NerveCheck: An inexpensive quantitative sensory testing device for patients with diabetic neuropathy

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ABSTRACT

Aims: Sensory neuropathy is central to the development of painful neuropathy, and foot ulceration in patients with diabetes. Currently, available QST devices take considerable time to perform and are expensive. NerveCheck is the first inexpensive ($500), portable QST device to perform both vibration and thermal testing and hence evaluate diabetic peripheral neuropathy (DPN). This study was undertaken to establish the reproducibility and diagnostic validity of NerveCheck for detecting neuropathy.

Methods: 130 subjects (28 with DPN, 46 without DPN and 56 control subjects) underwent QST assessment with NerveCheck; vibration perception and thermal testing. DPN was defined according to the Toronto criteria.

Results: NerveCheck’s intra correlation coefficient for vibration, cold and warm sensation testing was 0.79 (95% LOA: −4.20 to 6.60), 0.86 (95% LOA: −1.38 to 2.72) and 0.71 (95% LOA: −2.36 to 3.83), respectively. The diagnostic accuracy (AUC) for vibration, cold and warm sensation testing was 86% (SE: 0.038, 95% CI 0.79–0.94), 79% (SE: 0.058, 95% CI 0.68–0.91) and 72% (SE: 0.058, 95% CI 0.60–0.83), respectively.

Conclusions: This study shows that NerveCheck has good reproducibility and comparable diagnostic accuracy to established QST equipment for the diagnosis of DPN.

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1. Introduction

Quantitative sensory testing (QST) has been used for decades for diagnosing and quantifying the severity of DPN [1–5] and painful neuropathy [4,6–11]. Indeed several guidelines endorse the use of QST for the diagnosis of sensory abnormalities in diabetic neuropathy [2,12]. QST is an automated psychophysical method used to test vibration and thermal sensation which may help to risk stratify patients for the development of painful neuropathy, foot ulceration and amputation [13]. Easily deployed and inexpensive tests such as the tuning folk, pin-prick, VibraTip and 10 g monofilament can detect moderate to severe sensory loss but for early detection of sensory impairment, particularly in clinical trials, QST is required. It provides standardised and quantified stimuli which enable accurate quantification of sensory deficits [14] for vibration, a large fibre measure and thermal threshold testing for the detection of small fibre neuropathy [1–3,15].

Several QST devices are established and are primarily used in clinical research settings. The Neurothesiometer, VSA 3000 (Medoc), Vibrometer (Somedic), Vibration II (Physitemp) and Sensitometer are handheld devices but only perform vibration testing. CASE IV (WR Medical Electronics) measures the function of both vibration and thermal sensation but is large and expensive, provides a complex output in the form of just noticeable differences (JND) from a set of 25 standardised vibratory levels, and requires trained staff. The TSA-II-NeuroSensory Analyser (Medoc) and Sensor (Medoc) perform thermal testing only, are expensive and require a laptop to operate.

NerveCheck was designed to assess vibration (VFT), cold (CPT), warm (WPT) perception threshold and heat pain threshold (HPT). It costs ~$500 and is portable (size: 9.5 cm × 6.1 cm × 23.6 cm, weighted only 325 g including battery) as shown in Fig. 1. It applies a series of predefined stimuli over a broad range of intensities (i.e. vibration intensity, heat waveform and ramped stimuli (1 °C/s)) to the skin using the method of levels. For each stimulus, the subject reports whether the stimulus was perceived or not or whether it was painful or not. This method is not dependent on the reaction time of the subject. Thresholds for all four modalities are established within 9–13 min.

In the present study we have carefully validated the diagnostic ability of NerveCheck for assessing VFT, CPT and WPT in control subjects and patients with diabetes with a broad range of neuropathy. We have defined the thresholds and examined the reproducibility and diagnostic validity of NerveCheck against other established QST devices.

2. Subjects, materials and methods

The participants in the study were recruited from the Manchester Diabetes Centre, Manchester Royal Infirmary in Manchester, UK. The study was performed at the NIHR Wellcome Trust Clinical Research Facility between 7 January 2013 and 19 September 2014. Exclusion criteria included subjects with communication disorders, cognitive deficits, severe anxiety, severe depression or history of neuropathy due to a non-diabetic cause. Control subjects suffering from any acute or chronic pain condition were excluded. All subjects were without any pain medication for at least 24 h before the investigation. This study was approved by the Local Research Ethics committee and all patients gave informed consent to take part in the study. The research adhered to the tenets of the declaration of Helsinki.

