Medical Policy Manual

Topic: Automated Point-of-Care Nerve Conduction Studies  
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IMPORTANT REMINDER

Medical Policies are developed to provide guidance for members and providers regarding coverage in accordance with contract terms. Benefit determinations are based in all cases on the applicable contract language. To the extent there may be any conflict between the Medical Policy and contract language, the contract language takes precedence.

PLEASE NOTE: Contracts exclude from coverage, among other things, services or procedures that are considered investigational or cosmetic. Providers may bill members for services or procedures that are considered investigational or cosmetic. Providers are encouraged to inform members before rendering such services that the members are likely to be financially responsible for the cost of these services.

DESCRIPTION

Nerve conduction studies (NCS) and needle electromyography (EMG), when properly performed by a trained practitioner, are considered the gold standard of electrodiagnostic testing, and may be used to diagnose neuropathies (such as carpal tunnel syndrome [CTS] or peripheral neuropathy). However, the need for specialized equipment and personnel, along with additional time and cost, may limit the availability of electrodiagnostic testing for all patient populations. Automated point-of-care nerve conduction studies, also known as automated nerve conduction studies, are performed using portable devices with computational algorithms that are able to drive stimulus delivery, measure and analyze the response, and provide a report of study results. They have been proposed for use by nontechnical clinic personnel as an alternative diagnostic test for CTS and other neuropathies.

Regulatory Status

Several automated nerve conduction study devices have received clearance by the U.S. Food and Drug Administration (FDA). The majority of devices have 510(k) approval, a type of clearance which does not require data regarding clinical efficacy. Examples of devices currently on the market include:

- NeuroMetrix received specific FDA clearance to market the NC-stat® device “to measure neuromuscular signals that are useful in diagnosing and evaluating systemic and entrapment...
Neuropathies.” In addition, the approved application stated that “The NC-stat is intended to be used as an adjunct to and not a replacement for conventional electrodiagnostic measurements.”

- NeuroMetrix subsequently received FDA clearance to market newer models with biosensors and engineering changes that enable the NC-stat to be used for motor and sensory nerves of the wrist (median and ulnar) and foot (peroneal, tibial, and sural). The intended use as listed on the 510(k) approval from 2006 (K060584) is “to stimulate and measure neuromuscular signals that are useful in diagnosing and evaluating systemic and entrapment neuropathies.”

- The NeuroMetrix ADVANCE™ system received marketing clearance in 2008 (K070109). It is intended to perform nerve conduction studies using disposable surface electrodes (similar to NC-stat) with an additional module for invasive needle EMG. The ADVANCE™ system includes a real-time display of nerve conduction waveforms with a stylus for assignment of waveforms.

- The XLTek Neuropath (Excel-Tech) received clearance for marketing through the FDA’s 510(k) process in 2006; the indications are the same as those for NC-stat.

- The Neural-Scan™ NCS (Neuro Diagnostics) is a Class I diagnostic device (FDA clearance not usually required) that is being marketed “as part the [sic] neurological examination or for screening to detect peripheral neuropathies.”

- The Axon-II™ (PainDx) is an automated system that is being marketed for the detection of various sensory neurological impairments caused by various pathological conditions or toxic substance exposures, including signs of sympathetic dysfunction and detection of down-regulated A-delta function to locate injured nerve(s) The AXON-II software works with the Neural-Scan™ system (Neuro Diagnostics) and lists 7 automated studies (Cervical, Thoracic, Lumbar, Upper Extremities, Lower Extremities, Neuroma, Trigeminal) as well as a custom study. The Neural-Scan™ is a voltage-actuated sensory nerve conduction test device, which measures the voltage amplitude necessary to cause a discernible nerve impulse. Results are adjusted and compared to population means; the most severe hypoesthesia is considered the primary lesion.

- The Brevio® from Neurotron Medical received marketing clearance from the FDA in 2001. The Brevio® is intended “for use for the measurement of nerve response latency and amplitude in the diagnosis and monitoring of peripheral neuropathies.”

- The Mediracer® NCS (Mediracer Ltd.) is not currently available for marketing in the United States as it does not have FDA approval at this time.

