

## Towards viscoelastic characterisation of the human ulnar nerve

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1 **Towards Viscoelastic Characterisation of the Human Ulnar Nerve: an early assessment**  
2 **using embalmed cadavers**

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20

21

22 **Abstract**

23 Cubital tunnel syndrome is the most prevalent neuropathy of the ulnar nerve and its aetiology  
24 is controversial. Potential replacement materials should display similar viscoelastic  
25 properties. The purpose of this study was to assess the feasibility and merit of quantifying the  
26 frequency-dependent viscoelastic properties of proximal and distal sections of the human  
27 ulnar nerve. Four ulnar nerves ( $n = 4$ ) were dissected from the elbows of human cadavers and  
28 sectioned at the level of the cubital tunnel into proximal and distal sections. These eight  
29 sections of the ulnar nerve were sinusoidally loaded to induce stresses between 0.05 - 0.27  
30 MPa and the viscoelastic properties were measured between 0.5 - 24 Hz using Dynamic  
31 Mechanical Analysis. The nerves were found to exhibit frequency-dependent viscoelastic  
32 behaviour throughout this frequency range. The median storage moduli of the proximal  
33 nerves ranged between 7.03 and 8.18 MPa, and 8.85 to 10.19 MPa for distal nerves, over the  
34 frequency-sweep tested. The median loss moduli of the proximal nerves ranged between 0.46  
35 and 0.81 MPa and between 0.51 - 0.80 MPa for distal nerves. Ulnar nerves display frequency  
36 dependency viscoelasticity. Such characterisation is feasible with potential applications to  
37 suitable nerve grafts.

38 **Keywords:** Dynamic Mechanical Analysis; Frequency; Human; Ulnar nerve; Viscoelasticity.

39

## 40 **1. Introduction**

41 The ulnar nerve travels through the upper limb and cubital tunnel transmitting sensation from  
42 the skin overlying the hypothenar eminence, the corresponding area of skin posteriorly, the  
43 little finger and half of the ring finger as well as supplying motor function to numerous  
44 muscles of the forearm and hand [1]. Cubital tunnel syndrome is the most prevalent  
45 neuropathy of the ulnar nerve and the second commonest neuropathy of the upper limb [2].  
46 Its aetiology is controversial. Originally, it was thought to be due to a compressive or  
47 entrapment neuropathy [3–5]. However, more recently, it has been thought to be due to nerve  
48 strain [2,6–9].

49 Studies have found that at certain levels of strain (6-16%), blood flow to the nerve and  
50 conduction of impulses by the nerve were reduced or even arrested [10–13]. In terms of nerve  
51 conduction, it has been shown that a 6% increased nerve strain for longer than an hour led to  
52 70% decreased conduction velocity while a 12% increase in strain led to completely arrested  
53 nerve conduction in a study on rabbit nerves [13]. The nerve conduction returned once the  
54 above strains were removed [13]. In terms of blood flow, a 50% reduction was induced by  
55 8% strain in a rat's sciatic nerve while an 80% reduction in blood flow was caused by 15%  
56 nerve strain [12]. Blood flow was completely blocked by 16% strain in a rabbit sciatic nerve  
57 [11]. Therefore, for adequate nerve function, nerve strain must be minimised. It has  
58 previously been shown that during normal motion of the elbow and shoulder joints, strain is  
59 applied dynamically to the ulnar nerve to levels that could result in both impaired conduction  
60 and perfusion [8,14,15].

61 Human peripheral nerves are known to exhibit viscoelastic properties [16] and this has been  
62 demonstrated for the human ulnar nerve by performing *in vitro* stress relaxation tests [17].

