Shared genetic susceptibility between trigger finger and carpal tunnel syndrome: a genome-wide association study

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Summary

Background Trigger finger and carpal tunnel syndrome are the two most common non-traumatic connective tissue disorders of the hand. Both of these conditions frequently co-occur, often in patients with rheumatoid arthritis. However, this phenotypic association is poorly understood. Hypothesising that the co-occurrence of trigger finger and carpal tunnel syndrome might be explained by shared germline predisposition, we aimed to identify a specific genetic locus associated with both diseases.

Methods In this genome-wide association study (GWAS), we identified 2908 patients with trigger finger and 436579 controls from the UK Biobank prospective cohort. We conducted a case-control GWAS for trigger finger, followed by co-localisation analyses with carpal tunnel syndrome summary statistics. To identify putative causal variants and establish their biological relevance, we did fine-mapping analyses and expression quantitative trait loci (eQTL) analyses, using fibroblasts from healthy donors (n=79) and tenosynovium samples from patients with carpal tunnel syndrome (n=77). We conducted a Cox regression for time to trigger finger and carpal tunnel syndrome diagnosis against plasma IGF-1 concentrations in the UK Biobank cohort.

Findings Phenome-wide analyses confirmed a marked association between carpal tunnel syndrome and trigger finger in the participants from UK Biobank (odds ratio [OR] 11·97, 95% CI 11·1–13·8; p<1×10⁻⁶⁶). GWAS for trigger finger identified five independent loci, including one locus, DIRC3, that was co-localised with carpal tunnel syndrome and could be fine-mapped to rs62175241 (0·76, 0·68–0·84; p=5×10⁻¹¹). eQTL analyses found a fibroblast-specific association between the protective T allele of rs62175241 and increased DIRC3 and IGFBP5 expression. Increased plasma IGFB-1 concentrations were associated with both carpal tunnel syndrome and trigger finger in participants from UK Biobank (hazard ratio >1·04, p<0·02).

Interpretation In this GWAS, the DIRC3 locus on chromosome 2 was significantly associated with both carpal tunnel syndrome and trigger finger, possibly explaining their co-occurrence. The disease-protective allele of rs62175241 was associated with increased expression of long non-coding RNA DIRC3 and its transcriptional target, IGFBP5, an antagonist of IGF-1 signalling. These findings suggest a model in which IGF-1 is a driver of both carpal tunnel syndrome and trigger finger, in which the DIRC3-IGFBP5 axis directly antagonises fibroblastic IGF-1 signalling.

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Research in context

**Evidence before this study**
We did a literature review using Medline to determine the co-occurrence of trigger finger and carpal tunnel syndrome. We searched terms “trigger finger” and “carpal tunnel syndrome”, and included all studies published up to March 28, 2022. We included original research studies in all languages that quantitatively assessed co-occurrence of trigger finger and carpal tunnel syndrome, and excluded individual case reports. We identified three prospective observational studies and 24 retrospective studies. Studies report that trigger finger and carpal tunnel syndrome occur together more often than would be expected by chance.

**Added value of this study**
Our study is, to the best of our knowledge, the largest genome-wide association study (GWAS) for trigger finger.

We leveraged the results of this GWAS to identify a single locus (DIRC3) that is significantly associated with both trigger finger and carpal tunnel syndrome. We used multi-modal expression quantitative trait analysis to trace the mechanism by which DIRC3 modifies risk for these conditions, and provide evidence to show that IGF-1 signalling has a role in disease pathophysiology.

**Implications of all the available evidence**
The long-established co-occurrence of trigger finger and carpal tunnel syndrome might be at least partly explained by a shared germline predisposition, which acts to increase IGF-1 signalling in fibroblasts. Further research should determine whether this pathway might be a valid target for pharmacological management of trigger finger and carpal tunnel syndrome.

Methods

**Study design and participants**
Patients with carpal tunnel syndrome (n=16294) were defined as previously described. To maximise specificity, patients with trigger finger were defined by the intersection of patients who had International Classification of Diseases (ICD)-10 codes for trigger finger (M65.3, M65.30-39), and patients who had Office of Population Censuses and Surveys Classification of Surgical Operations and Procedures (OPCS) codes for tendon release (T723; figure 1A). In sensitivity analyses, we explored including patients who had either ICD-10 or OPCS codes alone (termed the extended cohort). We also included patients with self-reported trigger finger symptoms (UK Biobank self-reported illness code 1619), termed the mixed cohort. Further information for cohort definitions is provided in the appendix (p 2). To define a control group for all subsequent genome-wide analyses, we selected 488263 participants from UK Biobank with imputed genomic data, excluding 31657 participants who did not pass genomic quality control and 20027 participants who had coding of any nature for either carpal tunnel syndrome of trigger finger (appendix p 12).

