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**Are Genome-Wide Association Study Identified Single-Nucleotide Polymorphisms Associated With Sprint Athletic Status? A Replication Study With 3 Different Cohorts**

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### Article

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## Original Investigation

### Are GWAS-identified SNPs associated with sprint athletic status? A replication study with three different cohorts

**Running head:** Gene polymorphisms for top-level sprinters

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40 **Abstract**

41 **Purpose:** This study was aimed to replicate previously GWAS-identified sprint-related  
42 polymorphisms in three different cohorts of top-level sprinters and to further validate obtained  
43 results in functional studies. **Methods:** A total of 240 Japanese, 290 Russians and 593  
44 Brazilians were evaluated in a case–control approach. Of these, 267 were top-level  
45 sprint/power athletes. In addition, the relationship between selected polymorphisms and  
46 muscle fiber composition was evaluated in 211 Japanese and 287 Finnish individuals.  
47 **Results:** The G-allele of the rs3213537 polymorphism was overrepresented in Japanese (OR:  
48 2.07,  $P = 0.024$ ) and Russian (OR: 1.93,  $P = 0.027$ ) sprinters compared to endurance athletes  
49 and associated with increased proportion of fast-twitch muscle fibers in Japanese ( $P = 0.02$ )  
50 and Finnish ( $P = 0.041$ ) individuals. Meta-analysis of data from the four cohorts confirmed  
51 that the presence of the G/G genotype rather than G/A+A/A genotypes increased the odds  
52 ratio of being a sprinter compared to controls (OR: 1.54,  $P = 0.005$ ), endurance athletes (OR:  
53 1.79,  $P = 0.001$ ) or controls + endurance athletes (OR: 1.61,  $P = 0.001$ ). Furthermore, male  
54 sprinters with the G/G genotype were found to have significantly faster personal times in the  
55 100-m dash than those with G/A+A/A genotypes ( $10.50 \pm 0.26$  vs.  $10.76 \pm 0.31$ ,  $P = 0.014$ ).  
56 **Conclusion:** The rs3213537 polymorphism found in the *CPNE5* gene was identified as a  
57 highly replicable variant associated with sprinting ability and increased proportion of fast-  
58 twitch muscle fibers, in which the homozygous genotype for the major allele (i.e., the G/G  
59 genotype) is preferable for performance.

60

61 **Keywords:** athletes; copine-V; genetics; sprint performance; synaptic plasticity

62

## 63 **Introduction**

64 The sprint ability is a core capacity that underlies performance in many individual sports as  
65 well as team sports. Naturally, pure sprint athletes (e.g., 100-m runners) perform better on  
66 physiological and mechanical variables of sprint performance <sup>1</sup>. A velocity-oriented force–  
67 velocity profile is a major contributing factor for a better sprint performance <sup>2</sup>. The maximal  
68 sprint velocity and mean power produced over the event distance strongly influence  
69 performance <sup>3</sup>. During a sprint task, power output demand increases exponentially with  
70 velocity and the best sprinters accelerate over a longer distance than their lower performing  
71 counterparts <sup>4</sup>.

72 Sports performance is the combined result of numerous intrinsic and extrinsic factors,  
73 that is, the interaction between genetic factors and the environmental stimulus. Although  
74 training and other environmental stimulus are critical to performance achievement, individual  
75 performance thresholds can be determined by our genetic make-up. Twin studies have  
76 reported moderate to high heritability estimates for maximum movement speed as well as for  
77 other sprint and power phenotypes <sup>5, 6</sup> and so it has been proposed that elite sprint  
78 performance strongly depends on genetic characteristics.

79 Like other sports phenotypes, the sprint ability is a complex and polygenic  
80 phenomenon guided by the interaction of multiple genes and most likely gene variants. There  
81 are several polymorphisms that have been associated with elite power and sprint athletic  
82 status <sup>7</sup>. In particular, some of them were also associated with faster sprint times <sup>8,9</sup>; however,  
83 many of the polymorphisms suggested as favorable to sprinters were evaluated using case–  
84 control approaches that have not yet been replicated in subsequent studies or independent  
85 samples <sup>7</sup>. Replication studies are of paramount importance to better evaluate and characterize  
86 performance-relevant polymorphisms. The same association in independent samples indicates  
87 a greater relevance between the polymorphism and the target phenotype.