63 Unlike creep and stress relaxation, Dynamic Mechanical Analysis (DMA) is a dynamic

64 testing method used to determine the viscoelastic properties of a material or multi-component  
65 structure [18]. DMA involves the application of an oscillating force to a specimen and  
66 measuring the out-of-phase displacement [19]. This gives time-dependent strain,  $\varepsilon(t)$   
67 (equation 1), developed in response to the induced time-dependent stress,  $\sigma(t)$ , and the  
68 complex (dynamic) modulus,  $E^*(\omega)$  [20]:

$$69 \quad \varepsilon(t) = \frac{\sigma(t)}{E^*(\omega)} \quad (1)$$

70 The viscoelasticity of a material can be characterised in terms of storage and loss moduli [20–  
71 22]. The storage modulus ( $E'$ ) characterises the ability of the material to store energy that is  
72 then available for elastic recoil; while, the loss modulus ( $E''$ ) characterises the material's  
73 ability to dissipate energy. The storage and loss moduli are related to  $E^*$  and the phase angle  
74 ( $\delta$ ) by equation 2 and 3, respectively [20,22,23]:

$$75 \quad |E^*| = \sqrt{E'^2 + E''^2} \quad (2)$$

$$76 \quad \delta = \tan^{-1}\left(\frac{E''}{E'}\right) \quad (3)$$

77 To the authors' knowledge, the understanding of frequency-dependent viscoelastic properties  
78 of human ulnar nerve is currently absent. As the ulnar nerve is viscoelastic, and exposed to  
79 dynamic loading, its frequency-dependency requires characterisation. Furthermore, any  
80 potential replacement materials (allograft, synthetic grafts, etc.) should display similar  
81 viscoelastic properties. Moreover, frequency-dependent viscoelastic properties are important  
82 because if these measurements are used to infer the *in vivo* strain, then the strain itself would  
83 be highly sensitive to the rate of loading: of importance given the dynamic loading to which  
84 the ulnar nerve is exposed *in vivo*. Additionally, mechanical behaviour of viscoelastic  
85 biomaterials may differ considerably between physiological and sub-physiological loading  
86 rates [24].

87 The aim of this study was to assess the feasibility and merit of quantifying the frequency-  
88 dependent viscoelastic properties of proximal and distal sections of the human ulnar nerve.  
89 Furthermore, this study subsequently compared the ulnar nerve frequency-dependency  
90 viscoelastic properties of storage and loss moduli proximally and distally to the cubital  
91 tunnel. Given the limited availability of fresh human ulnar nerves for mechanical testing,  
92 embalmed human nerves have been used.

93

## 94 **2. Materials and Methods**

### 95 *2.1 Cadaver Information and Ulnar Nerve Specimen Preparation*

96 Four ulnar nerves were dissected and surgically removed from four elbows of three whole,  
97 intact embalmed cadavers (Table 1). Ethical approval was obtained from the Human Tissue  
98 Authority according to the Human Tissue Act (2004) under the University of Birmingham  
99 license (number 12236) with the donors consenting to the use of their cadavers for education  
100 and research. All tissues were obtained following the Declaration of Helsinki ethical  
101 principles.

102 The elbows were first marked and incised to expose the nerves. Sutures were then placed at  
103 approximately 20 mm or 30 mm (due to anatomical positioning). Biomechanical tests  
104 consisting of flexion and extension of the elbow at varying degrees of shoulder abduction  
105 were performed as part of a separate study [15]. The nerves were removed from the cadaver  
106 then wrapped and soaked in a damping down solution containing H<sub>2</sub>O, Poly(ethylene glycol)  
107 8000, biocleanse (Fisher Scientific, Loughborough, UK) and Industrial Methylated Spirits  
108 (IMS) (VWR International Ltd, Leighton Buzzard, UK). Next, the nerves were double  
109 bagged as whole nerves. Each nerve was approximately 20-30 cm in length. The nerves were  
110 then sectioned (Figure 1), at the level of the cubital tunnel into proximal and distal sections.

111 Three nerves were divided into 40 mm sections, (approximately 20 mm of a gauge and two  
112 10 mm shoulder sections used to grip the nerve for mechanical testing) and one nerve was  
113 divided into 50 mm sections, (approximately 30 mm of a gauge and two 10 mm shoulder  
114 sections). The difference in length was to maintain consistent suture positioning from a  
115 previous study [15]. Specimens were hydrated with the aforementioned damping down  
116 solution. Branches were removed with the nerves. The nerves were then mechanically tested  
117 the following day at room temperature.

