Complex Regional Pain Syndrome and Dysautonomia in a 14-Year-Old Girl Responsive to Therapeutic Plasma Exchange

Jeanne E. Hendrickson,1,2* Emma T. Hendrickson,3 Eric A. Gehrie,4 Davinder Sidhu,1 Gerd Wallukat,4,5 Ingolf Schimke,4 and Christopher A. Tormey1,6

1Department of Laboratory Medicine, Yale University School of Medicine, New Haven, Connecticut
2Department of Pediatrics, Yale University School of Medicine, New Haven, Connecticut
3Adams School, Guilford, Connecticut
4Berlin Cures GmbH, Berlin, Germany
5Max Delbrück Center for Molecular Medicine, Berlin, Germany
6VA Connecticut Healthcare System, West Haven, Connecticut

Reflex sympathetic dystrophy, also known as complex regional pain syndrome (CRPS), has recently been shown to be associated with autoantibodies against β2-adrenergic and muscarinic M2 receptors. In addition to pain and sudomotor/vasomotor symptoms, dysautonomia is also observed in a subset of CRPS patients. Despite its severity, there are few effective therapies for CRPS described to date. We report a case of a 14-year-old girl with CRPS of her right leg and dysautonomia (gastroparesis, postural tachycardia) refractory to multiple therapies, successfully treated with therapeutic plasma exchange (TPE) with albumin replacement. The patient, who has serum anti β2-adrenergic and muscarinic M2 receptor autoantibodies in addition to nicotinic acetylcholine receptor ganglionic autoantibodies, underwent an initial course of five TPEs over a 2-week period. She demonstrated a clinical response to TPE as manifested by a rapid improvement in her fatigue and gastroparesis, with a gradual yet significant improvement in her leg pain and sudomotor/vasomotor flares. Following the loading procedures, the patient was treated with rituximab. She continues to require periodic TPE to maintain a remission, with additional immunosuppression being considered long term. Although further studies are needed, TPE (in combination with immunosuppression) may be an appropriate therapy for CRPS patients with detectable autoantibodies, as it is for better characterized diseases with autoantibodies against neuronal surface receptors such as myasthenia gravis or Lambert Eaton myasthenic syndrome. J. Clin. Apheresis 00:000–000, 2015. © 2015 Wiley Periodicals, Inc.

Key words: CRPS; RSD; therapeutic plasma exchange; immunosuppression

INTRODUCTION

Complex regional pain syndrome (CRPS), previously referred to as Reflex Sympathetic Dystrophy (RSD), is a debilitating disease. Despite being described >100 years ago, the etiology of this disease has historically been poorly understood [1]. Symptoms include severe pain, allodynia, vasomotor, and trophic changes of the affected area. Upper extremities tend to be affected in adult cases of CRPS, whereas lower extremities are affected more often in childhood cases. A traumatic event, such as a soft tissue injury or a fracture, precedes many cases. Given the varying presentations and courses of CRPS, this disease has been categorized into “type I” or “type II,” with type I having no nerve lesion identified and type II being termed “causalgia” due to a specific nerve injury.

It is likely that there are further subcategories for CRPS, though they are not well defined. Many patients with CRPS have systemic symptoms, as reviewed by Schwartzman [2]. Such symptoms may affect multiple organ systems, including respiratory, cardiovascular, gastrointestinal, genitourinary, and others. Dysautonomic symptoms may be a prominent systemic feature in some CRPS patients. Like extremity CRPS symptoms, however, the etiology of these systemic symptoms has also historically been poorly understood.

CRPS is increasingly being appreciated to have an autoimmune etiology. Early studies suggesting a female preponderance and an HLA association [3,4] led to a search for autoantibodies. In 2004, Blaes et al. described autoantibodies against the autonomic nervous system in a subset of CRPS patients [5]. In 2009, these

*Correspondence to: Jeanne E. Hendrickson, Yale University Department of Laboratory Medicine and Pediatrics, 330 Cedar Street, Clinic Building 405, P.O. Box 208035, New Haven, CT 06520-0835, USA. E-mail: jeanne.hendrickson@yale.edu.

