Genetics and Sport Injuries: New Perspectives for Athletic Excellence in an Italian Court of Rugby Union Players

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Abstract: Several genes are involved in sport performance, especially in injuries incidence. The aim of this study was to investigate the association of ACE, ACTN3, COL1A1, and MCT1 genotypes and injuries in rugby players in order to find a genotype/phenotype correlation and provide useful information improving athletic performance. One-hundred male professional and semiprofessional rugby players were selected. Analysis was performed genotyping the genes ACE, ACTN3, COL1A1, and MCT1 as candidate gene of interest involved in athletic performance. A control group of non-athletic Italian male participants was analyzed to compare the results. We found statistical significance of MCT1 rs1049434 AA for total injuries ($\chi^2 = 0.115; p = 0.003$) and bone injuries ($\chi^2 = 0.603; p = 0.007$) in the rugby athlete population. No statistical significance was found between injury incidence and ACE, ACTN3, COL1A1 genotypes. The MCT1 AA genotype is associated with the incidence of total and bone injuries in the rugby player population. Although environmental factors such as lifestyle, diet, training, and stress can influence athletic performance, our data demonstrated the importance of genetic study in sport aimed at developing personalized training and achieving the best possible athletic excellence.

Keywords: rugby union; ACE; ACTN3; COL1A1; MCT1; injuries

1. Introduction

Sport-specific performance tests provide useful information to improve athlete performance at different levels, but more importantly to get a clearer picture of injury risk for each athlete. Several studies examined the incidence of injury in different contact sports [1]. Rugby union (RU) is an intermittent running-based team sport with high frequency of physical contacts [2] and its injury rate is higher than other team sports (but comparable to other collision sports) [3]. For a rugby athlete, injuries are one of the most disabling factors. Direct (traumatic) injuries result from collisions or physical impacts during sporting activities, while indirect (non-traumatic) injuries are caused by systemic failure due to overexertion during practice [4] such as a structural muscle injury or inappropriate warm-up [5]. Inadequate recovery after training and working beyond the pain threshold may be other factors contributing to the injury aggravation.
Different studies demonstrated genomic involvement in sports performance and the importance of correlation between genomics and sport excellence [6,7]. The correlation between genetics and sport susceptibility may provide information to improve athletic performance, including injury risk reduction [8]. Genetic component may explain approximately 66% of the variance in athlete status depending on sport discipline [9]. In fact, genotype-phenotype correlation could provide decisive information to guide the athlete toward better sports performance [10].

Angiotensin converting enzyme (ACE) gene encodes the enzyme responsible for the conversion of angiotensin I into form II expressed in various tissues including skeletal muscles. ACE I/D (rs1799752) involves an insertion (I) or a deletion (D) of a 287 bp Alu sequence in intron 16 of the gene, generating three different genotypes: II (Insertion in homozygosity), ID (heterozygosity of Insertion/Deletion), DD (Deletion in homozygosity). Blood level of ACE activity in people with DD genotype is twice as high as those with genotype II, while blood ACE activity is intermediate in people with ID genotype. I allele is associated with lower serum and tissue ACE activity and better performance in endurance sports, whereas D allele shows that its increased circulatory and tissue activity improve performances in power or sprint sports [11].

Alfa-actinin-3 (ACTN3) gene encodes α-actininin-3 protein typically expressed in rapidly contracting muscle fibers. R577X (rs1815739) genotype involves a substitution of the amino acid arginine with a stop codon. ACTN3 XX genotype is associated with a lower percentage of fast-twitch fibers [12], while ACTN3 RR genotype has been associated with athletic power/performance [13]. X allele carriers exhibit an increases risk of general sports injuries, whereas R allele carriers show a greater increase in strength and power following high-load endurance training [14].

Collagen Type I α 1 Chain (COL1A1) gene codes for the α1 chain of Col type I, which is responsible for the high tensile strength of tendons and ligaments [15]. Several studies indicated an association between the COL1A1 TT (rs1800012) genotype and a lower risk of soft tissue injuries such as cruciate ligament ruptures, Achilles’ tendinopathy and rupture, shoulder dislocation [16,17].

