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1 **TOTAL HEMOGLOBIN MASS, AEROBIC CAPACITY AND THE *HBB* GENE IN**

2 **POLISH ROAD CYCLISTS**

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26 **ABSTRACT**

27 The relationships between genes, amount of hemoglobin and physical performance are still
28 not clearly defined. The aim of this study was to examine the association between -551C/T
29 and intron 2, +16 C/G polymorphisms in the *HBB* gene and total hemoglobin mass (tHb_{mass})
30 and aerobic capacity in endurance athletes. tHb_{mass} and aerobic capacity indices, i.e. maximal
31 oxygen uptake (VO₂max), oxygen uptake at anaerobic threshold (VO₂AT), maximal power
32 output (Pmax), and power at anaerobic threshold (PAT), were determined in 89 young road
33 cyclists, female (n=39) and male (n=50), who were genotyped for 2 polymorphisms in the
34 *HBB* gene. The relative values of aerobic capacity indices differed significantly among intron
35 2, +16 C/G polymorphisms of the *HBB* gene only in female cyclists; athletes with GG
36 genotype had significantly higher values of VO₂max (P=0.003), VO₂AT (P=0.007), PAT
37 (P=0.015) and Pmax (P=0.004) than did C carriers. No relationships were found between the
38 C-carrier model (CC+CG vs GG in the case of intron 2, +16 C/G and CC+CT vs TT for -551
39 C/T polymorphisms of the *HBB* gene) and relative values of tHb_{mass}. Our results demonstrated
40 that the *HBB* gene could be related to aerobic capacity, but it seems that it does not result
41 from an increase in the amount of hemoglobin in the blood.

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43 **KEY WORDS: genetic polymorphism, *HBB* gene, hemoglobin, aerobic capacity,**44 **athletes**

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49 INTRODUCTION

50 In endurance athletes the most important factor of success is aerobic capacity, which is mainly
51 expressed by maximal oxygen uptake (VO_2max) (17,30). The important factors affecting
52 VO_2max are cardiac output, O_2 carrying capacity and oxygen utilization by muscle tissue
53 (30). However, among endurance athletes the main limiting factor of VO_2max is oxygen
54 supply (5). Hemoglobin is a protein responsible for efficient transport of oxygen to the
55 tissues; therefore it is an important factor contributing to aerobic capacity (25,30). Although it
56 has been estimated that an increase in total hemoglobin mass (tHb_{mass}) by 1 g causes a rise in
57 VO_2max of approximately 4 ml/min (25), it is worth emphasizing that the total hemoglobin
58 mass, rather than its concentration, shows a strong correlation with maximal oxygen uptake
59 (24,25). Variation in the amount of hemoglobin and indices of aerobic capacity is dependent
60 on many factors, e.g. iron status, illness, period of inactivity, altitude exposure as well as
61 length, duration, type, intensity and age of initiation of the training stimulus
62 (10,15,17,20,25,29), but may also be influenced by genetic parameters (8,15,17,26). Schmidt
63 and Prommer (24,25) reported that tHb_{mass} may be relatively stable in healthy adults (mostly
64 competitive athletes) over a very long period, despite changes in training and lifestyle.
65 To date, researchers have described over 300 genes that could be related to predisposition to
66 physical fitness and sports results (2, 7). One of them could be the beta hemoglobin (*HBB*)
67 gene (18). The impact of the *HBB* gene on physical performance is not well documented,
68 because so far this gene has been studied mainly in the context of genetic diseases (11). An
69 association between the *HBB* gene polymorphisms and running economy in the untrained
70 state and in response to aerobic training was described only in one study with recruits from
71 the Chinese military police (13).

72 Many authors have emphasized that genetic predisposition seems to be a prerequisite for high
73 tHb_{mass} and high endurance performance, but despite the many excellent scientific papers

74 about tHb_{mass} and performance parameters (15,24,25) the tHb_{mass}–performance–gene
75 relationship has not been clearly defined. Although Ahmetov et al. (3) showed recently that
76 the rs157231 CC genotype of the *NFIA-AS2* gene (involved in the regulation of expression of
77 the erythropoiesis inducing nuclear factor I A) was associated with high VO₂max and high
78 hemoglobin concentration, as well as a high number of reticulocytes and erythrocytes in
79 endurance athletes, they did not assess the total amount of hemoglobin. Therefore the aim of
80 our study was to examine the association between 2 polymorphisms of the *HBB* gene and
81 tHb_{mass} and indices of aerobic capacity in endurance athletes.

