An increase in copy number of Myosin Light Chain Kinase 1 associates with increased force production in Lithuanian athletes

By

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Declaration

I declare that this thesis and the work presented in it are my own and have been generated by me as the result of my research. It has not been submitted anywhere else for any award. Where other sources of information have been used they have been clearly acknowledged and referenced.
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I would like to dedicate this thesis to my mum who made me the man I am today and who I miss every day.
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Abstract

Copy number variation has been linked to the development of various diseases, however copy number variants have not been investigated for associations with athletic performance. The present study investigated copy number variation of the Myosin Light Chain Kinase (MYLK)1 and MYLK-Antisense 1 (MYLK-AS1) genes for associations with athletic performance in a cohort of Lithuanian athletes and controls. We hypothesised that increased MYLK1 copy number and decreased MYLK-AS1 copy number would be associated with strength and power athletes (STP) and measures of muscle performance. DNA was extracted from blood samples of 407 athletes and controls. Copy number of the target genes was determined using a multiplexed quantitative-polymerase chain reaction and the use of a multicopy reference assay. A higher MYLK1 copy number was overrepresented in the STP compared with controls (p=0.028[OR=9.97, 95% CI: 1.19-83.81]); however endurance athletes did not differ from controls or STP (p≥0.66; \(\chi^2\leq0.829\)). Positive correlations between MYLK1 copy number and strength and power performance were detected. Individuals with three copies were able to produce more power in isokinetic tests, jump higher and sprint faster than individuals with two copies in the whole cohort and these correlations were stronger in the STP. The strongest correlations found were with MYLK copy number and isokinetic flexion of the arm in STP (p=0.003, \(R^2=7.78\)). MYLK-AS1 did not associate with athlete status or measures of athletic performance (p>0.05). The improvements in performance with increased copy number indicate that copy number variants are associated with improved skeletal muscle phenotypes. These results, further current knowledge of how genetic variants underpin muscle phenotypes and indicate that MYLK1 potentially could be used as a target for improvements in athletic performance and treatment of muscle wasting disorders.
## Abbreviations

<table>
<thead>
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<tbody>
<tr>
<td>Deoxyribonucleic acid</td>
<td>DNA</td>
</tr>
<tr>
<td>Ribonucleic acid</td>
<td>RNA</td>
</tr>
<tr>
<td>Monozygotic</td>
<td>MZ</td>
</tr>
<tr>
<td>Dizygotic</td>
<td>DZ</td>
</tr>
<tr>
<td>Single nucleotide polymorphism</td>
<td>SNP</td>
</tr>
<tr>
<td>α-Actinin-3</td>
<td>ACTN3</td>
</tr>
<tr>
<td>Structural variant</td>
<td>SV</td>
</tr>
<tr>
<td>Polymerase chain reaction</td>
<td>PCR</td>
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<td>Fluorescence in situ hybridization</td>
<td>FISH</td>
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<tr>
<td>Quantitative PCR</td>
<td>qPCR</td>
</tr>
<tr>
<td>Copy number variant</td>
<td>CNV</td>
</tr>
<tr>
<td>Salivary amylase</td>
<td>AMY</td>
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<td>Survival of motor neuron</td>
<td>SMN</td>
</tr>
<tr>
<td>Myosin light chain kinase (gene)</td>
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</tr>
<tr>
<td>Regulatory light chain of myosin</td>
<td>RLC</td>
</tr>
<tr>
<td>Myosin light chain kinase (protein)</td>
<td>MLCK</td>
</tr>
<tr>
<td>Smooth muscle MLCK</td>
<td>smMLCK</td>
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</table>
Skeletal muscle MLCK  
Cardiac muscle MLCK  
MYLK-antisense1  
Strength and power athlete  
Endurance athlete  
Crossing point  
Odds Ratio  
Benjamini-Hochberg  
Post-activation Potentiation  

skMLCK  
cmMLCK  
MYLK-AS1  
STP  
END  
Cp  
OR  
BH  
PAP
Chapter 1 - Introduction

1.1 Background

Skeletal muscle is the largest organ in the human body making up approximately 30% of body mass (Kostek et al., 2010; Pescatello et al., 2013). The amount of force that can be generated (i.e. muscle strength), is the primary measure of skeletal muscle function and is often used as an indicator of health. The functional capacity of skeletal muscle is of critical importance because the maintenance of strength is essential in the prevention and treatment of muscle wasting diseases, such as sarcopenia (Roth et al., 2000; Roubenoff, 2000) and metabolic disorders, such as type II Diabetes (Boulé et al., 2001). Skeletal muscle underpins other processes such as locomotion and movement (Kostek et al., 2010). Since skeletal muscle is critically important for force production and movement, it is clear that the quality of muscle is a regulator of both sporting performance and health. The age related decline in both muscle mass and strength has been termed sarcopenia (Roth et al., 2000; Roubenoff, 2000; Metter et al., 2002; Goodpaster et al., 2006; Mitchell et al., 2012). Studies have shown the steady loss of skeletal muscle begins at 40-50 years of age and the rate of loss is of such a degree that by the age of 80 the typical loss of skeletal muscle fibres from the limbs is 50% (Faulkner et al., 2007; Mitchell et al., 2012). Despite the maintenance of a high level of physical fitness, the performance of ‘master’ athletes typically declines after the age of 40 and by the age of 80 can be reduced by up to 65% (Faulkner et al., 2007). However, ‘master’ athletes are stronger than master non-athletes of the same age (Hawkins et al., 2003). Muscle strength is lost at a greater rate than muscle mass(Faulkner et al., 2007; Mitchell et al., 2012). Despite a strong correlation between muscle mass and strength, in older individuals gains in lean mass can occur with decrements in muscle strength, indicating that the quality of the muscle is the important factor (Goodpaster et al., 2006; Mitchell et al., 2012). Sarcopenia is
an important process to understand because it can lead to disability, falls and the loss of independence in otherwise healthy elderly individuals (Roubenoff, 2000; Metter et al., 2002; Goodpaster et al., 2006; Mitchell et al., 2012). It has been shown that individuals with higher muscle strength, in large muscle groups, have a greater survival rate from a range of diseases compared to individuals with lower muscle strength (Metter et al., 2002; Goodpaster et al., 2006; Ruiz et al., 2009). Thus, understanding the determinants of muscle strength is an important area of scientific research from both a health and performance perspective.

Muscle strength is a highly variable trait and at the two opposing ends of the muscle strength scale there are elite athletes and individuals who suffer from muscle wasting diseases. Various factors determine muscle strength including environmental factors, such as nutrition, level of physical activity (Tiainen et al., 2009), and behavioural factors, such as motivation to train and level of pain tolerance (Sallis et al., 1992; Boutelle et al., 2004). Physiological factors also have been shown to influence strength through a myriad of different ways such as fibre composition, cross sectional area and size of the muscles (Kostek et al., 2010). There is a significant genetic component to these behavioural and physiological factors. The interaction between the environment and the genetics of an individual are just as important in phenotype determination as either factor alone (Tiainen et al., 2009). Individuals who possess advantageous genetics and control their environment correctly have the greatest chance of developing into elite athletes. Therefore, muscle strength is a biological process which is determined which is partially determined by an individual’s genetics and supported by both the environment and the interaction between the genetics and the environment (Kostek et al., 2010).
The human haploid genome is encoded on deoxyribonucleic acid (DNA) sequences which consist of approximately 3 billion nucleotides (Kruglyak & Nickerson, 2001). DNA encodes 20,687 protein coding genes in all individuals (Bernstein et al., 2012). The sequence of any two human genomes are highly similar with estimates of over 99% identical, which leaves 3 million potential nucleotides that determine the large phenotypic differences both within and across different populations (Kruglyak & Nickerson, 2001). The variation in the sequence of DNA causes the genetic differences in phenotypes. If a genomic variant is found at greater than 1% within a population then it is termed a common variant, whereas if the variant is found at less than 1% then it is known as a rare variant. Typically rare variants have a large impact on the function of a gene and are consequently less tolerable; therefore reducing the likelihood of them being passed on to future generations. Common variants tend to be more tolerable and have small to no effect on a gene, thus giving them a higher chance to be passed on to future generations than mutations (Pollex & Hegele, 2007).

Less than 2% of the human genome encodes for protein coding genes (Bernstein et al., 2012), and at least 5% of the genome has been shown to be under selection. Therefore it was thought regions other than protein coding regions were functional (Pheasant & Mattick, 2007). The remaining 95% of the genome was referred to as ‘junk’ DNA because it had no obvious functional role. The completion of ENCODE project discovered that the majority of the ‘junk’ DNA has some form of biochemical function. A total of 80% of the genome has been shown to have some form of functional role on either RNA or chromatin (Bernstein et al., 2012). Most of this percentage acts post transcriptionally to modify coding RNAs to alter regulation or expression of coding RNA (Palazzo & Gregory, 2014). One such type of non-coding gene which acts to control the level of mRNA is an antisense gene. Antisense transcripts have been shown to regulate the expression of sense transcripts through RNA-
RNA interactions. These interactions can cause gene silencing through chromatin modifications, such as DNA methylation and histone modification or interference where the sense and antisense transcripts hybridize and are subsequently destroyed (Werner, 2013).

1.2 Heritability of traits

Heritability is defined as the proportion of the variability of a phenotype which is explained by genetic factors within a population (Visscher et al., 2008). Twin and family studies are the two main study designs used to detect the heritable component of traits associated with athletic performance (Beunen & Thomis, 2006). Family studies examine the similarity between siblings and their parents to estimate the variation in the genetic component. These studies typically overestimate the genetic component due to conflation with shared environment. Twin studies examine monozygotic (MZ) and dizygotic (DZ) twins to determine the genetic and environmental impact of a trait. MZ twins have nearly identical genomes and DZ twins only share ~50% of their genetic material (Shih et al., 2004). If the phenotypes of MZ twins are more similar then it is said there is a genetic component, whereas if the phenotype of DZ twins is equally similar then it is said there is a purely environmental impact on the heritability of the trait.

Twin and family studies only show the heritability of a trait within the specific population which is being examined because factors such as race, sex and age can all affect the heritability differently depending on the population (Visscher et al., 2008). When the heritability of height is examined, the population specific aspect of heritability is highlighted. In Caucasian populations heritability of height in males has been shown to range from 87-93%, whereas in females the heritability is estimated to be 68-84% (Silventoinen et al.,...
2003). The heritability of height also has been examined in a range of different populations of African descent. In Nigerians the heritability of height is estimated at 62%, whereas in African Americans the heritability is estimated to be 87%, while in Jamaicans the heritability is 76% (Luke et al., 2001). The age of the individuals examined influences heritability. Typically the heritability of a trait will change over the course of a lifetime because of different environmental impacts at certain points in life. For example body mass at birth is dependent on the uterine environment, as an infant body mass is determined by the maternal milk production and as an adult there is no maternal component which influences body mass (Visscher et al., 2008). Twin and family studies indicate there are several aspects of athletic performance which have been shown to have large heritable aspects associated with them.

The heritability of aerobic performance is one of the most studied athletic performance traits. The early studies on heritability generally over or under estimated the genetic component because of a lack of control for bias such as, age or gender (Bouchard et al., 1986). The HERITAGE study is one of the recent studies which controlled for several factors (age, sex and body mass) and predicted the heritability of $\dot{V}O_2$ max and the trainability of $\dot{V}O_2$ max was approximately 50% in sedentary individuals (Bouchard et al., 1998, 1999). Several other traits of athletic performance have been studied including muscle fibre composition, anaerobic performance and muscle strength and power. More recent twin studies have estimated that the genetic element of an individual’s athletic performance is 31-85% (De Moor et al., 2007).

The heritability of muscle strength traits is not as clear because in the literature there are a wide range of heritability estimates. Twin and family studies have shown there is a large
heritable factor associated with muscle strength; however, inconsistencies in study design have led to wide estimates in the heritability estimates of muscle strength. Some studies show the upper heritability estimates to be 96% (Huygens et al., 2004b), whereas other studies predict heritability of strength to be as low as 14% (Beunen & Thomis, 2006). One reason for the lack of clarity in the hereditary estimates of muscle strength is that there are a lot of different measureable aspects of strength which are not consistent in the literature (Thomis et al., 1998). Elbow extension and flexion has been shown to produce a higher heritability estimate than knee extension and flexion (Huygens et al., 2004b, 2004a). Despite the wide range in heritability estimates of muscle strength there is a clear genetic component which explains the variance in the phenotype (Costa et al., 2012). Unfortunately heritability studies do not actually inform about the causal genetic variant (Pitsiladis et al., 2013). To allow the identification of the causal variant which underlie the heritability further studies, such as gene-association studies, must be conducted.

