Relationship Between Blood Lactate and Oxidative Stress Biomarkers Following Acute Exercise

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Abstract: We measured blood lactate, protein carbonyls (PC), and malondialdehyde (MDA) before and immediately after exercise (graded exercise treadmill test [GXT], Wingate cycle sprint, barbell squat, and barbell bench press) in active men. Data were analyzed using a 4 (exercise mode) x 2 (time) ANOVA with Tukey post hoc testing, and pairwise correlations were made. An interaction was noted for lactate (p = 0.009), with values increasing for all modes, and higher post exercise for bench press compared to Wingate and squat (p<0.05). No interaction or exercise mode effect was noted for PC or MDA (p>0.05). However, a time effect was noted for PC (p = 0.02), with values increasing from pre to post exercise. A positive correlation was noted between PC and MDA (r = 0.30; p = 0.0004), however no significant correlations were noted between PC and lactate (r = 0.06; p = 0.51) or between MDA and lactate (r = 0.10; p = 0.22). Blood lactate measured during the immediate post exercise period was not significantly correlated to biomarkers of oxidative stress in active men.

INTRODUCTION

Circulating lactate has been shown to rise to upper levels of 8-20mmol·L⁻¹ with acute anaerobic exercise, as lactate production from accelerated glycolysis overwhelms lactate utilization and clearance [1]. This 4 to 10 fold increase from resting levels results during and immediately following sprinting exercise and intense resistance exercise [1], in addition to graded exercise such as stress testing [2]. Specifically, lactate rises under conditions of increased glycolytic rate, decreased intramuscular oxygen pressure, and decreased lactate oxidation by the liver and kidneys [3].

Acute exercise also increases biomarkers of oxidative stress. Generation of reactive oxygen and nitrogen species (RONS) via normal metabolic pathways can overwhelm endogenous antioxidants and promote oxidation of proteins, lipids, and nucleic acids [4]. The extent of oxidative damage is commonly measured via quantitative assessment of circulating oxidized molecules, such as protein carbonyls (PC) and lipid peroxidation products such as malondialdehyde (MDA) [5]. During intense exercise, excessive RONS production via increased oxygen consumption and cellular metabolism, tissue ischemia/reperfusion, catecholamine autoxidation, as well as activation of RONS generating enzymes (e.g., xanthine oxidase, NADPH oxidase) may contribute to the rise in both PC and MDA [6].

Previous studies have measured the rise in both lactate and biomarkers of oxidative stress following sprint or resistance exercise [7-11]. Groussard et al. [7] suggest that exercise-induced metabolic acidosis contributes to oxidative stress, citing the earlier findings of Siesjo et al. [12]. While it is clear that circulating levels of both lactate and oxidative stress biomarkers rise with acute exercise, it is unknown whether the relationship between lactate and RONS (and subsequent elevation in oxidative stress biomarkers) is causative or merely coincidental.

Regarding oxidative stress, Groussard et al. [13] have shown that lactate ion is actually a scavenger of free radicals (hydroxyl radical and superoxide anion) in vitro. Yet, the authors caution that formation of lactate in vivo cannot be dissociated from subsequent metabolic acidosis, which may have a pro-oxidant effect. In vitro models of lactic acidosis have demonstrated generation of free radicals and lipid peroxidation [12,14,15]. Yet one must use caution when applying these findings to the matter of human exercise, in particular when considering that the extent of acidosis may not be as extreme within an in vivo system [16-18], and that multiple antioxidants exist within the human blood to minimize the extent of oxidation [4].

Currently, there is controversy regarding the contribution of lactate to exercise induced metabolic acidosis. Robergs et al. propose that glycolysis yields lactate (La⁻), not lactic acid (HLa), and explain that the generation of lactate actually attenuates intracellular acidosis [19, 20]. Yet proponents of the traditional concept of lactic acidosis question the veracity of Roberg’s stoichiometric calculations. They explain, among other criticisms, that the calculations do not recognize the falling intramuscular pH of exercising muscle [21-23]. Regardless of the stoichiometric methods used for intramuscular acid/base accounting, 80-100% of lactate escapes the sarcolemma via facilitated transport or diffusion as lactic acid, effectively resulting in blood lactic acidosis [24].

To date, only Lovlin et al. [11] have specifically analyzed a correlation between lactate formation and oxidative stress. They measured oxidative stress via the
thiobarbituric acid reactive substances (TBARS) assay, as well as blood lactate in a small sample of six, healthy young men at rest, 40% VO\textsubscript{2max}, 70% VO\textsubscript{2max}, and 100% VO\textsubscript{2max} on a cycle ergometer. While lactate markedly increased with exercise, TBARS was significantly lower at 40% VO\textsubscript{2max} and higher at 100% VO\textsubscript{2max}. The authors explain that exercise at 40% VO\textsubscript{2max} enhances antioxidant capacity, yet maximal exercise significantly increases blood lactate and plasma MDA. Overall, Lovlin et al. [11] found a correlation between lactate and MDA ($r^2 = 0.51$, $p<0.001$).

