Morphine Induces Synchronous Oscillatory Discharges in the Rat Locus Coeruleus

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The noradrenergic locus coeruleus (LC) plays a role in opioid dependence and withdrawal. In the present study, using a multiple-electrode recording technique that allowed several LC neurons to be recorded simultaneously over long time periods, LC neuronal activities were recorded before and after intracerebroventricular injection of morphine (26 nmol) under halothane anesthesia. We found that morphine did not simply decrease firing rates of LC neurons, as reported in earlier studies, but that it induced persistent oscillatory discharges in 49% (87 of 178) of the LC neurons recorded. Cross-correlation analysis revealed that almost all LC neurons (86 of 87) that exhibited oscillatory discharges were synchronized with at least one other neuron. When stated in terms of simultaneously recorded neuron pairs, 59% (292 of 492) of the oscillatory neuron pairs discharged synchronously. The morphine-induced synchronous oscillation began at ~10 min after morphine injection, reached its peak in ~20–30 min, persisted throughout the recording periods (up to 110 min after morphine injection, the longest recording time), and were reversed by an opioid receptor antagonist naltrexone. These data suggest that although the overall firing rate of LC neurons was reduced by morphine, the morphine-induced synchronous oscillatory activity may summate temporally and spatially at LC axon terminals and facilitate release of noradrenaline. Noradrenaline is an important neuromodulator and has been shown to induce and facilitate synaptic plasticity at LC target sites. We propose that the morphine-induced long-lasting synchronous oscillatory activity in the LC may be a neuronal signal that could induce synaptic plasticity leading to opioid addiction.

Key words: locus coeruleus; morphine; synchronous oscillation; multiple-electrode recording; noradrenaline; synaptic plasticity

The brain locus coeruleus (LC) is the largest cluster of noradrenergic neurons in the brain and projects broadly throughout the CNS (for review, see Foote et al., 1983). The LC is enriched with opioid receptors (Temple and Zukin, 1987) and plays a role in several effects of opioids, such as opioid dependence and withdrawal (for review, see Nestler et al., 1994; Nestler and Aghajanian, 1997; Zhu et al., 1998).

Systemic or intracerebroventricular administration of opioids, such as morphine, has been shown to have an inhibitory action on spontaneous LC neuronal activity (Korf et al., 1974; Bird and Kuhr, 1977; Aghajanian, 1978; Valentino and Wehby, 1988). These earlier electrophysiological studies have focused on the effects of opioids on the activity of individual LC neurons. Several recent studies indicate that temporal relationships among the activities of LC neurons can also be modulated under certain circumstances. For example, synchronous membrane potential in the LC was found in explant tissue cultures or brain slices prepared from neonatal (Finlayson and Marshall, 1988; Christie et al., 1989; Christie and Jelinek, 1993) and adult rats (Travagli et al., 1995; Ishimatsu and Williams, 1996). Synchronized LC firing was also observed in monkeys performing visual discrimination tasks (Usher et al., 1999). The synchronous activities in the LC have been proposed to have important implications in development (trophic role) (Christie et al., 1989) and regulation of cognitive performance (Usher et al., 1999). However, the effect of opioids on the temporal correlation between LC neurons has not been studied systemically in whole animals. In the present study, using a multiple-electrode recording technique that allowed several LC neurons to be recorded simultaneously over long time periods, we examined the effect of morphine, a classic opioid drug, on the temporal correlation between LC neurons.

MATERIALS AND METHODS

Surgery. All procedures were approved by the Institutional Animal Care and Use Committee at University of Mississippi Medical Center. Adult male Sprague Dawley rats (250–350 gm) were used in this study. A bundle of eight microwires (40 μm wire; NB Labs, Denison, TX) was stereotaxically implanted into the LC under sodium pentobarbital (50 mg/kg, i.p.) anesthesia. A 21 gauge guide cannula was implanted into the lateral cerebral ventricle for drug injection. The microwire bundle and guide cannula were secured in place with four stainless steel screws...
trempanied through the skull and adhered with dental acrylic. Rats were given at least 1 week to recover after the surgery before recording.

