Diffusion Tensor Imaging of the Median Nerve: A Systematic Review and Meta-Analysis of Normal Values in Asymptomatic Adults and How They Change in Carpal Tunnel Syndrome

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Research Article

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Abstract

**Background:** Carpal tunnel syndrome (CTS) leads to distortion of axonal architecture, demyelination and fibrosis within the median nerve. Diffusion tensor imaging (DTI) characterises tissue microstructure and generates reproducible proxy measures of nerve ‘health’ which are sensitive to myelination, axon diameter, fibre density and organisation. This meta-analysis summarises the normal DTI values of the median nerve, and how they change in CTS.

**Methods:** This systematic review included studies reporting DTI of the median nerve at the level of the wrist in adults. The primary outcome was to determine the normal fractional anisotropy (FA) and mean diffusivity (MD) of the median nerve. Secondarily, we show how the FA and MD differ between asymptomatic adults and patients with CTS, and how these differences are independent of the acquisition methods.

**Results:** 32 studies of 2643 wrists, belonging to 1575 asymptomatic adults and 1068 patients with CTS were included. The normal FA was 0·58 (95% CI 0·56, 0·59) and the normal MD was 1·138 x10^{-3} mm^2/s (95% CI 1·101, 1·174). Patients with CTS had a significantly lower FA than controls (mean difference 0·12 [95% CI 0·09, 0·16]). Similarly, the median nerve of patients with CTS had a significantly higher mean diffusivity (mean difference 0·16 mm^2/s x10^{-3} [95% CI 0·05, 0·27]). The differences were independent of experimental factors.

**Conclusion:** We provide summary estimates of the normal FA and MD of the median nerve in asymptomatic adults. Furthermore, we show that diffusion throughout the length of the median nerve becomes more isotropic in patients with CTS.

Introduction

Carpal tunnel syndrome (CTS) is the most common compressive neuropathy, affecting 10 million people annually. Consequently, CTS is the most expensive upper extremity musculoskeletal disorder, costing the USA health system over $2 billion annually and employers up to $114,000 per incident.\(^1\)

Compression of peripheral nerves leads to distortion of the axonal architecture, demyelination with or without poor remyelination, loss of the intrinsic vasculature and ultimately, fibrosis of the perineurial and epineurial connective tissue.\(^2,3\) Diffusion tensor imaging (DTI) characterises tissue microstructure and generates reproducible proxy measures of nerve ‘health’ which are sensitive to myelination, axon diameter, fibre density and organisation\(^4-7\) (Figure 1). DTI typically generates the following metrics: fractional anisotropy (FA), mean diffusivity (MD), axial diffusivity (AD) and radial diffusivity (RD). FA is a scalar value between zero and one – an FA of zero implies isotropic diffusion of water within a voxel, whilst a FA of one implies diffusion along a single axis (i.e., bidirectional diffusion along the length of the nerve). MD describes the average molecular diffusion rate within the voxel, whilst AD describes diffusion in the long axis and RD represents diffusion perpendicular to the long axis.
Several studies have shown that DTI metrics (FA and MD) are sensitive to microstructural changes which occur within the median nerve of patients with CTS (Figure 1). However, there are several uncertainties must be resolved before this technology could be used in clinical practice or as a reference standard in research studies. Firstly, the normal DTI values of the median nerve must be established and secondly, uncertainty around how experimental conditions (e.g. scanning parameters) influence DTI metrics need to be determined. These uncertainties, and how DTI metrics change in CTS, might be resolved through meta-analysis and represents the rationale for this study.

**Methods**

This review is registered with PROSPERO (CRD42020212378). It was designed and conducted in accordance with the Cochrane Handbook of Systematic Reviews and has been authored in accordance with the PRISMA 2020 statement.

**Types of Studies**

We included all studies which reported the findings of diffusion tensor magnetic resonance imaging of the median nerve, at the level of the wrist in asymptomatic adults or adults with CTS. There were no language restrictions. We excluded case reports and studies which did not report DTI metrics (e.g., studies which contained fibre tractography graphics only) of the median nerve.

**Participants**

This review considers 2 distinct populations:

1. Asymptomatic adults (aged ≥ 16 years) with no known pathology (past or present) affecting the peripheral nerves of the upper limb.
2. Adults with a diagnosis of carpal tunnel syndrome. For a study to be included, we did not impose any specific thresholds or criteria on the diagnosis of CTS, such as the presence of specific symptoms, provocative tests, aberrant electrophysiological parameters or imaging features.

