Ulnar neuropathy at the elbow (UNE) is a common site for compression in the arm, second only to median mononeuropathy at the wrist. Relying on sensory symptoms to secure a diagnosis is inherently problematic as nearly 40% of subjects with paresthesias in the ulnar distribution have been subsequently diagnosed with carpal tunnel syndrome. Current techniques to evaluate UNE are not as sensitive as diagnosing other mononeuropathies in the upper extremity, with a high risk of false negatives especially with mild nerve injury. The poor sensitivity can, at least partially, be explained by numerous pitfalls that have been described when assessing motor nerve conduction studies (NCS) of the ulnar nerve. Such aspects that limit utility include, measurement error, selective fascicular involvement, challenging data interpretation, and inability to adequately localize the lesion. Measurement variability (standard deviation) of the across elbow distance increases when the elbow is flexed. The absence of weakness or atrophy in muscles innervated by the ulnar nerve is another
important factor that potentially limits the functionality of electrodiagnostic testing as these patients have more subtle abnormalities.1,4,8,9 Interestingly, previous research indicates that subclinical axon loss can still occur despite patient’s description of purely sensory symptoms.4 Due to these numerous problems, a multiorganizational summary statement regarding identifying ulnar neuropathy recommends using multiple electrodiagnostic studies that agree to diagnose UNE rather than one isolated result.10

A unique NCS measuring a mixed compound nerve action potential (CNAP) across the elbow has been described using two different techniques.2,11 One technique still required elbow flexion, which did not solve issues with measurement error and only included 10 symptomatic patients.11 The goal of this present study was to establish a correlation between this uncommon NCS, mixed across the elbow, and other more common ulnar motor NCS to further aid diagnosing UNE. Specifically, the slower ulnar velocity across the elbow should correlate with a larger difference between the distal latencies of the ulnar and median nerves.

**Methods**

**Data Collection**

Prior to data collection, this study was approved by the Institutional Review Board at both Michigan State University and McLaren Healthcare with a waiver for informed consent and in accordance with The Code of Ethics of the World Medical Association (Declaration of Helsinki).

A retrospective analysis included all patients who presented to an academic-based electrodiagnostic laboratory for evaluation from January 2013 to June 2019. Inclusion criteria required three comparative NCS to be completed at the time of evaluation with associated sensory or sensorimotor symptoms in the ulnar nerve distribution (digit 4 and/or 5). The three acceptable NCS were ulnar motor to abductor digiti minimi (ADM), ulnar motor to first dorsal interossei (FDI), and mixed across the elbow. Upon chart review completion, 135 patients met the original inclusion criteria, however, two of those patients had absent motor responses above the elbow. This left 133 patient charts that would satisfy all data points for statistical analysis and began electrophysiologic testing with excluded UNE. Amplitudes for the mixed NCS have large standard deviations and thus not reliable to separate disease from control,1,2 therefore, although these data were collected, they were not thoroughly analyzed, other than a range from 1 to 40 microvolts for the median nerve and 1 to 42 microvolts for the ulnar nerve. Distal latencies, amplitudes, velocities were recorded for the previously mentioned NCS (FDI, ADM, and mixed across the elbow) as well as median motor to abductor pollicis brevis, sensory conduction to digits 1,2,5, and sural as applicable. None of the selected patients were diagnosed with median mononeuropathy at the wrist. Signs of acute denervation (fibrillation potentials) were differentiated between C8-T1 ulnar innervated muscles and C8-T1 nonulnar innervated muscles. There were six patients who had undergone ulnar nerve transposition prior to electrodiagnostic testing.

**Electrodiagnostic Protocol**

Although these data were gathered retrospectively, the electrodiagnostic laboratory follows the below protocols for obtaining the NCS discussed.

The ulnar motor to ADM NCS was completed using disk electrodes. The active electrode was placed over the belly of the ADM, the reference placed over the fifth metacarpophalangeal joint, and ground placed on the dorsal aspect of the hand. Stimulation sites were marked with a pen, 8 cm proximally from the active electrode (wrist), 5-cm proximal to the midpoint of the olecranon and medial epicondyle (above elbow), and 5-cm distal to the midpoint of the olecranon and medial epicondyle (below elbow). All stimulation points were measured along the course of the ulnar nerve. Distances between stimulation sites were measured to accurately assess nerve conduction velocity (NCV). The elbow was maintained in 90 degrees of flexion with across elbow distance of 10 cm as described in the multiorganizational summary statement.10

