Accelerated Muscle Deoxygenation in Aerobically Fit Subjects During Exhaustive Exercise Is Associated With the ACE Insertion Allele

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Introduction: The insertion/deletion (I/D) polymorphism in the gene for the major regulator of vascular tone, angiotensin-converting enzyme-insertion/deletion (ACE-I/D) affects muscle capillarization and mitochondrial biogenesis with endurance training. We tested whether changes of leg muscle oxygen saturation (SmO₂) during exhaustive exercise and recovery would depend on the aerobic fitness status and the ACE I/D polymorphism.

Methods: In total, 34 healthy subjects (age: 31.8 ± 10.2 years, 17 male, 17 female) performed an incremental exercise test to exhaustion. SmO₂ in musculus vastus lateralis (VAS) and musculus gastrocnemius (GAS) was recorded with near-IR spectroscopy. Effects of the aerobic fitness status (based on a VO₂peak cutoff value of 50 ml O₂ min⁻¹ kg⁻¹) and the ACE-I/D genotype (detected by PCR) on kinetic parameters of muscle deoxygenation and reoxygenation were assessed with univariate ANOVA.

Results: Deoxygenation with exercise was comparable in VAS and GAS (p = 0.321). In both leg muscles, deoxygenation and reoxygenation were 1.5-fold higher in the fit than the unfit volunteers. Differences in muscle deoxygenation, but not VO₂peak, were associated with gender-independent (p > 0.58) interaction effects between aerobic fitness × ACE-I/D genotype; being reflected in a 2-fold accelerated deoxygenation of VAS for aerobically fit than unfit ACE-II genotypes and a 2-fold higher deoxygenation of GAS for fit ACE-II genotypes than fit D-allele carriers.

Discussion: Aerobically fit subjects demonstrated increased rates of leg muscle deoxygenation and reoxygenation. Together with the higher muscle deoxygenation in aerobically fit ACE-II genotypes, this suggests that an ACE-I/D genotype-based personalization of training protocols might serve to best improve aerobic performance.

Keywords: cycling, aerobic metabolism, oxygen saturation, gene, exhaustive pedaling
INTRODUCTION

Exercise pronouncedly elevates the energy expenditure of working skeletal muscle due to increased ATPase activity of the actin-myosin filaments and ion pumps (Rolfe and Brown, 1997). The resultant energy demand is met by enhanced aerobic combustion of organic substrates in mitochondria (Tonkonogi and Sahlin, 2002; Ryan et al., 2014); giving rise to the commonly observed oxygen deficit with exercise (Lukin and Ralston, 1962; Nioka et al., 1998). The implicated genetic mechanism is mediated by the presence of a 287-basepair insertion (the I-allele) in intron 16 of the ACE gene silences tissue expression of ACE transcripts, reducing ACE activity and angiotensin 2 levels in the blood; whereas its absence (i.e., deletion, D-allele) is associated with the inverse effects (Jackman et al., 2008; Vaughan et al., 2013; Mathes et al., 2015; van Ginkel et al., 2016; Fluck et al., 2019). ACE I-allele carriers demonstrate exaggerated gains in the volume density of mitochondria in knee extensor muscle with repeated endurance training (Vaughan et al., 2013; Fluck et al., 2019); indicating that ACE-insertion/deletion (I/D) genotype-associated differences in aerobic metabolism develop with repeated sessions of endurance exercise.

Importantly, healthy ACE-DD genotypes demonstrate a lower muscle capillarization and a reduced capillary perfusion with endurance exercise than ACE I-allele carriers (van Ginkel et al., 2015; Valdivieso et al., 2017). Collectively, the observed ACE-I/D-related response to acute and repeated endurance exercise indicated either an elevated vasoconstriction or a lower potential for vasodilatation, in ACE D-allele carriers; giving rise to a lowered capacity for oxygen delivery to recruited skeletal muscles during and after exercise (Buikema et al., 1996; O’Donnell et al., 1998; van Dijk et al., 2000; Williams et al., 2000; Woods et al., 2002; Fluck and Hoppeler, 2003; Fluck et al., 2010).

Whether the reported influence of the ACE-I/D gene polymorphism on mitochondria and capillary processes and connected metabolic processes of skeletal muscle results in differences in muscle oxygen saturation during intense exercise is not known. Toward this end we tested the hypotheses, the ACE-I/D polymorphism is associated with (i) an accelerated and more extensive deoxygenation in aerobically fit subjects I-allele carriers compared to non-I-allele carriers, and whether (ii) the reoxygenation is accelerated in ACE I-allele carriers and may be further accelerated by a good fitness state.

METHODS

Study Design

This study had a cross-sectional design in which the subjects performed a ramp test to determine VO_{2peak} Peak Power Output (PPO) and SmO$_2$ in a knee extensor muscle (VAS) and an ankle extensor muscle (GAS). Prospective power analysis on the association of the ACE-I/D genotype with aerobic processes in skeletal muscle indicated that a total number of 24 replicates (distributing equally according to each of the four combinations of genotype and fitness type) is sufficient enough to reveal statistically significant associations between the ACE-I/D genotype × fitness state and differences in muscle deoxygenation (Supplementary Table 1). As the average age of participants was 31.81 ± 10.18 years in line with standard guidelines subjects were assigned as being aerobically fit or unfit based on whether their respective VO_{2peak} met the criteria of being above or below 50 mL·min^{-1}·kg^{-1} (Valdivieso et al., 2017; ACSM, 2021). ACE I/D gene polymorphism genotyping was performed retrospectively.
Subjects/Ethics
A total of 34 recreational active healthy subjects, i.e., 17 women and 17 men, participated in this study. Subject recruitment was by word-of-mouth and a public announcement with flyers in the professional or private environment of the research group. All subjects volunteered after their self-assessment that they met the requirements of presenting a good health and an age between 20 and 70 years. Upon the provision of an informed consent, the further specific inclusion criterion of an unobtrusive cardiovascular system was verified based on an inconspicuous ECG during exercise in a ramp test on a cycle ergometer. The exclusion criteria comprised evidence for a relevant valvular heart disease, arterial hypertension (blood pressure at rest > 140/90 mm Hg), arrhythmogenic cardiomyopathy, smoker, drug, or alcohol disease, known or suspected non-compliance with the protocol, or a contraindication for ethical reasons. All the 34 participating volunteers qualified for the inclusion in this study. The study has been approved by the Ethics Committee of the Canton of Zurich. All the investigations were conducted in accordance with the ethical standards of the Declaration of Helsinki of 1964.

