Background: Pain associated with Complex Regional Pain Syndrome (CRPS) is frequently excruciating and intractable. The use of botulinum toxin for relief of CRPS-associated pain has not been well described.

Objectives: To assess whether intramuscular botulinum toxin injections cause relief of pain caused by CRPS, and to assess the risks of this treatment.

Study Design: Retrospective chart review.

Setting: Outpatient clinic.

Methods: Thirty-seven patients with spasm/dystonia in the neck and/or upper limb girdle muscles.

Intervention: EMG-guided injection of Botulinum toxin - A (BtxA), 10-20 units per muscle. Total dose used was 100 units in each patient. Local pain score was measured on an 11-point Likert scale, 4 weeks after BtxA injections.

Results: Mean pain score decreased by 43% (8.2 ± 0.8 to 4.5 ± 1.1, P < 0.001). 97% of patients had significant pain relief. One patient had transient neck drop after the injections.

Limitations: This is a retrospective study. It lacks a control group and so the placebo effect cannot be eliminated. This study does not provide information on the efficacy of this treatment after 4 weeks.

Conclusions: Intramuscular injection of botulinum toxin in the upper limb girdle muscles was beneficial for short term relief of pain caused by CRPS. The incidence of complications was low (2.7%).

Institutional Review: This study was approved by the Institutional Review Board of the Drexel College of Medicine.

Key words: Complex regional pain syndrome, botulinum toxin, spasm, dystonia

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Comprehensive Pain Syndrome (CRPS) is an incompletely understood, frequently devastating condition characterized by severe neuropathic pain. Associated features include autonomic dysregulation (increased sweating, increased temperature of the affected area, edema), motor symptoms (weakness, tremor, dystonias) and trophic changes (e.g., loss of hair, nail changes, skin atrophy) (1,2). The exact pathophysiology of CRPS is not known. Sensitization of the nociceptive system,
both peripherally and centrally, is believed to be a key pathogenic process (3). The frequently excruciating pain is difficult to treat, and the natural history of CRPS is marked by unremitting worsening of the pain (1).

The most widely known action of Botulinum toxin A (BtxA) is its relaxing effect on skeletal muscles. BtxA is therefore widely used for the symptomatic relief of spasticity, dystonias, and other movement disorders. Recently there has been great interest in the use of BtxA for chronic pain. This interest stems from the fact that BtxA seems to have an early anti-nociceptive action that is independent of its muscle relaxing action and may be due to inhibition of central and peripheral sensitization (4-9). It has been previously hypothesized that the anti-nociceptive action of BtxA may be beneficial in CRPS (10).

We conducted a retrospective chart review of CRPS patients with pain and dystonia of neck and upper limb girdle muscles treated with intramuscular BtxA injections to ascertain the benefits and risks of this treatment.

**METHODS**

Thirty-seven patients were included in the study of whom 35 (95%) were females. All participants in this study were initially seen in the pain clinic, and had CRPS as their primary diagnosis. All patients met the IASP criteria for CRPS (11) with 26/37 (70%) of participants diagnosed with CRPS Type 1, and 11/37 (30%) of participants with CRPS Type 2. Ten (27%) of the participants had localized CRPS predominantly involving one or both upper limbs. CRPS involved the entire body in 27 (73%) of the patients.

Participants with spasm or dystonia in the upper limb girdle muscles were referred for BtxA treatment. Based on previous literature, it was believed that participants with spasm/dystonia in the upper limb girdle muscles were most likely to benefit from BtxA treatment. These participants subsequently had electromyography (EMG) guided BtxA injections into these muscles.

**Botulinum injections**

Participants were treated with intramuscular BtxA injections into specific upper limb girdle and neck muscles. EMG was utilized for injection of BtxA into the targeted muscles. Muscles were selected by patient complaints, hypertrophy, spasm and/or tenderness on palpation. The injecting needle also functioned as a monopolar recording electrode, and the reference electrode and ground electrode was a surface electrode placed close by. The needle was inserted into the targeted muscle using bony and soft tissue landmarks. The position of the needle in the targeted muscle was confirmed by asking the participants to activate the muscle and confirming the presence of motor unit potentials on the EMG monitor. Based on the size of the muscle, 10-20 units of BtxA were then injected through the needle. This procedure was repeated for all muscles injected. The total amount of BtxA used per participant was 100 units.