118

## 119 *2.2 Preliminary tests*

120 BOSE Electroforce DMA Grips (Bose Corporation, ElectroForce Systems Group, Minnesota,  
121 USA), were used to grip 10 mm on either side of the nerve. Preliminary ramp tests were  
122 conducted on two specimens from one cadaver (taken 10 cm proximal to and 10 cm distal to  
123 the cubital tunnel) of approximately 20 mm of a gauge of proximal and distal sections of all  
124 nerves. These samples were extended at a linear translational rate of 0.05 mm/s in accordance  
125 with a previous study [17] to characterise the quasi-static stress-strain curves of the human  
126 nerves (ulnar proximal and distal). Tensile tests were performed at an initial ramp up strain of  
127 10% [17]. A Vernier calliper was used to measure height and diameter of each nerve  
128 specimen. As the nerves were approximately elliptical in cross-sectional area, three sagittal  
129 (*a*) and three coronal (*b*) radii were measured and averaged, respectively, to calculate the  
130 elliptical cross-sectional area ( $A_e$ ) using equation 4 [17].

131

$$A_e = \pi ab \quad (4)$$

132 Force versus displacement of proximal and distal nerves showed differences in stiffness  
133 (gradient of the line in N/mm) between the two nerve specimens (see Figure 2). When  
134 comparing a linear region (often termed post-transitional), but avoiding any potential end-

135 stage plastic deformation, the proximal human nerve was stiffer than the distal nerve (see  
136 Figure 2). Calculating the stiffness of each nerve (as the force/extension within this linear  
137 range) led to values of 15.00 N/mm for the proximal human nerve and 8.07 N/mm for the  
138 distal nerve. Therefore, the DMA protocol devised included comparison of proximal and  
139 distal samples (Section 2.3).

140 Figures 3a and 3b show stress versus strain of the proximal and distal human nerves. For the  
141 proximal nerve, 2% (0.02) strain was equivalent to 0.04 MPa stress while 6% (0.06) strain  
142 was equivalent to 0.15 MPa of stress (see Figure 3a). However, 2% (0.02) strain, of the distal  
143 nerve, was equivalent to 0.05 MPa while 6% (0.06) strain was equivalent to 0.27 MPa of  
144 stress (see Figure 3b).

145 At approximately 7-8% strain, the distal nerve began to demonstrate signs of damage, as  
146 evidenced by a plateau of the induced stress (see Figure 3b), and may be associated with  
147 plastic deformation of the nerve and/or rupture. This plateau could mean that the  
148 microstructure of the nerve is rupturing. Therefore, the distal nerve's values of stress and  
149 strain were chosen to guide the DMA testing to avoid rupture in the actual experiment.

150

### 151 *2.3: Dynamic Mechanical Analysis (DMA)*

152 The viscoelastic properties of the nerve sections were characterised using a Bose  
153 ElectroForce 3200 testing machine running Bose WinTest 4.1 DMA software (Bose  
154 Corporation, ElectroForce Systems Group, Minnesota, USA). DMA has previously been used  
155 to quantify the storage and loss properties of a variety of biological tissues [22,25–28] and  
156 orthopaedic implants [18,29].

157 For DMA, each nerve was sinusoidally loaded to induce stresses between 0.05 MPa  
158 (equivalent to 2% strain of the distal nerve stress-strain curve; Figure 3) and 0.27 MPa. 2%



159 strain was chosen as the lower strain boundary to mimic the nerve *in vivo* conditions [30–32].  
160 As the elliptical area of the nerve varied, the applied force was calculated for each individual  
161 nerve specimen and the individual force ranges were applied to the individual specimens.  
162 Thus, the induced sinusoidal stress was consistent for all samples, varying from a trough of  
163 0.05 MPa to a peak of 0.27 MPa. Preliminary data (section 2.2), of the distal nerve (Figure 3),  
164 demonstrated that 6% strain was equivalent to 0.27 MPa of stress (see equation 5 where  $\sigma$  is  
165 stress,  $F$  is the applied force and  $A_e$  is the area of an ellipse).