**Search strategy**

The NICE Healthcare Databases (hdas.nice.org.uk) was searched according to Appendix 1 (Supplementary Materials) on 9th October 2020. The medRxiv and bioRxiv preprint archives were searched with the same strategy using the R package medrxivr. Later, CitationChaser was used for forward and backward citation chasing based on the final list of included studies (eFigure 1).

**Study selection**

Three review authors (DR, JR and FR) independently screened titles and abstracts for relevance, in accordance with the eligibility criteria. The full texts of potentially eligible articles were obtained and
again independently assessed by the same authors. Disagreements were resolved by discussion with RGW. The reasons for excluding studies are outlined in Appendix 2 (Supplementary Materials).

Data extraction

Three review authors (DR, JR and FR) independently double extracted all data. Thereafter, all datapoints were independently checked for accuracy by RGW. DTI parameters were extracted from the following anatomical levels of the median nerve: the distal radio-ulnar joint (DRUJ), the pisiform and the hook of the hamate, as these are three commonly used imaging landmarks which equate to the inlet, mid-point and outlet of the carpal tunnel. The nerve/hand was the unit of analysis. Many studies reported both the number of individuals and wrists scanned (as some studies involved bilateral scanning) but if not otherwise stated we assumed imaging was performed unilaterally. If data were missing, unclear or present in an unfavourable format then the authors were contacted by email with a request for more information. Four authors provided additional information upon request. When no reply was received, estimates were derived from graphs or imputed where possible.

Outcomes

The primary outcome was to estimate the normal DTI metrics (FA and MD) of the median nerve in asymptomatic adults. The secondary outcomes were to estimate the differences in DTI metrics (FA and MD) between asymptomatic adults and patients with CTS, and explore the associations between DTI metrics and: age, echo time (TE), repetition time (TR), resolution, the number of diffusion sensitising gradient directions (N\textsubscript{D}) sampled per shell, the b-value(s), different methods of k-space sampling and in-plane acceleration.

Methodological quality assessment

The risk of bias was independently assessed by three review authors (DR, JR and FR) using the ROBINS-I tool and displayed graphically using robvis. Disagreements were resolved by discussion with RGW.

Statistical analysis

The raw data are available via the Open Science Framework (https://osf.io/vqwkp/). The single study performed at 7 tesla was excluded from all meta-analyses given its clinical disparity. Using the meta suite of Stata v16 (StataCorp, Texas), the mean FA and MD from asymptomatic adults were pooled to estimate the normative values. We performed direct comparisons meta-analysis of the mean differences in FA and MD between asymptomatic adults and patients with CTS. Meta-analyses were subgrouped by anatomical location. Restricted maximum likelihood was used to estimate the between-study variance (\(\tau^2\)), with the Knapp and Hartung modification. Heterogeneity was quantified by \(I^2\). Using the metafor package, mixed-effects meta-regression was used to explore potential reasons for the observed heterogeneity in the direct comparisons meta-analysis of FA; the continuous covariates were age, in-plane resolution (mm\textsuperscript{2}), slice thickness (mm), echo time (TE in ms), b-value (mm\textsuperscript{2}/s) and number of diffusion-sensitising gradient directions (N\textsubscript{D}). TE and b-value were modelled as an interaction. Confidence intervals
(CI) were generated to the 95% level. To investigate the possibility of small-study effects for FA between asymptomatic adults and patients with CTS, a funnel plot was constructed with the pseudo CIs contoured by \( \text{tau}^2 \). EviAtlas was used to generate a map of the location of the 1st author’s institution.

## Results

Ultimately, 32 studies\(^{16-19,26-53}\) were included (eFigure 2).

### Study characteristics

Study characteristics are detailed in eTable 1. Overall, we included data from 2643 wrists belonging to 1575 asymptomatic adults and 1068 patients with CTS. Asymptomatic adults were a mean 6 years younger than patients with CTS (95% CI 3, 10). There were approximately twice as many females (1404:746) although this disparity was more pronounced in patients with CTS (660F:193M) than asymptomatic adults (737F:552M). The median number of authors was 6 (IQR 5-8) and studies were derived from 16 different countries (eFigure 3).

