Analgesic effects of intra-articular botulinum toxin Type B in a murine model of chronic degenerative knee arthritis pain

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Objective: To evaluate the analgesic effectiveness of intra-articular botulinum toxin Type B (BoNT/B) in a murine model of chronic degenerative arthritis pain.

Methods and materials: Chronic arthritis was produced in adult C57Bl6 mice by intra-articular injection of Type IV collagenase into the left knee. Following induction of arthritis, the treatment group received intra-articular BoNT/B. Arthritic control groups were treated with intra-articular normal saline or sham injections. Pain behavior testing was performed prior to arthritis, after induction of arthritis, and following treatments. Pain behavior measures included analysis of gait impairment (spontaneous pain behavior) and joint tenderness evaluation (evoked pain response). Strength was measured as ability to grasp and cling.

Results: Visual gait analysis showed significant impairment of gait in arthritic mice that improved 43% after intra-articular BoNT/B, demonstrating a substantial articular analgesic effect. Joint tenderness, measured with evoked pain response scores, increased with arthritis induction and decreased 49.5% after intra-articular BoNT/B treatment. No improvement in visual gait scores or decrease in evoked pain response scores were found in the control groups receiving intra-articular normal saline or sham injections. Intra-articular BoNT/B was safe, and no systemic effects or limb weakness was noted.

Conclusions: This study is the first report of intra-articular BoNT/B for analgesia in a murine model of arthritis pain. The results of this study validate prior work using intra-articular neurotoxins in murine models. Our findings show chronic degenerative arthritis pain can be quantitated in a murine model by measuring gait impairment using visual gait analysis scores (spontaneous pain behavior) and joint tenderness scores (evoked pain responses). Reduction of joint pain seen in this study is consistent with our hypothesis of inhibition of release of pain mediators by intra-articular BoNT/B, supporting further investigation of this novel approach to treatment of arthritis pain with intra-articular neurotoxins.

Keywords: intra-articular BoNT/B, osteoarthritis

Introduction

There is an ongoing need for new therapies for refractory arthritis pain. Osteoarthritis-related joint pain is a major cause of physical limitation, disability, morbidity, and increased health care utilization for the more than 27 million Americans affected by osteoarthritis.¹,² Up to 80% of patients with osteoarthritis have movement limitation and 25% cannot perform major activities of daily living.³ Up to 20% of the adult population is affected by chronic knee pain and associated disability, with 11% of adults with knee osteoarthritis needing help with personal care. Osteoarthritis of the knee is currently one of the five leading causes of disability among noninstitutionalized
adults. Additionally, 40% of adults with knee osteoarthritis reported their health as “poor” or “fair”.3

The primary focus of osteoarthritis care is joint pain management because there are no disease modifying agents available for osteoarthritis. Goals of osteoarthritis treatment include improved pain control and maintenance or improvement of joint function. Currently available systemic analgesics have a significant risk of potentially serious side effects. Insufficient joint pain relief, intolerable drug side effects, and adverse drug interactions are limitations of available oral analgesics.5,7 Localized therapies with intra-articular corticosteroids and viscosupplementation are alternative pain control options, but effects and duration may be variable. Minimally invasive surgical intervention for patients failing systemic and local therapies includes arthroscopic lavage and debridement. However, there is increasing evidence that arthroscopic debridement outcomes may be no better than placebo procedures or optimized physical and medical therapy.8,9 Joint replacement for severe disabling degenerative arthritis pain carries significant surgical risks, and is often not an option for many patients due to comorbid medical conditions or advanced age. As average life expectancy continues to increase, the burden of disabling arthritis pain is anticipated to increase as well.

Safe, effective, chronic arthritis pain treatment remains an unmet need for many patients, and represents a growing socioeconomic burden in an aging population.10