1.3 Genetic variation

The most commonly studied form of genetic variations studied are single nucleotide polymorphisms (SNPs). SNPs occur when one nucleotide is exchanged for another in the sequence of a gene. SNPs can occur in protein-coding exons, non-coding intronic regions of genes and intergenic regions (Barreiro et al., 2008). The majority of SNPs fall within the introns of genes and thus typically do not have an effect on the function of the gene and are typically viewed as silent. There are occasions when intronic SNPs affect the splicing or expression of a gene and these SNPs may have an effect on the function of a gene (Choi et al., 2008; Millar et al., 2010). When a SNP falls within an exon it can be further classified as either non-synonymous or synonymous. Synonymous SNPs do not cause a change in the
amino acid and typically are silent or cause differential splicing of the gene. Non-
synonymous SNPs lead to a change in amino acid and typically have the largest effects
because they can either cause premature truncation of the protein or change the class of the
amino acid potentially reducing the stability of the protein (Pollex & Hegele, 2007). For
example, if a SNP causes a change from a hydrophobic amino acid to a hydrophilic amino
acid, this can influence the stability of the protein depending on the cellular location (Strub et
al., 2004). The phenotypic consequence of a SNP will ultimately depend on the function of
the gene that is affected. For example, a SNP variation in a gene that is functionally involved
in the muscle contraction process may affect performance phenotypes.

The R577X polymorphism of the α-Actinin-3 (ACTN3) gene is one of the most
studied and well documented which is associated with skeletal muscle phenotypes (for a
detailed review see Berman & North, 2010). The X allele leads to a loss of function of
ACTN3 that is associated with a reduced performance in power and sprint activities (Moran
et al., 2007; Walsh et al., 2008). A study of elite athletes found a reduction of the X allele in
sprint athletes and the complete absence of the X allele in Olympic athletes (Yang et al.,
2003). Several other SNPs have been associated with muscle size, strength and power
phenotypes (for a detailed review see Ahmetov & Fedotovskaya (2012)). One of the key
studies which has driven associations between genotypes and muscle phenotypes is the
FAMuSS study (Thompson et al., 2004). The FAMuSS study recruited a total of 1300
individuals and genotyped them for around 500 polymorphisms and found associations with
several genetic loci and physical activity levels and muscle size, strength and body
composition at baseline and in response to resistance training (Pescatello et al., 2013).
Although SNPs are the most commonly studied class of genetic variant, they are not the only genetic variants that have been associated with functional consequences on phenotypes. The total heritability estimates of muscle performance traits have not been accounted for by SNP variation. Therefore, other classes of genetic variation, such as structural variants, may explain some of the missing heritability of muscle performance traits.

1.4 Structural Variants

Structural variation occurs when insertions, duplications or deletions of genetic material occur. The nature of a structural variant (SV) means it can not only impact several bases of a gene, but also several genes. Structural variants of the human genome can take the form of small deletions or insertions which impact up to 50 base pairs, through to large structural variants which can affect whole chromosomes (Boonpeng & Yusoff, 2013). The initial studies to detect genomic variation examined large structural variants because these studies were performed before the polymerase chain reaction (PCR) was invented. Without the technology to amplify the desired sequence of a gene, investigators relied on using microscope karyotyping and southern blot technology, which are limited to identifying large variants which affect large segments or whole chromosomes (Pollex & Hegele, 2007). The invention of PCR allowed the easy amplification of target gene sequences to identify SNPs, therefore the majority of studies moved into examining SNPs.

Several methods exist to identify SVs, each with associated advantages and disadvantages. These methods can be categorised into two main approaches, genome-wide scans for global identification of SVs and targeted scans of candidate genes. For genome-wide identification of SVs, the development of array based technology led to array-
Comparative Genomic Hybridization and SNP array based methods for detection of SVs. These array based methods use probes to compare the intensity of DNA samples to identify gains or losses of genetic material (Boonpeng & Yusoff, 2013). For the targeted scans approach, the traditional methods used are Southern blotting and fluorescence in situ hybridization (FISH) (Pollex & Hegele, 2007). Both southern blots and FISH are infrequently used because of the drawbacks associated with each of the methods. Southern blots require a high quantity of DNA and is a laborious process, while FISH is not suited to identifying a partial deletion of genes (Hoebeeck et al., 2007), because of resolution limitations (Boonpeng & Yusoff, 2013). The most common and simplest method for SV detection is to use quantitative PCR (qPCR) to compare the fluorescence of a target sequence compared to a reference assay (Boonpeng & Yusoff, 2013). One issue with qPCR based SV detection is the use of a single gene as the reference assay (e.g. RNASE P or TERT) because SVs could affect the assay which can lead to the incorrect determination of copy number of the target gene (Table 1).

<table>
<thead>
<tr>
<th>Reference Assay Copy Number</th>
<th>Target Gene Copy Number</th>
<th>Calculated Copy Number of Target</th>
<th>Correct copy Number for Target Gene</th>
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<tr>
<td>3</td>
<td>2</td>
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1.5 Copy number variation

Generally it is accepted that individuals have two copies of every gene, one inherited from their mother and one inherited from their father; however, it has been discovered that
the variation in copy number of a gene because of duplications and deletions of genetic material is widespread (Yang et al., 2013). Advances in genotyping and the human genome project allowed the investigation of gene copy numbers at a subchromosomal level (Lobo, 2008). A SV of DNA of 1000 nucleotides or more which differs in copy number from a reference genome is classified as a copy number variant (CNV) (Redon et al., 2006). In 2004, the first papers to identify the widespread extent of CNVs throughout the human genome were published (Sebat et al., 2004; Iafrate et al., 2004). Both of these studies identified approximately 250 regions of copy number losses or gains in two different cohorts of unrelated individuals. A larger study of 270 individuals identified 1447 CNV regions throughout the human genome which affected 360 mega bases of DNA; equivalent to 12% of the total human genome and between 6-19% of any chromosome (Redon et al., 2006). A more recent study identified that on average 5% of the genome is affected by CNVs (Conrad et al., 2010). These CNVs can either be inherited from parents or rise from de novo mutations, where both parents have a normal copy number at the region of DNA in question. Studies have shown that monozygotic twins can differ in copy number for genes (Hastings et al., 2009) and the same individuals can have a different copy number depending on the tissue examined (Piotrowski et al., 2008).

With the vast distribution of CNVs throughout the genome in a range of different organisms it is expected that they will contribute to the phenotypic differences between individuals. The main way CNVs will influence a phenotype will be through an alteration of gene dosage, causing a disturbance in coding sequences or disturbing the long-range regulation of genes through alteration of chromatin environment (Stranger et al., 2009). Between 85-95% of CNVs examined have been associated with changes in the expression of the affected genes (Stranger et al., 2009; Henrichsen et al., 2009). Deviation in gene copy
number associates with changes in the amount of the final protein levels. The proteins which do not show altered product tend to be components of complexes (Tang & Amon, 2013). On dosage sensitive genes, which have no compensatory mechanism, the presence of a duplication or deletion will directly increase or decrease the expression of the gene, respectively (Pollex & Hegele, 2007). When gene duplications occur there are several consequences for gene expression depending on the inclusion or absence of regulatory elements. If complete duplication of a gene including the regulatory region occurs there will be a greater increase in expression than if partial duplication of a gene occurs. Duplication of a gene can reduce expression level if the duplicated region ends up in different regulatory regions, such as near insulators or repressors, a reduction in expression will be observed. Duplications also can reduce expression if they prevents access to the regulatory machinery preventing efficient transcription (Henrichsen et al., 2009).

The association of CNVs with changes in phenotypes is not a recent development. One of the earliest functional CNVs to be identified occurred 78 years ago when a duplication in the Bar gene in Drosophila was discovered to narrow the eye field (Bridges, 1936). The change in copy number in any number of genes is not well tolerated and often leads to the development of disease (Hastings et al., 2009). Most studies examining CNVs in humans have examined the deleterious nature of CNV on phenotypes in diseased populations and have led to various associations between CNVs and disease phenotypes (Almal & Padh, 2012). Various neurological disorders including Parkinson’s disease (Singleton et al., 2003; Pankratz et al., 2011), Alzheimer’s disease (Sleegers et al., 2006) and Autism (Weiss et al., 2008) have been associated with CNVs. Numerous forms of cancers, such as prostate, breast and colorectal have also been associated with CNVs (Yang et al., 2013). Several chronic health related diseases have also been associated with CNVs including various cardiovascular
diseases (Pollex & Hegele, 2007) and metabolic disorders including both type I (Grayson et al., 2010) and type II (Jeon et al., 2010) Diabetes. There have been many CNVs found on known candidate genes for body mass. For example, the deletion of a 2.1Mb region which spans the UPC1 and IL15 genes increases the risk of obesity (Wang et al., 2010). CNV has previously been associated with lean body mass. Individuals with two copies of CNV2073 had 6.9% greater lean body mass than individuals with 3 copies and 11.2% more than individuals with 4 copies. One gene which is located in the region of CNV2073 is Gremlin1 which has already been reported as a candidate for lean body mass by linkage studies and molecular function studies (Hai et al., 2012). Gains of two CNVs (CNV1191 and CNV2580) have also associated with decrease in appendicular lean body mass (Ran et al., 2014).

Despite most of the associations with CNVs being with diseased phenotypes, CNVs have been shown to occur with healthy populations and have advantageous consequences. Salivary amylase (AMY)1 is the most common example of a gene which has CNV with beneficial effects. In populations with high starch diets it has been shown that there is an increase in the AMY1 copy number and elevated expression of the salivary amylase protein which is responsible for starch hydrolysis (Perry et al., 2008). Increased copy number for AMY1 also has been associated with a significantly reduced BMI and risk of obesity (Falchi et al., 2014). CNVs of other genes have been associated with beneficial phenotypes such as delayed progression of spinal muscular atrophy with an increase in the number of copies of the survival motor neuron (SMN)2 gene (Elsheikh et al., 2009). Patients with three or four copies of SMN2 have been shown to produce higher force for grip, elbow flexion, and knee extension and flexion compared to the patients with two copies; none of the increases reached statistical significance (Febrer et al., 2010). These studies show that CNVs can consequences in clinical populations and in the general population. To date no studies have investigated the
impact of CNVs with muscle performance in athletes. We would predict that because of the previous associations with disease, that CNVs in genes related to athletic performance will have phenotypic consequences for muscle function.

1.6 Myosin Light Chain Kinase

The Myosin Light Chain Kinase (MYLK) genes encode for the MYLK proteins which are responsible for the phosphorylation of the regulatory light chain of myosin (RLC). Western blot analysis has shown that the different MYLK proteins differ in the site of expression (Zhi et al., 2005). MYLK1 has been shown to be expressed ubiquitously throughout all tissues, however the main site of expression of the kinase is in smooth muscle and therefore it has been termed the smooth muscle MYLK (smMLCK). MYLK2 has been shown to express the skeletal muscle MYLK (skMLCK) specifically in skeletal muscle and predominantly in the type II fast twitch fibres (Zhi et al., 2005; Herring et al., 2010; Kamm & Stull, 2011). MYLK3 encodes for the cardiac MYLK (cmMLCK) protein which is expressed mainly in the cardiac myocytes (Takashima, 2009). The phosphorylation of the RLC of myosin is important in the contraction process for all three of the muscle types (smooth, skeletal and cardiac) (Takashima, 2009). In general MYLK controls the phosphorylation of the RLC of myosin to allow interaction between myosin and actin filaments in the contraction of muscle fibres (Stull et al., 2011).

