



Original research

Variants within the *MMP3* and *COL5A1* genes associate with soft tissue injury history in elite male rugby athletes

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ABSTRACT

Objectives: To investigate associations between genetic variants within *COLGALT1*, *COL1A1*, *COL3A1*, *COL5A1*, *KDR*, *MIR608*, *MMP3*, *NID1*, *TIMP2* and *VEGFA* and injury history in elite male rugby athletes.

Design: A case-control genetic association study was conducted on 184 elite male rugby athletes.

Methods: Participants were genotyped for 13 genetic polymorphisms previously associated with soft tissue injury using standard PCR assays. Injury data were collected via a self-reported injury-history questionnaire. Single-locus association and Total Genotype Score (TGS) analyses were conducted using χ^2 tests. In addition, multifactor dimensionality reduction and inferred haplotype analysis were used to identify genetic interactions.

Results: The TT genotype of *MMP3* rs679620 was underrepresented in the non-injured ligament group compared to the ligament sprain and ligament rupture groups (10 %, 32 %, 25 %; $P < 0.04$, respectively). The T allele of *MMP3* rs679620 was overrepresented in the non-injured tendon group compared to the tendinopathy group (50 %, 38 %; $P < 0.02$). The proportion of C allele carriers of *COL5A1* rs12722 was higher in the tendon rupture group than the non-injured tendon group (96 %, 75 %; $P < 0.02$). Furthermore, the T-C inferred haplotype frequency of *COL5A1* rs12722 and *COL5A1* rs3196378 was higher in the tendon rupture, ligament sprain and total injured athlete groups compared to their respective non-injured groups ($P < 0.02$).

Conclusions: This study is the first to identify associations between *MMP3* rs679620 and *COL5A1* rs12722 and soft-tissue injury history in elite male rugby athletes. These findings support the growing evidence that soft-tissue injury could be influenced by an athlete's genetic predisposition.

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Practical implications

- Elite rugby athletes carrying the TT genotype of *MMP3* rs679620 have an increased risk of ligament injury.
- Elite rugby athletes carrying the T-C inferred haplotype of *COL5A1* rs12722 and *COL5A1* rs3196378 have an increased risk of soft tissue injury.

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- In the future, these findings may help to develop injury prevention protocols in those with high-risk genetic profiles.

1. Introduction

Compared to other team sports rugby has one of the highest injury incidence and severity rates, most of these are soft tissue injuries which damage the ligaments, tendons and muscles.¹ This is likely due to the intermittent, collision-based nature of the sport. Furthermore, the increasing size and strength of elite rugby athletes² is likely contributing to the high injury rates in the modern game, due to associated changes in momentum during collisions and changes of direction.

Analysis of match injury data across 16 seasons of the English Premiership identified sprains and ligament injuries had the highest incidence and highest burden of injury (22.6/1000 h and 30 days, respectively).¹ The most recent available data from the men's rugby union World Cup (2019) found ligament injuries to be the most common, accounting for 21.7 % of all injuries reported during matches.³ Furthermore, knee ligament injuries were the most severe causing 935 days absence.³ Tendon injuries, although not as common or severe as ligament injuries, are a considerably debilitating injury for rugby players. Specifically, Achilles tendon (AT) injuries have been in the top five highest burden training injuries across multiple English rugby Premiership seasons (2018/19–2020/21), costing 5.5 days absence per 1000 h.⁴ Achilles tendon injury appears to be particularly debilitating for rugby union forward athletes causing 726 days absence across two seasons.⁵ Both ligament and tendon injuries are highly complex, multifactorial disorders that are determined by the interaction of several extrinsic and intrinsic factors.^{6,7} However, the growing body of evidence around the heritability of ligament and tendon injuries^{8,9} has prompted further research into possible genetic aetiology.

Anterior cruciate ligament (ACL) tears seem at least twice as likely in individuals with a family history of ACL tear compared to those with no family history.¹⁰ Indeed, a twin study found that the genetic contribution to ACL rupture was ~69 %.⁸ Most research into the genetics of ligament injury has utilised genetic association studies (GAS) to investigate the influence of single nucleotide polymorphism (SNP) (a DNA sequence variation when a single nucleotide alters between individuals) variation individually and collectively. From these studies, variants in several genes have been associated with altered risk of ligament injury such as collagen type I alpha I (*COL1A1* rs1800012),¹¹ collagen type III alpha I (*COL3A1* rs1800255),¹² and collagen type V alpha I (*COL5A1* rs12722; rs3196378).¹³ The $\alpha 1$ chains of type I, III and V collagen are coded by these genes, respectively, and are thought to potentially influence collagen repair and regulation.¹⁴ Genetic variants within matrix metalloproteinase-3 (*MMP3* rs591058, rs650108, rs679620) have also been associated with risk of ligament injury.^{15,16} The matrix metalloproteinase protein family aids in regulating the extracellular matrix (ECM), which affects the biomechanical properties of ligaments and tendon.¹⁷ Additionally, nidogen proteins aid in the development of the ECM¹⁸ and will likely affect its structure and function. The nidogen 1 gene (*NID1* rs4666048) was the strongest associated polymorphism identified in a fixed effect meta-analysis as part of a genome wide association (GWAS) study for ACL rupture.¹⁹ Furthermore, vascular endothelial growth factor A (*VEGFA* rs699947) and kinase domain receptor (*KDR* rs1870377) genes have previously been associated with ACL and Achilles tendon injury.^{20,21} These genes code for the VEGFA protein, an endothelial cell mitogen that stimulates angiogenesis, which is essential during the remodelling and repair of injured soft tissue, and its receptor protein KDR.²²