UK Biobank has approval from the North West Multi-Centre Research Ethics Committee (11/NW/0382), and this study was conducted under UK Biobank study ID 22572. The Oxford carpal tunnel syndrome cohort was derived from two clinical studies that were approved by the National Research Ethics Service (UK): the Pain in Neuropathy Study (PiNS; 10/H07056/35), and the Molecular Genetics of Carpal Tunnel Syndrome (MGCTS) study (16/LO/1920). Written informed consent was obtained from all participants recruited to these studies. Genomic data quality control steps for the UK Biobank trigger finger cohort is summarised in the appendix (p 12).

**Phenotypic association analysis**
To identify diagnoses associated with trigger finger, all UK Biobank first occurrence fields (a specific UK Biobank data resource) and cancer registry data (fields 40005 and 40006) were extracted, with ICD-10 codes mapped to Phecodes. Data entries were binarised to construct a matrix of 694 diagnosis codes (including trigger finger and carpal tunnel syndrome, as defined above) versus 502505 participants from UK Biobank. To determine the association between trigger finger diagnosis pairs, we constructed a 2×2 contingency table for each
Joint analysis (COJO) using a linkage disequilibrium reference derived from 1000 Genomes. Linkage disequilibrium score regression was implemented in the ldsc package for R using a UK Biobank European ancestry linkage disequilibrium reference derived from the PanUK Biobank project. Lead single-nucleotide polymorphisms (SNPs) were annotated using the OpenTargets Genetics portal, which considered three annotations: the nearest coding gene, genes with a cis-expression quantitative trait loci (eQTL) variant in linkage disequilibrium ($r^2>0.8$) with the lead SNP, and the Variant2Gene score.

Co-localisation analysis
Carpal tunnel syndrome summary statistics from our previous GWAS were used. To extract data that were specific to the DIRC3 locus, we filtered the merged summary statistics to a 1MB region centered at rs10203066. To extract a signed linkage disequilibrium correlation matrix for the SNPs in these regions, we used the ld_matrix_local function in the ieugwasr package in R using a linkage disequilibrium reference derived from 5000 randomly selected unrelated European participants from UK Biobank. Co-localisation analyses were implemented in the coloc package in R using the coloc.susie function, with default parameters. The posterior probability for hypothesis 4 (H4), reflecting the existence of a shared causal variant, was extracted. To determine the 95% credible set of co-localised variants, we extracted the posterior probabilities of each SNP, conditioned on H4 being true. These SNPs were functionally annotated using ensembl variant effect predictor in the Ensembl web server.

Replication in FinnGen cohort and meta-analysis
Summary statistics for trigger finger (M13) and carpal tunnel syndrome (G6) were downloaded from the FinnGen portal (release 4). The FinnGen analysis pipeline has been described previously. Plots were generated in LocusZoom using legacy mode and the 1000 Genomes European linkage disequilibrium reference, relative to the index SNP rs10203066. We did a sample size-weighted meta-analysis between our UK Biobank trigger finger summary statistics and FinnGen R4 trigger finger summary statistics, which we then implemented in METAL (version March 2011) using the analyse heterogeneity command.

Oxford–carpal tunnel syndrome cohort sample collection
Sample collection for both the PiNS and MGCTS has been described previously. Briefly, patients with clinically diagnosed carpal tunnel syndrome underwent carpal tunnel decompression surgery, during which tenosynovial specimens were collected. Samples were preserved in RNAlater (Thermo Fisher, MA, USA) before extraction using the High Pure RNA Isolation Kit (Roche, Basel, Switzerland). For paired whole-genome genotyping, DNA
was extracted from whole blood samples using the PureLink Genomic DNA Kit (Invitrogen, MA, USA).

