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# 1 Vitamin D metabolites are associated with physical performance in young healthy

- 2 adults
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#### 9 ABSTRACT

Purpose. To determine vitamin D metabolites and vitamin D receptor (VDR) single-10 nucleotide polymorphisms (SNPs) relationships with physical performance. 11 **Methods.** In 1205 men and 322 women (94.8% white Caucasian,  $22.0 \pm 2.8$  years) 12 commencing military training, we measured: serum vitamin D metabolites (25-13 hydroxyvitamin D (25(OH)D) and 24,25-dihydroxyvitamin D (24,25(OH)<sub>2</sub>D) by high-14 15 performance liquid chromatography tandem mass spectrophotometry, and 1,25dihydroxyvitamin D (1,25(OH)<sub>2</sub>D) by immunoassay); VDR SNPs (rs2228570, rs4516035, 16 17 and rs7139166 by polymerase chain reaction genotyping); and endurance performance by 2.4 km run, muscle strength by maximal dynamic lift, and muscle power by maximal vertical 18 jump. 19 **Results.** Serum 25(OH)D was negatively associated with 2.4 km run time and positively 20 associated with muscle power ( $\beta = -12.0$  and 90.1), 1,25(OH)<sub>2</sub>D was positively associated 21 with run time and negatively associated with strength and muscle power ( $\beta = 5.6, -1.06$ , and 22 -38.4), and  $24.25(OH)_2D$  was negatively associated with run time ( $\beta = -8.9$ ; P < 0.01), after 23 controlling for age, sex, smoking, alcohol, physical activity, time outdoors, season, and BMI. 24 Vitamin D metabolites (25(OH)D, 1,25(OH)2D, and 24,25(OH)2D) together explained 25 variances of 5.0% in run time, 0.7% in strength, and 0.9% in muscle power ( $\Delta F P < 0.001$ ). 26 All performance measures were superior with low  $1,25(OH)_2D:24,25(OH)_2D$  ratio (P < 0.05). 27 VDR SNPs were not associated with physical performance ( $\Delta F P \ge 0.306$ ). 28 **Conclusion.** Vitamin D metabolites accounted for a small portion of variance in physical 29 performance. Associations between vitamin D metabolites and run time were the most 30 consistent. VDR SNPs explained no variance in performance. Greater conversion of 31 25(OH)D to 24,25(OH)<sub>2</sub>D, relative to 1,25(OH)<sub>2</sub>D (*i.e.*, low 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio), 32

33 was favourable for performance, indicating  $24,25(OH)_2D$  may have a role in optimising

34 physical performance.

35 Key words: Vitamin D, exercise, endurance, muscle strength, muscle power,

- 36 polymorphisms.
- 37

### 38 INTRODUCTION

Serum 25-hydroxyvitamin D (25(OH)D) concentration is the recommended, and widely used, 39 indicator of an individual's vitamin D status  $(25(OH)D > 50 \text{ nmol} \cdot L^{-1})$  is deemed sufficiency 40 41 (1)) due to its abundance and longer half-life relative to other circulating vitamin D metabolites (1, 2). 1,25-dihydroxyvitamin D (1,25(OH)<sub>2</sub>D)—synthesized from 25(OH)D by 42 1α-hydroxylase—is the most biologically active vitamin D metabolite in humans. 43  $1,25(OH)_2D$  circulates in pmol·L<sup>-1</sup> concentrations, with its actions mediated by vitamin D 44 receptors (VDRs) (3). Despite their proximity in the metabolic pathway, there is no direct 45 correlation between serum 25(OH)D and 1,25(OH)2D due to the tight regulation of 46 hydroxylation enzymes (4). A dynamic relationship exists between 25(OH)D and 47 48  $1,25(OH)_2D$  when expressed as a relative ratio with 24,25-dihydroxyvitamin D (24,25(OH)<sub>2</sub>D) (4). 24,25(OH)<sub>2</sub>D is synthesized from 25(OH)D by 24-hydroxylase and like 49 25(OH)D, circulates in nmol·L<sup>-1</sup> concentrations. Although 24,25(OH)<sub>2</sub>D has been labeled as a 50 purely catabolic metabolite of vitamin D, potential biological roles and the possible existence 51 of a 24,25(OH)<sub>2</sub>D specific receptor have emerged (5-8). Individuals with low 25(OH)D, 52 53 normal 1,25(OH)<sub>2</sub>D, but increased 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio have higher parathyroid hormone (PTH) concentrations than those at the opposite end of the spectrum with high 54 55 25(OH)D, normal 1,25(OH)<sub>2</sub>D, and decreased 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio (4). High 24,25(OH)<sub>2</sub>D relative to 1,25(OH)<sub>2</sub>D (*i.e.*, low 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D) may reduce the 56 bioactivity of 25(OH)D and 1,25(OH)2D, downregulating PTH secretion, whilst maintaining 57

58 1,25(OH)<sub>2</sub>D within strict boundaries. Low 24,25(OH)<sub>2</sub>D relative to 1,25(OH)<sub>2</sub>D (*i.e.*, high

59  $1,25(OH)_2D:24,25(OH)_2D$ ) may upregulate PTH secretion and enhance the effects of vitamin

60 D (4). On the other hand, if  $24,25(OH)_2D$  is itself biologically active, low

61 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D may be beneficial. How this recently identified inverse

exponential relationship between serum 25(OH)D and  $1,25(OH)_2D:24,25(OH)_2D$  relates to

63 physiological outcomes, such as physical performance, remains unexplored.

