Predictive Value of Sympathetic Skin Response in Diagnosing Complex Regional Pain Syndrome: A Case-Control Study
Hyun Jung Kim, MD1,2, Hea Eun Yang, MD3, Dae Hyun Kim, MD1,2, Yoon Ghil Park, MD, PhD1,2

1Department of Rehabilitation Medicine, Gangnam Severance Hospital, Seoul; 2Rehabilitation Institute of Neuromuscular Disease, Yonsei University College of Medicine, Seoul; 3Veteran Health Service Medical Center, Seoul, Korea

Objective To investigate the predictive value of the sympathetic skin response (SSR) in diagnosing complex regional pain syndrome (CRPS) by comparing three diagnostic modalities—SSR, three-phasic bone scans (TPBS), and thermography.

Methods Thirteen patients with severe limb pain were recruited. Among them, 6 were diagnosed with CRPS according to the proposed revised CRPS clinical diagnostic criteria described by the International Association for the Study of Pain. SSR was measured in either the hands or feet bilaterally and was considered abnormal when the latency was prolonged. A positive TPBS finding was defined as diffuse increased tracer uptake on the delayed image. Thermographic findings were considered positive if a temperature asymmetry greater than 1.00°C was detected between the extremities.

Results Five of 6 CRPS patients showed prolonged latency on SSR (83% sensitivity). TPBS was positive in the 5 CRPS patients who underwent TPBS (100% sensitivity). Thermography was positive in 4 of 5 CRPS patients who underwent the procedure (80% sensitivity). The remaining 7 non-CRPS patients differed on examination. SSR latencies within normal limit were noted in 4 of 7 non-CRPS patients (57% specificity). Results were negative in 4 of 5 non-CRPS patients who underwent TPBS (80% specificity), and negative in 3 of 5 non-CRPS patients who underwent thermography (60% specificity).

Conclusion SSR may be helpful in detecting CRPS.

Keywords Complex regional pain syndromes, Galvanic skin response, Radionuclide imaging, Thermography

INTRODUCTION

Complex regional pain syndrome (CRPS) is a painful condition characterized by the presence of regional pain and sensory changes following a noxious event. The pain is accompanied by abnormal skin color, temperature change, abnormal sudomotor activity, or edema [1]. There are three stages of CRPS: stage 1, development of
limb pain; stage 2, progression of symptoms and signs; stage 3, limitation of motion, shoulder-hand syndrome, contractures of the digits, waxy and trophic skin changes, and brittle, ridged nails. Clinical experience suggests that treatment is most effective when initiated during stage 1 immediately after establishing a diagnosis but before radiographic changes appear [2]. The early clinical presentation of CRPS may resemble other disorders, such as cervical nerve root impingement, Pancoast syndrome, vasculitis, migratory osteolysis, arteriovenous fistulae, progressive systemic sclerosis, disuse atrophy, and angioedema. CRPS is usually diagnosed on clinical examination, but diagnosis during early CRPS development can be difficult due to the lack of objective measures.

CRPS encompasses disorders characterized by spontaneous or stimulus-induced pain that is disproportionate to the inciting event and accompanied by a wide variety of autonomic and motor disturbances in varying combinations. Harden et al. [3] proposed new diagnostic criteria for CRPS.

Several studies evaluated the diagnostic modalities for CRPS, but none have compared SSR, TPBS, and thermography for CRPS diagnosis. Thus, the aim of this study was to compare the predictive value of sympathetic skin response (SSR), three-phasic bone scan (TPBS), and thermography in the diagnosis of CRPS.

MATERIALS AND METHODS

Participants
A total 13 patients with severe upper or lower limb pain evaluated at the Gangnam Severance Hospital from December 2011 to February 2013 were included. The patients were diagnosed using the proposed revised CRPS clinical diagnostic criteria [3], according to the International Association for the Study of Pain (IASP). The CRPS type was determined using the IASP diagnostic criteria for CRPS [4]. Six patients were diagnosed with CRPS, according to the revised CRPS clinical diagnostic criteria proposed by IASP [3]. Among them, 4 patients were diagnosed with CRPS-I and 2 patients with CRPS-II. Seven patients reported similar upper or lower limb pain, but they did not fulfill the proposed clinical diagnostic criteria for CRPS at the time of evaluation and until 8 months follow-up post-symptom onset.