**RNA sequencing**

RNA extraction and library preparation were done as described previously. Reads were aligned to the GRCh37 reference with STAR using the Ensembl 87 gene annotation, with gene-level counts assigned using HTSeq. Count-level batch correction between MGCTS and PiNS cohorts was done using ComBat-seq. To facilitate inter-sample comparisons, count-level data was Trimmed Mean of the M-values (TMM)-normalised and log-transformed to generate log-transcripts per million (log-counts per million [CPM]) data.

**eQTL analysis**

Harmonised summary statistics from our analysis of Genotype-Tissue Expression (GTEx) project data were downloaded from the eQTL Catalogue. eQTL analysis for IGFBP5 in the cohort analysed by Neavin and colleagues was done as described previously. Briefly, for each individual and fibroblast cluster, the quantile-normalised pseudobulk average expression was extracted, and cis-eQTL association statistics were computed using a linear model implemented in MatrixEQTL with one PEER factor as covariate. eQTL analysis for DIRC3 and IGFBP5 in the Oxford-carpal tunnel syndrome cohort was implemented as a Kruskal-Wallis test for gene expression (log-CPM) against genotype.

**IGF-1 in UK Biobank**

IGF-1 plasma levels from the first recruitment visit were extracted from field 30770 and were normalised by Z-scoring, stratified by age (decile) and sex. For Cox regression of trigger finger syndrome-free survival or carpal tunnel syndrome-free survival against Z-scored IGF-1, the time variable used was time from blood sampling to ICD-coded diagnosis of carpal tunnel syndrome or trigger finger, or last follow-up date. Last follow-up date was determined through integration of death status, recruitment visits, and ICD coding dates. The model was adjusted for age at first recruitment visit, sex, and recruitment centre.

**Role of the funding source**

The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report.

**Results**

Phenome-wide analyses were conducted across the whole UK Biobank cohort (n=502 490), which recapitulated a highly significant association between trigger finger and carpal tunnel syndrome (p=1×10⁻⁵⁰; odds ratio (OR) 1·11; 95% CI 1·11–1·13; figure 1B). To explore the phenotypic association between trigger finger and carpal tunnel syndrome, we quantified the overlap between trigger finger and carpal tunnel syndrome (figure 1A) and explored the clinical characteristics of patients with trigger finger, carpal tunnel syndrome, and trigger finger–carpal tunnel syndrome overlap (appendix p 3). We identified evidence for a significantly increased prevalence of type 1 diabetes and type 2 diabetes, as well as significantly increased HbA1c levels (a routinely used biomarker for glycaemic control) in patients with trigger finger–carpal tunnel syndrome overlap compared with patients with carpal tunnel syndrome alone (p=1×10⁻⁴) and patients with trigger finger alone (p=0·01).

To investigate the genetic architecture of trigger finger, we conducted a GWAS in participants with European ancestry (n=456 606) from UK Biobank, incorporating 2908 patients with trigger finger (as previously defined [figure 1A]) who passed sample quality control (appendix p 12), and 436 579 controls (appendix p 2) using a mixed-model approach that accounted for unbalanced case-control ratios, population structure, and cryptic relatedness. There were insufficient numbers of patients with trigger finger in non-European ancestry groups to conduct a sufficiently powered analysis. There was also no evidence of significant confounding, with a genomic inflation factor (λGC 1·058), and we estimated the SNP-based heritability for trigger finger to be 0·8% (SE 0·1%). We identified five independent risk loci, comprising 419 variants, that met genome-wide significance (figure 2A). Conditional analyses showed no evidence of secondary signals at each locus. Using a multi-modal approach to gene-mapping, we identified 21 candidate genes at the five loci (figure 2B; appendix p 6).

In support of our stringent approach to trigger finger phenotype definition, we found that conducting the GWAS by including patients with either ICD-10 or OPCS codes (ie, the extended cohort) or additionally including patients with self-reported trigger finger (ie, the mixed cohort) markedly reduced the power to detect significant associations across the five loci (appendix p 2, 6).

