OPIATE ALKALOIDS ANTAGONIZE POSTSYNAPTIC GLYCINE AND GABA RESPONSES: CORRELATION WITH CONVULSANT ACTION

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SUMMARY

Opiate alkaloid and opioid peptide actions on spontaneous neuronal activity and postsynaptic amino acid responsiveness were assessed using intracellular recording techniques applied to murine spinal cord neurons in primary dissociated cell culture. Application of opiates was by superfusion and amino acids by iontophoresis. Glycine and GABA but not glutamate responses were antagonized by the opiate alkaloids. Since opiate effects on glycine and GABA responses were not naloxone-reversible, only weakly stereospecific, and not produced by the opioid peptide [d-Ala²]-Met-enkephalinamide, it is unlikely that these effects were mediated by opiate receptors. Opiate depression of glycine inhibition was correlated with the induction of paroxysmal depolarizations in cultured spinal cord neurons, suggesting that antagonism of inhibitory amino acid transmission may underlie the convulsant actions of high concentrations of the opiate alkaloids.

INTRODUCTION

Accumulating evidence indicates that opiates have several distinct inhibitory actions, including presynaptic depression of transmitter release²⁵,³⁰,³⁶, direct postsynaptic hyperpolarization²⁸,³⁹ and postsynaptic antagonism of amino acid responses⁶,¹⁴–¹⁶,²⁸,⁵³,⁵⁵. In addition, opiates have been reported to have excitatory actions⁶,¹¹,¹²,²¹. While several of these actions might be attributable to activation of different classes of opiate receptor, others might be non-specific, involving interaction with sites other than opiate receptors.

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Included among the excitatory actions of opiates is the production of convulsions by the opiate alkaloids\textsuperscript{13,14,20,41,47,48} and peptides\textsuperscript{19,20,46,48}. It is unclear if opiate induction of seizures is mediated by opiate receptors, since large concentrations of morphine are required\textsuperscript{20,47,48} and the resultant seizures are resistant to naloxone antagonism\textsuperscript{20,31,48}. The mechanism of opiate epileptogenesis is also unclear, but two mechanisms have been proposed: (1) presynaptic depression of GABA release\textsuperscript{4,37,54} and (2) postsynaptic modulation of amino acid responsiveness\textsuperscript{6,14}.

Opiates have been reported to antagonize glycine\textsuperscript{6,11,12,15,16,28}, GABA\textsuperscript{14–16} and glutamate\textsuperscript{15–17,23,53,55} effects on neuronal firing rate. Opiate antagonism of glycine- and GABA-induced inhibition of neuronal firing has generally not been naloxone reversible\textsuperscript{12,16}, suggesting the effect is not mediated by opiate receptors. Indeed, an interaction with glycine and GABA receptors rather than opiate receptors is likely, since the opiate alkaloids have been reported to displace \[^{3}H\]strychnine and \[^{3}H\]-GABA bound to synaptic membranes\textsuperscript{14,51}. In contrast, glutamate-induced firing has been reported to be depressed by opiates in a stereospecific\textsuperscript{53} and naloxone reversible\textsuperscript{17,23,53,55} manner, suggesting an opiate receptor mediated effect. However, naloxone did not reverse opiate depression of glutamate responses in several other studies\textsuperscript{16,18,45}.

We investigated opiate actions on spontaneous neuronal activity and postsynaptic amino acid responsiveness using intracellular recording techniques applied to murine spinal cord neurons in primary dissociated cell culture. We report that morphine antagonized postsynaptic glycine and GABA but not glutamate responses. Antagonism of glycine responses was correlated with the induction of paroxysmal depolarizations in cultured spinal cord neurons. Morphine actions on postsynaptic inhibitory amino acid responsiveness and spontaneous activity were not mediated by opiate receptors since the effects were not naloxone reversible, stereospecific, or produced by the opioid peptide \([D-Ala^2]-\text{Met-enkephalinamide}\).

**MATERIALS AND METHODS**

**Cell culture**

Neuronal cultures were prepared by dissecting spinal cords with attached dorsal root ganglia from 12–14-day-old fetal mice\textsuperscript{43}. The tissue was mechanically dissociated to a single cell suspension which was plated on 35 mm collagen-coated culture plates at a density of 1/4 to 1/2 spinal cord per plate. Initial culture medium consisted of 80% Eagle’s minimum essential medium, 10% horse serum, and 10% fetal calf serum. Approximately 4 days after initial plating, 5′-fluoro-2′-deoxyuridine and uridine were added to the cultures to inhibit the division of non-neuronal background cells after which growth medium was modified to 90% Eagle’s minimum essential medium and 10% horse serum. Four-to-twelve-week-old cultures were used in electrophysiological experiments.

