Oxygen uptake and blood metabolic responses to a 400-m run
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Abstract This study aimed to investigate the oxygen uptake and metabolic responses during a 400-m run reproducing the pacing strategy used in competition. A portable gas analyser was used to measure the oxygen uptake ($\dot{V}O_2$) of ten specifically trained runners racing on an outdoor track. The tests included (1) an incremental test to determine maximal $\dot{V}O_2$ ($\dot{V}O_{2\text{max}}$) and the velocity associated with $\dot{V}O_{2\text{max}}$ ($\dot{V} - \dot{V}O_{2\text{max}}$), (2) a maximal 400-m (400T) and (3) a 300-m running test (300T) reproducing the exact pacing pattern of the 400T. Blood lactate, bicarbonate concentrations $[HCO_3^-]$, pH and arterial oxygen saturation were analysed at rest and 1, 4, 7, 10 min after the end of the 400 and 300T. The peak $\dot{V}O_2$ recorded during the 400T corresponded to 93.9 ± 3.9% of $\dot{V}O_{2\text{max}}$ and was reached at 24.4 ± 3.2 s (192 ± 22 m). A significant decrease in $\dot{V}O_2$ ($P < 0.05$) was observed in all subjects during the last 100 m, although the velocity did not decrease below $\dot{V} - \dot{V}O_{2\text{max}}$. The $\dot{V}O_2$ in the last 5 s was correlated with the pH ($r = 0.86, P < 0.0005$) and $[HCO_3^-]$ ($r = 0.70, P < 0.05$) measured at the end of 300T. Additionally, the velocity decrease observed in the last 100 m was inversely correlated with $[HCO_3^-]$ and pH at 300T ($r = -0.83, P < 0.001$, $r = -0.69, P < 0.05$, respectively). These track running data demonstrate that acidosis at 300 m was related to both the $\dot{V}O_2$ response and the velocity decrease during the final 100 m of a 400-m run.

Keywords Long sprint running · Buffer capacity · pH · Bicarbonate

Introduction

The pacing strategy required for optimal performance in a 400-m running race consists of a fast start followed by an approximate 15% decrease in velocity, relative to peak velocity, during the last 100 m of the race (Hanon and Gajer 2009). The adoption of such a fast-start procedure induces a higher peak oxygen uptake ($\dot{V}O_{2\text{max}}$) compared with other pacing strategies (Bishop et al. 2002; Sandals et al. 2006). While athletes are able to reach their maximal oxygen uptake during a simulated 800-m race (Thomas et al. 2005), previous results have indicated that the peak oxygen uptake reached during a 400-m running race or maximal exercise of equivalent duration (≈1 min) is between 80 and 90% of $\dot{V}O_{2\text{max}}$ (Duffield et al. 2005; Nummela and Rusko 1995; Spencer and Gastin 2001). However, further research is required as these studies did not exactly replicate the competition race conditions (e.g. no start from a starting-block). In addition to the high aerobic contribution, the anaerobic contribution during a 400-m race is also important and has been reported to range from 57 to 65% of the total energetic needs during a race (Duffield et al. 2005; Spencer
The importance of the anaerobic system is reinforced by the strong relationship \((r = 0.85, P < 0.01)\) between anaerobic glycolysis, as estimated by the maximal lactate levels (above 20 mmol l\(^{-1}\)), and the average velocity sustained during the 400-m race in highly trained athletes (Lacour et al. 1990). This large anaerobic contribution, and subsequent accumulation of metabolites (Kindermann et al. 1977), may contribute to the decrease in velocity observed during the final 100 m of a 400-m race (Hanon and Gajer 2009). However, while this decrease in velocity has been reported to occur concomitantly with a decrease in \(\dot{V}O_2\) during a 800-m race (Thomas et al. 2005), it is not known if this also occurs during a 400-m race.

