Antiallodynic Effect of Thalidomide and Morphine on Rat Spinal Nerve Ligation-induced Neuropathic Pain

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Background:

Tumor necrosis factor-alpha and other proinflammatory cytokines are becoming well recognized as key mediators in the pathogenesis of many types of neuropathic pain. Thalidomide has profound immunomodulatory actions in addition to their originally intended pharmacological actions. There has been debate on the analgesic efficacy of opioids in neuropathic pain. The aim of this study was to investigate the effect of thalidomide and morphine on a spinal nerve ligation model in rats.

Methods:

Male Sprague-Dawley rats weighing 100−120 g were used. Lumbar (L) 5 and 6 spinal nerve ligations were performed to induce neuropathic pain. For assessment of mechanical allodynia, mechanical stimulus using von Frey filament was applied to the paw to measure withdrawal threshold. The effects of intraperitoneal thalidomide (6.25, 12.5, 25 and 50 mg/kg, respectively) and morphine (3 and 10 mg/kg, respectively) were examined on a withdrawal threshold evoked by spinal nerve ligation.

Results:

After L5 and 6 spinal nerve ligation, paw withdrawal thresholds on the ipsilateral side were significantly decreased compared with pre-operative baseline and with those in the sham-operated group. Intraperitoneal thalidomide and morphine significantly increased the paw withdrawal threshold compared to controls and produced dose-responsiveness.

Conclusions:

Systemic thalidomide and morphine have antiallodynic effect on neuropathic pain induced by spinal nerve ligation in rat. These results suggest that morphine and thalidomide may be alternative therapeutic approaches for neuropathic pain. (Korean J Pain 2010; 23: 172-178)

Key Words:
antinociceptive, cytokine, morphine, neuropathy, thalidomide.
INTRODUCTION

Neuropathic pain can be defined as pain caused or initiated by a primary lesion of the peripheral or central nervous system, manifesting with sensory symptoms and signs [1]. It is associated with allodynia, hyperalgesia and spontaneous pain with varying intensity, in which pain could be so severe that daily activities like wearing clothes might be difficult. Neuropathic pain can also decrease quality of life, cause patients to withdraw socially and emotionally, and cause problems with mood and sleep. Moreover, unrelieved pain inhibits respiration, decreases mobility, and decreases patient functional status, which may lead to iatrogenic events, including pneumonia, constipation, and deep vein thrombosis [2].

Medications primarily used for neuropathic pain generally fall into the categories of anticonvulsants, antidepressants, opioids, and topical agents. The current treatment options, however, are not effective enough to make the pain endurable and the response rate has been estimated at a 30% to 50% reduction in pain in up to 50% of patients [3].

Recently, accumulating evidence has demonstrated that both microglia and astrocytes release a variety of proinflammatory cytokines including interleukin (IL)-1β, IL-6, and tumor necrosis factor (TNF-α), which play a major role in mediating and/or maintaining hyperalgesia and allodynia in inflammatory, as well as neuropathic pain [4-6].

Thalidomide, a derivative of glutamic acid, was developed initially as a sedative and used for morning sickness until 1961 when its use was banned due to its association with major birth defects. It reappeared in clinical practice because of its immunomodulatory properties. Furthermore, investigations into the mechanisms underlying thalidomide’s pharmacotherapeutic action have increased [7,8], with its effects on the production and release of TNF-α being of notable importance. Thalidomide has been used in several cutaneous inflammatory disorders (erythema nodosum leprosum in lepromatous leprosy, cutaneous lupus erythematosus, severe aphthosis), cancers (relapsed/refractory multiple myeloma) and inflammatory conditions [9]. Previous studies have shown that thalidomide inhibits the inflammatory hyperalgesia in rats and the writhing nociceptive response in mice [10,11]. Thalidomide was found to be effective in animal models of neuropathic pain, such as chronic sciatic nerve injury (CCI) and Lumbar (L) 5 ventral root transection (VRT) neuropathy [12,13]; however, several human case reports and series have demonstrated mixed results, with some patients having dramatic responses, especially in chronic intractable conditions such as complex regional pain syndrome [14,15].