At the DIRC3 locus (encoding a long non-coding RNA [lncRNA], index SNP rs10203066 (p=6·73×10⁻¹³ [OR 0·75]; 95% CI 0·69–0·82) was shared with our GWAS on carpal tunnel syndrome (p=2·20×10⁻¹² [0·88]; 0·85–0·91; figure 3A, C). To confirm that this signal was not driven solely by patients with carpal tunnel syndrome in our trigger finger cohort, we undertook a further sensitivity GWAS analysis excluding any patients with carpal tunnel syndrome (appendix p 2) and confirmed that this signal retained genome-wide significance (p=1·09×10⁻¹⁰; appendix p 6). Consistent with the independent association of this locus with carpal tunnel syndrome and trigger finger, we confirmed an increase in statistical significance by merging all patients with trigger finger and carpal tunnel syndrome (p=1·28×10⁻⁴; appendix p 6).

We conducted a genetic correlation analysis using linkage disequilibrium score regression and identified
evidence of a high correlation between trigger finger and carpal tunnel syndrome traits (genetic correlation estimate 0.70). To identify the loci that were driving this genetic correlation, we conducted a co-localisation analysis with a multiple causal variant assumption (SuSiE coloc). This analysis identified a high (87%) posterior probability that trigger finger and carpal tunnel syndrome share a single causal variant at this locus. To replicate the association between the DIRC3 locus and both trigger finger and carpal tunnel syndrome, we extracted summary statistics for trigger finger and carpal tunnel syndrome from the FinnGen cohort (release 4; https://finngen.gitbook.io/documentation/v/r4/). The association with both trigger finger (1485 patients and 137185 controls; p=0.0055 [OR 0.88; 95% CI 0.81–0.96; appendix p 6]) and carpal tunnel syndrome (8576 patients and 158705 controls; p=1.90×10⁻⁵ [0.91]; 0.88–0.95) was confirmed using a Bonferroni-corrected threshold of p<0.01 (figure 3B, D).

A formal meta-analysis of UK Biobank and FinnGen summary statistics for trigger finger showed directional concordance at all loci, with low heterogeneity across three out of five loci (appendix p 7).

We leveraged the co-localisation analysis between the traits for carpal tunnel syndrome and trigger finger to fine-map the DIRC3 locus by extracting the 95% credible set (n=21) of co-localised variants (appendix pp 8–9). Next, using Ensembl Variant Effector Predictor, we functionally annotated the SNPs with their immediate regulatory environment. One SNP, rs62175241, had significant regulatory consequences by disrupting an enhancer site active in fibroblasts, as well as the binding motifs for a range of transcription factors, including KLF16 and KLF18 (appendix p 10).

To investigate how rs62175241 (p=5.03×10⁻¹⁰ [OR 0.76]; 95% CI 0.68–0.84) might modulate the expression of DIRC3, we conducted an eQTL analysis using data from 53 tissues that were examined as part of the GTEx project. This analysis provided evidence that the effect of rs62175241 (T allele, which is protective for carpal tunnel syndrome and trigger finger, allele fraction 0.14) on DIRC3 expression is highly tissue-specific, with positive regulation in the stomach and spleen, and negative regulation in the testes and amygdala (figure 4A).

In light of evidence that DIRC3 is able to directly activate expression of IGFBP5, with both genes found in the same topologically associating domain, we examined the effect of rs62175241 on IGFBP5 expression in GTeX. This analysis again provided evidence for tissue-specific eQTL associations and showed a discordant effect of rs62175241 on DIRC3 and IGFBP5 expression in the spleen (figure 4B). Considering that fibroblast proliferation is a histological feature of both trigger finger and carpal tunnel syndrome, we further investigated the effect of rs62175241 on IGFBP5 expression in fibroblasts. We re-analysed fibroblast single-cell eQTL data from 79 donors. Four of six fibroblast subtypes showed a significant positive association between the protective T allele of rs62175241 and IGFBP5 expression (appendix p 10), with the strongest association seen in...
LncRNAs inherently have markedly lower abundance than mRNAs; consistent with this and the shallow depth of sequencing in single cell RNA sequencing data, *DIRC3* was detected in fewer than 1% of cells, precluding further analysis.

Next, we analysed the association of rs62175241 on *DIRC3* and *IGFBP5* expression in diseased tenosynovium samples from patients with carpal tunnel syndrome (n=77). We confirmed that the protective T allele was associated with significantly increased *IGFBP5* expression (p=0.033, allele fraction 0.13; figure 4D). *DIRC3* expression levels were again low and did not show allele-specific differential expression (p=0.086, figure 4E). Because *IGFBP5* is a secreted protein, we investigated whether this variant might alter plasma concentration. We analysed a publicly available plasma proteomic GWAS dataset that was obtained from the SomaLogic platform, and found that this variant (tagged by rs10203066, A allele, $r^2=0.99$) was associated with a non-significant increased plasma *IGFBP5* (beta=0.017; p=0.11).