88 Recently, Pickering et al. <sup>10</sup> first performed a genome-wide association study (GWAS)  
89 to identify sprint-related genetic variants. These authors exposed a set of new polymorphisms  
90 associated with short-distance sprints in youth football players, some of which were replicated  
91 in an independent cohort of Polish women. The replication of these findings in top-level  
92 athletes of different ethnicities would be interesting, since only one cohort of Russian athletes  
93 validated the most associated polymorphisms.

94 Therefore, the purpose of this study was to replicate GWAS-identified sprint-related  
95 polymorphisms in three different cohorts of top-level sprinters. A secondary purpose of this  
96 study was to evaluate the relationship between these polymorphisms and the proportion of  
97 fast-twitch muscle fibers in two different cohorts. First, the selected polymorphisms were  
98 evaluated for sprinter athletic status and proportion of fast-twitch muscle fibers in a Japanese  
99 cohort. Subsequently, the most consistent polymorphism was evaluated for sprinter athletic  
100 status in two other cohorts from Russia and Brazil, and evaluated for proportion of fast-twitch  
101 muscle fibers in Finnish individuals. Since the target phenotype is the sprint ability, sprinters  
102 were compared to non-athletes (controls) or endurance athletes (the metabolic demands  
103 required to perform sprint or endurance events are opposites of each other).

## 104 **Methods**

105 Table 1 shows the polymorphisms selected for use in this study. All of them are single  
106 nucleotide polymorphisms (SNPs) and were selected based on a previous study <sup>10</sup> and  
107 according to the following criteria: biallelic polymorphisms located on autosomal  
108 chromosomes, two replications in the initial study and minor allele frequency > 1% in the  
109 Japanese population. Although the rs12688220 and rs8064257 polymorphisms also showed  
110 two replications in the initial study, they were not included because they did not meet the  
111 inclusion criteria.  
112

113 All cohorts included in this study had their procedures conducted according to the  
114 Declaration of Helsinki ethical principles for research involving human subjects. The  
115 Japanese studies were approved by the ethics committee of the Juntendo University and  
116 Fukuoka University. The Finnish study was approved by the coordinating ethics committee of  
117 the Hospital District of Helsinki and Uusimaa (this data was used with permission; Database  
118 of Genotypes and Phenotypes (dbGaP) Study Accession: phs000867.v1.p1). The Russian  
119 study was approved by the ethics committee of the Federal Research and Clinical Center of  
120 Physical-chemical Medicine of the Federal Medical and Biological Agency of Russia. The  
121 Brazilian study was approved by the ethics committee of the School of Physical Education  
122 and Sport, University of Sao Paulo, São Paulo, Brazil. A written informed consent was  
123 obtained from each participant.

124

### 125 **The Japanese Cohort**

126 The Japanese study involved 114 athletes (91 males and 23 females), of which 54 were  
127 sprint/power athletes (100-400 m runners, jumpers and throwers; mean age  $\pm$  SD:  $28 \pm 7$   
128 years) and 60 endurance runners (800 m to marathon; mean age  $\pm$  SD:  $24 \pm 3$  years). All of  
129 these athletes were international-level competitors. The control group comprised 126 healthy  
130 Japanese individuals.

131 Total DNA was isolated from saliva or venous blood using the Oragene • DNA  
132 Collection Kit (DNA Genotek, Ontario, Canada) or the QIAamp DNA blood Maxi Kit  
133 (QIAGEN, Hilden, Germany), respectively. The total DNA content was measured using the  
134 NanoDrop 8000 spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA).  
135 Subsequently, DNA samples were adjusted to a concentration of 50 ng/ $\mu$ L with Tris-EDTA  
136 buffer and stored at 4°C. Total DNA samples were genotyped using the HumanOmniExpress  
137 Beadchip (Illumina, San Diego, CA, USA) to genotype  $> 700,000$  SNPs, according to the  
138 manufacturer's instructions. Genotype calls were performed with Illumina GenomeStudio  
139 software and PLINK was used for quality control checks and association analyses.

140

### 141 *Evaluation of skeletal muscle fiber types*

142 First, a cohort of 203 Japanese healthy individuals (98 men and 105 women, with age range  
143 20-79 years) performed muscle biopsy was used for the association study between sprint-  
144 related polymorphisms and muscle fiber composition. Muscle samples were obtained from the  
145 belly of the vastus lateralis and myosin heavy chain (MHC) isoforms were determined by  
146 performing glycerol SDS-PAGE, as previously described <sup>11</sup>. These individuals had their DNA  
147 samples isolated from venous blood and the polymorphisms were genotyped using the  
148 Japonica SNP array <sup>12</sup>.