82

83 **METHODS**

84 **Experimental approach to the problem**

85 To the best of our knowledge, there is still no study concerning the association between the
86 *HBB* gene, amount of hemoglobin and aerobic capacity. To elucidate whether having specific
87 polymorphisms of the *HBB* gene could exert a positive effect on the amount of hemoglobin in
88 the blood and aerobic capacity, we analyzed relationships between *HBB* gene intron 2,
89 +16C/G and -551C/T polymorphisms and tHb_{mass} as well as maximal oxygen uptake
90 (VO₂max), oxygen uptake at anaerobic threshold (VO₂AT), maximal power output (Pmax),
91 and power at anaerobic threshold (PAT) in endurance athletes.

92

93 **Subjects**

94 Ninety-two road cyclists (male and female), aged 16-28 years, participated in the study. Most
95 of the study participants were members of national junior or senior teams. In order to exclude
96 individuals with symptoms of infectious or cardiovascular diseases, latent iron deficiency
97 (n=3) or iron deficiency anemia, the subjects were given a medical and biochemical
98 examination. Finally, the results obtained from 89 athletes (39 females and 50 males) were

99 analyzed. The physical characteristics of subjects, separated by gender, as well as basic data
100 concerning sports experience and training load, are shown in Table 1. (table 1 about here)
101 The results concerning *HBB* genotyping obtained in athletes were compared with those
102 observed in 119 Polish untrained persons (59 females and 60 males) aged 20-25 years (control
103 group). All athletes and untrained persons were Caucasians. The study was approved by the
104 Institute of Sport Committee of Ethics, and written informed consent was obtained from all
105 individual participants of the study.

106 **Design**

107 The study consisted of three steps performed on two days in the following order: first day – 1)
108 venous blood sampling and anthropometric measurements, 2) evaluation of aerobic capacity,
109 3) measurements of tHb_{mass}; second day – 1) measurement of body mass and venous blood
110 sampling, 2) measurements of tHb_{mass}.

111 **Procedures**

112 **Blood collection and analysis**

113 The blood samples were withdrawn from the cephalic vein in the morning in a preprandial
114 state after remaining for at least 15 min in a sitting position.

115 *Indices of iron status*

116 Hemoglobin concentration (Hb), hematocrit (Hct), and erythrocyte count (RBC) were
117 assessed using an ADVIA 120 hematological analyzer (Siemens, Germany). In serum the
118 following indices were measured: soluble transferrin receptor (sTfR) concentration by using
119 immunoenzymatic commercial kits (Ramco, USA); ferritin concentration by using the
120 immunoturbidimetric method (Pentra, USA), total iron binding capacity (TIBC) by using the
121 colorimetric method (BioMaxima, Poland), and C-reactive protein (CRP) by using the
122 immunoturbidimetric method (Pentra, USA).

123 **DNA isolation and *HBB* polymorphism typing**

124 Genomic DNA from athletes and untrained person was extracted from whole blood using the
125 GeneMATRIX Quick Blood DNA Purification Kit (Eurx, Germany). *HBB* gene intron 2,
126 +16C/G and -551C/T polymorphisms were analyzed as described previously (19) using pairs
127 of primers specific to DNA fragments containing the polymorphic site (13). Genotyping of
128 two SNPs was performed using the RFLP technique with 2 U of *AvaII* and 2 U of *RsaI*
129 restriction enzyme (Fermentas, USA) for intron 2, +16C/G and -551C/T typing, respectively.
130 All restriction cutting was performed for 2.5 h at 37°C and digested products were
131 electrophoresed on 3% agarose gel.

132 **Determination of tHb_{mass}**

133 tHb_{mass} was measured using a modified version of the CO rebreathing procedure, according to
134 Schmidt and Prommer (23). Briefly, the subjects inhaled a bolus of 99.9% chemically pure
135 CO (Linde Gas) in a dose of 1.0 ml/kg body mass for males and 0.8 ml/kg body mass for
136 females and rebreathed in a closed system (spirometer, SpiCo, Bayreuth, Germany) for 2 min.
137 The samples of the arterialized capillary blood were taken from the earlobe three times:
138 directly before the test and in the 6th and 8th minute after the respiration through the
139 spirometer was started. Analysis of the percentage value of carboxyhemoglobin (HbCO%)
140 (ABL 80 Flex, Radiometer, Denmark) was performed in triplicate samples before and in the
141 8th minute and in duplicate samples in the 6th minute of the study. A detailed description of
142 this method has been provided in publications by its authors (22,23). Based on the results of
143 tHb_{mass}, Hb and Hct, the blood (BV) and plasma volumes (PV) were also computed. In all
144 participants measurements of tHb_{mass} were made in duplicate. The typical error (TE) in our
145 laboratory with duplicated measures (24-48 h time lag between tests) in the cyclist group was
146 1.85%.