As *IGFBP5* is known to directly antagonise IGF-1 signalling, we hypothesised that higher IGF-1 plasma levels would be associated with significantly increased risk of both trigger finger and carpal tunnel syndrome in UK Biobank participants. We identified significant associations between plasma concentrations of IGF-1 and trigger finger (hazard ratio [HR] per 1 SD 1.04 [95% CI 1.01–1.07]; p=0.02) and carpal tunnel syndrome (HR per 1 SD 1.04 [1.02–1.05]; p=4.23×10⁻⁶), which was concordant with another carpal tunnel syndrome-specific analysis in UK Biobank. If the protective effect of rs62175241 was mediated via antagonism of IGF-1 signalling, we hypothesised that this variant would be associated with the attenuation of growth hormone-regulated phenotypes such as height and lean body mass. To investigate this hypothesis, we extracted growth phenotype summary statistics from UK Biobank and selected traits that met phenome-wide significance (p<1×10⁻⁵). This process identified 22 growth-related traits that were significantly associated with rs62175241 (appendix p 11), all of which had a negative beta, including standing height (p=3.57×10⁻¹⁸), weight (p=8.28×10⁻⁶), forced vital capacity (p=1.14×10⁻⁶), and appendicular lean mass (p=4.90×10⁻⁹). Altogether, the available data...
suggest that IGF-1 is associated with increased risk of both trigger finger and carpal tunnel syndrome, and that the T allele of rs62175241 might act to directly attenuate IGF-1 signalling, thus explaining its protective effect for trigger finger and carpal tunnel syndrome.

**Discussion**

This GWAS for trigger finger identified five risk loci, one of which was also identified in our previous GWAS of carpal tunnel syndrome. Hypothesising that a single genetic variant might contribute to the pathogenesis of both diseases, we fine-mapped this locus to a single putative causal SNP, rs62175241. Our single-cell eQTL analysis provided evidence to show that this variant was associated with tissue-specific modulation of the expression of DIRC3 and its known downstream effector, *IGFBP5*. Bulk RNA sequencing analysis of surgically resected tenosynovium samples from patients with carpal tunnel syndrome showed that this protective variant was associated with enhanced expression of *IGFBP5*. Considering that *IGFBP5* is an antagonist of IGF-1, we found that both trigger finger and carpal tunnel syndrome were positively associated with IGF-1 levels. These findings are important because they provide a direct biological insight into the shared pathophysiological mechanisms contributing to trigger finger and carpal tunnel syndrome (figure 5). Furthermore, our findings provide a starting point for investigating non-surgical interventions for these two common conditions.

To the best of our knowledge, the co-localised locus that mapped to the *DIRC3* gene has not been previously described in association with trigger finger. One previous GWAS was conducted to identify risk loci associated with trigger finger, finding a single genome-wide locus within *KLHL1* that did not replicate. *KLHL1* is an actin-binding protein, and the authors of this study speculated that this variant might lead to fibrocartilaginous metaplasia in tenocytes.

In this GWAS, the putative causal SNP, rs62175241, is located 3731 base pairs from the canonical transcription start site of the *DIRC3* gene. The *DIRC3* locus spans 450 kilobases between the *IGFBP5* and *TNS1* genes. By mapping the chromatin structure of the *IGFBP5-DIRC3-TNS1* gene territory in human keratinocytes, Coe and colleagues showed that *DIRC3* and *IGFBP5* are located within the same topologically-associated domain. They also found two DNA looping interactions between the *DIRC3* locus and *IGFBP5* promoter. In their study, *DIRC3* levels positively correlated with *IGFBP5* in melanoma RNA sequencing samples and they discovered that *DIRC3* acts in cis to control expression of *IGFBP5*