Ten studies (32%) were performed at a field strength of 1·5 tesla\(^{17,29,32,35,38,42,44,49,50,52}\), twenty one (65%) at 3 tesla\(^{16,19,26-28,30,31,33,34,36,37,39-41,43,45-48,51,53}\), and one at 7 tesla\(^{18}\). The median echo and repetition times were 87ms (IQR 65-91, range 21-103) and 7000ms (IQR 3800-7650, range 1470-10,254), respectively. Two studies used read-out segment echo-planar imaging (rsEPI)\(^{33,45}\), two did not specify the pulse sequence\(^{19,48}\) and the remainder used single-shot echo-planar imaging (ssEPI). Twelve studies described using the parallel imaging techniques (GRAPPA\(^{18,27,41}\), SENSE\(^{17,28,31,38,40,45,51}\) and CAIPIRINHA\(^{30}\)) and six studies used partial Fourier transformations\(^{30,32,34,38,40,43,45}\). The median slice thickness was 3·5mm (IQR 2·6-4·0mm, range 1·5-5mm). The median in-plane resolution was 1·09mm (IQR 0·7-1·5, range 0·4-1·88). Two studies investigated multiple b-values\(^{35,51}\) via discrete shells, although no studies reported whether acquisitions were half or whole shell and what sample scheme was used. The mean b-value was 1000 s/mm\(^2\) (SD 270, range 325-2000). The median \( N_D \) was 20 (IQR 15-25, range 6-32). A median of 3 signal averages (excitations) were obtained (IQR 2-5, range 1-12). When reported, the mean SNR of the b0 images was 25 (SD 12)\(^{18,28,30,32,35,38}\).

The risk of bias for the included studies is summarised in eFigure 4. The majority of studies were at low risk of methodological bias.

### Evidence Synthesis: Asymptomatic Adults

The FA of the median nerve in asymptomatic adults was reported in 29 studies\(^{16-19,26-35,37-41,43-51,53}\). Overall, the normal FA was 0·58 (95% CI 0·56, 0·59; \( I^2 \) 98%). The FA was highest at the level of the DRUJ (mean 0·61 [95% CI 0·58, 0·63; \( I^2 \) 96%], dropping at the level of the pisiform to 0·57 (95% CI 0·54, 0·61; \( I^2 \) 98%) and lowest at the level of the hook of the hamate (mean 0·54 [95% CI 0·51, 0·57; \( I^2 \) 95%).
The MD of the median nerve in asymptomatic adults was reported in 28 studies\textsuperscript{16–19,26–35,37–41,43–51}. Overall, the normal MD was $1.138 \times 10^{-3} \text{mm}^2/\text{s}$ (95% CI $1.101, 1.174$; $I^2$ 99%). The MD was lowest at the level of the DRUJ (mean $1.073 \times 10^{-3} \text{mm}^2/\text{s}$ [95% CI $1.019, 1.128$]; $I^2$ 93%), increasing at the level of the pisiform (mean $1.180 \times 10^{-3} \text{mm}^2/\text{s}$ [95% CI $1.115, 1.244$]; $I^2$ 96%) and highest at the level of the hook of the hamate (mean $1.151 \times 10^{-3} \text{mm}^2/\text{s}$ [95% CI $1.086, 1.217$]; $I^2$ 98%).

**Evidence Synthesis: Patients with CTS**

The FA of the median nerve in patients with CTS was reported in 19 studies\textsuperscript{16–19,26–28,31–33,37–39,44,46,48–50,52}. Overall, patients had a mean FA of 0.45 (95% CI 0.43, 0.47; $I^2$ 95%). The FA was lowest at the midpoint of carpal tunnel, at the level of the pisiform (mean 0.41 [95% CI 0.38, 0.43]; $I^2$ 86%), compared to the levels of the DRUJ (mean 0.48 [95% CI 0.44, 0.52]; $I^2$ 91%) or hook of the hamate (mean 0.45 [95% CI 0.42-0.48]; $I^2$ 93%).

The MD of the median nerve in patients with CTS was reported in 18 studies\textsuperscript{16–19,26–28,31–33,38,39,44,46,48–50,52}. Overall, patients with CTS had a pooled mean MD of $1.293 \times 10^{-3} \text{mm}^2/\text{s}$ (95% CI $1.227, 1.359$; $I^2$ 99%). The MD was highest at the level of the pisiform (mean $1.372 \times 10^{-3} \text{mm}^2/\text{s}$ [95% CI $1.245-1.500$]; $I^2$ 98%), $1.180 \times 10^{-3} \text{mm}^2/\text{s}$ at the level of the DRUJ (95% CI $1.064, 1.295$; $I^2$ 95%) and $1.335 \times 10^{-3} \text{mm}^2/\text{s}$ at the level of the hook of the hamate (95% CI $1.259, 1.411$; $I^2$ 93%).