Indeed, during 800- and 1,500-m running tests performed according to the pacing strategy of a competition, a decrease in \(\dot{V}O_2\) was observed in athletes during a 800-m race (Billat et al. 2009; Thomas et al. 2005) and during a 1,500-m race (Billat et al. 2009; Hanon et al. 2007). Furthermore, there was a significant positive relationship between the decrease in \(\dot{V}O_2\) and in velocity during the final stages of both the 800- (Thomas et al. 2005) and 1,500-m running races (Hanon et al. 2007) which suggests that a decrease in velocity may largely be responsible for the decrease in \(\dot{V}O_2\). It is important to note however, that the end velocity remained greater than the velocity associated with the \(\dot{V}O_2_{max}\) of each subject; thus, other mechanisms may also contribute. After maximal exercises, very low muscle and blood pH values (Nielsen 1999), an alteration in muscle buffer capacity (Bishop et al. 2009), and a large decrease in arterial blood bicarbonate concentration to values of 5 mmol l\(^{-1}\) (Kindermann et al. 1977) have been measured. While similar data have not been reported for a competitive 400-m race, it is likely that similar changes occur and that they may contribute to the large velocity decrease in the last 100 m of a 400-m race, and possibly also to the decrease in \(\dot{V}O_2\).

The purpose of this study therefore was to investigate the oxygen uptake response, and the subsequent metabolic and ionic perturbations in the blood, during a 400-m running test (400T) performed according to the optimal competition pace by highly trained 400-m specialists. Due to the pacing strategy (fast start notably) employed in a race (Han on and Gajer 2009), we hypothesized that athletes would be able to attain a high percentage of \(\dot{V}O_2_{max}\) and that there would be a \(\dot{V}O_2\) decrease during the last 100 m of the race when the velocity dramatically decreases. Because of the large ionic and metabolic perturbations previously observed at the end of supramaximal exercises (Hirvonen et al. 1992; Kindermann et al. 1977) and their established effects on O\(_2\) transport (Nielsen et al. 1999) and utilization (Jubrias et al. 2003), we also hypothesized that the end \(\dot{V}O_2\) response might be related to the acid–base status at the intermediate 300-m section of a 400-m running trial.

### Methods

#### Subjects

Ten 400-m runners (eight men and two women, age 21.8 ± 4.9 years, height 1.76 ± 0.06 m, and body mass 66.3 ± 6.6 kg) volunteered for the study. They trained three to five times per week for the 400-m race and were successful in regional and national running races (average performance of 50.9 ± 1.2 s for the men and 57.4 ± 3.7 s for the women). The study conformed to the recommendations of the Declaration of Helsinki, and participants gave voluntary written consent to participate in this experiment, which was approved by the local ethics committee.

#### Experimental protocol

Subjects performed three track-running tests during the summer competition period, on the same 400-m outdoor track, separated by at least 2 days. The first test was an incremental test aiming to determine \(\dot{V}O_2_{max}\). This was followed by a 400-m race performed according to the normal competition pacing strategy (400T) and a paced 300-m trial (300T). During this third trial, the subjects were asked to replicate the exact pacing strategy of the previously performed 400T.

For the three tests, oxygen uptake (\(\dot{V}O_2\)), minute ventilation (VE), volume of carbon dioxide released (V\(\dot{CO}_2\)), respiratory frequency (RF) and tidal volume (VT) were recorded continuously breath-by-breath with a portable gas exchange system (Cosmed K4b\(^2\), Roma, Italy). Calibration of both the Cosmed turbine (3-l syringe) and gas analysers was performed according to the manufacturer’s instructions before each test for each subject. Heart rate (HR) was measured and recorded continuously with a HR monitor (S810i and T61 electrode belt, Polar Electro, Kempele, Finland) for each athlete. Before and after the maximal sessions, arterialized capillary blood samples (85 \(\mu\)l) were taken from hyperemized ear-lobes in order to measure blood pH, arterial oxygen saturation (SaO\(_2\)) and bicarbonate concentration ([HCO\(_3\)]\(^{-}\)) with an i-STAT dry chemistry analyser (Abbott, Les Ulis, France). These measurements with the i-STAT portable analyser, anaerobically collected (cartridges closed hermetically), have been found to be reliable (ICC = 0.77–0.95 following maximal exercise) (Dascombe et al. 2007). In order to measure the blood lactate response to exercise, and because of the limited range of the i-STAT system (0.30–20 mmol l\(^{-1}\)), additional samples (20 \(\mu\)l) were taken for analysis using a Biosen Lactate analyser (Biosen C-line analyser, EKF Industrie, Elektronik GmbH, Barleben, Germany).
Incremental running tests