Monocarboxylate transporter 1 (MCT1) mediates the rapid transport of lactate, especially in skeletal muscle, where rapidly glycolytic muscle fibers produce high amounts of lactate. MCT1 (rs1049434) T1470A genotype is associated with symptomatic deficiency in lactate transport: the T allele correlates with a 35–40% reduction in lactate transport rate compared to the A allele [18].

Basing on the results of previous studies that correlated genes with the risk of injuries, we aimed to investigate ACE, ACTN3, COL1A1, and MCT1 as candidate genes of interest in order to evaluate a possible association between genotypes and the injury risk in an Italian cohort of rugby union players.

2. Materials and Methods
2.1. Participants and Study Design

A total of 100 male professional and semi-professional rugby players from Lazio Rugby 1927 and from G.S. Fiamme Oro Rugby of Polizia di Stato were enrolled. Except for 10 participants, the other participants were Italian (Caucasian). All athletes included in the study based on medical history and injury data, while athletes with personal and family history related to osteoporosis, collagenopathies and tendinitis were excluded. Finally, injuries were stratified according to the type of tissue involved: bone, muscle and tendon/ligament. Contusion and laceration were excluded from the study.

As control group, 100 Italian male no-athletic participants have been selected. All participants were older than 18 years and provided written informed consent. The present study is part of the investigation “Anthropometric and molecular profile of professional and semiprofessional athletes” and the study protocol was approved by the Ethics Committee of the Università Cattolica del Sacro Cuore in Rome (ID: 1858) in accordance with Declaration
of Helsinki for Human Research. In Figure 1 (A/B) we provided an overview of the study design.

![Sample Collection](image1.png)

**A**

![Sample Processing](image2.png)

**B**

**Figure 1.** The figure shows an overview of our study design. (A) The figure describes the collection of sample and injuries data in rugby union elite. (B) The figure represents a schematic laboratory workflow for the blood sample and buccal swabs.

### 2.2. Sample Preparation and DNA Analysis

All sample were analyzed at the Fondazione Policlinico Universitario Agostino Gemelli IRCCS in Rome, Italy. Blood or buccal swab samples were collected for each participant. Genomic DNA was extracted from blood or buccal swab samples using the automated MagCore Nucleic Acid Extractor (RBC Bioscience Corp., New Taipei City, Taiwan) instrument, in association with MagCore® Genomic DNA Whole Blood Kit. DNA concentration was determined using the NanoPhotometer P-Class (Implen, Westlake Village, CA, USA). Analysis by PCR was conducted using primers showed in Table 1. PCR and Sanger sequencing were used for MCT1 and COL1A1 genotyping. In detail, PCR products were sequenced on the 3500 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) in association with SeqScape™ software. For ACE (I/D) genotyping, PCR and 4% electrophoresis gel were used. PCR/RFLP fragments analysis was used for ACTN3 R577X genotyping. ACE and ACTN3 genotype assignment was determined using the Invitrogen™ iBright™ CL1500 Imaging System according to the expected amplicon patterns.

### Table 1. Sequence of primers used for the DNA analysis.

| Gene   | Primers Sequence                     | T Annealing |
|--------|--------------------------------------|-------------|
| **COL1A1** | Forward, 5'-GATGTCCTAGGTGCTGGAGTTAG-3' Reverse, 5'-TGGTAGAGACAGGAGGAGGG-3' | 58°         |
| **ACE**  | Forward F1, 5'-CCCATTTCCTCATGAGGGCTTCTG-3' Forward F2, 5'-TGGGATTACAGCGCTGAT-3' Reverse R1, 5'-AGAGCTGGAAATAAAAATGGGC-3' | 55°         |
| **ACTN3** | Forward, 5'-GGGCACACTGTGCCCCTTTC-3' Reverse, 5'-GATGTCCCTGGCGGGCGGAG-3' | 61°         |
| **MCT1**  | Forward, 5'-AGACCAGTATAGATGTTGCTGGG-3' Reverse, 5'-CCACTGTTGAGATTACAGGCCA-3' | 58°         |
2.3. Statistical Analysis and Data Interpretation

In population genetics studies, Hardy-Weinberg equilibrium is used for allele and genotype frequencies. We used the Statistical Package for Social Science (SPSS) for Windows version 22.0 (Statistical Package for Social Science, Chicago, IL, USA) in order to verify the correlation between injuries and polymorphism analyzed. Pearson’s Chi-square ($\chi^2$) tests was performed to compare genotype and allele frequencies between athletes and controls. The unpaired t-test was applied to compare injured and non-injured players for various genotypes using the dominant model of each single gene. The $p$-value for significance was set at $p < 0.05$.

