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HAL Id: hal-01835273
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Submitted on 11 Jul 2018

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Physiological demands of a simulated BMX competition

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Abstract

The aim of the present study was to investigate the physiological demands of Supercross BMX during a simulated competition in elite athletes. During a preliminary session athletes underwent an incremental cycling test to determine maximal oxygen uptake (VO₂max) and power at ventilatory thresholds. In a second phase, athletes performed alone a simulated competition, consisting of six races separated by 30 minutes of passive recovery on an actual BMX track. During this simulated competition, oxygen uptake, blood lactate, anion gap and base excess (BE) were measured. Results indicated that a simulated BMX competition induces in elite athletes a high solicitation of both aerobic (mean peak VO₂ (VO₂peak): 94.3 ± 1.2 % VO₂max) and anaerobic glycolysis (mean blood lactate: 14.5 ± 4.5 mmol.L⁻¹) during every race. Furthermore, the repetition of six races separated by 30 min of recovery led to a significant impairment of the acid-base balance from the third to the sixth race (mean decrease in BE: -18.8 ± 7.5 %, p<0.05). A significant relationship was found between the decrease in BE and VO₂peak (r = -0.73, p<0.05). These findings confirm the importance of anaerobic conditioning in BMX athletes, and highlight the importance of the oxidative pathway in this sport where repeated high-intensity efforts are required.

Keywords. Bicycle motocross, elite athletes, intermittent sprints, buffering capacity, aerobic demand, anaerobic glycolysis
**Introduction**

Supercross BMX consists in the repetition of races on an irregular ground with turns and jumps. The length of the course is of approximately 350 m, and each series lasts around 40 s. During international events, a BMX cyclist that reaches the final has to perform 6 races with a recovery period of about 30 min between each race.

Few studies have analyzed the constraints of elite BMX and have focused mainly on biomechanical characteristics [2, 24, 34]. It is well established that during exercises lasting 10 to 40 s all the metabolic pathways are involved in a non-sequential way, including at the same time the degradation of phosphagenes, the degradation of the carbohydrates, and also the aerobic metabolism [10, 29]. However, the power output developed during a BMX race is not constant and is characterized by peaks of maximum power and of short duration developed in particular during the initial phase of the race [2, 24]. For example, Matteo et al. [33] have observed that during a BMX race simulation, acyclic efforts (i.e. non-pedaling efforts) accounted for 86.3% and cyclic efforts (i.e. pedaling efforts) accounted for the remaining 16.7% of the overall performance, with differences relating to the difficulty of the track. Thus, BMX performance is characterized by the capacity of the cyclist to produce a high level of muscular power at the beginning of the race, while displaying excellent technical skills with quasi isometric phases of work and some phases of exercise for which the muscular power of the lower limb must be again maximal. Other characteristics of Supercross BMX greatly distinguish this sport from sprint exercise models; a BMX cyclist is required to perform six races throughout the competition, and the relative contributions of the varied metabolic pathways in this specific situation has yet to be examined.

In this context, the purpose of our study was to investigate the physiological demands (i.e. aerobic and anaerobic energetic contribution) of BMX cycling during a simulated competition on an actual outdoor BMX track in elite athletes, in order to better identify the nature of recovery strategies needed during competition, as well as training programs for these athletes.
Materials and methods

Subjects

Ten elite BMX cyclists (6 men and 4 women) who regularly compete at international level volunteered for this study. The characteristics of the athletes are shown in table 1. The experiment took place in March just before the first international events in April. All subjects were informed about the study protocol, the risks of tests and investigations, and their rights according to the Declaration of Helsinki. The study was approved by the local Ethics Committee before its initiation, and was performed in accordance with the ethical standards of the International Journal of Sports Medicine [21]

Experimental procedure

The study was divided into two phases; one preliminary testing session and, one week later, a simulated BMX competition on the usual BMX training track where the athletes trained.

