The \textit{ACTN3} R577X variant in sprint and strength performance

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INTRODUCTION

Muscle power is defined as high-intensity muscular performance in a short term and is composed of speed and strength. Muscle power and its components are determined by environmental and genetic factors. Both familial and twin studies suggest that genetic factors in power, speed and strength account for 35\%–80\% of inter-individual variation [6,28-30]. During the last two decades, there has been considerable interest in finding the genetic factors responsible for human performance. It has recently been reported that some candidate genes are related to human muscle power [19,20]. Among them, \textit{ACTN3} is one of the most interesting genes in that the \textit{ACTN3} variant (\textit{ACTN3} R577X SNP) directly determines the expression of \textalpha-actinin-3 protein that contributes to the construction of the contractile component in power-generating fast twitch fibers of the skeletal muscle [2,23]. In addition, it has been reported that the functional individual difference of \textalpha-actinin-3 protein expressed only in sarcomere of fast-twitch muscle fibers is not to be decided by multiple gene (170 counts) diversity coding \textalpha-actinin-3 but by \textit{ACTN3} R577X diversity such as dominant-recessive effect [16,18].

\textit{ACTN3}, the gene for encoding \textalpha-actinin-3, is located at 11q13-q14. This gene has a 1747C>T transition within exon 16, which results in the \textit{ACTN3} R577X SNP, giving three genotypes: the RR, RX and XX genotypes. In the \textit{ACTN3} variant (R577X SNP), the XX genotype deficits \textalpha-actinin-3 due to the premature stop codon without a pathological phenotype, whereas the RR and RX genotypes express \textalpha-actinin-3. As a result, the \textit{ACTN3} variant directly determines the expression of \textalpha-actinin-3 [1,18]. An important structural component of the Z disc is \textalpha-actinin-3, where it anchors actin thin filaments, helping to maintain the myofibrillar array only in fast twitch muscle fibers that are responsible for high velocity and force for power-generating contractions [1,2,18]. It has been suggested that the \textit{ACTN3} variant for the expression or deficiency of \textalpha-actinin-3 in fast-twitch muscle fibers should influence the power-generating muscle function [2,23].

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Several studies have found that the ACTN3 variant affects speed/power performance [5,7,17,20,24,27,32]. The R allele and the RR genotype, expressing α-actinin-3, are overrepresented whereas the XX genotype for α-actinin-3 deficiencies is underrepresented in sprint events, suggesting that α-actinin-3 is required for sprint/power performance. The α-actinin-3 deficiency has been associated with poorer sprinting performance in non-athletes. Experimental studies using ACTN3 KO mice have also shown that the absence of α-actinin-3 results in fast-twitch, glycolytic fibers developing slower-twitch and poorer power generation [4,15,19], supporting the responsiveness of the α-actinin-3 expression to optimal sprint and power performance. These results show that the ACTN3 variant is somewhat responsible for speed performance and also for determining some aspects of muscle power.

However, previous research on the ACTN3 variant focused mainly on speed performance without differentiating its influence on the power components of speed and strength. Because muscle power is the high-strength muscular performance that occurs over a short time period, it is composed of speed and strength each of which can be differentiated independently and can also be determined by environmental and genetic factors [6,28,29,30]. There is, however, a dearth of information on the association of strength, speed, or power with the ACTN3 variant. Only association of power with the ACTN3 variant and association of speed with the ACTN3 variant are reported [14,31]. Considering that the expression of sarcomeric α-actinin-3 is located at the fast-twitch fiber that generates strong, high-velocity muscle contraction, there is a strong need to understand whether α-actinin-3 is expressed or not and whether it should be associated with forceful contraction, i.e. muscle strength, for giving explosive power.

Therefore, the aim of this study is to examine the distribution of the ACTN3 R577X genotypes and alleles in power-oriented, speed-oriented and strength-oriented athletes and to search for differential associations of the power components with the ACTN3 R577X polymorphism.