3. Results

In the athlete group, we related each genotype with the data injuries (Supplementary Table S1). We reported a total of 573 injuries as follows: 129 concerning muscle tissue, 247 to the bone tissue and 170 as tendon injuries. Fifteen athletes reported no injuries during their competitive career. Statistical analysis was conducted in order to verify genotype-injuries association. For the athletes’ group, we compared gene individually with total injury and with each injury type (bone, muscle and tendon/ligament). No statistical significance was found between injury incidence and $ACE$, $ACTN3$, $COL1A1$ genotype ($p > 0.05$). However, a statistical significance between $MCT1$ AA genotype with total injuries ($\chi^2 = 0.115; p = 0.003$) and bones injuries ($\chi^2 = 0.603; p = 0.007$) was found, showing a possible increased susceptibility to injury for athletes carrying the AA genotype. Thus, our results showed a correlation between the $MCT1$ AA genotype and sport injuries, particularly bone injuries, in rugby athletes. The Figure 2 shows the genotype distribution in the rugby athletes and in the controls.

![Figure 2](image_url)

Figure 2. The figure shows the genotype frequency for all genes analyzed expressed as a percentage for both groups (athletes and control population).

The genotype distribution in the athletes’ population resulted as follows: $ACE$ rs1799752 (30% DD, 49% ID, 21% II); $ACTN3$ rs1815739 (30% CC, 48% CT, 22% TT); $COL1A1$ rs1800012 (56% GG, 40% GT, 4% TT); $MCT1$ rs1049434 (33% AA, 56% AT, 11% TT). For the control population: $ACE$ rs1799752 (47% DD 38% ID, 15% II); $ACTN3$ rs1815739 (54% CC, 31% CT, 15% TT); $COL1A1$ rs1800012 (64% GG, 36% GT, 0% TT); $MCT1$ rs1049434 (27% AA, 59% AT, 14% TT).
No statistically significant difference was detected for the allele frequency between the two groups \( (p > 0.05) \). The genotype distribution was in Hardy-Weinberg equilibrium \( (p < 0.05) \).

4. Discussion

This study showed the potential susceptibility to different types of injuries (muscle, tendon and bone) in the court of 100 athletes of Lazio Rugby 1927 and G.S. Fiamme Oro Rugby. Biostatistical calculations were performed for all individual genes on both total lesions and individual stratifications (bone, muscle, and tendon). ACE, ACTN3, and COL1A1 genes showed no significant associations for any type of injury \( (p > 0.05) \). For MCT1 gene, we found no significant results between genotype and tendon-muscle injuries, while some studies reported an association between MCT1 AA genotype and the higher incidence of muscle disorders in sportive players [19–21]. In contrast, our results showed a statistical correlation of the MCT1 AA genotype with total injuries \( (\chi^2 = 0.115; p = 0.003) \) and with bones injuries \( (\chi^2 = 0.603; p = 0.007) \). We emphasize that our results could provide important molecular information about the possible protective role of MCT1 TT genotype with respect to certain types of injuries. In fact, during training anaerobic glycolysis increases the concentration of lactate at the muscle level. Lactate is the result of a clinical manifestation of body fatigue during high-intensity exercise. Lactate transport across the plasma membrane is primarily mediated by monocarboxylate receptor MCTs, including MCT1. MCT1 is ubiquitously expressed in lactate consuming cells and it is involved in transport of lactate from rapidly contracting muscle fibers [22]. Lactate accumulation could be considered one of the events contributing to the aggravation of physical injuries in athletes and it could be the cause of long periods of physical inactivity associated with significant psychological, physical, and economic consequences. MCT1 AA genotype is associated with faster lactate transport rate than MCT1 TT genotype. Thus, our study could help explain the role of increased susceptibility of MCT1 AA genotype with injuries, especially bone lesions. In support of our results, data from literature indicate that MCT1 is also involved in the osteoblast differentiation; particularly MCT1 appears to be a negative regulator of osteoclast differentiation of bone marrow macrophages (BMMs) [23]. MCT1 has been shown to mediate passive bone resorption induced by lactic acid released from the anaerobic metabolism of metastatic malignant cells [24]. In recent years, many studies have been conducted on athletes in order to establish an association between genotype and athletic performance. Research is mainly focused on identifying molecular markers related to physical performance that may provide useful information to identify athletes who have a higher predisposition to the risk of certain types of injuries (bone, muscle, or tendon injuries). Predictive genomic DNA profiling allows to understand whether a genetic advantage could lead to a more effective rehabilitation in some individuals than others [25].

