EFFECTS OF SOME NARCOTIC ANALGESICS UPON THE MONOSYNAPTIC REFLEX INHIBITION FROM MUSCULAR AND CUTANEOUS AFFERENTS IN SPINAL CORD OF THE CAT

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It has been known that, at a small dose, narcotic analgesics depress polysynaptic reflex activity in the spinal cats (1) and nociceptive reflexes in the spinal rats (2), as well as in spinal cats and dogs (3-6), where they do not depress, or slightly enhance the monosynaptic reflex (3). In order to explain this depression induced by narcotic analgesics, it has been suggested that the interneurons responsible for mediation of the polysynaptic reflex are suppressed (3). Further, for a remarkable depression of nociceptive reflex discharges from high-threshold cutaneous afferents by morphine and pethidine, it has been suggested that some interneurons activated by such small fibers of the cutaneous nerve might be inhibited (7). On the other hand, large doses of morphine (15-20 mg/kg) has been reported to potentiate the Group Ia inhibition and depress the recurrent inhibition via Renshaw cell arcs in decerebrate cats (8).

However, it remains still unknown whether narcotic analgesics may depress spinal interneurons generally or special interneurons specifically. Therefore, it was attempted in the present study to find either one of these. As a test system, the effects of some narcotic analgesics upon the inhibitory influences to the extensor monosynaptic reflex from flexor reflex afferents were studied using unanesthetized low spinal cats.

METHODS

Twenty-four adult cats were used. All surgical operations were performed under ether anesthesia. After tracheal cannulation, the spinal cord was transected between T13 and L1 during local anesthesia with 1% procaine, then the lumbosacral spinal cord was exposed from L1 to S3. After cutting the ventral roots from L1 to S3, that of L1 or occasionally S1 was placed on the platinum wire electrode, and mono- and polysynaptic reflexes evoked by stimulating the peripheral nerves were recorded from this root. The responses were projected on the cathode-ray oscilloscope (Tektronix 565).

For stimulation, collar-type electrodes were placed to the following nerves of the ipsilateral hindlimb; medial gastrocnemius-soleus, quadriceps, sural, biceps-semitendinosus (BST) and flexor digitorum longus (FDL). In addition, two pairs of the similar electrodes were placed to the saphenous nerve for stimulating at proximal and for recording
at distal point. Electrical shock (0.01-0.1 msec, 0.3-0.5 Hz) was applied to these nerves from an electronic stimulator (MSE-40, Nihon Kohden).

The incoming volleys elicited by the conditioning stimulation of muscle nerves were monitored with a Ag-ball electrode placed on the adequate dorsal root entry zone. The testing monosynaptic reflexes were superimposed on a single frame at various intervals after the conditioning stimulation.

The dorsal root potentials from the quadriceps and saphenous nerves were recorded from L, or S, dorsal root filaments placed on a platinum wire electrode and cord dorsum potentials from these nerves were also recorded with Ag-ball electrode placed on the adequate dorsal root entry zone. These two potentials were projected on a cathode-ray oscilloscope through a preamplifier with a time constant of 0.3 second.

The exposed spinal cord was covered with warmed mineral oil and the temperature of both spinal pool and rectum were maintained between 37-39°C with a thermoregulating device made in our laboratory. Polyethylene catheters were inserted to the contralateral femoral vein for drug administration and femoral artery for monitoring the blood pressure. As soon as surgical operations were completed, ether anesthesia was discontinued and the animal was immobilized with gallamine triethiodide (Gallamine, Teikoku Chemical Industries) and artificially resired (30/min). Experiment was started at least 3-4 hours after discontinuing the anesthesia.

The drugs used in this report were as follows; morphine hydrochloride (Sankyo), fentanyl citrate (N-(1-phenethyl-4-piperidinyl) propionanilide dihydrogen citrate, Jansen Pharmaceutical Co.), oxymorphone (Sankyo), dimethylthiambutene (Ohton, Ono Pharmaceutical Co.) and nalorphine (Sankyo). Drugs were dissolved in physiological saline and diluted to 1 % (morphine, dimethylthiambutene, oxymorphone and nalorphine) and 0.008 % (fentanyl) in each and were administered intravenously slowly so as not to induce the blood pressure fall.