2.1. Demographic measures

All study participants underwent assessment of their glycated haemoglobin (HbA1c), body mass index (BMI), blood pressure and cholesterol.

2.2. Quantitative sensory testing using NerveCheck

Subject were familiarised with the procedure and allowed to acclimatise for 10 min in the examination room. NerveCheck (Phi Med Europe S.L. Barcelona, Spain) applies the method of levels where a series of predefined stimuli (in terms of vibration intensity, heat waveform and ramped stimuli (1 °C/ s)) were applied to the skin and for each stimulus the subject reported whether the stimulus was perceived or not, to
establish the vibration (VPT), cold (CPT) and warm (WPT) perception thresholds. There are four kinds of stimuli for vibration and thermal testing. The VPT has void, mild (2.7 V), moderate (4.2 V) and strong (6.4 V) with 9 stimuli in total. The CPT has void, mild (22.4 °C), moderate (17.8 °C) and strong (9.8 °C) with 5 stimuli in total. The WPT has void, mild (37 °C), moderate (39.4 °C) and strong (44.7 °C) with 5 stimuli in total. If the null stimulus answered yes constantly, the result was deemed invalid and was repeated. This method is not dependent on the reaction time of the subject.

The order of administration of stimuli was vibration followed by thermal testing. The stimulator was applied with a constant pressure to the area of skin to be tested. For vibration testing, the vibratory transducer was placed on the dorsal surface of the base of the nail of the great toe. For thermal testing, the thermode (thermolectric unit with a surface area of 5 cm × 2.5 cm) was placed on the dorsolateral surface of the foot. The thermode provides accurate controlled minute ramps of cooling and heating at the thermal-testing surface using the Peltier effect. The administration of cooling and heating stimuli involves gradual changes in temperature along a linear ramp to a preset value and after a specified time return to steady state following an inverse ramp.

The output is categorical in terms of degree of abnormality. The more the subject response correctly to the stimuli the higher the score gets. The normal and abnormal range for VPT is (12–8 and 7–0) and for CPT and WPT is (6–3 and 2–0). The higher the grading score, the more sensitive the participant is to the stimuli. The testing takes 3–13 min, depending on whether it is a single test or series of tests. More information about NerveCheck can be found online (http://www.phimedeurop.com/).

2.3. Quantitative sensory testing using established devices

Vibration testing was measured using a Neurothesiometer (Horwell, Scientific Laboratory Supplies, Wilford, Nottingham, UK) and was placed at the base of the left great toe. The test was repeated three times and the average value was recorded. Thermal testing including CPT and WPT was undertaken on the dorsum of the left foot using the MEDOC TSA-II-NeuroSensory Analyser (Medoc Ltd. Ramat Yishai 30095, Israel) and method of limits [16]. The test was repeated four times and the average value was recorded.

2.4. Neuropathy assessments

All patients underwent an assessment of neuropathy based on a standard protocol including: NDS to classify participants into without (NDS 0–2) and with (NDS 3–10) neuropathy [17,18]. Electro-diagnostic studies were undertaken using a Dantec “Keypoint” system (Dantec Dynamics Ltd. Bristol, UK) equipped with a DISA temperature regulator to keep limb temperature constantly between 32 and 35 °C. Peroneal Motor Nerve Conduction Velocity (PMNCV) was assessed in the right lower limb by a consultant neurophysiologist.

2.5. Study definition of diabetic peripheral neuropathy

The Toronto Diabetic Neuropathy Expert group recommendation was followed to define DPN: (a) abnormal PMNCV (<42 m/s) [19] and (b) abnormal symptoms (NSP) or signs of neuropathy, NDS (>2) [18].

2.6. Statistical analysis

We estimated that the minimum sample required to detect significant difference between the group with and without vibration sensation loss is 16 subjects and between the group with and without thermal sensation loss is 26. The sample size was calculated by means of an unpaired t-test and with a power of 85%. We examined the distribution of the data by means of relevant histograms and the Shapiro–Wilk test using StatsDirect statistical software, version 2.7.9. All data were expressed as median (5th percentile, 95th percentile). Mann–Whitney U test was performed to analyse differences between the medians. A P value <0.05 was considered statistically significant.

We examined the repeatability of NerveCheck with intraclass agreement using GraphPad Prism, version 6.05. The test–retest intervals were from 1 to 8 weeks references.