Each of the MYLK proteins have different roles specific to the type of muscle it is associated with. In smooth muscle, an increase in intracellular calcium leads to the activation of smMLCK by calmodulin (Kamm & Stull, 2001). The phosphorylation of the RLC by smMLCK has been shown to be sufficient to initiate the contraction process in smooth
muscles through the activation of myosin ATPase (Figure 1)(Kamm & Stull, 2001; Murthy, 2006). In both skeletal and cardiac muscle MYLK activation is not the key step in the initiation of the contraction process because it is not the calcium sensor, instead that role is played by troponin. The binding of calcium to troponin causes a conformational change, allowing interaction between the myosin cross-bridges and actin filaments which leads to the development of force (Farah & Reinach, 1995). In smooth muscle, RLC phosphorylation by smMLCK affects the myosin ATPase activity, whereas in skeletal and cardiac muscle there is no impact on ATPase activity by the specific MLCK. In skeletal muscle, the phosphorylation of RLC by skMLCK leads to changes in properties of the myosin cross-bridges (Figure 1)(Kamm & Stull, 2011). In mice with ablated MYLK2, it has been shown that there is a role for RLC phosphorylation in the potentiation of isometric twitch force. The knockout mouse displays a reduction in the phosphorylation of the RLC and produces significantly lower twitch force (Zhi et al., 2005; Gittings et al., 2011). There is also evidence of some RLC phosphorylation in the knockout mouse which suggests there could be some redundancy between skMLCK and smMLCK (Takashima, 2009). The functional role of MYLK1 and MYLK2 in the contraction of muscle and the suggestion of a potential functional redundancy for skMLCK by smMLCK makes both of these genes candidate genes for genetic variation which is functionally involved in muscle strength and power phenotypes.
Previous SNP association studies attempted to examine if variation in MYLK2 is associated with muscle function. However, the SNPs selected in these studies are found to encode for MYLK1 (C49T (rs2700352) and C37885A (rs28497577)) (Clarkson et al., 2005; Deuster et al., 2013). Despite both of the studies attempting to examine variation in MYLK2 but instead examining MYLK1, associations with strength phenotypes were found. The first study to examine SNP variation in MYLK1 and muscle phenotypes studied 157 patients who performed a muscle damage protocol of the elbow flexors to investigate the genetic determinants of exertional muscle damage (Clarkson et al., 2005). The two muscle strength phenotypes examined were baseline strength and greater strength loss following eccentric exercise of the elbow flexors. A key finding of the study was contrary to the hypothesis as they found patients homozygous for the T allele of C49T produced significantly higher baseline strength values. They also found that patients homozygous of the C allele of
C37885A showed significantly lower post exercise strength loss. A second study examined the risk of developing exertional rhabdomyolysis in 47 disease sufferers and 134 control individuals to identify polymorphisms associated with the disease (Deuster et al., 2013). Two of the polymorphisms that were investigated were the same MYLK1 SNPs as the Clarkson study. They did not find any association with C49T which is consistent with the previous study only finding a relationship with baseline strength. Individuals homozygous for the A allele of MYLK1 C37885A were determined to be 5 times more likely to suffer from exertional rhabdomyolysis than individuals who were CC homozygotes (Deuster et al., 2013), which is consistent with the findings that the C allele causes lower strength loss following exercise (Clarkson et al., 2005). The results of the two studies show that MYLK1 C49T and C37885A are both associated with muscle function changes in clinical populations (Clarkson et al., 2005; Deuster et al., 2013); however these SNPs have yet to be examined in an athletic cohort.

Unpublished data from our laboratory examined the same SNPs of MYLK1 as previous studies (Clarkson et al., 2005; Deuster et al., 2013) in a cohort of athletes and control individuals in an attempt to find if the associations with muscle performance phenotypes in clinical populations remain when examined in an athletic population. Three SNPs of MYLK2 were also examined to test if variation in the skeletal muscle specific gene also associated with muscle performance phenotypes. No muscle strength phenotypes were associated with the SNPs of MYLK2; however, associations were made with isokinetic test performance and the MYLK1 SNPs. Given that SNP variation has been associated with functional changes in muscle performance, we hypothesise that CNVs will affect the function of MYLK1 and therefore be associated with muscle strength phenotypes.
The antisense gene of MYLK1, called MYLK-antisense 1 (MYLK-AS1) is potentially another candidate gene for functional variation associated with athletic performance. Antisense genes are thought to functionally act to regulate gene expression, however the specific role of these mRNA genes is not fully understood (Werner, 2013). These antisense transcripts form duplex RNA through hybridization with the sense transcript. These interactions can reduce the gene expression and decrease the levels of protein synthesis by preventing the translational machinery access to the sense transcript (Kumar & Carmichael, 1998). Another potential role of antisense genes is to control the splicing of the associated sense gene, by preventing access to splice sites (Morrissy et al., 2011). Some antisense genes have been shown to stabilise the expression of the sense genes through competition for microRNA binding sites and RNA motifs (Werner, 2013). Several antisense genes have been shown to be functional however the mechanisms underlying the function have yet to be identified. It is thought that MYLK-AS1 may act to reduce the expression of MYLK1 leading to a reduction in RLC phosphorylation and decreased muscle performance.

The Database of Genomic Variants (MacDonald et al., 2014) identified the presence of CNVs which span the whole of the MYLK1 and MYLK-AS1 genes. There have been no CNVs detected which span the whole of the MYLK2 gene. The studies which identified the presence of CNVs on MYLK1 and MYLK-AS1 did not include any functional measures therefore the phenotypic consequences of changes in copy number in these genes is unknown (Wong et al., 2007; Itsara et al., 2009).
1.7 Aims & Objectives

The aim of the present study was to assess copy number variation in a cohort of athletes and control individuals for the first time and examine if it is associated with strength. MYLK1 and MYLK-AS1 have been selected as the target genes to examine if some of the missing heritability of muscle strength and power traits can be explained by CNVs in genes previously associated with functional SNP variation. There are three main objectives of the project: (1) to examine the extent of variation in MYLK1 and MYLK-AS1 copy number in athletes and control; (2) to examine if there is a different relationship between MYLK1 and MYLK-AS1 copy number between different groups of athletes (endurance and strength/power athletes) and controls; (3) to investigate how the copy number of the two target genes associates with quantitative phenotypes associated with strength and power. We hypothesise that increases in MYLK1 copy number will be overrepresented in the strength athletes and consequently also be associated with increased performance in tests of strength and power phenotypes. We expect to find the opposite in MYLK-AS1 with an underrepresentation in the strength athletes and decreased performance in measures of strength and power. The result of this study will further the understanding of genetic variation which influences muscle function.
Chapter 2 - Materials and Methods

2.1 Ethical Approval

Ethical approval was obtained for the genetics of Lithuanian athletes (GELA) study by the Lithuanian State Bioethics Committee and the University of Stirling, School of Sport Research Ethics Committee. The study abides by the principles outlined in the Declaration of Helsinki. Informed consent in writing was obtained from each participant prior to participation in the study.

2.2 Participant characteristics

The present study utilised the pre-established GELA cohort which consists of native Lithuanians (n=447) from the Lithuanian Sports University (Kaunas, Lithuania). All participants were males (17-37 years old) and where related individuals were recruited only the individual who competed at the highest level was retained for analysis. All participants underwent phenotype determination for a range of standard physiological tests (see: phenotype data). To confirm all participants were performing close to their peak at the time of phenotype determination, any participants unable to perform the physiological tests due to injury or disease were excluded from the study. Any athletes who could not be categorised as strength-power (STP), endurance (END) or they no longer performed at their top level were also excluded from the study. A total of 407 participants remained for further analysis following the exclusions of the participants who did not meet the necessary criteria (n=40).
The participants were divided into 3 separate groups: control individuals (n=197), strength and power athletes (STP; n=126) and endurance athletes (END; n=84). Descriptive characteristics of all participants can be found in Table 2. All athletes trained a minimum of twice a week, whereas the control participants did not take part in organised physical activity more than twice a week and did not compete in competitive sports unless it was a low impact sport. STP included weightlifters, combat athletes, field athletes (discus, javelin and long jump) gymnasts, team sport individuals (handball, volleyball, basketball and football), short distance swimmers and runners, a motocross-rally driver and a high-diver. END included long-distance runners, cyclists, triathletes, a rower, modern pentathletes, orienteers, skiers and walkers. Each of the athlete subgroups was further divided by highest level of competition. International athletes (n=14) were the participants who represented Lithuania at the Olympic Games or World Championships. National athletes (n=97) were ranked in the top 3 for their sport or have been included in a Lithuanian national sport squad but did not compete internationally. Regional athletes (n=63) were defined as athletes that were members of their local area squad. Other athletes (n=36) were a mixture of athletes who did not achieve any of the above levels or chose not to compete but trained regularly. The number of athletes who performed each sport and the level of performance of these athletes can be found in Appendix A.
Table 2 - Descriptive characteristics of the groups of participants. Values are means ± 95% CI. FFM; fat free mass, STP; strength and power athletes, END; endurance athletes

<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Control (n=197)</th>
<th>Athletes (n=210)</th>
<th>STP (n=126)</th>
<th>END (n=84)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yrs)</td>
<td>23.98 ± 0.60</td>
<td>21.77 ± 0.46</td>
<td>21.83 ± 0.57</td>
<td>21.68 ± 0.78</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>180.29 ± 0.83</td>
<td>181.65 ± 2.58</td>
<td>183.27 ± 4.23</td>
<td>179.27 ± 1.17</td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td>77.00 ± 1.50</td>
<td>76.90 ± 1.62</td>
<td>81.30 ± 2.26</td>
<td>70.35 ± 1.44</td>
</tr>
<tr>
<td>Body Fat (%)</td>
<td>15.80 ± 0.73</td>
<td>12.99 ± 0.48</td>
<td>14.12 ± 0.65</td>
<td>11.29 ± 0.51</td>
</tr>
<tr>
<td>FFM (kg)</td>
<td>64.28 ± 0.95</td>
<td>66.34 ± 1.63</td>
<td>69.07 ± 2.55</td>
<td>62.37 ± 1.20</td>
</tr>
<tr>
<td>Training experience (yrs)</td>
<td>N/A</td>
<td>8.04 ± 0.80</td>
<td>8.83 ± 1.05</td>
<td>6.87 ± 1.17</td>
</tr>
<tr>
<td>Training volume (hr/wk)</td>
<td>N/A</td>
<td>11.4 ± 1.13</td>
<td>10.72 ± 1.55</td>
<td>12.36 ± 1.6</td>
</tr>
</tbody>
</table>

2.3 Phenotype data

Participants were required to attend the laboratory for testing on three separate occasions having rested for at least 12 hours and fasted for at least 2 hours. On entry to the laboratory on the first day all participants were assigned a unique ID code to allow all data to be stored in an anonymous manner. All participants had a 10 ml venous blood sample extracted into an EDTA container and frozen at -80 °C prior to DNA extraction. During the laboratory visits the participants were required to fill out a number of questionnaires and complete a range of standardised physiological tests, which were carried out by trained individuals. The laboratory visits allowed detailed phenotypes to be collected including measures of anthropometrics, body composition, endurance performance, strength performance and cardiac size and function. For the current analysis a subset of the phenotypes which are indicative of strength and power performance. The selected tests included measures of isolated muscle group force production such as hand grip strength and isokinetic dynamometry for extension and flexion of the right and left legs and arms at a
range of speeds (30-180 °/s) and performance in whole body movements such as number of pull-ups, countermovement jump and static jump height and time to complete a 30 metre sprint (Details on the tests performed in Appendix B).

2.4 DNA Extraction

All extractions were performed at the University of Stirling. Genomic DNA was extracted from 200 µl of whole blood using silica columns (Macherey-Nagel GmbH & Co. KG, Germany) and extraction buffers (Qiagen Ltd, UK) using the QIAamp Spin DNA extraction protocol. DNA yield and quality was checked by absorbance on a Synergy HT microplate reader (BioTek, USA) with Gen5.0 software. Samples with a yield of less than 10 ng/µl were re-extracted (concentration of DNA samples can be found in Appendix C). All DNA samples were stored at -20 °C prior to analysis.