*IGFBP5* expression is altered in several fibrotic disease states. In lung tissue from patients with idiopathic pulmonary fibrosis, *IGFBP5* was upregulated and exogenous *IGFBP5* stimulated extracellular matrix secretion by idiopathic pulmonary fibrosis pulmonary fibroblasts. Furthermore, *IGFBP5* was upregulated in skin fibroblasts from patients with systemic sclerosis. In the present study, the protective T allele at our putative causal SNP had tissue-specific effects on expression of *DIRC3* and *IGFBP5*. In both skin fibroblasts and operative tenosynovium samples from patients with carpal tunnel syndrome, the protective allele was associated with enhanced expression of *IGFBP5*. *IGFBP5* is a highly conserved and multifunctional secreted protein that binds to IGF and can have complex and varying effects on IGF signalling depending on the tissue type and context. In bone, *IGFBP5* inhibits IGF-1 signalling by modulating binding to the IGF-1 receptor. Similarly, in mammary tissue, *IGFBP5* regulates involution by inhibiting IGF-1 signalling. Consistent with the hypothesis that overtactive IGF-1 signalling is important in trigger finger and carpal tunnel syndrome, we discovered that IGF-1 plasma concentrations were positively associated with both
conditions. Furthermore, despite carpal tunnel syndrome generally being associated with decreased height, the protective allele at our putative causal locus was associated with decreased height, suggesting a distinct pathophysiological mechanism.

Several other lines of evidence support the role of IGF-1 signalling in trigger finger and carpal tunnel syndrome. The prevalence of carpal tunnel syndrome\(^2\) and trigger finger\(^3\) is increased in patients with acromegaly, for whom raised IGF-1 levels are characteristic. By normalising levels of IGF-1, either through pituitary resection or somatostatin analogues, increased tendon thickness at the A1 pulley can be reversed, and symptoms of trigger finger ameliorated.\(^4\)

In healthy patients who do not have acromegaly, administering exogenous growth hormone stimulates a rise in IGF-1, and patients can subsequently develop carpal tunnel syndrome.\(^5\) Exogenous growth hormone is also known to increase the risk of type 2 diabetes, which, along with type 1 diabetes, we found to be significantly enriched in patients with both carpal tunnel syndrome and trigger finger\(^6\) (appendix p 3). Somatostatin analogues work not only by reducing pituitary growth hormone secretion but also by stimulating IGFBP5 secretion\(^7\) which, in turn, inhibits IGF-1 signalling. Of note, our phenotypic analysis highlighted a significant association between trigger finger and both rheumatoid arthritis and osteoarthritis, which was even stronger in the trigger finger-carpal tunnel syndrome overlap cohort (appendix pp 3–4). IGF-1 signalling has been implicated in synovitis in both osteoarthritis\(^8\) and rheumatoid arthritis,\(^9\) which would accord with our proposed pathophysiological mechanism underlying the observed phenotypic associations.

We recognise several limitations of the present study. Firstly, our GWAS of patients with trigger finger only included patients of European ancestry. It would be helpful to further characterise the genetic architecture of trigger finger in non-European groups, who are under-represented in GWASs. Although our GWAS had a greater power (81% power) to detect a hypothetical variant with a minor allele fraction of 50% and an OR of 1-2 than a previous study (3% power),\(^10\) our study was still relatively underpowered, especially for low-frequency variants, meaning that other relevant risk loci might not have met our pre-defined significance threshold. Although we were able to replicate our co-localised \(DIRC3\) locus in patients with trigger finger and patients with carpal tunnel syndrome from the FinnGen cohort, we were unable to replicate all five trigger finger loci. This might be partly explained by the size of the replication cohort (1485 patients with trigger finger) and the use of a different case definitions in FinnGen, but might also be consistent with the so-called winner’s curse phenomenon\(^11\) in GWASs. Regardless, these validation data strongly support a role for the \(DIRC3\) locus in both trigger finger and carpal tunnel syndrome. We have identified a biologically relevant mechanism that is likely to underpin the association between our co-localised risk locus and trigger finger and carpal tunnel syndrome. However, we recognise that the association between a protective haplotype at \(DIRC3\), \(IGFBP5\) and \(IGF-1\) are all correlative, and further studies are required to dissect these mechanisms and provide evidence for a causative effect.