Osteoarthritis is characterized pathologically by bony outgrowths (osteophytes), changes in subchondral and marginal bone, bone marrow edema, and damage to articular cartilage surfaces, leading to loss of joint space and joint misalignment. Soft tissue changes include variable degrees of synovial inflammation, capsular thickening, and ligament laxity.11,12 The periosteum and subchondral and marginal bone are richly innervated with sensory fibers, but our current understanding of the cause of arthritis pain remains limited. Most chronic arthritis research and treatment has focused on the degenerative mechanisms and immunologic processes associated with progressive joint damage, rather than the pathogenesis of arthritis-induced pain. Studies of the mechanisms of pain in arthritis have shown that inflammation within joints causes both peripheral and central sensitization of neurons, with spontaneous joint pain at rest and hyperalgesia.13 Given this peripheral sensitization, arthritis pain may be treated effectively by intra-articular neurotoxins. Reduction in arthritis pain following use of intra-articular botulinum toxin Type A (BoNT/A) in humans and in murine models of arthritis has been reported.14–17 We hypothesized that botulinum toxin Type B (BoNT/B) would also reduce chronic arthritic knee pain. Intra-articular BoNT/B is another option for arthritis pain control that may be superior to other BoNT serotypes. To test this hypothesis, we measured the effect of intra-articular BoNT/B on arthritis pain in a murine model of chronic degenerative arthritis.

**Methods**

**Animal subjects**

Forty C57Bl6 mice (Jackson Laboratories, Bar Habor, ME) aged 6–8 weeks old were used in this animal study which was approved by the Minneapolis Veterans Affairs Medical Center Institutional Animal Care and Utilization Committee. The animals were housed in groups of eight animals in the Animal Care and Research Facility at the Minneapolis Veterans Affairs Medical Center, a facility approved by the Association for Assessment and Accreditation of Laboratory Animal Care International. The care and studies of these animals were performed in accordance with the guidelines established in the Guide for the Care and Use of Laboratory Animals (The National Academies Press, USA).

**Collagenase-induced chronic degenerative arthritis model**

Chronic arthritis pain was produced in 40 C57Bl6 mice by intra-articular injection of 10 IU Type IV collagenase (Worthington Biomedical Corporation, Lakeville, NJ) in 10 µL normal saline into the left knee. We used a 30 gauge needle with a customized sheath that limited depth of needle penetration to 2.5 mm. The injection was performed through the midline of the patellar tendon just inferior to the patella to ensure accurate entry into the articular space of the knee. Prior to injection, the area was shaved and sterilized with alcohol and animals were anesthetized with isoflurane inhalation. Arthritis was evaluated four weeks after intra-articular collagenase injection. Mice were evaluated for spontaneous pain behavior, evoked pain behavior, and safety using a battery of standardized measures described below. The timepoints for behavioral testing were prior to and after induction of arthritis, and after treatment of arthritis pain.

**Spontaneous pain behavior: measurement of gait impairment**

Visual gait analysis was performed by walking the animals on a motorized treadmill (Columbus Instruments, Columbus, OH) at a constant speed of 17 cm/sec for a total time of 20 seconds. Gait was evaluated visually and graded...
Evoked pain behavior: measurement of joint tenderness

Evoked pain behavior (tenderness) was measured by tallying fights (kicks, attempts to break from restraint) and vocalizations for one minute in response to repeated firm palpation of the knee. A single examiner performed all examinations and was blinded as to treatment group assignment. A Palpometer® (Palpometer Systems, Inc., Victoria, BC) was used to train the examiner to apply consistent and precise firm pressure, defined as a level of 4 on the Palpometer (1100 gf/cm² = 15.6 psi). Pressure of this magnitude was high enough to elicit a significant pain response from arthritic joints, but not from normal joints. Both the right (normal) and left knee (arthritic) were examined, with the right knee serving as an internal control. The normal right knee was always examined first. In preliminary experiments with tenderness testing, our group found slightly elevated tenderness scores in the nonarthritic knee when the arthritic knee was examined first.

Safety

Systemic adverse effects were assessed by observing for anorexia, dehydration, hunched posture, poor grooming, coat changes, or other evidence of poor animal well-being. Given botulinum toxin’s known effects of muscle weakness, strength was measured at baseline before induction of arthritis pain, four weeks after intra-articular injection of collagenase (arthritic state), and after intra-articular treatments. Change in muscle strength was measured by the ability to grasp a wire grid against resistance and cling to it while inverted. Grasp ability was tested by applying traction to the animal’s tail parallel to the wire grid. Cling ability was tested by inverting the mouse on the wire grid three times with tail held down to wire grid. Both grasp and cling ability were graded on a 0–4 scale. A score of 0 represented inability to grasp or cling to the wire grid. A score of 4 represented a strong grip against resistance and no instability with inversion (Tables 2 and 3).