Opioids are among the most powerful analgesics in clinical use for the treatment of nociceptive pain, but data concerning their effectiveness in neuropathic pain has been controversial [16,17]. However, evidence now suggests that opioids are effective in relieving some pain symptoms in patients with neuropathic pain of primarily peripheral origin [18,19].

In the present study, we investigated the antiallodynic effects of morphine and thalidomide in a spinal nerve ligation rat model.

MATERIALS AND METHODS

1. Animal preparation

All experiments followed the Guidelines on Ethical Standards for Investigation of Experimental Pain in Animals [20]. Male Sprague–Dawley rats weighing 100–120 g were used in all experiments. Animals were acclimated to the laboratory environment for 5–7 days before being used in the study. While in the home cage environment, animals were fed a standard rat diet and tap water. Room temperature was maintained at 20–23°C with a 12:12 h light/dark cycle.

2. Animal model of neuropathic pain

Neuropathic pain was evoked by spinal nerve ligation of experimental rats, as previously described [21]. Briefly, the left L5 and L6 spinal nerves of rats were isolated adjacent to the vertebral column during sevoflurane anesthesia and tightly ligated with a 6-0 silk suture distal to the dorsal root ganglia. Care was taken to avoid injury of the L4 spinal nerve. Animals were considered to be in neuropathic pain when they exhibited mechanical allodynia i.e., paw flinching behavior response to the application of a bending force of less than 4 g. Sham rats underwent identical surgical procedures at the same level, but the spinal nerves were not ligated. Rats showing motor impairment after surgery were killed immediately using overdose of sevoflurane.

3. Drugs

The following drugs were used in this study: thalido-
mide (Sigma Chemical Co., St. Louis, MO, USA) and mor-
mphine sulfate (Sigma Aldrich Co., St. Louis, MO, USA).
Thalidomide was dissolved in dimethyl sulfoxide (DMSO),
and the solution was then diluted with saline to reduce the
concentration of DMSO to 10%. Morphine was dissolved in
normal saline. Both agents were intraperitoneally adminis-
tered. The dose of thalidomide and morphine were decided
based on previous studies using neuropathic pain models.

4. Assessment of mechanical allodynia

Paw withdrawal threshold (PWT) in response to me-
chanical stimulation was measured using the up and down
method [22] by applying calibrated von Frey filaments
(Stoelting, Wood Dale, IL, USA) from underneath the cage
through openings in the mesh floor to the hindpaw. Rats
were placed in a separate transparent plexus-glass cham-
ber with a wire mesh floor underneath and acclimated to
the test chambers for 30 min. A series of eight von Frey
filaments (0.4, 0.7, 1.2, 2.0, 3.6, 5.5, 8.5 and 15 g) were
applied vertically to the plantar surface of the hindpaw for
5 s while the hair was bent. Brisk withdrawal or paw
flinching was considered as positive responses. If rat
showed the positive response the less stiff filament was
used for the next trial, or the stiffer one for the case with
no withdrawal or licking. The cut-off value was determined
as 15 g, where the PWT was 15 g if the rat did not show
any withdrawal or licking response to the application of a
15 g von Frey filament. Rats which did not show allodynia
(less than 4 g) were excluded from this study.

5. Experimental paradigm

On the day of testing (day 10 after spinal nerve liga-
tion), rats were allowed to acclimate to the testing cham-
ber for 30 min then allocated into experimental and control
groups for the tested drugs. Control studies were per-
formed using intraperitoneal saline or DMSO according to
the solvent used for the tested drug. Animals were tested
only once in this study, and the investigator was blind to the
drug given. The effects of thalidomide (6.25, 12.5, 25,
and 50 mg/kg) and morphine (3 and 10 mg/kg) were inves-
tigated in the neuropathic pain state, and withdrawal
threshold was determined at 30 minutes after intra-
peritoneal administration of experimental drugs.