**Direct Comparisons Meta-Analysis: Asymptomatic Adults vs. Patients with CTS**

Fourteen studies reported direct comparisons between asymptomatic adults and patients with CTS\textsuperscript{16–18,27,28,31–33,38,39,44,46,49,50}. All studies reported a lower FA in patients with CTS compared to asymptomatic adults (mean difference 0·09 [95% CI 0·07, 0·11]; Figure 2). The largest difference between asymptomatic adults and patients with CTS was at the mid-point of the carpal tunnel, at the level of the pisiform (mean difference 0·12 [95% CI 0·09, 0·16]).

Patients with CTS had a higher mean diffusivity than asymptomatic adults (mean difference 0·12 mm$^2$/s $\times 10^{-3}$ [95% CI 0·08, 0·17], Figure 3). This difference was again most profound at the mid-point of the carpal tunnel, at the level of the pisiform (mean difference 0·16 mm$^2$/s $\times 10^{-3}$ [95% CI 0·05, 0·27]).

**Meta-regression**

Age was negatively associated with the FA in asymptomatic adults whereby each decade of life reduced the FA by approximately 0·003 (adjusted $\beta$ $-2·79 \times 10^{-3}$ [95% CI $-4·78 \times 10^{-3}, -8·12 \times 10^{-4}$]; $I^2$ 97%). However, there was no relationship between age and FA in patients with CTS (adjusted $\beta$ $9·70 \times 10^{-4}$ [95% CI $-2·89 \times 10^{-3}, 4·83x10^{-3}$]; $I^2$ 96%), Figure 4). Increasing age was also associated with MD whereby each decade of life increased MD by approximately 0·108 $\times 10^{-4}$ mm$^2$/s (95% CI $0·073 \times 10^{-4}, 0·140 \times 10^{-4}$; $I^2$ 99%, Figure 5) with no significant difference between asymptomatic adults and patients with CTS.
There was no relationship between $N_D$ and FA (eFigure 5) or MD (eFigure 6). The b-value was not associated with the FA (eFigure 7). There was an inverse relationship between the b-value and MD, whereby increments of 100 mm$^2$/s reduced the diffusivity by $0.04 \times 10^{-3}$ mm$^2$/s ($\beta = 3.849 \times 10^{-7}$ mm$^2$/s [95% CI -5.019 x10$^{-7}$, -2.678 x10$^{-7}$]; $I^2$ 98%, eFigure 8).

There were no significant differences between studies which used ssEPI or rsEPI. There was no association between the in-plane resolution (in square millimetres) and FA (eFigure 9) or MD (eFigure 10). Slice thickness was not associated with FA (eFigure 11) but was negatively associated with MD whereby increments of 1mm reduced the MD by $6.023 \times 10^{-5}$ mm$^2$/s (95% CI 9.754 x10$^{-5}$, 2.294 x10$^{-5}$; $I^2$ 99%, eFigure 12).

The TE was not associated with FA or MD (eFigures 13 and 14). The TR was not associated with FA (eFigure 15) but longer repetition times were associated with lower estimates of MD, whereby increasing the TR by 1 second decreased the MD by $2.990 \times 10^{-6}$ mm$^2$/s (95% CI 4.383 x10$^{-6}$, 1.598 x10$^{-6}$; eFigure 16).

Studies reporting the use of parallel imaging techniques (e.g. GRAPPA, SENSE or ASSET) yielded 5% higher estimates of FA ($\beta = 0.05$ [95% CI 0.02, 0.08]; $I^2$ 98%, eFigure 17) when compared to studies which did not report this information. Parallel imaging methods were not associated with differences in the MD. There was insufficient data to explore different partial Fourier settings. There was no association between the number of signal averages and FA or MD (eFigure 18 and eFigure 19).

Ultimately, mixed-effects multivariable meta-regression showed that having CTS was the strongest independent moderator of the observed heterogeneity in FA (Table 1). Age explained some of the residual between-study variance. The experimental factors we modelled did not explain the residual heterogeneity.

There was no evidence of publication bias (Eggers $\beta = 0.10$ [95% CI 0.06, 0.14]; $p=0.134$, eFigure 20).

**Discussion**

This study demonstrates that throughout the length of the median nerve, patients with CTS have more isotropic diffusion than asymptomatic adults. The largest differences were observed at the mid-point of the carpal tunnel, at the level of pisiform, where patients with CTS had lower fractional anisotropy and higher mean diffusivity. Of clinical importance, we demonstrate that these real-world differences in independent of age and experimental (acquisition) conditions.

