ORIGINAL RESEARCH

Efficacy of methylprednisolone versus other pharmacologic interventions for the treatment of central post-stroke pain: a retrospective analysis

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Correspondence: Anthony J Pellicane Rehabilitation Institute of Michigan, 261 Mack Avenue, Room 839G, Detroit, MI 48201, USA Tel +1 313 745 1190 Fax +1 313 746 1060 Email apellicane@hotmail.com **Purpose:** To determine if an oral, tapered methylprednisolone regimen is superior to other commonly used pharmacologic interventions for the treatment of central post-stroke pain (CPSP). **Patients and methods:** In this study, the charts of 146 stroke patients admitted to acute inpatient rehabilitation were retrospectively reviewed. Patients diagnosed with CPSP underwent further chart review to assess numerical rating scale for pain scores and as-needed pain medication usage at different time points comparing CPSP patients treated with methylprednisolone to those treated with other pharmacologic interventions.

Results: In the sample, 8.2% were diagnosed with CPSP during acute care or inpatient rehabilitation. Mean numerical rating scale for pain scores day of symptom onset did not differ between those patients treated with methylprednisolone versus those treated with other pharmacologic interventions (mean \pm standard deviation; 6.1 ± 2.3 versus 5.7 ± 1.6 , P = 0.77). However, mean numerical rating scale for pain scores differed significantly 1-day after treatment initiation $(1.7 \pm 2.1 \text{ versus } 5.0 \pm 1.9, P = 0.03)$ and 1-day prior to rehabilitation discharge $(0.3 \pm 0.9 \text{ versus } 4.1 \pm 3.2, P = 0.01)$ between the two groups. Compared to day of symptom onset, as-needed pain medication usage within the methylprednisolone group was marginally less 1-day after treatment initiation (Z = -1.73, P = 0.08) and 1-day prior to rehabilitation discharge (Z = -1.89, P = 0.06). No difference in as-needed pain medication usage existed within the non-steroid group at the same time points.

Conclusion: Methylprednisolone is a potential therapeutic option for CPSP. The findings herein warrant study in prospective trials.

Keywords: stroke, pain, central post-stroke pain, complex regional pain syndromes, therapeutics, neuralgia

Introduction

Central post-stroke pain (CPSP) can be defined as a central neuropathic pain condition occurring after stroke located in the body part(s) corresponding to a cerebrovascular lesion of the somatosensory system characterized by pain and sensory abnormalities where other causes of obvious nociceptive, psychogenic, or peripheral pain have been excluded.^{1,2} The prevalence and yearly incidence of CPSP has been reported as 7.3% and 8%, respectively.^{2,3} CPSP is considered challenging from a clinical management standpoint and existing treatment options do not result in optimal outcomes.⁴ One author describes CPSP as an under-recognized complication of stroke despite its potential to impair activities of daily living, deteriorate quality of life, and undermine rehabilitation efforts, and states that CPSP has an overall immense and devastating burden on patients and society.⁵ Post-stroke complex regional pain syndrome (CRPS) shares

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similar pain characteristics with CPSP, however, post-stroke CRPS also presents with extremity edema and dystrophic skin changes with temperature and color abnormalities.⁶ The pathophysiology underlying CPSP and post-stroke CRPS is not understood; however, these two diagnoses share the possible pathophysiologic process of central sensitization resulting in hyperexcitability of central nociceptive neurons.^{5,6}

Given that corticosteroids may regulate the levels of excitatory neuropeptides that contribute to central sensitization and neuronal hyperexcitability,⁷ Braus et al performed a randomized, non-blinded, placebo-controlled trial assessing the efficacy of an oral, tapered methylprednisolone regimen in the treatment of post-stroke CRPS.8 In this study, 91.2% of patients diagnosed with post-stroke CRPS became symptom free (in an average of 10 days after treatment initiation) and remained symptom free at 6-month follow-up after treatment with the steroid and physical therapy. Similarly, in a randomized, double-blind, active-placebo controlled trial, Kalita et al demonstrated significant improvement in 83.3% of post-stroke CRPS patients treated with an oral, tapered prednisolone regimen versus only 16.7% of patients treated with piroxicam.9 Given the positive results from these two studies, the overlap in pathophysiology between CPSP and post-stroke CRPS, and the difficulties clinicians face in the treatment of CPSP, the retrospective study described herein was initiated in an attempt to assess the efficacy of methylprednisolone versus other pharmacologic interventions for the treatment of CPSP.