$V_{O2\text{max}}$ and $v - V_{O2\text{max}}$ (minimal speed at which the athlete was running when $V_{O2\text{max}}$ occurred) were determined using an incremental test conducted on an outdoor track marked every 25 m. The running pace was given by sounds emitted through a speaker controlled by a computer software programme to ensure precise control of speed by setting an audible cadence. The initial velocity was 10 km h$^{-1}$ which was then increased by 1 km every min until exhaustion. Blood samples were taken from the ear lobe before and 0 and 3 min after the end of the test.

Maximal 400-m running test (400T)

The first maximal exercise was a 400-m running test. The warm-up was standardized according to a regular pre-event 400-m warm-up (15 min of jogging, stretching, three successive sprints: 2 × 50- and 1 × 80-m realized at the 400-m start velocity) and was followed by a 7-min recovery period before the onset of the test. In order to reproduce the competition situation, each athlete used starting-blocks for the start of the race. Athletes were asked to select their own pace in order to realize the best performance and to run as in a competition. After the warm-up, the Cosmed K4b$^2$ base harness and the HR monitor were put on the participants and the K4b$^2$ analyser and battery were attached to the runner’s torso. The time from the beginning of Cosmed K4b$^2$ measurement to the start of the time-trial was recorded to locate the exact start in the data files.

Mechanical characteristics

For each test, athletes were videotaped by a researcher standing on the top of terracing, who used a panoramic video system (Panasonic Super-VHS, sampling rate of 50 Hz) from a fixed spot to determine the athlete’s velocity at each 25-m interval along the track.

Velocities

Velocity was calculated by dividing the distance by the time taken to cover this distance. This calculation was performed for each 25 m in order to provide accurate audio feedbacks on 300T and for each 50 m to calculate stride frequency (SF).

Stride length and frequency

Stride length (SL) and SF were determined from the video recording (Dart-Fish Team Pro 4.5, Fribourg, Switzerland) between 125 and 175 m of running for 300- and 400-m run and also between 325 and 375 m of running for the 400-m run; we avoided the last metres before the finish line where the stride characteristics could be modified. A fatigue index $[(\text{peak velocity} - \text{final velocity})/\text{peak velocity}] \times 100$ was calculated from the peak and the final velocity being measured over a 25-m distance. SF was calculated from the stride numbers within the 50-m measurement section and from the time between the strides as follows: (number of strides) × (time between the strides)$^{-1}$. SL was then calculated as follows: (average velocity) × SF$^{-1}$.

Maximal 300-m running test (300T)

The warm-up was conducted as in the previous test but with the 2 × 50- and 1 × 80-m sprints performed at the preliminary 400-m start velocity. Athletes were asked to reproduce the identical pacing strategy realized during the previous 400-m running test. To assist the subjects to reproduce an identical pacing regulation, interval times for every 25-m between 0 and 300 m were determined for each athlete. An audio signal-generator (laboratory self-made) was programmed to produce individualized signals at each 25-m interval. After the warm-up, the K4b$^2$ system and the heart rate monitor were put on the subjects as described for the 400-m running test.

Blood samples

Blood samples were taken from the ear lobe just before the start of both the 400 and 300T (that was 4 min after the end of the warm-up) and then at 1, 4, 7 and 10 min during the passive recovery following each test.