$$166 \quad F = \sigma A_e \quad (5)$$

167 A preload condition, at 1 Hz for 28 cycles, was applied before the frequency sweep to ensure  
168 no stress relaxation affected the frequency sweep. Next, the storage ( $E'$ ) and loss ( $E''$ ) moduli  
169 were evaluated for 9 frequencies (0.5, 1, 1.5, 2, 5, 10, 15, 20 and 24 Hz).  $E'$  and  $E''$  were  
170 calculated using the WinTest DMA software. Following the application of the oscillating  
171 force, the out-of-phase displacement response is measured [19]. By performing a Fast Fourier  
172 Transform (FFT) of the sinusoidal load ( $F$ ) and displacement ( $d$ ) for each frequency, the  
173 magnitudes of the force ( $F^*$ ), magnitude of the displacement ( $d^*$ ), the phase lag ( $\delta$ ) and  
174 frequency ( $f$ ) were quantified [18].  $F^*$  and  $d^*$  were used to calculate the dynamic stiffness  
175 ( $k^*$ ) using equation 6.

$$176 \quad k^* = \frac{F^*}{d^*} \quad (6)$$

177 As the nerves were elliptical, a shape factor,  $S_c$  (equation 7), was used to calculate  $E'$  and  $E''$   
178 of the nerves using equations 8 and 9, respectively. Equation 7 uses a standard shape for a  
179 cylindrical sample [22,23], modified from a circular to an elliptical cross-section (see  
180 equation 4);  $h$  refers to the gauge length ('height') of the specimen. The procedure used for  
181 measuring the preliminary specimens, which is described above (Section 2.2), was used to  
182 measure the specimens tested with DMA. The test gauge length of the specimens was  $19.71 \pm$

183 1.26 mm with the exception of BM 172-14 in which a gauge length of  $27.83 \pm 2.61$  mm was  
184 used as sutures were placed differently due to anatomical positioning.

$$185 \quad S_c = \frac{\pi}{h}(ab) \quad (7)$$

$$186 \quad E' = \frac{k^* \cos \delta}{S_c} \quad (8)$$

$$187 \quad E'' = \frac{k^* \sin \delta}{S_c} \quad (9)$$

#### 188 2.4 Data analysis

189 All statistical analyses were performed using SigmaPlot 13.0 (SYSTAT, San Jose, CA,  
190 USA). To evaluate the frequency-dependent viscoelastic behaviour of the nerves, regression  
191 analysis, was performed for  $E'$  and  $E''$ . A logarithmic fit (equations 10 and 11) was found to  
192 best fit the data, and was evaluated in terms of the significance of the curve fit ( $p < 0.05$ ) and  
193 goodness of fit ( $R^2$ ).

$$194 \quad E' = A \ln(f) + B \quad (10)$$

$$195 \quad E'' = C \ln(f) + D \quad (11)$$

196 The 95% confidence intervals were calculated for proximal sections ( $n = 4$ ) and distal  
197 sections ( $n = 4$ ). For comparisons of all nerves, confidence intervals error bars were  
198 calculated with a sample size of 8 ( $n = 8$ ). A Wilcoxon ranked sum test was performed to  
199 evaluate the significant difference of the  $E'$ , of the proximal and distal nerves for each  
200 frequency tested. This test was also performed to compare  $E''$  of the proximal and distal  
201 nerves at each frequency tested. All statistical results with  $p < 0.05$  were considered  
202 significant.

203

204 **3. Results**

205 The nerves displayed viscoelastic behaviour throughout the tested frequency range. Figure 4  
206 shows the frequency dependent trend of the  $E'$  of the proximal and distal sections of ulnar  
207 nerves. The median  $E'$  of the proximal nerves ranged between 7.03 and 8.18 MPa for the  
208 different frequencies tested. This compared to the range of the distal nerves' median  $E'$  which  
209 was between 8.85 and 10.19 MPa for the same frequency range. The frequency-dependency  
210 of the  $E'$  (equation 10) was determined empirical to follow a logarithmic fit ( $p < 0.05$ ). No  
211 significant difference was observed for  $E'$  between the proximal and distal sections across all  
212 frequencies tested ( $p > 0.05$ ).