There are inherent problems with clinicians diagnosing CTS given that the available tests are largely unreliable. For example, nocturnal paraesthesias and many classical tests such as Phalen and Tinel, the scratch-collapse and sensory threshold testing have poor diagnostic value$^{54,55}$. Despite the widespread use of neurophysiology studies in patients with suspected carpal tunnel syndrome, its diagnostic accuracy is also poor$^{56}$. Moreover, surgeons still perform decompression surgery in the presence of
normal tests\textsuperscript{57}. It has been shown that DTI outperforms standard morphological imaging in patients with peripheral neuropathy\textsuperscript{34}. Building on this premise and the poor performance of other tests, we demonstrate that DTI could fill a diagnostic void in the management of (at least unclear or complex) patients with CTS given that DTI generates reliable, reproducible and standardised proxy measures of nerve health. In the UK, the cost of a non-contrast MRI of the extremity is less than a neurophysiology exam (£389 versus £444; $540 versus $609).\textsuperscript{58} Therefore, we suggest that DTI could be a cheaper and potentially more accurate method of both diagnosing CTS and guiding treatment choices. Incorporating DTI in the real-world management of CTS would be difficult and require significant training for clinicians, changes to infrastructure and clinical pathways but ultimately, we show that DTI yields unique diagnostic information which could of significant and ubiquitous clinical value. Moreover, DTI could add value to the assessment of patients who don't improve after treatment or have bilateral symptomatology, where current tests (e.g., electrical studies) are inherently insensitive principally because they have no normal values.

We observed high statistical heterogeneity which has many potential explanations. The majority of the (statistical) heterogeneity was explained by the presence of CTS and it is plausible that the remainder is explained by the ‘severity’ of disease, which we were unable to capture. For example, we speculate that patients with more severe CTS (e.g., symptoms for years cases, resulting in profound demyelination, axonal loss and fibrosis) are likely to have lower FA and higher MD than patients with recent-onset mild CTS. Age also explained some of the observed heterogeneity and this is unsurprising, given that FA is known to fall in aging peripheral nerves\textsuperscript{59}, just as it does in the white matter tracts of the brain\textsuperscript{60,61}. This is because aging axons lose integrity, undergo demyelination and there is a simultaneous increase in extra-cellular fluid. Importantly, we showed that DTI metrics were sensitive to CTS after adjusting for age. Finally, in highly controlled and extreme conditions, user-specified factors\textsuperscript{62} such as the SNR\textsuperscript{63}, b-value\textsuperscript{64,65}, $N_D$\textsuperscript{66,67}, distortion correction pipelines\textsuperscript{68,69}, tensor fitting methods\textsuperscript{63} and partial volume effects\textsuperscript{70} have been shown affect the DTI parameter estimates, which may explain some of the remaining heterogeneity. Although we could not explore the effects of all these factors, in general we observed that experimental conditions had little or no significant effect on the measured FA or MD. Therefore, despite the statistical heterogeneity, DTI appears to be reliably sensitive to the microstructural changes of the median nerve which occur in CTS.

There were no significant associations between FA and MD, and several core elements of the pulse sequence. Therefore, we suggest that clinicians and researchers wishing to acquire data to simply fit a tensor could optimise their sequence as follows. As tensors are robust to varying b-values (in the hindered range) we suggest a b-value to 300-800mm$^2$/s because this enables a shorter TE, which improves SNR and mitigates the effects of T2 shine-through. By reducing the TE might enable users to take advantage of other vendor-specific options to improve data quality and reduce distortions. Given that increasing the $N_D$ has little effect on tensor fitting\textsuperscript{71}, there are no crossing fibres to model and the median nerve is not tortuous (within or between voxels), we recommend limiting the $N_D$ to approximately 15. The normal
median nerve has a cross-sectional area of 9mm² (3.4mm diameter)\(^2\) and this increases with CTS\(^3\), so we recommend an in-plane resolution of approximately 1mm² to ensure that at least 1 voxel is not affected by partial voluming\(^70\) in all subjects. As that the median nerve is orthogonal to the imaging plane, the slice thickness can be tailored to achieve adequate SNR in the knowledge that it has little effect on the resultant FA. Until work is published to show the exact relaxation properties (T1, T2, T2\(^*\)) of the median nerve we suggest that TR is minimised (e.g., 4500ms) to reduce scan time. When planning the number of signal averages, balanced full datasets should be acquired in opposing phase-encoding directions (blip-up, blip-down) to yield the best corrections of various distortions, via the FMRIB FSL pipeline\(^74\). It should be noted that thicker slices, longer repetition times and more signal averages are associated with lower estimates of MD.