Data analysis

Breath-by-breath gas exchange values were averaged (Cosmed K4b$^2$ software) with a 5-s central moving average in order to obtain $V_{O2\text{max}}$ (incremental test) and $V_{O2\text{peak}}$ (400 and 300T). In order to compare with the literature, the highest incremental 30-s $V_{O2}$ value (30-s $V_{O2\text{max}}$) was also calculated. The end $V_{O2}$ ($V_{O2\text{end}}$) value was defined as the average of the last 5-s values. The $V_{O2}$/velocity, expressed in ml kg$^{-1}$ m$^{-1}$, was calculated for each 25 m and for each subject.

Statistical analysis

Data are reported as mean ± SD. The 400T response for gas-exchange variables, HR and running velocity was evaluated by a one-way analysis of variance with repeated-measures across each 25-m interval, followed by multiple comparisons (Student–Newman–Keuls) to test the effect of time on the variables. Relationships between variables (running velocity at different times of the race, stride characteristics, metabolic parameters, $V_{O2}$ kinetics and...
A significant decrease of 11.5 \% was observed when comparing 125 and 175 m after the start of the 400-m running test. The mean SL and SF between 125 and 175 m after the start of the 300T was equal to 0.13 m and 3.52 s, respectively. The maximal blood lactate value measured at the end of the test was 10.1 ± 2.7 mmol l⁻¹.

Performances and velocities during the 400 and 300T

The mean performance of the 400T was 52.2 ± 2.4 s for the men and 60.6 ± 4.5 s for the women, and corresponded to 97% of their best performance. The mean 400-m velocity was equal to 144.7 ± 9.8% of their v - VO₂ max. The peak start velocity (average of the individual distance at which peak start velocity was attained) was reached 67 ± 22 m after the onset of the race, and the velocities in the final 100 m were 8.37 ± 0.35 m s⁻¹ (i.e. 164.2% v - VO₂ max) and 6.93 ± 0.68 m s⁻¹ (i.e. 136.0 ± 10.5% v - VO₂ max) for the 300 and 400T. The intermediary time at 300 m (I-300) on the 400-m test averaged 39.6 ± 2.6 s and was not significantly different from the mean time of the 300-m running test (300T: 40.1 ± 2.1 s, i.e. a difference of 1.1%). During the 400-m running test, the mean velocity decrease in the last 100 m, expressed as a percentage of the velocity at 300 m, was 9.9 ± 5.5% (range from 2.2 to 20.4%) and the fatigue index was 22.5 ± 6.7%, (range from 11.8 to 34.1%).

Stride characteristics during the 400-m running test

The mean SL and SF between 125 and 175 m after the start of the race was 2.21 ± 0.11 m and 3.71 ± 0.12 Hz, respectively. At the end of the race (between 325 and 375 m after the onset of the 400T), these values were 1.95 ± 0.16 m and 3.52 ± 0.13 Hz which corresponded to a significant decrease of 11.5 ± 3.7 and 5.1 ± 3.8% for the SL and the SF, respectively during 400-m test (P < 0.01).

Cardio-respiratory parameters

Oxygen uptake response

The VO₂ peak reached during the 400T was equal to 54.2 ± 5.7 ml min⁻¹ kg⁻¹ and was not significantly different (P > 0.05) from the values reached during the 300T (53.6 ± 7.6 ml min⁻¹ kg⁻¹). Figure 1b displays the average VO₂ expressed relative to distance (each point being the average for each 25 m). During the 400T, the VO₂ peak value was detected at 192 ± 21 m (i.e. 24.4 ± 3.2 s after the onset of the test). This value corresponded to 93.9 ± 3.9% of VO₂ max. From 200 m until the end of the test, a significant decrease in VO₂ was observed (P < 0.0001). In the last 100 m, this significant decrease in VO₂ (P < 0.05) was equal to 8.5 ± 3.9 ml min⁻¹ kg⁻¹ (15.6 ± 6.5% of VO₂ peak) and was observed in all subjects.