RESULTS

Effects of fentanyl and morphine upon the cutaneous inhibition of the extensor monosynaptic reflex

As was previously known (11) and shown in Figs. 1 and 2, the monosynaptic reflex (MSR) evoked by the stimulation of the medial gastrocnemius-soleus nerve (mG-S-MSR) was inhibited by the preceding conditioning shock to either the saphenous or sural nerve at a supramaximal intensity (cutaneous inhibition). This inhibition began about 5 msec after the shock, attained the maximum at about 10-15 msec and gradually decayed. It took about 30-40 msec until the complete recovery (Fig. 1A, C). When either fentanyl, a potent narcotic analgesic (9, 10) (40 μg/kg, Fig. 1B), or morphine (3 mg/kg, Fig. 1D) was administered intravenously, the duration of the cutaneous inhibition was shortened from about 25 msec to about 15 msec. However the onset latency and its peak time were almost un influenced. In most cases the rebound facilitation appeared after the cutaneous inhibition (Fig. 1B), the MSR often being twice as large as the control.
Fig. 1. Effects of fentanyl and morphine upon the cutaneous inhibition. Monosynaptic reflex of the L7 ventral root was evoked from the medial gastrocnemius-soleus nerve (0.01-0.1 msec, 10 V) at various intervals after the conditioning shock to the saphenous nerve at a supramaximal intensity for high threshold cutaneous afferents (0.1 msec, 50 V). A and C represent the control state for the B and D, respectively. B: 5 minutes after fentanyl administration (40 µg/kg), D: 5 minutes after morphine injection (3 mg/kg). Arrow in A indicates a timing of the cutaneous nerve shock. Time, 10 msec.

The effect of fentanyl (40 µg/kg) disappeared in 40-60 minutes after administration (see below). When the same amount of fentanyl was injected after the complete recovery, the blockade of the late phase of the cutaneous inhibition was again observed with a similar time course and potency as with the previous injection. Thus, it is clear that the late phase of the cutaneous inhibition is depressed reversibly by a small dose of fentanyl and therefore it is considered that there is little, if any, tachyphylaxis upon repeated administration of fentanyl.

These effects were confirmed without exception in 29 trials in 13 cats with fentanyl (12 trials with 80 µg/kg, 10 trials with 40 µg/kg, 3 trials with 20 µg/kg, 2 trials with 8 µg/kg and 2 trials with 0.8 µg/kg) and 21 trials in 21 cats with morphine (one trial with 10 mg/kg, 5 trials with 5 mg/kg, 11 trials with 3 mg/kg, 2 trials with 1 mg/kg and 2 trials with 0.5 mg/kg).

As already known, this inhibition is evoked from low and high threshold afferents of cutaneous nerves, presumably from fibers of Group Aα and Aδ range (11-13).

Fig. 2A shows the time course of the cutaneous inhibition when the saphenous nerve was stimulated at a supramaximal intensity for the high threshold fibers, together with two components of the nerve potential (Aα and Aδ) recorded from the distal end (Fig. 2A, insetted figure) (13). The onset latency of the first component of the nerve potential was about 0.3 msec and that of the second was about 1.0 msec. Since the distance from the stimulating electrode (cathode) to the recording one was about 20 mm, the calculated values for the conduction velocity of these fibers were about 60 and 20 m/sec, respectively.
In order to determine the nerve component concerned in the cutaneous inhibition, MSR at 12 msec (B) and 20 msec (C) after a conditioning shock of varied intensity were plotted in Fig. 2B and C, respectively, against the shock intensity together with the nerve potential size. The Aα potential reached its maximum at an intensity of about 2 times and Aδ component appeared at an intensity of about 1.8 times and reached maximum at about 2.5 times the threshold for Aα fibers (Fig. 2B, C). As shown in Fig. 2B, the mG-S-MSR at 12 msec after conditioning stimulus started to be depressed at near threshold intensity and depression became maximum at 1.5 times of the threshold. Whereas in

![Fig. 2. Cutaneous inhibition sensitive to fentanyl and morphine.](image-url)
C, the inhibition observed at 20 msec after conditioning stimulus developed not smoothly but with a break at about 1.7 times of the threshold. The MSR inhibition was about 20% up to this point, and then developed more, reaching a plateau level at about 2.3 times of the threshold. This breaking point roughly corresponded to the threshold intensity for the Aδ elevation, and the secondary increase of the inhibition paralleled with the development of Aδ elevation of nerve action potential. Thus, it is clear that the MSR depression is induced by both Aγc and Aδ components of the cutaneous fibers and further the later part of the depression is produced when the shock intensity is more than 2 times of the nerve threshold, presumably by Aδ fibers.