64

Beyond regulating calcium and phosphate homeostasis and augmenting bone mineralisation 65 66 (1), extra-skeletal functions of vitamin D and its metabolites have emerged following the discovery of the VDR in almost all human tissues (3, 9). 1,25(OH)<sub>2</sub>D stimulates skeletal 67 muscle protein synthesis by VDR-mediated signaling (9), and may improve cardiac, skeletal 68 69 muscle, and endothelial function (10-14). Avoiding low serum 25(OH)D and achieving 70 vitamin D sufficiency (1) may, therefore, be important for muscle strength and endurance type exercise (15, 16). Cross-sectional studies investigating the influence of vitamin D on 71 72 physical performance in young healthy adults have reported both positive and no associations between circulating 25(OH)D and physical performance (17-20), when controlling for 73 variables that influence performance (e.g., sex, body composition, smoking, physical activity, 74 and season (21-23)). In contrast, improving vitamin D status by increasing serum 25(OH)D 75 76 with oral vitamin D<sub>3</sub> supplementation or increased sunlight exposure has not enhanced 77 physical performance in randomized controlled trials (24, 25). This inconsistency between observational and interventional studies may be due to a focus on serum 25(OH)D as a 78 measure of vitamin D status, and not examining the relative concentrations of vitamin D 79 80 metabolites. Rather than simply increasing serum 25(OH)D, shifting 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D from high to low may be necessary for a beneficial effect on 81

82 performance to emerge.

Several single-nucleotide polymorphisms (SNPs) within vitamin D pathway-related genes are
associated with circulating 25(OH)D and may be responsible for some inter-individual
variability in the vitamin D endocrine system (26). SNPs in the gene that encodes for the
VDR have been studied in relation to muscle strength and function in mostly elderly and
sedentary adults with equivocal results (26). The relationship between rs2228570, rs4516035,
and rs7139166 VDR polymorphisms and physical performance remains to be determined in
young, physically active adults.

90

The purpose of the study was to examine the relationship: i) between vitamin D metabolites (serum 25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D) and physical performance; and ii) between VDR polymorphisms (rs2228570, rs4516035, and rs7139166) and physical performance. We hypothesized a three-dimensional model of vitamin D metabolites incorporating serum 25(OH)D and 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D would have a dynamic relationship with physical performance; and SNPs in the VDR would be associated with variance in performance.

57

#### 98 METHODS

#### 99 **Participants**

1527 British Army recruits (1205 men and 322 women, 94.8% white Caucasian; Table 1) 100 101 voluntarily participated in the study, after providing informed written consent and passing a 102 physician-screened military medical assessment. All experimental procedures were completed during week one of initial Army training. Participants were recruited during week 103 one of initial Army training between April 2013 and May 2017 from three military training 104 populations: male infantry recruits at Infantry Training Centre, Catterick; standard entry 105 female recruits at Army Training Centre, Pirbright; and male and female officer cadets at 106 107 Royal Military Academy, Sandhurst-thereby providing a representative sample of all

individuals commencing Army training in the UK. A subset of these data have been
published (25). The present study includes unpublished vitamin D metabolite and SNP data,
and is from a larger sample, more representative of all individuals commencing Army
training. The study received ethical approval from the UK Ministry of Defence Research
Ethics Committee (protocol number 165/Gen/10) and was conducted in accordance with the
Declaration of Helsinki (2013).

114

# 115 Study design

116 A cross-sectional, observational study design was used to determine whether serum vitamin D metabolites and VDR SNPs were associated with physical performance in young healthy 117 adults. All assessments were performed during week one of initial Army training and are 118 listed here. Venous blood samples were obtained for analysis of: serum 25(OH)D, and its 119 metabolites 1,25(OH)<sub>2</sub>D and 24,25(OH)<sub>2</sub>D; and VDR SNPs in whole blood (rs2228570, 120 rs4516035, and rs7139166). Physical performance was assessed by a maximal effort 2.4 km 121 run, and tests of maximal dynamic lift strength and vertical jump peak power output. Body 122 mass and height (Seca, Hamburg, Germany) were measured in light clothing and without 123 shoes. Participants self-reported their alcohol intake; smoking habits; physical activity levels; 124 and typical time spent outdoors, using questionnaires. 125

126

#### 127 Experimental procedures

#### 128 Blood collection and handling

129 Whole blood samples were obtained by venipuncture from a prominent vein in the

130 antecubital fossa into one serum vacutainer and one EDTA vacutainer (Becton Dickinson,

131 Oxford, UK). Whole blood in the EDTA vacutainer was immediately frozen at -80°C for

132 later analysis. Whole blood in the serum vacutainer was left to clot in a vacutainer rack at

room temperature for 1 h before being centrifuged at 1500 g for 10 min at 4°C, with serum aliquots immediately frozen at  $-80^{\circ}$ C for later analysis.