VT, RF, VE and HR responses

The peak values for these variables were observed at 200 (VT), and 275 m (RF, VE) after the onset of the 400-m running test. These peak values were 1.8 ± 0.4 l min⁻¹, 79.5 ± 18.9 breaths min⁻¹ and 128.2 ± 16.2 l min⁻¹ for VT, RF and VE, respectively. A significant decrease in these ventilatory variables was noted in the last 100 m (P < 0.05). The HR reached 98.2 ± 1.7% of the maximal value 35 ± 6 s after the onset of the race and remained constant until the end of the race.

Blood metabolic variables

The metabolic results measured before and after the running tests are presented in Table 1. The metabolic values, measured post 400T, are significantly different from those measured post 300T except for the SaO₂ values which were measured at 1 min. The pre-test values were not significantly different between the tests.

Relationships between ventilatory, biomechanical and metabolic variables

Table 2 shows that the 400T end VO₂ value was positively correlated with both the 300T pH value (r = 0.85, P < 0.005) (Fig. 2a) and the 300T [HCO₃⁻] (r = 0.70, P < 0.05). In the same way, the magnitude of the 400T VO₂ drop was greater when 300T lactate concentration was high (r = 0.61, P < 0.05). On the other hand, the 400T VO₂ velocity and SL decreases were greater when the 300T pH and [HCO₃⁻] were low (P < 0.05 and 0.001, respectively) (Fig. 2b). The overall time-courses of VO₂ and velocity were correlated (r = 0.60, P < 0.01) and this correlation becomes higher (r = 0.92, P < 0.0001) when considering only the last 200 m. Finally, end VO₂ values were correlated with both VO₂ peak (r = 0.77, P < 0.01) and VO₂ max (r = 0.66, P < 0.05).
Discussion

The main finding of the present study was that trained athletes were able to reach a high percentage of VO2max in less than 25 s from the start of a 400-m race. Furthermore, a decrease in VO2 corresponding to 15% of VO2peak was observed in the last 100 m for all subjects. The end VO2 and the velocity decrease were positively correlated with both the blood pH and [HCO3−] measured after the 300T.

The present study indicated that during a 400-m race well-trained athletes are able to reach 94% of their VO2max in less than 25 s. This VO2peak value was higher than that observed in previous field studies where subjects reached 52 (Reis et al. 2004), 82 (Duffield et al. 2005) and 89% (Spencer and Gastin 2001) of VO2max during a 400-m race. To our knowledge, this is the first time that it has been reported that high relative values of VO2max can be attained so quickly during running exercise. As reviewed by Gastin (2001), it has previously been demonstrated that VO2 can be as high as 90% of the athlete’s maximum after 30–60 s (Gaesser and Poole 1996; Gastin and Lawson 1994; Granier et al. 1995; Poole and Richardson 1997). However, these previous studies all utilized intensive cycling exercise of short duration and with a maximal starting power (Wingate test or all-out exercise). Furthermore, manipulating pacing strategies, previous studies have indicated that VO2 during maximal exercise can be increased by employing an all-out start (Bishop et al. 2002; Gardner et al. 2003; Gastin et al. 1995), or a competition-start strategy (Hanot et al. 2008; Thomas et al. 2005). This may be attributable to the greater phosphocreatine (PCr) breakdown at the onset of exercise when using a very fast start procedure (Hirvonen et al. 1992). Similarly, the greater percentage of VO2peak values in the present study, compared to previous research may be related to intensive start performed by our trained subjects. A peak velocity as high as 8.37 m s−1 (i.e. 164.2% v VO2peak), explained by the use of starting-blocks, was reached between 50 and 75 m after the onset of the race.

Our higher relative VO2peak with respect to values observed in previous 400-m studies may also be due to the differences in the VO2 sampling window. For example, previous authors (Duffield et al. 2005; Spencer and Gastin 2001) aiming to determine the energy system contributions need not use as small a sampling window as that was used in the present study. As the velocity and therefore the O2 uptake are never steady during a 400-m running race, the use of a large sampling window will tend to smooth and decrease peak values. It can then be concluded that oxygen uptake can attain a level close to the VO2max values during brief and intense running exercise, provided that both the start velocity and the duration are sufficient.