149 Second, muscle fiber composition in 287 Finnish individuals (167 men, age  $59.5 \pm 8.1$   
150 years; 120 women, age  $60.7 \pm 7.4$  years) from the FUSION study was estimated based on the  
151 expression of the myosin heavy chain 1 (*MYH1*), myosin heavy chain 2 (*MYH2*), myosin  
152 heavy chain 7 (*MYH7*), Ca<sup>2+</sup> ATPase A1 and Ca<sup>2+</sup> ATPase A2 genes, as previously described  
153 <sup>13</sup>. Muscle samples were obtained from the vastus lateralis using a conchotome, under local  
154 anesthesia with 20 mg·ml<sup>-1</sup> lidocaine hydrochloride without epinephrine <sup>14</sup>. DNA samples  
155 were extracted from the blood and the polymorphisms were genotyped using the  
156 HumanOmni2.5-4v1\_H BeadChip array (Illumina, San Diego, CA, USA).

157

### 158 **The Russian Cohort**

159 The Russian study involved 173 athletes (99 males and 74 females; mean age  $\pm$  SD:  $31.3 \pm$   
160  $7.5$  years), of which 70 were elite sprinters (100-400 m runners, 500-1000 m speed skaters, 50  
161 m swimmers) and 103 elite endurance athletes (biathletes, rowers, cross-country skiers, 3-10  
162 km runners, 800-1500 m swimmers and triathletes). All of these athletes were international-

163 level competitors, of which 30 (13 sprinters and 17 endurance athletes) were highly elite  
164 athletes (i.e., prize winners in international competitions). The control group comprised 117  
165 healthy unrelated citizens (66 males and 51 females, mean age  $\pm$  SD:  $47.9 \pm 4.8$  years),  
166 without any competitive sport experience. This Russian cohort is independent of the one  
167 previously published<sup>10</sup>.

168 Molecular genetic analysis was performed with DNA samples obtained from  
169 leukocytes (venous blood). Four millilitres of venous blood was collected in tubes containing  
170 EDTA (Vacuette EDTA tubes; Greiner Bio-One, Kremsmünster, Austria). Blood samples  
171 were transported to the laboratory at 4°C, and DNA was extracted on the same day. DNA  
172 extraction and purification were performed using a commercial kit according to the  
173 manufacturer's instructions (Technoclon, Moscow, Russia), which included chemical lysis,  
174 selective DNA binding on silica spin columns and ethanol washing. Extracted DNA quality  
175 was assessed by agarose gel electrophoresis. The genotyping process was performed using  
176 HumanOmni1-Quad BeadChips or HumanOmniExpress BeadChips (Illumina, San Diego,  
177 CA, USA) to genotype > 900,000 SNPs. The assay required 200 ng of DNA sample as input  
178 with a concentration of at least 50 ng/ $\mu$ l. Exact concentrations of DNA in each sample were  
179 measured using a Qubit Fluorometer (Invitrogen, Waltham, MA, USA). All further  
180 procedures were performed according to the instructions of the Infinium High-Density Assay.

181

### 182 **The Brazilian Cohort**

183 The Brazilian study involved 305 athletes (200 males and 105 females; mean age  $\pm$  SD:  $25.4$   
184  $\pm 6.9$  years), of which 143 were elite sprinters (100-400 m runners, 50-200 m swimmers,  
185 canoeing and cycling) and 162 endurance athletes (rowers, > 1.5 km runners, 400-1500 m  
186 swimmers and triathletes). While 36% of these athletes were nationally prominent  
187 competitors, 64% were international-level competitors. The control group comprised 288  
188 healthy Brazilian individuals (187 males and 101 females, mean age  $\pm$  SD:  $29.6 \pm 8.1$  years),  
189 without any competitive sport experience.

190 Genomic DNA of the Brazilian participants was isolated from buccal epithelial cells  
191 obtained from mouthwashes with a 0.9% saline solution prepared with DNA- and DNase-  
192 free water as previously described<sup>8</sup>. Briefly, the DNA samples were extracted using  
193 chloroform, precipitated using ethanol and resuspended with 1 $\times$  Tris-EDTA buffer. DNA  
194 quantification and quality assessment were performed using the NanoDrop 2000  
195 spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA). The A260/A280 ratio  
196 was used to evaluate the quality of the sample, which values between 1.7 and 2.1 were  
197 considered acceptable. The genotyping process was performed using a pre-designed specific  
198 TaqMan<sup>®</sup> SNP Genotyping Assay (Applied Biosystems, Foster city, CA, USA), according to  
199 the manufacturer's instructions and using the Rotor-Gene Q PCR cycler (Qiagen, Hilden,  
200 Germany). A scatter plot showing the endpoint fluorescence signals (i.e., an increase in VIC  
201 or FAM fluorescent signal) was used to discriminate the genotypes. The transcript alleles  
202 were used similarly to that previously used.