It was hypothesized that methylprednisolone would be superior to other pharmacologic interventions for the treatment of CPSP in the acute rehabilitation inpatient with regard to reduction in numerical rating scale for pain scores (NRS) and as-needed pain medication usage. We have no knowledge of any previously published studies that describe the treatment of CPSP with an oral steroid.

Material and methods

This study was approved by the Wayne State University Institutional Review Board in Detroit, Michigan. All patients admitted to an acute inpatient rehabilitation facility after acute stroke between January 7, 2010 and June 30, 2011 were studied retrospectively. Patients documented as being diagnosed with CPSP during acute care or acute inpatient rehabilitation underwent chart, vital signs, and medical administration record review in order to obtain demographic information (age, gender, race), stroke type, past medical history, symptom onset (days post-stroke), pharmacologic intervention(s), 11-point NRS on day of symptom onset, NRS 1-day after treatment initiation, subjective pain reports 1-day after treatment initiation, NRS 1-day prior to rehabilitation discharge, as-needed pain medication usage on day of symptom onset, as-needed pain medication usage 1-day after treatment initiation, and as-needed pain medication usage 1-day prior to rehabilitation discharge. Additionally, those patients documented as being diagnosed with CPSP during acute care or acute inpatient rehabilitation underwent chart review of the initial outpatient follow-up appointment after rehabilitation discharge in order to obtain data on subjective pain reports.

The primary outcome measure was mean NRS during inpatient care and the secondary outcome measure was mean as-needed pain medication usage during inpatient care. For each inpatient, the mean NRS was calculated from all pain scores documented in the electronic medical record from 6:00 am to 11:59 pm on the specified day (without regard to documentation of painful area). Similarly, for each patient, mean number of as-needed pain medications used was calculated from all documented as-needed pain medication administrations per the medication administration record from 6:00 am to 11:59 pm on the specified day (without regard to type of as-needed pain medication administered).

In addition to the aforementioned analysis of those patients diagnosed with CPSP during acute care and acute inpatient rehabilitation, the remainder of the study population also underwent additional investigations. Patients documented as being diagnosed with post-stroke CRPS during acute care, during acute inpatient rehabilitation, or at the initial outpatient follow-up appointment after rehabilitation discharge and patients documented as being diagnosed with CPSP at the initial outpatient follow-up appointment after rehabilitation discharge were also studied. These patients (combined with those patients diagnosed with CPSP during acute care or acute inpatient rehabilitation) were studied in order to obtain prevalence and symptom onset data for CPSP and post-stroke CRPS. These patients' demographic information (age, gender, race), stroke type, and past medical history were also recorded.

Statistical methods

Independent *t*-tests were used for the between group comparisons for NRS at different time points. The Mann–Whitney U test was used for the between group comparisons for as-needed pain medication usage at different time points. Paired *t*-tests were used for the within group comparisons for NRS at different time points. The Wilcoxon signed rank test was used for the within group comparisons for as-needed

pain medication usage at different time points. Criterion for declaring a statistically significant difference was P < 0.05. Criterion for marginal significance was P < 0.10.

Results

During the study period, 146 patients were admitted to an acute inpatient rehabilitation facility after acute stroke. For the entire sample, demographic data, stroke type, and stroke history is presented in Table 1. Additionally, for those patients diagnosed with CPSP and post-stroke CRPS during the study period, demographic data, stroke type, stroke history, prevalence, symptom onset (days), timing of symptom onset, and location of patient at diagnosis (inpatient versus outpatient), is presented in Table 1. Note that two patients diagnosed with CPSP as an inpatient were later diagnosed with post-stroke CRPS at outpatient follow-up after rehabilitation discharge.