**METHODS**

**Participants**

All participants in this study were recruited from August, 1996 to December, 2011. All athletes in this study were members of the Korean National team and university elite athletes and the control group consisted of non-athletes participating in national physical fitness survey and national health promotion business. Athletic participants in this study consisted of 849 elite athletes that play 11 different sports including fencing, skating, track & field, swimming, wrestling, boxing, cycling, weightlifting, gymnastics, field hockey and judo. Only weightlifters, sprinters (≤400 m), speed skaters (≤1,500 m) and swimmers (≤100 m) were selected for this study since these sports are mainly focused on strength, speed and power. The weightlifters are considered to be strength-oriented athletes while the speed skaters and swimmers are considered to be speed-oriented athletes [3,21,26]. All three groups (weightlifters, speed skaters and swimmers) belong to the power-oriented category [3,21,26]. As many as 121 top-level athletes participated in this study and their athletic performance records were confirmed by the Korea Sports Council.

A total of 975 Koreans aged 18-39yrs, and composed of healthy controls (n = 854), top-level strength-oriented athletes (n = 63), and top-level speed-oriented athletes (n = 58) were included in this study (see Table 1). Power-oriented top-level athletes (n = 121) were collectively assigned by merging the top-level strength-oriented athletes and the top-level speed-oriented athletes. The 854 healthy controls (435 men, 419 women) met the following inclusion criteria: (1) apparently healthy people (2) no known diseases such as neuromuscular disease, cardiovascular disease or metabolic problems (3) native Korean aged 18-39yrs and (4) no participation in athletic events. The top-level strength-oriented athletes were selected based on their having represented Korea at the international level or on their having won an international championship in weightlifting. Of the final 63 weightlifters (37 men, 26 women) selected as top level strength-oriented athletes for this study, 23 had won medals at the World Championships and Olympic Games. The top-level speed-oriented athletes were selected along the same lines i.e. outstanding performance in sprint events and participation in top international competitions and/or Olympic Games as

**Table 1. Characteristics of the top-level elite athletes, and controls**

| Sport Event | age | Career* |
|-------------|-----|---------|
| Running     | 22.2 ± 3.6 | 21.2 ± 4.3 |
| Speed skating | 3.4 ± 2.9 | 2.9 ± 2.1 |
| Swimming  | 20.8 ± 4.6 | 21.2 ± 4.3 |
| Strength-oriented athletes | 63 | ≤100 m, n = 58 | 2.9 ± 2.1 | Running (n = 58) |
| Speed-oriented athletes | 58 | ≤400 m, n = 20 | 3.4 ± 2.9 | Running (n = 20) |
| Power-oriented athletes | 121 | ≤22.2 ± 3.6 | 2.9 ± 2.1 | Above (n = 121) |
| Control (n = 854) | 32.6 ± 4.8 | - |

Data are mean and standard deviation (within parentheses). *P-Career as the international athlete, a: the power-oriented athletes are composed of strength-oriented athletes and speed-oriented athletes.
Table 2. The distribution of the ACTN3 R577X genotype and the allele frequency in the top-level power-oriented athletes, and controls

| Group                          | Genotype | Allele |
|-------------------------------|----------|--------|
|                               | RR       | RX     | XX     | \(P^2\) | R      | X     | \(P^2\) |
| Control (\(n = 854\))         | 255 (29.9) | 436 (51.1) | 163 (19.1) | 946 (55.4) | 762 (44.6) |
| Power-oriented (\(n = 121\))  | 49 (40.5)  | 58 (47.9)  | 14 (11.6)  | .025 | 156 (64.5) | 86 (35.5) | .008 |

Data are absolute and relative values (within parentheses). \(P\)-value for the \(\chi^2\) test or Fisher’s Exact Test in comparison to controls. a: the power-oriented athletes are composed of strength-oriented athletes and speed-oriented athletes.