The majority of participants were injected on only one side - the right side in 15 (41%) and the left in 17 (46%). Eight (22%) had bilateral injections. The specific sites of injection are listed in Table 1.

### Table 1. Muscles injected with Botulinum Toxin A

<table>
<thead>
<tr>
<th>Muscle</th>
<th>Number</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Trapezius</td>
<td>37</td>
<td>100</td>
</tr>
<tr>
<td>2. Splenius Capitis</td>
<td>37</td>
<td>98</td>
</tr>
<tr>
<td>3. Levator Scapulae</td>
<td>35</td>
<td>95</td>
</tr>
<tr>
<td>4. Longissimus Capitis</td>
<td>17</td>
<td>46</td>
</tr>
<tr>
<td>5. Obliqus Capitis</td>
<td>13</td>
<td>35</td>
</tr>
<tr>
<td>6. Scalene</td>
<td>12</td>
<td>32</td>
</tr>
<tr>
<td>7. Semispinalis</td>
<td>12</td>
<td>32</td>
</tr>
<tr>
<td>8. Obliqus Capitis</td>
<td>13</td>
<td>35</td>
</tr>
<tr>
<td>9. SCM</td>
<td>4</td>
<td>11</td>
</tr>
<tr>
<td>10. Paraspinals, not specified</td>
<td>4</td>
<td>11</td>
</tr>
</tbody>
</table>

Pain scores were recorded based on the participant’s report of local pain on an 11-point Likert scale, where a score of zero meant the participant had no pain, and a score of 10 was the worst pain imaginable. Pre-treatment pain scores were collected immediately prior to the injections. Post-treatment pain scores were collected 4 weeks after the BtxA injections. None of the study participants had significant medication changes in the 4 weeks after BtxA injection. Participants who had significant medication changes, or had possible disease modifying treatments such as ketamine infusions, were excluded from this study. A participant was classified as “improved” if the decrease in pain scores was 2 or more. This was based on the results of a recent meta-analysis of pain studies by Farrar et al (12) that demonstrated a 2 point decrease to be clinically significant.
**Statistical Analysis:**

All statistical analysis was done using STATA 9 (Stata-Corp, College Station, TX). The student t-test was used for comparing means, the rank-sum test was used for comparing continuous variables for small samples, and the chi-square test was used for comparing proportions.

**IRB Approval:**

This study was approved by the Institutional Review Board of the Drexel College of Medicine.

**Results**

**Pain relief**

All participants reported severe local pain at baseline, which was 8.2 ± 0.8 (range 7 to 10). The mean local pain score after the BtxA was 4.5 ± 1.1 (range 2 to 8). On an average, there was a 43% (range 11% to 78%) decrease in local pain scores. This difference was statistically significant (P <0.001). These results are illustrated in Fig. 1.

By Farrar’s criterion (i.e., a decrease in local pain score by 2 or more points), 36/37 participants (97%) reported significant improvement in local pain.

**Side-effects:**

One out of the 37 participants (2.7%) developed a transient neck drop after BtxA injection. This participant required a non-invasive plastic neck brace after BtxA injections. The neck drop resolved spontaneously 2 weeks after the BtxA injection without any further complications.

None of the participants reported new onset dysphagia after the injections. No other serious side-effects developed after the intramuscular BtxA injections.

**Factors affecting improvement:**

Age, sex, extent of CRPS, CRPS type, number of extremities affected, and the duration of disease were not found to affect the magnitude of pain decrease seen after BtxA injections.