When performing ulnar motor to FDI, disk electrodes were again utilized, with the active electrode placed along the belly of the FDI muscle at approximately the midpoint between the first and second metacarpals, the reference electrode placed at the first metacarpophalangeal joint, and the ground on the dorsal aspect of the hand. The stimulator was then placed at the same stimulation sites as previously described for ADM with NCV again calculated. This technique was consistent with that described previously in the literature.12

To complete the mixed NCS across the elbow, the technique described by Heise and Toledo was used. The bar electrode was applied with the active electrode placed 10-cm proximal to the medial epicondyle between the bellies of biceps and triceps muscles and the ground placed in the mid forearm2 (Fig. 1). Both median and ulnar nerves were respectively stimulated at the wrist with the elbow extended. The difference between the peak distal latencies for the ulnar and median nerves created the ulnar-median mixed latency difference (U-MLD). This was the number employed to examine for correlation between the NCV across the elbow with the motor NCS to ADM and FDI.

**Results**

Of the 135 patient charts, the age range was 18 to 84 years with a mean age of 54 ± 14 years. The sex distribution slightly favored females to males with a ratio of 3:2. Two patients had absent responses to ADM and FDI above the elbow and were subsequently removed for statistical analysis. Of the remaining 133 patients, only 121 of the patients had present responses for the ulnar mixed NCS across the elbow. One patient had absent responses at both ulnar and median mixed NCS across the elbow; this patient was ultimately diagnosed with a length-dependent axonal sensorimotor peripheral polyneuropathy.

When assessing for correlation between the two NCV for ADM and FDI, Pearson correlation revealed an $r = 0.739$, $p < 0.0001$ (Fig. 2). There were two different sets of Pearson correlations run for the U-MLD. One comparing U-MLD to across elbow NCV for both ADM and FDI excluding those patients with absent responses for the mixed NCS and the
other set that substitutes the largest U-MLD for the absent ulnar mixed responses. Excluding the absent responses, the Pearson correlation between U-MLD to ADM and FDI NCV across the elbow was $r = -0.479$ (<Fig. 3>) and $r = -0.543$ (<Fig. 4>), respectively ($p < 0.00001$).

When the absent responses are replaced with the largest U-MLD 2.6 ms, the adjusted $r = -0.60$, $p < 0.00001$ compared with ADM and adjusted $r = -0.642$ to FDI, $p < 0.00001$. In both instances, the negative r-value indicates the inverse relationship between ulnar velocity across the elbow and U-MLD. That is, the larger U-MLD is associated with a slower ulnar NCV across the elbow.

Attempting to diagnose UNE using NCV across the elbow, leaves the clinician to classify the patient into three groups: clearly normal NCV, clearly abnormal NCV, and borderline NCV. The AANEM (American Association of Neuromuscular and Electrodiagnostic Medicine) criteria of $>50$ m/s is normal and updated criteria suggests 43 m/s and slower is clearly abnormal.$^{10,13}$ Therefore, we considered NCV between 44 and 50 m/s to be borderline. This borderline or indeterminate group comprised 36% of the ulnar nerves assessed in this retrospective study. Of this indeterminate group, 64.5% had an abnormal U-MLD of 0.9 milliseconds or more, only two of which were absent responses (6%). This criterion of 0.9 ms (used by this laboratory) is the midpoint between the less stringent 0.69 ms proposed most recently by Di Virgilio et al$^{14}$ and the strict 1.1 ms suggested by Heise and Toledo.$^{2}$ We do not attempt to know exactly what number is “abnormal” or truly clinically significant.

