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El Khoury, LY, Rickaby, R, Samiric, T & Raleigh, SM

Author post-print (accepted) deposited by Coventry University’s Repository

Original citation & hyperlink:
https://dx.doi.org/10.1016/j.jsams.2017.08.016

DOI 10.1016/j.jsams.2017.08.016
ISSN 1440-2440
ESSN 1878-1861

Publisher: Elsevier

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PII: S1440-2440(17)31021-6
DOI: http://dx.doi.org/10.1016/j.jsams.2017.08.016
Reference: JSAMS 1606

To appear in: Journal of Science and Medicine in Sport

Received date: 8-2-2017
Revised date: 3-8-2017
Accepted date: 22-8-2017

Please cite this article as: El Khoury Louis, Rickaby Rebecca, Samiric Tom, Raleigh Stuart M.Promoter methylation status of the TIMP2 and ADAMTS4 genes and patellar tendinopathy. Journal of Science and Medicine in Sport http://dx.doi.org/10.1016/j.jsams.2017.08.016

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Promoter methylation status of the *TIMP2* and *ADAMTS4* genes and patellar tendinopathy

Louis El Khoury¹,², Rebecca Rickaby¹, Tom Samiric³ and Stuart M Raleigh¹,

¹The Centre for Physical Activity and Chronic Disease, The Institute of Health and Wellbeing, University of Northampton, Northampton, UK, ²Department of Biochemistry and Molecular Biology, Mayo Clinic, Rochester, MN, USA ³School of Human Biosciences and the Musculoskeletal Research Centre, La Trobe University, Melbourne, Victoria, Australia and

Correspondence to: Dr S Raleigh, Division of Health and Life Sciences, University of Northampton, Park Campus, Boughton Green Road, Northampton NN2 7AL, UK; stuart.raleigh@northampton.ac.uk

Abstract

**Objectives** Patellar tendinopathy (PT) is a debilitating and prevalent condition that tends to affect those who are physically active or engaged in jumping sports. Although tendinopathies are known to have a genetic basis, the role of DNA methylation as an epigenetic factor and risk determinant for human PT has never been described.

We sought to determine whether differences existed between the methylation profiles of both the *TIMP2* and *ADAMTS4* gene promoter sequences in a cohort of males having undergone surgery for patellar tendinopathy compared to controls.

**Design** Case-control epigenetic study using DNA from 10 males with PT and 10 males with healthy tendons.

**Methods** We used PCR and targeted pyrosequencing to interrogate the methylation profiles of CpG sites upstream of both the *TIMP2* (4 sites) and *ADAMTS4* (6 sites) genes. We compared methylation differences between the two groups using t-tests.
**Results** We report no significant (p>0.05) methylation differences within the TIMP2 gene promoter between the PT group and controls across the 4 CpG sites investigated. In contrast, we detected a significant (p=0.016) difference in the methylation status of 1 CpG site, approximately 3 Kb upstream of the ADAMTS4 gene between the PT group and controls.

**Conclusions** To our knowledge, this is the first study to investigate how DNA methylation impacts on the risk of human tendinopathy. Our data indicate that the methylation status of the ADAMTS4 gene is altered in patellar tendinopathy and we speculate on how this change might modify the patellar tendon extra-cellular matrix environment.

**KEY WORDS:** Tendon, genetics, epigenetics, sports injury, tendinopathy

**Introduction**

Patellar tendinopathy (PT), also known as jumper’s knee, is an activity-dependant chronic condition characterised by anterior pain to the knee\(^1\). Injuries to the patellar tendon are common\(^2,3\). They account for nearly 30% of knee injuries in US high school athletes\(^4\) and are prevalent amongst volleyball (45%) and basketball (32%) players alike\(^1\). PT is also found in military cohorts and the injury can lead to prolonged periods of disability\(^5\). Both tendon neovascularisation and thickening are associated with PT injuries\(^6,7\). Furthermore, some authors report the injury to associate with an increase in collagen type III fibres along with elevated levels of glycosaminoglycans\(^8\). A number of risk factors for PT have been tentatively reported in the literature such as increased weight and BMI\(^9\) and there is a growing body of evidence to suggest that genetics can play a role in the predisposition to human tendinopathy\(^9,10\). However, to date, and to our knowledge, no single study has
investigated the role of human DNA methylation (as an epigenetic factor) in sports injury phenotypes like PT.