Similarly, it has been suggested that there is a genetic component to the aetiology of tendon injury. Indeed, in a twin study of tennis elbow (epicondylitis) in women,⁹ heritability was estimated at ~40 %. Numerous GAS report associations between Achilles tendinopathy and several

genetic variants across a variety of genes such as *COL5A1* (rs12722),²³ MicroRNA 608 (*MIR608* rs4919510), a small non-coding RNA that mediates gene silencing and translational repression,²⁴ *MMP3* (rs591058, rs650108, rs679620)^{25,26} and tissue inhibitors of metalloproteinases-2 (*TIMP2* rs4789932), which aids in the regulation of the ECM with MMP proteins.²⁶ Furthermore, the Collagen beta(1-O) galactosyltransferase 1 (*COLGALT1* rs8090) gene, which may contribute to the pathogenesis of connective tissue disorders, potentially through aberrant post-translational modifications that impair the function of collagen-modifying enzymes,²⁷ was the strongest associated SNP identified in a fixed-effect meta-analysis as part of a GWAS for Achilles tendon pathology.¹⁹ In addition, Achilles tendon rupture has been associated with variants in the *MMP3* (rs679620) and *TIMP2* (rs4789932) genes.²⁶

Genetic variation may have a strong influence on tendon and ligament structure and function, which could alter an individual's risk of injury. Inter-individual variability of tendon and ligament properties is likely to cause micro and macro-trauma at differing strain levels amongst individuals, thus similar injury-inciting events amongst rugby players and/or playing positions may have vastly different outcomes. This could influence individual injury incidence and severity rates, thus affecting time lost from matches and training as well as potentially affecting early retirement. Ficek et al.,²⁸ found potential evidence of this within male professional footballers, identifying that the *COL1A1* G-T haplotype (rs1107946 and rs1800012, respectively) was associated with reduced risk of ACL injury. Further evidence of this was found in male academy soccer players with a maturation status of pre-peak height velocity with variants of *COL5A1* rs12722 and *VEGFA* rs2010963 associated with a higher prevalence of ligament and tendon injuries.²⁹ However, no association between risk of ACL rupture and several collagen gene variants was found in a cohort of elite female athletes from high-risk team sports.³⁰ This could indicate potential sex and maturation differences in these associations, so further elucidation is necessary.

Past research has shown that for genetic variants previously associated with soft-tissue injury, elite male rugby athletes possess more 'favourable' variants both individually and collectively compared to a non-athlete population (*COLGALT1* rs8090, *COL3A1* rs1800255, *COL5A1* rs12722 and rs3196378, *MIR608* rs4919510, *MMP3* rs591058 and rs679620 and *NID1* rs4660148; three-SNP model of *COL5A1* rs12722, *COL5A1* rs3196378 and *MIR608* rs4919510; 13-SNP total genotype score model (TGS)).^{31,32} However, there is no previous research directly investigating genetic associations with the history of soft tissue injury within an elite rugby athlete population. Thus, the aims of this study were to (1) investigate whether gene variants previously associated with soft-tissue injury risk are associated with history of soft-tissue injury in elite rugby athletes and (2) compare polygenic characteristics between athletes with a history of soft-tissue injury and those with no history of injury. It was hypothesised that the genotypes and alleles associated with soft-tissue injury risk would be overrepresented in elite rugby athletes with a history of injury compared to those with no previous history. Furthermore, it was hypothesised that athletes with no history of soft-tissue injury would have a higher TGS,³³ i.e. a greater combination of favourable genotypes, than athletes with a history of injury.