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autoantibodies were further determined to bind to surface receptors of primary cultured autonomic sympathetic and myenteric plexus neurons, including differentiated neurons with a cholinergic phenotype [6]. A landmark study published in 2011 by Kohr et al. identified the autoantibody target as β2-adrenergic and muscarinic M2 receptors, with binding against the second extracellular loop described [7]. These antibodies were further shown to be functionally active via a cardiomyocyte bioassay [7]; another laboratory also reported functionally active autoantibodies against alpha-1a receptors in patients with longstanding CRPS [8]. Passive transfer experiments of sera from humans with CRPS into mice led to clinical and laboratory features resembling the human disease [9,10], further supporting the pathologic nature of these detected autoantibodies.

Historically, few treatments for CRPS have been effective. An extensive review of potential treatments is summarized by Harden et al. [1]. However, few randomized controlled trials have been conducted on CRPS patients. In brief, a 2004 randomized trial of gabapentin showed a mild effect on pain in some CRPS patients [11]. In 2013, a randomized trial of the amino-bisphosphonate neridronate showed improvement in pain in some participants [12,13]. Ketamine infusions, nerve blocks, spinal cord stimulators, systemic analgesia, and other treatments have been shown to be partially effective in a subset of CRPS patients, but these therapies represent case series or anecdotal reports.

Immunomodulatory therapies for CRPS have only recently begun to be investigated. Steroid treatment has been show to help CRPS symptoms in case reports and small studies [14–16]. Anti-tumor necrosis factor (TNF) treatment has been reported to result in subtle improvements in pain in some CRPS patients [17,18]. Intravenous immunoglobulin (IVIG) has also recently been investigated: a case series of patients with chronic pain (including some with CRPS) treated with a trial of IVIG was published in 2002, with 50% of patients being shown to have a 30% reduction in their pain [19]. A case report published in 2005 showed transient improvement in CRPS symptoms in a single patient following IVIG infusion [20], and another case report published in 2013 showed significant improvement in a patient following high dose (2 g/kg) IVIG infusion [21]. A small randomized controlled crossover trial showed partial improvement in pain and autonomic limb symptoms in patients with refractory CRPS following low dose (0.5 g/kg) IVIG [22]; a larger randomized controlled trial of low dose IVIG in CRPS patients is currently ongoing [23].

Therapeutic plasma exchange (TPE), which serves as an effective adjunctive therapy for other neuroimmune disorders that are mediated in part by autoanti-

CASE REPORT

A 14-year-old girl, with a 10-year history of worsening right knee pain and gastrointestinal dysmotility, presented to an urgent care clinic with severe (9 out of 10 on the numerical rating scale) [27] right knee pain after a minor fall in school gym class. Imaging, including a plain X-ray (XR) and an magnetic resonance imaging (MRI) scan, was normal. The patient was placed in a knee immobilizer and put on crutches, due to an inability to put any weight on her leg without severe pain. Intermittent color changes of her entire right leg from the upper thigh to the mid portion of the dorsum of her right foot were observed (Figs. 1A–1C), as were intermittent temperature changes ranging from “ice cold” to episodes of isolated sweating. The location of her most severe pain, the medial aspect of her right knee, was noted to be bruised approximately 2 weeks after the injury (Fig. 1D). The patient was referred to a tertiary care center, where her exam showed allodynia in the distribution of her right leg where she was also experiencing pain and vasomotor symptoms. Her reflexes were normal, evaluation of right leg strength was limited by severe pain, and fine bilateral hand tremors were noted. Her pupils were dilated but responsive to light. Based on her history and clinical presentation, she was diagnosed with CRPS.