Receiver operating characteristic (ROC) curve analysis was used to compare the diagnostic accuracy of NerveCheck against established QST devices using GraphPad Prism, version 6.05. ROC curve analysis established the area under the curve (AUC) to determine the optimal sensitivity and specificity of the NerveCheck test.

3. Results

3.1. Clinical data

130 subjects (74 with diabetes mellitus (DM) (59 Type 1 DM and 15 Type 2 DM) with median age 55.7 (interquartile range – IQR: 42.9–66.1) and 56 control subjects with median age 43.6 (interquartile range – IQR: 35.7–53.1)) were studied. Of the diabetic subjects 28 were diagnosed with and 46 without diabetic peripheral neuropathy (DPN) based on the Toronto criteria. The demographic and clinical characteristics of the participants with and without DPN and controls are presented in Table 1. BMI, HbA1c and cholesterol levels did not differ between the groups with and without DPN, but age (P < 0.0001), duration of diabetes (P < 0.0001) and systolic blood pressure (P = 0.0005) were significantly greater in those with DPN. The group with DPN had a significantly higher Neuropathy disability score (NDS) (P < 0.0001) and significantly lower peroneal motor nerve conduction velocity (PMNCV) (P < 0.0001). In the NerveCheck tests, the group with DPN had a significantly lower score for vibration perception threshold (VPT) (P < 0.0001), cold perception threshold (WPT) (P < 0.0001) and warm perception threshold (WPT) (P < 0.0001) compared to those without DPN.

3.2. NerveCheck defined threshold values

To define a threshold value of the NerveCheck grading score for VPT, CPT and WPT we have used a mean minus 2 standard
deviation (SD) cut-off, based on our control population (n = 56).
The normal range of the NerveCheck for VPT, CPT and WPT is 4–12 for VPT, 3–6 for CPT and 3–6 for WPT. The total grading score for VPT, CPT and WPT is 0–12, 0–6 and 0–6, respectively.

3.3. NerveCheck reproducibility

Controls and subjects with diabetes (n = 16) were tested on two separate occasions to examine the intraclass agreement with test-retest intervals ranging from 1 to 8 weeks. The NerveCheck has good reproducibility as the intraclass agreement for VPT, CPT and WPT is 0.79 (95% limits of agreement: –4.20 to 6.60), 0.86 (95% limits of agreement: –1.38 to 2.72) and 0.71 (95% limits of agreement –2.36 to 3.83), respectively.

3.4. NerveCheck diagnostic validity for diabetic peripheral neuropathy

The diagnostic performance of the NerveCheck in detecting sensory loss measured against established QST devices is expressed in AUC using ROC curve analysis (Fig. 2). The VPT and thermal testing were compared against the Neurothesiometer and TSA-II-NeuroSensory Analyser (Medoc), respectively. The AUC for VPT is 86% (SE: 0.038, 95% CI 0.79–0.94, P < 0.0001), for CPT 79% (SE: 0.058, 95% CI 0.68–0.91, P < 0.0001) and for WPT 72% (SE: 0.058, 95% CI 0.60–0.83, P < 0.0004).

The VPT of the NerveCheck displayed high sensitivity 84% (95% CI 63.92–95.46%) and high specificity 81% (95% CI 72.07–87.66%) with a likelihood ratio of 4.36 for the diagnosis of DPN. The CPT exhibited high sensitivity 89% (95% CI 81.72–94.23%) and moderate specificity 67% (95% CI 46.04–83.48%) with likelihood ratio 2.67 for the diagnosis of DPN. The WPT exhibited high sensitivity 75% (95% CI 65.86–83.14%) and moderate specificity 66% (95% CI 45.67–82.06%) with likelihood ratio 2.18% for the diagnosis of DPN.
categorical output which can easily be interpreted in relation to the severity of neuropathy and hence risk stratification. It is the first inexpensive ($500), portable (9.5 cm × 6.1 cm × 23.6 cm) QST device to perform both vibration and thermal testing. We have demonstrated good reproducibility and validity of NerveCheck for assessing sensory loss compared to established devices and give its far lower cost we would suggest much wider use in the clinic. Specifically, NerveCheck has good reproducibility and its ability to detect sensory loss is highly comparable to established QST devices.