MEDICAL POLICY CRITERIA

Automated nerve conduction tests are considered investigational for the diagnosis of all indications, including but not limited to carpal tunnel syndrome, peripheral neuropathy, and lumbosacral radiculopathy.
Automated nerve conduction tests have been proposed for use as a diagnostic tool for carpal tunnel syndrome (CTS), a pressure-induced entrapment neuropathy of the median nerve as it passes through the carpal tunnel, resulting in sensorimotor disturbances. CTS is defined by its characteristic clinical symptoms, which may include pain, subjective feelings of swelling, and nocturnal paresthesia. Automated portable nerve conduction device studies have also been proposed for the diagnosis of diabetic neuropathy, a condition which is relatively common in patients with diabetes mellitus and which can lead to important morbidity including pain, foot deformity, and foot ulceration. Lumbosacral radiculopathy, another common neurologic disorder, wherein damage to disks in the lumbar and or sacral region of the back causes nerve irritation and damage, is another proposed indication for these types of devices.

Validation of any new diagnostic technique focuses on three parameters:

1. Demonstration of technical feasibility

   Normally conducted in the pre-clinical setting, the focus of this parameter is on test reproducibility and establishment of test protocol. Technical feasibility of a device is typically assessed with two types of studies, those that compare test measurements with a gold standard and those that compare results taken with the same device on different occasions (test-retest).

2. Demonstration of diagnostic performance

   Diagnostic performance is evaluated by the ability of a test to accurately diagnose a clinical condition in comparison with the gold standard. For accurate interpretation of study results, sensitivities, specificities, and positive and negative predictive values compared to a gold standard must be known. The sensitivity of a test is the ability to detect a disease when the condition is present (true positive), while specificity indicates the ability to detect whether disease exists in patients who are suspected of disease but who do not have the condition (true negative). Evaluation of diagnostic performance, therefore, requires independent assessment by the two methods in a population of patients who are suspected of disease but who do not all have the disease. Studies that do not meet these criteria (broad patient population and comparison of point-of-care use with the standard laboratory NCS-EMG) may be considered relevant to the technical feasibility of the device, but are inadequate for evaluation of its diagnostic performance.

Within this context, the comparators for automated point-of-care nerve conduction studies are as follows:

- While there is no absolute gold standard for diagnosis of CTS, electrodiagnostic studies are commonly used to quantify the amount of nerve damage, and a positive response to conservative management (steroid injection, splints, and modification of activity) can confirm the clinical diagnosis.[1]

- Diagnosis of peripheral neuropathy is often made clinically through the physical examination, combined with simple sensory tools such as the 10-g Semmes-Weinstein monofilament or the 128-Hz vibration tuning fork, although there is no standard protocol for the diagnosis of this condition.[2] Electrodiagnostic studies may also be used to confirm the
presence or absence of diabetic neuropathy.

- Diagnosis of lumbosacral radiculopathy is normally made through patient history and physical examination and can be quantified and confirmed with MRI or nerve conduction studies (NCS) and needle electromyography (EMG).[3]

3. Evaluation of clinical outcomes

Also known as clinical utility, this parameter evaluates how the results of the new test can be used to benefit patient management and its impact on health outcomes. The clinical utility of both positive and negative tests must be assessed as it relates to any added benefits which cannot be achieved by the standard of care. While in some cases, new diagnostic tests can be adequately evaluated using technical and diagnostic performance, when a test identifies a new or different group of patients with a disease, randomized trials are needed to demonstrate the impact of the test on net health outcomes.

**Literature Appraisal**

Several case series have been conducted on the technical feasibility and diagnostic accuracy of automatic nerve conduction studies, mainly those performed with NC-stat® technology, and one case series reports on the clinical utility of the device. There are no randomized studies comparing automated nerve conduction studies with nerve conduction and needle electromyography (NCS-EMG) studies.

**Diabetic Peripheral Neuropathy**

- A study of motor nerve function compared NC-stat with standard nerve conduction tests of the wrist in a small study of 17 subjects with diabetes mellitus who had clinical evidence of peripheral neuropathy in either the upper or lower extremity.[4] Correlations between the tests results were high (ranging from 0.70 for ulnar distal motor latency (DML) to 0.96 for median nerve DML).