Preliminary session

Subjects underwent an incremental cycling test at a self-selected cadence on an electromagnetically-braked ergocycle (SRM, Schoberer Rad Messtecnik, Jülich, Welldorf, Germany). The test was performed in similar standard environmental conditions for all cyclists. The test began with a warm-up lasting 6 min at 100 W, after which the power output was increased by 30 W every minute until volitional exhausted. During this incremental cycling exercise, oxygen uptake (VO₂), minute ventilation (VE), and respiratory exchange ratio (RER) were continuously measured with a breath by breath gas exchange analyzer (Cosmed K4b2, Roma, Italy). Gas and flow analyzers were calibrated prior to each test using gases of known concentration and a 3 L syringe (Hans Rudolph, Kansas City, MO). The criteria used for the determination of VO₂max were a plateau in VO₂ despite an increase in power output, a RER above 1.1, and a heart rate (HR) above 90% of the predicted maximal HR (Howley et al. 1995). VO₂max was determined from the four highest VO₂ values recorded when VO₂ reached a plateau, whilst the peak power output (PPO) was determined as the mean cycling power
output recorded over the 1min period equating with VO2max. The first and the second ventilatory threshold (VT1 and VT2) were determined according to the method described by Wasserman et al. [31].

Simulated BMX competition

The cycling protocol was designed to reproduce as most as possible the physiological demand of a standard BMX international competition. The race course was a short outdoor track similar than that classically retrieved by participants in competition. Each race started from a 6 m high ramp followed by a 342-m hilly course with jumps, bumps and tightly banked corners. All cyclists individually performed 6 consecutive races separated by 30 min during which they were instructed not use any recovery strategy. Physiological measurements were performed immediately after each cycling race. Since the exercise was conducted in a challenging context for athletes (i.e. one month before the first international events), every athlete was well motivated to perform maximally the competition simulation. Competition was realized with a mean outdoor temperature of 17 ± 0.7 °C and wind lower than 3 knots.

Physiological assessment during the competition

Expired gases and oxygen uptake

Expired gases and oxygen uptake were measured immediately after every race using the K4b2 gas exchange system (Cosmed, Italy). The mask was applied on the face during 1min immediately after the end of the race. The oxygen uptake recovery curve during the first 20 s was used to predict peak oxygen uptake (VO2peak) according to the method described by Carre et al. [13] and recently used to assess aerobic demand of sprint swimming [23].

Perceived exertion

Immediately after the expired gases measurement, perceived exertion was recorded using the 6-20 Borg scale ranging from very very light (6) to exhaustion (20) [14].

Blood sampling
Blood lactate concentration (mmol.L⁻¹) was measured before (baseline value) and 3 and 25 min after every race using the Lactate Pro analyzer (Arkray) validated by Tanner et al. [30]. The capillary blood samples (5 μl) were collected from the finger.

Analysis of electrolytes changes (anion gap) and base excess was assessed using a portable clinical blood analyzer (i-STAT™, Abbott, Princeton, USA) [15]. Prior to each testing session, the i-STAT analyser was calibrated according to manufacturer’s specifications by an electronic stimulation. Cartridges were stored prior to use as per manufacturers instructions (2–8°C), and were removed to cold approximately 5 min prior to use. Capillary samples were collected from each subject by fingerstick with a lancing device and balanced 100μL heparin capillary tubes at the end of the first, the third and the sixth race and during the recovery period at 5 min and 25 minutes. Prior to sampling, the puncture site was cleaned with alcohol and carefully dried by using surgical gauze. Each sample was immediately expelled from the capillary tubes into the sample wells of two CG4⁺ cartridges. Anion Gap was reported as the difference between measured cations sodium and potassium and anions chloride and bicarbonate. Base excess was calculated from pH, partial pressure of carbon dioxide (PCO₂), and hemoglobin concentration (Hb).

