Effects of high-intensity training on muscle lactate transporters and postexercise recovery of muscle lactate and hydrogen ions in women

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Submitted 3 December 2007; accepted in final form 29 September 2008

Bishop D, Edge J, Thomas C, Mercier J. Effects of high-intensity training on muscle lactate transporters and postexercise recovery of muscle lactate and hydrogen ions in women. Am J Physiol Regul Integr Comp Physiol 295: R1991–R1998, 2008. First published October 1, 2008; doi:10.1152/ajpregu.00863.2007.—The purpose of this study was to investigate the effects of high-intensity interval training (3 days/wk for 5 wk), provoking large changes in muscle lactate and pH, on changes in intracellular buffer capacity (βmin in vitro), monocarboxylate transporters (MCTs), and the decrease in muscle lactate and hydrogen ions (H+) after exercise in women. Before and after training, biopsies of the vastus lateralis were obtained at rest and immediately after and 60 s after 45 s of exercise at 190% of maximal O2 uptake. Muscle samples were analyzed for ATP, phosphocreatine (PCr), lactate, and H+. MCT1 and MCT4 relative abundance and βmin in vitro were also determined in resting muscle only. Training provoked a large decrease in postexercise muscle pH (pH 6.81). After training, there was a significant decrease in βmin in vitro (−11%) and no significant change in relative abundance of MCT1 (96 ± 12%) or MCT4 (120 ± 21%). During the 60-s recovery after exercise, training was associated with no decrease in muscle lactate, a significantly smaller decrease in muscle H+, and increased PCr resynthesis. These results suggest that increases in βmin in vitro and MCT relative abundance are not linked to the degree of muscle lactate and H+ accumulation during training. Furthermore, training that is very intense may actually lead to decreases in βmin in vitro. The smaller postexercise decrease in muscle H+ after training is a further novel finding and suggests that training results in a decrease in H+ accumulation and an increase in PCr resynthesis which can actually reduce the decrease in muscle H+ during the recovery from supramaximal exercise.

buffer capacity; monocarboxylate transporter 1; monocarboxylate transporter 4; phosphocreatine resynthesis; females

REMOVAL OF HYDROGEN IONS (H+) during intense skeletal muscle contractions occurs via intracellular buffering and via a number of different transport systems (27). The intracellular buffer capacity (βmin in vitro) measures the contribution to physicochemical buffering by proteins, dipeptides (e.g., carnosine), and phosphates within the muscle but excludes “dynamic” metabolic buffering, such as the rephosphorylation of ADP by phosphocreatine (PCr). Sarcolemmal transporters also have an important role in the regulation of intracellular pH during high-intensity exercise (29). In particular, lactate-H+ cotransport during exercise is largely mediated by monocarboxylate transporters (MCTs), which are stereoselective for L-(-)-lactate and dependent on the pH gradient (29). Although 14 MCTs are known to exist, lactate-H+ cotransport in skeletal muscle is attributed to two isoforms (MCT1 and MCT4) (29, 38, 51). Although both βmin in vitro (13, 43) and MCT1 and MCT4 (6, 11, 30) have been reported to be altered in response to different types of exercise training, little is known about the stimulus for such muscle adaptations.

The local formation of H+ and/or lactate within the skeletal muscle during exercise has been proposed to be an important stimulus for adaptations of the muscle pH-regulating systems (13, 31, 50). There is some support for this hypothesis, as βmin in vitro has been reported to increase in response to high-intensity, but not moderate-intensity, training (14). In addition, Hashimoto et al. (23) recently reported that the addition of lactate anions to a tissue culture can increase the expression of MCT1 mRNA and protein in L6 cells. However, in vivo, greater accumulation of lactate and H+ during training has not always been shown to be associated with greater increases in MCT relative abundance (31, 35).

An important difference between in vivo and tissue culture studies is that, in vivo, lactate production occurs concurrently with the formation of H+ (25, 42). Furthermore, research suggests that too large an accumulation of H+ during training may have a detrimental effect on adaptations to the pH regulatory systems within the muscle (15, 47). Recent findings from our laboratory have shown that, despite being matched for training intensity and total work performed, interval training with short (1-min) rest periods (end-exercise pH ~6.81) resulted in a consistent decrease in βmin in vitro, whereas interval training with longer (3-min) rest periods (end-exercise pH ~6.90) resulted in a consistent increase in βmin in vitro (15). In addition, decreasing the H+ accumulation during interval training (via sodium bicarbonate ingestion) results in significantly greater improvements in MCT4 relative abundance in rats (47). Therefore, we hypothesized that, similar to our previous findings for βmin in vitro, high-intensity interval training interspersed with short rest periods, inducing a low muscle pH, may not increase MCT relative abundance.