**Intracellular recording**

For electrophysiological studies, cultures were placed on the heated stage (35 °C)
of an inverted phase contrast microscope which allowed neuronal impalement by a recording electrode under visual control. Intracellular recordings were obtained using 4 M potassium acetate or 3 M potassium chloride filled micropipettes which had a resistance of 25–50 MΩ. A modified bridge circuit allowed simultaneous current passage and voltage recording by single microelectrodes. Data were recorded on a 6-channel polygraph.

Prior to recording of spontaneous electrical activity, growth medium was poured from the culture and replaced by Dulbecco’s phosphate-buffered saline (DPBS). A thin coat of mineral oil over the DPBS retarded evaporation. Recordings were obtained from each culture over a 3 h period. The initial hour consisted of recording spontaneous activity in control medium. At the end of both the first and second hours, opiates were added to the culture using a microliter pipetter. Five to ten neurons were impaled during each hour.

**Iontophoresis of amino acids during superfusion**

Experiments were performed in DPBS containing 10 mM magnesium ion to suppress spontaneous activity. GABA, glycine and glutamate (all 0.5 M; pH 3.2, 1.8, and 9.1, respectively) were applied by iontophoresis using constant current pulses (500 ms) of 0-40 nA.

Morphine (Mallinckrodt), levorphanol and dextrorphan (Hoffmann-LaRoche), naloxone (Endo Laboratories), and [d-Ala²]-Met-enkephalinamide (Calbiochem) were superfused through the culture at a rate of 0.6 ml/min by a peristaltic pump. Neurons were impaled in control medium, the peristaltic pump was started and the superfusion rate was then unaltered for the duration of the recording. Iontophoretic current was adjusted to obtain amino acid responses of 10–20 mV in control medium with responses evoked every 0.5–1.0 ml of superfusate. Opiate containing buffer was superfused through the culture until 2–3 successive amino acid responses of similar amplitude were obtained after which the opiate was rinsed from the culture. Cells were often held for 3–4 h allowing up to 6 morphine concentrations to be tested on single neurons. The opioid peptide [d-Ala²]-Met-enkephalinamide was dissolved in ammonium acetate buffer (pH = 4.5) and then frozen in aliquots to be used in a single experiment. To prevent loss of peptide during superfusion, the medium contained 0.1% bovine serum albumin and 50 μg/ml bacitracin.

Recording micropipettes were filled with 3 M potassium chloride; thus, chloride ions were injected into neurons, thereby altering the reversal potential for GABA and glycine responses from about --60 to about --20 mV. Since neurons were hyperpolarized to a potential of --80 mV, approximately 60 mV separated membrane potential from GABA and glycine reversal potentials and all amino acid responses were depolarizing.

**RESULTS**

**Opiate alkaloids produced paroxysmal depolarizing events**

Spontaneous activity recorded from neurons in control medium consisted of
randomly occurring action potentials and inhibitory and excitatory postsynaptic potentials (Fig. 1A). Fewer than 10% of the neurons, however, showed a pattern of activity consisting of a burst of action potentials riding on a depolarization greater than 8 mV with a duration of 800 ms to 2 s. The incidence of these paroxysmal depolarizing events (PDE) was increased by morphine (Fig. 1B). Naloxone, administered at 0.2–2 times the concentration of morphine, did not antagonize morphine-induced PDE, although it may have slightly attenuated the amplitude of the depolarizations (Fig. 1C). The opiate alkaloids produced PDE stereospecifically with 100 μM levorphanol inducing PDE in 60% of the neurons (Fig. 1D, Fig. 2), but the

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L(+) enantiomer dextrophan not altering spontaneous baseline activity (Figs. 1E, 2). The opioid peptide [D-Ala²]-Met-enkephalinamide, did not increase the occurrence of PDE at concentrations of 200 nM to 20 μM (Fig. 1F). The incidence of PDE was increased by morphine at concentrations of 10–125 μM in a dose-dependent manner (Fig. 2). Morphine induced PDE in 50% of neurons (ED₅₀), corrected for baseline occurrence of PDE in control medium, at a concentration of 25 μM (Fig. 2). Recordings were obtained from 16 to 27 neurons in two to three cultures at each morphine concentration.