213 Figure 4b shows the frequency dependent trend of the  $E''$  of the proximal and distal sections  
214 of ulnar nerves. The  $E''$  was lower than the  $E'$  for both proximal and distal sections of nerves  
215 at all tested frequencies. Over the same frequency range tested, the median value for  $E''$  of  
216 the proximal nerve specimens ranged between 0.46 and 0.81 MPa while the range of median  
217 for the distal nerves was 0.51 and 0.80 MPa. No significant difference was observed between  
218 proximal and distal sections for  $E''$  ( $p > 0.05$ ). With the exception of the  $E''$  for proximal  
219 BM 172-14, the frequency-dependency of the  $E''$  (equation 11) was empirically described by  
220 a logarithmic fit (Table 2). Individual fits for  $E'$  and  $E''$  have been provided as  
221 supplementary data.

222 Figure 5 shows the frequency dependent trend of the  $E'$  of all proximal and distal sections of  
223 the ulnar nerves combined. The confidence interval error bars approximately halve between  
224  $E'$  and  $E''$  of proximal and distal nerves and  $E'$  and  $E''$  of all nerves due to doubling of the  
225 sample size. Figure 5b shows the frequency dependent trend of the  $E''$  of all proximal and  
226 distal sections of the ulnar nerves combined. The  $E''$  was less than the  $E'$  for all sections of  
227 the nerves combined at all tested frequencies.

228

#### 229 **4. Discussion**

230 This study has, for the first-time, demonstrated that human ulnar nerves display frequency-  
231 dependent viscoelastic properties. Embalmed nerves have been used to demonstrate the  
232 feasibility of characterising their viscoelastic properties throughout a physiologically relevant  
233 frequency range. Except for BM 172-14  $E''$ , all nerves  $E'$  and  $E''$  followed an empirical  
234 logarithmic frequency-dependent trend. Preliminary data, of the distal nerve, demonstrated  
235 that 6% strain was equivalent to 0.27 MPa of stress. This induced stress was selected as the  
236 maximum induced stress for dynamic mechanical analysis to ensure no rupture occurred  
237 under dynamic loading. The median storage moduli of the proximal nerves ranged between  
238 7.03 and 8.18 MPa for the different frequencies tested. This compared to the range of the  
239 distal nerves' median storage modulus which was between 8.85 and 10.19 MPa for the same  
240 frequency range. Over the same frequency range, the median loss moduli of the proximal  
241 nerves ranged between 0.46 and 0.81 MPa while the range of the distal nerves' median loss  
242 modulus was 0.51 and 0.80 MPa. In this preliminary study, no significant differences in  
243 viscoelasticity were identified between proximal and distal samples, however, this finding  
244 would require confirmation with a larger data set. A larger data set would also allow  
245 meaningful comparisons to assess of any gender differences in nerve viscoelasticity.

246 No consensus exists regarding the critical limit of elongation with various studies ranging  
247 from 6% to 100% [16]. From the preliminary test of the distal nerve, the nerve began to  
248 rupture at approximately 7-8% strain; this can be seen by a plateau of the induced stress with  
249 increased strain. This maximum stress (0.27 MPa) at 6% strain was used to ensure no  
250 rupturing occurred during DMA while the stress at 2% strain (0.05 MPa) was used to ensure  
251 the nerve specimens were always under tension. A comparison was undertaken to investigate

252 whether the strain measured, from the preliminary ramp test, was comparable with the  
253 dynamic “estimated” strain measured by using the complex modulus and induced peak and  
254 trough stresses (Equation 1; see Table 3).

255 The estimated strain at 0.05 MPa ranged from  $0.65 \pm 0.18\%$  (0.5 Hz) to  $0.56 \pm 0.16\%$  (24 Hz)  
256 while at 0.27 MPa the estimated strain ranged from  $3.49 \pm 0.99\%$  (0.5 Hz) to  $3.01 \pm 0.85\%$   
257 (24 Hz). This estimated strain is different to the preliminary strain (2%, for 0.05 MPa, and  
258 6% for 0.27 MPa). This variation may be due to differences in testing procedure (quasi-static  
259 versus dynamic) or may also be due to the linearity assumption of using the complex  
260 modulus for the estimated strain [20]. In relation to *in situ* strain of human cadavers,  
261 numerous studies have quantified a wide range of strains; 0-17% [15], 0-14% [7], 29% [8], 9-  
262 69% [33]. The values estimated in this present study are within these ranges; thus, the  
263 viscoelastic measurements provided are within a range which corresponds to existing  
264 measures of strain.