**Discussion**

In this retrospective study, we found that intramuscular injection of botulinum toxin in the upper limb girdle muscles was beneficial for short term relief of pain caused by CRPS. Participants in this study were diagnosed with CRPS by strict application of IASP criteria by a single expert (RJS), minimizing diagnostic variability. All EMG-guided BtxA injections were administered by a single individual, who was well acquainted with the procedure. There was a 43% decrease in local pain scores 4 weeks after intramuscular BtxA injections. The incidence of serious side effects associated with this therapy was low (2.7%). There was no clinical prognostic marker for success of treatment. A previous study of the natural history of CRPS demonstrated little change of symptom severity after 1 year (1).

In most patients, CRPS is initiated by trauma to the body, a surgical procedure or fracture (1). The inciting event can produce neuropathic pain by peripheral and central sensitization. Peripheral sensitization lowers the threshold for nociceptor discharge by inducing changes in the receptors themselves (autosensitisation) and/or by increasing the excitability of the pain terminal membrane (heterosensitisation) (13). In CRPS, there is evidence of immune mediated inflammation, a process which is likely to be self-sustaining (3). Substance P and CGRP peptides are the vasoactive neuropeptides thought to be most active in the process of neurogenic inflammation (2,14) which might also enhance the “afferent barrage” to the dorsal root ganglia (DRG) and dorsal horn of the spinal cord. This induces central sensitization of pain transmission neurons which is pivotal to the maintenance of neuropathic pain (3,13).

Botulinum Toxin A acts by cleaving the SNAP-25 (Synaptosome-associated protein of 25 kd) complex in the presynaptic terminal, which prevents formation of the SNARE (soluble N-ethyl maleimide sensitive factor-attachment protein-receptor) system. As a consequence, neurotransmitter vesicles do not fuse with the presynaptic membrane, which decreases the release of neurotransmitters at the synaptic cleft. This mechanism decreases the release of acetylcholine, CGRP, substance-P and glutamate which may decrease the nociceptive fiber discharge (9,15-17).

Dystonia occurs in approximately 20% of CRPS patients (18,19). This abnormal, sustained contraction of skeletal muscles can reasonably be assumed to be a significant source of pain in CRPS patients. In the past, intrathecal baclofen and intrathecal glycerine have been evaluated for relief of CRPS associated dystonia. In larger case series, the former was found to be effective (20,21), but was associated with a high complication rate in at least one study (21); while the latter was ineffective in relieving either pain or dystonia (22). In our study, we postulate that the relief of CRPS induced pain by intramuscular BtxA injection is multifactorial: 1) relief of neurogenic inflammation (23-25), 2) relaxation of dystonic muscles that may decrease the afferent nociceptive barrage from sensitised A-delta and C fibers, 3) a distinct antinociceptive action which is distinct from these above mentioned mechanisms (4-6).

There are a significant number of clinical studies on the use of BtxA in neuropathic pain, that includes 2 recent randomized controlled trials. Yuan et al (26) demonstrated in a placebo-controlled randomized trial of 18 patients that intradermal BtxA is effective in relieving painful diabetic neuropathy with results showing 44% of patients receiving BtxA reporting significant relief of pain, as compared to 0% in the placebo group. Ranoux et al (6), in another placebo-controlled randomized trial of 29 patients, showed that intradermal BtxA was effective in post-herpetic neuralgia or posttraumatic/postoperative neuropathic pain. In this study, 40% of patients receiving BtxA reported significant improvement at the end of 2 weeks, as compared to 14% of the patients receiving placebo. The relief of pain was sustained for 6 months after the initial injection. Multiple other non-randomized studies demonstrating the antinociceptive action of BtxA in neuropathic pain caused by diverse clinical conditions have been summarized (27).

In contrast, the use of BtxA for pain relief in CRPS has only been described in 5 small studies, which are summarized in Table 2.

In contrast to earlier studies of BtxA in CRPS, a recent randomized trial failed to find improvement in pain after intradermal BtxA injections (29). The authors postulated that failure was due to the very severe/advanced nature of CRPS in their study participants, and the mode of administration of BtxA (intradermal-subcutaneous rather than intramuscular). This study was followed up quickly by the same group by a small case series describing 2 patients who were treated with intramuscular instead of intradermal BtxA (28). Both patients had improvement in pain, skin discoloration and swelling which is in agreement with the earlier study by Argoff (10).
Botulinum Use in CRPS

Table 2. Previous literature on botulinum use for pain relief in CRPS.