2.5 Copy Number Assays

To analyse the copy number of MYLK1 and MYLK-AS1, a multiplexed quantitative real-time polymerase chain reaction (qPCR) method was used. Primer and hydrolysis probe assays (Integrated DNA Technologies Inc, USA) were designed to amplify a region which lies in the introns of the genes of interest. Care was taken to ensure that no SNPs had been reported previously in European populations within the primer and probe sequences. The Basic Local Alignment Search Tool (BLAST, National Centre for Biotechnology Information) was used to ensure that the primers and probes were specific to the genes of interest and did not recognise any other sequences. The MYLK1 assay contains the forward primer (5’-ACATACACAAGGTCAGTCACG-3’), reverse primer (5’-
GAGTCTGAACACTGAGGCTAG -3’) and Cy5 labelled probe (5’-CAAGCCACTGAGGATGAGGCA-3’) which amplifies a 99 base pair region. The MYLK-AS1 assay contains the forward primer (5’-TCCAGAGAGAGAGAAACCTTAGTT-3’), reverse primer (5’- GAGATGGETCCAGGATAGTAAAGATAAAG-3’) and 6-FAM labelled probe (5’-TGTCAGGAAGGGAGGAAGAT-3’) which amplifies a 101 base pair region. Figure 2 displays the location of these two amplicons. For determination of gene copy numbers the Type-it CNV probe PCR kit + qC kit was used including the multicopy Type-it CNV Reference Probe Assay Solution (Qiagen Ltd, UK) of a known copy number labelled with a MAX fluorescence probe. The use of this reference assay was selected as the control assay because it has a copy number of 36 which makes it more robust to CNVs than traditional assays which use single copy genes (Table1). The fluorescence emission spectra of the 6-FAM (494-518), MAX (524-557) and Cy5 (643-667) dyes are all close together, therefore, colour compensation was performed to optimise fluorescence detection to prevent any crossover fluorescence from the other probes being picked up.
Figure 2 – Location of the MYLK1 and MYLK-AS1 sequences on Chromosome 3 according to the Genome Reference Consortium Human Build 38. Bp; base pair location on chromosome 3, MYLK1; myosin light chain kinase1, MYLK-AS1; myosin light chain kinase-antisense1.

All samples were run in triplicate and each plate contained a non-template control also run in triplicate. A reaction volume of 12.5 µl was used for all reactions, containing 6.25 µl of 2x Type-it CNV Probe PCR Master Mix (Qiagen Ltd, UK), 0.5 µl of 25x Ref assay, 0.5 µl of 25x MYLK1 assay, 0.5 µl of 25x MYLK-AS1 assay, 2.75 µl of nuclease-free water and 2 µl of DNA normalised to 5 ng/µl. All reactions were performed on the same LightCycler 480 (Roche Diagnostics Ltd, UK) system and included activation at 95 °C for 5 min followed by 40 cycles of alternative 30 s periods at 95 °C and 60 °C as recommended by the Type-it CNV probe protocol.

2.6 Copy number determination

The median absolute deviation method was used to identify any outlier crossing point (Cp) within each triplicate for each assay. The mean of the remaining Cps in each triplicate
was calculated to give the corrected Cp for each sample in the cohort. The median absolute deviation was also used to identify any samples from the cohort which needed to be repeated because of outlying corrected values. The difference between the Cp of genes of interest and the reference assay was calculated to allow determination of the copy number for each participant.

In qPCR reactions, primers which amplify at an efficiency of 90-110% are acknowledged as acceptable for the amplification of targets (D’haene et al., 2010). There are a number of statistical methods which can be used to determine qPCR efficiency all of which can produce a different result because of the different algorithms used (Ruijter et al., 2013). Small changes in reaction efficiency can dramatically alter copy number estimates (Yuan et al., 2007). Standard curves of serial dilutions and LinRegPCR (Ramakers et al., 2003) were used to calculate the efficiencies of the MYLK1 and MYLK-AS1 assays. Different values (range 85-107%) were obtained for both of these measures which would change the number of copies of these genes. The normal copy number for most genes is two and in previous studies only four out of ~1700 individuals have been detected with alterations (duplications or deletions) of copy number (Wong et al., 2007; Itsara et al., 2009). In the present cohort, we calculated that efficiencies of 105.7% and 99.8% for MYLK1 and MYLK-AS1 respectively would lead to a median copy number of two. These efficiencies are within the range that we calculated; therefore we scaled the obtained Cp values using these efficiencies. It should be noted that the scaling of these data will not affect the associations with quantitative phenotypes. All statistical analyses were performed with, and without, the scaling of data and similar results were obtained. To generate the copy number for each participant, copy number was grouped into the nearest integer as has been previously described (Gonzalez et al., 2005; Cantsilieris et al., 2012). The grouping of copy number may reduce the power to detect
associations, however it leads to clearer interpretation of associations and similar results were obtained when correlations were performed on the raw copy number or integer copy number.

2.7 Statistical Analysis

All statistical analysis was carried out using Minitab (version 16, Minitab, State College, PA). All phenotype data were tested for normality using the Ryan-Joiner test (Ryan, 1974). Any data that were not normal were transformed using the Box-Cox transformation to give better approximation of the normal distribution. Data were back-transformed for display in figures and tables, thus all values are means ± 95% confidence intervals unless otherwise stated.

2.7.1 Comparison of copy number / genotype between groups

Chi-squared contingency tables (corrected using the Yates correction for continuity (Yates, 1934) where appropriate) were used to determine if the athletic groups differed from each other or from the control group in copy number for each of the target genes. Differences between the groups were examined using odds ratios (OR) to identify the likelihood of an individual being assigned to a particular athlete group or performance level achieved based on copy number. The significance of the OR was determined using chi squared contingency tables and 95% confidence intervals (CI). Significance was accepted as chi-squared p<0.05.
2.7.2 Association of copy number with physiological variables

All phenotype data were z-scored using the overall mean and standard deviation. Phenotype data were separately z-scored within each of the subgroups to account for group differences. Pearson’s correlation analysis was performed using the z-scored phenotype data with and without group corrections to identify associations between copy number and physiological variables. The Benjamini-Hochberg (BH) correction for multiple testing was applied to control for false discovery rate separately for isolated muscle and whole body performance tests (Benjamini & Hochberg, 1995). Pearson’s correlations were accepted as significant where p-value < α-value determined in BH correction. For the significant correlations, two tailed T-tests were used to compare the means in quantitative phenotypes for each copy number. T-test significance was accepted as p<0.05.
Chapter 3 - Results

3.1 Copy number detection in each group

Copy number determination was achieved for 405 of the individuals (99.5% success). The same two individuals failed for both target genes and the reference assay. Out of the 405 individuals examined, 12 (3%) individuals have variations of MYLK1 away from the normal copy number of two, whereas 18 (4.5%) individuals have variations of copy number for MYLK1-AS1. Only one control had a deviation of copy number for MYLK1 which is at the frequency expected (Wong et al., 2007; Itsara et al., 2009); however, controls with deviations in copy number for MYLK-AS1 are found at a greater frequency than anticipated. It should be noted, however, that previous studies used to determine expected frequencies used a range of ethnicities which could differ in allele frequencies (Wong et al., 2007; Itsara et al., 2009). In the group of athletes (N=208), 11 and 12 have alterations away from the normal copy number of two for MYLK1 and MYLK-AS1 respectively. Deviations away from a copy number of two have been shown to be a more regular occurrence in STP than END for both of the target genes. The number of deletions and duplications within each of the groups can be found in Table 3.
Table 3 – The copy number of the individuals within each of the groups. Values are the total number with the percentage of the group in brackets. CN; copy number, STP; strength and power athlete, END; endurance athlete

<table>
<thead>
<tr>
<th></th>
<th>MYLK1 CN (%)</th>
<th>MYLK-AS1 CN (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Total (n=405)</td>
<td>3 (0.74)</td>
<td>393 (97.04)</td>
</tr>
<tr>
<td>Athlete (n=208)</td>
<td>3 (1.44)</td>
<td>197 (94.71)</td>
</tr>
<tr>
<td>STP (n=124)</td>
<td>2 (1.61)</td>
<td>116 (93.55)</td>
</tr>
<tr>
<td>END (n=84)</td>
<td>1 (1.19)</td>
<td>81 (96.43)</td>
</tr>
<tr>
<td>Control (n=197)</td>
<td>0 (0.00)</td>
<td>196 (99.49)</td>
</tr>
</tbody>
</table>

3.2 Copy number differences between groups

A trend exists where the copy number of MYLK1 in the athletes differs to the controls ($\chi^2 = 5.150$, $p=0.076$). When athlete subgroups are examined, the distribution of copy numbers is significantly different between STP and the controls ($\chi^2 = 6.103$, $p=0.047$). END did not differ from the controls ($\chi^2 = 0.828$, $p=0.661$) or STP ($\chi^2 = 0.436$, $p=0.804$) for MYLK1 copy numbers. OR display that athletes are less likely to have two copies ($p=0.005$) of MYLK1 and there is a trend where athletes are more likely to have three copies ($p=0.051$) compared to the controls. STP are less likely to have two copies ($p=0.003$) and more likely to have three copies ($p=0.028$) than controls (Table 4). There were no associations between MYLK-AS1 copy number and any of the groups ($\chi^2 = 0.114-0.268$; $p=0.874-0.945$).
Table 4 - Odds Ratios for athlete group vs copy number for MYLK1. Odds Ratio of N/A is because none of the controls have a copy number of one for MYLK1. STP; strength and power athletes, CI; Confidence interval

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Copy number</th>
<th>p-value</th>
<th>Odds Ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Athletes vs Controls</td>
<td>1</td>
<td>0.264</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.005</td>
<td>0.09 (0.01-0.71)</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.051</td>
<td>7.84 (0.97-63.27)</td>
</tr>
<tr>
<td>STP vs Controls</td>
<td>1</td>
<td>0.287</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>0.003</td>
<td>0.07 (0.01-0.60)</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.028</td>
<td>9.97 (1.19-83.81)</td>
</tr>
</tbody>
</table>

In the next stage of analysis, the highest level of competition was investigated for associations between copy numbers of the target genes. For MYLK1, the only association found with level of performance was between Regional level athletes and the controls ($\chi^2 = 6.618; p=0.037$). The Regional athletes are less likely to have two copies ($p=0.003$ [OR 0.06; 95% CI:0.01-0.5]) and more likely to have three copies ($p=0.013$ [OR 13.75; 95% CI:1.51-125.52]) than the controls. No associations were found for highest level of competition and MYLK-AS1 copy number ($\chi^2 = 0.055-1.644, p=0.440-0.973$).

### 3.3 Copy number associations with quantitative measurements

To account for the difference in training of the participants, the relationship between copy number and quantitative phenotypic measurements has been investigated with and without corrections for subgroup (Table 5 and 6). To test for associations which could potentially confound other analyses, correlations were made between MYLK1 / MYLK-AS1 copy number and age, height, body mass, training history and volume. No significant
correlations existed between any of these variables and MYLK1 / MYLK-AS1 copy number before or after correction for subgroup (p=0.091-0.878).