The patient was serially treated with the following medications: nonsteroidal anti-inflammatory drugs (ibuprofen, naproxen), acetaminophen, codeine, gabapentin, amitryptaline, tramadol, prednisone, and pregabaline. Despite these therapies, she continued to rate her pain as a 9 out of 10 and was unable to put any weight on her right leg. She was also treated with twice weekly physical therapy, including desensitization therapy, without any improvement in her pain or vasomotor limb symptoms. Likewise, cognitive behavior therapy did not impact her pain. She had no improvement in her symptoms with local therapy, including
transcutaneous electrical nerve stimulation (TENS), diclofenac gel, and capsaicin cream.

In addition to her right leg pain, she began to experience worsening dysautonomic symptoms 1 to 2 months after her injury, including gastroparesis, constipation, tachycardia, and orthostatic hypotension. She also developed significant fatigue and dizziness, persistent discoloration of her leg during flares, and lack of hair and toenail growth in the affected limb. She was ultimately hospitalized for one week due to an inability to tolerate any solid foods, with an upper endoscopic examination revealing redundant antral tissue (Fig. 2) and delayed opening of the pylorus to the endoscope. An upper gastrointestinal study showed delayed emptying of barium through the pylorus, with no physical obstruction noted. Erythromycin therapy slightly improved her GI symptoms, although she continued to experience stomach pain with solid food intake as well as constipation. Neither metoclopramide nor pyridostigmine bromide improved her GI motility. An electrocardiogram completed during her hospitalization, ordered to evaluate new chest pain, showed sinus tachycardia with a baseline heart rate of 110; an echocardiogram showed a structurally normal heart with continuous tachycardia. At that time, her blood pressure when supine was 110/70 with a heart rate of 110 beats/min; within 1 min of standing her blood pressure was 90/60 with a heart rate of 130 to 140 beats/min. A brain MRI with and without contrast was normal. A dual-energy X-ray absorptiometry (DEXA) scan was normal for age.

Given worsening dysautonomic symptoms, the patient’s serum was sent to Mayo Medical Laboratories (Rochester, MN) for an Autoimmune Dysautonomia evaluation. The testing revealed an acetylcholine receptor ganglionic neuronal antibody (0.08 nmol/L; reference value ≤0.02).

Serum was also sent to Berlin Cures GmbH (Berlin, Germany), and the immunoglobulin fraction was tested at a dilution of 1:40. In the bioassay performed, functional autoantibodies against G-protein coupled receptors change the rate of spontaneously beating neonatal rat cardiomyocytes like the corresponding agonists; the effects of autoantibodies can be blocked by the corresponding receptor antagonists. Two different

![Fig. 1. Lower extremity images. A–C, Depict discoloration of the patient’s right lower extremity during sudomotor/vasomotor flares. D, Depicts bruising of the medial aspect of the patient’s right knee (the location of her most severe pain).](image1)

![Fig. 2. Upper endoscopy, with evidence of a thickened antrum/pylorus.](image2)
Autoantibodies were identified in this patient’s serum. One was directed against the β2-adrenergic receptor, and increased the beating rate of the cardiomyocytes (Δbeats/min = 26.0). This effect was blocked by the specific antagonist ICI118.551 (0.1 μM), (Δbeats/min = 0.0). The other autoantibody detected was directed against the muscarinic M2 receptor and caused a negative chronotropic effect (Δbeats/min = −20.7). This effect was blocked by 1 μM atropine (Δbeats/min = 0.0). The calculated cut off value of normal was Δbeats/min = 7.2. Moreover, the functional activity of the detected autoantibodies was neutralized by short overlapping peptides corresponding to the second extracellular loop of both receptors; the autoantibodies recognized in both receptors an epitope localized in the N-terminal portion. Detailed information concerning IgG preparation, preparation and culture of neonatal rat cardiomyocytes, bioassay handling, and antibody differentiation has previously been described [28].