2. Materials and methods

2.1. Participants

This study was conducted in accordance with the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) guidelines for a case-control observational study. Manchester Metropolitan University ethics committee granted approval of this study, which complies with the Declaration of Helsinki (Ethics code 24048). The participants were from the RugbyGene project (led by Manchester Metropolitan University with national and international collaboration which is ongoing³⁴) and comprised of elite male rugby

athletes of European ancestry (n = 184 (165 rugby union (RU), 19 rugby league (RL); mean (standard deviation) height 1.86 (±0.07) m, mass 102 (±12) kg, body mass index 29.5 (±4.12) kg/m², age 26 (±5) yr) including 37.3 % British, 33.0 % Italian, 20.0 % Irish, and 9.7 % of other nationalities, having given written informed consent. For TGS and SNP-SNP epistasis interaction analyses, 177 elite rugby athletes were utilised as 7 athletes did not have a full data set for all 13 polymorphisms investigated. All participants were considered elite rugby players, as they had competed regularly (at least 5 matches) since 1995 (when rugby became professional) in the highest professional league in the UK, Ireland, or South Africa for RU and the highest professional league in the UK for RL.³¹ Furthermore, 55 % of the RU athletes had competed at international level for a “high performance union” (Regulation 16, <http://www.worldrugby.org>), and 50 % of RL athletes had competed at international level.

2.2. Procedures

The procedures are consistent with those reported previously.³¹ Blood, buccal swab or saliva samples were attained via the following procedures. Blood (36.7 % of all samples) was drawn from a superficial forearm vein into an EDTA tube, saliva (63.2 %) samples were collected into Oragene DNA OG-500 collection tubes (DNA Genotek, Ottawa, Ontario, Canada) and sterile buccal swabs (Omni swab; Whatman, Springfield Mill, UK) were rubbed against the buccal mucosa of the cheek for ~30 s.

DNA isolation was performed with the QIAamp DNA Blood Mini kit and standard spin column protocol (Qiagen, West Sussex, UK). Briefly, 200 µL of whole blood/saliva, or one buccal swab, was lysed and incubated, the DNA washed, and the eluate containing isolated DNA stored at 4 °C.

Genotyping for 13 polymorphisms (see list below) was performed using two protocols. Protocol one: Approximately 20 % of the DNA samples were genotyped via real-time PCR using a StepOnePlus (Applied Biosystems, Paisley, UK) as previously described,³¹ with adjustment of thermocycling conditions depending on reagents utilised. Protocol two: Approximately 80 % of the DNA samples were genotyped by combining 2 µL GTXpress Master Mix (Applied Biosystems), 0.2 µL Fast GT Sample Loading Reagent (Fluidigm, Cambridge, UK), 0.2 µL H₂O and 1.6 µL of purified DNA, for samples derived from blood and saliva. Furthermore, 1.78 µL assay (Applied Biosystems), 1.78 µL Assay Loading Reagent (Fluidigm, Cambridge, UK) and 0.18 µL ROX reference dye (Invitrogen, Paisley, UK) were combined per assay inlet. An integrated fluid circuit controller RX (Fluidigm) was used to mix samples and assays using a Load Mix (166x) script. PCR was performed using a real-time FC1 Cycler (Fluidigm, Cambridge, UK) GT 192X24 Fast v1 protocol. The 192X24 microchip plate was then placed into the EP1 Reader (Fluidigm) for end-point analysis. Duplicates of all samples were in 100 % agreement for both protocols. For both protocols, the appropriate

TaqMan assays were utilised (Applied Biosystems, Paisley, UK) and assay context sequences for each polymorphism are shown in Table 1.

2.3. Soft-tissue injury history

Soft tissue injury history in elite rugby athletes was collected utilising a self-reported injury-history questionnaire (Supplementary document). This was developed by the current investigators using prior literature³⁵ and in consultation with medical practitioners and experienced researchers. Athletes were asked to provide details of their geographic ancestry, playing position, playing history, highest level of play, tendon and ligament injury incidence independent of mechanism, and whether each injury was medically diagnosed. An investigator assisted participants with completion of the questionnaire to maximise accuracy. To reduce athlete-response bias, medical staff and coaches were not present during questionnaire completion.

2.4. Calculation of TGS

To quantify the combined influence of the candidate polymorphisms (Table 1) an additive TGS algorithm was utilised,³³ based on the assumption of codominance effects of the alleles. The homozygote genotypes with the lower soft tissue injury risk, according to prior literature, were allocated a ‘genotype score’ of 2, heterozygote genotypes were scored 1 and the higher soft tissue injury risk homozygote genotypes were scored 0.