265 To the authors’ knowledge, no other studies have investigated the viscoelastic properties  
266 (storage modulus and loss modulus) of the ulnar nerve through DMA. Therefore, there is no  
267 other literature with which to compare the current results directly. Ma et al. [17] investigated  
268 *in vitro* mechanical properties (tensile ramp and stress relaxation tests) of cadaveric nerves as  
269 well as measuring *in vivo* stress and deformation intraoperatively. At the same strain, the  
270 authors found that the *in vivo* induced stress was over seven times higher than the measured  
271 induced stress from the *in vitro* tests [17]. This highlights the different biomechanical  
272 properties of a nerve *in situ*, when it is surrounded by connective tissue and still has branches  
273 and blood vessels attached, to when it is removed from the body. Further, at 10% strain, Ma  
274 et al. [17] calculated that the *in vitro* induced stress, of the ulnar nerve, was approximately  
275 0.18-0.19 MPa while the present study calculated an induced stress of 0.37 MPa (distal) and  
276 0.43 MPa (proximal); approximately 2.0-2.4 times greater. This difference may be due to

277 multiple factors which includes the variability of human tissues, the inconsistency across the  
278 testing methodologies and storage/preservation techniques (fresh-frozen [17] versus  
279 embalmed (present study).

280 A potential limitation of the present study is the use of embalmed nerves instead of fresh  
281 nerves. Embalmed cadavers were the only type available to use at the time of testing. It is  
282 unethical and, therefore, impossible to obtain live human nerves for *in vitro* mechanical  
283 testing. Thus, all intact nerves would have had some form of treatment. However, while there  
284 is a difference in absolute values between *in situ* biomechanical properties of unembalmed  
285 and embalmed ulnar nerves, a correlation in strain values has been previously demonstrated  
286 [34]. Another limitation of this study is that only 4 cadavers were available at the time of  
287 testing which likely explains the variability seen in the results of this study. This sample size  
288 might preclude generalizability. In this study, all samples were obtained from only 4 nerves;  
289 thus, a large difference in means would be necessary, and minimal standard deviation, to  
290 detect a difference with significance ( $p < 0.05$ ) when comparing proximal and distal samples.  
291 However, our results are consistent with literature where appropriate, and furthermore, clear  
292 and consistent trends were obtained.

293 In this current study, frequency-dependent viscoelasticity has been assessed over a range of  
294 0.5-24 Hz. While much of this range of frequencies may not appear physiological,  
295 characterisation of natural tissues should consider not only physiological rates of loading, but  
296 also loading associated with exercise, other daily activities, pathophysiology and/or trauma  
297 [23,24,35]. However, loading rates and equivalent frequencies associated with loading of the  
298 upper-limb/elbow, and of potential relevance to the ulnar nerve are less well understood than,  
299 say, for natural tissues such as for heart valves [35-37] or lower limbs [23,38,39]. However,  
300 there are upper-limb studies which suggest that frequencies of 20 repeats/min (0.33 Hz) are  
301 associated with discomfort levels within a physiological loading range [40], providing a

302 lower range for an experimental loading frequency. Whereas, hand-transmitted vibration for  
303 steering wheels have been calculated as having a weighting factor (from an ergonomic  
304 perspective) which is greatest between 6-25 Hz [41]; peaking at 12.5 Hz. The range of  
305 loading frequencies identified from the above studies (0.33 – 25 Hz) is consistent with the  
306 range assessed in our study (0.5 – 24 Hz). However, it is recognised there may be conditions  
307 which might expose the nerve to higher loading frequencies not assessed in our study, e.g.  
308 300 Hz [42]. Furthermore, the frequencies used to guide this current study are estimates, as  
309 the strain rate of the ulnar nerve itself associated with loading *in vivo* is not currently known.  
310 Thus, it is the trend across a range of frequencies (0.5 – 24 Hz) which is viewed as important  
311 in our current study, indicating a frequency range for future studies.