147 **Aerobic capacity**

148 A graded exercise test to exhaustion was performed on a cycle ergometer (Cyclus2, Leipzig,
149 Germany) to determine maximal aerobic capacity ($\text{VO}_{2\text{max}}$), maximal power output (P_{max}),
150 as well as anaerobic threshold (AT). The tests were performed using the participant's personal
151 bike. The test started at workload 1.50 W/kg of body mass and was increased every 3 minutes
152 by 0.75 W/kg for males and 0.70 W/kg for females. The test was terminated when the subject
153 could no longer complete the desired workload despite verbal encouragement. Additional
154 maximal exercise performance criteria were: a heart rate close to age predicted maximum,
155 respiratory exchange ratio (RER) value of >1.1 , blood lactate concentration >10 mmol/L. The
156 test was preceded by a 10-minute warm-up at workload of 1 W/kg and thereafter a 5-minute
157 rest.

158 During the exercise test expiratory air was analyzed using a portable measuring system
159 (MetaMax, Cortex, Germany). Prior to each test this system was calibrated with a known
160 volume syringe and gas concentration (O_2 , CO_2). Heart rate was monitored using the Polar
161 Sports Tester device.

162 At the end of each workload capillary blood samples were taken from the fingertip in order to
163 determine changes in lactate concentration (Super GL2 analyzer, Dr. Muller, Germany).

164 The anaerobic threshold was assumed as power output (PAT) and corresponding oxygen
165 uptake ($\text{VO}_{2\text{AT}}$) at threshold (4 mmol/L) blood lactate concentration (14) and was estimated
166 by the method of interpolation.

167 **Anthropometric measurements**

168 Anthropometric measurements comprising assessment of body height, body mass and
169 skinfold thickness were performed. The percentage of body fat was calculated using the
170 equation of Durnin and Womersley (9).

171 **Statistical analysis**

172 All the data are presented as means and standard deviations, and were analyzed using the
173 Statistica 10 software package (StatSoft Inc. Tulsa, USA).
174 Owing to the low number of CC homozygotes for intron 2, +16 C/G and of -551 C/T
175 polymorphisms, they were combined with heterozygotes (C-carrier model) and compared to
176 GG and TT homozygotes, respectively. Differences between mean values of tHb_{mass} in groups
177 of athletes (males and females separately) possessing different genotypes of the *HBB* gene
178 were tested by the Kruskal-Wallis test, whereas the Mann-Whitney U test was used for
179 comparison of mean values of tHb_{mass}, oxygen consumption, and power output in groups
180 distinguished according to genotype variants. The significance of differences in genotype and
181 allele frequencies as well as conformity with the Hardy–Weinberg principle was estimated
182 using the χ^2 test. A Pearson correlation test was used to analyze the relationship between two
183 quantitative variables. The statistical significance was set at $P < 0.05$.

184

185 **RESULTS**

186 Both polymorphisms were in Hardy-Weinberg equilibrium in male and female athletes and
187 controls. No differences were found in the *HBB* genotype and allele frequencies between male
188 and female athletes, as well as between athletes and controls (Table 2). (table 2 about here)

189 The *HBB* genotypes had no significant effect on relative values of tHb_{mass} both for female and
190 male athletes (Table 3).

191 Moreover, there were no associations between PV, BV and Hb concentrations and genotype
192 variants of the *HBB* gene (data not shown). Also no relationships were found between
193 genotype models, i.e. CC+CG vs GG in the case of intron 2, +16 C/G polymorphism and
194 CT+CC vs TT for -551 C/T polymorphism of the *HBB* gene and relative values of tHb_{mass}
195 (Table 3). (table 3 about here)

196 The relative values of aerobic capacity indices differed according to intron 2, +16 C/G
197 polymorphism of the *HBB* gene in female cyclists; athletes with GG genotype had
198 significantly higher values of $VO_2\text{max}$ ($P=0.003$), $VO_2\text{AT}$ ($P=0.007$), P_{max} ($P=0.004$) and
199 PAT ($P=0.015$) than did C carriers (CC + CG genotypes) (Table 4).