<table>
<thead>
<tr>
<th>Study</th>
<th>Treated CRPS patients</th>
<th>Controls</th>
<th>Treatment</th>
<th>Results</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Safarpour 2010</td>
<td>2 participants. One with right hand, and the other with left hand and forearm CRPS</td>
<td>None</td>
<td>200 units BtxA in 4 muscles in one participant, and 240 units BtxA in 3 muscles in the second participant injected into tender &quot;trigger points&quot; of affected muscles.</td>
<td>Pain score in the local area decreased from 10 to 2 in the first participant, and from 9 to 4 in the second participant. Pain relief sustained in the first participant for 3 years with quarterly injections. Both had improvement in discoloration, swelling and temperature changes in the affected limb.</td>
<td>Results are different than, and complementary to, earlier results by the same group with intradermal BtxA</td>
</tr>
<tr>
<td>Safarpour 2010</td>
<td>8 participants with severe CRPS affecting all 4 limbs</td>
<td>6 similar CRPS participants treated with intradermal injections with normal saline</td>
<td>Intradermal and subcutaneous injections, dose: 40-100 units</td>
<td>No improvement in pain scores, or secondary measures (quantitative sensory testing, sleep quality, pain impact, global satisfaction)</td>
<td>see discussion</td>
</tr>
<tr>
<td>Carroll 2009</td>
<td>9 participants with lower limb CRPS</td>
<td>Same 9 participants (double-blinded randomized crossover study) Placebo treatment was 10 mL of 0.5% bupivacaine</td>
<td>Cervical sympathetic blocks using 10 mL of 0.5% bupivacaine and 75 units of BtxA.</td>
<td>Pain relieved for a longer duration in treatment group (median 71 days) as compared to controls (median 10 days).</td>
<td>Primary endpoint was defined as return to baseline pain level.</td>
</tr>
<tr>
<td>Lauretti 2005</td>
<td>2 CRPS participants (extent not noted) with marked hand dystonia</td>
<td>None</td>
<td>75 IU of BtxA into intrinsic finger flexors and wrist</td>
<td>All participants had improvement in dystonia and pain. Motor relief sustained for up to 8 months.</td>
<td>Participants also received a series of 5 ipsilateral stellate ganglion blocks. Effect of BtxA cannot be discerned separately, especially in the absence of a control group.</td>
</tr>
<tr>
<td>Argoft 2002</td>
<td>11 participants with CRPS affecting only one upper extremity</td>
<td>None</td>
<td>Intramuscular, shoulder girdle muscles. Dose: 25-50 units</td>
<td>All patients reported relief in pain and improvement of skin color and edema at 6 and 12 weeks</td>
<td>Only study which documented changes in sudomotor symptoms</td>
</tr>
</tbody>
</table>

Conclusions

There are several limitations of this study. These include its retrospective nature, the lack of a control group, and the possibility of a placebo effect. The study does not provide information on the efficacy of this treatment after one month. The current study was unable to identify predictors of pain relief after BtxA injection. Further studies are essential to determine these patient characteristics, and define subgroups of CRPS patients that are most likely to benefit from intramuscular BtxA injections.

This study is the largest study to date that documents a potentially useful therapy for a disease characterized by devastating and frequently intractable pain. In view of its limitations, this retrospective study provides data to justify a larger prospective randomized control trial. Further studies need to a) confirm these results, b) identify the subgroups of CRPS patients and modes of BtxA administration that optimize pain relief, c) assess...
changes in other symptoms of CRPS (sudomotor, vasomotor, trophic) and associated conditions (migraine), d) assess its use as an adjunct with other treatments such as ketamine to increase the efficacy of these medications, or to prolong their effect by decreasing the chronic pain stimuli that lead to central sensitization (32,33).

References