Moving forward, we now focus solely on those patients diagnosed with CPSP during acute care or acute inpatient rehabilitation. Of the sample, 8.2% (N = 12) was diagnosed with CPSP during acute care or acute inpatient rehabilitation. Mean onset of symptoms was

Table I Descriptive data

	Entire sample	CPSP	Post-stroke
	(N = 146)	(N = I 4)	CRPS (N = 8)
Prevalence		9.6%	5.5%
Mean age \pm SD (years)	60.3 ± 14.3	$\textbf{49.7} \pm \textbf{16.9}$	52.9 ± 6.5
Gender			
Male	47.9% (70)	42.9% (6)	40.0% (3)
Female	52.1% (76)	57.1% (8)	60.0% (5)
Racial background			
White	19.9% (29)	7.1% (1)	12.5% (1)
Black	77.4% (113)	71.4% (10)	75.0% (6)
Other	2.7% (4)	21.4% (3)	12.5% (1)
Stroke type			
Ischemic	78.8% (115)	85.7% (12)	62.5% (5)
Hemorrhagic	17.8% (26)	14.3% (2)	25.0% (2)
Hemorrhagic conversion	3.4% (5)	0.0% (0)	12.5% (1)
History of prior stroke			
Yes	21.2% (31)	7.1% (1)	50.0% (4)
No	78.8% (115)	92.9% (13)	50.0% (4)
Mean \pm SD (days)		20.9 ± 27.2	55.9 ± 28.7
symptom onset			
post-stroke			
Timing of symptom			
onset post-stroke			
≤30 days		92.9% (13)	12.5% (1)
31-90 days		0.0% (0)	75.0% (6)
91–180 days		7.1% (1)	12.5% (1)
Diagnosis made as inpatient		85.7% (12)	12.5% (1)
Diagnosis made as outpatient		14.3% (2)	87.5% (7)

Abbreviations: CPSP, Central Post-Stroke Pain; SD, standard deviation.

 12.3 ± 6.5 days (mean \pm standard deviation) post-stroke. Patients presented to acute inpatient rehabilitation 8.8 ± 5.1 days after admission to acute care. Two of the twelve patients developed symptoms during their acute care course. Excluding these two patients, acute rehabilitation inpatients developed CPSP 5.0 ± 4.4 days after admission to rehabilitation.

After symptom onset, 7 of the 12 patients diagnosed with CPSP were treated with an oral, tapered methylprednisolone regimen (Medrol Dosepak®; Pfizer Inc., New York, NY, USA) while 5 of the 12 patients received other pharmacologic interventions (these included amitriptyline, fluvoxamine, gabapentin, pregabalin, and lamotrigine in varying combinations). Five of the seven patients who received methylprednisolone did not receive any other treatment for CPSP. The other two methylprednisolone patients received pregabalin and amitriptyline with lamotrigine, respectively. Moving forward, those patients treated with the methylprednisolone taper will be termed the steroid group and those patients treated with other pharmacologic interventions will be termed the nonsteroid group. Table 2 details the methylprednisolone taper used. Due to the short duration of the taper, no patients were on methylprednisolone when discharge NRS or discharge as-needed pain medication usage data was collected.

Mean NRS day of symptom onset for the steroid group and non-steroid groups did not differ significantly (6.1 ± 2.3 versus 5.7 ± 1.6, t[9] = 0.30, P = 0.77). Mean NRS 1-day after treatment initiation for the steroid group was significantly lower than the mean NRS 1-day after treatment initiation for the non-steroid group (1.7 ± 2.1 versus 5.0 ± 1.9, t[9] = -2.55, P = 0.03). Mean NRS 1-day prior to rehabilitation discharge for the steroid group was also significantly lower than the non-steroid group (0.3 ± 0.9 versus 4.1 ± 3.2, t[9] = -3.06, P = 0.01). Figure 1 provides