Intra-articular neurotoxin and controls

Four weeks following intra-articular injection of 10 IU collagenase into the left knee, 17 animals were treated with intra-articular BoNT/B (Myobloc®, Solstice Neurosciences Inc., South San Francisco, CA) 0.02 IU in 5 µL of normal saline into the arthritic left knee. Gait assessment, joint tenderness, and strength examinations were performed three days following intra-articular BoNT/B to allow time for the toxin to take effect. Control groups consisted of arthritic animals treated with either intra-articular normal saline or a sham injection to the left knee at the four-week time point. Seven animals received 5 µL or 10 µL intra-articular normal saline and eight animals received sham injections. Gait impairment assessment, joint tenderness, and strength examinations were performed three days following intra-articular normal saline or sham injections.

Histologic examination of normal and arthritic knees

Following conclusion of the study, right (normal) and left (arthritic) knees of representative animals were examined for histologic evidence of degenerative arthritis. The animals were humanely euthanized using C02 gas and secondary exsanguination. Right and left lower extremities were dissected. Articular specimens were fixed in 10% buffered formalin for 24 hours and decalcified in 10% ethylenediamine

<table>
<thead>
<tr>
<th>Table 1 Gait impairment criteria</th>
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<tr>
<td>0 Mouse refuses to walk on treadmill</td>
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<tr>
<td>1 Mouse walks up to 30% of the allotted time without touching dividers, jumping, splaying its legs, or dragging hindquarters; displays severe inconsistencies in gait</td>
</tr>
<tr>
<td>2 Mouse walks up to 60% of the allotted time without touching dividers, jumping, splaying its legs, or dragging hindquarters; displays moderate inconsistencies in gait</td>
</tr>
<tr>
<td>3 Mouse walks up to 90% of the allotted time without touching dividers, jumping, splaying its legs, or dragging hindquarters; displays minor inconsistencies in gait</td>
</tr>
<tr>
<td>4 Mouse walks the entire time without touching dividers, jumping, splaying its legs, or dragging hindquarters; displays no inconsistencies in gait</td>
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<th>Table 2 Grasp impairment criteria</th>
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<tbody>
<tr>
<td>0 Mouse gives no resistance to being pulled across screen; unable to grasp the screen</td>
</tr>
<tr>
<td>1 Minimal force needed to pull the mouse across the screen; significant difficulty grasping the screen</td>
</tr>
<tr>
<td>2 Moderate force needed to pull the mouse across the screen; moderate difficulty grasping the screen</td>
</tr>
<tr>
<td>3 Significant force needed to pull the mouse across the screen; minimal difficulty grasping the screen</td>
</tr>
<tr>
<td>4 Full force needed to pull the mouse across the screen; no difficulty grasping the screen.</td>
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tetraacetic acid for two weeks before paraffin embedding. Paraffin-embedded specimens were then sectioned and stained with hematoxilyn and eosin.

Statistical methods
The unpaired Student’s t-test was used to compare groups, ie, prior to induction of arthritis, arthritic, treated, and normal right knee values. Comparisons were made between normal, arthritic, BoNT/B-treated arthritic, and saline-treated arthritic controls using unpaired Student’s t-tests. The significance level was selected at a P value of 0.05.

Results
Pain behaviors following induction of arthritis by intra-articular collagenase
Arthritis was successfully induced in 40 mice by intra-articular injection of 10 IU collagenase in 10 µL normal saline into the left knee. Animals were examined four weeks after intra-articular collagenase for development of arthritis. Significant alterations in gait due to arthritis pain was demonstrated by decline in visual gait score from 3.50 (SEM = 0.076) to 2.36 (SEM = 0.112), P < 0.0001. Evoked pain behavior scores induced by palpation of the painful arthritic knee were increased significantly, from a baseline mean of 0.83 (SEM = 0.225) to 2.65 (SEM = 0.0083), P < 0.0134) following intra-articular BoNT/B (Figure 1).

Effects of intra-articular normal saline and sham injections following induction of arthritis
Treatment control groups included seven arthritic mice receiving either 5 or 10 µL intra-articular normal saline into the left knee, and eight arthritic mice receiving sham injections into the left knee at four weeks following intra-articular collagenase. Animals were examined three days following either normal saline or sham injections and compared with untreated arthritic animals (n = 40). There were no significant changes in spontaneous pain behavior scores or evoked pain behavior scores in these control animals. No significant change in visual gait analysis score was noted following either intra-articular normal saline or sham injections (P = 0.225 and P = 0.1921, respectively). Evoked pain response scores following intra-articular normal saline or sham injections did not change significantly (P = 0.9043 and P = 0.5355, respectively, Figure 1).