SSR
Patients were informed on the SSR method prior to examination. Patients were positioned supine and relaxed on a secured bed in a silent examination room kept at 22°C–24°C or higher [5-10]. The skin temperature was maintained at 32°C–36°C [11-17]. The skin was cleansed and covered with electrolyte gel; active electrodes were then placed on the palm or sole and the reference electrode over the dorsum of the respective body part. The median nerve contralateral to the recorded site was stimulated and latency was measured. SSR was performed using EMG equipment (Medelec Synergy; Oxford Instruments Medical Ltd., Surrey, UK) that included a built-in monitor, bipolar stimulator, and discoid electrodes at the following settings: 10–30 mA stimulation intensity, 0.1 ms stimulation duration, 0.5–1,000 Hz frequency filter, 0.5 mV/div sensitivity, 0.5 sec/div sweep speed, 1 minute minimal interstimulus interval, irregular stimulus, and 3 [18]. The stimulus was delivered irregularly a 1-minute or greater interstimulus interval to avoid habituating the sympathetic skin response. Latency was measured at the initiation of the negative phase, and the shortest latency was chosen among three results. Knezevic and Bajada [11] previously reported that the mean palmar peripheral autonomic surface potential latency was 1.52±0.13 seconds and the mean plantar peripheral autonomic surface potential latency was 2.07±0.16 seconds in 30 normal subjects. We considered the latency was prolonged if it was longer than 1 SD (>1.65 seconds in palmar and >2.23 seconds in plantar).

TPBS
TPBS was performed following an intravenous bolus of 20 mCi of Tc-99m methylene diphosphonate. Sequential radionuclide blood flow (first phase) images were obtained, blood pool (second phase) images were obtained 2 to 3 minutes later, and delayed (third phase) images were obtained 3 to 4 hours later. Diffuse increased tracer uptake during the third phase, confirmed by a nuclear medicine physician, was considered a positive finding.

Thermography
Patients were informed of factors affecting the thermography result and were forbidden to receive any treatment or examination that could affect the thermography findings, such as physical activity, electromyography,
alcohol for 24 hours previously, and tobacco for 2 hours previously. Before undergoing thermography, the patient was acclimated for 15 minutes unclothed in an isolated room maintained at a mean temperature of 23°C±2°C and a relative humidity of 50%. A temperature asymmetry between the affected and unaffected extremities greater than 1.00°C was considered a positive finding [19].

Statistical analysis
The sensitivity and specificity was calculated for SSR, TPBS, and thermography.

RESULTS

Participants
Subject demographics were shown in Table 1. Most of the patients were examined by SSR, TPBS, and thermography as part of the diagnostic confirmation. However, 1 CRPS patient and 1 non-CRPS patient underwent SSR alone; one non-CRPS patient did not undergo TPBS, and another non-CRPS patient did not undergo thermography.

Sympathetic skin response
The SSR findings in 5 of 6 CRPS patients indicated prolonged latency (83% sensitivity). SSR latencies within normal limit were noted in 4 of 7 non-CRPS patients (57% specificity) (Table 2).

Three-phasic bone scan
Positive findings were noted in all 5 CRPS patients who underwent TPBS (100% sensitivity). The findings in the 5 non-CRPS patients who underwent TPBS were negative in 4 of 5 subjects (80% specificity) (Table 2).

Thermography
Thermography was positive in 4 of 5 CRPS patients (80% sensitivity). In the 5 non-CRPS patients who underwent thermography, negative findings were noted in 3 of 5 subjects (60% specificity) (Table 2).

DISCUSSION
This was the first comparative study of the diagnostic...
utility of SSR, TPBS, and thermography with the predictable value of SSR in CRPS diagnosis.