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of the muscle pH-regulating systems (13, 31, 50), one might also expect that high-intensity training, inducing a low muscle pH, might lead to a greater decrease in muscle lactate and H⁺ after intense exercise. However, although many studies have measured changes in blood lactate after different forms of endurance training, we are aware of few studies that have reported changes in muscle lactate and H⁺ during the recovery from intense exercise (35, 43). Interpretation of these previous findings is compromised by the methodological approach taken, in that muscle biopsies were obtained after more work was performed (and greater changes in metabolites were produced) after training. Furthermore, these studies measured muscle metabolite changes after 3–5 min of recovery, and we are aware of no previous studies that have investigated more rapid (i.e., ≤1-min) changes in muscle metabolites during the recovery from intense exercise.

The aim of this study, therefore, was to investigate postexercise recovery of lactate and H⁺ and changes in βm in vitro and MCTs after high-intensity interval training (3 days/wk for 5 wk). As the accumulation of H⁺ has been shown to affect oxidative phosphorylation, enzyme activity, and ion regulation during some exercise tasks (18, 26, 44, 46), it is important to understand the stimuli responsible for training-induced changes in factors that regulate H⁺ accumulation during and after exercise. A secondary aim was to investigate changes in MCT relative abundance in women after training, as we are aware of only one study that has recruited women (17), and this study recruited elite, cross-country skiers. We hypothesized that high-intensity interval training, inducing a low muscle pH, would not increase βm in vitro, MCT relative abundance, or the decline in muscle lactate and H⁺ during the recovery from intense exercise.

**METHODS**

**Subjects**

Six active, female, sport science students [20 ± 2 yr of age, 63.9 ± 7.8 kg body mass, 43.2 ± 4.9 ml·kg⁻¹·min⁻¹ maximal O₂ uptake (V̇O₂max)] volunteered to participate in the study. Each subject was involved in an intermittent sport (hockey, netball, tennis, basketball, or football) at club level (1–2 training sessions per week plus 1 competition). Subjects were informed of the study requirements, benefits, and risks before giving written informed consent. Approval for the study procedures was granted by the Institutional Research Ethics Committee.

**Experimental Overview**

An overview of the experiment is shown in Fig. 1. All subjects completed a familiarization trial for all tests before the start of the study. During pre- and posttraining testing, each subject performed a graded exercise test (GXT) and a 45-s constant-intensity exercise test (190% V̇O₂max, CIT₄₅); in addition, a muscle biopsy from the vastus lateralis muscle was obtained at rest, immediately after the CIT₄₅, and 60 s after the CIT₄₅. After the pretesting period, subjects performed supervised, high-intensity interval training (6–12, 2-min intervals at 100% V̇O₂max, with 1 min of rest between intervals) 3 days/wk for 5 wk. Pre- and posttraining tests were conducted at the same time of day. Subjects were asked to maintain their normal diet and training throughout the study. They were also required to consume no food or beverages (other than water) 2 h before testing and were asked not to consume alcohol or perform vigorous exercise in the 24 h before testing. Each subject was asked to record food and fluid consumption 2 days before each test and to replicate this food and fluid consumption during posttraining testing.

**GXT**

A GXT was performed on an air-braked, track-cycle ergometer (Evolution, Adelaide, Australia) for determination of V̇O₂max and the lactate threshold. For the test, which consisted of graded exercise steps (4-min stages), we used an intermittent protocol (with 1-min break between stages). The GXT commenced at 50 W, and, thereafter, intensity was increased by 30 W every 4 min until volitional exhaustion (4).

**Gas Analysis During the GXT**

During the GXT, expired air was continuously analyzed for O₂ and CO₂ concentrations with use of Ametek gas analyzers (models SOV S-3A11 and COV CD-3A, Applied Electrochemistry, Pittsburgh, PA). Ventilation was recorded every 15 s using a turbine ventilometer (model 225A, Morgan, Kent, UK). The gas analyzers were calibrated immediately before and verified after each test with use of three certified gravimetric gas mixtures (BOC Gases, Chatswood, Australia); the ventilometer was calibrated before exercise and verified after exercise with use of a 1-liter syringe in accordance with the manufacturer’s instructions. The ventilometer and gas analyzers were connected to an IBM personal computer, which measured and displayed variables every 15 s. The sum of the two highest consecutive 15-s values was recorded as the subject’s V̇O₂max.

**CIT₄₅**

The high-intensity cycle test consisted of 45 s of continuous cycling at a set power output (395 ± 47 W) on an air-braked, front-access cycle ergometer (model Ex-10, Repco, Mulgrave, Victoria, Australia). Toe clips and heel straps were used to secure the feet to the pedals, and the test was performed in the seated position. Strong verbal encouragement was provided to each subject during the test. On the basis of pilot work, the power output for the CIT₄₅ was 190% of the mean power output at V̇O₂max (to enable a workload that was severe but could be completed by all subjects).