200 Among the male athletes these indices did not differ significantly between the C-carrier model
201 and GG genotypes in intron 2, +16 C/G polymorphism of the *HBB* gene (Table 5).

202 The -551 C/T polymorphism of the *HBB* gene had no significant effect on relative values of
203 $VO_2\text{max}$, $VO_2\text{AT}$, P_{max} and PAT in both female and male athletes (Tables 4 and 5). (tables 4
204 and 5 about here)

205 In female athletes there was an association between tHb_{mass} and $VO_2\text{max}$ ($P=0.00002$),
206 $VO_2\text{AT}$ ($P=0.00000$), P_{max} ($P=0.00001$) and PAT ($P=0.00000$) in relative values. In men a
207 relationship was observed between relative values of tHb_{mass} and $VO_2\text{max}$ ($P=0.00008$),
208 $VO_2\text{AT}$ ($P=0.0006$) and PAT ($P=0.0012$) (Figure 1). (figure 1 about here) Additionally, there
209 was a significant association between absolute values of tHb_{mass} and $VO_2\text{max}$, $VO_2\text{AT}$, P_{max} ,
210 and power output at $4 \text{ mmol}\cdot\text{l}^{-1}$ blood lactate concentration in both male and female athletes
211 (data not shown).

212

213 **DISCUSSION**

214 In athletes, hematological traits are important not only in the clinical and health aspect but
215 also with respect to their physical performance. The regulation of erythropoiesis takes place
216 on several levels and depends on many factors such as cytokines, hormones, transcription
217 factors, and miRNA, which in turn have an effect on gene expression (12), while training has
218 only small effects on the total amount of hemoglobin in the blood (24). On the other hand,
219 many studies, including the one presented here, indicate a strong relationship between tHb_{mass}
220 and maximal oxygen uptake (15,25). Moreover, very high values of tHb_{mass} were observed in

221 elite Polish endurance athletes, as well as in young athletes who had just begun professional
222 training (unpublished results). These results confirm that hemoglobin is the principal
223 transporter of oxygen, and therefore a high total amount of it could, to a large extent,
224 determine aerobic capacity (1,25). One of the genes responsible for the production of red
225 blood cells and hemoglobin is the *HBB* gene. It should be noted that hundreds of variations
226 have been identified in the *HBB* gene, and many polymorphisms may be related to
227 hematological traits (11). For example, Auer et al. (4) reported that one polymorphism of the
228 *HBB* gene (rs33971440) was associated with lower hemoglobin concentration, hematocrit
229 level and clinical anemia. It is more likely that several polymorphisms of the *HBB* gene are
230 responsible for the amount of hemoglobin and hence for aerobic capacity. In addition, it is
231 often emphasized that genetics is an important factor influencing physical performance,
232 although it is still not known which gene variants have an impact on it (2,3,16,21). So far in
233 sport genetics the *HBB* gene has been examined only for three polymorphisms (intron 2+16
234 C/G, -551 C/T and +340 A/T polymorphisms) (13). He et al. (13) observed the relationship
235 between homozygosity for the C allele of -551C/T and intron 2, +16 C/G (rs10768683)
236 polymorphisms and running economy training response, but not with VO_2max . In our study in
237 male athletes there was no relationship between the *HBB* gene polymorphisms and VO_2max ,
238 as well as other aerobic capacity indices. However, in female athletes we observed a strong
239 relationship between relative values of VO_2max , P_{max} and PAT and the *HBB* gene variants,
240 but only in the case of G homozygotes of the intron 2, +16 C/G polymorphism. One might
241 suggest that the same association was not replicated in male athletes due to relatively small
242 sample size, differences in factors affecting hemoglobin levels between genders and the fact
243 that within-person variation from day to day of hemoglobin values are higher in men than in
244 women (6). However, the results of our study show no differences in the *HBB* genotype and

245 allele frequencies between male and female athletes, which is in accordance with earlier
246 results obtained in Polish cross-country skiers and runners (19).