- Another study assessed the validity of NC-stat to diagnose diabetic peripheral neuropathy through sural nerve testing in patients from diabetes and diabetic neuropathy outpatient practices.[5] Seventy-two consecutive patients (64 with type 2 diabetes) who completed a clinical evaluation, a conventional nerve conduction study, and a point-of-care NC-stat assessment were enrolled. The point-of-care assessment was independently conducted by non-technologist research staff following a single 1-hour lesson in the NC-stat protocol. The amplitude potential of the sural nerve was tested as an early indicator of diabetic neuropathy. Using a threshold of 6 µV, the authors report that the sensitivity and specificity of NC-stat for diagnosis of diabetic sensorimotor polyneuropathy, as defined by clinical and conventional electrophysiological evaluation, was 92% and 82%, respectively. The Spearman correlation coefficient (compared with the reference standard) was 0.95. As noted by the authors, further study is needed in a broad spectrum of patients, including those who present with atypical neuropathy in a clinical setting. The authors also note that further investigation is needed into specific approaches that include the point-of-care nerve conduction study as a component of the clinical care of those with polyneuropathy.

- Pambianco and colleagues compared the accuracy of two automated point-of-care nerve conduction study devices, the NC-stat and the Neurometer R-CPT (Neurotron, Inc.), along with sensory threshold devices, the Michigan Neuropathy Screening Index (MNSI) and the 10-g
Semmes-Weinstein monofilament test in a cohort of 195 patients with type 1 diabetes. The accuracy of each diagnostic test was compared with a standard clinical exam protocol for the diagnosis of two outcomes: diabetic peripheral neuropathy and amputation, ulcer or neuropathic pain. Estimates of sensitivity and specificity of the NC-stat device were 79% and 48%, respectively, for detection of diabetic peripheral neuropathy, and 77% and 38% for the detection of amputation, ulcer, or neuropathic pain. The MNSI had the highest combination of sensitivity (87% and 80%) and specificity (49% and 36%) for each outcome. The authors conclude that the reduced specificity of the point-of-care nerve conduction study devices limits their use as a diagnostic tool for individuals with type 1 diabetes, and that the MSNI presents the best combination of sensitivity and specificity out of the diagnostic tools considered.

In summary, estimates of diagnostic accuracy of automated point-of-care nerve conduction study devices vary. The largest study of diagnostic accuracy recommended the use of the Michigan Neuropathy Screening Index over point-of-care nerve conduction studies, citing superior estimates of sensitivity, specificity, time, and cost. Additional studies are needed to establish a consensus in regards to the diagnostic performance and clinical utility of this test.

Carpel Tunnel Syndrome (CTS)

- In an early report of the NC-stat technology using DML to diagnose CTS, Leffler and colleagues reported that in 248 symptomatic hands (apparently a combination of an initial and validation group), compared with conventional diagnosis, testing using this device had a sensitivity of 86% and specificity of 90%. [7]

- Another study compared results from NC-stat and standard nerve conduction studies in a previously diagnosed patient population. [8] This study compared distal motor latency of the median nerve in 72 patients (of 400 treated) with established CTS before and after surgical intervention, finding a correlation coefficient of 0.88 for the median nerve DML. However, a scatter plot indicates a poor correlation for longer latencies.

- A Pearson correlation coefficient of 0.944 was reported for DML for 46 patients with CTS who had a nerve conduction study at a different time (average of 28 days difference). [9]

- In the report by Rotman, the NC-stat DML was shown to have a sensitivity of 89% “at the predetermined specificity of 95%” for the diagnosis of CTS for “70 hands” that met the standardized CTS case definition. [9] However, in a point-of-care study evaluating industrial workers for possible CTS using distal motor latency, many individuals who were identified with prolonged DML by NC-stat fell within the normal range (using 95% cutoff point) as defined by this study population. [10] This study also comments on the importance of sensory nerve findings in the diagnosis of CTS, suggesting a need to better define “normal” values.

- The NeuroMetrix data registry was analyzed for all NC-stat studies performed over a period of 10 days that were coded for CTS and performed by a primary care provider. [11] The initial data set consisted of studies on 1,190 patients performed by 613 different physician practices; studies that met CTS testing guidelines (82% met strict guidelines and 93% met less restrictive guidelines) were further analyzed. Thus, in nearly 1 of 5 patients (18.4%), the studies did not meet strict CTS testing guidelines. From the limited set, 31% were identified as normal, 53% exhibited CTS, 5% demonstrated an ulnar neuropathy, and 11% showed a nonspecific neuropathy. No comparison was made with standard nerve conduction testing nor was an
assessment made of the impact of this testing on relevant clinical outcomes.