TGS model

TGS = (100/26)*COLGALT1_{rs8090} + COL1A1_{rs1800012} + COL3A1_{rs1800255} + COL5A1_{rs12722} + COL5A1_{rs3196378} + KDR_{rs1870377} + MIR608_{rs4919510} + MMP3_{rs679620} + MMP3_{rs591058} + MMP3_{rs650108} + NID1_{rs4660148} + TIMP2_{rs4789932} + VEGFA_{rs699947}

A TGS of 100 represents the ‘perfect’ polygenic profile for soft tissue injury risk and 0 represents the ‘worst’ possible outcome for the variants examined in this study.³²

2.5. Data analysis

Pearson's Chi-square (χ²) tests were used to measure Hardy-Weinberg equilibrium and to compare genotype (using three analysis models: additive, recessive, and dominant) and allele frequencies between injured and non-injured groups. The group comparisons were as follows: total injured athletes (TIA) vs total non-injured athletes (TNIA), then sub-group analysis of athletes who had a history of; tendon rupture (TR) vs no tendon injury (NIT); tendinopathy (TD) vs NIT; ligament rupture (LR) vs no ligament injury (NIL); and ligament sprain (LS) vs NIL. With 80 % statistical power, analyses of total injured

Table 1 Genotype score of each polymorphism.

Gene name	Gene abbreviation	rs number	Assay context sequence	Polymorphism	Genotype score (GS)
Collagen beta(1-O) galactosyltransferase 1	COLGALT1	8090	COLGALT1 rs8090: CTCCC[A/G]GTCCC	A/G	AA = 2, GA = 1, GG = 0
Collagen type I alpha 1	COL1A1	1800012	COL1A1 rs1800012: CGCCC[A/C]CATTCC	A/C	AA = 2, AC = 1, CC = 0
Collagen type III alpha 1	COL3A1	1800255	COL3A1 rs1800255: GTGGA[A/G]CTGGT	A/G	GG = 2, GA = 1, AA = 0
Collagen type V alpha 1	COL5A1	12722	COL5A1 rs12722: ACCCA[C/T]GCGCC	C/T	CC = 2, CT = 1, TT = 0
		3196378	COL5A1 rs3196378: ACCCC[A/C]GCCCT	C/A	CC = 2, CA = 1, AA = 0
Kinase Domain Receptor	KDR	1870377	KDR rs1870377: ACAGC[A/T]TGGCT	A/T	TT = 2, TA = 1, AA = 0
MicroRNA 608	MIR608	4919510	MIR608 rs4919510: CAGCT[C/G]CGTTT	G/C	GG = 2, GC = 1, CC = 0
Matrix metalloproteinase-3	MMP3	591058	MMP3 rs679620: TTTT[C/T]GAGGT	T/C	TT = 2, TC = 1, CC = 0
		650108	MMP3 rs591058: GAAAT[C/T]GAGAA	G/A	GG = 2, GA = 1, AA = 0
		679620	MMP3 rs650108: TTAGA[A/G]GTAGC	T/G	TT = 2, TC = 1, CC = 0
Nidogen 1	NID1	4660148	NID1 rs4660148: TTTT[C/G]TITGGG	T/G	TT = 2, TG = 1, GG = 0
Tissue inhibitors of metalloproteinases-2	TIMP2	4789932	TIMP2 rs4789932: TATCT[A/G]CTGTA	G/A	GG = 2, GA = 1, AA = 0
Vascular endothelial growth factor A	VEGFA	699947	VEGFA rs699947: TGGCA[A/C]GATCT	C/A	CC = 2, CA = 1, AA = 0

Tendon and ligament injury-associated alleles in previous literature are underlined.³² Assay context sequences with alleles corresponding to VIC/FAM, respectively, highlighted in bold.

athletes compared with total non-injured athletes were able to detect a small-moderate effect size (w) of 0.25 and analyses between smallest injury subgroups (LR vs NIL) were able to detect a moderate effect size (w) of 0.32. For each polymorphism, 20 tests were subjected to Benjamini–Hochberg corrections to control for false discovery rate and probability values are reported. Where appropriate, odds ratios (ORs) were calculated to estimate effect size. Total genotype scores for all groups were not normally distributed, therefore Mann–Whitney U tests were utilised to compare TGS between injured athlete groups and non-injured. Means and extent of kurtosis were calculated to describe the distribution of TGS within groups. χ^2 tests were used to compare the frequency of injured athletes and non-injured athletes in the top and bottom thirds of TGS scores. Bonferroni adjustment was utilised where appropriate to control for false discovery. We also evaluated the ability of the TGS to correctly distinguish injured from non-injured athletes across all groups by receiver operating characteristic (ROC) curves, calculating the area under the curve (AUC) and 95 % confidence intervals (95 % CI). Multifactor dimensionality reduction (MDR) software (<https://sourceforge.net/projects/mdr/>) was used to identify any possible SNP–SNP epistasis interactions. Haplotypes were inferred using SNPStats. SPSS for Windows version 29 (SPSS, Chicago, IL) software was used for analysis. P values < 0.05 were considered statistically significant.

