

Simple Screening Tests for Peripheral Neuropathy as a Prediction of Diabetic Foot Ulceration

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Background and Objectives: Diabetes mellitus is one of the major health problems, and one of its commonest complications is peripheral neuropathy. The aim of this study is to evaluate four standard simple screening tests (10-g Semmes-Weinstein monofilament examination {SWME}, vibration sensation by a 128Hz tuning fork, superficial pain test and ankle reflex) for detection of neuropathy using the nerve conduction study (NCSs) as the standard criterion in diabetic patients.

Methods: The study included 120 patients with diabetes mellitus, all patients had complete clinical assessment including presence or absence of neuropathy, exclusion of other causes of neuropathy, NCS, 10 g Semmes-Weinstein monofilament examination, vibration sensation by a 128 Hz tuning fork, superficial pain sensation and ankle reflex were done for all patients.

Results: Foot care practices were followed by 80 (66.6%) of the study population. When compared with NCSs, the monofilament was the most specific at 91%, less sensitive 57%, superficial pain, ankle reflex had lower specificity (36%, 41%) respectively, sensitivity (62%, 57%) respectively. Tuning fork had specificity 90%, sensitivity 56%.

Conclusion: The study findings show that the simple screening tests (10-g SWME, vibration testing, superficial pain test and ankle reflex) can be used confidently for annual screening of diabetic neuropathy in diabetic patients.

Key Words: Monofilament-tuning fork-ankle reflex-superficial pain-neuropathy.

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Diabetes mellitus is considered as a major health problem and one of its commonest complications involves the feet. Diabetic foot problems may lead to lower limb amputation which occurs commonly in uncontrolled diabetics. The chronic peripheral neuropathy which is associated with diabetes presents progressively and insidiously with poor symptoms related to the pathological severity.^{1,2}

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Peripheral neuropathy initiates the pathophysiology to leg ulceration and may be amputation and it is the main cause for sensory ataxia, painful paresthesia, and Charcot deformity.³ Early identification and screening of neuropathy offer an important opportunity for the diabetic patient to actively alter the suboptimal glycemic control and improve foot care prior to significant morbidity.⁴ Electrophysiological studies have shown the relation between glycosylated hemoglobin and the presence and also the severity of neuropathy.⁵



Figure 1 10 gram sensory monofilament.

The aim of this study is to evaluate four standard simple screening tests (10-g Semmes-Weinstein monofilament examination {SWME}, vibration sensation by a 128 Hz tuning fork, ankle reflex and superficial pain test) for detection of neuropathy using the NCSs as the standard criterion in diabetic patients.

Patient and Methods

Our study included 120 diabetic patients between March 2010 to July 2011, history taking of duration of diabetes, associated diseases, presence or absence of neuropathy (e.g. foot pain, tingling, numbness, imbalance, weakness and upper limb symptoms) and its duration if present, and they were examined as follows:

1. Exclusion of other causes of neuropathy (e.g. familial, nutritional, uremic, and alcoholic) by comprehensive examination medically and neurologically.
2. Standardized NCSs (bilateral) including motor (tibial, peroneal, median, and ulnar) and sensory (sural, median, and ulnar) nerves –performed by a blinded technicians to the status of the patient.
3. A 10-g SWME (Fig. 1) superficial pain and vibration sense by a 128 Hz tuning fork (Fig. 2a,b) and ankle reflex, all were performed by a third examiner who is blinded to the history, physical examination and the results of NCSs.



Figure 2a and 2b 128 hz tuning fork.

All data were entered on standardized forms, subjects identification was by number, date of birth, and initials.

Sensory testing methods

The patient was given a reference sensation by application of the stimulus to the sternum and then asked the nature of the sensation perceived. When the nature of the sensation was perceived accurately on the sternum, the patient was asked, with eyes closed, to describe the sensations experienced sequentially at the sites described below:

The SWME was conducted using a 10-g monofilament applied to a non-callused site on the dorsum of the first toe just proximal to the nail bed. It was repeated four times on both feet in an arrhythmic manner. The SWME threshold was defined as the total number of times the application of 10-g monofilament was not perceived by the patient, and it varied from 0 to 8.

Vibration testing by a 128-Hz tuning fork applied to the bony prominence bilaterally situated at the dorsum of the first toe just proximal to the nail bed. The patient was asked to report the time at which vibration diminished beyond perception. The tuning fork was then applied to the dorsal aspect of the distal phalanx of the examiner's thumb. The time (in seconds) at which vibration sensation diminished beyond the examiner's perception was then recorded on a standardized form. The values from both sides were added to provide a single score for statistical analyses.