Table 2 Met	hylprednisolone	taper	details
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Timing of	Dose and regimen
prescription	
Methylprednisolone	Day 1: 12 mg with lunch, 4 mg with dinner, and 8 mg at night
taper prescribed	Day 2: 4 mg with each meal and 8 mg at night
before	Day 3: 4 mg with each meal and 4 mg at night
12:00 noon	Day 4: 4 mg with each meal
	Day 5: 4 mg with breakfast and dinner
	Day 6: 4 mg with breakfast
Methylprednisolone	Day 1: 24 mg one time only
taper prescribed	Day 2: 4 mg with each meal and 8 mg at night
after 12:00 noon	Day 3: 4 mg with each meal and 4 mg at night
	Day 4: 4 mg with each meal
	Day 5: 4 mg with breakfast and dinner
	Day 6: 4 mg with breakfast

a visual representation of these results. Table 3 details the percent reduction in NRS for the steroid and non-steroid groups 1-day after treatment initiation and 1-day prior to rehabilitation discharge compared to day of symptom onset. Evaluation of subjective pain reports in the daily progress notes 1-day after treatment initiation revealed that five of seven (71.4%) patients in the steroid group reported no pain compared to none (0.0%) of the patients in the non-steroid group. As-needed pain medication usage one day after treatment initiation for the steroid and non-steroid groups did not differ significantly (U = 10.5, P = 0.27). The steroid group demonstrated marginally significantly lower as-needed pain medication usage 1-day prior to rehabilitation discharge compared to the non-steroid group (U = 6.0, P = 0.07).

For the steroid group, mean NRS day of symptom onset compared to 1-day after treatment initiation differed significantly $(6.1 \pm 2.3 \text{ versus } 1.7 \pm 2.1, t[6] = 4.98, P = 0.002)$ and mean NRS day of symptom onset compared to 1-day prior to rehabilitation discharge also differed significantly (6.1 ± 2.3) versus 0.3 ± 0.9 , t[6] = 5.88, P = 0.001). For the non-steroid group, mean NRS day of symptom onset compared to 1-day after treatment initiation (5.7 \pm 1.6 versus 5.0 \pm 1.9, t[3] = 1.20, P = 0.32) and compared to 1-day prior to rehabilitation discharge $(5.7 \pm 1.6 \text{ versus } 4.1 \pm 3.2, t[3] = 1.85,$ P = 0.16) did not differ significantly. For the steroid group, as-needed pain medication usage day of symptom onset compared to 1-day after treatment initiation (Z = -1.73, P = 0.08) and compared to 1-day prior to rehabilitation discharge (Z = -1.89, P = 0.06) were marginally significantly different. For the non-steroid group, as-needed pain medication usage day of symptom onset compared to 1-day after treatment initiation (Z = -1.13, P = 0.26) and compared to 1-day prior to rehabilitation discharge (Z = -1.07, P = 0.29) did not differ significantly.

Six of the 12 patients diagnosed with CPSP during acute care or acute inpatient rehabilitation followed-up as outpatients after discharge from acute inpatient rehabilitation. Of these six, three patients were in the steroid group and three patients were in the non-steroid group. Two of the three steroid group patients reported no pain at outpatient follow-up. The other steroid group patient's pain was deemed musculoskeletal. All three patients in the non-steroid group reported pain at outpatient follow-up. One of the non-steroid group patient's pain was deemed musculoskeletal. The other two non-steroid group patients were diagnosed with poststroke CRPS at outpatient follow-up.

Discussion

The prevalence and yearly incidence of CPSP has been reported as 7.3% and 8%, respectively^{2,3} and the majority of patients are diagnosed with CPSP within 3 months of stroke with immediate and delayed onset (>1 year) being possible but atypical.^{2,3,10–12} Fitting with existing literature, the study described here reports a 9.6% prevalence of CPSP with 92.9% of CPSP patients developing symptoms 30 days or less poststroke (Table 1). The occurrence and timing of post-stroke CRPS is less clearly defined. For example, McLean studied stroke rehabilitation inpatients admitted over a 1-year period and diagnosed post-stroke CRPS in only 1.5% of patients.13 However, in this study, diagnosis required a positive bone scan which may have resulted in under-diagnosis. Similarly, in another study by Davis et al, 12.6% of ischemic stroke rehabilitation inpatients were diagnosed with CRPS with most patients developing signs and symptoms between