Ramp Test
Subjects performed a test of incremental exercise on an electrically braked cycle ergometer (Ergoselect 200, Ergoline, Bitz, Germany, UK) being accompanied with spiroergometric and NIRS measurements. Prior to the test, anthropometric data (height and body mass) were collected, and the body mass index (BMI) was calculated. A resting ECG was recorded and verified by a physician to ensure that the subjects did not demonstrate a contraindication to conduct the ramp test.

The test protocol was conducted in an air-conditioned laboratory at a standardized temperature of 20°C according to a modified version of a published protocol (Whipp et al., 1981). In brief, the test started with a 3 min period of rest, when subjects sat still on the cycle ergometer while maintaining a normal breathing pattern. Subsequently, subjects started pedaling at an initial power (75 W for women and 100 W for men). Target power was increased every 20 s (18 W·min⁻¹ for women and 30 W·min⁻¹ for men). The subjects were asked to keep a constant self-chosen pedal cadence throughout the test (optimally between 70 and 100 rpm). The test was stopped when the subjects experienced volitional exhaustion and/or were not able to maintain the target pedal cadence. Subsequently, recordings continued during a period of 8 min, when subjects rested in a seated position on the cycle ergometer.

Pulmonary gas exchange (oxygen uptake) was measured after proper calibration through a breath-by-breath spiroergometry system (MetaLyzer® 3B-R2, CORTEX Biophysics, Leibzig, Germany). VO₂peak and PPO were defined as the last achieved, and peak, values before exhaustion manifested. To non-invasively measure SmO₂ during the ramp test, a muscle NIRS
monitor (Moxy, Fortiori Design LLC, Minnesota, USA) was used as established (Fitze et al., 2019). The device uses four different light sources covering wavelengths in the range of 630 to 850 nm and a modified Beer–Lambert law to perform measurements of SmO2. For measurements, two NIRS monitors were positioned on specific sites of the skin that were shaved using a disposable razor (Gallant, Dynarex, Orangeburg, USA) and cleaned with an alcohol swab (WebcoTM, Covidiem™, Dublin, Ireland). One monitor was placed on the lower third of the VAS in the middle of the muscle belly on the left leg of the subjects. The other monitor was placed on the GAS, onto a fictive line between the Malleolus medialis and medial plateau of the tibia. Sensors were attached using the recommended tape (Moxy Adhesive Attachments, Fortiori Design LLC, Minnesota, USA). In order to protect the NIRS device from ambient light, it was covered with an adhesive non-woven fabric (Hypafix®, BSJ medical, Hamburg, Germany). Recording started synchronized with the start of the ramp test (with 3 min of rest).

Genotyping

Buccal swabs were collected with an ear-bud, air-dried in a laboratory fume cupboard (Seculow 1500, Waldner, Wangen, Germany, UK) for 2 h and then stored at +4°C. Subjects were told not to consume any food or to drink in the 30 min prior to sample collection.

Deoxyribonucleic acid extraction was performed according to a commercially available protocol (QIAamp® DNA Mini Kit, Qiagen, Hilden, Germany, UK). In brief, the cotton swab was separated from the stick with scissors, followed by incubation steps in a 2 ml microcentrifuge tube to degrade contaminating ribonucleic acids, lyse the cells with QIAGEN® proteinase K, and enrich the contained genomic DNA with QIAamp Mini spin columns with the help of an air-cooled microcentrifuge (Prysm™, Labnet International, Edison, USA). The resulting sample (150 µl) elute was stored at +4°C until genotyping was performed.

Genotyping was carried out by polymerase chain reactions in 48 well-plates followed by high-resolution melt analysis using a real-time PCR system (EcoTM, illumina®, San Diego, USA) according to the instructions. The reaction mix per well-included for each sample 2 µl of the DNA solution, 1 µl distilled H2O2, 1 µl of MgCl2 (25 mmol), 5 µl of KAPA HRM FAST Master Mix (2×) and 1 µl of the 1- or D-allele-specific primer mix (2 µmol). The primer mix for the detection of the 66 bp amplex, which is specific for the I-allele, contained the primer ACE2 (5′-TGGGATACAGGCAGTATACAG-3′) and the primer ACE3 (5′-ATTCCAGCTGGATAAAATT-3′). The primer mix for the detection of the 83 bp amplex, which is specific to the D-allele, contained primers ACE1 (5′-CATCCTTTCTCCCATTTCTC-3′) and ACE3 (5′-ATTTCAGAGCTGGATAAAATT-3′). The sealed plate was centrifuged to remove any bubbles and was submitted to a standardized thermal protocol as published (Valdivieso et al., 2017). Genotype analysis was carried out using a genetic variation analysis software (EcoStudy Version 5.0, Illumina®, San Diego, California, USA). The respective genotype was verified based on the presence of an allele-specific melting curve for the amplified products in the respective PCR reactions, as established by microsequencing of PCR reactions from reference samples at a commercial provider (Microsynth, Balgach, Switzerland, UK) (Valdivieso et al., 2017).