3. Results

3.1. Injury analysis

Twenty-nine athletes reported no relevant soft tissue injury, whilst 155 athletes had suffered from a relevant form of soft tissue injury. A breakdown of the sub-group injury totals for each genotype can be seen in Supplementary Table 1. Fifty-four athletes had suffered from both ligament and tendon injuries, 13 athletes had both ligament and tendon ruptures; and four athletes had injuries across all four categories (tendon and ligament ruptures, tendinopathy and ligament sprains). Six athletes had more than one area of tendinopathy, 18 athletes suffered more than one ligament rupture, and 21 athletes had more than one area of ligament sprain. A breakdown of injury location for each sub-group can be found in Supplementary Table 2.

3.2. Genotype and allele frequencies

Genotype frequencies were in Hardy–Weinberg equilibrium for 96 of the 104 possible values across all polymorphisms in the injured and non-injured groups apart from *COLGALT1* rs8090 (LS group), *COL5A1* rs12722 (NIT and TR groups), *COL5A1* rs3196378 (NIT, LS and TIA groups), *MIR608* rs4919510 (TNIA group) and *TIMP2* rs4789932 (TR group) (Supplementary Table 1).

For *MMP3* rs679620, the TT genotype and T allele were underrepresented, whilst the proportion of C allele carriers was overrepresented in TD (13.7 %, 38.3 % and 86.3 % respectively) compared to NIT (28.1 %, 50.0 % and 71.9 %; $P < 0.04$, Fig. 1). However, the TT genotype and T allele were underrepresented, whilst the proportion of C allele carriers was more common in NIL (10 %, 36.3 % and 90 %, respectively) compared to LR (32.4 %, 52.1 % and 67.6 %, $P = 0.03$, Fig. 1). Similarly, the TT genotype was underrepresented, whilst the proportion of C allele carriers was more common in NIL (10 % and 90 %) compared to LS (25.2 % and 74.5 %, $P = 0.04$). The LR and LS groups had over three times the odds of carrying the TT genotype compared to NIL (LR odds ratio (OR): 4.3, 95 % confidence intervals (CI): 1.4–13.6; LS OR = 3.0 (1.0–9.2)). For allele/genotype frequency data for all SNPs please refer to Supplementary Table 1.

For *COL5A1* rs12722, the TC genotype and proportion of C-allele carriers were more common in TR (73.1 % and 96.2 %, respectively) compared to NIT (40.5 % and 75.0 % $P < 0.02$). The TR group had eight times the odds of carrying the C allele compared to NIT (OR = 8.3 (1.1–64.3)). There were no differences in genotype or allele frequencies for *COL5A1* rs12722 between any other groups.

There were no differences in genotype or allele frequencies between any groups for the other 11 genetic variants.

3.3. Haplotype and SNP epistasis analysis

The T-C inferred haplotype frequency of *COL5A1* rs12722 and *COL5A1* rs3196378, respectively, was lower in NIT, NIL and TNIA (11.7 %, 6.7 % and 2.2 %, respectively) compared to TR, LS and TIA (37.7 %, 17.8 % and 16.2 %, $P < 0.02$, Fig. 2). Additionally, the C-A inferred haplotype frequency of *COL5A1* rs12722 and *COL5A1* rs3196378 was higher in TR and LS than their respective non-injured groups ($P < 0.05$). Furthermore, the C-C inferred haplotype was higher in NIT than TR ($P < 0.01$). There were no inferred haplotype frequency differences for *MMP3* rs591058, rs650108 and rs679620 between any injured and non-injured groups. Multifactor dimensionality reduction analysis could not identify a model to discriminate between any injury group or sub-group and the respective non-injured group ($P \geq 0.35$).

3.4. Total genotype score

There was a significant difference between the TGS of LS compared to NIL ($P = 0.02$), with the frequency distribution for LS and NIL shown in Fig. 3. There were no differences in TGS between any other injured and non-injured groups (TR vs NIT, TD vs NIT, LR vs NIL and TIA vs TNIA; $P > 0.05$). Mean (standard deviation) TGS and kurtosis statistics are reported in Table 2. When the numbers of injured and non-injured athletes in the upper and lower thirds of the TGS were compared, there were no differences between any injured and non-injured groups