312 Repeatable characterisation of samples with DMA requires a dynamic “steady-state” [38] to  
313 be reached using preconditioning loading cycles. For some natural soft tissues (e.g. articular  
314 cartilage) there is evidence that this can require in excess of 1000 loading cycles [43].  
315 However, a minimal number of preconditioning cycles is recommended to avoid the risk of  
316 fatigue. In our current study, 28 preconditioning loading cycles were found to enable  
317 repeatable viscoelastic characterisation with DMA. Therefore, while 28 cycles may appear  
318 high as compared to quasi-static material’s characterisation studies (typically employing less  
319 than 10 preconditioning loading cycles), it is low as compared to preconditioning used for  
320 DMA of natural soft tissues.

321 Nerves are non-homogenous in nature and structure varies throughout and between individual  
322 nerves [16], so the conclusions from this study should be extrapolated only with caution to  
323 other nerves, as the measurements may be specific to the ulnar nerve in the region of the  
324 cubital tunnel. However, determining the viscoelastic properties of nerves is crucial for  
325 choosing suitable nerve grafts, either in manufacturing synthetic grafts or in checking the  
326 suitability of allografts. Knowledge of viscoelastic properties is also important in designing

327 and manufacturing diagnostic, surgical and surgical training devices as well as for making  
328 computational models for research [25] and for the multi-physics modelling of nerves.  
329 Furthermore, a deeper understanding of the mechanical properties of peripheral nerves allows  
330 a greater appreciation of mechanisms of nerve injury and repair. It is hoped that such  
331 knowledge and equipment will lead to better patient outcomes.

332

## 333 **5. Conclusion**

334 The human ulnar nerves display frequency-dependency viscoelasticity. Both the median  
335 storage and loss moduli increased logarithmically as the frequency increased, with the storage  
336 modulus consistently greater than the loss modulus. Such characterisation is feasible with  
337 potential applications to suitable nerve grafts.

338

## 339 **DECLARATIONS**

### 340 **Ethics approval and consent to participate**

341 Ethical approval was obtained from the Human Tissue Authority according to the Human  
342 Tissue Act (2004) under the University of Birmingham license (number 12236) with the  
343 donors consenting to the use of their cadavers for education and research. All tissues were  
344 obtained following the Declaration of Helsinki ethical principles.

345

### 346 **Consent for publication**

347 All donors consented to the use of their cadavers for education and research. This study  
348 reports age and gender of donors only.



349

350 **Competing interests and/or conflicts of interest**

351 None declared.

352

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499

500 FIGURE CAPTIONS

501 **Figure 1:** BM 171-14 left ulnar nerve with (a) Five sutures marked in red numbers. b) Left  
502 ulnar nerve with black arrow marking where it was sectioned at the cubital tunnel. c) Left  
503 ulnar nerve proximal (left) and distal (right) sections. One section had 30 mm of a gauge with  
504 10 mm for gripping at either end. d) Final nerve sections for testing (lengths are  
505 approximate).

506

507 **Figure 2:** Force (N) versus displacement (mm) of proximal and distal human nerves.

508

509 **Figure 3:** Stress versus strain of proximal (a) and distal (b) sections of the human ulnar  
510 nerve. Stress is measured in MPa while strain is dimensionless. Red lines show 2% and 6%  
511 (0.02 and 0.06) strain which corresponds to 0.05 and 0.27 MPa stress.

512

513 **Figure 4:** The proximal and distal ulnar nerve frequency dependent (a) storage modulus ( $E'$ )  
514 ( $\text{N}/\text{mm}^2$ ) and (b) loss modulus ( $E''$ ) ( $\text{N}/\text{mm}^2$ ) (median  $\pm$  95% confidence intervals).

515

516 **Figure 5:** The ulnar nerve (combined proximal and distal sections) frequency dependent (a)  
517 storage modulus ( $E'$ ) ( $\text{N}/\text{mm}^2$ ) and (b) loss modulus ( $E''$ ) ( $\text{N}/\text{mm}^2$ ) (median  $\pm$  95%  
518 confidence intervals).