Data Processing

A representative example of the recorded and processed timeline of SmO2 during the ramp test, as well as the extracted parameters,
TABLE 2 | Association between fitness status and angiotensin-converting enzyme-insertion/deletion (ACE-I/D) genotype on parameters of muscle oxygen kinetics during the ramp test of cycling exercise.

| Phase         | Parameter          | Statistical size | Fitness | Genotype | Fitness x genotype | Muscle | Muscle x fitness | Muscle x genotype | Fitness x genotype |
|---------------|--------------------|------------------|---------|----------|-------------------|--------|-----------------|-------------------|-------------------|
| Rest          | SmO2baseline       | p-value          | 0.018   | 0.027    | 0.151             | 0.004  | 0.355           | 0.812             | 0.305             |
|               |                    | η²               | 0.090   | 0.079    | 0.034             | 0.123  | 0.014           | 0.001             | 0.018             |
| Exercise      | SmO2min            | p-value          | ≤0.001  | 0.597    | 0.098             | 0.052  | 0.208           | 0.031             | 0.296             |
|               |                    | η²               | 0.240   | 0.005    | 0.045             | 0.058  | 0.026           | 0.075             | 0.018             |
| Exercise      | Δdeoxygenation     | p-value          | ≤0.001  | 0.141    | 0.020             | 0.321  | 0.891           | 0.174             | 0.920             |
|               |                    | η²               | 0.302   | 0.036    | 0.087             | 0.015  | 0.000           | 0.031             | 0.000             |
| Exercise      | tdeoxygenation     | p-value          | 0.011   | 0.174    | 0.151             | 0.586  | 0.834           | 0.519             | 0.472             |
|               |                    | η²               | 0.104   | 0.031    | 0.034             | 0.005  | 0.001           | 0.007             | 0.009             |
| Exercise      | Slopedeoxygenation | p-value          | 0.006   | 0.598    | 0.006             | 0.175  | 0.902           | 0.306             | 0.575             |
|               |                    | η²               | 0.125   | 0.005    | 0.118             | 0.029  | 0.000           | 0.017             | 0.005             |
| Stop          | SmO2max            | p-value          | 0.013   | 0.364    | 0.357             | ≤0.001 | 0.526           | 0.945             | 0.451             |
|               |                    | η²               | 0.098   | 0.014    | 0.014             | 0.674  | 0.007           | 0.000             | 0.000             |
| Stop          | t1/2deoxygenation  | p-value          | ≤0.001  | 0.705    | 0.101             | ≤0.001 | 0.105           | 0.131             | 0.765             |
|               |                    | η²               | 0.270   | 0.002    | 0.044             | 0.392  | 0.043           | 0.038             | 0.001             |
| Stop          | t1/deoxygenation   | p-value          | 0.262   | 0.818    | 0.571             | 0.123  | 0.822           | 0.659             | 0.828             |
|               |                    | η²               | 0.021   | 0.001    | 0.005             | 0.037  | 0.001           | 0.004             | 0.001             |
| Stop          | Slope1tdeoxygenation| p-value          | 0.031   | 0.598    | 0.365             | 0.823  | 0.682           | 0.607             | 0.592             |
|               |                    | η²               | 0.076   | 0.005    | 0.014             | 0.001  | 0.003           | 0.004             | 0.005             |
| Stop          | SmO2overshoot      | p-value          | 0.838   | 0.273    | 0.687             | ≤0.001 | 0.182           | 0.793             | 0.842             |
|               |                    | η²               | 0.001   | 0.020    | 0.003             | 0.473  | 0.030           | 0.001             | 0.001             |

Values refer to the calculated level of statistical significance (p) and effect size (η²) for the effect of the aerobic fitness state, ACE-I/D genotype (ACE-II vs. ACE-ID/ACE-DD), and muscle type on assessed parameters of deoxygenation and reoxygenation in the leg muscles of the studied 34 subjects. P-values that were deemed statistically significant (i.e., p < 0.05) are underlined. ANOVA with post-hoc test of least significant difference.

is shown in Figure 1. Data pre-processing and analysis were performed as previously described using a data processing program (MATLAB 2015a, The Mathworks, Natick, USA) (Fitze et al., 2019). In short, SmO2 data were filtered using a 2nd order zero-phase shift Butterworth low-pass filter with a cutoff frequency of 0.03 Hz. Extraction of the values for relevant parameters was performed as previously described: SmO2baseline was declared as the mean value of the t3-min prerest period. The minimum SmO2 value during the ramp test (SmO2min) was determined based on the last local minimum of the filtered SmO2 prior to reoxygenation. Δdeoxygenation was set as the difference between SmO2baseline and SmO2min. tdeoxygenation was the time from the beginning of the ramp test until SmO2min was reached. slopedeoxygenation was calculated of the values of Δdeoxygenation over tdeoxygenation. SmO2max was defined as the highest value achieved within the period between the start of reoxygenation and test termination. SmO21/2reoxygenation was defined as 50% of the difference between SmO2max and SmO2min. t1/2reoxygenation was defined as the time between SmO2min and SmO21/2reoxygenation. Slope 1/2reoxygenation was calculated using Δ1/2reoxygenation over t1/2reoxygenation. SmO2overshoot represented the difference between SmO2max and SmO2baseline. Concerning the assumption of a linear process over the ramp protocol before exhaustion of SmO2 and Power a high correlation was detected (R = −0.981 ± 0.214). For the display of the average course of SmO2 in VAS and GAS during the ramp test, the recorded raw values were averaged over 9 s intervals, and the mean and SE over all 34 samples calculated for each time point/9 s interval. For the values from the exercise phases, the “time coordinates” were scaled to the duration of a reference data set for a study participant which ceased pedaling nearest to the duration of the lower 25% quartile of tdeoxygenation, i.e., 358 s.

Statistical Analysis
An online calculator was used to determine whether the observed genotype frequency is consistent with the Hardy–Weinberg equilibrium (Rodriguez et al., 2009). Prospective and retrospective power analyses were conducted with G-Power (version 3.1.9.6, http://www.gpower.hhu.de/) and the Statistical Package for the Social Sciences (SPSS version 23, IBM, Armonk, USA), respectively. Variance homogeneity for sub-samples was analyzed with Levene Test, whereby only for t1/2reoxygenation (p = 0.027) variance inhomogeneity was deduced. Therefore, a multivariate ANOVA was used to assess the effects of the ACE I/D genotype and aerobic fitness status, and their interaction, and the influence of the muscle type and gender, on parameters of muscle oxygenation (SmO2) during the ramp test. A co-dominant genetic model was applied to calculate the effects of the ACE I/D genotype, i.e., carriers vs. non-carriers of the D-allele. A post-hoc test of the least significant difference was applied to localize effects. Statistical analyses and graphical presentations were calculated with SPSS (SPSS version 23, IBM, Armonk,
USA) and assembled for presentation using MS-Office Excel and Powerpoint (Microsoft Office Professional Plus 10, Kildare, Ireland, UK). Significance was declared depending on whether a \( p \)-value below 0.05 was met.