In the present study, a VO2 decrease was observed during the final 100 m in all athletes, confirming previous results obtained during other maximal, exhaustive running exercises realized at a constant pace (Nummela and Rusko 1995; Perrey et al. 2002) or in field conditions (Hanot et al. 2007; Thomas et al. 2005). Although undiscussed by the respective authors, this phenomenon also appears in the figures of other studies incorporating paddling or cycling exercise (Bishop et al. 2002; Yamamoto and Kaneshia 1995; Zamparo et al. 1999; Williams et al. 2006). Nevertheless, in previous studies, no decrease has been observed at the end of 400-m running tests. Once more, this could be
explained by the sampling window used by the authors [200 m (Duffield et al. 2005) and 30 s (James et al. 2007)] which do not allow this observation during such intense and brief exercise. Nummela and Rusko (1995) who observed a $\dot{V}O_2$ decrease at the end of a maximal 50-s treadmill exercise used a 5-s sampling window.

In the present study, this decrease in $\dot{V}O_2$ appeared to be related to the peak [lactate]. The peak blood lactate concentration measured in the present study (22.0 ± 1.9 mmol l$^{-1}$), which was similar to values traditionally observed after a competitive 400-m race (Kindermann et al. 1977; Lacour et al. 1990), but was higher than all recent experimental data on 400-m races (Duffield et al. 2005; Reis et al. 2004) (≈ 13 to 16 mmol l$^{-1}$); these differences may have contributed to the absence/presence of this phenomenon. In the last 100 m, additional significant decreases in pH (7.18 to 7.00) and $[\text{HCO}_3^-] (10.7$ to $4.9$ mmol l$^{-1}$) were observed, and the pH and $[\text{HCO}_3^-]$ values measured at the onset of the $\dot{V}O_2$ decrease (300 m) were significantly related to the 400T end $\dot{V}O_2$ values. This final metabolic state aggravation, observed at the end of the 400T, is in accordance with the depletion revealed in blood bicarbonate after competitive rowing (Nielsen 1999), as well as the decrease in muscle buffering capacity observed after 45 s of exhausting exercise (Bishop et al. 2007), with the result being that the organism is unable to prevent an additional acidosis. Then, the decrease in $\dot{V}O_2$ and in velocity could be related to acidosis-induced inhibition of oxidative phosphorylation in contracting muscles (Jubrias et al. 2003) and consequently, with the large decrease in ATP observed at the end of a 400-m race (Hirvonen et al. 1992). It should be noted however, that the decrease in $\dot{V}O_2$ observed in the present study was not correlated to the pH values at 300-m ($r = -0.56, P = 0.10$). Furthermore, the correlations observed between end $\dot{V}O_2$ and both $\dot{V}O_2$peak and $\dot{V}O_2$max could indicate that athletes with a high $\dot{V}O_2$max (and therefore high end $\dot{V}O_2$) may have presented higher pH values at 300 m due to a greater use of aerobic energy resources during the 400 m test than athletes with a lower $\dot{V}O_2$max.

Blood metabolic perturbations, which could reflect changes in the muscle, could contribute to greater muscle fatigue during the last 100 m since the velocity strongly decreased (SL in particular) until values above the velocity associated with $\dot{V}O_2$max in each athlete. This velocity drop was negatively and significantly correlated to blood pH post 300T and to the $[\text{HCO}_3^-]$. The progressive reduction in running speed could then be the result of a combination of changes occurring in the muscle resulting in an incapacity to compensate for fatigue (Nummela et al. 1992). The significant correlation between the metabolic values at 300 m (blood pH and $[\text{HCO}_3^-]$) and the subsequent velocity decrease suggests that the ability of the athletes to finish strongly during this type of exhausting exercise may be linked to the level of acidosis when approaching the finish line.