203

### 204 **Association with sprint performance**

205 To further investigate the influence of sprint-related polymorphisms on sprint performance, a  
206 sample of 37 top-level 100-m runners (28 Brazilians and 9 Russians) had their personal best  
207 sprint running time in the 100-m dash at official events compared between the genotypes of  
208 the selected polymorphism. Athlete's personal records were acquired using the International  
209 Association of Athletics Federations (IAAF) database, available online at  
210 <https://www.worldathletics.org/athletes>. Only athletes with performance data available on the  
211 IAAF database were included in the study.

212

## 213 **Statistical analysis**

214 First of all, the Chi-square test ( $\chi^2$ ) was used to test for the presence of the Hardy-Weinberg  
 215 equilibrium (HWE) in each control group. A departure from HWE was observed when  $\chi^2 >$   
 216 3.84 (i.e.,  $P > 0.05$ ). Thereafter, the frequencies of genotypes or alleles were compared  
 217 between sprinters and ethnically-matched controls or ethnically-matched endurance athletes  
 218 using the  $\chi^2$  test or Fisher's exact test when appropriate. Differences in the proportion of  
 219 muscle fiber types between groups with different genotypes were analyzed using unpaired  $t$ -  
 220 test and one-way ANOVA. The unpaired  $t$ -test was also used to evaluate the influence of the  
 221 selected polymorphism on 100-m sprint performance. The significance level was established  
 222 at  $P < 0.05$ .

223 For the pooled analysis of the Japanese, Russian and Brazilian cohorts, meta-analysis  
 224 was conducted using the Review Manager (RevMan) computer program version 5.3  
 225 (*Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2014*). Data from  
 226 a previously published Russian case-control study<sup>10</sup> were also used in the meta-analysis, that  
 227 is, data from 99 Russian highly elite athletes (i.e., Winners of World Championships, World  
 228 Cups or Olympic Games; 43 sprinters and 56 endurance athletes) and 173 controls were also  
 229 included in the meta-analysis. The DerSimonian and Laird random-effects model was used to  
 230 calculate weighted odds ratio (OR) and its 95% confidence interval (95% CI). The test of  
 231 overall effect was assessed using the Z score with the significance level established at  $P <$   
 232 0.05. Heterogeneity between studies was assessed using the standard  $\chi^2$  test (Cochran Q test)  
 233 and the  $I^2$  statistic.

234

## 235 **Results**

### 236 **Case-control association study**

237 Of the three polymorphisms evaluated in the Japanese case-control study, only the rs3213537  
 238 of the copine 5 (*CPNE5*) gene was found to be significant. The G-allele of the rs3213537  
 239 polymorphism was overrepresented in Japanese sprinters compared to endurance athletes  
 240 (80.6 vs. 66.7%;  $P = 0.024$ ) and associated with increased proportion of glycolytic fast-twitch  
 241 (IIx) muscle fibers in Japanese male controls (G/G (n = 69)  $24.7 \pm 9.4\%$ , G/A (n = 22)  $22.1 \pm$   
 242  $6.9\%$ , A/A (n = 7)  $16.7 \pm 7.0\%$ ;  $P = 0.02$ ). A strong trend towards an increase in the  
 243 proportion of type IIx fibers was observed in Japanese males even when the analysis was  
 244 adjusted for age ( $P = 0.061$ ). In addition, the G-allele of the rs3213537 polymorphism was  
 245 also associated with increased proportion of fast-twitch muscle fibers in 287 Finnish  
 246 individuals adjusted for sex and age (G/G (n = 189)  $55.8 \pm 14.9\%$ , G/A (n = 91)  $54.1 \pm$   
 247  $14.2\%$ , A/A (n = 7)  $43.1 \pm 16.6\%$ ;  $P = 0.041$ ). Based on these associations, the rs3213537  
 248 polymorphism was selected for replication in the Russian and Brazilian cohorts. Given the  
 249 low frequency of the homozygous genotype for the minor allele (i.e., the A/A genotype), the  
 250 rs3213537 polymorphism was analyzed only under the dominant model (G/G vs. G/A+A/A).  
 251 Of note, the transcript alleles instead of the genomic alleles were used to facilitate the link  
 252 between this study and previous data (discovery stage)<sup>10</sup>, that is, the G/A alleles represent the  
 253 genomic C/T alleles.