135

#### 136 Biochemical analysis

Total serum 25(OH)D (25(OH)D<sub>2</sub> and 25(OH)D<sub>3</sub>) and total 24,25(OH)<sub>2</sub>D (24,25(OH)<sub>2</sub>D<sub>2</sub> and 137 24,25(OH)<sub>2</sub>D<sub>3</sub>) were measured with high-performance liquid chromatography tandem mass 138 spectrophotometry using a Micromass Quattro Ultima Pt electrospray ionisation mass 139 spectrometer, as described previously (27). Serum 1,25(OH)<sub>2</sub>D was measured by 140 141 chemiluminescent immunoassay using a DiaSorin LIAISON® XL analyser (Stillwater, Minnesota, USA). The measurement ranges of the assays were  $0-200 \text{ nmol}\cdot\text{L}^{-1}$  for 25(OH)D<sub>2</sub> 142 and 25(OH)D<sub>3</sub>, 0–25 nmol·L<sup>-1</sup> for 24,25(OH)<sub>2</sub>D<sub>2</sub> and 24,25(OH)<sub>2</sub>D<sub>3</sub>, and 12–480 pmol·L<sup>-1</sup> for 143 1,25(OH)<sub>2</sub>D. The mean coefficient of variation (CV) for intra-assay imprecision across the 144 145 measuring range of the assays was 4.9% for 25(OH)D<sub>2</sub>, 8.3% for 25(OH)D<sub>3</sub>, 7.7% for 24,25(OH)<sub>2</sub>D<sub>2</sub>, 9.0% for 24,25(OH)<sub>2</sub>D<sub>3</sub>, and 7.4% for 1,25(OH)<sub>2</sub>D. The cumulative inter-146 assay CVs were <7.4% for 25(OH)D<sub>2</sub>, <9.6% for 25(OH)D<sub>3</sub>, <10.6% for 24,25(OH)<sub>2</sub>D<sub>2</sub>, 147  $\leq$ 8.9% for 24,25(OH)<sub>2</sub>D<sub>3</sub>, and  $\leq$ 9.3% for 1,25(OH)<sub>2</sub>D. All biochemical analyses were 148 undertaken by the Good Clinical Laboratory Practice and Vitamin D External Quality 149 Assessment Scheme certified Bioanalytical Facility at the University of East Anglia. 150 151

# 152 Single-nucleotide polymorphisms

Whole blood samples in EDTA vacutainers were defrosted and resuspended for 15 min on a
rotating wheel. Genomic DNA was isolated from whole blood using the ReliaPrep<sup>TM</sup> Blood
gDNA Miniprep System (Promega, Southampton, UK) according to the manufacturer's
instructions. Using samples of DNA, Kompetitive Allele Specific PCR (KASP<sup>TM</sup>, LGC

Genomics, Teddington, Middlesex, UK) genotyping was used for SNP genotyping of
rs2228570, rs4516035, and rs7139166 in the VDR gene.

159

#### 160 *Endurance performance*

Endurance performance was assessed as the time to complete a maximal effort 2.4 km run, 161 recorded to the nearest second. After an 800 m warm up, the 2.4 km run was performed on a 162 163 standardized running course at each training site. The time to complete a 2.4 km run is indicative of maximal aerobic capacity (28) and is assessed during selection, training, and 164 165 throughout a military career. All participants were accustomed to performing this test from selection before commencing military training. Faster 2.4 km run times indicated better 166 endurance performance. Therefore, negative associations with run time indicated improved 167 endurance performance, and positive associations with run time indicated worsened 168 endurance performance. 169

170

#### 171 *Muscle strength*

Maximal dynamic lift strength was determined as the maximal weight lifted using an 172 incremental lift machine that simulates a power clean weightlifting movement, as described 173 previously (29). The device consisted of a vertically moving carriage with handgrips 174 positioned 0.30 m above the ground. Participants lifted the weight (20 kg starting mass) to a 175 176 height where the handgrips were 1.45 m from the ground, the height of a British Army four tonne truck. With each successful lift, the weight was increased by 5 kg. The test was 177 terminated when participants failed to lift the weight to 1.45 m on their second attempt. 178 Differences in body height may have affected the participants ability to lift weight to the 179 same absolute height, however, this measure of maximal dynamic lift strength was chosen 180 because it correlates with and predicts success in military and functional tasks (28). 181

182 *Muscle power* 

Vertical jump peak power output was assessed by countermovement vertical jump using a 183 jump mat (Takei Scientific Instruments, Tokyo, Japan) and validated equation (30): peak 184 power (W) = (51.9 x maximal vertical jump height (cm)) + (48.9 x body mass (kg)) - 2007, 185 as described previously (29). We analyzed this estimate of muscle power rather than jump 186 height because lower body power is important for the performance of military specific tasks 187 188 (28). A belt was fitted around the waist of each participant and secured to a rubber mat. Participants were instructed to jump as high as possible three times, with their hands placed 189 190 on their hips. A fourth jump was performed if jump height increased across the three attempts, indicative of a learning effect. Maximal vertical jump height was recorded as the 191 highest score achieved. Test-retest reliability of  $r \ge 0.90$  has been reported for these 192 performance tests (29). 193

194

### 195 Statistical analysis

Hierarchical multiple linear regression was used to examine the association between vitamin 196 D metabolites and physical performance. Age, sex, smoking, alcohol intake, physical activity, 197 time spent outdoors, season, and body mass index (BMI) were included in regression models 198 as covariates (21-23, 31). The association between vitamin D metabolites and physical 199 performance was analyzed in two steps. Serum 1,25(OH)<sub>2</sub>D and 24,25(OH)<sub>2</sub>D were included 200 201 in step one, and 25(OH)D was added in step two, so the relationship between serum 1,25(OH)<sub>2</sub>D and 24,25(OH)<sub>2</sub>D, and physical performance could be examined, with and 202 without 25(OH)D. Given the large inter-individual differences in metabolites, these variables 203 204 were standardized by scaling them relative to their standard deviation to improve the interpretation of beta coefficients. Cohen's  $f^2$  effect sizes were calculated using a standard 205 formula (32). No signs of strong heteroscedasticity or deviations from a normal distribution 206