The purpose of the current study was to extend the findings of Lovlin and colleagues [11], by using a larger sample size, and measuring blood lactate and oxidative stress biomarkers in response to acute anaerobic exercise in order to determine if correlations exist between these variables.

**MATERIALS AND METHODOLOGY**

**Subjects**

Data collected from 56 recreationally active men were used in this study. These data are one component of larger data sets associated with studies for which manuscripts have previously been published [8, 25, 26], inclusive of other variables. Subjects completed a detailed health history questionnaire and underwent a physical examination, including anthropometric testing, prior to enrollment. All subjects were healthy and recreationally active, regularly involved in resistance and aerobic exercise. Subjects did not use medications or nutritional supplements (e.g., antioxidants) that may have affected the dependent variables being measured. All experimental procedures were performed in accordance with the Helsinki Declaration and approved by the University Human Subjects Review Board. Subjects provided both verbal and written consent prior to participating. Subject characteristics are presented in Table 1.

**Table 1. Characteristics of 56 Recreationally Active Men**

<table>
<thead>
<tr>
<th>Variable</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yrs)</td>
<td>24 ± 5</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>176 ± 8</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>86 ± 16</td>
</tr>
<tr>
<td>Body fat (%)</td>
<td>14 ± 6</td>
</tr>
<tr>
<td>Resting Heart Rate (bpm)</td>
<td>68 ± 5</td>
</tr>
<tr>
<td>Resting SBP (mmHg)</td>
<td>120 ± 9</td>
</tr>
<tr>
<td>Resting DBP (mmHg)</td>
<td>77 ± 10</td>
</tr>
<tr>
<td>Hours per week exercise</td>
<td>4±3</td>
</tr>
</tbody>
</table>

Note: Values are mean ± SD.

**Exercise Protocols**

After completion of all screening procedures, a familiarization session for the relevant exercise mode was completed. Then, on a separate day, the following acute bouts were performed: 1) graded exercise treadmill test using the Bruce treadmill protocol (GXT; $n = 15$), Wingate cycle sprint ($n = 13$), barbell squat ($n = 13$), or barbell bench press ($n = 28$). Please note that the Wingate cycle sprint and the barbell squat were performed by the same subjects on two separate days. For all exercise modes, subjects reported to the laboratory in a fasted (4+ hour) state. They were instructed not to perform any strenuous physical tasks during the 48-hour period preceding the exercise bout. For the GXT, the Bruce treadmill protocol was performed on a motorized treadmill until exhaustion. The average duration of the GXT was 10 minutes. Further details are provided elsewhere [26]. For the Wingate cycle sprint test, subjects pedaled on a Monark friction braked cycle ergometer modified to conduct Wingate testing at a load equal to 7% of body weight. Prior to the 30-second test, subjects performed a warm-up of light cycling exercise interspersed with 5-second sprints. For the barbell squat, subjects performed 15 repetitions using a load equal to 70% of subjects’ pre-determined 1 repetition maximum (1RM) + body weight (system mass), using a Smith machine. Further details for the Wingate and squat test are provided elsewhere [8]. For all exercise modes, subjects were instructed and encouraged to exercise until exhaustion, and heart rate (HR) and perceived exertion (RPE) were measured before and immediately after the exercise bout.

**Blood Collection and Analysis**

Venous blood samples (~10mL) were obtained via needle and collection tube, before and within one minute following exercise. Samples were immediately analyzed for whole blood lactate as an indication of anaerobic metabolism and participant effort (Accutrend; Roche Diagnostics, Mannheim, Germany). The remainder of whole blood was separated immediately to plasma and stored in multiple aliquots at -80°C to be used for the measurement of protein carbonyls (PC) and malondialdehyde (MDA).

Protein carbonyls were analyzed using an enzyme linked immunosorbent assay (ELISA) (Zentech Technology, Dunedin, New Zealand) following the procedures as outlined by the manufacturer. Malondialdehyde was analyzed using a commercially available colorimetric assay (Northwest Life Sciences Specialties, Vancouver, WA), using the modified method described by Jentzsch and coworkers [27]. Assays were performed in duplicate on first thaw.

**Statistical Analysis**

Data were analyzed using a 4 (exercise mode) x 2 (time/intervention) analysis of variance (ANOVA). Tukey post hoc testing was done as needed. Multivariate, pairwise correlations were made between all biochemical variables, in addition to heart rate and RPE. All analyses were performed using JMP statistical software version 4.0 (SAS Institute, Cary, NC). Statistical significance was set at $p<0.05$. The data are presented as mean ± SEM, except for subject characteristics which are presented as mean ± SD.

**RESULTS**

An exercise mode by time interaction was noted for HR ($p = 0.002$), RPE ($p = 0.0002$), and blood lactate ($p = 0.009$), with values increasing for all variables from pre to post exercise for all modes. Heart rate was higher post exercise for the GXT compared to bench press ($p<0.05$), RPE was
higher post exercise for the GXT compared to bench press and squat (p<0.05), and lactate was higher post exercise for bench press compared to Wingate and squat (p<0.05). No interaction or exercise mode effect was noted for PC or MDA (p>0.05). A time effect was noted for PC (p = 0.02), with values increasing from pre to post exercise. However, no time effect was noted for MDA (p = 0.28). Data for all exercise modes, pre and post exercise are presented in Table 2.