Statistical analysis

All data presented are means ± SD (tables and figures). Each dependant variable was compared between the different cycling periods using an ANOVA with repeated measures (time). Newman-Keuls post-hoc tests were applied to determine between-means differences if the analysis of variance revealed a significant main effect for periods. Relationships between dependent variables (VO₂ peak, blood lactate, Anion Gap or Base excess at different period of the competition) were analyzed by a Pearson’s correlation coefficient. In order to evaluate the magnitude of changes in recorded data for the different cycling periods, the coefficient of variation (CV) was also calculated. All statistical analyses were conducted using Statistica software version 7.0 (Statsoft, France). For all statistical analysis, a P < 0.05 value was accepted as the level of significance.

Results
Physiological characteristics of elite BMX cyclists recorded during the preliminary session are presented in Table 1.

Performance

In both experimental sessions, no significant effect of time was observed among the 6 races with values above 97% of best individual performance recorded during the 6 races and mean exercise duration of 32 ± 4 s (figure 1a). Furthermore, a good reproducibility of performance was observed across races with mean coefficient of variation (CV) values of 2.7 ± 0.2%.

Physiological assessment during the competition

Parameters measured for each series are presented in Table 2.

Oxygen uptake response

No significant effect of time was observed on VO_{2peak} values among the 6 series. Results indicated during a single BMX race lasting 32 ± 4 s, an important contribution of aerobic metabolism with VO_{2peak} values higher than 90% of VO_{2max} (mean: 94.3 ± 1.2% VO_{2max}) and for some subjects values reaching 100% VO_{2max} (figure 1b).

Perceived exertion

No effect of time was observed on perceived exertion among the series. The mean perceived exertion was 16.2 ± 0.9, indicating a hard to very hard effort.

Blood metabolic variables

A high contribution from the anaerobic glycolysis assessed from blood lactate values was also observed with mean values of 14.5 ± 4.5 mmol.L^{-1}. Furthermore, after 25 min of recovery mean values of blood lactate were 5.4 ± 1.2 mmol.L^{-1}. No significant effect of time was observed on lactate values as well after races than at the end of recovery with intra-individual coefficient of variation (CV) values of: 8.3 ± 4% (Figure 1c).
A significant main effect of time was observed for Anion Gap and BE (p<.05) with an increase in Anion Gap and decrease in Base Excess (BE) among races. Compared with resting baseline values, the increase becomes significant for Anion Gap after the third race whereas Base Excess values were significantly lower than baseline values for BE at the end of the first race and at the end of all the recovery periods. A further decrease in BE was observed at the end of the sixth race, when compared with the first one (Figure 2). A significant relationship (r = -0.73, p<0.05) was found between VO_{2peak} (expressed in % VO_{2max}) recorded during the last race and the decrease in BE from the first to the sixth series (delta BE).

**Discussion**

The main findings of the present study are that a simulated BMX competition in elite athletes elicits a high contribution from both aerobic and anaerobic glycolysis during every race despite the relatively short duration and non-constant power development during a race. Furthermore during a simulated BMX competition the repetition of six maximal sprints leads to a significant impairment of the acid-base balance from the third to the sixth race without any clear effect on performance. The use of backward extrapolation to predict peak oxygen uptake was the only one that let the cyclists entirely free of their movements and realize all the technical elements of a BMX race (start, turns…). Using the same method for estimating VO_{2peak}, Jalab et al. [23] recently studied the dynamics of the aerobic response during a 100-m freestyle swim in well trained swimmers. Their results showed that VO_{2} increased very quickly at the beginning of race to reach 80 % of the maximal aerobic power at 25 m, 94 % at 50 m and 100 % at 75 m. Furthermore, high values of VO_{2} have been reached after 30 s of intensive cycling with a maximal starting power [18] and previous pacing studies have indicated that VO_{2} during maximal exercise can be increased by employing an all-out start [6] or a competition-start strategy [20]. This effect could be related to the greater phosphocreatine (PCr) breakdown at the onset of exercise when using a very fast start procedure [22]. BMX effort is mainly acyclic [24], and power output is developed by the lower limb predominantly during the first 10s of the race. Thus, the relatively high percentage of VO_{2max} recorded at end-exercise suggests that the initial high metabolic demands have been carried over, presumably due to the energy required for technical work and
isometric work of the upper limb. Such finding should be taken into account to design optimal training
and recovery strategies. Furthermore, our result could also be related to the specificity of BMX
training which is composed mainly of repeated sprints. Indeed, several studies have shown that the
oxidative metabolism is critically taxed during repeated-sprint exercises, which influences
performance [4, 28]. Furthermore, Bailey et al., [1] have shown that short-term training program
involving repeated, all-out sprint training resulted in an enhanced fractional muscle O₂ extraction
(examined via near-infrared spectroscopy), faster VO₂ kinetics, and an increased tolerance to high-
intensity exercise. Overall these results indicate for the first time an important aerobic contribution to
the total energy provision in BMX, suggesting that VO₂max could be one determinant of performance.
However, a BMX race is also characterized by consecutive short-duration sprints probably performed
at supra-maximal intensity, and involving other energetic pathways.