254 Table 2 shows the genotype distribution and allele frequency of the rs3213537  
 255 polymorphism in the three cohorts evaluated. Similar to that observed in the Japanese cohort,  
 256 the G-allele was overrepresented in Russian sprinters compared to endurance athletes (86.4  
 257 vs. 76.7%;  $P = 0.027$ ) or controls (86.4 vs. 77.8%;  $P = 0.042$ ). Indeed, the G/G genotype was  
 258 overrepresented in Russian sprinters compared to endurance athletes (74.3 vs. 56.3%;  $P =$   
 259 0.017). In the Brazilian cohort, there was a difference of  $\approx 5\%$  of G/G genotype carriers  
 260 between sprinters and the two other groups (controls and endurance athletes), but this



261 difference was not statistically significant. However, the direction of effect observed in the  
 262 Brazilian cohort was the same as in the Japanese and Russian cohorts (OR > 1.2).

263

### 264 **Meta-analysis**

265 Meta-analysis showed that, in the pooled data of the Japanese, two Russian (including data  
 266 from the previous study <sup>10</sup>) and Brazilian cohorts, the G-allele frequency was significantly  
 267 higher in sprinters compared with controls ( $P = 0.004$ ), endurance athletes ( $P = 0.002$ ) or  
 268 controls + endurance athletes ( $P = 0.002$ ), as shown in Table 3. Indeed, presence of the G/G  
 269 genotype rather than G/A+A/A genotypes increased the chance of being a top-level sprinter  
 270 compared to controls (OR: 1.49, 95% CI: 1.10–2.01;  $P = 0.005$ ), endurance athletes (OR:  
 271 1.79, 95% CI: 1.26–2.55;  $P = 0.001$ ) or controls + endurance athletes (OR: 1.58, 95% CI:  
 272 1.19–2.10;  $P = 0.002$ ). There was no evidence of heterogeneity between studies.

273

### 274 **Sprint performance**

275 Figure 1 shows the comparison of the personal best times in 100-m performance between  
 276 male sprinters with the G/G genotype ( $n = 26$ ) and male sprinters with the G/A+A/A  
 277 genotypes ( $n = 11$ ). Male sprinters with the G/G genotype have been found to have  
 278 significantly faster personal times ( $10.50 \pm 0.26$  s vs.  $10.76 \pm 0.31$  s,  $P = 0.014$ ).

279

### 280 **Discussion**

281 This study aimed to replicate potential sprint-related polymorphisms recently identified by a  
 282 GWAS in three independent cohorts of top-level sprinters, as well as to evaluate their  
 283 relationship with the proportion of fast-twitch muscle fibers. The main finding of this  
 284 investigation involving 1,875 subjects was that the G-allele of the rs3213537 polymorphism  
 285 was more frequent in sprinters and associated with increased proportion of fast-twitch muscle  
 286 fibers in Japanese and Finnish individuals and the 100-m sprint performance in Brazilian and  
 287 Russian sprinters, particularly the homozygotes (i.e., carriers of the G/G genotype). Meta-  
 288 analysis of 310 sprinters compared with 694 non-athletes and 381 endurance athletes showed  
 289 that carriers of the G/G genotype were  $\approx 1.6$  times more likely to be a sprinter.

290 The rs3213537 is an intronic polymorphism found in the *CPNE5* gene located at the  
 291 6p21.2 region of the chromosome 6. Copines are a family of calcium-dependent, membrane-  
 292 binding proteins that are evolutionary conserved from protozoans to humans <sup>15</sup>. Present in all  
 293 major mammalian organs, copines may play fundamental roles in eukaryotic cell processes <sup>16</sup>.  
 294 Copine proteins contain two N-terminal C2 domains that involve residues important for  
 295 calcium and phospholipid binding and a C-terminal A domain that may be involved in  
 296 protein–protein interactions <sup>15</sup>. This well characterized structure, especially the C2 domains,  
 297 suggests their involvement in processes of signal transduction or membrane trafficking, which  
 298 occurs in a calcium-dependent manner <sup>16</sup>. However, their biological roles have not yet been  
 299 fully defined.