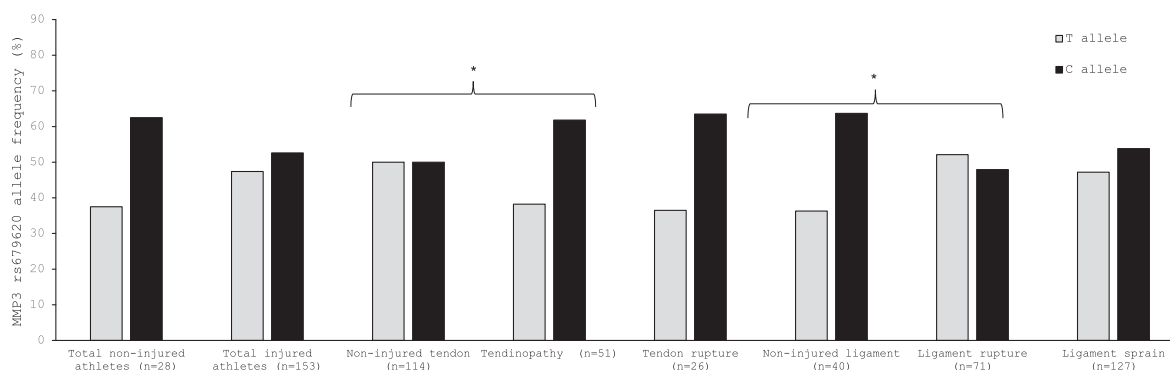


Fig. 1. Allele frequency of *MMP3* rs679620 for all injured and non-injured groups. Asterisks (*) indicate a difference in allele frequency between the injured group and their respective non-injured group ($P < 0.05$).

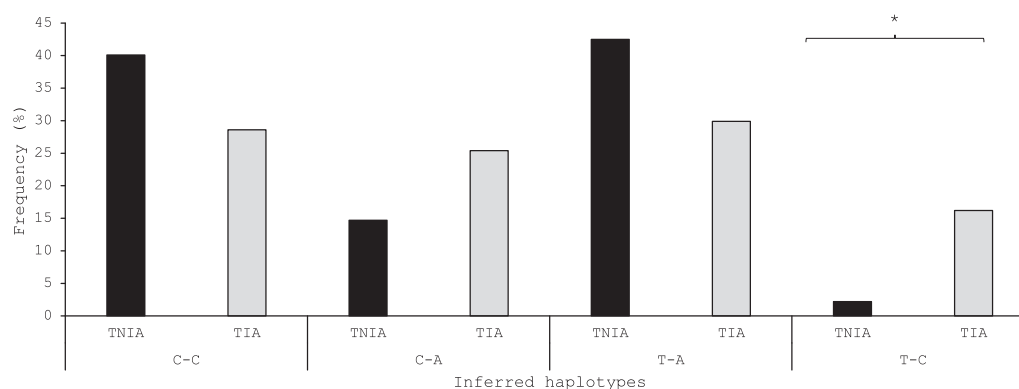


Fig. 2. Inferred haplotypes derived from *COL5A1* rs12722 and rs3196378. TNIA = total non-injured athletes, TIA = total injured athletes. *Different from TNIA ($P < 0.001$).

($P > 0.05$). Finally, the TGS was able to distinguish between LS and NIL (AUC = 0.62; 95 % CI = 0.52–0.72, $P < 0.02$, Fig. 4), but could not distinguish between any other groups.

4. Discussion

This study is the first to identify associations between *MMP3* rs679620 and *COL5A1* rs12722 and soft-tissue injury history in elite male rugby athletes, thus indicating a likely inherited advantage from carrying protective genetic variants involved in collagen and ECM structure and function. Furthermore, the T-C inferred haplotype frequency of *COL5A1* rs12722 and *COL5A1* rs3196378 was higher in the tendon rupture, ligament sprain and total injured athlete groups compared to their respective non-injured groups. As such, the TGS was able to distinguish between the non-injured ligament group and the ligament sprain group. These combined findings suggest a likely polygenic influence on soft tissue injury risk in rugby. As hypothesised, elite male rugby athletes with a history of soft-tissue injury mostly carried more of the apparent injury-risk genotype/alleles than non-injured athletes, although this was not consistent for all polymorphisms.

The TT genotype and T allele of *MMP3* rs679620 were overrepresented in the non-injured tendon athlete group compared to the tendinopathy group which aligns with previous findings.^{25,36} Conversely, the TT genotype and T allele were underrepresented in the non-injured ligament group compared to the ligament rupture group. The TT genotype was also underrepresented in the ligament sprain group. Indeed, the ligament rupture and ligament sprain groups had over three times the odds of carrying the TT genotype compared to non-injured athletes. Similar findings have been seen within sporting populations for non-contact ACL ruptures¹⁶ and knee injuries.²⁹ The

mRNA expression profile of *MMP3* appears to contrast between ACL injury and Achilles tendinopathy.^{37,38} Therefore, the opposing findings within this study could be due to underlying differences in the pathophysiology of tendon and ligament injuries. Alterations in ECM homeostasis are thought to play a role in soft tissue injury risk. Specifically, *MMP3*, which regulates ECM homeostasis via proteolytic activity, is considered to be an essential regulator of matrix degradation and remodelling.³⁸ The present findings suggest that the rs679620 missense polymorphism may, along with other variants, play a role in tendon and ligament injury risk, possibly via the regulation of ECM homeostasis.