247 Because both tHb_{mass} and *HBB* genotypes (13) demonstrated relationships with indices of
248 aerobic capacity, it was suggested that tHb_{mass} may depend on the *HBB* gene. However, we
249 did not confirm this hypothesis, because regardless of gender none of the *HBB* variants
250 (genotypes and genotype models) showed an association with tHb_{mass} or Hb concentration.
251 Similar results were observed in Polish cross-country skiers and middle and long distance
252 runners (19). In accordance with this, the *HBB* gene effect is opposite to other genes, because
253 higher hemoglobin and hematocrit levels were observed in some polymorphisms of *EPO*
254 (erythropoietin), *TFR2* (transferrin receptor 2), *NFIA-AS2* (nuclear factor I A antisense RNA
255 2) and *HIF1A* (hypoxia-inducible factor 1 alpha) genes (3,4,27). Despite the fact that the *HBB*
256 gene is one of the primary genes in hemoglobin synthesis (28), there is still too little
257 information concerning relationships of this gene's polymorphisms with amount of
258 hemoglobin, so this issue requires further investigations.

259 Moreover, we did not find any differences in the *HBB* genotype distribution and allele
260 frequencies between athletes and control groups. However, such a phenomenon has been
261 observed for other "sport genes", and it has been suggested that genetic factors may
262 predispose to successful sport performance (7). There is no study concerning the distribution
263 of genotypes of the *HBB* gene among athletes and control groups, so we cannot compare our
264 results with others.

265 The only study on this issue was carried out on Chinese non-athletes (13). However,
266 comparing the frequencies of genotypes in Polish and Chinese populations can be difficult
267 due to the ethnic origin, because certain alleles could be overrepresented in some ethnic
268 groups (32). This is especially evident in the frequency of CC genotype for intron 2 +16 C/G
269 polymorphism, which in the Polish male population was 6.0% and 2.0% in athletes and

270 controls, respectively, in contrast to 24.5% in the Chinese male population (13). Moreover,
271 the racial differences in impact of specific polymorphisms on exercise capacity is strongly
272 suggested (31,32). As described by He et al. (13), in the Chinese cohort +16CC genotype was
273 associated with better physical performance, while in Polish athletes GG genotype benefits
274 endurance capacity. It seems that this discrepancy is due to ethnic origin rather than selection
275 for endurance disciplines, which is confirmed by similar results obtained in our earlier study
276 (19), as well as the lack of differences in distribution of both polymorphisms between athletes
277 and the control group, regardless of sex, in the present study. Therefore, we cannot clearly
278 determine whether this gene may be considered as a "sports gene" and be helpful in the
279 selection of athletes for sport.

280 To our knowledge this study is the first to determine the association between the *HBB* gene,
281 tHb_{mass} and parameters of aerobic capacity in athletes. The main finding of our study was the
282 significant correlation of aerobic capacity indices with one polymorphism of the *HBB* gene
283 intron 2, +16 C/G in the female group, so the impact of the *HBB* gene on aerobic capacity
284 may be connected with gender. We also found that neither of the studied polymorphisms of
285 the *HBB* gene was associated with total hemoglobin mass.

286

287 **PRACTICAL APPLICATIONS**

288 Our results suggest that the *HBB* gene intron 2, +16 C/G polymorphism may be related to
289 aerobic performance, but it seems that it is not due to an increase in the amount of
290 hemoglobin in the blood. Therefore, the *HBB* GG genotype can be considered as one of the
291 genetic markers associated with predisposition to endurance performance in females.
292 However, further research including tHb_{mass}, genes and aerobic performance indices on a
293 larger population of athletes and using different ethnic cohorts is necessary to better

294 understand the relationship between hemoglobin amount, genetic predisposition and physical
295 performance.

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400 **Figure legend**

401 Figure 1

402 Relationships between relative values of total hemoglobin mass (tHb_{mass}) and (A) relative
403 values of maximal oxygen uptake (VO₂max), (B) oxygen uptake at anaerobic threshold
404 (VO₂AT), (C) maximal power output (Pmax), and (D) power output at anaerobic threshold
405 (PAT) in female and male cyclists; circles – females, triangles – males.

406

407 **Titles of tables:**

408 Table 1. Characteristics of study participants (mean ± SD)

409 Table 2. Genotype and allele frequencies of intron 2,+16 G/C and -551C/T polymorphisms of
410 *HBB* gene in male and female athletes

411 Table 3. Relative values of total hemoglobin mass according to *HBB* genotypes in male and
412 female athletes (mean ± SD)

413 Table 4. Aerobic capacity indices according to *HBB* genotypes in female athletes (mean ± SD)

414 Table 5. Aerobic capacity indices according to *HBB* genotypes in male athletes (mean ± SD)