- In 2011, Bourke et al. reported a non-randomized comparison of clinic-based NC-stat versus referral to standard electrodiagnostic testing that evaluated efficiency of work-up and costs. The study included 142 patients being considered for decompression surgery for CTS at a hand clinic.\[12\] Seventy-one patients who accepted nerve conduction studies in a nurse-led clinic were compared with 71 historical controls who had been sent for nerve conduction studies at the regional neurophysiological unit. Patients with known or suspected complex neurological conditions were excluded from the study. Outcome measures were time from presentation to carpal tunnel decompression, the cost of each pathway, and the practicalities of using the device in the clinic. In the NC-stat group, 43 patients (61%) had a diagnosis of CTS confirmed by NC-stat and underwent decompression surgery and 28 patients (39%) had normal or inconclusive tests. Of the 28, 12 were referred for electrodiagnostic testing and 2 of the 12 were recommended for decompression surgery (3% false negative). In the referred group, 44 patients (62%) had confirmation of CTS and underwent decompression surgery. Use of NC-stat in the clinic reduced the time from presentation to surgery from 198 days to 102 days. Cost saving for NC-stat was reduced by the need to refer nearly 20% of patients for standard electrophysiological testing, but still favored the clinic-based approach. Health outcomes for the two approaches were not assessed.

In summary, the published literature on the use of the NC-stat device for the diagnosis of CTS is focused on the technical feasibility and diagnostic performance of the device. Questions remain regarding the optimal range of the device and the target population. Further studies are required to demonstrate technical feasibility, diagnostic accuracy, and clinical utility of this device.

Lumbosacral Radiculopathy

- Fisher and colleagues explored the relationship between NC-stat and routine NCS/needle electromyography (EMG) in 34 consecutive patients with a clinical history and/or examination consistent with lumbosacral radiculopathy.\[13\] Inclusion in the study was based on chart review of symptoms from clinical history and/or examination (including low back pain or buttock pain, numbness, and/or paresthesias of one or both lower extremities) and having undergone testing with both NC-stat and routine electrodiagnostic studies. Of the 34 patients included in the study, 28 had magnetic resonance imaging (MRI) of the lumbosacral spine within 6 months of electrodiagnosis, 2 had a post-myelogram computed tomography (CT) scan, and 3 had lumbosacral spine radiographs. A neuroradiologist who was blinded to the clinical evaluation and electrodiagnostic results determined from MRI or CT that lumbosacral root injury was likely at the L4-5 and/or L5-S1 levels in 18 patients (60%). The study found some correlation between the electrodiagnostic testing and NC-stat. However, 6 of 10 patients who had unremarkable routine electrodiagnostic results had abnormal F-wave and compound muscle action potential (CMAP) amplitude abnormalities with NC-stat testing. The clinical implications of this finding are uncertain.

- A 2011 report by Schmidt and colleagues assessed the accuracy of NC-stat diagnosis of lumbosacral radiculopathy in 50 patients and 25 controls with no prior history of lumbosacral radiculopathy.\[14\] The patient cohort included patients referred to a tertiary referral EMG laboratory for testing of predominantly unilateral leg symptoms (pain, numbness, or weakness). Control subjects were recruited from clinic employees and from patients referred to the EMG laboratory for upper limb symptoms. All patients underwent focused history and physical
examination and both standard and automated electrodiagnostic testing. Automated testing was performed by experienced technicians who were unaware of the electrodiagnostic test results. In the patient cohort, the sensitivity of NC-stat was found to be 0% for L4 radiculopathy, 69% for L5 radiculopathy, and 64% for S1 radiculopathy compared with standard electrodiagnostic testing. By standard electrodiagnostic evaluation, 22 of the 50 symptomatic patients had findings consistent with L4, L5 or S1 radiculopathy and 28 patients were found to be normal or to have a diagnosis other than lumbosacral radiculopathy; NC-stat identified only 4 of these 28 cases (specificity of 14%). Standard electrodiagnostic testing also identified other important diagnoses in 9 patients (18%) that were not identified by the automated test, while NC-stat reported 6 other diagnoses in patients found to be normal by standard electrodiagnostic testing. All standard electrodiagnostic tests in the control group were normal, but the automated test found that 18 of these subjects were abnormal (specificity of 32%). The study found that the raw nerve conduction data were comparable for the 2 techniques; however, computer-generated interpretations by the automated device showed low specificity (numerous false positives) in both symptomatic patients and normal control subjects. An accompanying editorial by England and Franklin states that the use of automated nerve conduction devices is controversial, and that the use of NC-stat for lumbosacral radiculopathy would likely lead to a high misdiagnosis rate and potentially inappropriate treatment, including surgery. England and Franklin also conclude that an overly sensitive but not very specific test for carpal tunnel syndrome, or other mono- or polynuropathies, cannot replace expert use and interpretation of conventional electrodiagnostic testing.