**Discussion**

This study confirms our hypothesis that there are strong correlations between the three tests for ulnar nerve slowing across the elbow in patients with suspected UNE. The traditional motor studies to ADM and FDI have a stronger correlation. Calculating a $R^2$ of approximately 0.54 suggests that these two motor tests agree (or disagree) approximately 50% of the time. When including the absent ulnar mixed nerve responses, the respective $R^2$ values between ADM and
FDI to U-MLD are 0.36 and 0.41. This suggests that these studies disagree or measure differences approximately 59 to 64% of the time. Although these tests are strongly correlated, they are not identical. This difference provides evidence to use an additional test that measures different aspects of the ulnar nerve. We purposefully did not attempt to calculate sensitivity, specificity, or determine which test was more accurate. There was no reliable way to define a “Gold Standard” or “reference standard” (clinical, electrodiagnosti-
elbow. Updated criteria indicates that <43 m/s is abnormal for across elbow NCV. By definition, that designates NCV between 44 and 49 m/s are abnormal according to the older criteria but normal by the new criteria. This predicament leaves many patients with indeterminate criteria for diagnosing UNE. In such cases where the data remain inconclusive, NCS to FDI, ulnar inching, and/or needle examination of FDI, and ulnar innervated forearm muscles are recommended. Despite these guidelines, diagnosing ulnar neuropathy remained more challenging than other peripheral nerves. The difficulty may at least in part be explained in that NCV slowing >10 m/s across the elbow can be as common as 32 to 36% in asymptomatic individuals. Needle electromyography does not always clarify this because forearm flexors have been demonstrated to be spared despite clear evidence of UNE. Others have warned of dangers diagnosing nerve disease using only a single test. Balancing sensitivity and specificity remains problematic. A Danish task force created a novel criterion for diagnosing UNE, hoping to improve accuracy. Scores were assigned based on degree of NCV across elbow and forearm, presence or absence of a difference exceeding 20 m/s for across elbow NCV compared with forearm, and presence or absence of conduction block across the elbow. To improve clinical certainty, additional ulnar motor conduction studies were recommended including adductor pollicis, FDI, and sensory near nerve needle technique. Neither the Danish task force nor the AANEM appear to have considered using the mixed across the elbow NCS to enhance diagnostic accuracy. Given that patients with mainly sensory symptoms are more difficult to diagnose with electrodiagnostic testing; further tests to evaluate sensory and motor function should be useful.

Robinson et al described the value of a summary index when diagnosing carpal tunnel syndrome and not relying on a solitary abnormal test. Our study suggests the U-MLD is frequently abnormal in patients with UNE and likely contributes additional information to the electrodiagnostic evaluation of UNE. The potential to identify UNE early in the disease process, prior to more significant motor involvement has the capability to improve surgical outcomes.

Few studies assessing the sensory component of nerve function across the elbow have been completed; those that have, illustrate wide variations in amplitudes of controls, which was thought to limit their utility. Merlevede et al was the first to describe the mixed CNAP across the elbow to further evaluate UNE. However, two major flaws in this research arose; a lack of power with only ten symptomatic patients and continuing to require elbow flexion as part of the study protocol. Elbow flexion continues to contribute to measurement error. Heise and Toledo later adapted this technique to again measure the mixed CNAP but with the elbow in full extension, forearm supinated, and a sample size of 100 symptomatic patients. In this study, four patients with normal ulnar motor NCS had abnormal CNAP latency difference and motor inching studies, suggesting that ulnar motor studies in isolation are missing the diagnosis of UNE in some patients that could be identified using alternative techniques.

In our laboratory, we have incorporated the U-MLD into the routine evaluation of UNE. The U-MLD is relatively easy and quick to obtain, especially in thin patients. This test allows for direct comparison to the median nerve, in the same person, at the same time, and limits some intersubject variability (temperature, genetic physiological differences, and arm length). Anecdotally, we have noted that the ulnar response is frequently absent in patients with more severe slowing across the elbow, axon, loss, or significant conduction block. At this point, there is no clear evidence to suggest one test is superior to the others.
This current study has several limitations. The retrospective nature of analysis causes multiple limitations. These include:

- There was likely powerful selection bias of patients that had all three NCS performed. Many patients that were clearly normal or clearly indicative of UNE on motor NCS were not included because the U-MLD was not completed. The U-MLD was most likely recorded when the diagnosis was in doubt and/or it was “convenient.”
- There is no way to reliably calculate sensitivity, specificity, or determine which of the tests had the best accuracy. We intentionally did not attempt to define a “Gold Standard” or “reference standard” for the diagnosis of UNE.
- Past medical history including diabetes or other medical problems, was not collected or considered as a part of exclusion criteria. Without further research, it is unclear how these disease processes would affect the U-MLD.
- Although all data were collected from the same electrodiagnostic laboratory, not all NCS were completed by the same physician. This has both advantages and disadvantages regarding generalizability. To date, there has been no prospective study examining interrater reliability with the mixed across the elbow NCS.

Although these limitations exist, this research solidifies a strong relationship between the mixed across the elbow (U-MLD) and UNE. The negative r-value describes the inverse relationship between ulnar NCV across the elbow and U-MLD, that is, the slower the velocity across the elbow the larger the U-MLD. Mixed across the elbow has moderate to strong correlation with both ADM and FDI NCV. This gives construct validity to the U-MLD. All three tests measure ulnar nerve function slightly differently and without further prospective data, it is unclear which single test is best, creating a state of clinical equipoise. The authors propose that some combinations of ulnar motor to ADM, ulnar motor to FDI, and mixed across the elbow tests may be more accurate than relying on motor studies alone.

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Conflict of Interest
None declared.