Morphine antagonized glycine- and GABA-mediated postsynaptic responses

Morphine reversibly antagonized postsynaptic glycine (19 of 19 cells) and GABA (12 of 12 cells) responses (Fig. 3A). Glutamate responses (9 cells) were unaltered by morphine concentrations as high as 1.5 mM. Antagonism of glycine responses by morphine occurred over 1.5–750 μM with an ED₅₀ of 35 μM (Fig. 4). Approximately ten-fold higher concentrations of morphine, 75 μM to 1.5 mM (ED₅₀ = 475 μM), were required to antagonize GABA responses (Figs. 3A, 4). No changes in resting membrane potential or conductance were apparent during superfusion of opiate alkaloids through the cultures at concentrations which completely attenuated glycine and GABA response amplitudes.

At a concentration of 20 μM, the opioid peptide [D-Ala²]-Met-enkephalinamide did not affect the amplitude of postsynaptic glycine (n = 8), GABA (n = 6), or...
Fig. 4. Morphine attenuation of glycine and GABA responses was dose dependent. Morphine depressed glycine and GABA responses at concentrations exceeding 1.5 μM (ED$_{50}$ = 35 μM) and 75 μM (ED$_{50}$ = 475 μM), respectively. Glutamate responses were unaffected by morphine over the concentration range tested (1.5 μM to 1.5 mM). Mean data from 3–9 neurons are given for each concentration tested. Bars represent standard error of the mean.

glutamate (n = 8) responses (Fig. 3B). Although opioid peptide actions have been found to desensitize rapidly$^{21}$, it is unlikely that desensitization explains the observed lack of effect by [d-Ala$^2$]-Met-enkephalinamide. Data were obtained from neurons in multiple (5) cultures, and amino acid responses were not antagonized in neurons which had not previously been exposed to the opioid peptide. Furthermore, application of 1.5 mM morphine almost totally reduced GABA and glycine responses in neurons which had been unaffected by [d-Ala$^2$]-Met-enkephalinamide.

To determine if opiate antagonism of glycine and GABA responses was stereospecific, 100 and 500 μM concentrations of both levorphanol and dextrorphan were tested. Determination of relative levorphanol and dextrorphan potency in antagonizing the responses of neurons to glycine and GABA was obtained by testing equal concentrations of the enantiomers during each recording. Levorphanol antagonized glycine responses with potency similar to morphine while dextrorphan was less potent (Fig. 5, 6). Glycine responses were reduced by mean values of 72% by levorphanol and 12% by dextrorphan at enantiomer concentrations of 100 μM (n = 5) (Fig. 6). The effects were dose-dependent with 500 μM concentrations of levorphanol and dextrorphan (n = 3) reducing glycine responses with mean values of 92% and 64%, respectively (Fig. 6). Glycine responses were significantly more reduced by levorphanol than dextrorphan at both 100 μM (paired t-test, P < 0.005, one-tailed) and 500 μM (paired t-test, P < 0.01, one-tailed) concentrations. At 100 μM concentrations, levorphanol was at least 3 times more potent than dextrorphan in attenuating glycine responses (paired t-test, P < 0.05, one-tailed). The difference in the effectiveness of the enantiomers to attenuate glycine responses was not as marked at 500 μM concentrations, since levorphanol concentrations were approached which maximally depressed glycine responses.

In contrast to effects on glycine responses, 100 μM concentrations of levor-
Fig. 5. Opiate alkaloid effects on glycine, but not GABA responses, were weakly stereospecific. The enantiomers, levorphanol and dextrorphan, were tested at concentrations of 100 μM and 500 μM on a single neuron. At 100 μM concentrations, levorphanol but not dextrorphan attenuated the glycine response and neither enantiomer affected the GABA response amplitude. The enantiomers at 500 μM concentrations were equipotent in attenuating GABA responses. The depolarization associated with iontophoresis of glycine during superfusion of 500 μM levorphanol was a coupling artifact, not a glycine response.