Table 1 Mean ± SD values for blood parameters measured during the 400- and 300-m running sessions

<table>
<thead>
<tr>
<th></th>
<th>Pre-run</th>
<th>1</th>
<th>4</th>
<th>7</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>pH</strong></td>
<td>300</td>
<td>7.40 (0.04)$^a$</td>
<td>7.19 (0.05)$^{a,b}$</td>
<td>7.16 (0.05)$^{a,b}$</td>
<td>7.18 (0.05)$^a$</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>7.39 (0.04)</td>
<td>7.08 (0.04)$^b$</td>
<td>7.00 (0.07)$^b$</td>
<td>7.00 (0.04)</td>
</tr>
<tr>
<td><strong>[Lac]</strong></td>
<td>300</td>
<td>3.5 (1.2)</td>
<td>14.6 (1.5)$^{a,b}$</td>
<td>16.6 (1.3)$^{a,b}$</td>
<td>16.8 (1.5)$^a$</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>3.8 (0.8)</td>
<td>16.4 (0.9)$^b$</td>
<td>21.2 (2.1)$^b$</td>
<td>22.0 (1.9)</td>
</tr>
<tr>
<td><strong>[HCO$_3^-$]</strong></td>
<td>300</td>
<td>20.7 (2.3)</td>
<td>13.5 (1.9)$^b$</td>
<td>10.5 (1.7)$^{a,b}$</td>
<td>10.0 (2.2)$^a$</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>19.5 (1.4)</td>
<td>9.9 (1.2)$^b$</td>
<td>6.4 (1.1)$^b$</td>
<td>5.5 (1.4)$^b$</td>
</tr>
<tr>
<td><strong>SaO$_2$</strong></td>
<td>300</td>
<td>97.8 (0.4)</td>
<td>96.0 (1.6)$^b$</td>
<td>96.9 (1.2)</td>
<td>97.3 (0.5)</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>97.9 (0.3)</td>
<td>95.4 (1.0)$^b$</td>
<td>95.5 (1.9)</td>
<td>95.8 (1.0)</td>
</tr>
</tbody>
</table>

The blood parameters were collected 3 min before the test at the end of the warm-up (pre-race), and at 1, 4, 7 and 10 min of recovery following both tests. The blood variables are pH, lactate concentration [Lac] and bicarbonate concentration [HCO$_3^-$] in mmol l$^{-1}$, oxygen saturation (SaO$_2$) in %, partial pressure for O$_2$ (PO$_2$) and CO$_2$ (PCO$_2$) in mmHg. n = 10 except for [Lac] 400 where n = 8

$a$ For the same time analysis, 300 data is statistically different from the 400 data (300 pre-race is different from 400 pre-race,..., 300 post 10 is different from 400 post 10). $P < 0.05$

$b$ For the same parameter, statistically different from the previous result (post 4 is different from post 1,..., post 10 from post 7). $P < 0.05$

Table 2 Pearson correlation coefficients ($r$) between the end $\dot{V}O_2$ (last 5 s), the velocity decrease (last 100 m) measured during the 400T (400-m running test) and the metabolic parameters measured after the 300T (300-m running test)

<table>
<thead>
<tr>
<th></th>
<th>300, 400T</th>
<th>pH</th>
<th>[HCO$_3^-$]</th>
<th>SaO$_2$</th>
<th>[Lac]</th>
</tr>
</thead>
<tbody>
<tr>
<td>End $\dot{V}O_2$</td>
<td>0.85**</td>
<td>0.70*</td>
<td>0.45</td>
<td>0.54</td>
<td></td>
</tr>
<tr>
<td>VO$_2$ decrease (%)</td>
<td>-0.56</td>
<td>-0.57</td>
<td>-0.31</td>
<td>0.61*</td>
<td></td>
</tr>
<tr>
<td>Velocity decrease</td>
<td>-0.69*</td>
<td>-0.83**</td>
<td>-0.29</td>
<td>0.52</td>
<td></td>
</tr>
<tr>
<td>Stride length decrease</td>
<td>-0.85**</td>
<td>-0.68**</td>
<td>-0.50</td>
<td>0.37</td>
<td></td>
</tr>
</tbody>
</table>