6. General behavior

For the purpose of examining motor impairment by
thalidomide and morphine, the highest dose of each drug
used was given intraperitoneally to 10 additional rats.
Motor function was assessed by the righting reflex and
placing—stepping reflex. The former was evaluated by
placing the rat horizontally with its back on the table,
which normally gives rise to an immediate coordinated
twisting of the body to an upright position. The latter was
evoked by drawing the dorsum of either hind paw across
the edge of the table. Normally rats try to put the paw
ahead into a position to walk. Pinna reflex and corneal re-
flex were also evaluated and judged as present or absent.

7. Statistical analysis

Data are expressed as mean ± SEM. Time response
data are presented as paw withdrawal threshold to me-
chanical stimulation, while dose—response data are pre-
seated as the percent of maximum possible effect (%MPE).
Withdrawal threshold data from von Frey filament testing
were converted to %MPE, according to the formula: %MPE
= [(postdrug threshold – post—injured baseline thresh-
old) / (cutoff threshold – post—injured baseline threshold)]
× 100. To analyze the 50% probability PWT data and
dose—responsiveness, one—way analysis of variance (ANOVA)
with Scheffe multiple comparison test was used. P values
of < 0.05 were considered statistically significant.

RESULTS

1. Time course for development of mechanical allodynia in
spinal nerve—ligated rats

Before spinal nerve ligation, PWTs ranged from 12 g
to 14 g. After L5, 6 spinal nerve ligation, withdrawal
threshold of the nerve—ligated group was significantly re-
duced in comparison with that of the sham—ligated group,
and this reduced threshold persisted for 21 days. With-
drawal threshold of the sham group did not change during
the observation period (Fig. 1).

2. Effect of intraperitoneal thalidomide on mechanical
withdrawal thresholds

Thalidomide significantly increased PWT compared to
controls (Fig. 2A), and produced dose—responsiveness (Fig.
2B). The lower dose of thalidomide (6.25 mg/kg) produced
no significant reversal of the tactile allodynia, however the
higher dose of thalidomide (12.5, 25 and 50 mg/kg, re-
spectively) effectively increased mechanical withdrawal
3. Effect of intraperitoneal morphine on mechanical withdrawal thresholds

Morphine (3 and 10mg/kg, respectively) significantly increased PWT compared to controls (Fig. 3A), and produced dose-responsiveness (Fig. 3B).

4. General behavioral effects

None of the experimental drug-treated rats showed any motor dysfunction, guarding or obvious change in motor activity when observed in their home cages. All rats remained alert during the experiment.

DISCUSSION

Neuropathic pain encompasses a myriad of painful disease states that are often hard to treat, especially with one single medication. There is no 'gold standard' in regard to one medication for neuropathic pain. Some new medications have emerged during the past few years that help to augment the armamentarium of medications used in neuropathic pain [3]. Recently, increasing evidence indicates that glial cells, one of non-neuronal cells, also play an important role in the pathogenesis of neuropathic pain and the mechanism of neuropathic pain by nerve injury. Neuropathic pain is newly regarded as complex and integral in the interaction of neuronal and non-neuronal cells [4-6]. Not only neurotransmitters from neurons at the synapses, but also many cytokines released from glial cells are involved in the development of hyperalgesia and allodynia in inflammatory as well as neuropathic pain. In addition, much research indicates that proinflammatory cytokines such as TNF-α, IL-1β and IL-6 are major elements contributing to development and maintenance of the pathological state of pain [6,23]; consequently, they have
Fig. 3. Effects of intraperitoneal morphine on the hindpaw-withdrawal response to von Frey filament stimulation after spinal nerve ligation. Data are presented as withdrawal threshold (A) or percentage of maximal possible effect (%MPE, B). Each line or bar represents mean ± SEM of 5−7 rats. Control studies were performed using intraperitoneal saline. Morphine produced a dose-dependent increase in withdrawal threshold. *P < 0.05, †P < 0.01 vs. control.