**RESULTS**

**Subjects Characteristics**

Table 1 summarizes selected physiological and characteristics of the 34 volunteers per genotype and fitness status. Aerobically fit subjects demonstrated on average a 14.9 ml O\(_2\) min\(^{-1}\) kg\(^{-1}\) higher specific VO\(_2\) peak and a 112.6 W higher PPO than the unfit subjects. Body mass, BMI, height, and weight did not differ between the aerobically fit and unfit subjects.

The studied population was found to stand in Hardy-Weinberg equilibrium \((p = 0.261)\). The ACE-I/D genotype was not associated with differences in VO\(_2\)peak or PPO independent of whether assessed as single effect \((p = 1.00, p = 0.26)\) or interaction effect with the aerobic fitness state \((p = 0.85, p = 0.44; \text{Table 1})\). BMI was 2.2 kg/m\(^2\) higher in non-carriers than carriers of the ACE D-allele.

**Muscle Oxygen Saturation During the Ramp-Incremental Pedaling Exercise**

Figure 1 visualizes an example of the alterations in SmO\(_2\) in *musculus vastus lateralis* during the ramp-incremental pedaling exercise. SmO\(_2\) was fairly stable during the first phase of rest (SmO\(_2\) baseline) and then fell with the onset of contraction (deoxygenation) at an average rate of \(-0.060 \pm 0.028% \text{SmO}_2\) s\(^{-1}\) to a minimal value (SmO\(_2\) min) until voluntary exhaustion manifested. SmO\(_2\) rapidly recovered with a rate of 0.736 ± 0.459% SmO\(_2\) s\(^{-1}\), to, or above, baseline values with the cessation of exercise (reoxygenation).

Muscle deoxygenation was similar between VAS and GAS muscle \((p = 0.321, \text{Table 2; Figure 2})\). Muscle type differences resolved for parameters of reoxygenation (\(\Delta_1/2\) reoxygenation, SmO\(_2\) max, SmO\(_2\) overshoot; \text{Supplementary Figure 1}), and SmO\(_2\) at baseline \((59.9 vs. 52.9%; p = 0.004), all being higher in VAS than GAS.

**Aerobic Fitness Affects Muscle Deoxygenation and Reoxygenation During Ramp-Incremental Exercise**

Parameters that characterized the deoxygenation and reoxygenation kinetics in the two-leg muscles with ramp-incremental pedaling exercise demonstrated associations with aerobic fitness (Table 2). Figure 3 depicts the differences for the corresponding effect of the aerobic fitness status for the combined values for both the leg muscles.

The values for SmO\(_2\) min \((25.6\%), \Delta\text{deoxygenation (59.1\%)}\) and the slope\(_\text{deoxygenation} (22.1\%)\), were lower in aerobically fit compared to unfit subjects, when the \(t\)\(_\text{deoxygenation} \) was higher in fit than unfit subjects \(+24.2\%). Conversely, the values for \(\Delta_1/2\text{reoxygenation (44.2\%)}\), slope\(_{1/2}\text{reoxygenation (43.9\%)}\) and SmO\(_2\) max \(+9.4\%)\) were higher in aerobically fit than unfit subjects.

When assesses separately for VAS and GAS, the values for SmO\(_2\) min \((24.8 vs. 35.1\%, 22.8 vs. 28.9\%\), \(\Delta\text{deoxygenation (34.9 vs. 21.7\%, 31.1 vs. 19.8\%)}\) and the slope\(_\text{deoxygenation (i.e., 0.065 vs. 0.054 SmO}_2\) s\(^{-1}\), 0.056 vs. 0.045% SmO\(_2\) s\(^{-1}\)) were all lower in aerobically fit compared to unfit subjects. Conversely, the values for \(\Delta_1/2\text{reoxygenation (27.8 vs. 18.4\%, 15.5 vs. 11.6\%)}\) were higher in aerobically fit than unfit subjects. Additionally, the values for baseline SmO\(_2\) min in GAS \((53.9 vs. 48.7\%\), SmO\(_2\) max in VAS \((80.2 vs. 71.8\%\) and \(t\)\(_\text{deoxygenation in VAS (543.7 s vs. 430.3 s)}\) were higher in fit than unfit subjects.

**Differences in Muscle Deoxygenation During Ramp-Incremental Exercise Are Associated With the Interaction Between Aerobic Fitness \(\times\) ACE-I/D Genotype**

Baseline values for SmO\(_2\), alone, were associated with the ACE-I/D genotype \((p = 0.027, \text{Table 2})\). The values of two kinetic parameters resuming muscle deoxygenation, i.e., \(\Delta\text{deoxygenation and slope deoxygenation, demonstrated interactions between the aerobic fitness status and the ACE-I/D genotype (Table 2)}\).

For both parameters, the interaction was localized to the lowest values in fit non-carriers of the ACE D-allele \((\text{Figure 4})\). For \(\Delta\text{deoxygenation, the values in the fit ACE-II genotypes were 29.4\% lower than in fit non-carriers of the D-allele and 56.4\% lower than in unfit ACE-II genotypes. Alike for the slope deoxygenation, the values in fit ACE-II genotypes were 52.5\% lower than in unfit ACE-II genotypes and 31.3% lower than in fit carriers of the D-allele.**

The interaction effect of aerobic fitness status and the ACE-I/D genotype for \(\Delta\text{deoxygenation and slope }\text{deoxygenation was not affected by the muscle type (p = 0.920, p = 0.575), and sex (Supplementary Table 2), where }\Delta\text{deoxygenation and slope }\text{deoxygenation in VAS and GAS were the lowest in the fit ACE-II genotypes (Supplementary Figure 2)}\).

Supplementary Table 3 depicts the average values for the observed parameters of deoxygenation and reoxygenation in both studied leg muscles for each genotype and fitness status.