DNA methylation occurs when a methyl group binds to a cytosine nucleotide base converting it to 5-methylcytosine\(^1\). This takes place predominantly in CpG rich regions (CpG islands) which are typically located in the promoter regions of genes. DNA methylation tends to down regulate gene expression by interfering with the accessibility of genes to transcription factors\(^2\). Hence, methylation can profoundly affect transcription levels and an epigenetic mode of inheritance might significantly modify the effect of genotype upon phenotype\(^\_\). Furthermore, methylation results from either directionally programmed changes, part of the ageing process, or from stochastic and spontaneous alterations attributed to environmental factors \(^3,4\). It is difficult to predict, as of yet, the reason why and at which rate some non-age-associated CpG sites undergo changes in methylation.

The \textit{TIMP2} (tissue inhibitor of metalloproteinase 2) gene encodes a protein that has an important role in extra-cellular matrix (ECM) homeostasis as it degrades a number of metalloproteinases (MMPs)\(^5\). Recently, a single nucleotide polymorphism (SNP) within the \textit{TIMP2} promoter (rs4789932) was found to associate with Achilles tendon pathology (ATP)\(^6\) where \textit{TIMP2} expression levels are altered. For example, Jones and colleagues (2006) reported a decrease in \textit{TIMP2} expression in tendinopathic tissue compared to healthy tendons\(^7\) while Karousou et al. (2010) reported the opposite effect\(^8\). Interestingly, \textit{TIMP2} expression is under tight epigenetic control and hypermethylation has been shown to switch off transcription\(^9\). However, the effect of \textit{TIMP2} methylation on the risk of PT has not been investigated.

The \textit{ADAMTS4} (A disintegrin and metalloproteinase with thrombospondin motifs 4) gene encodes a protein that degrades aggregcan and a number of other components of the ECM\(^10\). \textit{ADAMTS4} expression has been found to increase in knee osteoarthritis and this might
exacerbate the condition by contributing to the elevated degradation of proteoglycans\(^{19}\). 

*ADAMTS4* mRNA expression levels have been shown to differ in ruptured Achilles tendons compared with normal and tendinopathic tendons\(^{20}\). Furthermore, in an equine tendon model *ADAMTS4* expression level decreased in response to surgical transection. The decrease paralleled an increase in aggregan, versican and lumican\(^{21}\). Hence, the expression of *ADAMTS4* likely plays an important role in the tendinopathic process but, as yet, our understanding of how *ADAMTS4* expression is controlled in tendinopathy is unclear. Furthermore, like *TIMP2*, *ADAMTS4* is under epigenetic control and loss of methylation has been shown to be responsible for the upregulation of *ADAMTS4* protein in osteoarthritic cartilage\(^{22}\).

It is known that an individual’s methylation profile may change in response to physical activity\(^{23}\). However, somewhat surprisingly, the role that DNA methylation plays, as an epigenetic factor, has never been investigated in relation to human tendinopathy. With this in mind, the aims of the present study were to establish, for the first time, whether epigenetics, in particular DNA methylation, was changed in tissue isolated from patellar tendinopathy compared to healthy tissue. Our study specifically focused on the DNA methylation statuses of the *TIMP2* and *ADAMTS4* genes in a targeted approach as both, as discussed above, undergo changes in expression during tendinopathy but the mechanisms underlying these changes are unknown.

**Material and methods**

Patellar tendon tissue was obtained from 10 male participants with healthy patellar tendons (CON) undergoing ACL reconstruction surgeries using a patellar tendon graft. We also obtained 10 patellar tendinopathy (PT) samples from males selected for surgical debridement for recalcitrant overuse patellar tendinopathy. All tissue was immediately stored at \(-80^\circ\text{C}\) until processing. The DNA samples used in this study were collected from
participants recruited from 2003-2005 that were originally described elsewhere in studies focusing only on proteoglycan and RNA levels\textsuperscript{24}. Participant information was obtained from surgery theatre lists and was limited to age, sex and ethnicity. Unfortunately, additional demographic information such as height, weight, BMI and other relevant medical history were not made available to us. Hence our cohort should be considered as a convenience sample. All participants were otherwise healthy Caucasians aged from 19 to 41 years and gave written informed consent. Approval was obtained from the University of Northampton’s School of Health Research Ethics Committee and La Trobe University Human Ethics Committee (Application number 12-086).

