Cutaneous Silent Period in the Evaluation of Small Nerve Fibres

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ABSTRACT

Introduction: High intensity cutaneous stimulus transiently suppresses tonic voluntary muscle activity resulting in cutaneous silent period (CSP). Aim: The aim of our study was to evaluate the normal values of an onset latency L1, a late latency L2 and a duration of CSP after stimulating sensory fibres of the median nerve. Material and Methods: This prospective study was performed at the Neurology Department, Clinical Center of Sarajevo University in period from January 1st 2013 to December 1st 2013. In our study we examined 61 subjects. The group included our relatives, coworkers and friends. The informed consent from testing subjects was obtained. Results: The origin of silent period is stimulation of small A-delta nerve fibres. The pre-synaptic or post-synaptic interruption of the electrical volley to motor neurons is discussed. Median values of muscle activity suppression in healthy female is 55.0 ms (45.0-74.0) and 59.0 ms (52.0-67) male subjects. There is a correlation between the onset latency L1 and the late L2 latency (p<0.03). In the on-going study it seems that delay of L1 and shorter muscle activity suppression might provide a sign of small nerve fibres involvement. Conclusion: The use of CSP improves the value of neurophysiology examination.

Key words: small nerve fibres, cutaneous silent period.

1. INTRODUCTION

Routine neurophysiology nerve conduction study and needle electromyography (EMG) reflect the function of fastest, large-diameter conducting motor and sensory nerve fibres (1). EMG examination is insufficient assessing the integrity of the whole nerve, to distinguish between axonotmesis and neurotmesis. To rule out the possible interruption of nerve, an additional neurophysiology method is needed. Cutaneous silent period (CSP) is a non-invasive technique which provides the insight to the function of the small-diameter nerve fibres and completes the neurophysiology nerve examination (2).

A sufficient nociceptive cutaneous stimulus exits A-delta nerve fibres and evokes a transient inhibition of voluntary muscle activity occurring in muscles ipsilateral and contralateral to the electrical stimulus (3, 4). The CSP is often used to assess the pathophysiology of ocular (5), oromandibular (5), cervical (6) and brachial dystonia (7). The abnormal CSP was also observed in Parkinson’s disease (8) and intramedullary lesion of the spinal cord (9). Quantitative sensorymetry is performed in many laboratories to evaluate the small nerve fibres (10). Recently to evaluate nerve integrity also ultrasound examination is recommended (11).

2. AIM

The aim of our study was to evaluate the normal values of an onset latency L1, a late latency L2 and a duration of CSP after stimulating sensory fibres of the median nerve. This procedure might be helpful in assessing the function of small diameter nerve fibres.

3. MATERIALS AND METHODS

This prospective study was performed at the Neurology Department, Clinical Center of Sarajevo University in period from January 1st 2013 to December 1st 2013. In our study we examined 61 subjects. The group included our relatives, coworkers and friends. The informed consent from testing subjects was obtained. The subjects with diabetes, alcoholism, polyneuropathy, kidney dysfunction, systemic inflammatory and malignant diseases and the subjects receiving psychotropic drugs respectively, were excluded. The study was approved by the local ethics committee.
All subjects sat in a comfortable chair in a calm room. Using a Synergy EMG machine, the CSP by single electrical stimulation (0.5 ms duration and 80-100 mA intensity, sweeps 250 ms, filters 30 and 10 kHz) at the tip of digit II by bipolar electrode placed on the palmar part of the digit was elicited. Correct placement of the bipolar stimulating electrode on the palmar side of the digit is very important to avoid a possible activation of radial sensory fibres. The superficial electrodes (Care Fusion, Middleton, WI, USA) on the muscle belly of abductor pollicis brevis were placed (Figure 1). During near-maximum activated APB muscle electrical stimulus was delivered. At least 4 individual responses were recorded and superimposed. The onset latency (L1) was recorded at the beginning of muscle activity suppression and the second–late latency (L2) at the start of new muscle activity. The difference between two latencies indicates the duration of CSP.

The data is evaluated by descriptive statistics and determination of Spearman’s correlation coefficient, Mann-Whitney U and Wilcoxon W tests. The onset latencies L1, the late latencies L2 and the duration of CSP—inhibition of muscle activity—were statistically analyzed.

4. RESULTS

In our study 61 subjects were enrolled. The demographic data and CSPs with the onset, the late latencies – L1, L2 and the duration of muscle activity suppression are depicted in Table 1. In our group of healthy subjects significantly more female (74%) than male subjects (26%) were included but the sex didn’t affect the onset (L1), late latency (L2) and the duration of CSP—inhibition of muscle activity—were statistically analyzed.