**Electrophysiological recordings.** LC neuronal activities were recorded before and after intracerebroventricular injection of morphine solution (26 nmol, 5 μl, in saline) under halothane anesthesia (1.25%, mixed with oxygen). Body temperature was maintained at 37°C with a heating pad. Online isolation and discrimination of neuronal activity was accomplished using a commercial multichannel neuronal acquisition processor (MNAP) system (Plexon, Dallas, TX) that allows one to monitor groups (up to four neurons per wire) of neurons simultaneously. Identifying different neurons on a single wire was accomplished by real-time discrimination of individual waveforms using template analysis procedures provided by the MNAP system. To ensure that neurons recorded by different wires were distinct, we compared the shape of their waveforms, firing rates, and patterns (e.g., interspike interval histograms) before further analysis. LC neurons were identified using previously established criteria, i.e., low spontaneous firing rates, responses to noxious stimuli, and changes in firing rates in response to morphine (Korf et al., 1974; Bird and Kaahr, 1977; Aghajanian, 1978; Valentin and Wehby, 1988). At the end of recording, currents (30 μA, 15 sec) were passed through the microwires to create lesions that were verified histologically.

**Data analysis.** Mean firing rates, autocorrelograms, and cross-correlograms were analyzed using Nex (Nex Technologies, Lexington, MA) and Matlab (Mathworks, Natick, MA) software programs. The degree of oscillation was quantified by an oscillatory index, which was computed as the ratio of the amplitude of the first satellite peak to the offset of the autocorrelogram (König, 1994). The strength of synchrony was quantified by a synchrony index, which was computed as the ratio of the amplitude of the central peak to the offset of the cross-correlogram (König, 1994). The mean firing rates, oscillatory indexes, and synchrony indexes before and after morphine administration were compared using paired t test. Data are presented as mean ± SEM.

**RESULTS**

A total of 408 neurons were recorded from 19 adult rats before and after intracerebroventricular injection of morphine. One hundred and seventy-eight neurons in 10 rats were identified as LC neurons using established criteria (see Materials and Methods). The mean spontaneous firing rates of the LC neurons decreased 48%, from 4.0 ± 0.5 spikes/sec to 2.1 ± 0.3 spikes/sec (n = 178; p < 0.0001; paired t test) 10 min after intracerebroventricular injection of morphine (26 nmol, 5 μl). We found that a subpopulation of the LC neurons (87 of 178; 49%) not only exhibited decreases in their mean firing rates, but also exhibited repeated bursts of discharge activity. The discharge patterns of two representative LC neurons recorded simultaneously are shown in Figure 1. In addition to a decreased firing rate, the two LC neurons exhibited bursts of action potentials 20 min after morphine injection (Fig. 1a). At higher temporal resolution, we observed that the bursts occurred regularly (~90 sec/cycle) (Fig. 1b,c, right panels). The regularity was confirmed by their autocorrelograms, which showed distinctive satellite peaks after morphine injection (Fig. 1d, filled histogram). The degree of regularity was quantified by an oscillatory index. Twenty minutes after morphine injection, 23.3 ± 2.4% (n = 87) of the LC neuronal activity was deemed oscillatory, compared with 0.3 ± 0.1% of the activity (n = 87; p < 0.0001; paired t test) before morphine. The length of the average cycle of the morphine-induced oscillation was 109.3 ± 3.2 sec (n = 87).

An earlier study showed similar burst discharges in the LC after morphine injection in an awake monkey (Aston-Jones et al., 1992). However, because single electrodes were used in that study, it was unable to determine the temporal correlations between LC neurons. In the present study, using the multiple-channel recording technique, we were able to examine if the morphine-induced bursts in LC neurons were temporally correlated. Figure 1, b and c (right panels), shows that the morphine-induced oscillatory bursts were temporally correlated between the two LC neurons that were recorded simultaneously, as indicated by the dotted vertical lines. This apparent synchrony was confirmed statistically by computing the cross-correlation of the discharge activity of the pair of LC neurons. Their cross-correlograms showed a significant central peak (centered at 0.1 msec time lag) (Fig. 1e, filled histogram) after morphine, indicating that this pair of LC neurons discharged synchronously.