Thalidomide has profound immunomodulatory actions in addition to its originally intended pharmacological actions. Thalidomide is currently approved by the US Food and Drug Administration in conjunction with dexamethasone in newly diagnosed multiple myeloma, and in the acute treatment of cutaneous manifestations of erythema nodosum leprosum, as well as in maintenance therapy for its prevention and suppression.

In the present study, intraperitoneal thalidomide significantly increased PWT compared to control and produced dose-responsiveness. The lower dose of thalidomide (6.25 mg/kg) produced no significant reversal of tactile allodynia, but the higher dose of thalidomide (12.5, 25 and 50 mg/kg, respectively) effectively increased mechanical withdrawal threshold compared to vehicle.

Although thalidomide and its analogues have broad-spectrum properties, their major mode of action is the inhibition of TNF-α synthesis. Thalidomide reduces synthesis of TNF-α in cultures of lipopolysaccharide-stimulated human microglial cells [25], and also modulates TNF-α production by several mechanisms, for example, by enhancing TNF-α mRNA degradation [26] or by binding to α1-acid glycoprotein [27]. Thalidomide was also found to be effective in animal models by inhibiting TNF-α production [10–13]. Previous studies have shown that thalidomide inhibits inflammatory hyperalgesia in rats and the writhing nociceptive response in mice as a possible consequence of inhibition of TNF-α production [10,11]. It has also been reported that thalidomide reversed and prevented the hyperalgesia by inhibiting TNF-α production in chronic sciatic nerve injury and L5 VRT neuropathy, respectively [12,13]. However, in other studies [13,28], thalidomide was only effective when given preemptively, i.e., before manifestation of the pain-related symptom. However, in our study intraperitoneal injection of thalidomide effectively attenuated mechanical allodynia when it was applied at day 10 after surgery. This discrepancy may be related to different animal models and experimental methods. Also, it has also been reported that when applied 7 days after surgery thalidomide was still capable of inhibiting up-regulation of TNF-α in L5 VRT neuropathy [13]. Thus, it is most likely that the antinociceptive effect of thalidomide in this present study comes from the interruption of TNF-α production. The pharmacological properties of thalidomide aside from their immunomodulatory capacity could also explain the effects observed in our study. The sedative-hypnotic effect of thalidomide was suggested as mediated via interruption or modulation of serotonin’s actions within the CNS. However, rats injected with thalidomide did not show any sedative state at the dose of maximum analgesic effect during the behavioral test.

Various studies have reported reduced antinociceptive efficacy of morphine in animal models of peripheral or
central nerve injury. However, conflicting results regarding the efficacy of opioids on neuropathic pain in different neuropathic animal models have been reported. While systemic administration of morphine attenuates allodynia and hyperalgesia in chronic constriction injury and spinal nerve ligation (SNL) models [29,30], intrathecal morphine is apparently ineffective [29,31]. In contrast, intrathecal morphine dose-dependently reversed mechanical allodynia in a rat model of central pain, whereas systemic morphine had little effect on this measure [32]. Recently, these findings have been challenged by Zhao et al. [33] who reported antiallodynic effects of both systemic and intrathecally administered morphine in the SNI, SNL and SCI animal models of neuropathic pain. These results indicate the efficacy of opioids in neuropathic pain is variable and seems to depend on several factors (e.g., the kind of nerve injury and the route of drug administration). In this study, the 3 and 10 mg/kg doses of systemic morphine significantly increased PWT compared to vehicle. Our study indicates that systemic morphine produces antiallodynic effects in SNL rats, which is consistent with previous reports [30,33]. Our results further support the efficacy of opioid receptor agonists in alleviating signs of neuropathic pain in animal models of peripheral nerve injury.

In this present study, intraperitoneal thalidomide and morphine significantly increased PWT compared to control and produced dose-responsiveness. However, similar studies on various animals are required to obtain a reliable overview of the effect of thalidomide on peripheral and central nerve injury–induced neuropathic pain in a clinical setting.

In conclusion, systemic morphine and thalidomide have antiallodynic effects on neuropathic pain induced by SNL in rats. These results suggest that morphine and thalidomide may be an alternative therapeutic approach for neuropathic pain.

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