300 There are at least eight different human copine proteins, which were referred to using  
 301 roman numerals. Some of them (copine-I, -II and -III) are ubiquitously expressed, while the  
 302 others have a more restricted expression profile <sup>17</sup>. As an example, copine-VI is a cytosolic  
 303 protein strongly expressed in hippocampal excitatory neurons that has been shown to affect  
 304 the structural plasticity of the dendritic spine in response to presynaptic activity <sup>18</sup>. Synaptic  
 305 calcium signals lead to copine-VI translocation from the cytosol to the postsynaptic spine  
 306 membranes, where they can serve as a calcium sensor that links neuronal activity to the  
 307 subsequent long-term changes in synaptic structure by altering actin cytoskeleton morphology  
 308 <sup>19</sup>. It was shown that copine-VI is responsible for the recruitment and local activation of the  
 309 Rac family small GTPase 1 (Rac1) protein, which, in turn, activates the Rac1-PAK-LIMK1-  
 310 Cofilin pathway and cause actin re-arrangement in favor of the long-lasting, stable

311 strengthening of excitatory synapses<sup>19</sup>. The molecular events underlying copine-V (encoded  
312 by *CPNE5* gene) are less understood, however, there may be some resemblance to other  
313 copine proteins, such as copine-VI, as they are structurally highly similar. Nonetheless, they  
314 can be expressed in different brain regions or tissues and interact with different proteins.

315 Based on animal research, copine-V has been shown to play a key role in the  
316 development of the central nervous system as it is highly expressed during the embryonic  
317 brain development<sup>20</sup>. Its expression decreases dramatically in the adult brain, remaining  
318 expressed in some non-neural tissues such as the heart, lung and muscles<sup>21</sup>. Nevertheless,  
319 although its expression may be low in the cortex and almost undetectable in the cerebellum of  
320 the adult brain, *CPNE5* is moderately expressed in the striatum of adult mice that have  
321 learned a complex motor task<sup>22</sup>. Alterations in neuronal ensemble activity and synaptic  
322 plasticity of the striatum are highly relevant for efficient human motor actions because it is  
323 the foundation for long-term motor learning or motor memory<sup>22, 23</sup>.

324 There is evidence supporting that a lack of motor memory may be detrimental to  
325 power and sprint performance<sup>24</sup>. Individuals with superior working memory are able to  
326 perform faster and more accurate in motor tasks due to a better neural efficiency<sup>25</sup>. Although  
327 with training, both neural activity and performance can be improved. Repetitive activation of  
328 the same neuronal circuit induces the clustering of new spines in postsynaptic membranes,  
329 favoring motor performance as it strengthens the dynamics of synaptic transmission<sup>26</sup>. Thus,  
330 the most effective neural communication favors sprint performance. There are synaptic inputs  
331 at the central and peripheral levels, directly influencing the rapid activation of muscles<sup>27</sup>. The  
332 ability of the neuromuscular system to increase contractile activity when muscle activation is  
333 intended to be performed as quickly as possible, referred to as Rate of Force Development  
334 (RFD), is considered vital for athletes requiring high-speed motor actions such as sprinters.  
335 Cross-sectional studies have shown that top-level sprint/power athletes are characterized by a  
336 markedly greater RFD<sup>28</sup>. Moreover, athletes with a higher RFD demonstrated faster sprint  
337 times<sup>29</sup>. Additional contributions may also occur due to differences in muscle fiber type  
338 composition—the RFD is faster in type II fibers<sup>27</sup>.

339 Whether *CPNE5* rs3213537 mutant carriers have impaired motor memory or muscle  
340 recruitment ability remains to be established, but the homozygous genotype for the major  
341 allele (i.e., the G/G genotype) was associated with fast-twitch muscle fibers and faster times  
342 in the 100-m event, which is considered the standard measure of the sprint ability of human  
343 bipedal locomotion<sup>2</sup>. Based on its role in the central nervous system, *CPNE5* polymorphisms  
344 were previously associated with alcohol dependence and obesity<sup>20</sup>. In particular, the mutant  
345 allele of the *CPNE5* rs3213537 polymorphism was strongly associated with alcohol abuse<sup>20</sup>,  
346 which adversely impacts athletic performance in a number of different ways, including mood  
347 instability and sensory-motor system dysfunction<sup>30</sup>.