Normal right knee controls
Throughout all stages of this study the right knee was a normal, nonarthritic internal control. After induction of arthritis, evoked pain behavior response score in the contralateral knee increased from a baseline mean of 0.83 (SEM = 0.208) to 2.65 (SEM = 0.728, P = 0.0083).

Safety
Grasp and clinging strength were measured to monitor safety because of the known effects of muscle weakness caused by botulinum toxins. Strength was measured at baseline, after development of arthritis, and in the post-treatment state. Strength was evaluated by measuring ability to grasp and cling. Forty mice were examined at baseline and four weeks following intra-articular collagenase (arthritic state). Strength assessment showed a significant decline in both measures following induction of arthritis pain, prior to treatment with intra-articular BoNT/B. Grasp scores declined 34% from a baseline score of 3.83 (SEM = 0.120) to 2.53 (SEM = 0.155, P < 0.0001). Cling scores also declined 34% from a baseline score of 3.67 (SEM = 0.129) to 2.43 (SEM = 0.168, P = 0.0003). After induction of arthritis and three days following intra-articular BoNT/B into the left knee, grasp scores improved 22% (n = 17, P = 0.0704), clinging scores

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**Table 3 Cling impairment criteria**

<table>
<thead>
<tr>
<th>Score</th>
<th>Description</th>
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<tr>
<td>0</td>
<td>Mouse cannot hold on when the screen is tilted vertically</td>
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<tr>
<td>1</td>
<td>Mouse has shown that it can grip the screen when vertical but falls during every inversion</td>
</tr>
<tr>
<td>2</td>
<td>The mouse falls off the screen during two inversions or falls off the screen during one inversion and displays instability during the other two inversions</td>
</tr>
<tr>
<td>3</td>
<td>The mouse falls off the screen once or shows instability during two inversions</td>
</tr>
<tr>
<td>4</td>
<td>The mouse does not display instability with any inversion</td>
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improved 23% (n = 17, P = 0.2752), although not reaching statistical significance in this small study. The seven animals that received intra-articular normal saline injections had no significant change in grasp or cling scores compared with the arthritic state (P = 0.3964 and P = 0.7457 respectively, Figure 2). Eight animals that received sham injections had no change in grasp (P = 0.5637), but did have a significant decrease in ability to cling (P = 0.0019). No signs of anorexia, dehydration, hunched posture, poor grooming, coat changes, or other evidence of poor animal well-being were noted in any animals at any point during the study.

Histologic examination of normal and arthritic knees
Following conclusion of the study, right (normal) and left (arthritic) knees of representative animals were examined for
histologic evidence of degenerative arthritis. Hematoxylin and eosin staining of knees revealed irregularities and thinning of articular cartilage and early osteophyte formation compared with normal knees, ie demonstrating changes consistent with osteoarthritis (Figures 3 and 4).

Discussion
This study is the first report of intra-articular BoNT/B for analgesia in a murine model of arthritis pain. The results of this study validate prior work using intra-articular neurotoxins in murine models.16 Our findings show chronic degenerative arthritis pain can be quantified in a murine model by measuring gait impairment with visual gait analysis scores (ie, spontaneous pain behavior) and joint tenderness scores (ie, evoked pain responses). Visual gait analysis showed significant impairment of gait in arthritic mice that improved 43% after intra-articular BoNT/B, demonstrating a substantial articular analgesic effect. Joint inflammation is not a prominent feature of degenerative arthritis, but joint tenderness, measured with evoked pain response scores, increased with arthritis induction and decreased 49.5% after intra-articular BoNT/B treatment. These changes in joint tenderness were clinically and statistically significant even with the relatively small animal numbers used in this study. Reduction of joint pain seen in this study is consistent with our hypothesis of inhibition of release of pain mediators by intra-articular BoNT/B.