Table 3. The distribution of the ACTN3 R577X genotype and the allele frequency in the top-level sprinters, weightlifters, and controls

| Group                          | Genotype | Allele |
|-------------------------------|----------|--------|
|                               | RR       | RX     | XX     | \(P^2\) | R      | X     | \(P^2\) |
| Control (\(n = 854\))         | 255 (29.9) | 436 (51.1) | 163 (19.1) | 946 (55.4) | 762 (44.6) |
| Strength-oriented (\(n = 63\)) | 25 (39.7)  | 28 (44.4)  | 10 (15.9)  | .262 | 78 (61.9) | 48 (38.1) | .155 |
| Speed-oriented (\(n = 58\))   | 24 (41.4)  | 30 (51.7)  | 4 (6.9)    | .026 | 78 (67.2) | 38 (32.8) | .013 |

Data are absolute and relative values (within parentheses). \(P\)-value for the \(\chi^2\) test or Fisher’s Exact Test in comparison to controls.

representative Korean sprinters. Of the 58 participants (37 men, 21 women) selected as top-level speed-oriented athletes, 18 were speed skaters (≤1,000 m), 21 were sprinters (≤400 m) and 19 were swimmers (≤100 m). Additionally 19 of them had won medals in their special area either in the Asia or World Championships or the Olympic Games. Written informed consent was obtained from all subjects under protocols approved by the Institutional Review Boards of the Eulji University School of Medicine.

**Genotype analysis**

Genomic DNA was isolated from peripheral blood cells using a QIAamp blood kit (Qiagen, Valencia, CA, USA) according to the manufacturer’s protocols. The SNP of the ACTN3 R577X (rs1815739) was analyzed with custom-designed primers and probes (Assay ID: C_590093_1_) for the MGB TaqMan® SNP genotyping assay (Applied Biosystems, Foster City, CA, USA). The primers are Forward: 5’-ACGATCGTTCAAGGGAACACT-3’ and Reverse: 5’-ACCCCTGGATGGCATGATG-3’. Allele-specific probes were labeled with the fluorescent dyes VIC (5’-TCGTCTCGGTCAGC-3’) and FAM (5’-CGCTCTCAGTCAGC-3’). RCR was carried out in a total reaction volume of 20\(\mu\)L containing 200ng of template, 10\(\mu\)L TaqMan Universal PCR Master Mix No AmpErase UNG (2x) and 0.5\(\mu\)L 40xSNP Genotyping Assay with the following amplification protocol: denaturation at 95°C for 10 minutes, followed by 40 cycles of denaturation at 92°C for 15 seconds and finally annealing and extension at 60°C for 1 minute. Post PCR, the genotype of each sample was attributed automatically by measuring the allele-specific fluorescence on the ABI PRISM® 7900 Sequence Detection System using SDS 2.3 software for allelic discrimination (Applied Biosystems, Foster City, CA, USA). Duplicate samples and negative controls were included to ensure accuracy of the genotyping.

**Statistical analysis**

PAWE (Power for Association with Error) software program was used to test sample size validity and the sample size validity was confirmed by having type I error set at 0.05 and type II error as 80% power level. The ACTN3 allele and genotype frequencies were obtained by direct count. The Hardy-Weinberg equilibrium (HWE) for genotype distribution of each group was then estimated by the chi-square test. Differences in the allele frequencies and genotypes between the controls and the power-oriented athletes, between the controls and the top-level strength athletes and between the controls and the top-level sprinters were compared using the chi-square test or Fisher’s exact test based on the presence of less than five frequencies in one cell. Trend analysis for the genotype distribution between the controls and the power-oriented, top-level strength and speed groups was conducted by the linear-by-linear association analysis. All analyses were conducted using SPSS statistical software (Version 13.0; SPSS Inc., Chicago, IL). A two-tailed significance at .05 level was chosen to test for type I errors.