Factors affecting repeated sprint ability (RSA) have been widely studied in the last decade but for
short-duration exercise (<10 seconds), interspersed with brief recoveries (<60 seconds) [3, 19]. In
these studies one of most often suggested factor of RSA is the accumulation of H⁺ (acidosis) that may
impair performance through numerous mechanisms including inhibition of glycolytic enzymes such as
phosphofructokinase, impairment of the contractile machinery through inhibition of H⁺ on CA²⁺
binding to troponin or reduction of CA²⁺ reuptake and release. This hypothesis was supported by first
studies demonstrating a correlation between RSA and both muscle buffer capacity and changes in
blood pH [7, 9]. However, there are still many questions regarding the relationship between muscle
performance and pH decrease or H⁺ accumulation [25-26]. For example based on the notion that
intracellular acidification is one of the major causes of fatigue several studies have tried to analyze the
effect of ingested bicarbonate on exercise performance (e.g., Carr et al. [12]) and numerous studies
were unable to validate the ergogenic effect of bicarbonate loading. Recently, in a laboratory-
simulated BMX exercise composed by three Wingate tests separated either by 30 min recovery [33] or
by 15 min recovery [32], the authors did not observe any effect of sodium bicarbonate ingestion on
performance and perceived exertion. Furthermore, in our study despite a significant increase in anion
gap after the third race and a significant decrease in base excess between the first to the last race, no
effect was observed on performance in our elite athletes (figure 1a). The lack of any effect of time on performance recorded in this study could indicate that, in elite BMX cyclists, 30 min of recovery is enough to preserve maximal performance capacity throughout the competition. In the present study, the decrease in BE could be mainly related to the high contribution from anaerobic glycolysis assessed by blood lactate concentration values. These values measured after each series in the present study (14.5 ± 4.5 mmol.l⁻¹), are similar to experimental data recorded on 400-m runs [16, 27] (13-16 mmol l⁻¹). The lack of any period effect on blood lactate despite the decrease in base excess is in agreement with previous results indicating that the decrease in base excess is often larger than the increase in blood lactate [11]. Furthermore in our study we have recorded a significant relationship (r = -0.73, p<0.05; figure 3) between peak VO₂ (expressed in % VO₂max) recorded during the last race and the decrease in base excess from the first to the sixth series (delta BE). Changes in VO₂ explained 54% of the variation in BE, which is in agreement with previous researches identifying a relationship between RSA and aerobic fitness [7, 17].These findings confirm the importance of aerobic conditioning in sports, where repeated high-intensity efforts are required, such as BMX [5, 8]

Conclusion

These data, collected during a simulated competition on an outdoor BMX track, demonstrate for the first time that elite cyclists reach a very high relative VO₂ during every race. Moreover, even in these world-class athletes, we observed a significant impairment in acid-base balance inversely related with aerobic metabolism solicitation. Thus, while specific training to maximize energy production from anaerobic sources should remain a significant part of the yearly plan, strategies to develop the oxidative pathway should certainly not be neglected in order to develop the full metabolic potential of BMX cyclists.