were detected for model residuals. Sensitivity analyses (data not shown) conducted with 207 vitamin D metabolites log transformed or categorized into tertiles, as described previously 208 209 (33), resulted in no substantive changes to the null-hypotheses tests or effect sizes. Variance inflation factor for all multiple regression models was <4.2, indicating no presence of 210 multicollinearity (34). The association between vitamin D metabolites and physical 211 performance was also explored by clustering participants into groups based on two 212 213 dimensions of serum 25(OH)D and 1,25(OH)2D:24,25(OH)2D ratio. Clustering was performed using a k-means technique and the Bayesian information criterion to select the 214 215 number of clusters, with the *n* in each cluster determined by the algorithm (35). Pairwise comparisons of the mean differences in physical performance across clusters were made 216 using the t-distribution. Multiple linear regression models were also used to investigate the 217 association between VDR SNPs and physical performance. Separate models were fitted for 218 each of the SNPs whilst controlling for the same variables used to assess the association 219 between vitamin D metabolites and physical performance. Associations were evaluated by 220 conducting F-tests for nested linear models. Pairwise comparisons of the mean differences 221 between vitamin D metabolites across seasons were made using the t-distribution. All 222 statistical tests were conducted within a general linear model framework and using R 3.6.2 (R 223 Foundation for Statistical Computing, Vienna, Austria). Statistical significance was accepted 224 225 at *P* < 0.05.

226

#### 227 **RESULTS**

#### 228 Vitamin D metabolites and season

There was some seasonal variation in vitamin D metabolites (P < 0.001, Table 2). Across all seasons, 66.7% of participants were vitamin D sufficient, 21.4% were insufficient, and 11.9%

- were deficient. During winter, 30.7% were vitamin D sufficient, 38.6% insufficient, and
  30.7% deficient.
- 233

# 234 Vitamin D metabolite predictors of physical performance

235 *Endurance performance* 

236 Serum 1,25(OH)<sub>2</sub>D was positively associated, and 24,25(OH)<sub>2</sub>D was negatively associated

with 2.4 km run time after controlling for age, sex, smoking, alcohol intake, physical activity,

time spent outdoors, season, and BMI (P < 0.01 and P < 0.001, respectively; Table 3). These

relationships remained following the addition of serum 25(OH)D as a predictor (P < 0.001

and P < 0.01, respectively), with 25(OH)D negatively associated with 2.4 km run time (P < 0.01)

241 0.001). Vitamin D metabolites (25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D) together

explained 5.0% of the variance in 2.4 km run time (significant  $\Delta F P < 0.001$ ).

243

# 244 Muscle strength

Serum 1,25(OH)<sub>2</sub>D was negatively associated, and 24,25(OH)<sub>2</sub>D was positively associated

246 with maximal dynamic lift strength (muscle strength) after controlling for age, sex, smoking,

247 alcohol intake, physical activity, time spent outdoors, season, and BMI ( $P \le 0.01$  and  $P \le 0.01$ 

248 0.001, respectively; Table 3). Following the addition of serum 25(OH)D as a predictor,

- 249  $1,25(OH)_2D$  remained negatively associated with muscle strength (P < 0.001), but neither
- 250 24,25(OH)<sub>2</sub>D nor 25(OH)D were associated with muscle strength (P = 0.126 and P = 0.093,
- respectively). Vitamin D metabolites (25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D) together
- explained 0.7% of the variance in muscle strength (significant  $\Delta F P < 0.001$ ).

253

254

256 *Muscle power* 

- 257 Serum 1,25(OH)<sub>2</sub>D was negatively associated, and 24,25(OH)<sub>2</sub>D was positively associated
- with vertical jump peak power output (muscle power) after controlling for age, sex, smoking,
- alcohol intake, physical activity, time spent outdoors, season, and BMI (P < 0.05 and P < 0.05
- 260 0.001, respectively; Table 3). Following the addition of serum 25(OH)D as a predictor,
- 261  $1,25(OH)_2D$  remained negatively associated (P < 0.01), 24,25(OH)<sub>2</sub>D was not associated (P
- 262 = 0.791), and 25(OH)D was positively associated with muscle power (P < 0.001). Vitamin D
- 263 metabolites (25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D) together explained 0.9% of the
- 264 variance in muscle power (significant  $\Delta F P < 0.001$ ).
- 265

# 266 Relative concentrations of vitamin D metabolites and physical performance

- 267 *Endurance performance*
- Run times were faster in participants within clusters 3 to 6 vs cluster 1 and 2, and participants

within clusters 4 to 6 vs cluster 3 (P < 0.05, Fig. 1A).

270

271 *Muscle strength* 

- 272 Maximal dynamic lift strength was higher in participants within clusters 4 to 6 vs cluster 1,
- and participants within clusters 4 and 6 vs cluster 2 (P < 0.05, Fig. 1B).
- 274

275 *Muscle power* 

- 276 Vertical jump peak power output was higher in participants within clusters 3 to 6 vs cluster 1
- 277 (*P* < 0.05, Fig. 1C).
- 278
- 279
- 280

#### 281 Vitamin D receptor polymorphisms and physical performance

Vitamin D receptor SNPs did not explain any of the variance in 2.4 km run time, muscle strength, or muscle power after controlling for age, sex, smoking, alcohol intake, physical activity, time spent outdoors, season, and BMI (significant  $\Delta F P \ge 0.306$ , Table 4). There were no between genotype differences in 2.4 km run time, muscle strength, or muscle power when no confounding factors were controlled for ( $P \ge 0.086$ ).