Early reports of diagnostic accuracy of this device are suggestive that the NC-stat device may have a broader sensitivity and lower specificity when compared with conventional electrodiagnostic testing. Additional studies of diagnostic accuracy are needed, along with randomized controlled trials to ensure that new patients identified with this test benefit from treatment decisions based upon test results.

General Neuropathy

- A 2010 publication by NeuroMetrix reported test-retest reproducibility with the ADVANCE™ system in 30 subjects with symptoms suggestive of neuropathies; 29 subjects completed the study. Coefficients of variation ranged from 4.2% to 9.8% for tests measured 3-7 days apart. Between session intraclass correlation coefficients (ICCs) ranged from 0.98 for F-wave latency to 0.77 for sural sensory conduction velocity.

- Another study evaluating technological performance compared results for sensory nerve testing from NC-stat and the reference standard in median and ulnar nerves in 60 patients referred to an EMG laboratory for neck and shoulder pain who also volunteered to undergo testing with NC-stat. The reported correlations (Pearson correlation) between the NC-stat and the reference standard were high (0.91 for median nerve distal sensory latency [DSL], 0.70 for ulnar DSL, and 0.88 for the median ulnar difference of the distal sensory latency). However, this final correlation was calculated only with the responses obtained for 81 of 120 possible nerve pairs. The authors of this study report systematic differences between the two techniques and indicate that use of the NC-stat would require applicable reference ranges.

- Another NeuroMetrix-sponsored trial compared NC-stat and standard EMG results for peroneal and posterior tibial nerve conduction in 60 patients referred to an EMG laboratory. The report indicated that all patients referred to the laboratory were offered the opportunity to participate, but did not provide the total number of referrals. F-wave latency (FLAT) was found to have the
highest correlation (0.91, 0.90 Spearman correlation coefficient for peroneal and posterior tibial nerves, respectively), with moderate correlations for amplitude (0.86, 0.73) and distal motor latency (0.70, 0.45). Although NC-stat results were significantly correlated with standard EMG tests in the study population as a whole, in a subgroup analysis of the most abnormal half of responses, the correlation coefficient for amplitude of the peroneal response was 0.62, and the correlation coefficient for distal motor latency was reduced to 0.32 for the posterior tibial nerve and 0.10 for the peroneal nerve. Thus, in this pathological subgroup analysis, criterion validity was lost for the peroneal distal motor latency and decreased from “excellent” to “acceptable” for the other parameters. The authors note that “this study did not address interpretations performed by physicians using NC-stat data, nor the validity of the reference ranges used or the way these were collected.”

- In 2010, a case series report was published on the use of the NC-stat technology for the diagnosis of any neuropathy within a primary care clinic in Utah. Diagnostic results of the NC-stat device were compared with pre-test diagnosis (methodology of diagnosis not specified). In 59 of 100 tests, results from the NC-stat agreed with pre-test diagnosis (Kappa statistic not reported). 7 of 100 tests were not reported because of incomplete data or issues relating to the technology. There was no patient selection criteria, limiting conclusions about which type of patients may benefit from this diagnostic test. In addition, it was not specified how the pre-test diagnosis was made, with the reference standard diagnostic test (NCS with EMG) or otherwise.