Copy number of the target genes was investigated for associations with selected phenotypes associated with strength and power performance for isolated muscle groups and whole body movements. To assess the correlation between these phenotypes and copy number, Pearson’s correlations were performed with BH correction for multiple testing was applied to reduce the risk of type 1 statistical errors. Overall in the whole cohort MYLK1 copy number was associated with strength and power phenotypes, in isolated muscle groups and whole body movements (Table 5). The average variance explained by these significant correlations is 1.5% with the highest variance being explained by isokinetic extension of the elbow (2.76%). After correction for subgroup the associations for counter movement jump and sprint performance are lost whilst the remaining associations are weakened (Table 5). The average variance explained by the group corrected correlations is 1.35 %. The highest percentage of the variance is still found with the isokinetic extension of the elbow, however the variance drops down to 2.34 %. None of the strength and power phenotypes correlated with copy number for MYLK-AS1 with or without the correction for subgroup (Table 6).
Table 5 - Pearson's correlation coefficient results for MYLK1 copy number and quantitative variables before and after correction for subgroup. Significant (p<0.05) values are highlighted in bold. Values marked by * have a p-value lower than the acceptable alpha value according to the BH correction. kgFFM; kilogram of fat free mass, IKR, isokinetic right; IKL, isokinetic left

<table>
<thead>
<tr>
<th>Trait</th>
<th>MYLK1</th>
<th>Before Subgroup Correction</th>
<th>After Subgroup Correction</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>p-value</td>
<td>Coefficient</td>
</tr>
<tr>
<td>Number of pull-ups</td>
<td>0.08</td>
<td>0.089</td>
<td>0.792</td>
</tr>
<tr>
<td>Counter movement jump height (cm)</td>
<td>0.049</td>
<td>0.099</td>
<td>0.980</td>
</tr>
<tr>
<td>Static jump height (cm)</td>
<td>0.009*</td>
<td>0.132</td>
<td>1.742</td>
</tr>
<tr>
<td>Sprint 0-10 (s/kgFFM)</td>
<td>0.022*</td>
<td>-0.116</td>
<td>1.346</td>
</tr>
<tr>
<td>Sprint 10-20 (s/kgFFM)</td>
<td>0.018*</td>
<td>-0.121</td>
<td>1.464</td>
</tr>
<tr>
<td>Sprint 20-30 (s/kgFFM)</td>
<td>0.028</td>
<td>-0.112</td>
<td>1.254</td>
</tr>
<tr>
<td>Sprint Total (s/kgFFM)</td>
<td>0.020*</td>
<td>-0.118</td>
<td>1.392</td>
</tr>
<tr>
<td>Total Handgrip (kg)</td>
<td>0.029</td>
<td>0.111</td>
<td>1.232</td>
</tr>
<tr>
<td>IKR knee extension torque (30 °/s)</td>
<td>0.081</td>
<td>0.091</td>
<td>0.828</td>
</tr>
<tr>
<td>IKL knee extension torque (30 °/s)</td>
<td>0.026</td>
<td>0.116</td>
<td>1.346</td>
</tr>
<tr>
<td>IKR knee extension torque (90 °/s)</td>
<td>0.024</td>
<td>0.117</td>
<td>1.369</td>
</tr>
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<td>IKL knee extension torque (90 °/s)</td>
<td>0.007*</td>
<td>0.139</td>
<td>1.932</td>
</tr>
<tr>
<td>IKR knee extension torque (180 °/s)</td>
<td>0.045</td>
<td>0.104</td>
<td>1.082</td>
</tr>
<tr>
<td>IKL knee extension torque (180 °/s)</td>
<td>0.020</td>
<td>0.121</td>
<td>1.464</td>
</tr>
<tr>
<td>IKR knee flexion torque (30 °/s)</td>
<td>0.220</td>
<td>0.064</td>
<td>0.410</td>
</tr>
<tr>
<td>IKL knee flexion torque (30 °/s)</td>
<td>0.199</td>
<td>0.067</td>
<td>0.449</td>
</tr>
<tr>
<td>IKR knee flexion torque (90 °/s)</td>
<td>0.137</td>
<td>0.077</td>
<td>0.593</td>
</tr>
<tr>
<td>IKL knee flexion torque (90 °/s)</td>
<td>0.156</td>
<td>0.074</td>
<td>0.548</td>
</tr>
<tr>
<td>IKR knee flexion torque (180 °/s)</td>
<td>0.064</td>
<td>0.096</td>
<td>0.922</td>
</tr>
<tr>
<td>IKL knee flexion torque (180 °/s)</td>
<td>0.084</td>
<td>0.090</td>
<td>0.810</td>
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<td>IKR elbow extension torque (30 °/s)</td>
<td>0.009*</td>
<td>0.136</td>
<td>1.850</td>
</tr>
<tr>
<td>IKL elbow extension torque(30 °/s)</td>
<td>0.009*</td>
<td>0.135</td>
<td>1.823</td>
</tr>
<tr>
<td>IKR elbow extension torque (90 °/s)</td>
<td>0.177</td>
<td>0.071</td>
<td>0.504</td>
</tr>
<tr>
<td>IKL elbow extension torque (90 °/s)</td>
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<td>0.111</td>
<td>1.232</td>
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<td>IKR elbow extension torque(180 °/s)</td>
<td>0.001*</td>
<td>0.166</td>
<td>2.756</td>
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<td>0.028</td>
<td>0.114</td>
<td>1.300</td>
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<td>IKR elbow flexion torque (30 °/s)</td>
<td>0.014*</td>
<td>0.128</td>
<td>1.638</td>
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<td>IKL elbow flexion torque (30 °/s)</td>
<td>0.026</td>
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<td>1.323</td>
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<td>IKR elbow flexion torque (90 °/s)</td>
<td>0.292</td>
<td>0.055</td>
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<td>IKL elbow flexion torque (90 °/s)</td>
<td>0.121</td>
<td>0.080</td>
<td>0.640</td>
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<tr>
<td>IKR elbow flexion torque (180 °/s)</td>
<td>0.456</td>
<td>0.039</td>
<td>0.152</td>
</tr>
<tr>
<td>IKL elbow flexion torque (180 °/s)</td>
<td>0.009*</td>
<td>0.135</td>
<td>1.823</td>
</tr>
</tbody>
</table>
Table 6 - Pearson's correlation coefficient results for MYLK1-AS1 copy number and quantitative variables before and after correction for subgroup. Significant (p<0.05) values are highlighted in bold. Values marked by * have a p-value lower than the acceptable alpha value according to the BH correction.

<table>
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<tr>
<th>Trait</th>
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<th>With Group Correction</th>
</tr>
</thead>
<tbody>
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<td></td>
<td>p-value</td>
<td>Coefficient</td>
</tr>
<tr>
<td>Number of pull-ups</td>
<td>0.807</td>
<td>0.012</td>
</tr>
<tr>
<td>Counter movement jump height (cm)</td>
<td>0.795</td>
<td>0.013</td>
</tr>
<tr>
<td>Static jump height (cm)</td>
<td>0.583</td>
<td>0.028</td>
</tr>
<tr>
<td>Sprint 0-10 (s/kgFFM)</td>
<td>0.913</td>
<td>-0.006</td>
</tr>
<tr>
<td>Sprint 10-20 (s/kgFFM)</td>
<td>0.849</td>
<td>-0.010</td>
</tr>
<tr>
<td>Sprint 20-30 (s/kgFFM)</td>
<td>0.731</td>
<td>-0.017</td>
</tr>
<tr>
<td>Sprint Total (s/kgFFM)</td>
<td>0.843</td>
<td>-0.010</td>
</tr>
<tr>
<td>Total Handgrip (kg)</td>
<td>0.272</td>
<td>0.056</td>
</tr>
<tr>
<td>IKR knee extension torque (30 °/s)</td>
<td>0.911</td>
<td>0.006</td>
</tr>
<tr>
<td>IKL knee extension torque (30 °/s)</td>
<td>0.094</td>
<td>0.087</td>
</tr>
<tr>
<td>IKR knee extension torque (90 °/s)</td>
<td>0.768</td>
<td>0.015</td>
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<tr>
<td>IKL knee extension torque (90 °/s)</td>
<td>0.949</td>
<td>0.003</td>
</tr>
<tr>
<td>IKR knee extension torque (180 °/s)</td>
<td>0.699</td>
<td>0.020</td>
</tr>
<tr>
<td>IKL knee extension torque (180 °/s)</td>
<td>0.571</td>
<td>-0.029</td>
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<tr>
<td>IKR knee flexion torque (30 °/s)</td>
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<td>IKL knee flexion torque (30 °/s)</td>
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<td>-0.013</td>
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<tr>
<td>IKR knee flexion torque (90 °/s)</td>
<td>0.995</td>
<td>0.001</td>
</tr>
<tr>
<td>IKL knee flexion torque (90 °/s)</td>
<td>0.859</td>
<td>-0.009</td>
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<tr>
<td>IKR knee flexion torque (180 °/s)</td>
<td>0.537</td>
<td>0.032</td>
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<tr>
<td>IKL knee flexion torque (180 °/s)</td>
<td>0.619</td>
<td>0.026</td>
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<tr>
<td>IKR elbow extension torque (30 °/s)</td>
<td>0.967</td>
<td>-0.002</td>
</tr>
<tr>
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<td>-0.007</td>
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<tr>
<td>IKR elbow extension torque (90 °/s)</td>
<td>0.986</td>
<td>-0.001</td>
</tr>
<tr>
<td>IKL elbow extension torque (90 °/s)</td>
<td>0.869</td>
<td>0.009</td>
</tr>
<tr>
<td>IKR elbow extension torque(180 °/s)</td>
<td>0.407</td>
<td>0.043</td>
</tr>
<tr>
<td>IKL elbow extension torque (180 °/s)</td>
<td>0.429</td>
<td>0.041</td>
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<tr>
<td>IKR elbow flexion torque (30 °/s)</td>
<td>0.924</td>
<td>0.005</td>
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<td>IKL elbow flexion torque (30 °/s)</td>
<td>0.984</td>
<td>0.001</td>
</tr>
<tr>
<td>IKR elbow flexion torque (90 °/s)</td>
<td>0.968</td>
<td>-0.002</td>
</tr>
<tr>
<td>IKL elbow flexion torque (90 °/s)</td>
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<td>-0.003</td>
</tr>
<tr>
<td>IKR elbow flexion torque (180 °/s)</td>
<td>0.628</td>
<td>-0.025</td>
</tr>
<tr>
<td>IKL elbow flexion torque (180 °/s)</td>
<td>0.841</td>
<td>-0.010</td>
</tr>
</tbody>
</table>
3.4 Subgroup copy number associations with quantitative variables.

The subgroup corrected correlations of MYLK1 (Table 5) indicates that there are differences between the subgroups therefore, all of the subgroups were individually examined for correlations between MYLK1 copy number and phenotypes associated with strength and power performance. An increased copy number in STP is associated with increases in the jump performance, handgrip, isokinetic test performance and sprinting performance. However, after the BH correction only static jump and isokinetic performance remain significant (Table 7). The strength of the correlations is stronger in STP compared to when the whole cohort is examined together as seen by a higher percentage of the variance being explained (Table 7). In the control group, only one individual has a deviation in copy number away from two therefore accurate correlations could not be performed and no associations are found in END (data presented in Appendix D). There were no significant subgroup corrected or uncorrected correlations for MYLK-AS1, therefore the subgroups were not investigated for correlations between copy number and performance in the strength and power tests.
Table 7 - Pearson's correlation coefficient results between STP for MYLK1 copy number and quantitative variables. Correlations with p<0.05 are highlighted in bold and values marked by * have a p-value lower than the acceptable alpha value according to the BH correction. kgFFM; kilogram of fat free mass IKR, isokinetic right; IKL, isokinetic left

<table>
<thead>
<tr>
<th>Trait</th>
<th>MYLK1 CN SSP associations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>p-value</td>
</tr>
<tr>
<td>Number of pull-ups</td>
<td>0.125</td>
</tr>
<tr>
<td>CMJ height (cm)</td>
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</tr>
<tr>
<td>Static jump height (cm)</td>
<td>0.022*</td>
</tr>
<tr>
<td>Sprint 0-10 (s/kgFFM)</td>
<td>0.032</td>
</tr>
<tr>
<td>Sprint 10-20 (s/kgFFM)</td>
<td>0.038</td>
</tr>
<tr>
<td>Sprint 20-30 (s/kgFFM)</td>
<td>0.091</td>
</tr>
<tr>
<td>Sprint Total (s/kgFFM)</td>
<td>0.043</td>
</tr>
<tr>
<td>Total Handgrip (kg)</td>
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</tr>
<tr>
<td>IKR knee extension torque (30 °/s)</td>
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</tr>
<tr>
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</tr>
<tr>
<td>IKR knee extension torque (90 °/s)</td>
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<tr>
<td>IKL knee extension torque (90 °/s)</td>
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<tr>
<td>IKR knee extension torque (180 °/s)</td>
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<tr>
<td>IKL knee extension torque (180 °/s)</td>
<td>0.038</td>
</tr>
<tr>
<td>IKR knee flexion torque (30 °/s)</td>
<td>0.721</td>
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<tr>
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<tr>
<td>IKR knee flexion torque (90 °/s)</td>
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</tr>
<tr>
<td>IKL knee flexion torque (90 °/s)</td>
<td>0.496</td>
</tr>
<tr>
<td>IKR knee flexion torque (180 °/s)</td>
<td>0.217</td>
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<tr>
<td>IKL knee flexion torque (180 °/s)</td>
<td>0.325</td>
</tr>
<tr>
<td>IKR elbow extension torque (30 °/s)</td>
<td>0.029</td>
</tr>
<tr>
<td>IKL elbow extension torque (30 °/s)</td>
<td>0.06</td>
</tr>
<tr>
<td>IKR elbow extension torque (90 °/s)</td>
<td>0.223</td>
</tr>
<tr>
<td>IKL elbow extension torque (90 °/s)</td>
<td>0.087</td>
</tr>
<tr>
<td>IKR elbow extension torque (180 °/s)</td>
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</tr>
<tr>
<td>IKL elbow extension torque (180 °/s)</td>
<td>0.065</td>
</tr>
<tr>
<td>IKR elbow flexion torque (30 °/s)</td>
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</tr>
<tr>
<td>IKL elbow flexion torque (30 °/s)</td>
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<td>IKR elbow flexion torque (90 °/s)</td>
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<tr>
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<tr>
<td>IKR elbow flexion torque (180 °/s)</td>
<td>0.597</td>
</tr>
<tr>
<td>IKL elbow flexion torque (180 °/s)</td>
<td>0.007*</td>
</tr>
</tbody>
</table>

The phenotypes associated with strength and power which were found to be correlated to MYLK1 copy number were examined in the overall cohort (see Appendix E) and STP (see Appendix F) for the percentage difference between copy numbers. The strongest correlations
were found for performance in isolated muscles. Overall in the cohort and in STP, the highest amount of variance explained is for the isokinetic extension of the elbow at 180 °/s. Overall individuals with three copies of MYLK1 were able to produce 38% more force than individuals with two copies, while STP with three copies are able to produce 28% more force than STP with two copies (Figure 3).