Patients with CRPS may experience sudomotor disturbances, such as hyperhidrosis or hypohidrosis [2,20-23]. The characteristic clinical signs and symptoms of autonomic dysfunction, including edema, skin temperature and color fluctuations, and hyperhidrosis, indicate a role of the sympathetic nervous system in the CRPS pathophysiology [24]. Sympathetic dysfunction is considered the main pathophysiological mechanism of CRPS [25-29].

The SSR is used to measure electrical activity changes occurring in the sweat glands following sympathetic stimulation [5,30,31]. Following an emotional or noxious stimulus, changes in sudomotor activity mediated by the sympathetic nervous system alters the skin’s resistance to an electrical current.

There are few articles investigating the use of SSR for CRPS diagnosis. Pankaj et al. [32] reported the diagnostic utility of SSR for diagnosing CRPS and compared SSR to TPBS. The study was conducted on 60 patients with CRPS affecting the hand; an absent response or a response with an peak-to-peak amplitude <50% compared to the contralateral hand was considered abnormal. Selcuk et al. [33] evaluated SSR in stroke patients with and without CRPS and healthy volunteers; and the onset latency, amplitude, and the number of phases was recorded. The SSR amplitude was significantly higher in patients with CRPS than in patients without it. Pankaj et al. [32] considered the reduction of amplitude of SSR as abnormal, while Selcuk et al. [33] considered the increment of amplitude of SSR as abnormal. Previous studies have shown conflicting results.

Mondelli et al. [34] found that the latency of SSR was prolonged in patients with primary Raynaud’s phenomenon. Uncini et al. [6] reported that the SSR amplitude varied between people and that several factors influence the SSR amplitude. SSR latencies remained consistent even among significantly different recording sites, but not among different stimulation sites; therefore, measuring the SSR latency is important. We evaluated the latency of SSR for abnormality.

Several studies evaluated diagnostic modalities for CRPS. Park et al. [35] demonstrated that the sensitivity of CRPS-I diagnosis was 79.5% and 36.4% in thermography and TPBS, respectively. However, TPBS alone cannot provide an accurate diagnosis; therefore, it is imperative that TPBS findings be integrated with clinical evaluation and other relevant findings.

According to Schurmann et al. [36], the sensitivity of plain radiography, TPBS, thermography, and MRI in diagnosing early post-traumatic CRPS-I was poor, but TPBS and MRI performed at 8 and 16 weeks later, and bilateral radiography at 8 weeks showed a high specificity (TPBS, 96% and 100%; MRI, 78% and 98%, respectively; radiography, 94%). Moon et al. [37] reported that the diagnostic value of a positive TPBS result is low for CRPS, based on the Budapest research criteria. These results suggest that TPBS, thermography, MRI, and radiography cannot be used as screening tests, as they are unable to reliably differentiate between normal post-traumatic changes and those caused by CRPS-I. Clinical findings remain the diagnostic gold standard, while imaging serves as an auxiliary tool to establish diagnosis in suspected cases.

Pankaj et al. [32] found that TPBS is a very sensitive corroborative test to confirm a clinical suspicion of CRPS during the initial disease stages, but not in advanced cases. Alternatively, the SSR can detect sympathetic dysfunction in cases associated with a sweating abnormality and may have assistive diagnostic value in advanced cases of CRPS when TPBS is less reliable. Selcuk et al. [33] showed that SSR could be performed in stroke patients with CRPS even during early CRPS.

The present study was limited by small number of patients (n=13). Further studies using a larger number of patients are necessary. Another limitation is that we added 1 standard deviation to the mean of the latency of SSR to judge the abnormality of SSR. The latency of SSR was shorter than 1650 ms in patient #2; however, the latency of SSR of the affected side was longer than the unaffected side. If we consider the result of SSR as abnormal in patient #2, the sensitivity of SSR is improved to 100%.

The signs and symptoms of the patient are used in proposed clinical diagnostic criteria for CRPS, therefore it is highly subjective. Considering low cost, non-invasiveness and low possibility of complication from SSR, using SSR in diagnosing CRPS may be helpful.

In conclusion, SSR may be a helpful diagnostic test for CRPS. Although additional study is needed, SSR may serve as an objective tool to diagnose CRPS.
CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

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