Other laboratory values, including thyroid function tests, a complete metabolic panel, a complete blood count, a PT/PTT, a vWF panel, an antinuclear antibody (ANA), a rheumatoid factor, a C-reactive protein (CRP), and a cyclic citrullinated peptide, were all within normal limits. Anti-phosphatidylserine IgA and IgG [29,30] were both negative.

TPE was initiated in light of the patient’s refractory leg symptoms, worsening dysautonomia, the detection of autoantibodies, and two recently published case series [25,26]. All other previous therapies had been discontinued at the time of initiation of TPE, including physical therapy as well as cognitive behavioral therapy. An initial intensive schedule of five TPE procedures over a 14-day period were completed using a Cobe Spectra Apheresis System (TerumoBCT, Lakewood, CO), with each procedure exchanging one plasma volume. Albumin with calcium gluconate was used as the replacement fluid and the total fluid balance was set to 110% to address the patient’s low baseline blood pressure. After the second TPE, the patient’s fatigue and nausea began to improve. Additionally, the patient’s right leg stopped flaring (i.e., sweating or turning bright red or ice cold) after the second procedure. After the third procedure, she could eat solid foods without difficulty and she began diuresing and having regular bowel movements. After the fifth procedure, her dizziness began to improve. After completion of the induction TPE the patient’s right leg hair and toenails, which had not grown at all in 4 months, began growing again. Furthermore the bruising of her right knee, which had been present for 4 months, resolved. The patient tolerated all loading TPEs without significant incident, although some perioral tingling was noted at the completion of procedures one and two.

Maintenance procedures were initially planned to be monthly, though 8 days after the patient’s last TPE her nausea and GI dysmotility returned as did her dizziness and fatigue. Her leg pain, which had just begun to subside slightly, also worsened again. The symptoms were significantly improved once the intervals between TPE treatments were shortened. This pattern of partial symptom recurrence within 7 to 8 days of the past procedure and significant improvement within 12 to 24 h after the next procedure persisted for 2 months. Over the course of time, the patient’s right leg pain slowly but very significantly improved (Fig. 3, right). Of note, she had rated her pain as a 9 out of 10 for a full 120 days before the initiation of TPE (Fig. 3, left).

Rituximab, at a dose of 500 mg/m², was administered twice (2 weeks apart), and the intervals between TPE procedures were again attempted to be spaced further. At the time of this report (5 weeks after the first rituximab dose), TPE was still being required weekly to control CRPS and GI dysmotility symptoms. The patient’s cardiovascular dysautonomic symptoms improved somewhat after rituximab, with less hypotension but persistent tachycardia. Autoantibody testing has not been repeated as of this writing.

**DISCUSSION**

TPE was effective in symptomatic relief for this 14-year-old girl with CRPS, dysautonomia, and...
autoantibodies against β2-adrenergic and muscarinic M2 receptors. Her CRPS was longstanding, likely having been present for over a decade. However, it was not diagnosed until a minor injury led to severe pain and classic sudomotor/vasomotor findings. This patient’s disease, which involved not only her right leg (with pain rated as a 9 out of 10 for 4 months) but also her cardiovascular system (postural tachycardia/fatigue), and her GI system (gastroparesis/dysmotility), was refractory to all other therapies attempted before plasmapheresis. TPE significantly improved each of her symptoms, just as it has recently been described to improve the symptoms of other patients with CRPS [25,26].

It is not surprising that TPE is effective for CRPS with β2-adrenergic and/or muscarinic M2 receptor autoantibodies, given that TPE is currently a first or second line therapy for other autoimmune diseases involving surface binding autoantibodies such as myasthenia gravis or Lambert-Eaton myasthenic syndrome [24]. Removal of the patient’s autoantibodies presumably led to her symptomatic improvement, though repeat autoantibody testing was not completed given logistic difficulties. It cannot be ruled out, however, that removal of other substances such as inflammatory cytokines [31–33] may also have contributed to her improvement.