In Fig. 2D and E, drug effects upon the cutaneous inhibition were compared at a low (1.6 times of Aγc threshold) and a high (4.0 times of Aγc threshold, supramaximal intensity for Aδ fibers) conditioning shock intensities. In the control with a low intensity conditioning shock the cutaneous inhibition was observed from about 5 msec to 20 msec after the conditioning and followed by a rebound facilitation (Fig. 2D, open circles), while with a high intensity one, the cutaneous inhibition was prolonged to about 35 msec as shown in A and E (open circles). After administration of fentanyl (40 μg/kg), the time course of the cutaneous inhibition at a weak conditioning stimulation was little affected (Fig. 2D, crosses), whereas that at a supramaximal intensity was markedly shortened from about 30 msec to 15 msec in duration (Fig. 2E, crosses). It is noteworthy that the remaining inhibition after fentanyl is almost the same regardless of the intensity of the conditioning stimulation of the saphenous nerve, and followed by a rebound facilitation.

Morphine showed similar effects. As illustrated in Fig. 2F in another preparation, a small dose (1.0 mg/kg) of morphine blocked the late phase of the cutaneous inhibition and the total duration was shortened from about 25 msec to 15 msec, and the rebound facilitation appeared.

This morphine effect—shortening of the total duration of the MSR depression—was immediately abolished by nalorphine (0.5 mg/kg, filled circles).

In some cases, the MSR was also recorded from S1 ventral root. The time courses of the cutaneous inhibition in this ventral root with low and high conditioning shock were similar to those recorded in L7. A large dose of fentanyl (80 μg/kg) often blocked not only the late phase but also the early phase of the cutaneous inhibition recorded in S1 ventral root, in contrast to the selective block of the late phase in L7. However, relatively small doses of fentanyl (less than 40 μg/kg) and morphine (less than 10 mg/kg) blocked the late phase of the cutaneous inhibition selectively in this root as well.

**Time courses of fentanyl effect at various doses upon the cutaneous inhibition**

Time courses of the effect of fentanyl at various doses upon the cutaneous inhibition were studied. Testing and conditioning shocks were given at a fixed interval of 15 msec. In Fig. 3, three different doses, 40, 8 and 0.8 μg/kg, were sequentially administered in a spinal cat. The second and the third injection of fentanyl were applied after the complete recovery of the cutaneous inhibition, i.e., 2 hours and one hour respectively after the previous injection. In control, the cutaneous inhibition was about 65%. Forty μg/kg
of fentanyl began to depress the cutaneous inhibition immediately after the administration and blocked it completely in less than one minute. The MSR was rather enhanced (two filled circles under abscissa of Fig. 3) by the conditioning stimulation between 1.5 minutes and 8 minutes after administration. Thereafter, this fentanyl effect gradually decayed and the inhibition recovered in about 40 minutes (filled circles). When a dose of 8 \( \mu \text{g/kg} \) was administered about 2 hours later, the cutaneous inhibition was suppressed, though weak, with a similar time course. Peak of the fentanyl effect was seen at about 5 minutes after administration. One hour later, 0.8 \( \mu \text{g/kg} \) was injected. The depression was slight, as much as about 7% (open circles).

Effects upon the MSR inhibitions from other afferent inputs

A. Group Ia and Ib inhibition: The Group Ia inhibition of BST-MSR by the conditioning stimulation of the quadriceps nerve was examined at an intensity less than 2 times of the nerve threshold. In accordance with the literature (14), the onset latency, peak time and duration of this inhibition were about 2, 3 and 1.5 msec, respectively. This inhibition was followed by MSR facilitation which is considered to be due to Group Ib afferent fibers (15). To these inhibition and facilitation of the MSRs, neither morphine (3 mg/kg) nor fentanyl (40 \( \mu \text{g/kg} \)) induced any visible change.

When the quadriceps nerve was stimulated at an intensity 1.15 times the nerve threshold, the mG-S-MSRs were inhibited over a period from about 0.7 msec to 35 msec, with the maximal inhibition at about 8 msec after the peak of the nerve potential evoked by the conditioning stimulation and recorded at the dorsal root entry. This inhibition of mG-S-MSR is considered primarily due to the Group Ib and II afferent fibers (15). Another Group Ib inhibition of mG-S-MSR was obtained by the conditioning stimulation.
of FDL nerve at an intensity less than 2 times the nerve threshold (15). As shown in Fig. 4G for the quadriceps nerve stimulation, neither fentanyl (40 μg/kg, filled circles) nor morphine (5 mg/kg, crosses) caused a significant effect on these Ib and II inhibitions. This was confirmed in 6 trials with quadriceps nerve stimulation and 7 trials with FDL nerve stimulation.