Role for intra-articular botulinum therapy
Botulinum neurotoxin (BoNT) is produced by Clostridium botulinum as a complex of proteins containing the neurotoxic
moiety associated with nontoxic components. There are seven serologically distinct BoNT serotypes that all act by inhibiting release of signal chemicals packaged in neuronal vesicles.10 The exocytosis of neuronal signal chemicals is dependent on the function of the N-ethylmaleimide-sensitive factor attachment protein receptor complex, collectively called the soluble N-ethylmaleimide-sensitive factor activating protein receptor (SNARE) proteins.19,20 All serotypes of BoNT cleave SNARE proteins. The specific target site within the SNARE complex is dependent on the BoNT serotype.19 Although all BoNTs act by disabling SNARE-associated exocytosis, the potencies and characteristics of their actions vary.19–21 Botulinum toxins affect striated muscle by creating a chemical denervation that is temporary and reversible through highly potent inhibition of acetylcholine release at the neuromuscular junction.20 Acetylcholine is not the only neurotransmitter affected by BoNTs. BoNTs have also been found to affect the release of multiple SNARE-dependent neuropeptides, including substance P, glutamate, and calcitonin gene-related peptide, all important mediators of articular pain transmission.15,22

BoNTs are the most potent neurotoxins known. However, small doses are successful as cosmetic and musculoskeletal therapies. Currently, BoNT/A and BoNT/B are the best characterized and most used clinically. BoNT/A injections are analgesic for painful muscle contractions associated with cervical dystonia, migraine/tension headaches, and myofascial pain syndromes.23 In BoNT/A treatment of painful soft tissue syndromes, pain relief preceded the resolution of muscle contractions, suggesting that BoNTs may have antinociceptive effects independent of known effects on neuromuscular junctions.24 BoNT/A inhibited capsaicin-stimulated release of substance P from embryonic rat dorsal root ganglia neurons in culture.25 Subcutaneous BoNT/A paw injections in a formalin-induced rat model of pain reduced electrical excitations and c-fos expression in the spinal cord, and reduced edema and tissue glutamate release.26

Efficacy of intra-articular BoNT/A for refractory arthritis pain in humans, and in murine models of arthritis joint pain, has been reported recently. Intra-articular BoNT/A reduced lower extremity arthritis pain by an average of 55%, and shoulder pain by an average of 71% in a study of 11 patients with chronic arthritis pain refractory to intra-articular corticosteroids, with no noted adverse effects.14 In another study of patients with refractory axial skeletal pain, eight of 11 reported a decrease in pain score, improved activities of daily living, and range of motion following BoNT injections. These BoNT injections for axial skeletal pain provided longer lasting pain relief than corticosteroid injections.27 Similar results were found in a randomized placebo-controlled trial of BoNT/A in chronic severe shoulder pain.17 Another randomized controlled trial found that intra-articular BoNT/A was as effective as intra-articular corticosteroids for chronic knee pain.28 Krug et al have reported significant analgesic effects of intra-articular BoNT/A in murine models of chronic inflammatory arthritis.16 Analgesic effects were not found in the acute carrageenan arthritis pain model.

Intra-articular BoNT/B was safe, with no weakness of limb muscles or systemic effects noted. This finding of safety confirms our prior work with intra-articular BoNT/A in murine arthritis models and small human studies. It is possible that higher doses of intra-articular BoNT/B could be used to optimize analgesic effects. The current study does have the noted weakness that our visual gait analysis system may not be sensitive enough to measure full treatment effect, therefore future studies will incorporate computerized digital gait analysis. Additionally, in this preliminary study, the duration of action of intra-articular BoNT/B has not been fully explored. Evaluation of dose response and duration of effect are future research directions for our group. Mixtures of various intra-articular botulinum serotypes may provide faster onset of action and longer duration of effects. Such mixtures may prove useful in other types of articular pain.

This study supports the hypothesis that chronic arthritis pain may be amplified by neuropeptide release in the periphery. Inhibition of neuropeptide release may have altered nociceptor function, and reduced pain generation and neurogenic inflammation. This selective chemodenervation of articular pain fibers with intra-articular injection of neurotoxins is a novel local approach to treatment of arthritis joint pain. Interruption of neuropeptide release by intra-articular BoNT/B appeared to decrease pain responses in the joint and improve gait abnormalities. The results of this study support further investigation of this novel approach to treatment of arthritis pain with intra-articular neurotoxins.

Disclosure

The authors have no conflicts of interest in this work.

References