Table 1. Demographic data and the values of cutaneous silent period

<table>
<thead>
<tr>
<th>Sex</th>
<th>Number (%)</th>
<th>age</th>
<th>Cutaneous silent periods—Median (25th–75th percentiles)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Onset latency L1 (ms)</td>
</tr>
<tr>
<td>female</td>
<td>45/61 (74%)</td>
<td>49</td>
<td>66 (49-73)</td>
</tr>
<tr>
<td>male</td>
<td>16/61 (26%)</td>
<td>51</td>
<td>69 (42-79)</td>
</tr>
<tr>
<td>P</td>
<td>0.0005</td>
<td>0.687</td>
<td>0.173</td>
</tr>
</tbody>
</table>

Table 2. Correlation between the onset (L1), the late latencies (L2), age and sex. CSP (L1) – the onset latency L1 of cutaneous silent period. CSP (L2) – the late latency L2 of cutaneous silent period

<table>
<thead>
<tr>
<th>CSP (L1)</th>
<th>Spearman’s rho</th>
<th>p</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-.240</td>
<td>.030</td>
<td>61</td>
</tr>
<tr>
<td></td>
<td>-.176</td>
<td>.063</td>
<td>61</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>CSP (L2)</th>
<th>Spearman’s rho</th>
<th>p</th>
<th>N</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>-.281(*)</td>
<td>.028</td>
<td>61</td>
</tr>
<tr>
<td></td>
<td>-.133</td>
<td>.305</td>
<td>61</td>
</tr>
</tbody>
</table>

Table 3. Sex and age correlation of CSP duration

The age of both groups was close together and median values of CSP duration were comparable (Table 1). The onset latency (L1) showed a mild correlation with the late latency (L2). By a longer L1 a longer L2 is expected. There is a mild negative correlation between L2 latency and age. The greater age of subjects makes L2 latency slightly shorter (Table 2). In a patient with a severe carpal tunnel syndrome a routine EMG revealed a prolonged distal motor latency and a smaller amplitude of compound muscle action potential – M wave (Figure 2). The sensory nerve action potential of the median nerve on the ring finger was absent, while sensory nerve action potential of the ulnar nerve was clearly recorded (Figure 3). In a case with median nerve entrapment neuropathy a shorter suppression of voluntary muscle activity (31 ms) was recorded comparing to the suppression duration of a healthy subject (60 ms). The delay of L1 latency (27 ms longer) was observed in patient with the median nerve entrapment than in a healthy subject (Figure 4).
Cutaneous Silent Period in the Evaluation of Small Nerve Fibres

4). A prolonged L1 latency in patients with carpal tunnel syndrome was described (12).

5. DISCUSSION

The current study demonstrated a method of CSP measurement. CSP was recorded stimulating the sensory fibres of the median nerve. The decision to evaluate CSP of the median nerve was based on the fact that median nerve entrapment is the most common entrapment neuropathy in humans (13). The focal demyelination of larger – diameter fibres is the primary mechanism of entrapment neuropathy. The second step of nerve entrapment leading to nerve ischemia affects predominantly smaller-diameter fibres producing paraesthesias and pain (14,15). The routine EMG cannot assess the function of small-diameter fiber function which can be evaluated by CSP. Different studies demonstrated the role of CSP in evaluation of corticospinal impairment (16), spinal cord lesion (9), syringomyelia (17) and rigidity in Parkinson’s disease (18).

The sex of subjects didn’t influence the onset (L1) and late (L2) latencies as well as the duration of muscle activity suppression (Table 1). To obtain an optimal CSP it is very important to achieve near maximal contraction of the thenar muscles. At least 4 recordings of CSPs are recommended. The superimposed CSP recordings improve the exact measurement of latencies and duration of muscle activity suppression. The correct position of the stimulating electrode on the palmar side of the second finger enables avoiding the stimulation of radial sensory fibres (Figure 1). An optimal CSP by higher single electrical pulses was delivered. The importance of considerable stimulus intensity more than voluntary muscle contraction is reported (19).

There is a mild correlation between L1 and L2 (Table 2). The shorter duration of CSP and delayed L1 latency indicate the involvement of smaller-diameter median nerve fibres but preserved median nerve integrity (Figure 3). Additional involvement of small nerve fibres might influence the improvement of entrapment neuropathy after surgery.

6. CONCLUSION

CSP is an inhibitory spinal reflex produced by small A-delta nerve fibres and provides an important information of nerve fibre continuity. To obtain reliable CSP a proper muscle contraction and particularly suitable strong stimulus intensity of the target sensory nerve fibres is required. This neurophysiology method is useful in assessing severe entrapment neuropathy, traumatic nerve injury or different polyneuropathies. The measurement of CSP enables a new information of nerve function and increases the sensitivity of routine EMG.

CONFLICT OF INTEREST: NONE DECLARED

REFERENCES