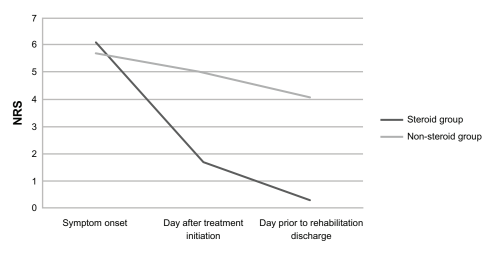


Figure I NRS for pain at different time points. Abbreviation: NRS, Numerical Rating Scale for pain scores.

Table 3 Percentage of patients with different percent reductions in

 NRS at different time points for the steroid and non-steroid groups

	Steroid	Non-steroid
	group	group
NRS day of symptom onset t	o I-day after treatme	nt initiation
\geq 30% reduction	100.0%	25.0%
\geq 50% reduction	71.4%	0.0%
100% reduction	42.9%	0.0%
NRS day of symptom onset t	o I-day prior to reha	bilitation discharge
\geq 30% reduction	100.0%	25.0%
\geq 50% reduction	100.0%	25.0%
100% reduction	85.7%	25.0%

Abbreviation: NRS, Numerical Rating Scale for pain scores.

the second and fourth months post-stroke.¹⁴ Conversely, Kocabas et al reported a higher incidence of post-stroke CRPS (48.8%) in patients followed for 28 weeks with 70% of cases developing 6 weeks post-stroke or later.¹⁵ Also, Gokkaya et al reported that 30.5% of stroke rehabilitation inpatients developed post-stroke CRPS.16 However, it should be noted that these patients were admitted to rehabilitation 67.8 ± 38.9 days after stroke due to considerable patient load at the study site. In the study described here, we report 5.5% prevalence for post-stroke CRPS with 87.5% of patients developing symptoms 31 days or more post-stroke (Table 1). The variability in the reported occurrence of post-stroke CRPS is typically considered to be the result of differences in diagnostic criteria coupled with overlapping signs and symptoms in the stroke patient both with and without a CRPS diagnosis.6 However, we feel the timing of patient assessment should also be considered as a contributor to this variability as well. For example, Kocabas et al¹⁵ and Gokkaya et al¹⁶ demonstrated higher percentages of poststroke CRPS; however, they followed their patients longer and first assessed their patients later (respectively) than the studies that demonstrated lower percentages of post-stroke CRPS. Further, in the study described here, patients were diagnosed with post-stroke CRPS 55.9 \pm 28.7 days after stroke and 87.5% were diagnosed as outpatients. These figures are in stark contrast to the CPSP patients who were diagnosed 20.9 ± 27.2 days after stroke with 85.7% diagnosed as inpatients (Table 1). As a result, it seems that post-stroke CRPS is a complication after stroke that may develop later than CPSP and studies who fail to follow patients for an adequate amount of time may underestimate the occurrence of post-stroke CRPS.

It has been suggested that that amitriptyline, lamotrigine, and pregabalin are all reasonable first-line treatment options for CPSP.^{1,4,17,18} The seminal works that helped to