519



520 TABLES

521 **Table 1.** Ulnar nerve specimens.

<b>Cadaver ID</b>	<b>Donor Age</b>	<b>Gender</b>	<b>Side</b>
<b>Cadaver 1</b>	90	Male	Right
<b>Cadaver 1</b>	90	Male	Left
<b>Cadaver 2</b>	89	Male	Left
<b>Cadaver 3</b>	75	Female	Left

522

523 **Table 2.** Logarithmic regression of storage modulus (E') and loss modulus (E'') for proximal  
 524 and distal sections of nerves. The units of coefficients (A and C) and constants (B and D) are  
 525 N/mm<sup>2</sup>. Regression with a p < 0.05 were deemed significant.

<b>Specimen ID</b>	<b>A</b>	<b>B</b>	<b>R<sup>2</sup></b>	<b>p value</b>	<b>C</b>	<b>D</b>	<b>R<sup>2</sup></b>	<b>p value</b>
Proximal BM 176-14	0.32	7.80	0.98	<0.001	0.03	0.59	0.69	0.006
Proximal BM 172-14	0.25	6.07	0.65	0.009	0.09	0.42	0.41	0.063
Proximal BM 171-14 Left	0.33	9.99	0.98	<0.001	0.05	0.63	0.60	0.014
Proximal BM 171-14 Right	0.26	6.54	0.96	<0.001	0.03	0.41	0.72	0.004
<b>Median of all proximal</b>	<b>0.29</b>	<b>7.17</b>	<b>0.98</b>	<b>&lt;0.001</b>	<b>0.05</b>	<b>0.51</b>	<b>0.53</b>	<b>0.026</b>
Distal BM 176-14	0.33	7.94	0.97	<0.001	0.03	0.51	0.70	0.005
Distal BM 172-14	0.30	5.42	0.97	<0.001	0.02	0.40	0.64	0.009
Distal BM 171-14 Left	0.41	12.66	0.97	<0.001	0.10	0.73	0.68	0.006
Distal BM 171-14 Right	0.35	10.12	0.98	<0.001	0.06	0.62	0.63	0.010
<b>Median of all distal</b>	<b>0.34</b>	<b>9.03</b>	<b>0.98</b>	<b>&lt;0.001</b>	<b>0.04</b>	<b>0.67</b>	<b>0.67</b>	<b>0.007</b>
<b>Median all proximal and distal</b>	<b>0.33</b>	<b>7.87</b>	<b>0.98</b>	<b>&lt;0.001</b>	<b>0.04</b>	<b>0.54</b>	<b>0.51</b>	<b>0.031</b>

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528

529 **Table 3.** Estimated strain (%) calculated from the complex (dynamic) modulus ( $E^*$ ). The  
 530 estimated strain is calculated at the maximum (0.27 MPa) and minimum (0.05 MPa) induced  
 531 stress (median  $\pm$  standard deviation).

Frequency (Hz)	$E^*$ (MPa)	Strain at 0.05 MPa (%)	Strain at 0.27 MPa (%)
0.5	7.73 $\pm$ 2.39	0.65 $\pm$ 0.18	3.49 $\pm$ 0.99
1	7.89 $\pm$ 2.46	0.63 $\pm$ 0.18	3.42 $\pm$ 0.97
1.5	8.02 $\pm$ 2.48	0.62 $\pm$ 0.18	3.37 $\pm$ 0.96
2	8.09 $\pm$ 2.50	0.62 $\pm$ 0.18	3.34 $\pm$ 0.95
5	8.30 $\pm$ 2.53	0.60 $\pm$ 0.17	3.25 $\pm$ 0.92
10	8.54 $\pm$ 2.54	0.59 $\pm$ 0.16	3.16 $\pm$ 0.85
15	8.78 $\pm$ 2.58	0.57 $\pm$ 0.15	3.08 $\pm$ 0.81
20	8.96 $\pm$ 2.48	0.56 $\pm$ 0.14	3.01 $\pm$ 0.77
24	8.97 $\pm$ 2.70	0.56 $\pm$ 0.16	3.01 $\pm$ 0.85

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