In conclusion, we provide evidence for a phenotypic association between trigger finger and carpal tunnel syndrome in patients from the UK Biobank cohort. We identified a putative causal variant in our GWAS of trigger finger that overlaps with carpal tunnel syndrome, which possibly accounts for some of the phenotypic overlap between these two conditions. Through multimodal expression eQTL analyses, we directly linked a protective causal variant in \(DIRC3\) to increased expression of \(IGFBP5\), an \(IGF-1\) antagonist.\(^12\) Finally, we provide evidence to show a significant association between plasma IGF-1 concentrations and both trigger finger and carpal tunnel syndrome, altogether implicating IGF-1 signalling in the pathophysiology of both conditions. Collectively, our findings indicate that the protective variant at rs62175241 acts by enhancing expression of IGFBP5, via \(DIRC3\) which, in turn, inhibits

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**Figure 5: Proposed model for the role of \(DIRC3\) in carpal tunnel syndrome and trigger finger**

The disease-protective T allele of rs62175241 (\(DIRC3\) locus) is associated with increased expression of long non-coding RNA (IncRNA) \(DIRC3\) and increased RNA expression of its direct transcriptional target \(IGFBP5\). \(IGFBP5\) protein is a secreted antagonist of IGF-1 signalling that acts to suppress growth signalling. Increased plasma IGF-1 is associated with both carpal tunnel syndrome and trigger finger, suggesting that IGF-1 signalling is a driver of both conditions. We propose that the association between the T allele of rs62175241 with carpal tunnel syndrome, trigger finger, and growth phenotypes (eg, lean mass) is mediated by increased antagonist of fibroblast IGF-1 signalling, altogether directly implicating IGF-1 signalling in carpal tunnel syndrome and trigger finger pathophysiology. \(DIRC3\)=Disrupted In Renal Carcinoma 3; eQTL=expression quantitative trait. \(IGF-1\)=insulin-like growth factor-1; \(IGFBP5\)=insulin-like growth factor-binding protein-5.
IGF-1 signalling (figure 5). Further studies are required to fully characterise this pathway and delineate whether it might be a valid target for pharmacological management of trigger finger and carpal tunnel syndrome.

**Contributors**

BP, DF, and AW conceived and designed the study. BP constructed the statistical analysis plan and extracted case and control cohorts. SOK conducted the computational analyses. GB assimilated and processed raw RNA sequence data. Wu-Ra extracted DNA for genotyping, and MN conducted quality control and imputation of the genotype data. ABS and DLB recruited and phenotyped the Pain in Neuropathy Study (PiNS) cohort, and conducted the RNA extraction with AW. DN completed single-cell analyses with supervision from JP. DF and AW supervised the project and guided all data analyses. SOK, BP, and AW accessed and verified all data reported in the manuscript. BP, SOK, DF, and AW wrote the manuscript with input from all authors. All coauthors approved the final version of the manuscript.

**Declaration of interests**

DLB reports grants from Lilly and AstraZeneca, has acted as a consultant on behalf of Oxford Innovation for Angen, Biointervene, Bristows, LatigoBio, GSK, Ionis, Lilly, Olink Biosciences, Orion, Regeneron, and Theranexus, and has a patent application “a method for the treatment or prevention of pain, or excessive neuronal activity, or epilepsy” (16/337,428). GB reports consultancy fees from Biocoding and Ivy Farm. JP is chair of Oz Single Cells and has stock or stock options in Sonic Healthcare. All other authors declare no competing interests.

**Data sharing**

RNA sequencing data from the Pain in Neuropathy Study (PiNS) cohort has been reported previously and is available at accession GEO:080283 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE108023). The relevant raw data (count matrix and genotype calls at rs62175241) and code that are necessary to replicate analyses in the expanded Oxford-CTS RNA sequencing cohort are provided on Github (https://github.com/samkleeman1/cts_tf/). A Jupyter Notebook summarising the genotype data quality control and genome-wide association analysis implemented in Regenie is provided on Github (https://github.com/samkleeman1/cts_tf/). Single-cell fibroblast expression quantitative trait (eQTL) summary statistics and applicable source data are published alongside the original manuscript. UK Biobank data can be requested through the application process detailed at https://www.ukbiobank.ac.uk/. Summary statistics for the primary trigger finger GWAS (patients with both International Classification of Diseases[ICD]-10 and Office of Population Censuses and Surveys Classification of Surgical Operations and Procedures[OPCS]-4 codes for trigger finger) have been uploaded to the GWAS Catalog (accession code GCST90104907). Summary statistics for the sensitivity analyses described in the manuscript are available from the corresponding author on request.

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