A positive correlation was noted between HR and RPE (r = 0.97; p<0.0001), HR and lactate (r = 0.73; p<0.0001), HR and PC (r = 0.25; p = 0.003), RPE and lactate (r = 0.71; p<0.0001), RPE and PC (r = 0.25; p = 0.003), and PC and MDA (r = 0.30; p = 0.0004). However, no significant correlations were noted between PC and lactate (r = 0.06; p = 0.51) or between MDA and lactate (r = 0.10; p = 0.22). When analyzing correlations for each exercise mode independently, it was noted that no differences existed as compared to when data from the four exercise modes was pooled. Therefore, only pooled data are included here. A correlation matrix for all variables is presented in Table 3.

**DISCUSSION**

In the present analysis, differences in blood lactate and oxidative stress biomarkers were noted between the exercise modes (see Table 2); this may be attributed to differences in exercise intensity, mechanics, and duration. Of the exercise modes, the GXT produced the greatest increase in HR (196.60±6.79bpm) and RPE (18.85±0.65), as the GTX likely requires the largest whole body energy expenditure [28]. The bench press produced the lowest HR (162.21±4.80bpm) and RPE (14.93±0.46). This may be expected, because the bench press exercise involved pure eccentric muscle actions and had a relatively limited cardiovascular demand. However, the bench press exercise did produce the greatest increase in lactate from 1.52±0.35mmol·L⁻¹ to 8.71±0.34mmol·L⁻¹.

Additionally, the venous blood samples were obtained from the median cubital vein, which may have a higher concentration of lactate immediately following upper body exercise compared to lower body exercise. Unlike the other exercise modes involving larger muscle groups and/or whole body movement (squats, Wingate, GXT), the bench press alone failed to cause a statistically significant difference in pre and post PC. This may suggest that formation of PC is more closely associated with exercise that challenges larger muscle groups and/or the cardiovascular system.

The strong correlation between lactate, HR, and RPE across all exercise modes is expected. The Borg scale quantifying RPE has been validated as a measure of exertion for both lower extremity and upper extremity exercise [29]. Exertion correlates well with HR and blood lactate concentration [30].

Interestingly, there was no relationship noted between lactate and PC or lactate and MDA. This is contradictory to the work of Lovlin and colleagues who found an association between MDA and lactate at maximal exertion [11]. The current study used a more specific MDA analysis technique than that of Lovlin et al. [11], which utilized thiobarbituric acid (TBA), a reagent that reacts with other molecules not associated with oxidative stress and has been criticized for its lack of specificity [5]. Additionally, blood was drawn for the current study within one minute of exercise completion, whereas samples were collected at the midpoint of a 5 minute rest period between each stage by Lovlin et al. [11]. Additionally, Lovlin et al. [11] selected a range of exercise intensity from 40-100%VO₂max, which differed from the specific exercise performed in the present study. With such a range of intensity, Lovlin et al. [11] reported a much wider range and peak (up to approximately 19mmol·L⁻¹) for blood lactate values, as opposed to our values of only approximately 6-9nmol·L⁻¹ (Table 2). All of the above mentioned disparities may have contributed to the opposing findings between our work and that of Lovlin and colleagues [11].

The current study does not support the application of in vitro findings by Siesjo et al. [12], Bralet et al. [14], or Fauconneau et al. [15] to explain exercise-induced oxidative stress. These respective investigations involved in vitro models of acidosis at lower pH levels than that found with anaerobic exercise. The models of Siesjo, Bralet, and Fauconneau and colleagues involved pH levels of 6.0-6.5 [12], 5.0-6.0 [14] and 3.0 [15], respectively. In an investigation of maximal human exercise, Hermansen et al. [31] found that whole blood pH dropped from 7.42 to an average low level of 7.11 following a single bout of maximal exercise. The corresponding intramuscular pH fell from an average of 6.92 to 6.41±0.04. Hence, the extent of exercise-
induced whole blood or intramuscular acidosis may not be sufficiently intense to induce lipid peroxidation or protein oxidation, as represented by increased MDA and PC, respectively. This notion supports our finding of no correlation between blood lactate and oxidative stress response to acute exercise.

CONCLUSION

Evidence clearly supports an increase in blood lactate with acute exercise, with mixed findings for markers of oxidative stress [5]. While in vitro models of lactic acidosis have demonstrated the prooxidant capacity of lactate at high concentrations, there is not sufficient evidence to assume that either blood lactate or intramuscular lactate at exercise-induced concentrations is associated with oxidative stress, in particular in healthy, trained men. Physiological buffers, antioxidant capacity, and the hemodynamics of human exercise lessen application of the findings of Siesjo et al. [12], which imply a correlation between lactate and exercise-induced oxidative stress. However, it is possible that untrained subjects, who generally have lower antioxidant capacity, may also clear lactate at a slower rate and thus, present with a greater metabolic acidosis and subsequent susceptibility to oxidative damage.

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REFERENCES