287

#### 288 DISCUSSION

289 Serum 25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D were associated with 2.4 km run time; 1,25(OH)<sub>2</sub>D was associated with muscle strength; and 25(OH)D and 1,25(OH)<sub>2</sub>D were 290 associated with muscle power, in young healthy adult men and women. Other factors 291 292 contributing to physical performance (age, sex, smoking, alcohol intake, physical activity, 293 time spent outdoors, season, and BMI) were controlled for as covariates using hierarchical multiple linear regression. Vitamin D metabolites (25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D) 294 together explained variances of 5.0% in 2.4 km run time, 0.7% in muscle strength, and 0.9% 295 in muscle power. In terms of practical significance, the magnitude of the association between 296 vitamin D metabolites and physical performance can be considered small (Cohen's  $f^2$  effect 297 sizes <0.15). Nevertheless, in real-world terms, for every 1 SD increase in 25(OH)D (+28.0 298 nmol·L<sup>-1</sup>), 2.4 km run time was 12 s faster and vertical jump peak power output 90 W higher; 299 for every 1 SD increase in 24,25(OH)<sub>2</sub>D (+3.3 nmol·L<sup>-1</sup>), 2.4 km run time was 9 s faster; and 300 for every 1 SD increase in 1,25(OH)<sub>2</sub>D (+36.5 pmol·L<sup>-1</sup>), 2.4 km run time was 6 s slower, 301 maximal dynamic lift strength 1 kg lower, and vertical jump peak power output 38 W lower. 302 303 As hypothesized, serum 25(OH)D and 1,25(OH)2D:24,25(OH)2D had a dynamic relationship 304

305 with physical performance: 2.4 km run times were faster, and muscle strength and muscle

power were greater in men and women with proportionally greater conversion of 25(OH)D to
24,25(OH)<sub>2</sub>D relative to 1,25(OH)<sub>2</sub>D (*i.e.*, low 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio). Examining
this relationship between vitamin D metabolites provides a unique insight into how the
vitamin D metabolic pathway is related to physical performance and suggests that
24,25(OH)<sub>2</sub>D may have role in optimising physical performance. Contrary to our hypothesis,
VDR SNPs (rs2228570, rs4516035, and rs7139166) were not associated with physical
performance.

313

# 314 Vitamin D metabolite predictors of physical performance

The negative association between serum 24,25(OH)<sub>2</sub>D and 2.4 km run time, and positive 315 associations between 24,25(OH)<sub>2</sub>D and muscle strength and muscle power, were weaker or 316 317 absent when 25(OH)D was added as a predictor because of the tight correlation between these metabolites (4). Serum 25(OH)D was itself negatively associated with 2.4 km run time 318 and positively associated with muscle power. In contrast, 1,25(OH)<sub>2</sub>D was positively 319 associated with 2.4 km run time and negatively associated with muscle strength and muscle 320 power, even when 25(OH)D was included as a predictor. However, serum 1,25(OH)2D alone 321 does not reflect vitamin D reserves or status because it is tightly regulated by the 322 hydroxylation enzymes expressed by CYP27B1 and CYP24A1, with 1,25(OH)2D production 323 upregulated by PTH and downregulated by fibroblast growth factor 23 (FGF23) (36). No 324 325 metabolites were associated with muscle strength when controlling for age and sex in the only published study to examine 25(OH)D,  $1,25(OH)_2D$ , and  $24,25(OH)_2D$  relationships with 326 muscle function (37). This non-significant finding may be explained by the small sample size 327 328 (116 adults, 20–74 years) (37), relative to the present study.

329

330

**Relative concentrations of vitamin D metabolites and physical performance** 331 An inverse exponential relationship exists between serum 25(OH)D and 332 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio (4). As the availability of 25(OH)D as a precursor 333 diminishes, the conversion of 25(OH)D to 24,25(OH)<sub>2</sub>D is reduced, resulting in a 334 proportional increase in 1,25(OH)<sub>2</sub>D (38). Superior physical performance in adults with low 335 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio suggests 24,25(OH)<sub>2</sub>D is not a purely catabolic metabolite. 336 337 Given 1,25(OH)<sub>2</sub>D is the most biologically active vitamin D metabolite, greater circulating concentrations of 1,25(OH)<sub>2</sub>D relative to 24,25(OH)<sub>2</sub>D (*i.e.*, high 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D) 338 339 might be expected to be favourable for physical performance (16). Our novel finding that proportionally greater conversion of 25(OH)D to 24,25(OH)2D relative to 1,25(OH)2D (*i.e.*, 340 low 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio) was better for endurance performance, muscle strength, 341 and muscle power, suggests 24,25(OH)<sub>2</sub>D might influence physical performance. Rather than 342 being a catabolic waste product, emerging evidence indicates 24,25(OH)<sub>2</sub>D has a role in 343 osteoblastic differentiation and bone development (39), promotion of fracture healing (7, 8, 344 40), and protection against cartilage damage (6). The existence of 24,25(OH)<sub>2</sub>D receptors, 345 and their possible function relevant to physical performance needs to be examined. 346 347 Vitamin D metabolites may improve physical performance by increasing the delivery of 348 oxygenated blood to muscle through improved endothelial function (13, 14) and maintenance 349

350 of normotension (41). Vitamin D metabolites can also increase mitochondrial oxidative

function, potentially attenuating the development of skeletal muscle fatigue (12). By

potentially increasing aerobic capacity, these mechanisms could account for why the

associations between vitamin D metabolites and 2.4 km run time were the most consistent.