348 As mentioned earlier, the *CPNE5* rs3213537 is a gene variant occurring within an  
349 intron (genomic position and change: g.36748144C>T based on the Genome Reference  
350 Consortium Human Build 38). Introns harbour polymorphisms that can influence the  
351 expression of the genes that host them and modulate the genotype–phenotype relationship.  
352 Thus, this polymorphism may modulate *CPNE5* expression and its calcium-modulated signal  
353 transduction. Interestingly, the interaction between copines and membranes occurs at  
354 concentrations of calcium that are likely to occur in the cytosol of stimulated cells but not in  
355 resting cells<sup>16</sup>, and therefore, calcium-regulated phenotypes may be affected by mutations in  
356 the *CPNE5* gene. Of particular interest, during neuromuscular junction formation, muscle  
357 fibers are intrinsically pre-specialized by clustering postsynaptic proteins, whereas the proper  
358 patterning of postsynaptic protein clusters in the center of developing muscle fibers and the  
359 subsequent innervation by the motor nerve critically depend on calcium signals<sup>31</sup>.

360 Collectively, we speculate that the G/G genotype may be involved in synaptic plasticity and  
361 muscle fiber specificity in a way that favors sprint performance.

362 The present study has some limitations. Our muscle fiber composition study included  
363 only non-athlete individuals of a wide age range. However, if the polymorphism is associated  
364 with increased proportion of fast-twitch muscle fibers in untrained individuals, these  
365 individuals (carriers of the associated variant) are expected to respond better to sprint training.  
366 Power training, like that used by sprinters, seems to conserve the pre-training number of fast-  
367 twitch fibers while increasing their fiber cross-sectional area, particularly type IIx fibers <sup>32</sup>,  
368 favouring a higher RFD <sup>33</sup>. Type IIx fibers have the highest muscle fiber conduction velocity  
369 <sup>34</sup> and are considered key determinants of the RFD, especially in power-trained individuals <sup>33</sup>.  
370 Power output in type IIx fibers was 2-fold higher than type IIa fibers and 14-fold greater than  
371 type I fibers <sup>35</sup>. In line with this, the G-allele of the rs3213537 polymorphism was previously  
372 associated with the 10-m performance in a cohort of untrained Polish women <sup>10</sup>, as well as  
373 associated with the 100-m performance in elite athletes. Although our case-control study  
374 included metabolically similar athletes, the performance association study evaluated only  
375 runners. Additional studies evaluating other sprint-oriented disciplines will be interesting,  
376 given that there may be differences between sports disciplines.

### 377 **Practical Applications**

378 The GWAS represents a promising and productive way to study sports-related phenotypes by  
379 providing a number of new candidate polymorphisms—that need to be evaluated in  
380 independent cohorts of different ethnicities and using different methodological approaches to  
381 better assess the relationship between the polymorphisms and traits of interest. In this regard,  
382 collaborative efforts involving well characterized athlete cohorts of different ethnic  
383 backgrounds will be of critical importance for further progress. In the present study, based on  
384 data from different cohorts, it is plausible to assume that the rs3213537 polymorphism (G/G  
385 genotype) may be part of a favorable genetic profile for sprinters. Notwithstanding, it is  
386 important to emphasize that sports phenotypes are complex and polygenic phenomena and  
387 should therefore be interpreted with caution.

### 388 **Conclusion**

389 The G/G genotype of the *CPNE5* gene rs3213537 polymorphism was associated with sprint  
390 athletic status and performance. While the G-allele was associated with the proportion of fast-  
391 twitch muscle fibers in Japanese and Finnish individuals, the G/G genotype was associated  
392 with faster personal times in the 100-m sprint performance among elite athletes from Brazil  
393 and Russia. It is worth mentioning that a complex network of genes contributes to sports  
394 performance, and the *CPNE5* rs3213537 is just one of several variants that can make-up the  
395 genetic profile of the elite athlete.

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509 **Figure caption**

510

511

512 **Figure 1** Association between the rs3213537 polymorphism and the best 100-m personal time  
513 in Brazilian and Russian male sprinters. The dashed line represents the qualifying time for the  
514 Tokyo 2020 Olympic Games (10.05 s).

