Strength-duration curve: a measure for assessing sensory deficit in peripheral neuropathy

WG FRIEDLI, M MEYER

From the Department of Neurology, University Hospital, Zürich, Switzerland

SUMMARY By using an isolated constant current stimulator producing true square-wave pulses, sensory strength-duration curves were obtained at various sites by percutaneous electrical stimulation. Strength-duration curves derived from normal groups were compared to those of patients with peripheral neuropathy. Stimulus strength at sensory threshold was shown to be a reproducible measure of sensory deficit, increasing parallel to the degree of axonal failure found by conventional methods. This may be useful as a complementary method in assessing peripheral neuropathy.

The fact that both muscle and peripheral nerve characteristically respond to electrical stimulation allows a number of electrodiagnostic procedures. The relationship between the strength of a stimulus and its duration for producing minimal excitation is expressed by the strength-duration (S-D) curve.1 The standard type of S-D curve utilising visible muscle contraction has been applied to traumatic peripheral nerve injuries and lower motor neuron lesions.2-4 While electromyography provides extensive information as to the state of innervation of skeletal muscles, sensory conduction velocity often does not mirror the clinical state of the patient. For several reasons the quantitative examination of cutaneous sensation encounters serious problems.5 Both afferent and efferent fibres have been studied by various electrodiagnostic techniques: nerve excitability has been determined by the strength of the true square-wave stimuli required to elicit a barely visible nerve action potential at a recording site proximal to the stimulus.5-11 Stimulus strength at sensory threshold has been shown to be more reproducible than the one just evoking a sensory action potential.12 Sensory S-D curves obtained with behavioural psychophysical techniques have matched those obtained by electrophysiologic procedures.13 However, few clinicians have used cutaneous electrical stimulation for assessing peripheral nerve function14 and clinical feasibility has not been apparent in the past. According to our findings the sensory S-D curve offers, to a certain extent, a useful method for assessing peripheral neuropathies.

Methods

Silver surface electrodes of 8 mm diameter filled with electrode cream were taped about 2.5 cm apart over the skin areas of interest. Skin preparation consisted of shaving the skin if necessary and cleaning it with ether and alcohol. The distal electrode was connected to the cathode of an isolated stimulation system consisting of an impulse generator (W-P Instruments Inc.) and a constant current stimulator (DISA 15 E 07). The latter was modified for external pulse width control and for extended pulse width of 50 ms (with approximately 10% pulse drop for 50 ms pulse duration). The amount of current in milliamperes required to produce a minimally perceptible sensation was determined for various durations of square-wave pulses ranging from 0-05 to 50 ms (0-05, 0-06, 0-08, 0-10, 0-20, 0-50, 1-0, 2-0, 5-0, 10-0, 20-0, 50-0 ms). For each of the 12 pulse durations detection thresholds were obtained by a method of descending limits, continuously decreasing from a level at which the subject had a minimal sensation. Electrical stimuli were given at a rate of 0-5/s and were displayed on an oscilloscope screen in front of the subject. Usually three independent measurements were performed on each subject for each pulse duration tested, the series starting with 50 ms stimuli. S-D curves were obtained by plotting current intensities producing a just perceptible sensation against stimulus duration.

For each stimulation site studies were performed on a control group and a patient group. Each control group consisted of 10 to 12 healthy subjects without history or clinical evidence of neuromuscular disease. To ascertain the reproducibility of normal curves the same subject was tested in 10 independent series of measurements on successive days. During the measurements skin temperature was controlled by means of a digital thermometer and kept constant by an infrared lamp within the range of 1°C.
Results

S-D curves were surprisingly reproducible in healthy subjects on successive trials. The task was more difficult for some of the patients depending on the degree of sensory impairment, especially if paresthesiae were present. The subjects compared the sensation produced by electrical stimuli to the sensation of being tapped, that is, a stimulus tactile in quality.

In a first series of experiments normal S-D curves were obtained at various stimulation sites. In one group of 12 healthy subjects sensory thresholds were measured bilaterally in the cutaneous field of the superficial peroneal nerve. The electrodes were placed on the proximal dorsum of the foot following the tendon of extensor hallucis longus. Average thresholds across trials and different subjects are presented separately for the two sides of the body in fig 1A. A rapid decrease of the threshold current with increasing stimulus duration up to 0-5 to 1-0 ms was observed while there was little change of threshold current with longer stimuli. The minimal current required to produce cutaneous sensation also with very long current (rheobase) was achieved by pulse durations of approximately 10 ms. Sensory thresholds were found to be lower for the left compared to the right side of right-handed people in both upper and lower extremities. Although the difference in absolute values may be disregarded, this asymmetry was a consistent finding in our normal subjects.