In individual rats, several LC neurons (2–21 neurons per rat) that exhibited oscillatory discharges were recorded simultaneously. The degree of synchrony between possible pairs of oscillatory LC neurons in each rat was analyzed. We analyzed a total of 492 neuron pairs from nine rats. Fifty-nine percent of the LC neuron pairs (292 of 492) exhibited synchronous oscillation after morphine. When stated in terms of neurons rather than pairs, however, almost all neurons (86 of 87) were involved in synchronous oscillation with at least one other neuron after morphine. It is not surprising that the percentage of neurons involved in synchrony is larger than that of pairs showing synchrony, because oscillatory LC neurons recorded in individual rats did not always discharge synchronously as a single group. In six rats, the oscillatory LC neurons formed a single synchronous group in each rat. In the other three rats, however, the oscillatory LC neurons formed two or three subgroups in each rat, and synchrony was only present between neurons within the same subgroup. The strength of the synchrony between a pair of LC neurons was quantified by a synchrony index. Twenty minutes after morphine injection, 31.5 ± 2.3% (n = 292 pairs) of the LC neuronal activity was synchronous, compared with 6.4 ± 0.6% (n = 292; p < 0.0001; paired t test) before morphine.

To examine the time course of the morphine-induced synchronized oscillation, the oscillatory indexes of the 87 LC neurons and the synchrony indexes of the 292 LC neuron pairs were computed and averaged every 10 min, 30 min before morphine injection and 110 min afterward (Fig. 2a). The morphine-induced oscillation began at ~10 min after morphine injection, reached its peak in ~20–30 min, and persisted throughout the recording periods (up to 110 min after morphine injection, the longest recording time). The morphine-induced synchrony showed a strikingly similar time course. Moreover, the oscillatory indexes were highly correlated to the synchrony indexes (Fig. 2b) (r² = 0.87). Because there was a strong correlation between the time course of synchrony and that of oscillation in LC neurons (Fig. 2), they may share a common mechanism (Singer, 1993).

In addition to the 87 LC neurons that exhibited both decreases in firing rates and synchronous oscillations, another subpopulation of LC neurons (91 of 178, 51%) exhibited sustained decreases in firing rates, but neither oscillatory discharges nor synchrony after morphine injection. These two subpopulations of LC neurons that responded to morphine in the two different ways were found in almost every individual rat (9 of 10). However, these two subpopulations of LC neurons were indistinguishable in terms of mean firing rates before (4.1 ± 0.5 spikes/sec, n = 87 vs 3.9 ± 0.5 spikes/sec, n = 91; p > 0.5; t test) and 10 min after morphine (2.2 ± 0.3 spikes/sec, n = 87, vs 2.0 ± 0.3 spikes/sec, n = 91; p > 0.7; t test) (Fig. 3). Their mean firing rates did not recover 110 min after morphine.

Effect of systemic administration of morphine on the firing pattern of LC neurons was also examined. Among the 27 LC neurons recorded after intravenous administration of morphine (1.25 mg/kg), 17 of them exhibited both decreases in mean firing rates and synchronous oscillatory discharges. Their oscillatory indexes were 23.6 ± 2.6% after morphine compared with 2.5 ±
0.2% before morphine ($p < 0.001$; paired $t$ test; $n = 17$). Their synchronous indexes were 35.9 ± 2.7% after morphine compared with 1.3 ± 0.04% before morphine ($p < 0.001$; paired $t$ test; $n = 136$ pairs). The other 10 LC neurons showed decreases in firing rates but showed neither oscillatory discharges nor synchrony after systemic administration of morphine. Thus, the systemic administration of morphine produced the same actions as did the intracerebroventricular administration of morphine. These data suggested that the synchronous oscillation observed after intracerebroventricular administration of morphine was not attributable to an unexpected effect of morphine that was injected by the intracerebroventricular route. Furthermore, the synchronous oscillation observed after morphine administration was reversed by an opioid antagonist naltrexone (0.1 mg/kg, i.v.), indicating that the morphine-induced synchronous oscillation is opioid receptor-specific. The oscillatory indexes were 2.2 ± 0.1% after naltrexone compared with 23.6 ± 2.6% before naltrexone ($p < 0.001$; paired $t$ test; $n = 17$). The synchronous indexes were 1.7 ± 0.1% after naltrexone compared with 35.9 ± 2.7% before naltrexone ($p < 0.001$; paired $t$ test; $n = 136$ pairs). In addition, LC neurons exhibited neither oscillation nor synchrony after saline injection (data not shown), suggesting that the observed synchronous oscillation was not attributable to halothane anesthesia or to mechanical disturbance of the tissue caused by the electrodes.