- A 2008 report assessed the diagnostic performance of NC-stat against the gold standard NCS in patients who had been referred for electrodiagnostic testing at one of several academic medical centers. Of 47 patients who were invited to participate in the study, 14 patients declined to participate or were excluded due to missing records, resulting in data analysis on 33 patients. The goal of the study was to compare the measurements of the two methods of nerve conduction testing as they would be used in standard practice, thus, patients were not excluded on the basis of the particular diagnosis for which they were referred. The diagnosis being tested was carpal tunnel syndrome in 25 patients (76%), with the remaining 8 patients having 8 other potential diagnoses, including ulnar neuropathy, upper extremity paresthesias, and C6 radiculopathy. NC-stat results could not be obtained for 2 patients for median motor studies and 3 patients for median sensory studies (15%). Based on the manufacturer’s suggested cutoff for abnormal nerve conduction, sensitivity was 100% for both the motor and sensory median-ulnar difference; specificity was 62%–69% for the motor median-ulnar difference and 41% to 47% for the sensory median-ulnar difference. Pearson correlation coefficients ranged from 0.40 for the ulnar nerve to 0.91 for the median dorsal motor nerve. The authors concluded that the recommended cutoff values for NC-stat may need to be adjusted, although the specific study results were limited by the small sample size. In addition, the authors noted that the study did not evaluate how well physicians can assign clinical relevance to the results, and that while the device may be suited for research studies or screening of symptomatic patients, “in many clinical situations referral to a specialist for a more comprehensive evaluation would be prudent.”

In the majority of studies on the use of the NC-stat device for the diagnosis of unspecified neuropathies, patient selection criteria were not defined, meaning conclusions from these studies may not apply to a broad population of patients with suspected neuropathy. Results from the single study of diagnostic performance should be interpreted with caution as subject attrition may have inflated estimates of accuracy (if more complicated cases were dropped from the analysis). Additional studies on patients suspected with specific types of neuropathy are needed to establish technical feasibility, diagnostic accuracy, and clinical utility.
**Healthy Patients**

- NeuroMetrix reported intra-operator reliability in 15 healthy subjects who underwent measurements 7 days apart, evaluating technical feasibility of the test.\[^{21}\] The report states that “each upper and lower extremity nerve was tested twice by the same technician,” and that 9 subjects participated in both upper and lower extremity studies. It is not clear from the report whether the upper and lower extremities were designed as separate studies, or if 12 of 42 (29%) measurements did not provide usable data. Of the data reported, the coefficient of variation ranged from 0.013 for F-wave latency to 0.298 for the compound muscle action potential amplitude of the peroneal nerve.

- In 2009, NeuroMetrix published a study of reference ranges for key nerve conduction parameters in healthy subjects.\[^{22}\] Data analyzed in the paper were pooled from 5 studies, including from 92 to 848 healthy subjects with data on the median, ulnar, peroneal, tibial, and sural nerves. Subject age and height were found to affect the parameters. In addition to providing reference ranges for clinicians to use (providing that NCS techniques are consistent with those described in the paper), the authors stated that clinicians could use the same method to develop their own reference ranges. At this time, the proposed reference ranges have not been validated in a clinical patient population.

Conclusions about the technical feasibility, diagnostic performance and clinical outcomes of this device as used on healthy subjects may not be valid and reliable for subjects with neuropathy.

**Clinical Practice Guidelines**

Several clinical practice guidelines have addressed the use of automatic nerve conduction studies:

In 2006, the American Association of Neuromuscular & Electrodiagnostic Medicine (AANEM) issued a position statement that maintains standardized nerve conduction studies performed independent of needle EMG studies may miss data essential for an accurate diagnosis, and nerve disorders are far more likely to be misdiagnosed or missed completely if a practitioner without the proper skill and training is interpreting the data, making a diagnosis, and establishing a treatment plan.\[^{23}\] The organization states that, “the standard of care in clinical practice dictates that using a predetermined or standardized battery of NCSs for all patients is inappropriate,” and concludes that, “It is the position of the AANEM that, except in unique situations, NCSs and needle EMG should be performed together in a study design determined by a trained neuromuscular physician.”

**Summary**

Studies on portable automated point-of-care nerve conduction test results have shown a correlation with standard electrodiagnostic testing; however, questions remain about the diagnostic performance and clinical utility (i.e., impact on health outcomes) of point-of-care automated testing. Particularly needed are data on the sensitivity and specificity of automated nerve conduction tests performed by non-specialists at the point-of-care in comparison with the “gold standard” of laboratory nerve conduction studies/electromyography (NCS/EMG). There is no peer-reviewed published medical literature on the use of these portable tests for clinical decision-making and their impact on clinical outcomes. Overall, evidence remains insufficient to evaluate the effect of automated point-of-care nerve conduction tests on health outcomes. Therefore, automated point-of-care nerve conduction tests are considered...
investigational.

REFERENCES


**CROSS REFERENCES**

*Quantitative Sensory Testing*, Medicine, Policy No. 91

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