DNA was obtained from all patellar samples using the PureLink® Genomic DNA Kit (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s recommendations. The DNA quantity was determined using a NanoDrop 2000 (Thermo Fisher Scientific, Wilmington, DE). The DNA samples (50 ng), in addition to a universal non-methylated control DNA sample from the EpiTect® PCR Control DNA Set (Qiagen, Hilden, Germany), were bisulfite treated using the EpiTect® fast DNA bisulphite Kit (Qiagen, Hilden, Germany) according to the manufacturer’s recommendation. The Pyrosequencing assays for the \textit{TIMP2} (Hs\_TIMP2\_01\_PM) and \textit{ADAMTS4} (Hs\_NDUFS\_01\_PM) genes used in this study were selected from the pre-designed PyroMark® CpG Assays (Qiagen, Hilden, Germany). The single assay designed for the CpG island within the \textit{TIMP2} promoter region measured methylation at 4 CpG sites. (-488 bp, -478 bp, -467 bp, and -455 bp from the first exon). The assay chosen for the CpG sites within the promoter region of the \textit{ADAMTS4} gene was selected to cover a wider area of the promoter region and spanned 6 CpG sites that were -2964 bp, -2973 bp, -2995 bp, -2998 bp, -3000 bp and -3002 bp from the first exon. We took a pragmatic approach to the choice of CpG sites that we investigated. In some cases it was not possible to study certain regions of the promoters as sequencing assays did not meet quality control standards. However, as \textit{ADAMTS4} is regulated by a large promoter
sequence, we were especially interested to determine how methylation of CpG sites that are distant to the transition start site impacted on injury risk.

All bisulfite converted DNA samples in addition to hypermethylated and hypomethylated controls were PCR amplified using the pre-designed primer sets of the PyroMark® CpG Assays (Qiagen, Hilden, Germany). A Techne TC-512 thermocycler (Bibby Scientific Ltd, Staffordshire, UK) was used to amplify the target sequences according to the manufacturer protocol. To confirm successful PCR amplification the products were run on a 1x agarose gel alongside a GeneRuler™ 100 bp ladder (MBI Fermentas, UK). The biotinylated PCR products were then immobilised to Streptavidin-coated Sepharose high-performance beads (GE Healthcare, Buckinghamshire, UK) according to the PyroMark® Q24 manufacturer’s recommendations (Qiagen, Hilden, Germany). The immobilised PCR products were captured using the PyroMark® Q24 vacuum (Qiagen, Hilden, Germany) and released into a 24 well PyroMark® Q24 plate containing 1x sequencing primer diluted in 25 μl PyroMark® Annealing Buffer. Before placing the Q24 plate into the pyrosequencer, the sequencing primer was allowed to anneal on a preheated hot-plate at 80 °C for 2 min followed by room temperature incubation for 10 min. Following pyrosequencing, pyrograms were visualised and percentage methylation of the CpG sites was determined using the PyroMark® Q24 v2.0.6 software. Data analysis was performed using SPSS Version 20 program (IBM Corp. Armonk, NY). A t-test was used to compare the mean percentage methylation of each CpG site between the CON and the PT samples for both the TIMP2 and ADAMTS4 CpG sites. Significant differences in methylation were called when p<0.05. Correction for multiple testing was not conducted for reasons previously described by Posthumus et al (2011)

Results

PCR products for the TIMP2 and ADAMTS4 promoter sequences investigated were detected as single bands with the expected amplicon lengths (154 bp and 244 bp respectively) as shown in figure 1. Typical pyrograms obtained showing percent methylation
at the 4 CpG sites for *TIMP2* (panels A and B) and the 6 CpG sites for *ADAMTS4* (panels C and D) can be seen in figure 2. In our cohort the CON and PT groups were similarly matched for age (p=0.449) and gender. For *TIMP2* we found no significant difference (p=0.885) in the mean methylation percentages across all 4 sites between the CON and the PT groups. Furthermore, there was no significant difference in methylation for any of the 4 CpG sites when analysed independently between the CON and the PT groups as shown in figure 3. However, for the *ADAMTS4* gene, CpG site 3 was significantly (P=0.016) more methylated in the PT group compared to the CON group (figure 3). Further data analysis showed no significant difference between CON and PT in the remaining 5 sites when analysed independently (figure 3), or for the mean percentages of methylation across the 6 sites (p=0.294).