Furthermore, medical and technical staff could develop personalized training programs to optimize the player’s athletic potential.

5. Conclusions

Recent data reported the association between MCT1 AA genotype and higher incidence of muscle injury in sports people [19], while our data indicated a strong association between MCT1 AA genotype and higher incidence of total and bone injuries in the group of 100 rugby players.

The study of the relationship between genetics related to injury phenotype seems to be increasingly useful in developing personalized training that allows athletes to focus on their genetic predispositions and achieve the best possible athletic performance.

Environmental elements such as training, lifestyle, and both physiological and psychological factors may cause an epigenetic change altering genes expression and affecting athletic performance.
Our future aim is to better clarify the association of genetics and sports by Next Generation Sequencing using multigene panels in a larger population of athletes.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/genes13060995/s1: Table S1. DNA genotyping and injuries collection in the athletes' group. Table 1 shows the sequence of Sanger homemade primers designed using “Primers 3 software” (http://primer3.ut.ee/, accessed on 1 January 2022). The folding characteristic of PCR products and primers were determined using the secondary structure profiling software DINAMelt (http://unafold.rna.albany.edu/?q=mfold, accessed on 1 January 2022). BLAST software was used in order to test the specificity of primers sequences (http://blast.ncbi.nlm.nih.gov/Blast.cgi, accessed on 1 January 2022). PCR was set up in 25 µL obtained by adding 12.5 µL of MasterMix 2X (Promega, Madison, WI, USA), containing Buffer (PH = 8.5), dNTPs (400 µM), Taq Polymerase (50 units/µL) and Mg+ (3 mM), 3 µL of genomic DNA as template (about 40–55 ng), 0.5 µL of primers (0.2 µM each) and ultrapure H2O until reaching the final volume. The PCR program was set on an initial denaturation at 95 °C for 2’, followed by 35 cycles of 95°C for 30”’, a step 30’’ of Annealing Temperature conforming to the homemade primers, 72 °C for 30’’ and a final extension at 72 °C for 5’. For ACE I/D variant, after the PCR step, total volume was loaded on a 4% agarose gel and subjected to electrophoresis for 30–35’. For ACTN3 variant, PCR amplification products were loaded on an electrophoresis 2% agarose gel to verify the PCR efficiency. Following RFLP was performed adding 15 µL of PCR product, 0.5 µL of Ddel Restriction Enzyme associated to 2 µL of Buffer D (Promega, Madison, WI, USA) and 0.5 µL of BSA. Total volume was incubated at 37 °C for 3 h. After electrophoresis of 30–35’ on a 4% agarose gel. In Table S1 we reported all results of genotyping and injuries data collection in the athletes group. The Figure 1 shows the overview of experimental design and samples workflow, from collection to statistical results. The Figure 2 represents the comparison of genotypic frequency between the athletes and the control groups. The abscissa represents the genotype investigated for each gene and the ordinate presents the genotype frequency in percentage found for the two different group.

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Institutional Review Board Statement: This study complied with the Ethical Principles for Medical Research Involving Human Subjects according to the Declaration of Helsinki of World Medical Association and it was certified by the Committee of the Applicable Institution of the Fondazione Policlinico Universitario Agostino Gemelli IRCCS, Rome (ID: 1858).

Informed Consent Statement: All participants involved in the study were identified and informed consent was obtained. Written informed consent has been obtained from the participants to publish this paper.

Data Availability Statement: The data presented in the study are available on request from the corresponding author due to restrictions (privacy).

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