354 Whether similar relationships exist between vitamin D metabolites and performance in

355 endurance events of longer duration is an interesting area for future study. Vitamin D

metabolites may exert beneficial downstream effects on physical performance by modulating
muscle remodelling, since the VDR/retinoid X receptor signaling pathway is upregulated
during the early stages of hypertrophy (42). Vitamin D can enhance skeletal muscle repair
following damaging exercise and help to protect against infection by supporting aspects of
innate and acquired immunity (16). By doing so, vitamin D may help to minimise the number
of training sessions missed by athletes and military personnel, and thus potentially lead to
improved physical performance.

363

# 364 Vitamin D receptor polymorphisms and physical performance

This is the largest study to examine the relationship between VDR SNPs (rs2228570, 365 rs4516035, and rs7139166) and physical performance in young adults, with no associations 366 observed. Previously, the rs2228570 allele related to increased VDR function was associated 367 with weaker muscle strength in older adults (men and women, mean 62 years (43); men, 58-368 93 years (44)). Increased VDR function may increase CYP24A1 expression, leading to the 369 degradation and decreased availability of 1,25(OH)<sub>2</sub>D (9). Associations between the 370 rs2228570 polymorphism and quadriceps strength were no longer statistically significant 371 after controlling for fat-free mass in women (mean 42 years) (45) and men (58–93 years) 372 (44), suggesting this polymorphism may influence muscle mass rather than strength per se. In 373 374 contrast to the present study, strength differed between groups of children (mean 10 years) 375 with different rs4516035 alleles, but no differences emerged for rs2228570 alleles (46). The range of muscle or other performance assessments used, and differences in participants' 376 fitness and age (skeletal muscle VDR expression decreases with age (47)) have contributed to 377 378 these equivocal findings.

379

#### 381 **Perspectives**

382 This study provides a unique insight into the dynamic relationship between vitamin D

- metabolites and demonstrates that the relative circulating concentrations of  $1,25(OH)_2D$  and
- 384 24,25(OH)<sub>2</sub>D are related to physical performance. Randomized controlled trials have shown
- vitamin D<sub>3</sub> supplementation does not improve physical performance (24), however, in these
- studies vitamin D status has been assessed using 25(OH)D in isolation and the relative
- concentrations of 1,25(OH)<sub>2</sub>D and 24,25(OH)<sub>2</sub>D have not been considered. Shifting
- $1,25(OH)_2D:24,25(OH)_2D$  ratio from high to low may be necessary for a beneficial effect on
- 389 performance to occur. Whether oral vitamin D<sub>3</sub> supplementation can correct high

 $1,25(OH)_2D:24,25(OH)_2D$  ratio and achieve high 25(OH)D and low

- 391 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio—and thereby enhance physical performance—remains to be
- determined. How much and how often oral vitamin  $D_3$  is needed to achieve a steady state of
- vitamin D metabolites needs to be examined. Avoiding high serum 24,25(OH)<sub>2</sub>D has been

recommended because 24,25(OH)<sub>2</sub>D may act to block the activity of the VDR (48). This

- recommendation, however, assumes 24,25(OH)<sub>2</sub>D is purely a catabolic waste product—a
- 396 hypothesis that warrants further evaluation given that relatively high 24,25(OH)<sub>2</sub>D was

397 associated with superior performance.

398

Avoiding a relative increase in serum  $1,25(OH)_2D$  may be beneficial for performance,

400 therefore, supplementation with alfacalcidol (1-hydroxyvitamin D<sub>3</sub>) or calcitriol

401 (1,25(OH)<sub>2</sub>D) are unlikely to be effective for enhancing physical performance—especially

402 because serum 1,25(OH)<sub>2</sub>D is maintained within a tight range, despite fluctuations in

- 403 25(OH)D and  $24,25(OH)_2D$  (4, 36). Supplementation that increases  $1,25(OH)_2D$  beyond its
- 404 normal plateau, thereby increasing the 1,25(OH)<sub>2</sub>D:24,25(OH)<sub>2</sub>D ratio, could be detrimental.
- 405

406	Genome wide studies have identified CYP24A1 as one of the major genetic determinants of
407	variability in vitamin D metabolism (49). Increased CYP24A1 activity increases
408	$24,25(OH)_2D$ and decreases $1,25(OH)_2D$ and PTH (36). Whether supplementation, dietary or
409	lifestyle interventions can be used in individuals with genetically lower CYP24A1 activity (as
410	indicated by increased 1,25(OH) <sub>2</sub> D:24,25(OH) <sub>2</sub> D ratio), to manage their vitamin D
411	metabolism and enhance physical performance requires future study.
412	
413	The present study is limited by its cross-sectional design. Associations between vitamin D
414	metabolites and physical performance could be explained by reverse causation, <i>i.e.</i> , fitter,

415 more physically active individuals spend more time outdoors exposed to sunlight and, in-turn,

416 had higher serum concentrations of vitamin D metabolites. However, we included

417 participants' self-reported physical activity levels and typical time spent outdoors as

418 covariates in our regression models. Almost all of the participants in the present study were

419 white Caucasian. Whether the vitamin D metabolites and SNPs we examined are associated

420 with physical performance in other ethnic groups is unknown and warrants further study.