**Discussion**

To our knowledge, this is the first study to investigate DNA methylation differences in clinically resected human tendinopathy samples compared to healthy tendon tissue. Unlike the recent study of Trella and co-workers\(^{27}\) who used a genome wide design in a mouse tendinopathy study, we employed a hypothesis-driven, targeted approach to study methylation profiles of two genes (*TIMP2* and *ADAMTS4*) that have been associated with the tendinopathic process in human\(^{15,20}\).

*TIMP2* is known to be under epigenetic control with a CpG island in the promoter region of the gene extending from the transcription initiation site upstream for approximately 900 bp\(^{18}\). The methylation of the CpG sites located between positions -295 and -145 bp was found to reduce the expression of *TIMP2* in prostate cancer samples\(^{18}\). Likewise, the overlapping region ranging from -350 to -240 bp was also found to be over-methylated in cervical cancer\(^{28}\). However, our work showed that there were no methylation changes within the CpG Island ranging from position -488 to -455 between the CON and PT tissue samples.
Moreover, all analysed samples showed very low methylation levels which might infer that this region of the TIMP2 promoter is not involved in regulating TIMP2 expression or deregulation during the process of patellar tendinopathy. Furthermore, the lack of change in TIMP2 methylation status between PT tissue and controls shown here is consistent with the absence of TIMP2 expression level changes between PT and control tissue previously shown by Parkinson et al (2010)\textsuperscript{24}. The fact that we found no difference in TIMP2 methylation status between PT tissue and controls it interesting. This might suggest that previously observed differences in TIMP2 expression in Achilles tendinopathy\textsuperscript{17} are not an active part of the tendinopathic process but possibly no more than an artefact of the injury. Of course, we only interrogated a small number of CpG sites for TIMP2 in this study and investigating a larger area might well have revealed significant differences between the PT and CON samples.

Of the 6 CpG sites investigated 3 kb upstream of the ADAMTS4 gene we found one site, located at position -2995, that was significantly (p=0.016) more methylated in the PT group compared to the CON group. Although only one of the 6 CpG sites was more methylated in the tendinopathic tissue compared to controls, the impact of this when considering the risk of patellar tendinopathy might be far reaching. To be specific, the ADAMTS4 gene is known to be under epigenetic control\textsuperscript{22} and regulated by a large promoter (-4109 to +406) that consists of two NFAT and several Runx binding elements\textsuperscript{25}. One of the NFAT binding regions resides only 103 bp away from the CpG site that we investigated and it is plausible that a change in methylation status at that site, as measured by us, could affect the binding of NFAT transcription factors and alter the expression of ADAMTS4. Interestingly, an increase in the accumulation of proteoglycans including aggrecan was found previously in PT tissue taken from the participants in this study\textsuperscript{24} and others\textsuperscript{29}. However, somewhat surprisingly, ADAMTS4 levels in this patient cohort did not differ compared to controls\textsuperscript{24}. Recently, the expression of ADAMTS4 was shown to decrease, in a region specific manner, in an ovine model of tendinopathy\textsuperscript{30}. The reduction in ADAMTS4 expression was found to be
commensurate with an increase in aggrecan\textsuperscript{30}. Therefore, we speculate that in some situations the accumulation of proteoglycans such as aggrecan may be explained, at least in part, by reduced levels of ADAMTS4 caused by hypermethylation at CpG sites within the ADAMTS4 gene promoter. It is also worth noting that methylation change at single CpG sites can have profound effects on phenotypes. For example, Fürst and colleagues\textsuperscript{31} have observed this phenomenon with respect to the expression of estrogen receptor alpha (ESR1) and methylation changes at single sites upstream of the SLC23A2 and NCOR2 genes have been shown to influence the severity of spinal muscular atrophy\textsuperscript{32}.