B. High threshold muscle afferents: When the quadriceps nerve was conditioned at supramaximal intensity, the mG-S-MSR was inhibited over a period from about 5 msec to more than 50 msec with a marked inhibition between about 7 msec and 30 msec (Fig. 4A). Effects of fentanyl and morphine were investigated in 15 cats. Although fentanyl (40 μg/kg) did not induce any change in the latency and the maximal inhibition of the MSR (40% of control in amplitude), it shortened the duration of the inhibition remarkably to about 25 msec, the maximal inhibition being between 7 and 20 msec (Fig. 4B). This effect disappeared almost completely in about 40 minutes after administration (Fig. 4C). One hour later, after taking the control record shown in Fig. 4D, 3 mg/kg of morphine was administered. Reduction of the duration of the depression was similarly obtained (E), which was antagonized by nalorphine (0.5 mg/kg, F).

C. Recurrent inhibition: The mG-S-MSRs recorded in split half of L7 ventral root were inhibited immediately after the conditioning shock of the other split half of the

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**Fig. 4.** Effects of fentanyl and morphine upon the inhibitory pathways from Group Ib, Renshaw and high threshold muscle afferents to the medial gastrocnemius-soleus monosynaptic reflex.

Quadriceps nerve was stimulated for a conditioning at supramaximal intensity (A-F) and at an intensity 1.15 times the nerve threshold (G). H: split half of the L7 ventral root was conditioned and the MSR evoked by the medial gastrocnemius-soleus nerve shock was recorded from the other half of split L7 ventral root. A-C depict effect of fentanyl and D-F that of morphine. A and D: control, B: 2 minutes after fentanyl (40 μg/kg), C: 60 minutes after fentanyl (40 μg/kg) administration. E: 3 minutes after morphine (3 mg/kg). F: 2 minutes after nalorphine (0.5 mg/kg) given at the top of morphine effect, about 10 minutes after morphine. Note the restoration of prolonged inhibition in C and F. Time, 10 msec.

Open circles in G and H represent control state, filled circles during fentanyl effective state, (3-6 minutes after administration of 40 μg/kg), and crosses during morphine effective state (3-6 minutes after administration of 5 mg/kg).

Ordinates: amplitude of the monosynaptic reflex relative to the control size. Abscissae: conditioning-testing intervals in msec.
L7 ventral root at a supramaximal intensity for α fibers. This inhibition reached maximum at about 10 msec and continued for about 50 msec (Fig. 6H). This time course was exactly the same as previously reported by Renshaw (16). Neither fentanyl (40 μg/kg, crosses) nor morphine (5 mg/kg, filled circles) induced any change in it.

**D. Dorsal root potentials:** In order to see the effects upon the presynaptic inhibitory mechanism which could possibly explain the observed MSR depression by the conditioning stimulation, effects of morphine, fentanyl upon the dorsal root potentials were investigated. The dorsal root potentials were recorded from L7 or S1 dorsal root filament upon stimulation of the quadriceps and the saphenous nerves at an intensity either supramaximal or less than 2 times the nerve threshold. With a supramaximal shock to the muscle nerve dorsal root potentials with onset latency of about 5 msec, peak time of about 40 msec and duration of about 90-100 msec were observed. With a supramaximal stimulation to the cutaneous nerve, the dorsal root potentials of the similar time course were obtained. When the stimulus intensity was weakened (about 1.5 times of the nerve threshold of the muscle nerve fibers), the amplitude became smaller. Cord dorsum potentials were also recorded simultaneously. In these potentials evoked by stimuli of either intensities, neither morphine (5 mg/kg) nor fentanyl (40 μg/kg) produced any change. Although excitability of the terminals of cutaneous afferents, of low threshold and high (threshold), as well, was not examined, it seems that presynaptic mechanism is not involved so that the drug may influence the dorsal root potentials. Therefore, this would not be a primary cause for the observed MSR depression by the conditioning stimulation of these nerve fibers.

**Effects of other narcotics upon cutaneous inhibition**

From the results described above, it was shown that a small dose of both fentanyl and morphine blocked the late phase of the cutaneous inhibition. Attempts were made to see possible effects of other narcotic analgesics upon the cutaneous inhibition. It has been reported that dimethylthiambutene (17) has analgesic actions as potent as morphine and oxymorphone has that about 10 times as potent as morphine (18). Intravenously administered oxymorphone (0.25 mg/kg) and dimethylthiambutene (5 mg/kg) were also found to block the late phase of the cutaneous inhibition similarly to fentanyl and morphine.

**DISCUSSION**

In the present experiments, such narcotic analgesics as morphine, fentanyl, dimethylthiambutene and oxymorphone showed a common characteristic effect at small doses of specifically blocking the late phase of the MSR inhibition from cutaneous and muscular nerves. This portion of MSR inhibition is presumably due to an excitation of an Aβ component of the cutaneous nerves and high threshold muscle afferents. It is considered that C component of the afferent fibers is not involved in the cutaneous inhibition described above, since it should take more than 100 msec until the afferent volley of C fibers reach the entrance of the spinal cord, and the cutaneous inhibition described in this paper was observed over a period from about 5 to 30-35 msec after the cutaneous
nerve stimulation.