Phenol and dextrorphan (n = 4) only minimally attenuated GABA response amplitude and did so with similar potencies; each depressed GABA responses with a mean of 5% (Fig. 6). Levorphanol and dextrorphan at 500 μM concentrations (n = 5) reduced GABA responses with mean values of 65% and 58%, respectively. Thus, the enantiomers were equipotent in reducing GABA response amplitude at 500 μM concentrations (paired t-test, P > 0.1 one-tailed). The effects of levorphanol and dextrorphan at 100 μM concentrations were not tested for a statistical difference since most neurons did not respond to either of the enantiomers at this concentration.

Naloxone was tested for its ability to antagonize depression of glycine responses by morphine. Morphine at 75 μM was chosen since it produced unambiguous but

Fig. 6. Opiate effects on glycine responses were weakly stereospecific. Mean glycine and GABA response amplitudes during superfusion of 100 μM and 500 μM concentrations of levorphanol and dextrorphan expressed as a percentage of response amplitude observed in control medium. Levorphanol was more potent than dextrorphan in depressing glycine-responses at 100 μM (n = 5) and 500 μM (n = 3) concentrations. GABA responses were depressed equipotently by the enantiomers at 100 μM (n = 4) and 500 μM (n = 5) concentrations. Bars represent standard error of the mean. A statistically significant difference in enantiomer potency as determined by a paired t-test is denoted by an asterisk.
Fig. 7. Naloxone did not reverse morphine depression of glycine responses. Glycine response amplitude is plotted as a function of the volume superfused through the culture. Control glycine responses were about 13 mV (1). Superfusion of medium containing 75 μM morphine attenuated glycine response amplitude to about 4 mV (2). Morphine-induced attenuation was not reversed by addition of either 50 μM (3) or 200 μM (4) naloxone to the superfusion medium. Superfusion with control medium restored the control glycine amplitude (5). Application of 75 μM morphine again antagonized the glycine responses (6).

incomplete diminution of glycine responses (mean reduction 70%). Addition of naloxone in concentrations of 50 μM to 1 mM to superfusion medium containing 75 μM morphine did not reverse depression of glycine responses by morphine (n = 4) (Fig. 7). Subsequent rinsing of morphine and naloxone from the culture restored glycine responses to control amplitude.

DISCUSSION

Convulsant actions of opiates

We have demonstrated that opiate alkaloids altered spontaneous neuronal activity, producing large amplitude, long-duration depolarizations with associated action potentials (PDE). Convulsants such as penicillin, pentylenetetrazol, bicuculline, and picrotoxin have been reported to produce a similar pattern of activity in cultured spinal cord neurons29. PDE may be an in vitro equivalent of paroxysmal depolarization shifts (PDS) observed during intracellular recording from neurons in cortical epileptic foci23,33,42 and of bursting recorded from spinal motoneurons following convulsant application44. Therefore, opiate alkaloid-induced PDE in cultured spinal cord neurons might serve as a model of morphine produced convulsions in vivo.

To determine if opiate alkaloids produced PDE by interaction with opiate receptors, the effect was tested for stereospecificity, naloxone reversibility, and agonist properties of the opioid peptides. The production of paroxysmal activity in the culture
was stereospecific, but neither naloxone-reversible nor [d-Ala²]-Met-enkephalinamide-induced. These data indicate that the convulsant actions of opiates on cultured spinal cord neurons were not mediated by opiate receptors and, hence, were non-specific. This contrasts with findings implicating opiate receptors in seizure induction. Intracerebroventricular administration of the enkephalins or morphine has been reported to produce convulsions which were antagonized by high concentrations of naloxone⁴⁹,⁵⁰,⁵ⁱ. Opioid peptides have also been shown to induce naloxone-reversible PDS in hippocampal neurons⁵². However, the opiate alkaloids have been reported to induce seizures by a non-opiate-receptor-mediated mechanism, suggested by the agonist actions of naloxone and the equipotency of opiate enantiomers as convulsants¹⁴,¹⁵. Thus, it is likely that opiate-mediated convulsions are produced by two mechanisms: an interaction of the opiate alkaloids and peptides with opiate receptors and by an action of the opiate alkaloids on neutral amino acid (glycine and GABA) receptors.