At the present time it is not known what percentage of CRPS patients have detectable autoantibodies, with few patients [7] having been tested for β2-adrenergic or muscarinic M2 receptor autoantibodies. To the authors’ knowledge, and after extensive investigations among CLIA-certified reference centers, there appear to be no laboratories in the United States that offer these tests in a clinical setting. The assays [34] are technically challenging and labor intensive, with the bioassay platform utilized for this patient’s serum involving beating rat cardiomyocytes. Of note, our patient also tested positive for nicotinic ganglionic receptor autoantibodies; this is a radioimmunoprecipitation assay using iodine 125-labeled epibatidine complexed to the α3 acetylcholine receptor solubilized from a human neuroblastoma cell line. It is plausible that these autoantibodies may also be contributing to our patient’s GI dysmotility and dysautonomia.

A subset of CRPS patients exhibit dysautonomic symptoms, and there is an increasing appreciation for the potential contributor of autoimmunity to dysautonomia in postural tachycardia (POTS), GI dysmotility, and autonomic ganglionopathy [2,35]. Li et al. recently described β1-adrenergic autoantibodies in 14/14 patients with POTS, with β2-adrenergic autoantibodies being detected in approximately half of these patients [36]; previous studies have also suggested a potential autoimmune component to this disease with some patients also having ganglionic acetylcholine receptor antibodies [37,38]. These same ganglionic receptor autoantibodies, primarily to the α3 subunit of nicotinic acetylcholine receptors, have also been described to contribute to GI dysmotility [37,39–41] and autoimmune ganglionopathy [40,42,43]. Of 15,000 patients tested by the Mayo Clinic Laboratories, ganglionic receptor autoantibodies were detected in <1%. High levels (≥1 nmol/L) were associated with pandysautonomia. Medium levels (0.10–0.99 nmol/L) were associated with peripheral neuropathies, dysautonomia, or encephalopathy. Low levels (0.03–0.09 nmol/L), such as those detected in this patient, have typically been associated with nonautoimmune disorders or peripheral neuropathies [44].

Our patient has required weekly TPE to control her symptoms, thus prompting a trial of rituximab. To the authors’ knowledge, utilization of rituximab for CRPS has never previously been published in the medical literature, though it has been used successfully in dysautonomia [42,45] and other autoimmune and autoantibody-mediated disorders [46,47], including those responsive to plasmapheresis such as myasthenia gravis [48,49]. Longer term follow-up is needed before the efficacy of rituximab in this case can be commented on. Additional immunosuppression, potentially in combination with adjunctive therapies, will likely be necessary for this patient should frequent plasmapheresis continue to be needed post-rituximab for symptomatic relief.

As with any case report, it should be emphasized that a limitation of our manuscript is that we investigated only a single patient. It remains to be determined whether TPE may be an ideal first or second-line therapy for patients with a diagnosis of CRPS, or perhaps may be most applicable to those with CRPS and dysautonomic features. Identification of circulating autoantibody(ies) before TPE initiation would be ideal, though such an approach remains logistically challenging in the US. The role of long-term TPE for CRPS, both in this patient and in others, is not known at this point in time.

In sum, TPE with albumin replacement, with an initial intensive course of five procedures over 14 days followed by a periodic maintenance schedule, provided impressive symptomatic relief for our patient with severe, refractory CRPS and dysautonomia. This case, in combination with two recently published case series documenting symptomatic relief in adult CRPS patients treated with TPE [25,26], suggest that this intervention—though not risk free—may be one of the more successful treatments described to date for this disease. Additional studies of the autoimmune nature of CRPS, including larger studies of TPE, are warranted.

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**AUTHOR CONTRIBUTIONS**

J.E.H., E.T.H., and C.A.T. wrote the manuscript, and all authors edited the manuscript and approved of the final version. G.W. and I.S. also contributed time and reagents for antibody testing.

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