**DISCUSSION**

The aim of this study was to investigate whether a NIRS-based measure of the balance between supply and use of oxygen during exhaustive pedaling exercise is associated with the ACE-I/D genotype and stands in dependence of the fitness state (Casey and Joyner, 2015; Ross et al., 2019). The mechanism underpinning muscle deoxygenation involves local reactions in skeletal muscle, such as contraction-induced elevations in capillary perfusion and an increased mitochondrial respiration (Bartke, 1987; Zoll et al., 2002; Clifford and Hellsten, 2004; Korthuis, 2011; Grassi and Quaresima, 2016).

The observed higher slopes and minima/maxima of muscle oxygen saturation during and after exhaustive ramp exercise are in line with previous observations on the enhanced and accelerated capacity for muscle deoxygenation of aerobically fit individuals during exhaustive pedaling exercise and subsequent
reoxygenation with the offset of pedaling (Figure 2) (Ding et al., 2001; Brizendine et al., 2013; Casey and Joyner, 2015; Perrey and Ferrari, 2018; Ross et al., 2019). These observations support the view that NIRS-based measures of oxygen saturation are a proxy of mitochondrial activity (Pilegaard et al., 2002; Ryan et al., 2014). The resulting difference in the proxy for the local respiratory capacity between fit and unfit subjects indicated that this study would allow to identify whether the hypothesized association between the ACE-I/D genotype and kinetic aspects of oxygen saturation with exhaustive muscle work would depend on the aerobic fitness state.

Based on previous observations on ACE-I/D associated differences on angiotensin-modulated vasoconstriction (Korthuis, 2011; van Ginkel et al., 2015, 2016; Valdivieso et al., 2017), variability in exercise-induced changes in muscle oxygen saturation during the exhaustive type of pedaling exercise were expected to be associated, with the ACE-I/D genotype and the aerobic fitness status. We observed a 1.5-fold increased
FIGURE 4 | Muscle deoxygenation during exercise is associated with the interaction between the aerobic fitness state and the angiotensin-converting enzyme-insertion/deletion (ACE-I/D) genotype. Box Whisker plots for the Δ_deoxgenation (A) and the slope of deoxygenation (B) in VAS and GAS combined in the function of the aerobic fitness status and the ACE-I/D genotype. Lines connect conditions demonstrating significant differences at *p < 0.05 and **p < 0.01. ANOVA for the factor “aerobic fitness” with post-hoc test of least significant difference.

Muscle deoxygenation in the aerobically fit than unfit subjects (Figure 3). Accordingly, the identified slopes of deoxygenation during exhaustive ramp exercise emphasized that this difference was due to an accelerated rate of deoxygenation for both investigated leg muscles in the aerobically fit compared to the unfit subjects, i.e., VAS: −3.5 vs. 2.9% per minute; GAS: −4.1 vs. −3.3% per minute). The slope 1/2 reoxygenation was overall 1.5-fold enhanced in aerobically fit compared to unfit subjects relating to the reportedly 2-fold faster recovery rates of oxygen consumption in musculus vastus lateralis of endurance athletes than inactive controls (Brizendine et al., 2013).

We explain the former observations in terms of the model that the precipitous drop of muscle oxygen saturation with contraction is due to enhanced oxygen consumption in mitochondria, which is not matched by the increase in muscle perfusion (Baker et al., 2010; Jones et al., 2017). Conversely, muscle reoxygenation postexercise arises because the respiratory activity of mitochondria levels off, while muscle perfusion continues to be elevated due to the fact that muscle capillaries remain maximally perfused for a longer time (Egginton and Hudlicka, 1999; Korzeniewski, 2003; Clifford and Hellsten, 2004). Anatomical factors that modify quantitative aspects of muscle oxygen saturation, such as the content of mitochondria and the capillarization, are an integral part of the underpinning processes. Accordingly, as muscle oxygen saturation is related to mitochondrial respiratory capacity (Ryan et al., 2014), the increased muscle deoxygenation and steeper negative slope of deoxygenation with pedaling exercise in the aerobically fit subjects would be explained by the functionally more developed capacity for mitochondrial respiration due to typical increases in mitochondrial volume density in trained skeletal muscles (Hoppeler et al., 1985). By contrast, the increased slope of reoxygenation for Musculus vastus lateralis during recovery from exhaustive exercise in the aerobically fit compared to the unfit individuals would be driven by the total capacity for capillary perfusion because the arterioles are in a fully dilated state in the recruited knee extensor muscles (Egginton and Hudlicka, 1999; Clifford and Hellsten, 2004). This suggestion would be supported by the observation that the estimated rate of reoxygenation was 9–15-fold accelerated in the studied leg muscles for both the fit and unfit subjects, compared to the corresponding rate of deoxygenation. Consequently, our results imply that capacitive differences in capillary perfusion exist between the aerobically fit and unfit participants of our investigation. Interestingly, the slope 1/2reoxygenation was 3-fold higher in VAS than GAS muscle (i.e., 69 vs. 21%), inditing higher capacities for reperfusion in the knee extensor than ankle extensor muscle.

The observed interaction effect between the aerobic fitness status × ACE-I/D genotype for the slope of deoxygenation and trend for such an effect on SmO2 min in the leg muscles during ramp-incremental exercise (Table 2, Figure 3) meets the expectation emanating from the aerobic fitness-associated influence of the ACE-I/D gene polymorphism on mitochondrial volume density (van Ginkel et al., 2015). In this article, we had reported that the volume density of mitochondria in musculus vastus lateralis is elevated in aerobically fit non-carriers compared to carriers of the ACE D-allele (i.e., endurance athletes) (van Ginkel et al., 2015). As well we had identified that the increase in the volume density of subsarcolemmal mitochondria in musculus vastus lateralis with cycling endurance training is amplified in non-carriers compared to carriers of the D-allele (Vaughan et al., 2013). As the content of mitochondria sets the capacity for muscle oxygen consumption, it appears reasonable to expect that the capacity for oxygen consumption, and, conversely, the
reduction in muscle oxygen saturation during exercise (Ryan et al., 2014), was largest in the knee extensor muscles of the aerobically fit subjects that did not carry the ACE D-allele.