establish these assertions were performed by Leijon et al,¹⁹ Vestergaard et al,²⁰ and Kim et al.²¹ For amitriptyline, a randomized, blinded, crossover, placebo-controlled trial demonstrated that 75 mg daily of the tricyclic antidepressant yielded significantly lower mean daily pain ratings on a 10-step verbal scale compared to placebo (4.2 ± 1.6) versus 5.3 ± 2.0 by week 4 of the treatment period); and, perhaps more importantly, 67% of the amitriptyline patients reported improvement in pain on global assessment compared to only 7% of the placebo group.¹⁹ For lamotrigine, a randomized, blinded, crossover, placebo-controlled trial demonstrated that a 200 mg daily dose of the anticonvulsant resulted in significantly lower median daily pain ratings on an 11-point Likert scale compared to placebo (5 versus 7) after 8 weeks of medication titration; however, only 44% of patients were deemed clinical responders (defined by a lamotrigine pain score ≥ 2 points lower than the corresponding placebo value).20 For pregabalin, a randomized, blinded, parallel group, placebo-controlled trial assessed the efficacy of 150 mg to 600 mg per day of the anticonvulsant in 219 patients with CPSP.²¹ Mean pain score on the Daily Pain Rating Scale decreased in both groups; however, there was no significant difference between the two groups at baseline or endpoint (6.5 to 4.9 in the pregabalin group; 6.3 to 5.0 in the placebo group). Additionally, the majority of patients treated with pregabalin did not achieve a 30% or 50% reduction in mean pain score compared to baseline. However, the pregabalin group did improve significantly over the placebo group in some of the secondary outcome measures including those regarding sleep and anxiety, and on the Clinician Global Impression of Change rating scale. In comparison to these studies, a short, oral, tapered methylprednisolone regimen appears to be far superior for the treatment of CPSP with regard to absolute reduction and rate of reduction in a pain scale score. Further, the percentage of methylprednisolonetreated patients with \geq 30%, \geq 50%, and 100% reductions in NRS at different time points is remarkable compared to: (1) the non-steroid patient treated patients in this study, (2) the aforementioned pregabalin study, 21 and (3) existing literature commenting on clinically meaningful percent reduction in pain scores.²²⁻²⁶ Additionally, the NRS scores 1-day prior to rehabilitation discharge and the subjective pain reports noted at outpatient follow-up suggests longterm benefit for CPSP patients treated with methylprednisolone. The authors are unaware of any existing literature investigating changes in as-needed pain medication usage in response to treatment in patients with CPSP. Here, within

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the steroid group (but not within the non-steroid group) at different time points, and between the steroid and nonsteroid group at discharge, there was less as-needed pain medication usage noted.

Currently, CPSP and post-stroke CRPS are considered distinct diagnoses. Perhaps CPSP and post-stroke CRPS could be considered one diagnosis that exist on a continuum with CPSP representing a more limited and post-stroke CRPS representing a more florid manifestation of the same syndrome. Given that two patients initially diagnosed with CPSP were later diagnosed with post-stroke CRPS in this study may speak to this possibility. Relatedly, the differences in mean symptom onset and timing of symptom onset for CPSP and post-stroke CRPS described in this study (Table 1) may not demonstrate different onset times for two different post-stroke complications. Instead, these differences may represent the time required for a patient to transition from the more limited to the overt presentation of the same diagnosis. The consideration that CRPS possesses a variety of phenotypes is not without precedent and is applicable to this discussion. Specifically, Bruehl et al postulated CRPS as existing in subtypes; namely, a limited neuropathic pain/sensory syndrome, a limited vasomotor syndrome, and a florid syndrome.²⁷ While their study did not focus on stroke exclusively, it is apparent that their limited neuropathic pain/sensory syndrome might exist on a spectrum that closely approximates CPSP. Given the overlap in proposed pathophysiology underlying these two diagnoses, 5,6,17,28 considering that they may be one diagnosis existing on a continuum is not unreasonable. Additionally, since autonomic dysfunction post-stroke can occur in the absence of CRPS²⁹ this further blurs the line between CPSP and post-stroke CRPS.

Conclusion

This study introduces methylprednisolone as a potential therapeutic option for patients suffering from CPSP. The findings herein warrant study in large prospective clinical trials. Additionally, this study proposes that CPSP and post-stroke CRPS might exist on a continuum given the two diagnoses overlap in pathophysiology and treatment, and since two patients first diagnosed with CPSP later developed post-stroke CRPS. Importantly, it should be noted that both CPSP patients who later developed post-stroke CRPS were not treated with methylprednisolone. This, coupled with the abrupt improvement in NRS noted in the steroid group, might suggest that methylprednisolone has abortive properties in the treatment and prevention of progression of central pain phenomena after stroke.

The authors report no conflicts of interest in this work.

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