421

# 422 Conclusions

423 Serum 25(OH)D, 1,25(OH)<sub>2</sub>D, and 24,25(OH)<sub>2</sub>D were associated with 2.4 km run time;
424 1,25(OH)<sub>2</sub>D was associated with muscle strength; and 25(OH)D and 1,25(OH)<sub>2</sub>D were

425 associated with muscle power, after controlling for covariates. Vitamin D metabolites

426 accounted for a small portion of variance in physical performance. Polymorphisms in the

427 VDR were not associated with physical performance. Faster 2.4 km run times and greater

428 muscle strength and muscle power in adults with proportionally greater conversion of

429 25(OH)D to  $24,25(OH)_2D$  relative to  $1,25(OH)_2D$  (*i.e.*, low  $1,25(OH)_2D:24,25(OH)_2D$  ratio)

430 indicates  $24,25(OH)_2D$  may have a role in optimising physical performance.

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- 435

# 436 CONFLICT OF INTEREST

- 437 The authors have nothing to disclose. The results of the present study do not constitute
- 438 endorsement by the ACSM. The results of the study are presented clearly, honestly, and
- 439 without fabrication, falsification, or inappropriate data manipulation.

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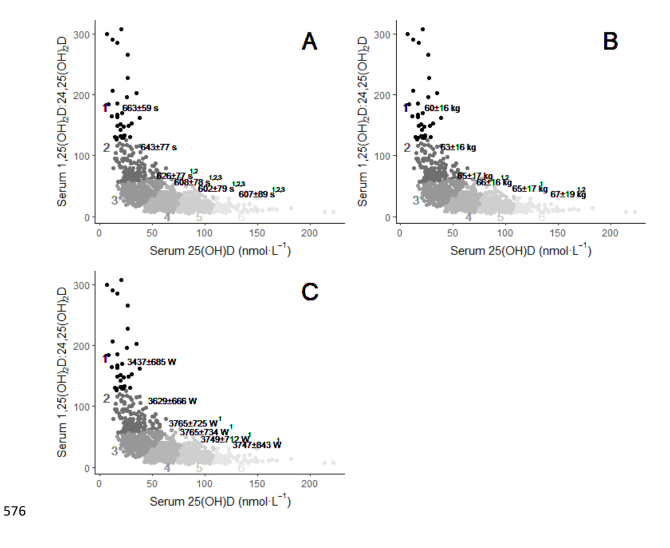
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#### 570 **FIGURE CAPTIONS**

Figure 1. Dynamic relationships between vitamin D metabolites and physical performance. 571

Participants (1 per filled circle) are categorized into one of six clusters, with each cluster's 572

- mean  $\pm$  SD physical performance shown. Panel A, 2.4 km run time; Panel B, maximal 573
- dynamic lift strength; and Panel C, vertical jump peak power output.  $^{1}P < 0.05$  vs cluster 1; 574
- $^{2}P < 0.05$  vs cluster 2;  $^{3}P < 0.05$  vs cluster 3. 575



**Table 1.** Demographic, anthropometric, lifestyle behavior, and physical performance

 characteristics.

Demographics				
Age (years)	$22.0\pm2.8$			
Sex (% men)	78.9			
Ethnicity				
White Caucasian (%)	94.8			
Other (%)	5.2			
Anthropometrics				
Body mass (kg)	$74.1\pm10.4$			
Height (m)	$1.75\pm0.08$			
BMI (kg·m <sup>-2</sup> )	$24.1\pm2.5$			
Lifestyle behaviors				
Smoker (%)	36.6			
Alcohol user (%)	89.5			
Physical activity (h-week-1)	$9.6\pm11.2$			
Time spent outdoors (h·week <sup>-1</sup> )	<1	1–3.5	3.5–6	>6
April-September (%)	2.3	28.7	33.1	35.9
October-March (%)	6.7	39.0	26.7	27.6
Physical performance				
2.4 km run time (s)	$617\pm79$			
Maximal dynamic lift strength (kg)	$65\pm17$			
Vertical jump peak power output (W)	$3702\pm748$			

BMI, body mass index. Data are mean  $\pm$  SD or precent.

	Spring	Summer	Fall	Winter	All Seasons	
	<i>n</i> = 358	<i>n</i> = 472	<i>n</i> = 394	<i>n</i> = 303	<i>n</i> = 1527	
Vitamin D status						
Sufficient (%)	66.8	85.6	71.8	30.7	66.7	
Insufficient (%)	22.9	11.9	18.0	38.6	21.4	
Deficient (%)	10.3	2.5	10.2	30.7	11.9	
25(OH)D (nmol·L <sup>-1</sup> )	$62.3\pm26.8^{\text{ a}}$	$76.8\pm25.8$	$65.2\pm26.0^{\text{ a}}$	$43.5 \pm 22.7$ a,b,c	$63.8\pm28.0$	
1,25(OH)2D (pmol·L <sup>-1</sup> )	$141.8\pm34.8$	$142.9\pm36.5$	$132.1 \pm 37.1$ <sup>a,c</sup>	$131.1 \pm 35.8$ <sup>a,c</sup>	$137.5 \pm 36.5$	
24,25(OH)2D (nmol·L <sup>-1</sup> )	$5.1\pm3.0^{\rm \ a,b}$	$6.6\pm3.0$	$6.5 \pm 3.2$	$3.8\pm3.2$ <sup>a,b,c</sup>	$5.6\pm3.3$	
1,25(OH)2D:24,25(OH)2D	$40.2\pm32.1~^{d}$	$28.3\pm21.6^{\text{ c,d}}$	$26.9\pm20.6~^{\text{c,d}}$	$52.8\pm41.3$	$35.6\pm30.5$	
25(OH)D:24,25(OH)2D	$13.8\pm4.3$	$12.7\pm3.8^{c,d}$	$11.1 \pm 3.3$ <sup>a,c,d</sup>	$13.7\pm5.4$	$12.8\pm4.3$	

Table 2. Seasonal variation in vitamin D status and serum vitamin D metabolites.