515

516

517 **Table 1** Description of the polymorphisms evaluated in the present study

Polymorphism	Location (position <sup>†</sup> )	Consequence	REF/ALT	Sprint Allele	Previous association
rs3213537	Chromosome 6 (36748144)	Intron variant	G/A	G (major allele)	SFP, SPW, AS
rs1929877	Chromosome 9 (78799771)	Intergenic variant	A/G	G (minor allele)	SFP, AS, MF
rs17347590	Chromosome 20 (48525214)	Intergenic variant	C/A	C (major allele)	SFP, SPW, MF

518 Legend: REF, Reference allele; ALT, Alternate allele; SFP, Associated with sprint performance in young  
519 British football players; SPW, Associated with sprint performance in healthy young Polish women; AS,  
520 Associated with sprint/power athlete status; MF, Associated with proportion of fast-twitch muscle fibers in  
521 Russian physically active subjects. <sup>†</sup>Genomic position based on GRCh38 (Genome Reference Consortium  
522 Human Build 38).  
523

524 **Table 2** Genotype distribution and allele frequency of the rs3213537 polymorphism in the  
 525 Japanese, Russian and Brazilian cohorts

Group	<i>n</i>	Genotypes (%)			G allele	Comparisons: <i>P</i> -value (Effect Direction)	
		G/G	G/A	A/A		G/G vs. G/A+A/A	Alleles (G vs. A)
Japanese sprint/power athletes	54	63.0	35.2	1.9	80.6	1.000	1.000
Japanese endurance athletes	60	45.0	43.3	11.7	66.7	<u>0.062 (OR: 2.08)</u>	<u>0.024 (OR: 2.07)</u>
Japanese controls	116	57.8	35.3	6.9	75.4	0.615 (OR: 1.24)	0.334 (OR: 1.35)
Russian sprinters	70	74.3	24.3	1.4	86.4	1.000	1.000
Russian endurance athletes	103	56.3	40.8	2.9	76.7	<u>0.017 (OR: 2.24)</u>	<u>0.027 (OR: 1.93)</u>
Russian controls	117	62.4	30.8	6.8	77.8	0.110 (OR: 1.74)	<u>0.042 (OR: 1.82)</u>
Brazilian sprinters	143	73.4	23.8	2.8	85.3	1.000	1.000
Brazilian endurance athletes	162	68.5	28.4	3.1	82.7	0.378 (OR: 1.27)	0.439 (OR: 1.21)
Brazilian controls	288	67.7	29.5	2.8	82.5	0.266 (OR: 1.32)	0.331 (OR: 1.23)

526 Underlined values indicate an association trend ( $0.05 < P < 0.07$ ), and double underlined values

527 indicate nominal associations ( $P < 0.05$ ). Legend: OR, Odds Ratio.

528



529 **Table 3** Meta-analysis of the association between the rs3213537 polymorphism and sprinter athlete status

Comparison	Model	OR (95% CI)	Heterogeneity	Test for overall effect
Sprint/Power athletes vs. Controls	G/G vs. G/A+A/A	1.49 (1.10–2.01)	$\chi^2 = 1.94$ ( $P = 0.58$ ); $I^2 = 0\%$	$Z = 2.57$ ( $P = 0.01$ )
	Alleles (G vs. A)	1.47 (1.13–1.92)	$\chi^2 = 2.91$ ( $P = 0.41$ ); $I^2 = 0\%$	$Z = 2.85$ ( $P = 0.004$ )
Sprint/Power athletes vs. Endurance athletes	G/G vs. G/A+A/A	1.79 (1.26–2.55)	$\chi^2 = 3.30$ ( $P = 0.35$ ); $I^2 = 9\%$	$Z = 3.25$ ( $P = 0.001$ )
	Alleles (G vs. A)	1.70 (1.22–2.35)	$\chi^2 = 3.70$ ( $P = 0.30$ ); $I^2 = 19\%$	$Z = 3.16$ ( $P = 0.002$ )
Sprint/Power athletes vs. Controls + Endurance athletes	G/G vs. G/A+A/A	1.58 (1.19–2.10)	$\chi^2 = 2.48$ ( $P = 0.48$ ); $I^2 = 0\%$	$Z = 3.13$ ( $P = 0.002$ )
	Alleles (G vs. A)	1.55 (1.18–2.03)	$\chi^2 = 3.35$ ( $P = 0.34$ ); $I^2 = 11\%$	$Z = 3.16$ ( $P = 0.002$ )

530 Comparisons are expressed as Odds Ratio (OR) and 95% Confidence Interval (95% CI). Heterogeneity between studies was assessed using the Cochran  
531 Q test ( $\chi^2$ ) and the  $I^2$  statistic.

532

533