Previously, the C allele of *COL5A1* rs12722 was identified as protective from ligament injury in females¹³ and Achilles tendinopathy in males and females.²³ However, we found the proportion of C allele carriers to be higher in the tendon rupture group, with the TT genotype overrepresented in the non-injured tendon group. This aligns with Hall et al.,²⁹ who found that male academy soccer players carrying the CC genotype had a higher prevalence of musculoskeletal and ligament injury than T allele carriers. Furthermore, when *COL5A1* rs12722 and rs3196378 were combined, the T-C inferred haplotype was overrepresented in the tendon rupture, ligament sprain and total injured athlete groups compared to their respective non-injured groups. The *COL5A1* 3' untranslated region, where rs12722 and rs3196378 are located, has been shown to affect mRNA stability,³⁹ which may lead to altered *COL5A1* mRNA secondary structure – possibly influencing type V collagen production. Although this proposed mechanism has still to be fully elucidated, our data, which align in part with previous findings, suggest that variants within *COL5A1* may influence the soft-tissue injury incidence of elite rugby athletes.

COLGALT1 rs8090 and *NID1* RS4660148 were the strongest SNPs associated with Achilles tendon pathology and ACL rupture in a fixed-

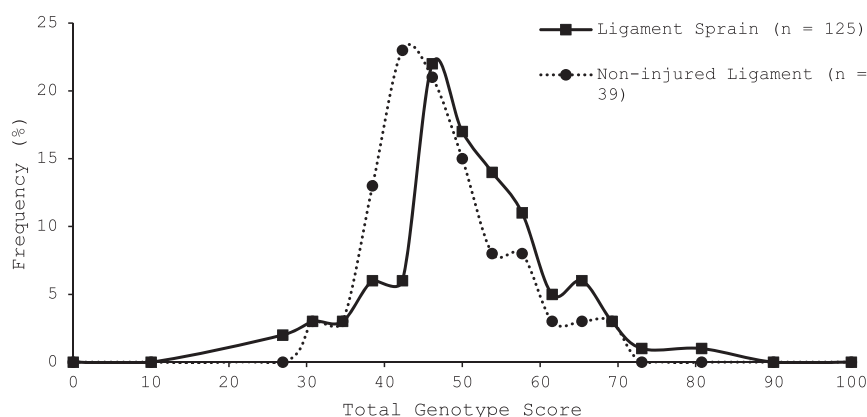


Fig. 3. Frequency distribution of TGS. There was a significant difference in TGS between LS and NIL ($P < 0.02$).

Table 2
TGS mean and kurtosis.

Group	Mean (SD) TGS	Mean (SE) kurtosis statistic
Total non-injured athletes	47.4 (8.0)	1.6 (0.9)
Total injured athletes	49.8 (10.2)	−0.0 (0.4)
Non-injured tendon	49.7 (10.3)	0.1 (0.5)
Tendon rupture	49.3 (9.6)	0.3 (0.9)
Tendinopathy	48.2 (8.5)	−0.2 (0.7)
Non-injured ligament	47.1 (8.2)	0.6 (0.7)
Ligament rupture	50.3 (11.3)	−0.7 (0.6)
Ligament sprain	50.3 (9.8)	0.4 (0.4)

effect meta-analysis within a GWAS,¹⁹ though they did not reach genome-wide significance ($P > 6 \times 10^{-5}$; $P > 5 \times 10^{-6}$, respectively). *COL3A1* rs1800255 and *MIR608* rs4919510, have also previously been linked with ligament and tendon injuries.^{12,24} However, no associations were observed between these variants and tendon or ligament injury within this study. *COL3A1* encodes type III collagen, which plays a key role in type I collagen fibrillogenesis and is influenced by *MIR608*.²⁴ Elevated collagen III levels during fibrillogenesis reduce collagen I content, resulting in smaller, disorganised fibrils and diminished tensile strength.¹⁷ Additionally, *COLGALT1* has been shown to affect collagen I function, with mouse models linking its mutation to musculoskeletal abnormalities.⁴⁰ Thus, it was considered that these variants would influence soft tissue injury risk. Additionally, prior research has found all four variants to be associated with elite status within a male rugby population,³¹ suggesting that they may influence performance-related traits such as muscular strength, particularly relevant for rugby athletes. However, based on our present findings they appear to have limited effect on soft tissue injury.