Vitamin D sufficient, serum  $25(OH)D \ge 50 \text{ nmol} \cdot L^{-1}$ ; insufficient, serum 25(OH)D 30 - <50

nmol·L<sup>-1</sup>; and deficient, serum 25(OH)D <30 nmol·L<sup>-1</sup>. Data are mean  $\pm$  SD or percent. a,

lower than summer; b, lower than fall; c, lower than spring; d, lower than winter, P < 0.001.

**Table 3.** Serum 1,25(OH)<sub>2</sub>D, 24,25(OH)<sub>2</sub>D, and 25(OH)D predictors of 2.4 km run time (endurance), muscle strength (maximal dynamic lift), and muscle power (vertical jump peak power output).

	Serum vitamin D					
	metabolites	Beta	R <sup>2</sup>	$\Delta R^2$	Sig. ΔF	$f^2$
2.4 km run time	1,25(OH) <sub>2</sub> D	4.1**	0.488	0.044	< 0.001	0.09
	24,25(OH) <sub>2</sub> D	-18.2***				
	1,25(OH) <sub>2</sub> D	5.6***	0.494	0.050	< 0.001	0.10
	24,25(OH) <sub>2</sub> D	-8.9**				
	25(OH)D	-12.0***				
Muscle strength	1,25(OH) <sub>2</sub> D	-0.95**	0.668	0.007	< 0.001	0.02
	24,25(OH) <sub>2</sub> D	1.41***				
	1,25(OH) <sub>2</sub> D	-1.06***	0.668	0.007	< 0.001	0.02
	24,25(OH) <sub>2</sub> D	0.75				
	25(OH)D	0.86				
Muscle power	1,25(OH) <sub>2</sub> D	-27.8*	0.672	0.006	< 0.001	0.02
	24,25(OH) <sub>2</sub> D	63.7***				
	1,25(OH) <sub>2</sub> D	-38.4**	0.675	0.009	< 0.001	0.03
	24,25(OH) <sub>2</sub> D	-5.5				
	25(OH)D	90.1***				

After controlling for covariates (age, sex, smoking, alcohol intake, physical activity, time spent outdoors, season, and BMI) serum 1,25(OH)<sub>2</sub>D and 24,25(OH)<sub>2</sub>D were entered in step one, and 25(OH)D was entered in step two as predictors of physical performance. Beta, standardized beta coefficient; Sig.  $\Delta$ F, significant F change *P* value; *f*<sup>2</sup>, Cohen's *f*<sup>2</sup> effect size, *f*<sup>2</sup> ≥0.02, ≥0.15 and ≥0.35 represent small, medium and large effect sizes, respectively (32). \**P* < 0.05, \*\**P* < 0.01 and \*\*\**P* < 0.001.

**Table 4.** VDR SNP (rs2228570, rs4516035, and rs7139166) predictors of 2.4 km run time (endurance), muscle strength (maximal dynamic lift), and muscle power (vertical jump peak power output).

	VDR SNP	Beta	Beta	R <sup>2</sup>	$\Delta R^2$	Sig. <b>A</b> F	$f^2$
		CC vs alternate	CC vs alternate				
		allele 1	allele 2				
2.4 km run time	rs2228570	0.48	-4.0	0.429	0.000	0.610	< 0.001
	rs4516035	0.59	-0.23	0.429	0.000	0.716	< 0.001
	rs7139166	0.71	0.56	0.429	0.000	0.978	< 0.001
Muscle strength	rs2228570	0.39	0.52	0.650	0.000	0.743	< 0.001
	rs4516035	0.57	0.86	0.650	0.000	0.575	< 0.001
	rs7139166	-0.29	-0.88	0.650	0.000	0.599	< 0.001
Muscle power	rs2228570	19.4	19.0	0.666	0.000	0.610	< 0.001
	rs4516035	39.2	51.1	0.666	0.000	0.306	0.002
	rs7139166	-12.2	-50.8	0.666	0.000	0.312	0.002

After controlling for covariates (age, sex, smoking, alcohol intake, physical activity, time spent outdoors, season, and BMI), VDR SNP genotypes were entered as predictors of physical performance. rs2228570: 40%, 44%, and 16% of participants had CC, CT (alternate allele 1), and TT (alternate allele 2) genotypes; rs4516035: 18%, 46%, and 36% of participants had CC, TC (alternate allele 1), and TT (alternate allele 2) genotypes; rs7139166: 36%, 46%, and 18% of participants had CC, GC (alternate allele 1), and GG (alternate allele 2) genotypes. VDR, vitamin D receptor; SNP, single-nucleotide polymorphism; Beta, standardized beta coefficient; Sig.  $\Delta$ F, significant F change P value;  $f^2$ , Cohen's  $f^2$  effect size,  $f^2 \ge 0.02$ ,  $\ge 0.15$  and  $\ge 0.35$  represent small, medium and large effect sizes, respectively (32).