**Limitations**

The main limitation of this study is the inherent and pervasive problem of CTS diagnosis which may have biased the findings. At present there is no internationally agreed diagnostic criteria for CTS and as such, there is clinical variation which is likely to be present in the includes studies. We planned to capture disease severity from the original studies, but this information was not available. As a matter of urgency, the community should work towards a consensus on objective criteria which constitute a diagnosis of CTS. Given the present absence of a reliable test for CTS, we speculate that DTI could fill the void and serve as the reference standard in future studies.

It is widely known that diffusion metrics in the brain are strongly dependent on the preprocessing distortion correction pipelines (i.e., software) used\(^74\) but still there is no consensus on the minimum or indeed ideal suite of corrections to perform. This issue is compounded in the limb owing to an absence of research on the topic and hardware limitations. The majority of the included studies did not describe any form of distortion correction, how the diffusion data were reconstructed or how metrics were extracted from the median nerve. Before DTI can be used clinically, variations in these pipeline should be tested and a universal pipeline and standards for reporting diffusion data should be agreed by consensus.

Some readers will criticise our choice to pool estimates of FA and MD in the presence of high statistical heterogeneity. This was done because forest plots provide an important graphical representation of the variability of measurements in relation to experimental conditions (e.g., b-values and N\(_D\)) which are easy to interpret and moreover, meta-regression in this situation facilitates the exploration of heterogeneity.

**Conclusions**

We provide summary estimates of the normal FA and MD of the median nerve in asymptomatic adults. Furthermore, we show that diffusion throughout the length of the median nerve becomes more isotropic in patients with CTS.

**Declarations**
Conflicts of interests

None.

Data sharing

The raw data are available via the Open Science Framework (https://osf.io/vqwkp/). The statistical syntax is available from the senior author (RGW) upon request.

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Ethical approval

Not applicable

Contributions

DR co-authored and registered the protocol, performed the searches, screened articles, extracted data, performed the risk of bias assessments and co-authored the manuscript. FR screened articles, extracted data, performed the risk of bias assessments and co-authored the manuscript. JR screened articles, extracted data, performed the risk of bias assessments and co-authored the manuscript. RGW conceived the study, co-authored the protocol, supervised the searches and citation chasing, checked all the extracted data for accuracy, performed the statistical analyses, created the artwork and co-authored the manuscript. All authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

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**Tables**

**Table 1.** Mixed-effects meta-regression

<table>
<thead>
<tr>
<th>Covariates</th>
<th>Adjusted Change in Fractional Anisotropy (β)</th>
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*Adjusted R² = 46%, τ² = 0.0047, I² = 97%

*Echo time is a function of the b-value as larger b-values mandate relatively longer echo times, so these variables are modelled as the product to minimise the number of covariables, mitigate collinearity and model the interaction between the two variables

**Figures**
A diagram of nerve fibres (top) and in cross-section (bottom) demonstrating how diffusion tensor imaging metrics change in CTS. In healthy nerves, the axons are enveloped by thick myelin sheaths and arranged relatively tightly, which restricting the diffusion of water to the long axis of the nerve. Chronic compression leads to distortion of the axonal architecture, demyelination and as such, degradation of physiological barriers to the diffusion of water diffusion. Consequently, more diffusion occurs perpendicular to the long axis of the nerve and water is more free to diffuse around the fibres, reducing the factional anisotropy (FA) and increasing the magnitude of diffusion (mean diffusivity, MD).
Figure 2

A forest plot of the fractional anisotropy of the median nerve, at 3 anatomical levels, between asymptomatic adults and patients with carpal tunnel syndrome.
A forest plot of the mean diffusivity of the median nerve, at 3 anatomical levels, between asymptomatic adults and patients with carpal tunnel syndrome.
Figure 4

A scatterplot of study-level estimates of fractional anisotropy in asymptomatic adults and patients with carpal tunnel syndrome, against age in years. The size of the points corresponds to the precision (inverse variance) of the study.
Figure 5

A scatterplot of study-level estimates of mean diffusivity in asymptomatic adults and patients with carpal tunnel syndrome, against age in years. The size of the points corresponds to the precision (inverse variance) of the study.

Supplementary Files

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- SupplementaryMaterials.docx