Reproducibility of S-D curves for the same stimulation site is shown in fig 1B for the same subject. While being in the same range, the threshold values showed smaller standard deviations when compared with the whole group, even if independent measurements were performed on different days. A similar degree of reproducibility was observed in individual subjects for the cutaneous field of the median or ulnar nerve.

Temporal summation properties are represented by the product of current strength and duration at threshold levels (stimulus energy in microcoulombs) as a function of stimulus duration (fig 1B). According to the Bunsen-Roscoe law or Bloch’s law stimulus energy at threshold levels remains constant for stimulus durations less than the “critical duration” specific for a sensory modality. For vision, reciprocity between luminance and time at threshold has been found to be limited to durations of about 10 ms in the paramacular region5 while the critical duration was found to be considerably longer for other sensory modalities. In agreement with Rollman’s findings our data suggest a very small critical duration of less than 0-10 ms for cutaneous electrical stimulation. The absence of a total temporal summation in our graphs was probably due to the fact that minimum pulse duration was limited to 0-05 ms. Different degrees of partial temporal summation were found for successive periods, the first extending to about 0-10 ms, and a second to about 1-0 ms. There was no temporal summation for pulse durations beyond 10 ms and the function had a slope of 1-0.

To allow comparison between different stimula-
Fig 2 S-D curves (superficial peroneal nerve) in neuropathic disease compared to normals of fig 1 (hatched area: mean ± 1 standard deviation). Individual curves from patients with angiopathic neuropathy (A), alcoholic (B), uraemic (C) and diabetic (D) polyneuropathy (details in table). Compare marked differences of rheobases and the similar chronaxy values for the different conditions. Upper graph: threshold energy as a function of pulse duration for patient B and healthy group showing differences in temporal integration properties between the two.

Sensory S-D curves were obtained from 25 patients with various kinds of peripheral neuropathy. Some examples are given for stimulation in the cutaneous field of the superficial peroneal nerve in fig 2 (see table for case descriptions). S-D curves of the pathological subject groups are compared with the group of normal subjects in fig 1. The rate of climb in the pathological curves was larger from right to left compared to that of the normal curves. However, even if present, this variation of slope was too small to be of any practical relevance. In general, the sensitivity threshold in terms of stimulus current was raised over the whole range of
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Table  Case descriptions (patients of figure 2). Rating of pathological conditions: 0 normal, 1 mild, 2 moderate, 3 severe

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<th>Type of neuropathy</th>
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<th>Weakness</th>
<th>Arephey</th>
<th>Sensory impairment</th>
<th>Vibratory sense</th>
<th>Further symptoms</th>
<th>Motor conduction defect</th>
<th>Amplitude evoked response</th>
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Fig 4  (A) 41-year-old male. Traumatic partial lesion of the right median nerve in the distal forearm 4 months before examination. Severe sensory impairment to all modalities in the index (●), markedly less in the middle finger (○). No sensory action potentials from digit to wrist. Note the difference between S-D curves on the two fingers which is in accordance with the clinical findings. (B) 24-year-old female, suffering from symptoms of a moderate carpal tunnel syndrome (sensory conduction of 39 ms, small sensory action potential of 4 μV). S-D curve on the right index finger before (○) and after (●) paraneural infiltration with an anaesthetic into the carpal tunnel (when no action potential could be recorded any more). (C) S-D curves (index finger) of a 60-year-old male patient with severe carpal tunnel syndrome before (○) and 3 months after (●) surgical decompression of the right median nerve. Note the difference of rheobase before (no sensory action potential, distal motor latency of 6·3 ms) and after surgery (markedly delayed action potential of 2·5 μV, motor latency of 5·1 ms).
pulse duration as is shown by the curve shifting upward and to the right. This shift was also found for stimulus energy at threshold levels (fig 2, upper graph): the patient curve shows an overall rise of threshold energy. A comparison of the slopes of the two energy functions revealed changes in temporal summation properties. In contrast to healthy subjects, a reciprocity between stimulus strength and time at threshold levels was found for pulse durations up to 0-10 ms in subjects with severe neuropathy. Hence, not only partial temporal summation was extended to longer pulse durations but also the critical duration was prolonged in the pathological case.