Morphine antagonism of glycine and GABA responses

We have found that morphine antagonized postsynaptic glycine and GABA, but not glutamate responses. These data are in agreement with observations in vivo that morphine antagonized glycine⁶,¹¹,¹²,¹⁵,¹⁶,²⁸ and, at higher iontophoretic currents, GABA¹⁴–¹⁶ induced depression of unit firing. In contrast to our findings, morphine and the opioid peptides have been reported to depress glutamate excitatory actions on neuronal firing in a naloxone-reversible¹,¹⁷,²³,⁵³ and stereospecific⁵⁵ manner. Opiate depression of glutamate-induced firing was not reversed by naloxone in several other studies¹⁶,¹⁸,⁴⁵. Our results suggest that opiates do not directly antagonize glutamate action on glutamate receptors, but do not rule out the possibility of an opiate receptor interaction with glutamate receptors.

It is unlikely that antagonism of amino acid responses was mediated by opiate receptors since the effect was not produced by the opioid peptide [d-Ala²]-Met-enkephalinamide, antagonized by naloxone, or highly stereospecific. While levorphanol was three-fold more potent than dextrophan in attenuating glycine responses, the observed difference was 10–100 times less than predicted for an opiate-receptor-mediated phenomenon³,⁵,⁴⁰. Furthermore, concentrations of morphine required for antagonism of glycine- and GABA-mediated responses were higher than expected for an opiate-receptor-mediated effect. High levels of analgesia occur at 1 μM brain concentrations of morphine²²,³⁵, whereas a similar concentration of morphine attenuated glycine response amplitude by less than 10%. In preparations where opiate concentration could be accurately determined and correlated with a physiological action, opiates have been found to exert effects in the low nanomolar concentration range⁴⁸,⁴⁹,⁵⁰,⁵². Therefore, there is a 100–1000-fold difference in potency between morphine antagonism of inhibitory amino acid responses and opiate-receptor-mediated effects. These data are consistent with antagonism of inhibitory amino acid responses underlying opiate-mediated convulsions and not analgesia since 100–1000 times the analgesic dose of morphine is required for the induction of seizures⁴¹,⁴⁷.

Morphine antagonized postsynaptic glycine and GABA responses in the absence
of an effect on resting membrane potential or conductance. Both amino acids increase membrane conductance to chloride ions. Therefore, morphine might simultaneously depress neuronal responsiveness to GABA and glycine by binding to a site which modulates chloride channel conductance. However, such a mechanism is unlikely since GABA and glycine responses were depressed with differing potency by the opiates: the dose response curves for depression of GABA and glycine responses were separated by one log unit, and antagonism of glycine but not GABA responses was stereospecific. Furthermore, opiate alkaloids have been shown to displace \[^{3}H\]GABA and \[^{3}H\]strychnine with low affinity from GABA and glycine receptors\(^{14,51}\), respectively. Thus, it is likely that opiate alkaloids bind to glycine and GABA receptors as do strychnine\(^{10}\) and bicuculline\(^{7,8}\), rather than to chloride channels.

**Antagonism of inhibitory amino acids: correlation with paroxysmal activity**

Morphine produced PDE over a concentration range similar to that producing antagonism of glycine responses (Fig. 8), while morphine dose–response curves for production of PDE and antagonism of GABA responses were separated by one log unit with PDE occurring at lower morphine concentrations. Since GABA and glycine are inhibitory transmitters with a ubiquitous distribution in the central nervous system\(^{9,24,26}\), it is likely that alteration of the actions of these amino acids would have profound effects on neuronal activity. Indeed, a decrease in amino acid mediated postsynaptic inhibition has been proposed as a mechanism of convulsant action\(^{29,34}\).

The good correlation between antagonism of postsynaptic glycine responses and increased PDE in spinal cord neurons was not surprising since glycine is the principal inhibitory transmitter in spinal cord\(^{50}\). Our data suggest that glycine-mediated inhibition must be reduced by approximately 20\% for the production of paroxysmal activity in the culture. At supraspinal levels, GABA is the principle inhibitory transmitter\(^{37}\) and, therefore, the dose–response curve for morphine-mediated PDE might be expected to be shifted to the right.

![Fig. 8. Morphine attenuation of glycine responses and induction of PDE occurred over similar concentration ranges. Glycine responses were approximately 20\% attenuated at threshold morphine concentration for PDE. ED\(_{50}\) for attenuation of glycine responses and induction of PDE were 35 \(\mu\)M and 25 \(\mu\)M, respectively.](image-url)
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