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<table>
<thead>
<tr>
<th>Variables</th>
<th>Males (n=6)</th>
<th>Female (n=4)</th>
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<tbody>
<tr>
<td>Age (years)</td>
<td>20.8 ± 2.2</td>
<td>19.0 ± 2.2</td>
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<tr>
<td>Height (cm)</td>
<td>177.3 ± 9.7</td>
<td>169.3 ± 6.9</td>
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<tr>
<td>Body mass (kg)</td>
<td>75.4 ± 3.3</td>
<td>61.2 ± 3.9</td>
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<tr>
<td>Body mass index (kg.m⁻²)</td>
<td>24.2 ± 1.6</td>
<td>21.4 ± 1.4</td>
</tr>
<tr>
<td>VO₂max (mL.min⁻¹.kg⁻¹)</td>
<td>54.7 ± 4.7</td>
<td>41.8 ± 1.4</td>
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<tr>
<td>Peak Power Output (W)</td>
<td>400.0 ± 57.7</td>
<td>280.0 ± 10.0</td>
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<tr>
<td>Power at VT1 (W)</td>
<td>212.5 ± 14.4</td>
<td>150.0 ± 34.2</td>
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<tr>
<td>Power at VT2 (W)</td>
<td>331.3 ± 55.4</td>
<td>200.0 ± 25.2</td>
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Values are means ± SD

VT1 and VT2, 1<sup>st</sup> and 2<sup>nd</sup> ventilatory threshold
Table 2 Mean ± SD values for parameters measured during the six sessions (S1 to S6) of the simulated BMX competition.

<table>
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<tr>
<th></th>
<th>rest</th>
<th>R1</th>
<th>R2</th>
<th>R3</th>
<th>R4</th>
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<tr>
<td>VO₂peak (%VO₂max)</td>
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<td></td>
<td>93.3 ± 2.5</td>
<td>94.2 ± 3.1</td>
<td>97.4 ± 1.5</td>
<td>94.3 ± 2.8</td>
<td>93.1 ± 3.2</td>
<td>94.5 ± 3.1</td>
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<td>Blood Lactate (mmol.L⁻¹)</td>
<td>1.1 ± 0.2</td>
<td>14 ± 1.2*</td>
<td>15.8 ± 1.4*</td>
<td>15.3 ± 0.8*</td>
<td>14.6 ± 0.7*</td>
<td>14.2 ± 1.1*</td>
<td>13.9 ± 0.8*</td>
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<tr>
<td>RPE (6-10)</td>
<td>16.1 ± 1.1</td>
<td>16.2 ± 0.8</td>
<td>16.3 ± 1.2</td>
<td>16.4 ± 0.7</td>
<td>16.2 ± 1.2</td>
<td>16 ± 0.9</td>
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<td>Base excess (mmol.L⁻¹)</td>
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<td>-1.8 ± 0.3</td>
<td>-15.2 ± 3.2*</td>
<td>-6.8 ± 2.5*</td>
<td>-17.7 ± 3*</td>
<td>-8.1 ± 3.2*</td>
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<tr>
<td>Anion Gap (mmol.L⁻¹)</td>
<td>14 ± 4.5</td>
<td>16.7 ± 5.2</td>
<td>15.1 ± 4.8</td>
<td>22.7 ± 4*</td>
<td>15.8 ± 3.2</td>
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* Significantly different from rest, $ significantly different from the first race (p<.05)
Figure 1: Performance (a), oxygen uptake (b), and blood lactate concentration (c) recorded at the end of the 6 sessions (S1-S6) of the simulated BMX competition. Values are means ± SD.
Figure 2: Changes in base excess (a) and anion gap (b) among the 6 sessions (S1-S6) of the simulated BMX competition. Values are means ± SD.

* Significantly different from rest, $ significantly different from the first session (S1) (p<.05)
Figure 3: Relationship between delta BE (Base Excess) calculated (in %) between the last and the first session and peakVO₂ (in % VO₂max) recorded during the last session of the simulated BMX competition.

![Graph showing the relationship between Delta BE in % and VO₂peak in % VO₂max](image)

\[ R^2 = 0.5398 \]