COL1A1 rs1800012 and *KDR* rs1870377 have previously been associated with ligament injury,^{11,20} whilst *TIMP2* rs4789932 and *VEGFA* rs699947 have been linked to Achilles tendon pathology and ligament injury.^{20,21,26} Based on this prior evidence, all four variants were investigated for potential associations with soft-tissue injury. However, no significant associations were identified. These results may reflect inconsistencies in the existing literature, as only findings related to *COL1A1* rs1800012 have been consistently replicated. Therefore, further research is required to clarify the functional relevance of these polymorphisms.

When polygenic analysis was performed, there was a higher mean TGS in the ligament sprain group compared to the non-injured ligament group. Additionally, ROC analysis could significantly discriminate the non-injured ligament group from the ligament sprain group. This

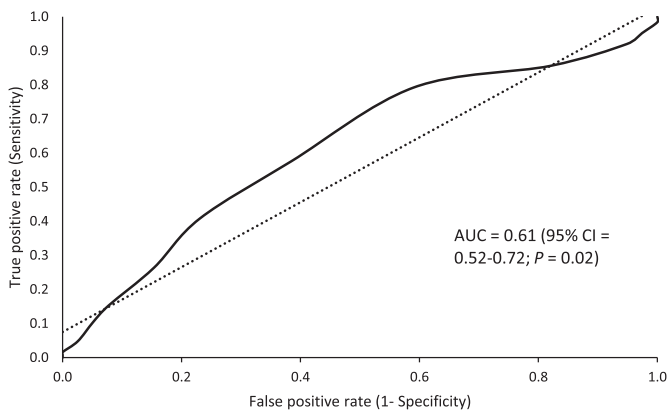


Fig. 4. Receiver operating characteristic curve (ROC) summarising the ability of TGS to classify ligament sprain from non-injured ligament rugby athletes. AUC indicates the area under the curve (95 % confidence intervals).

suggests that rugby athletes with no history of ligament sprain do not appear to carry ‘preferable’ soft-tissue injury associated polygenic profiles. A possible reason for this could be due to the equivocal evidence base of the prior literature regarding the ‘risk’ allele of each SNP, with only four (*COL1A1* rs1800012, *COL3A1* rs1800255, *COL5A1* rs12722 and *MMP3* rs679620) of the thirteen polymorphisms studied having had their ‘risk’ allele replicated in a comparable cohort. There were no other differences in TGS between any other injured group and their respective non-injured groups. Furthermore, when the top and bottom thirds of the TGS were compared, there were no differences found between any groups. The results could in part be down to the relatively small sample sizes of the individual TGS groups, reducing their statistical power. However, it is most likely that at this present time the evidence base for allocating ‘risk’ alleles for TGS analysis on soft-tissue injury is limited due to the lack of consistent findings across polymorphisms.

The present study is not without limitations. The retrospective nature of injury data collection is susceptible to recall bias due to reliance on memory. However, retrospective studies are time- and resource-efficient and promote greater participation from elite sporting populations, thus it was deemed appropriate for this cohort. Furthermore, this study did not account for mechanism of injury such as via contact or non-contact which could influence the results and should be considered when interpreting the present data. Finally, the aetiology of soft-tissue injury is extremely complex including multiple intrinsic and extrinsic factors. Therefore, the data provided in this study should be regarded within that context, particularly due to the relatively small number of genetic variants studied.

The practical application of genetic data for injury risk management remains limited at this time. Whilst findings from this and other studies strongly suggest a heritable component to soft tissue injury susceptibility, the specific genetic variants, both individually and in combination, have yet to be fully identified and understood. Moreover, mechanistic evidence detailing how these variants influence injury risk is currently sparse. Nonetheless, as additional variants are discovered and functional pathways are better characterised, such data may eventually contribute to more personalised approaches to injury prevention and management amongst athletes.

5. Conclusion

We have presented the first genetic associations between *MMP3* rs679620 and *COL5A1* rs12722 and soft-tissue injury history in elite male rugby athletes. Furthermore, the T-C inferred haplotype of *COL5A1* rs12722 and *COL5A1* rs3196378, respectively, was also associated with soft-tissue injury. The functionality of these genetic variants needs further elucidation to identify how soft-tissue injury risk may be affected. Nevertheless, the current data suggest that soft-tissue injury could be influenced by an athlete’s genetic predisposition. This study provides further insight into the detailed aetiology of soft tissue injuries within elite male rugby and may, in future, be worthy of consideration for managing the interindividual variability of injury risk in rugby.

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Confirmation of ethical compliance

Manchester Metropolitan University ethics committee granted approval of this study, which complies with the Declaration of Helsinki (Ethics code 24048).

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CRediT authorship contribution statement

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Declaration of interest statement

The authors declare that they have no competing interests.

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