**DISCUSSION**

These results provide the first evidence that central or systemic administration of morphine does not simply decrease firing rates of LC neurons, as reported in earlier studies (Korf et al., 1974; Bird and Kuhar, 1977; Aghajanian, 1978; Valentino and Welbey, 1988), but that it induces long-lasting synchronous oscillatory discharges in a subpopulation of LC neurons. The morphine-induced synchronous oscillation can be reversed by an opioid receptor antagonist naltrexone. These results also suggest that the
responses of LC neurons to morphine were not homogeneous. There was another subpopulation of LC neurons that exhibited sustained decreases in firing rates, but did not exhibit synchronous oscillatory discharges after morphine injection.

There are two possible mechanisms underlying the morphine-induced synchronous oscillatory discharges in the LC. On one hand, electrotonic couplings in the LC, which have been shown in brain slices from neonatal rats (Christie et al., 1989; Christie and Jelinek, 1993) and adult rats (Travaglri et al., 1995; Ishimatsu and Williams, 1996), could mediate the morphine-induced synchronous activity. On the other hand, central administration of morphine could induce synchronous oscillations in other brain areas that provide excitatory inputs to the LC, such as paragigantocellularis (Aston-Jones et al., 1986).

The morphine-induced synchronous oscillatory discharges in the LC may have a powerful influence on noradrenaline release in the widespread LC target areas. A microdialysis study has shown that noradrenaline output in the prefrontal cortex was inhibited initially by morphine injection, but progressively returned to baseline (Rossetti et al., 1993). This phenomenon cannot be simply explained by the changes in the mean firing rate of LC neurons that showed a sustained inhibition after morphine (Fig. 3) (Aston-Jones et al., 1992). However, the progressive return of noradrenaline release could be explained by the morphine-induced synchronous oscillatory discharges reported here. Although the overall firing rate of LC neurons was reduced by morphine injection, as a result of temporal and spatial facilitation, the morphine-induced synchronous oscillatory activity could periodically induce increases in the release of noradrenaline in LC target areas.

Numerous studies have demonstrated that noradrenaline is a modulatory transmitter and can induce and facilitate long-lasting synaptic plasticity in LC target sites (Neuman and Harley, 1983; Lacaille and Harley, 1985; Winson and Dahl, 1985; Hopkins and Johnston, 1988; Huang and Kandel, 1996; Kirkwood et al., 1999; Huang et al., 2000) (for review, see Bailey et al., 2000). For example, a brief application of norepinephrine induced long-lasting potentiation in the dentate gyrus in the absence of tetanic stimulation (Neuman and Harley, 1983; Lacaille and Harley, 1985; Winson and Dahl, 1985). In both the mossy fiber hippocampal pathway and cortico-amygdala pathway, β-adrenergic agonists facilitate the late phase of long-term potentiation (Hopkins and Johnston, 1988; Huang and Kandel, 1996; Huang et al., 2000). Synaptic plasticity has been hypothesized to be involved in mechanisms underlying the development of opioid addiction (for review, see Nestler, 2001, Williams et al., 2001). It has been demonstrated that chronic morphine treatment facilitates long-term potentiation of Schaffer collateral to CA1 synapses in hippocampus (Mansouri et al., 1999). Furthermore, blockades of NMDA receptors or nitric oxide have been shown to prevent the development of opioid dependence (Trujillo and Akil, 1991; Fundytus and Coderre, 1994; Majeed et al., 1994; Dambisya and Lee, 1996; Zhu and Ho, 1998). The opioid-induced synaptic plasticity may not only result from the local effect of opioid on neurons with opioid receptors, but also may result from the indirect effects of opioid. We propose that the morphine-induced persistent synchronous oscillatory discharges in the LC may periodically induce increases in noradrenaline release in the LC target areas, which may facilitate and/or initiate long-lasting synaptic plasticity leading to opioid addiction.
REFERENCES