**RESULTS**

A combination of control individuals and top-level power-oriented athletes, divided into top-level weightlifters (strength-oriented) and sprinters (speed-oriented), was analyzed in this study. Table 1 shows the ACTN3 allele and genotype frequen-
cies from the control and power-oriented groups. The ACTN3 genotype distributions in the control group and the power-oriented group were in Hardy-Weinberg equilibrium ($\chi^2_{[df=1]} = 0.934$ for the control, $\chi^2_{[df=1]} = 0.258$ for the power-oriented group; $p > 0.10$). The ACTN3 genotype and allele distribution in the power-oriented athletes showed significant differences when compared with the controls ($\chi^2_{[df=2]} = 7.347$, $P < 0.05$ for genotype; $\chi^2_{[df=1]} = 7.105$, $P < 0.01$ for allele). The power-oriented group had an underrepresented frequency of the XX genotype (11.6% vs 19.1%) and the X allele (35.5% vs 44.6%) in comparison to the control group.

To test the association of power types (strength and speed performance) with the ACTN3 genotype, the power-oriented group was divided into the strength-oriented group (the top-level sprinters) and the speed-oriented group (the top-level sprinters). Both of them were in HWE ($\chi^2_{[df=1]} = 1.758$ for speed-oriented athletes, $\chi^2_{[df=1]} = 0.210$ for strength-oriented athletes; $p > 0.10$). As shown in Table 1, no statistical differences in the ACTN3 genotype and allele distributions were found between the strength-oriented athletes and the controls ($\chi^2_{[df=2]} = 2.682$, $P < 0.262$ for genotype; $\chi^2_{[df=1]} = 2.022$, $P = 0.155$ for allele). Only the speed-oriented athletes showed significant differences in the frequency distributions of the ACTN3 genotype (Fisher’s Exact Test = 7.252, $P < 0.05$) and the allele ($\chi^2_{[df=1]} = 6.200$, $P < 0.05$) from that of the controls. In Fig. 1, the frequency of the α-actinin-3 non-expression genotype (XX) was significantly reduced while the frequency of the α-actinin-3 expression genotype (RR + RX) was significantly increased in the speed-oriented athletes compared with the controls (Fisher’s Exact Test = 5.395, $P < 0.05$; Fig. 1). No significant difference in the α-actinin-3 expression genotype distribution was found in the strength-oriented athletes compared with the controls ($\chi^2_{[df=1]} = 0.396$, $P < 0.529$).

**DISCUSSION**

The aim of this study is to explore an association of the ACTN3 genotype with speed and/or strength performance. The ACTN3 genotype distribution of Korean top-level sprinters for speed, weightlifters for strength and mixed power-oriented athletes for power were compared to the ACTN3 genotype distribution of Korean adults. There was a significant association of the ACTN3 genotype with power performance as was expected. When dividing power performance into speed and strength performances, the ACTN3 genotype was only associated with speed performance but a relationship with strength performance was not identified. The ACTN3 XX genotype in deficit of the α-actinin-3 protein in fast-twitch muscle fibers had a significantly low distribution in the speed performance group, while the distribution of the ACTN3 XX genotype showed a somewhat low ratio compared to the control group in the strength performance group with no statistical significance. Therefore, speed performance among the power factors showed preference for the ACTN3 RR + RX genotype but not for the ACTN3 XX genotype.

The association of the ACTN3 genotype with speed performance in this study is similar to previous study results. Yang et al. [32] reported reduced distribution of the ACTN3 XX genotype in sprint/power groups, composed of 7 Australian Judo elite athletes and 42 sprinters. Niemi and Majamaa [17] and Papadimitriou et al. [20] reported remarkably low levels of the ACTN3 XX genotype in Greek sprinters. In these studies, the distributions of the ACTN3 XX genotype in the control groups were about 18% similar to the 19% distribution for the control group in our study. Also, the authors reported low XX genotype distributions for sprinters (6%, 0%, and 8.8%) compared to the control groups which agreed with our results of 6.9%. The ACTN3 genotype is favorable to sprint performance while the ACTN3 XX genotype lacking α-actinin-3 is unfavorable to sprint performance.

