisms that produce the anhedonic properties of drug withdrawal (Koob et al. 1989). Previous studies have found that the accumbens is a site that mediates the reinforcing properties of drugs (Dworkin et al. 1988; Pettit et al. 1984; Zito et al. 1985). In this study, the fact that fluoxetine was equally effective in decreasing the preference for the morphine environment in morphine withdrawn subjects when given in the accumbens as when given systemically provides support for the view that the accumbens may also contribute to the anhedonic properties of drug withdrawal.

chronic morphine treated rats significantly decreased the preference for the morphine-paired environment. Morphine (C) given on the test day significantly enhanced preference in acute morphine animals. The symbol * indicates a significant difference between control and drug groups (p < .01).

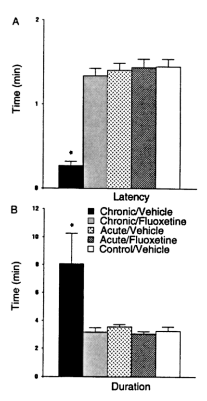


Figure 3. Chronic morphine rats exhibited increased anxiety in a conditioned defensive burying paradigm. This figure shows the burying time latency (A, the time to begin burying after the shock) and duration (B, time spent burying) for chronic morphine rats pretreated with vehicle, chronic morphine rats pretreated with fluoxetine, and chronic saline rats pretreated with vehicle. Enhanced anxiety in chronic morphine treated rats was evidenced by significantly shorter latencies to begin burying as well as a significant increase in burying duration relative to chronic saline-treated animals. The withdrawal-induced increase in anxiety evidenced by burying behavior was blocked by pretreatment with fluoxetine (10 mg/kg). The symbol * indicates a significant difference between chronic/vehicle and all other groups (p < .01).

For the acute morphine rats, there was no long-term change in brain 5-HT systems and no negative state produced. In these rats, the 5-HT enhancing treatments may have resembled the acute effects of morphine, which also enhances 5-HT function (Tao and Auerbach 1994, 1995) thereby cueing the animals to the morphine environment. The fact that similar enhancements in preference were produced by morphine pretreatment on the test day in the acute morphine treated rats supports this possibility. In these rats the 3.5 mg/kg dose of fluoxetine was slightly more potent in producing an enhanced preference than the 10 mg/kg dose. This may reflect a sedative effect that is known to occur at high doses of this drug (Gupta and Masand 1999).

In this study, as in a previous report (Harris and Aston-Jones 1993), morphine withdrawn rats showed pro-

found increases in anxiety in the conditioned burying paradigm. Here, fluoxetine administration markedly decreased this morphine withdrawal-induced anxiety. Similar anxiolytic results for fluoxetine have been shown in rats using the elevated plus maze task (Griebel et al. 1999). In addition, fluoxetine is also known to have anxiolytic effects in the treatment of human anxiety disorders (Nutt et al. 1999). Because anxiety is a potent stimulator of drug cravings in human heroin addicts (Childress et al. 1994), it may also play a role in the enhanced preference for morphine cues seen in the chronic morphine rats.

Serotonin is a potent stimulator of DA release (Benloucif et al. 1993; Parsons and Justice 1993). Although speculative, it is possible that by increasing brain 5-HT levels, the fluoxetine and 5-HTP treatments used here may also stimulate the DA system, thereby enhancing the levels of both transmitters. This increased DA may attenuate some aspects of morphine withdrawal associated with depressed accumbal DA (Harris and Aston-Jones 1994; Pothos et al. 1991; Rossetti et al. 1992). Increased DA release may also contribute to the cueing by fluoxetine and 5-HTP which leads to the enhanced preference for the morphine environment seen in the acute morphine treated rats (discussed above). Although a number of reports have shown that fluoxetine in the doses used in this study has no effect on DA levels in the accumbens (Clark et al. 1996; Ichikawa and Meltzer 1995; Perry and Fuller 1992), one recent study found that similar doses of fluoxetine significantly increased DA levels in the frontal cortex (Gobert et al. 1997). The frontal cortex is one area shown to be hypofunctional during opiate abstinence and withdrawal in human opiate addicts (Danos et al. 1998; Gerra et al. 1998; Krystal et al. 1995) and it is possible that the treatments given in this study may be effective in changing preference by altering activity in this brain region.

The present data indicate that our animals exhibited emotional signs of morphine withdrawal at the time of preference testing that mimic those seen in human addicts, i.e. desire for morphine associated cues and anxiety (Jaffe 1990). Furthermore, these findings indicate that decreased 5-HT function may be associated with these subjective symptoms of morphine withdrawal. To date no treatment for morphine addiction has been satisfactory. Our results indicate that 5-HT enhancing agents may be promising treatments for morphine abuse. Understanding the role of altered 5-HT function in morphine withdrawal could be important in the design of pharmacological treatments for morphine addiction.

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