When interpreting individual QST results based on the normal range in the NerveCheck grading score, age should be taken into consideration [22]. The normal range of the NerveCheck for VPT, CPT and WPT in the current study was based on our control population with a median age of 43.6. The normal range in the NerveCheck grading score applies to the age range 30–72. Indeed there are very few large-scale reference data sets for QST in older adults [23]. Both vibration and thermal testing have been reported to be independent of gender [24]. In the current study we did not evaluate thermal allodynia or hyperalgesia, as the available normal range for these heat pain thresholds is highly variable [6,25].

The reproducibility of QST has been a challenge with a significant variability between sessions in the same patient. Indeed in a very early study over 30 years ago, Fagius et al. reported a difference of 150% between assessments using the method of limits [26]. This problem appears lesser marked though with the method of levels [27] and NerveCheck uses the method of levels where it applies a series of predefined stimuli over a broad range of intensities. For each stimulus, the subject reports whether the stimulus was perceived or not or whether it was painful or not and is not dependent on the reaction time of the subject. Null stimuli have been included in the algorithms to reduce bias related to a false response. The vibration testing of NerveCheck produces low variability and a factor that may be relevant is that the vibration stimulus is applied with a constant pressure to the area of skin to be tested. Additionally, NerveCheck runs the vibration testing before the thermal testing as assessment of thermal sensation before vibration testing has been found to increase the risk of vibration hyperalgesia [28]. Of relevance, several studies have shown good reproducibility of VPT (intraclass correlation >0.55) in control subjects and in various patient populations [3,29]. All said, the thermode of the NerveCheck is a highly engineered device that can provide cooling and heating stimuli along a linear ramp to a preset value at the thermode testing surface using the Peltier mechanism as described by Dyck et al. [14].

QST is an effective technique for the diagnosis of sensory neuropathy and also provides a composite of quantitative measures, which can be deployed to define the severity of neuropathy. Vibration deficits in the feet suggest large fibre dysfunction [21] and cold and warm deficits indicate small fibre dysfunction [21]. Dysfunction of small nerve fibres is thought to be responsible for many painful peripheral neuropathies [30]. These small fibre neuropathies cannot be evaluated using standard electrophysiological testing [1–3,15]. Our study shows that NerveCheck has both high sensitivity 84% and high specificity 81% for vibration testing and high sensitivity and moderate specificity for thermal testing. Of relevance, studies have reported variable sensitivity of thermal testing depending on the severity of neuropathy, 27–98% for cold and 22–98% for warm deficits [3], while vibration testing has 58–84% sensitivity and 51–86% specificity [31,32].

NerveCheck provides a cost effective means to identify deficits in vibration and thermal sensation. Both Medoc TSA-II NeuroSensory Analyser and NerveCheck detect sensory deficits. The former test provides the patient’s perception threshold for cold and warm. However, unlike TSA-II NeuroSensory Analyser, the NerveCheck indicates whether the results are normal or abnormal and stratifies the severity of sensory loss. Additional evaluation may include an assessment of Neuropathic Impairment Score (NIS), autonomic dysfunction via the Neuropad or the assessment of heart rate variability to deep breathing [33,34] and small fibre structural damage using corneal confocal microscopy (CCM) or skin punch biopsy for intra-epidermal nerve fibre density (IENFD) [35,36].

In conclusion this is a small but detailed study showing that NerveCheck has good reproducibility and good diagnostic accuracy for assessing sensory loss compared to established QST devices. Clearly larger, prospective studies confirming the diagnostic ability of NerveCheck are required in diabetic neuropathy and in other neuropathies. NerveCheck is the first inexpensive, portable QST device to perform both vibration and thermal testing and therefore provides new opportunities for use in the clinic. It could therefore be deployed as an inexpensive but accurate diagnostic test for diabetic neuropathy in primary care in the developed world and throughout the developing world; which is set to see an explosion in diabetes and hence diabetic neuropathy. It could also be deployed in countries such as India and Brazil for conditions such as leprosy.

Author contributions


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Conflict of interest

We wish to draw the attention of the Editor to the following facts which may be considered as potential conflicts of interest and to financial contributions to this work. Phi Med Europe S.L. provided the Nervecheck device, funded half of the
study and covered some of the travel expenses and conference registration fees for presenting the results of this study. M.N. Odrioza, S. Odrioza, M.B. Odrioza and A. Odrioza are the owners and inventors of the NerveCheck.

We confirm that the manuscript has been read and approved by all named authors and that there are no other persons who satisfied the criteria for authorship but are not listed. We further confirm that the order of authors listed in the manuscript has been approved by all of us.

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