It is clear from the present results that these narcotic analgesics at small doses do not depress spinal interneurons generally but act specifically at some neurons activated directly or indirectly by such small fibers. As for the mode of action concerned in the abolishment of the MSR inhibition produced by these small fibers, there may be at least two possibilities; 1) spinal interneuronal chain activated by small fibers and having inhibitory influences to the mG-S motoneurons may be blocked by these narcotic analgesics, 2) spinal interneuronal chain activated by small fibers and having excitatory influences to mG-S motoneurons may be facilitated by these drugs. However, there is no evidence available for selecting either one of these possibilities at present. In view of our present results, it is interesting that morphine reportedly inhibits activity of some dorsal horn cells which are repetitively activated after a long delay by a single shock to dorsal roots at supramaximal intensity (21) and small doses of morphine and pethidine depress the nociceptive reflex discharges evoked by the stimulation of Aδ, post Aδ and C fibers (7).

The blocking action of the late phase of the cutaneous inhibition by fentanyl was seen at a dose more than 0.8 μg/kg (see Fig. 4) and the blockade by morphine was seen at a dose more than 0.5 mg/kg. According to Janssen (9) fentanyl had analgesic action 100-300 times as potent as morphine in Haffner’s method. This was confirmed by us, i.e., fentanyl is about 90 times as potent as morphine in Haffner’s method in mice and about 300 times as potent in tooth pulp stimulation of chronic rabbit (22). In the present study, the minimal dose of fentanyl for the blockage of the late phase of the cutaneous inhibition was about 1/600 of that of morphine. The difference of the dose between morphine and fentanyl in this blocking action may be due to that of the sensitivity of neurons. The effective doses for this blocking action of other narcotic analgesics tested also ranged in their respective analgesic doses (17, 18). This quantitative accordance suggests that the underlying mechanism for the block of MSR inhibition may be in common with that for the analgesic action.

Jurna et al. (19) demonstrated that morphine and pethidine depressed the post-tetanic potentiation of the spinal monosynaptic reflex and also repetitive discharges of α-motoneurons caused by stretching of extensor muscles (20) and suggested that these narcotic analgesics may act at presynaptic site. However, in our present experiments, the dorsal root potentials evoked by both weak and supramaximal stimulus intensity were not influenced and accordingly presynaptic mechanism at the primary afferent terminals appears uninfluenced by either fentanyl or morphine.

It was reported that morphine depressed the release of acetylcholine not only in isolated guinea pig ileum (23, 24) but also in the cat brain (25-28). However, the recurrent inhibition via Renshaw cell arcs, relevant transmitter being considered as acetylcholine (29), was almost unchanged in our experiment by either morphine (5 mg/kg) or fentanyl (40 μg/kg). Kruglov (8) previously demonstrated that the recurrent inhibition was little influenced by intravenous administration of morphine at a single dose of 5 mg/kg, but it was gradually inhibited by adding the dose up to 15-20 mg/kg. At a small dose (0.5-5
mg/kg) by which the late phase of the MSR inhibition from both Aδ of cutaneous nerves and high threshold muscle afferents was blocked, the recurrent inhibition was uninfluenced. From this, it appears that a small dose of morphine does not cause the reduction of the release of acetylcholine in lumbosacral spinal cord sufficient to influence the Renshaw inhibition. Therefore, a possibility that a cholinergic mechanism is involved in the late phase of the cutaneous inhibition may be excluded.

**SUMMARY**

The effect of fentanyl and morphine upon the MSR depression induced by the low and high threshold cutaneous and muscular afferents in spinal cord were investigated in unanesthetized low spinal cat. Morphine (0.5–10 mg/kg, i.v.) and fentanyl (0.8–80 μg/kg, i.v.) reduced the inhibition of the spinal monosynaptic reflex from the medial gastrocnemius-soleus nerve by a conditioning shock to Aδ fibers of the cutaneous nerve and high threshold muscle afferents. While no significant influences were observed upon the MSR inhibitions from low threshold cutaneous and muscle nerves, recurrent collaterals, and on the dorsal root potentials.

Other narcotic analgesics tested, oxymorphone (0.25 mg/kg, i.v.) and dimethylthiambutene (5 mg/kg, i.v.), also depressed the MSR inhibition due to Aδ fibers of the cutaneous nerve.

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