By contrast, we did not find an aerobic fitness-associated interaction effect of the ACE-I/D-genotype on parameters of muscle reoxygenation in *Musculus vastus lateralis* during ramp-incremental exercise (Table 2). The ACE-I/D genotype has been found to be associated with altered capillary perfusion of distal limbs (i.e., the fingers) during exhaustive pedaling leg exercise and the capillarization of *Musculus vastus lateralis* (Valdivieso et al., 2017; Fluck et al., 2019) in not-specifically endurance-trained subjects. We observed however a trend (p = 0.10) for an interaction between ACE-I/D genotype and fitness state for Δ1/2 reoxygenation, which resolved for VAS in unfit subjects to a trend (p = 0.07) for a 10.7% higher Δ1/2reoxygenation in carriers compared to non-carriers of the D-allele (Supplementary Table 3). As well we observed a shallower slope of muscle deoxygenation during ramp-incremental exercise in unfit non-carriers compared to unfit D-allele carriers. This observation may in addition to differences in mitochondria respiration be reflective of the before mentioned better capillary perfusion of D-allele non-carriers during strenuous exercise, overriding of angiotensin 2-mediated vasoconstriction (van Ginkel et al., 2015). Concomitantly, our data relate to findings in ACE-DD genotypes on an altered balance between biochemical reactions that replenish and deplete intermediates of the TCA cycle, which releases energy through the oxidation of acetyl-CoA (Mathes et al., 2015). Nevertheless, an increased respiratory capacity of skeletal muscle is understood to contribute in an over-proportional manner to gains in maximal oxygen uptake with endurance training (Hoppeler et al., 1985) and add together with cardiovascular and pulmonary factors to the systemically measurable maximal oxygen uptake (di Prampero, 2003). Intriguingly, although the slope of muscle deoxygenation during exhaustive ramp exercise (i.e., −0.081 vs. −0.034% SmO₂ s⁻¹) and VO₂ peak (i.e., 54.4 vs. 40.2 mlO₂ min⁻¹ kg⁻¹) differed substantially between the aerobically fit and unfit ACE-II genotypes, the slope of muscle deoxygenation did not differ between the aerobically fit and unfit carriers of the ACE D-allele, despite a different VO₂ peak (i.e., 54.9 vs. 39.8 ml O₂ min⁻¹ kg⁻¹). Thus, the fitness × genotype-related effects on muscle deoxygenation were not reflected at the level of statistical significance for VO₂ peak. ACE-I/D genotypes have been found to demonstrate different hemodynamic responses during maximal exercise (Hellsten and Nyberg, 2015). Collectively our results, therefore, support the view that the association between ACE-I/D × fitness state for oxygen saturation in working skeletal muscle may not always manifest in statistically different maximal oxygen uptake due to a lower effect size at the system level and a possibly different contribution of knee extensor muscle between ACE-I/D genotypes to systemic oxygen uptake (Hoppeler et al., 1985; Jones et al., 2002; Fluck et al., 2010; Valdivieso et al., 2017; Williams et al., 2017).

The identified effects should be viewed in terms of the limitations of this study. First, NIRS-based measures only allow to compute values of oxygen concentration from a rather small volume of tissue which may bear the risk of being confounded by subcutaneous tissue material (Richardson et al., 2001; Grassi and Quaresima, 2016; Jones et al., 2017; Perrey and Ferrari, 2018). The recorded values, however, have been reported to be reliable, especially at the moderate exercise intensities used in this investigation, thus allowing to conduct real-time measurements in a non-invasive manner (Grassi and Quaresima, 2016; Crum et al., 2017; Jones et al., 2017; Perrey and Ferrari, 2018). Furthermore, within our investigation, we also identify considerable reductions for the computed levels of oxygen saturation that are in line with the reported larger degrees for muscle deoxygenation and reoxygenation between the aerobically fit and unfit subjects (Brizendine et al., 2013; Perrey and Ferrari, 2018). As well, for the purpose of data interpretation, it needs to be considered that oxygen saturation reflects the difference between oxygen supply and demand, thus providing only an indirect estimate of the possibly larger changes in the flux of oxygen with exercise (Collins et al., 2011; Rosenberry et al., 2019). This may especially come into account with the onset, and offset, of exercise when metabolic processes are not in a steady-state. Also, we note that in order to avoid other vasodilatation-related influences that may camouflage fitness state and genotype effects, we carried out the ramp-incremental exercise in the mere absence of a warm-up or muscle work to avoid the activation of mitochondrial respiration by unloaded contractions (Nioka et al., 1998; Boone et al., 2012; Perrey and Ferrari, 2018). Finally, we note that our observations are based on a rather heterogeneous population of volunteers, which were not matched for physical fitness between the ACE-I/D genotypes. Both the men and women were recruited to achieve the prospectively determined number of observations to reach the statistical significance of effects, although it had been shown that gender affects muscle deoxygenation during incremental ramp exercise (Murias et al., 2013).

**CONCLUSION**

Our measurements resolve that the ACE-I/D genotype affects aerobic fitness state-related differences in muscle oxygen saturation in recruited skeletal muscle during exhaustive pedaling exercise. Our findings corroborate the view that aerobic metabolism in exercised muscle importantly contributes to the variability of the systemically assessed values for aerobic capacity.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Materials.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethics Committee of
University of Zürich. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

DN, WF, and MF: study design. MF: funding. MVE, SR, AE, SC, and WF performed experiments. MVE, AF, WP, SC, and MF analyzed experiments. AF, WP, and MF analyzed the data. BG, DN, and MF interpreted the results. BG, MVE, and MF drafted the manuscript. BG, SR, MVE, DN, and MF revised the manuscript. All authors contributed to the article and approved the submitted version.

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REFERENCES

ACSM (2021). Guidelines for Exercise Testing and Prescription, 11th Edn. Philadelphia, PA: Wolters Kluwer Health.

Badtke, G. (1987). Sportmedizinsche Grundlagen der Körpererziehung und des sportlichen Trainings. Leipzig.