![Figure 3 – Mean peak force produced (N*m) for isokinetic right arm extension of the elbow at 180 °/s for each copy number in the whole cohort and the STP alone. Copy numbers which do not share the same letter are significantly different from each other with each group. CN; copy number, IKR Ext; isokinetic right arm Extension, STP; strength and power athletes.](image-url)

For performance in the whole body tasks, overall in the whole individuals with three copies were able to jump 12% and 32% higher in the static jump than individuals with two copies and one copy respectively. Static jump performance indicates that STP with two copies are able to jump 36% higher than STP with one copy and a trend existed for STP with three copies jumping higher than STP with one copy (p=0.064; Figure 4). Overall in the whole cohort, individuals with three copies are 11% quicker per unit of fat free mass at sprinting 30m compared to individuals with only two copies. There is a trend for STP with
three copies being quicker per unit of fat free mass than STP with one (p=0.062) or two (p=0.0791) copies (Figure 5).

Figure 4 - Mean static jump height (cm) for each copy number in the whole cohort and the STP. Copy numbers which do not share the same letter are significantly different from each other with each group. CN; copy number, STP; strength and power athletes.

Figure 5 - Mean 30 m sprint time (presented as s/kgFFM) as for each copy number in the whole cohort and the STP alone. Copy numbers which do not share the same letter are significantly different from each other with each group. CN; copy number, kgFFM; kilograms of fat free mass, STP; strength and power athletes.
Chapter 4 - Discussion

4.1 Main Findings

The current study aimed to investigate if higher MYLK1 copy number and lower MYLK-AS1 copy number is associated with athlete status and performance in physiological tests associated with strength and power performance in a cohort of athletes and non-athlete controls. We report, for the first time, increased copy number of MYLK1 is associated with athlete status and improved muscle performance. The main findings of the current study are: 1) individuals with higher MYLK1 copy number are more likely to be strength and power athletes than controls; 2) the highest level of competition achieved is not associated with MYLK1 copy number; 3) an increase in MYLK1 copy number is associated with better performance in tests indicative of strength and power, across the whole cohort and for STP alone; 4) copy number of MYLK-AS1 does not influence athlete status or performance in tasks associated with strength and power.

4.2 MYLK1

4.2.1 Copy number and athlete status

In the current study, we hypothesised that there would be an overrepresentation of individuals with a higher copy number in STP. We demonstrated that individuals with an additional copy of MYLK1 are more likely to be a STP than a control (Table 4). A deviation of copy number away from two does not confer an added chance of being an END compared to control. Taken together, these data suggest that having three copies of MYLK1 provides an advantage which predisposes individuals to becoming a STP. Although no other studies have examined
the nature of CNVs in athletes and associations with athlete status, several studies have associated CNVs with the chance of developing several diseases. There are several diseases which have been associated with the alterations of copy number, including neurological disorders (Singleton et al., 2003; Sleegers et al., 2006; Weiss et al., 2008; Pankratz et al., 2011), various forms of cancer (Yang et al., 2013) and several chronic health diseases (Pollex & Hegele, 2007; Grayson et al., 2010; Jeon et al., 2010). There are also examples of CNVs reducing the risk of health disorders such as obesity (Falchi et al., 2014) and delaying the onset of spinal muscular atrophy (Elsheikh et al., 2009). The associations between CNVs and disease highlight the fact that alterations in gene copy number can be functional for a range of different traits and that consequences can be beneficial or deleterious. The mechanism by which CNVs have phenotype consequences and alter disease susceptibility remains to be fully understood but is thought to occur through a change in the expression of the affected gene (Pollex & Hegele, 2007; Grayson et al., 2010; Yang et al., 2013). Despite this being the first study to associate CNVs with athlete status, other forms of genetic variation, namely SNPs have been associated with athlete status.

When SNP genotype frequencies of athletes are examined there is an indication that there is a genetic predisposition to the trade-off between endurance and power capacity. Certain genotypes are over-represented in either strength or endurance athletes which indicates individuals are genetically predisposed to perform better in either strength/power or endurance-type activity (Yang et al., 2003). ACTN3 is a muscle structural protein involved in the stabilization of muscle contractions and is predominantly expressed in fast twitch muscle fibres (Yang et al., 2003; Moran et al., 2007; Walsh et al., 2008). The X allele of the R577X ACTN3 polymorphism has been shown to render the protein non-existent (Berman & North, 2010). When the genotype frequencies of R577X are examined, it is found that there is a lack
of the X allele and over-representation of the R allele in strength and power athletes including Russian power athletes (Druzhevskaya et al., 2008), Greek power athletes (Papadimitriou et al., 2008) Finnish sprinters (Niemi & Majamaa, 2005), American strength athletes (Roth et al., 2008) and Australian sprint/power athletes (Yang et al., 2003). These studies show that genetic variants which affect fast twitch fibres can be overrepresented in a cohort of power and strength athletes but not in endurance based athletes as was found in the present study.

In humans, the majority of muscles are comprised of a mixed fibre type composition, therefore it has been suggested that individuals may be predisposed to either power or endurance phenotypes (Van Damme et al., 2002; Yang et al., 2003). Animal models have extensively been used to determine if there is a pre-determined evolutionary trade-off in muscle capacity towards either power or endurance phenotypes. In mice a negative correlation exists between maximum power output and fatigue resistance of muscles suggesting an evolutionary trade-off (Wilson & James, 2004). In a study examining the performance of decathletes, it has been found that performance in the 1500m race is negatively correlated with the performance in the 100m, 100m hurdles, shot put and long jump (Van Damme et al., 2002). These findings suggest that the performance of decathletes is either favoured towards power or endurance and indicate that in humans a trade-off between endurance and power capacity does in fact exist. Genetic variants, such as, MYLK1 copy number may influence the sport in which athletes perform as is indicated by the results of the present study.

A potential reason for the overrepresentation of higher MYLK1 copy number in the STP but not in the END is because MYLK is predominately expressed in the fast twitch type
II muscle fibres rather than slow twitch type I muscle fibres (Zhi et al., 2005; Herring et al., 2010; Kamm & Stull, 2011). The fibre type distributions of athletes who compete in different events have been examined in previous studies (Costill et al., 1976; Zierath & Hawley, 2004). It has been found that athletes who perform endurance based events have a higher relative percentage of slow twitch fibres and sprinters have a higher percentage of fast twitch fibres (Costill et al., 1976). The increased percentage of fast twitch fibres in sprinters is possibly because they are the fibres which can produce higher force and velocity, which are required to compete in short distance, high power output events (Zierath & Hawley, 2004). Therefore, it is expected that in our cohort the greatest distribution of fast twitch fibres will be found in the STP and these individuals will be the most affected by functional genetic variation in the MYLK1 gene.

4.2.2 Level of Competition

In our cohort of athletes when the highest level of competition achieved is examined for associations with MYLK1 copy number, only the regional athletes differ from the controls for increased copy number. If MYLK1 copy number was associated with the highest level of competition achieved then we would expect to find that the higher classifications of athletes (international and national standard) would differ from the lower levels of athletes (regional and other) and controls. Given the lack of associations across different competition levels, our results indicate a gain in copy number is associated with the selection of sport because of natural ability, although there might not be any further advantage gained from training.
Genetic variants for power and endurance based sports have shown increased frequencies of one allele in higher level athletes. In a study looking at 10 common genotypes which have been associated with endurance performance it is found that individuals with a higher number of ‘endurance alleles’ are more likely to be an elite athletes than non-elite athletes (Ahmetov et al., 2009). Another study has found the frequency of the Gly482 allele of the PPARGC1A gene was 20% higher in top-level athletes compared to national level athletes (Eynon et al., 2010). Associations also have been made with genotype frequencies in elite and non-elite power orientated athletes. In a cohort of elite Australian power athletes (Yang et al., 2003) and a cohort of Finnish sprinters (Niemi & Majamaa, 2005), the XX genotype of ACTN3 R577X is not detected. These studies strengthen the theory that certain genotypes may predispose athletes to certain athletic disciplines since higher level athletes show an increased allele frequency compared to both controls and lower level athletes. Despite these previous findings that a range of genes have been associated with elite athlete status; MYLK1 copy number does not associate with elite athletic performance in the present study.

A potential reason for the lack of findings in the level of competition of the athletes is that both higher and lower MYLK1 copy number are rare variants and therefore found at a much lower frequency than the genetic variants examined in other studies. A gain in copy number was only found in our cohort at 2% (5% in STP) and only 12 individuals (3%) in the cohort had deviations away from a normal copy number of two (Table 3). In previous studies considering the highest level of competition achieved, the minor allele frequency has been greater than 5% in all cases and is typically around 30% (Yang et al., 2003; Niemi & Majamaa, 2005; Ahmetov et al., 2009; Eynon et al., 2010). Another potential reason for MYLK1 copy number associating with STP but not level of competition is that the overall
ability of an athlete is determined through various genetic polymorphisms and environmental factors (Pitsiladis et al., 2013). Environmental factors such as opportunity, economic possibility and social support are all involved in the determination of elite level athletes (Lucía et al., 2010). The sport in which the athlete competes also needs to be considered for the level of competition achieved; regional athletes of one sport may actually be a better overall calibre athlete than a national level athlete of a different sport but competition for places in the national squad may be the limiting factor. Therefore highest level of competition achieved may not be the best measure for the effect of the genetic variants because it may not be representative of overall ability.

4.2.3 Quantitative data

The inclusion of the quantitative phenotype data strengthens the study because it allows the investigation of the impact of copy number on different aspects of performance which provides more conclusive findings than the qualitative nature of athlete status and level of competition data. Positive correlations were found between MYLK1 copy number and measures of strength and power in both isolated muscles and dynamic movements. The correlations were absent or weakened after correction for subgroup. Therefore, the differences between groups explain some of the phenotypic consequences between copy number associations. Between-group differences are not surprising because of the different training strategies required for STP and END and a lack of training performed by the controls. Each of the athletic subgroups was examined individually for the correlations between copy number and quantitative traits. Only one of the controls had a deviation away from a copy number of two. Therefore, correlations were not performed within this group. Several correlations were found in the STP for both isolated and dynamic muscle
performance (Figure 3-5), whereas no associations were found in END (Appendix D). The correlations in STP were expected because of the overrepresentation of STP with three copies of MYLK1. The existence of correlations in STP and not END suggest that it may be the type II muscle fibres which are affected or that the effect of the CNV is increased through training specific to STP.