Blood samples taken 4 min after the end of the 300 m test except for SaO$_2$ (1 min)

$* P < 0.05, ** P < 0.01$
Dempsey et al. (1982) have also demonstrated that physiological perturbations in the O₂ transport system, such as ventilatory regulation, could be induced by metabolic H⁺ accumulation. Indeed, during the 400-m race, our subjects presented particularly extreme peak values of RF (81 ± 22 breaths min⁻¹). These results could indicate that our subjects hyperventilated in order to partially compensate for their metabolic acidosis and to maintain an effective alveolar O₂ partial pressure (Miyachi and Katayama 1999) since VE continued to raise whereas \( \dot{V}O_2 \) peak had already been reached (Dempsey et al. 1982). Furthermore, a development of fatigue in respiratory muscles has been demonstrated during exhausting exercise (Gallagher et al. 1985), and a decrease in VT, which could indicate a reduction of inspiratory muscles strength, has been reported in the present study and in studies which observed a decrease in oxygen uptake (Billat et al. 2009; Hanon et al. 2007; Perrey et al. 2002; Thomas et al. 2005). During maximal whole-body exercise, the combination of a very low pH value and a poor vital capacity (unverified in this study, but demonstrated by Nielsen- et al. 1999) becomes critical for arterial oxygen pressure (PaO₂) and consequently for \( \dot{V}O_2 \) (Dempsey et al. 1982). In the present study however, no significant relationship was observed between the magnitude of the \( \dot{V}O_2 \) decrease and the recovery SaO₂ values. Nevertheless, a SaO₂ value of 95% measured 1 min after the end of the trial has been reported to correspond to a final SaO₂ value of less than 90% (Rasmussen et al. 1991). Therefore, the relation between end \( \dot{V}O_2 \) and SaO₂ needs to be more accurately tested.

The time-courses of \( \dot{V}O_2 \) and velocity have been shown to be highly correlated in the last 200 m and both the velocity decrease and the end \( \dot{V}O_2 \) observed during 400T have been shown to be related to the acidosis measured at 300 m. We can then, therefore, hypothesize that a common mechanism may explain the decrease in both \( \dot{V}O_2 \) and velocity. However, the question remains: does the \( \dot{V}O_2 \) decrease the consequence or the cause of the velocity decrease? It is also possible that both are the result of the metabolic perturbations and unrelated to each other.

A possible implication of this study for training is that improving the capacity of the buffer systems and the aerobic power may result in a smaller decrease in both pH and velocity at the end of the race. This advantage should be obtained without decreasing the capacity of the athlete to run very fast in the first 300 m. However, the absence of muscle data and the delay between the end of the exercise and SaO₂ measurement have to be considered as limitations in the interpretation of our results.

**Conclusion**

These track-running data demonstrate that a highly relative \( \dot{V}O_2 \) can be reached within 25 s from the start of a 400-m trial and that the end \( \dot{V}O_2 \) values may be the consequence of metabolic perturbations correlated to the acidosis. The accurate physiological hypotheses explaining the \( \dot{V}O_2 \) decrease during a maximal running exercise performed above the velocity associated to \( \dot{V}O_2 \) max need to be specifically investigated in order to demonstrate the central or (and) peripheral cause of this phenomenon.

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**References**


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**Fig. 2**

a Relationship between the 400-m end velocity (last 100 m expressed in % of the velocity at 300 m) and the [HCO₃⁻] values measured 4 min after the 300T. \( n = 10, R = -0.83, P < 0.001 \).

b Relationship between the 400-m end \( \dot{V}O_2 \) (last 5 s) and blood pH values measured 4 min after the 300T. \( n = 10, R = 0.86, P < 0.001 \).

The broken lines indicate the 95% confidence intervals.


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