Baker, J. S., McCormick, M. C., and Robers, R. A. (2010). Interaction among skeletal muscle metabolic energy systems during intense exercise. J. Nutr. Metab. 2010, 905612. doi: 10.1155/2010/905612

Boone, J., Bouckaert, J., Barstow, T. J., and Bourgois, J. (2012). Influence of priming exercise on muscle deoxy[Hb + Mb] during ramp cycle exercise. Eur. J. Appl. Physiol. 112, 1143–1152. doi: 10.1007/s00421-011-0166-2

Brizendine, J. T., Ryan, T. E., Larson, R. D., and McCully, K. K. (2013). Skeletal muscle metabolism in endurance athletes with near-infrared spectroscopy. Med. Sci. Sports Exerc. 45, 869–875. doi: 10.1249/MSS.0b013e31827e0eb6

Buikema, H., Pinto, Y. M., Rooks, G., Grandjean, J. G., Schunkert, H., and van Gilst, W. H. (1996). The deletion polymorphism of the angiotensin-converting enzyme gene is related to phenotypic differences in human arteries. Eur. Heart J. 17, 787–794. doi: 10.1093/oxfordjournals.eurheartj.a014947

Casey, D. P., and Joyner, M. J. (2015). Regulation of increased blood flow (hyperemia) to muscles during exercise: a hierarchy of competing physiological needs. Physiol. Rev. 95, 549–601. doi: 10.1152/physrev.0035.2013

Chuang, M. L., Ting, H., Otsuka, T., Sun, X. G., Chiu, F. Y., Hansen, J. E., et al. (2002). Muscle deoxygenation as related to work rate. Med. Sci. Sports Exerc. 34, 1614–1623. doi: 10.1097/00005768-200210000-00013

Clifford, P. S., and Hellsten, Y. (2004). Vasodilatory mechanisms in contracting skeletal muscle. J. Appl. Physiol. 97, 393–403. doi: 10.1152/japplphysiol.00179.2004

Collins, J. A., Rudenski, A., Gibson, J., Howard, L., and O’Driscoll, R. (2011). Relating oxygen partial pressure, saturation and content: the haemoglobin–oxygen dissociation curve. Breathe 11, 194–201. doi: 10.1183/20734735.001415

Crum, E. M., O’Connor, W. J., Van Loo, L., Valeckx, M., and Stannard, S. R. (2017). Validity and reliability of the Moxy oxygen monitor during incremental cycling exercise. Eur. J. Sport Sci. 17, 1037–1043. doi: 10.1080/17461393.2017.1330899

Dimitriou, G., Papakonstantinou, D., Stavrou, E. F., Tsafas, S., Vervenioti, A., Athanasiadou, A., et al. (2010). Angiotensin-converting enzyme gene polymorphism and respiratory muscle function in infants. Pediatr. Pulmonol. 45, 1233–1239. doi: 10.1002/ppul.21316

Ding, H., Wang, G., Lei, W., Wang, R., Huang, L., Xia, Q., et al. (2001). Non-invasive quantitative assessment of oxidative metabolism in quadriceps muscles by near infrared spectroscopy. Br. J. Sports Med. 35, 441–444. doi: 10.1136/bmj.35.6.441

Egginton, S., and Hudlicka, O. (1999). Early changes in performance, blood flow and capillary fine structure in rat fast muscles induced by electrical stimulation. J. Physiol. 515(Pt 1), 265–275. doi: 10.1111/j.1469-7793.1999.265ax.x

Flitie, D. P., Franchi, M., Popp, W. L., Ruoss, S., Catuogno, S., Camenisch, K., et al. (2019). Concentric and eccentric pedaling-type interval exercise on a soft robot for stable coronary artery disease patients: toward a personalized protocol. J. Physiol. 515(Pt 1), 265–275. doi: 10.1111/j.1469-7793.1999.265ax.x

Fluck, M., Kramer, M., Fitze, D. P., Saper, S., Franchi, M. V., and Valdivieso, P. (2019). Cellular aspects of muscle specialization demonstrate genotype - phenotype interaction effects in athletes. Front. Physiol. 10, 526. doi: 10.3389/fphys.2019.00526

Fluck, M., Vaughan, D., and Westerblad, H. (2010). Linking genes with exercise: where is the cut-off. Eur. J. Appl. Physiol. 110, 1095–1098. doi: 10.1007/s00421-010-1662-9

Grassi, B., and Quaresima, V. (2016). Near-infrared spectroscopy and skeletal muscle oxidative function in vivo in health and disease: a review
from an exercise physiology perspective. J. Biomed. Opt. 21, 091313. doi: 10.1117/1.JBO.21.9.091313

Hellsten, Y., and Nyberg, M. (2015). Cardiovascular adaptations to exercise training. Compr. Physiol. 6, 1–32. doi: 10.1002/cphy.c140080

Hoppeler, H., Howald, H., Conley, K., Lindstedt, S. L., Claassen, H., Vock, P., et al. (1985). Endurance training in humans: aerobic capacity and structure of skeletal muscle. J. Appl. Physiol. 59, 320–327. doi: 10.1152/jappl.1985.59.2.320

Jackman, M. R., Ravussin, E., Rowe, M. J., Pratley, R., Milner, M. R., and Willis, W. T. (2008). Effect of a polymorphism in the ND1 mitochondrial gene on human skeletal muscle mitochondrial function. Obesity 16, 363–368. doi: 10.1038/oby.2007.40

Jones, A., Montgomery, H. E., and Woods, D. R. (2002). Human performance: a role for the ACE genotype? Exerc. Sport Sci. Rev. 30, 184–190. doi: 10.1097/00003677-200210000-00008

Jones, S., D’Silva, A., Bhua, A., Lloyd, G., Manisty, C., Moon, J. C., et al. (2017). Improved exercise-related skeletal muscle oxygen consumption following uptake of endurance training measured using near-infrared spectroscopy. Front. Physiol. 8, 1018. doi: 10.3389/fphys.2017.01018

Korthuis, R. (2011). Skeletal Muscle Circulation. San Rafael, CA: Morgan & Claypool Life Sciences.