Different polymorphisms have been associated with different aspects of performance, such as response to training or maximal force production (Kostek et al., 2010; Ahmetov & Fedotovskaya, 2012). Previous MYLK1 SNP studies have identified a potential role of MYLK1 variation in the determination of baseline strength levels (Clarkson et al., 2005) which could suggest that MYLK1 may make individuals stronger and therefore more likely to perform sports related to power rather than endurance. The previous associations with MYLK1 have only been examined in isolated muscle groups therefore the effect on physiological performance is yet unknown (Clarkson et al., 2005). The current study associates increased copy number with both performance in isolated muscle groups and performance in whole body movements physiologically relevant to performance. The findings with phenotypes associated with isolated muscles (Figure 3), such as dynamometry measures, allow the determination of peak force, however these are not physiologically relevant measures of performance. Individuals with three copies of MYLK1 were able to jump higher than individuals with one or two copies. An advantage also is seen in sprinting performance over 30m where the individuals with three copies are quicker per unit of fat free mass than the individuals with only two copies. MYLK1 copy number is shown to explain a lower percentage of the variance in these whole body traits; however, this is expected because these whole body movements consist of the contractions of several muscles, therefore external factors have more of a chance to affect them.
Two SNPs in MYLK1 have previously been associated with muscle performance in diseased cohorts (Clarkson et al., 2005; Deuster et al., 2013) which supports the findings of the present study. The T allele of C49T of MYLK1 has been associated with increased baseline strength in a clinical cohort (Clarkson et al., 2005). Unpublished work in this cohort of athletes reports associations with the C allele of C49T and increased force production. The differences in the allele which is associated with strength could be explained by the different cohorts that were examined. The other SNP which has been associated with muscle performance phenotypes is C37885A. The C allele of C37885A has been shown to have a protective effect on strength loss following muscle damaging exercise (Clarkson et al., 2005). Unpublished work in the present cohort has associated the C allele with increased peak force in isokinetic arm tests which supports the findings in the literature of the C allele producing an advantageous muscle performance phenotype. The SNP studies support our findings that genetic variants of MYLK1 are associated with performance in quantitative muscle performance traits.

With SNPs typically only explaining 1-2% of the variance in muscle strength between individuals, a large percentage of the variance remains unexplained (Kostek et al., 2010). The CNV of MYLK1 in this study has been shown to explain at most 3% of the variance between individuals in the cohort. When the variance in the STP is examined, we demonstrate that over 7% of the performance in high speed elbow extension between individuals is explained by differences in copy number. The results indicate that CNVs may contribute more too phenotypic traits than SNPs which is expected because SNPs only affect one nucleotide; whereas CNVs can influence millions of nucleotides and thus can have more drastic effects. More studies on CNVs and athletic performance are required to confirm the associations before a statement on the size of the effect compared to SNPs can be made.
4.2.4 Mechanisms

Increased MYLK1 copy number is thought to associate with increased performance in strength and power phenotypes through an increase in the expression of the MYLK1 gene. Despite MYLK1 being expressed in skeletal muscle, skeletal muscle RLC is not a good substrate for MYLK1 because of differences in substrate properties (Kamm & Stull, 2001). In MYLK2 ablated mice the phosphorylation of the RLC still occurs which indicates there may be a redundancy mechanism between skMLCK and smMLCK despite a lack of increased MYLK1 expression as a result (Takashima, 2009). The phosphorylation of the RLC in MYLK2 ablated mice supports this because it indicates that other kinases are able to phosphorylate skeletal muscle RLC and will have the same effect on enhanced rate of force development as MYLK2 (Zhi et al., 2005). In cardiac tissue the smMLCK is expressed at a lower level than cmMLCK and despite differences in substrate properties, smMLCK is still capable of phosphorylating the cardiac muscle RLC. This has led to the suggestion that following the decline in cmMLCK expression smMLCK may act as a backup kinase (Josephson et al., 2011). We speculate that the increased copy number of MYLK1 leads to increased expression of smMLCK which acts together with the skMLCK to produce increases in the rate of force development and force potentiation.

In humans, RLC phosphorylation has been linked to post-activation potentiation (PAP) of type II muscle fibres (Lorenz, 2011). The previous contractile history of the muscle has a large effect on the ability of the muscle to generate force and enhanced contractile response is thought to occur following high intensity contraction (Docherty & Hodgson, 2007). PAP results in increased force and rate of force development which has led to
suggestions that the molecular mechanism responsible for the enhanced contractile response is increased phosphorylation of the RLC. The phosphorylation of the RLC by MLCK occurs at a rapid rate and the dephosphorylation is a slower process, therefore it is thought that prior contractions function to prime the muscle for future contractions and subsequently increase the rate of force development and peak isometric twitch force (Zhi et al., 2005). Greater PAP has been consistently observed in trained individuals compared to untrained individuals (Chiu et al., 2003) and amongst individuals with a higher percentage of type II muscle fibres (Hamada et al., 2000). Therefore we speculate that the increase in MYLK1 copy number will lead to increased expression of MYLK1 and subsequently increased phosphorylation of the RLC in skeletal muscle leading to elevated PAP. A potential reason that increased copy number is found more often in STP is that they are the thought to be the individuals with the greater distribution of type II muscle fibres and therefore will gain the greatest advantage from PAP.

4.3 MYLK-AS1

It was hypothesised that an increase in the MYLK-AS1 gene would lead to detriments in athletic performance. No associations were found between MYLK-AS1 copy number and athlete status or highest level of competition achieved. These findings indicate that alterations in MYLK-AS1 copy number do not confer any advantage or disadvantage to athletic performance. When the quantitative phenotype data for traits associated with strength and power are examined, it is shown that there are no positive or negative correlations with MYLK-AS1 copy number and these phenotypes. Taken together, these data indicate that alterations in MYLK-AS1 copy number is not involved in athletic performance related to strength and power. It has been shown that most CNVs have some form of effect on
expression which usually translates to alterations in the final protein content (Stranger et al., 2009; Tang & Amon, 2013). In the present study, we do not find any associations between MYLK-AS1 copy number and function; however that does not mean that there is no functional consequence. The results suggest that alterations in copy number of MYLK-AS1 does not influence skeletal muscle phenotypes related to strength and power. The lack of functional consequences of CNVs of MYLK-AS1 suggests that antisense genes may not always influence the function of the associated sense genes.

Antisense genes are found across a range of different organisms (Dahary et al., 2005) and at a higher frequency with genes which have multiple isoforms. It has been suggested that antisense transcription has been conserved through evolution to influence the splicing of the associated sense gene (Morrissy et al., 2011). There is some evidence that antisense genes can function to alter the splicing pattern of sense transcripts (Hastings et al., 1997; Annilo et al., 2009). Sense and antisense transcript interaction usually affects both the expression and splicing of the sense gene; however it has been shown for a small number of genes that antisense can affect only expression or splicing alone (Morrissy et al., 2011). The formation of double stranded RNA is the expected mechanism which leads to alternative splicing patterns. Antisense transcripts have been found to bind to the sense transcript to mask the splice sites of the THRA gene (Hastings et al., 1997). The findings suggest that it is a consequence of sequence overlap between the sense and antisense transcripts which causes the alternative splicing. To determine the importance of sequence overlap for antisense mediated splicing, it has been examined if regions with antisense overlap are more likely to reveal antisense mediated splicing than regions without overlapping sequences. There is a 2.5 fold increase in regions with overlapping sense and antisense sequences suggesting that overlapping sequences are an important factor in antisense mediated splicing (Morrissy et al.,
2011). As is shown in Figure 2, there is a region of overlap between the MYLK1 and MYLK-AS1 genes. Therefore, variation in MYLK-AS1 may not affect the expression of MYLK1 rather regulate the splicing pattern; hence there may not be any phenotypic consequence on muscle strength and power of CNVs of MYLK-AS1.

The MYLK1 gene encodes at least three products, including nonmuscle MYLK1, smooth muscle MYLK1 and Telokin, which are produced through the use of different promoters rather than alternative splicing (Herring et al., 2006). Alternative splicing has been shown to create multiple isoforms of nonmuscle MYLK1 transcripts (Lazar & Garcia, 1999; Hong et al., 2011). If MYLK1 sense and antisense gene interactions function to only affect the splicing of nonmuscle MYLK1 then this may explain why there are no functional impacts on muscle strength and power phenotypes. It is possible that the findings with MYLK1 copy number are because of increased expression of smooth muscle MYLK1 and interactions with skeletal muscle to affect the strength and power phenotypes. The alteration of MYLK-AS1 expression may influence the splice pattern of the nonmuscle MYLK1 isoform, therefore potentially having no impact on skeletal muscle strength and power phenotypes.

4.4 Summary

4.4.1 Combined findings

Overall, we have demonstrated that higher copy number of MYLK1 leads to associations with athlete status and improved performance in physiological tests associated with strength and power. The associations between copy number and quantitative phenotypes are weakened upon correction for subgroup which suggests that the type of training
performed by the athletes has an effect on the phenotypic consequences of MYLK1 copy number alterations. On assessment of the subgroups, we find only one control participant had a deviation in copy number and that there are no associations in END. Higher copy number in the STP is shown to associate with increased performance in tests which isolate specific muscle groups and tests which rely on whole body movements. Increased MYLK-AS1 copy number was hypothesised to be detrimental to muscle performance, however we fail to find any associations between copy number in this gene and either athlete status or performance in physiological tests associated with strength and power. The lack of association with athletic performance in MYLK-AS1 suggests that the function of the antisense gene is not only the regulation of MYLK1 expression.

The current study has found that there are different phenotypic consequences between CNVs of MYLK1 and MYLK-AS1. Despite the sequence of the two genes overlapping (Figure 2), an alteration in copy number in one of the genes is not always met with alteration in copy number in the other. The different phenotypic consequences of copy number on the target genes indicates different copy number events must be detected in the two genes, otherwise the associations between athlete status and quantitative phenotypes would be identical. It was originally hypothesised that if there was a gain of MYLK1 or loss of MYLK-AS1 that this would increase MYLK1 expression and result in increased strength and power and the opposite would be true of a loss in MYLK1 or gain in MYLK-AS1 copy number. We only detect these associations with MYLK1 which suggests that the expected alteration of MYLK1 expression in skeletal muscle by deviations of copy number does not occur in MYLK-AS1. These findings suggest that the copy number variants investigated affect separate mechanisms for MYLK1 and MYLK-AS1.
It is thought that higher MYLK1 copy number exerts functional effects on athletic performance through increased expression of MYLK1. We suggest that the increased expression of MYLK1 will lead to increases in strength and power phenotypes through a compensatory mechanism between MYLK1 and MYLK2. This compensation is thought to exist because of the ubiquitous expression of MYLK1 in all types of tissue and evidence there is still RLC phosphorylation in skeletal muscle of mice with ablated MYLK2. It has been shown that phosphorylation of the RLC leads to increased rate of force production and force potentiation. The lack of association with MYLK-AS1 is thought to be explained by the fact that not all antisense genes are involved in gene expression. It has been shown that some antisense genes function to alter the splicing of the associated sense gene (Morrissy et al., 2011). With evidence of alternative splicing of MYLK1 to create several isoforms of the nonmuscle MYLK1 protein (Hong et al., 2011), we suggest that potentially the role of the MYLK-AS1 gene may be to control the splicing and therefore CNVs of this gene would be unlikely to affect skeletal muscle phenotypes.

The results of the present study have identified, for the first time, associations between CNVs and muscle performance in an athletic cohort. These results taken with previous SNP data on MYLK1 (Clarkson et al., 2005)suggest that variation in this gene is functionally involved in the determination of muscle strength and power in both athletic and clinical populations. Replication of these findings in other cohorts and determination of the mechanisms behind these associations are required to understand the health implications of MYLK1 genotypes.
4.4.2 Strengths

The inclusion of different athletic subgroups in the GELA cohort allowed the investigation of the impact of CNVs on different types of training. The quantitative phenotype data both of whole body movements and isolated muscles allowed the examination of the functional effects these variants can have on performance. Identification of specific aspects of performance that are affected by CNVs allows the inference of potential mechanism underlying the associations between gene copy number and performance. The use of the multicopyType-it CNV Reference Probe assay instead of a single reference gene adds to the validity of the copy number determination in the current study. The Database of Genomic Variants (MacDonald et al., 2014) has shown that the typical reference genes TERT and RPPH1 have whole gene spanning CNVs which could affect copy number determination in target genes (Conrad et al., 2010; Xu et al., 2011). If there is a CNV on one of the loci of the Type-it CNV reference assay then the copy number determination will only be altered by a fraction, therefore, the effect will be greatly reduced compared to traditional reference assays. The use of fluorescently labelled probes rather than fluorescent dyes increases the reliability of copy number determination because it has been determined that fluorescent dyes can overestimate copy number in long amplicons and AT rich sequences (Colborn et al., 2008).