Although our study was the first, to our knowledge, to report DNA methylation changes associated with human tendinopathy there were some limitations. Firstly, although our participants were all male with no significant age differences between the PT cases and controls we were unable to control for height, weight or level of physical activity in our dataset as this information was not available to us. The acquisition of such data in future experiments would improve confidence in the interpretation of the results. Secondly, our study was limited to a small number of clinical samples that were available from Caucasian men. Therefore, the changes we observed may not represent methylation changes that might occur in other populations. Hence, this work should be validated in larger, non-Caucasian cohorts and in female patients. Finally, this study has revealed that an over-methylation event in the promoter sequence of the ADAMTS4 gene associates with patellar tendinopathy. The hypermethylated CG site sits close to an important DNA regulatory element and might influence risk of tendinopathy by modifying ADAMTS4 levels leading to structural or regulatory changes within the ECM components of the tendon. Such changes might predispose an individual to develop the injury.

Conclusions
We provide the first evidence that DNA methylation (an epigenetic factor) within the promoter sequence of the human *ADAMTS4* gene might be important in patellar tendinopathy. Our data have clear implications as set out below:

**Practical implications**

- As DNA methylation changes are modified by the environment it might be possible, in the future, to alter methylation status, through adaptations to an athlete’s training programme, to reduce the risk of tendinopathy.

- DNA methylation changes might well explain discrepant gene association findings that have been observed in some studies on human tendinopathy.

- DNA methylation, as an epigenetic factor, should be considered as a potential modifier of genotype and other biochemical factors when considering risk of tendinopathy.

**Acknowledgements**

This study was funded by grants from the Rosetrees Trust (M183) and the University of Northampton. The authors would also like to acknowledge Mr Julian Feller (Consultant Orthopaedic Surgeon) and his team for the provision of the samples.

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FIGURE 1. Agarose gel (1%) electrophoresis images of the TIMP2 and ADAMTS4 gene promoter sequences investigated. Panel A (TIMP2): Well 1, 100 bp ladder; well 2, no DNA (negative control); well 3 and 4, control samples; wells 5 and 6, DNA from PT; well 7, hypermethylated control; well 8, hypomethylated control; well 9, no DNA (negative control) and well 10 bisulphite converted hypomethylated control. Panel B (ADAMTS4): Well 1, 100 bp ladder; wells 2 and 4, control DNA samples; wells 5 to 7, DNA from PT; well 8, hypermethylated control; well 9, hypomethyated control; well 10, bisulfite converted hypomethylated control and well 11, no DNA (negative control). Gels were run as described in the Materials and Methods section.

FIGURE 2. Typical pyrograms showing the methylation status of the TIMP2 and ADAMTS4 gene promoter sequences investigated from both control and PT patients. Panel A: Percent methylation at each of the four CpG sites within the TIMP2 gene promoter from healthy tendon. Panel B: Percent methylation at each of the four CpG sites within the TIMP2 gene promoter from a patient with patellar tendinopathy. Panel C: Percent methylation at each of the six CpG sites within the ADAMTS4 gene promoter from healthy tendon. Panel D: Percent methylation at each of the six CpG sites within the ADAMTS4 gene promoter from a patient with patellar tendinopathy. In each figure the grey boxes indicate the percentage of each CpG site that is methylated in the sample that is run.

FIGURE 3. Bar graphs showing the mean (± SD) percent methylation status at each of the CpG sites investigated within the TIMP2 (Panel A) and ADAMTS4 (Panel B) gene promoter sequences. For both genes the CpG sites for healthy tendons are shown as white bars and those for tendinopathic tendons are shown as black bars. *p=0.016.
FIGURE 1 Panel A

FIGURE 1 Panel B
FIGURE 2 Panel A

FIGURE 2 Panel B
FIGURE 2 Panel C

A1: GTATGAGCGYGTTAGGGGAAAGGGAGTAGGGAAAGGAGYGTGGYGT

FIGURE 2 Panel D

B4: GTATGAGCGYGTTAGGGGAAAGGGAGTAGGGAAAGGAGYGTGGYGT
FIGURE 3

A. *TIMP2*

B. *ADAMTS4*