S-D curves for the median and ulnar nerve were obtained by stimulating the distal palanges of the index and middle finger or of the little finger respectively. Normal curves were compared with those found with various nerve lesions. Even if there was clear clinical and electromyographic evidence of entrapment neuropathy, sensitivity thresholds often did not reflect the pathological condition. However, S-D curves were significantly altered with a severe carpal tunnel syndrome, characterised by marked conduction deficits, electromyographic abnormalities in the thenar muscles or loss of the sensory action potential (fig 3). In these cases, serial S-D curves after surgical decompression of the median nerve displayed a functional improvement (fig 4C). There was also a clear relationship between sensory deficits due to traumatic nerve lesions and the behaviour of sensitivity thresholds as a function of stimulus duration (fig 4A).

Discussion

Together with the clinical and electrodiagnostic evaluation of a peripheral neuropathy, the systematic examination of sensory strength-duration curves provides useful help in diagnosis.

The characteristics of psychometric functions for electrocutaneous stimuli have suggested that current pulses bypass the receptors and directly excite sensory fibres as supported by other results. Moreover, temporal integration properties of psychophysical S-D curves were consistent with stimulation of large A-fibres by electrocutaneous pulses. According to previous findings fibres with large diameter have faster velocities and lower thresholds than fibres with small diameter.

The correlation of the S-D curve with pathological clinical and electrodiagnostic findings in the same patient provides further information as to the anatomical substrate tested by the psychophysical technique. Both patients A and B of fig 2 had a symmetrical motor and sensory polyneuropathy with subnormal to normal nerve conduction and the electromyographic equivalents of a severe axonal lesion. On the other hand, the findings in patients C and D are characterised by motor and sensory conduction defects with little evidence of denervation. This is in agreement with the fact that S-D curves are hardly impaired in chronic entrapment neuropathies (for example, the median nerve in fig 4B) but affected in proportion to the incidence of axonal lesion (for example, patients of fig 3).

Discontinuities in the classical S-D curve result from the difference in excitability threshold between muscle and nerve tissue. Therefore, sensory S-D curves representing properties of afferent nerve fibres are expected to be smooth also in case of partial denervation. Pathological conditions are characterised by an over-all increase in threshold excitability providing both a shift of the curve and changes in temporal integration properties. There are various possible explanations for this behaviour; it may be due to a threshold increase of the A fibres, or stronger stimuli may be required to recruit an increasing number of A fibres for minimal sensation. Another possibility is that stronger pulses may provide excitation of other types of afferent fibres in addition to A fibres. This is suggested by the changes in temporal summation properties. However, our results cannot give additional information about the elements of sensory S-D curves.

Assessment of sensory impairment is part of the clinical evaluation of peripheral neuropathies. For several reasons quantification of cutaneous sensation by clinical examination is inaccurate and calls for methods of measurement using “adequate” stimuli. Both sensory S-D curves and quantitative sensory tests are complementary to the clinical evaluation of peripheral nerve disorders. The fact that the sensory threshold is mostly affected with axonal lesions suggests a valuable complement to conventional electrodiagnostic techniques.

The simplicity of the method (no complex recording and averaging equipment) represents a further advantage in comparison with electrophysiographic procedures. However, the isolated constant-current stimulator has to be capable of producing true square-wave stimuli over a wide range of pulse durations. Moreover, the subject tested must be attentive and cooperative during the behavioural assessment of peripheral nerve function. The examination can be reduced to a few representative pulse-durations. The amount of current of infinite duration required to produce minimal sensation (rheobase) represents a valuable measure and approximates stimulus durations of 10 to 50 ms. The chronaxy, that is the duration of a threshold stimulus of twice the rheobase, proved to be less informative, especially because it
is based on calculation. Temporal integration properties and critical duration are useful parameters but require several measurements in the low range of stimulus duration. For each laboratory S-D curves must be based on normal values (specific for age and stimulation site). However, bilateral comparison within the same subject might be indicative of mononeuropathy.

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References