Druzhkevskaya et al. [7] and Papadimitriou et al. [20], studied Russian and Greek populations respectively and reported that the distributions of the ACTN3 XX genotype were low in the power-oriented performance group compared to their control groups. Santiago [27] also reported a reduced distribution of the XX genotype in Spanish soccer players who needed explosive power through a combination of...
strength and speed. In contrast, neither Nigerian nor Greek Track and Field athletes skilled in power-oriented performance [20,33] showed any significant differences in their ACTN3 XX genotype distribution compared to the control groups. Thus, research on the association of ACTN3 with power-oriented performance has not shown consistent results. This may come from differences in the genotype distribution between human populations [33]. It is well-known that the genetic structures of Caucasians and Africans are different, leading to some functionally different characteristics. Actually the XX genotype of the ACTN3 in Caucasian is ~19%, higher than the 10% found in Africans. In addition, the differences in the composition of speed and strength athletes skilled in power-oriented performance may be another cause of the inconsistent association of ACTN3 with power performance [16,18,33]. This study divided the ACTN3-associated power-oriented performance into speed performance and strength performance but the ACTN3 genotype distribution showed association only with speed performance and none with the strength performance. The coordination level of speed and strength affecting power-oriented performance may play an important role in determining the association between ACTN3 and power. As a result, the association of the ACTN3 R577X variant with power-oriented performance would not show consistency and a variant that distinguishes the components of power into speed and strength would be needed to find the association of the ACTN3 variant with athletic performance.

As there was no significant difference in the ACTN3 genotype distribution of Korean top-level weightlifters with that of the control group in this study, the ACTN3 variant had no significant association with strength performance. Because weightlifting requires a single maximal contraction of the skeletal muscle, it is essential for the skeletal muscle, especially for fast-twitch muscle fibers, to be hypertrophic [8,9,11,12]. It is expected that the distribution of the ACTN3 XX genotype was low because in the ACTN3 XX genotype, α-actinin-3 is structured in the Z-line of the fast-twitch muscle fiber that functions during rapid speed and strong power [13,31]. However, in contrast with this hypothesis, the ACTN3 XX genotype was not significantly reduced in weightlifters. Such a result corresponded to previous study results with isokinetic strength performance of active young men and women. There was no difference in the isokinetic torque and rate of decline of torque between ACTN3 genotypes [10]. These results including ours could support the contention that the lack of α-actinin-3 in fast-twitch muscle fibers by the XX genotype only slightly affects the muscular strength of a single maximum contraction. On the other hand, the null type of the α-actinin-3, ACTN3 XX genotype, showed a significantly low distribution in speed performance as consistently reported in previous studies and also in this study. Therefore, it is thought that the α-actinin-3 deficient genotype, the ACTN3 XX genotype, may affect speed performance that needs rapidly repeated muscular contractions. In the future, it is necessary to explore the mechanism that causes the difference in the contraction of fast-twitch muscles that is due to the lack of α-actinin-3. In the case where the lack of α-actinin-3 has no relation with the maximum muscular strength, there is also the possibility that another protein can compensate for this lack. This mechanism also requires further study.

CONCLUSIONS

The ACTN3 XX genotype that is in deficit of α-actinin-3 is confirmed to be strongly associated with speed performance that requires multiply repeated rapid muscle contractions. The ACTN3 XX genotype had no association with strength performance for single maximum muscle contractions among power components. Therefore, we found suggestive evidence that the ACTN3 R577X polymorphism plays a role in discriminating speed performance athletes from strength athletes and sedentary controls. In other words, the ACTN3 R577X variant is the main gene for speed performance but not for strength performance.

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