4.4.3 Limitations

As with all other studies there are limitations associated with the present study. It has been shown that somatic mosaicism of CNVs can occur where within different tissues there is a different copy number (Piotrowski et al., 2008). The CNVs were examined in blood leucocytes, however the tissue that we are interested in is skeletal muscle. If there are
discrepancies in copy number of MYLK1 between leucocytes and skeletal muscle cells, then it would be difficult to detect associations with strength and power phenotypes. Such discrepancies could also suggest that a non-skeletal muscle mechanism causes the associated increases in power and strength. While the investigation of muscle specific mechanisms would undoubtedly offer an interesting insight to the relationship between CNV and performance, the invasive nature of muscle biopsies can lead to the loss of training, which can make it difficult to recruit athletes. Thus, collection of skeletal muscle biopsies was not possible with the current cohort. Nonetheless, by using, blood samples we were able to study a much larger cohort than would otherwise have been possible which provides greater power to detect genomic variants.

4.4.4 Future work

Future studies should aim to replicate the findings of this study in a different cohort of athletes and controls to validate the findings in MYLK1 copy number with associations to strength and power phenotypes. These studies should also examine other CNVs in genes which have been previously associated with muscle phenotypes to identify if other CNVs are associated with athletic performance. Studies should also include measures of the expression of target genes in skeletal muscle to confirm if increases in copy number are met with concurrent increases in expression. The inclusion of the expression data will provide some insight to potential mechanisms which underlie the athletic performance phenotypes. Once replication of MYLK1 and CNV of other genes have been investigated for associations with muscle performance, these polymorphisms can potentially be used to identify individuals at risk of developing muscle wasting disorders and metabolic diseases and provide them with personalised plans to maximise strength to improve quality of life.
Chapter 5 - Conclusion

In conclusion, CNVs of MYLK1 are found at a higher frequency in STP than in non-athlete controls. These variants of MYLK1 copy number were found to associate with quantitative phenotypes associated with strength and power. These associations explain more of the variance in the STP than in the cohort as a whole (~7% and ~3% respectively for variance in elbow extension peak torque). Conversely, it was thought that MYLK-AS1 would associate with decrements in performance; however, the presence of CNVs of MYLK-AS1 does not associate with any of the measures of strength and power phenotypes in this study. It is thought that the increases with MYLK1 copy number will lead to concurrent increases in the expression of MYLK1. The mechanisms underlying the functional changes in muscle strength and power phenotypes cannot be identified in the present study. The mechanism could potentially be an increased expression of smMLCK which could affect RLC phosphorylation in skeletal muscle to increase the rate of force development and peak force. The unknown function of the antisense gene in the regulation of protein-coding genes could be the reason that the alteration of MYLK-AS1 copy number does not influence strength and power phenotypes as was hypothesised. The result of the present study furthers the current knowledge of how genetic variants underpin muscle performance. Further studies are needed to confirm that variation in MYLK1 is associated with increased strength and power. If these findings can be replicated, MYLK1 could potentially be used as a target to maintain an individual’s strength for improvements in athletic performance and the treatment of muscle wasting disorders.
References


Boonpeng H & Yusoff K (2013). The utility of copy number variation (CNV) in studies of hypertension-related left ventricular hypertrophy (LVH): rationale, potential and challenges. Mol Cytogenet 6, 8.


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**Appendix A** – The total number of athletes and number of athletes within each level of competition who perform each of the sports.

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Appendix B – Details on the physiological tests selected for phenotype determination

**Handgrip strength**

General upper body strength was tested with a commonly used handgrip test. A Hydraulic Hand Dynamometer (Model J00105, Lafayette Instrument Company, Lafayette, IN) was used to measure grip strength. Subjects were seated at the table with knees and elbow of the tested arm flexed to 90°. Each subject was permitted three trials with each hand. The dynamometer was adjusted to the individual hand size, and before each trial subjects used a towel to dry their palms to ensure a good grip. On each attempt, subjects were required to squeeze the dynamometer with maximal effort for two seconds. A rest period of 30 s was allowed between trials, and the best of the trials was recorded for each hand and totalled together to generate the total handgrip strength.

**Isokinetic dynamometry.** Knee and elbow flexion and extension peak torques were measured on isokinetic dynamometer (Biodex Pro3, USA) at 30, 90 and 180 °/s. Each set consisted of 3 repetitions of consecutive flexion/extensions cycles of maximal efforts throughout the range of motion. Sets always started with 30 °/s, and were followed by 90 °/s and then 180 °/s. Each of the sets was separated by a 90 s rest period, subjects started with right leg, followed by left leg, then right arm and left arm.

**Pull-ups.** A standard rigid gymnastic bar (2.5 cm diameter) at a height of 2.6 m from the ground was used. Participants were instructed to give their maximum efforts to perform as
many pull-ups as possible at their preferred speed but without passively hanging on the bar. The test was carefully supervised by a trained individual.

**Counter movement jump and static jump.** Performing counter movement jump, the participant were required to squat down quickly to 90° angle at the knee and immediately jump up as high as possible. Performing static jump, after a 3–5 s at still squat with 90° knee angle the participant had to jump as high as possible without any additional drop in the center of gravity. Counter movement jump attempts always preceded static jumps attempts. During jumps of both modes, arms were held akimbo, and the knee angles as well as the vertical modes of the jumps were inspected visually by a single researcher. A rest period of at least 30 s between jumps was used. Jumps were performed on a portable multicomponent force platform (Kistler, type 9286AA, Switzerland), which measured the vertical component of ground reaction force at a frequency of 1000 Hz. Kistler BioWare software, with Performance module (version 3.06c), was used to collect and analyze force data during the jumps. Height (cm) of the jump was calculated using the following formula: Height = 122.625 \( \times (T_f)^2 \), where \( T_f \) = flight time (s). After practicing for 2 trials on each jump mode, three test trials of each mode were allowed, and the highest jump of each mode was taken for subsequent analysis.

**30 m sprint.** The sprint run was performed on a smooth hard surface. Four fixed field (without reflectors) electronic photocells (NewTest Powertimer; Oulu, Finland) for time measurement were positioned exactly at 0 m, 10 m, 20 m and 30 m at a height of 1 m, i.e., time for three consecutive 10 m sprint runs were recorded with an accuracy of 0.001 seconds. Three trials were allowed, with the recovery of 2-3 min in between. Data of the best 30 m performance were taken for the analysis.
Appendix C – Concentrations of the DNA samples used in the present study

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Appendix D - Pearson's correlation coefficient results between END for MYLK1 copy number and quantitative variables. IKR; isokinetic right, IKL; isokinetic left

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<td>0.79</td>
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<tr>
<td>IKR elbow extension torque (90 °/s)</td>
<td>0.537</td>
<td>-0.073</td>
<td>0.53</td>
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</tr>
<tr>
<td>IKL elbow extension torque (90 °/s)</td>
<td>0.509</td>
<td>0.078</td>
<td>0.61</td>
<td></td>
</tr>
<tr>
<td>IKR elbow extension torque (180 °/s)</td>
<td>0.731</td>
<td>-0.041</td>
<td>0.16</td>
<td></td>
</tr>
<tr>
<td>IKL elbow extension torque (180 °/s)</td>
<td>0.766</td>
<td>0.035</td>
<td>0.12</td>
<td></td>
</tr>
<tr>
<td>IKR elbow flexion torque (30 °/s)</td>
<td>0.704</td>
<td>0.045</td>
<td>0.20</td>
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<tr>
<td>IKL elbow flexion torque (30 °/s)</td>
<td>0.805</td>
<td>-0.029</td>
<td>0.09</td>
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<tr>
<td>IKR elbow flexion torque (90 °/s)</td>
<td>0.542</td>
<td>-0.072</td>
<td>0.52</td>
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<tr>
<td>IKL elbow flexion torque (90 °/s)</td>
<td>0.366</td>
<td>-0.107</td>
<td>1.14</td>
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<tr>
<td>IKR elbow flexion torque (180 °/s)</td>
<td>0.701</td>
<td>-0.045</td>
<td>0.21</td>
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</tr>
<tr>
<td>IKL elbow flexion torque (180 °/s)</td>
<td>0.562</td>
<td>-0.068</td>
<td>0.47</td>
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</tbody>
</table>
Appendix E – Increases in performance per copy number in the whole cohort for quantitative phenotypes which displayed significant correlations after BH corrections. IKR: isokinetic right, IKL; isokinetic left.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Copy number test</th>
<th>p-value</th>
<th>Percentage Change (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Static jump height (cm)</td>
<td>2 vs 1</td>
<td>0.248</td>
<td>17.98</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.036</strong></td>
<td><strong>11.81</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td><strong>0.041</strong></td>
<td><strong>31.92</strong></td>
</tr>
<tr>
<td>Sprint 0-10 (s/kgFFM)</td>
<td>2 vs 1</td>
<td>0.315</td>
<td>-2.41</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.015</strong></td>
<td><strong>-10.04</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.113</td>
<td>-12.21</td>
</tr>
<tr>
<td>Sprint 10-20 (s/kgFFM)</td>
<td>2 vs 1</td>
<td>0.722</td>
<td>-1.16</td>
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<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.008</strong></td>
<td><strong>-11.54</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.139</td>
<td>-12.52</td>
</tr>
<tr>
<td>Sprint Total (s/kgFFM)</td>
<td>2 vs 1</td>
<td>0.661</td>
<td>-1.11</td>
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<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.010</strong></td>
<td><strong>-10.97</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.123</td>
<td>-11.96</td>
</tr>
<tr>
<td>I KL knee extension torque (90 °/s)</td>
<td>2 vs 1</td>
<td>0.447</td>
<td>13.97</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.016</strong></td>
<td><strong>14.97</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.148</td>
<td>31.04</td>
</tr>
<tr>
<td>I KR elbow extension torque (30 °/s)</td>
<td>2 vs 1</td>
<td>0.994</td>
<td>0.18</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.002</strong></td>
<td><strong>28.86</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.220</td>
<td>29.10</td>
</tr>
<tr>
<td>I KL elbow extension torque(30 °/s)</td>
<td>2 vs 1</td>
<td>0.676</td>
<td>7.43</td>
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<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.006</strong></td>
<td><strong>26.42</strong></td>
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<tr>
<td></td>
<td>3 vs 1</td>
<td>0.110</td>
<td>35.81</td>
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<tr>
<td>I KR elbow extension torque(180 °/s)</td>
<td>2 vs 1</td>
<td>0.879</td>
<td>-2.30</td>
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<tr>
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<td>3 vs 2</td>
<td><strong>0.001</strong></td>
<td><strong>37.62</strong></td>
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<tr>
<td></td>
<td>3 vs 1</td>
<td>0.181</td>
<td>34.45</td>
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<tr>
<td>I KR elbow flexion torque (30 °/s)</td>
<td>2 vs 1</td>
<td>0.949</td>
<td>0.90</td>
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<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.005</strong></td>
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<tr>
<td></td>
<td>3 vs 1</td>
<td>0.195</td>
<td>22.78</td>
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<tr>
<td>I KL elbow flexion torque (180 °/s)</td>
<td>2 vs 1</td>
<td>0.810</td>
<td>-1.41</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td><strong>0.002</strong></td>
<td><strong>27.47</strong></td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.205</td>
<td>25.67</td>
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</table>
Appendix F – Increases in performance per copy number in STP for quantitative phenotypes which displayed significant correlations after BH corrections.

<table>
<thead>
<tr>
<th>Trait</th>
<th>Test</th>
<th>p-value</th>
<th>Percentage Change (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Static jump height (cm)</td>
<td>2 vs 1</td>
<td>0.012</td>
<td>35.49</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td>0.233</td>
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<tr>
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<td>3 vs 1</td>
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<td>IKRA, extension 180d/sec., N*mLOG10</td>
<td>2 vs 1</td>
<td>0.350</td>
<td>14.54</td>
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<tr>
<td></td>
<td>3 vs 2</td>
<td>0.004</td>
<td>28.33</td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.106</td>
<td>46.99</td>
</tr>
<tr>
<td>IKLA, flexion 180d/sec., N*mLOG10</td>
<td>2 vs 1</td>
<td>0.354</td>
<td>14.45</td>
</tr>
<tr>
<td></td>
<td>3 vs 2</td>
<td>0.010</td>
<td>25.07</td>
</tr>
<tr>
<td></td>
<td>3 vs 1</td>
<td>0.069</td>
<td>43.14</td>
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</tbody>
</table>