Test	Sensitivity %	Specificity%	PPV%	NPV%	P-value
Monofilament	57	91	94	44	P<0.001
Superficial Pain	62	36	49	50	P=0.99
Tuning fork	56	90	93	44	P<0.001
Ankle reflex	57	41	53	44	P<0.10

Table 1 Sensitivity, specificity and P-value of the four tests. PPV (positive predictive value), NPV (negative predictive value).

The vibration test threshold was defined as the total number of times the application of the vibrating tuning fork and the dampening of vibration was not felt, with scores varying between 0 and 8.

Superficial pain sensation was conducted using a sterile Neurotip™ (Owen Mumford) applied four times to the two sites described in SWME. The superficial pain threshold was defined as the total number of times the application of the pain sensation was not perceived, with scores varying from 0 to 8. Ankle reflex was assessed with a tendon hammer and was recorded as either present or absent.

Criterion standard

Standardized techniques for NCSs were applied with temperature control and fixed distances. Measurements of latencies, distances, and amplitudes were assessed in a standard fashion using onset latencies and baseline to peak amplitudes. Initial positive peak (if present) to negative peak measurements were conducted for sensory responses. F waves were generated for all motor nerves, and minimal, reproducible latencies were measured. Conduction velocities were calculated for motor and sensory nerves.

All conduction velocity and distal amplitude values for the NCSs were given a score of 0 for normal and 1 for abnormal. The mean reference values $\pm 2SD$ were taken as the normal range. The maximum NCS score if all parameters were abnormal was 28 points (16 motor and 12 sensory). The total NCS score was defined as the sum of the number of abnormal values.

Statistical analyses

The data were analyzed and we used the X2 test to detect the difference of results across the four tests compared to NCSs, $P < 0.01$ was considered statistically significant. By constructing Receiver operating characteristic curve, sensitivity, specificity, positive and negative predictive values were calculated for the various tests using NCS as the gold standard definition of neuropathy.

Results

The study included 120 diabetic patients. The mean age was 52.2 ± 4.8 (range 22-85 years), 59 males (49.2%), 61 females (50.8%). In the study all patients had type 2 diabetes, the mean duration of diabetes was 8.21 ± 7.81 years and mean FPG 202.15 ± 50.44 mg/dl, 80.2% were receiving oral hypoglycemic drugs, 19.8% were receiving insulin. Foot care practices were followed by 80 (66.6%) patients of the study population by optimization of the glycemic control, cessation of smoking, prescription of adequate proper fit foot wear with wide deep box and debridement of calluses with follow up at 6 months and 12 months.

Evaluation of neuropathy by nerve conduction study (NCSs) showed peripheral neuropathy in 75 patients (62.5%), using other testing modalities neuropathy was found in 56 (46.6%) patients with monofilament, in 45 (37.5%) with Superficial pain test, in 57 (47.5%) with vibration test and in 56 (46.6%) with ankle reflex.

Table 1 presents the sensitivity, specificity, PPV, NPV and P-value of each diagnostic test compared with NCSs which was taken as the gold standard.

Discussion

In this study we used NCS as a standard criterion for the diagnosis of peripheral neuropathy. Since peripheral neuropathy is a main element in causing of both foot ulceration and amputation so selection of rapid, simple and accurate testing method for diagnosis of peripheral neuropathy in diabetic patients is so important and apart from NCS we select monofilament, superficial pain, tuning fork and ankle reflex for evaluation of peripheral neuropathy.

The most frequently used modality for peripheral neuropathy detection is the nylon Semmes-Weinstein monofilament.⁷ Inability to perceive 10 g of force 5.07 monofilament applies is associated with clinically significant large-fiber neuropathy. In our study monofilament showed a sensitivity 57% and a specificity 91% compared to other studies with 95% sensitivity and 82% specificity,^{8,9} other studies showed sensitivity of 77% and specificity 96%⁶, also another study showed sensitivity 66% and 34% specificity.¹⁰

The 128 Hz tuning fork in our study showed sensitivity of 56% and specificity of 90% compared to other study sensitivity and specificity was 53% and 99% respectively. The superficial pain test in our study showed sensitivity and specificity of 62% and 36% respectively 59% and 97% respectively.⁶ The ankle reflex in our study showed sensitivity and specificity of 57% and 41% respectively compared to other studies showed sensitivity and specificity of 75% and 89% respectively.¹¹

Conclusion

In our study we found that the simple screening tests (10g SWME, vibrating test, superficial pain test and ankle reflex) can be used confidently for annual screening of diabetic neuropathy in diabetic patients.

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