Korzeniewski, B. (2003). Regulation of oxidative phosphorylation in different muscles and various experimental conditions. Biochem. J. 375, 799–804. doi: 10.1042/bj20030882

Lukin, L., and Ralston, H. J. (1962). Oxygen deficit and repayment in exercise. Int. Z. Angew. Physiol. 19, 183–191. doi: 10.1007/BF01037947

Mathe, S., van Ginkel, S., Vaughan, D., Valdivieso, D., and Flück, M. (2015). Gene–pharmacological effects on exercise-induced muscle gene expression in healthy men. Anatomy Physiol. Curr. Res. 55, 2161–0940. doi: 10.4172/2161-0940.55-005

Muiras, J. M., Keir, D. A., Spencer, M. D., and Paterson, D. H. (2013). Sex-related differences in muscle deoxygenation during ramp incremental exercise. Respir. Physiol. Neurobiol. 189, 530–536. doi: 10.1016/j.resp.2013.08.011

Nioka, S., Moser, D., Lech, G., Evangelistti, M., Verde, T., Chance, B., et al. (1998). Muscle deoxygenation in aerobic and anaerobic exercise. Adv. Exp. Med. Biol. 454, 63–70. doi: 10.1007/978-1-4615-4863-8_8

O’Donnell, C. J., Lindpaintner, K., Larson, M. G., Rao, V. S., Ordovas, J. M., Schaefer, E. J., et al. (1998). Evidence for association and genetic linkage of the angiotensin-converting enzyme locus with hypertension and blood pressure in men but not in women in the Framingham Heart Study. Circulation 97, 1766–1772. doi: 10.1161/01.CIR.97.18.1766

Perrey, S., and Ferrari, M. (2018). Muscle oximetry in sports science: a systematic review. Sports Med. 48, 597–616. doi: 10.1007/s40279-017-0820-1

Piegaard, H., Keller, C., Steensberg, A., Wulff Helge, J., Klarlund Pedersen, B., Saltin, B., et al. (2002). Influence of pre-exercise muscle glycogen content on exercise metabolism assessed by near-infrared diffuse correlation spectroscopy during incremental handgrip exercise. J. Appl. Physiol. 127, 698–706. doi: 10.1152/japplphysiol.00273.2019

Ross, R., Goodpaster, B. H., Koch, L. G., Sarzynski, M. A., Kohrt, W. M., Johannsen, N. M., et al. (2019). Precision exercise medicine: understanding exercise response variability. Br. J. Sports Med. 53, 1141–153. doi: 10.1136/bjsports-2018-100328

Ryan, T. E., Brophy, F., Lin, C. T., Hickner, R. C., and Neuffer, P. D. (2014). Assessment of in vivo skeletal muscle mitochondrial respiratory capacity in humans by near-infrared spectroscopy: a comparison with in situ measurements. J. Physiol. 592, 3231–3241. doi: 10.1113/jphysiol.2014.274456

Tonkonogi, M., and Sahlin, K. (2002). Physical exercise and mitochondrial function in human skeletal muscle. Exerc. Sport Sci. Rev. 30, 129–137. doi: 10.1097/00003677-200207000-00007

Valdivieso, P., Vaughan, D., Laczko, E., Brogioli, M., Waldron, S., Rittweger, J., et al. (2017). The metabolic response of skeletal muscle to endurance exercise is modified by the ACE I/D gene polymorphism and training state. Front. Physiol. 8, 993. doi: 10.3389/fphys.2017.00993

van Dijk, M. A., Kroon, I., Kamper, A. M., Boomsma, F., Danser, A. H., and Chang, P. C. (2000). The angiotensin-converting enzyme gene polymorphism and responses to angiotensins and bradykinin in the human forearm. J. Cardiovasc. Pharmacol. 35, 484–490. doi: 10.1006/jcph.2000.3344-20003000-00002

van Ginkel, S., de Haan, A., Woerdeman, J., Vanhees, L., Serne, E., de Koning, J., et al. (2015). Exercise intensity modulates capillary perfusion in correspondence with ACE I/D modulated serum angiotensin II levels. Appl. Transl. Genom. 4, 33–57. doi: 10.1016/j.atg.2015.03.002

van Ginkel, S., Ruoss, S., Valdivieso, P., Degens, H., Waldron, S., de Haan, A., et al. (2016). ACE inhibition modifies exercise-induced pro-angiogenic and mitochondrial gene transcript expression. Scand. J. Med. Sci. Sports 26, 1180–1187. doi: 10.1111/sms.12572

Vaughan, D., Huber-Abel, A. F., Graber, F., Hoppeler, H., and Fluck, M. (2013). The angiotensin converting enzyme insertion/deletion polymorphism alters the response of muscle energy supply lines to exercise. Eur. J. Appl. Physiol. 113, 1719–1729. doi: 10.1007/s00424-012-2583-6

Whipp, B. J., Davis, J. A., Torres, F., and Wasserman, K. (1981). A test to determine parameters of aerobic function during exercise. J. Appl. Physiol. Respir. Environ. Exercise Physiol. 50, 217–221. doi: 10.1152/jappl.1981.50.1.217

Williams, A. G., Rayson, M. P., Jubb, M., World, M., Woods, D. R., Hayward, M., et al. (2000). The ACE gene and muscle performance. Nature 403, 614. doi: 10.1038/3501141

Williams, C. J., Williams, M. G., Eynon, N., Ashton, K. J., Little, J. P., Wisloff, U., et al. (2017). Genes to predict VO2max trainability: a systematic review. BMC Genomics 18, 831. doi: 10.1186/s12864-017-4192-6

Wood, D. R., Pollard, A. J., Collier, D. J., Jamshidi, Y., Vassiliou, V., Hawe, E., et al. (2002). Insertion/deletion polymorphism of the angiotensin I-converting enzyme gene and arterial oxygen saturation at high altitude. Am. J. Respir. Crit. Care Med. 166, 362–366. doi: 10.1164/rccm.2103060

Zoll, J., Sanchez, H., N’Guessan, B., Ribera, F., Lampert, E., Bigard, X., et al. (2002). Physical activity changes the regulation of mitochondrial respiration in human skeletal muscle. J. Physiol. 543, 191–200. doi: 10.1113/jphysiol.2002.019661

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