**Training Intervention**

Within 4–7 days of baseline testing, subjects began a high-intensity interval-training program (6–12, 2-min intervals at 100% V̇O₂max, with 1 min of rest between sets). The subjects performed three training sessions per week (Monday, Wednesday, Friday) for 5 consecutive weeks. All training sessions were completed on mechanically braked cycle ergometers (model 818E, Monark, Stockholm, Sweden) and were preceded by a 5-min warm-up at 50 W. Progression was controlled by increasing the workload (by altering the resistance) and the number of intervals performed in a training session. Training programs followed a periodized plan (session-to-session of altering training volume) so as to simulate athletic training programs, allow progression, and prevent overtraining. Pilot work performed on five subjects not involved in the present study (recruited from the same...
population) revealed that muscle H⁺ concentration ([H⁺]) 1 min after the sixth interval of a typical interval training session was 155 ± 15 mmol/l (pH 6.81).

Capillary Blood Sampling and Heart Rate

Glass capillary tubes were used to collect 50 μl of blood during the GXT (D957G-70-35, Clinitubes, Radiometer Copenhagen) and 100 μl of blood during the CIT45 (D957G-70-125, Clinitubes). A hyperemic ointment (Finalgon, Boehringer Ingelheim) was applied to the earlobe 5–7 min before initial blood sampling. Capillary blood samples were taken at rest and immediately after each 4-min stage of the GXT for the determination of the lactate threshold [using the modified Dmax method (4)]. Capillary blood samples were also taken at rest, immediately after the CIT45, and 3 and 5 min after the CIT45. Blood H⁺ and lactate concentrations were determined using a blood-gas analyzer (ABL 625, Radiometer Copenhagen). The blood-gas analyzer was regularly calibrated using precision standards and routinely assessed by external quality controls.

Muscle Sampling and Analysis

On the day of the CIT45, local anesthesia [5 ml, 1% lidocaine (Xylocaine)] was administered, and two incisions were made into the vastus lateralis of each subject. The first incision was used for the resting biopsy, and the second was closed with a Steri-Strip and subsequently used for the biopsies taken after exercise; manual suction was applied for all samples. The first muscle sample was taken (before warm-up) during supine rest. Samples (50 – 80 mg) were also suctioned for all samples. The first muscle sample was taken immediately after cessation of the CIT45, and 3 and 5 min after the CIT45. Blood H⁺ and lactate concentrations were determined using a blood-gas analyzer (ABL 625, Radiometer Copenhagen). The blood-gas analyzer was regularly calibrated using precision standards and routinely assessed by external quality controls.

Muscle Buffering Capacity: Titration Method ([βm viro] and [H⁺])

Freeze-dried, resting muscle samples (3–5 mg) were enzymatically assayed for ATP, PCr, and lactate. ATP, PCr, and lactate were extracted from muscle samples by the addition of 6% perchloric acid before centrifugation (10,000 g for 10 min). The supernatant was removed and neutralized by the addition of 2.4 mol/l KOH and 3 mol/l KCl. Samples were centrifuged again, and the supernatant was stored at −80°C. ATP, PCr, and lactate were measured using a previously described method (1).

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Sample Preparation for Western Blotting

Proteins were isolated from resting muscle samples for Western blotting by a standard method previously described by McCullagh et al. (34) and previously used in our laboratory (48). Muscle protein concentrations were determined in duplicate by bicinchoninic acid assay (Pierce, Interchim, Montluçon, France) with the use of BSA as a standard. There was no significant difference in the concentration of protein isolated from pre- and posttraining resting muscle samples (3.6 ± 0.9 vs. 3.4 ± 0.8 g/μl, P > 0.05).

Western Blotting of MCT1 and MCT4

MCT abundance was determined using previously described methods (34). Human antibodies specified for MCT1 and MCT4 detection resulted from specific antigens produced with synthetic peptides against the COOH-terminal region of MCT1 and MCT4, respectively, linked with their cysteine residue at KLH EDC-activated carrier protein. These antibodies were injected in solutions for immunization of New Zealand White rabbits. Polyclonal antibodies yielded a single band on a Western blot that corresponded to 43 kDa, consistent with the molecular mass reported previously (34). Human antibody specificities were confirmed in preliminary experiments in which the peptides blocked the detection of MCT1 and MCT4 (48). Membranes were washed as previously described, and MCT1 or MCT4 expression was detected by enhanced chemiluminescence (Biomax MR films, Kodak, Reuil-Malmaison, France). Films were developed and processed using a hyperprocessor (model RNP 1700, Amersham, Les Ulis, France). MCT1 and MCT4 protein band densities were determined by scanning the blots on a scanner (Duoscan T1200, AGFA, New York, NY) with Scion Image software (Scion, Frederick, MD). Results are expressed in arbitrary optical density units, as used by others (48).

Citrate Synthase Activity

Homogenates for citrate synthase (CS) activity were prepared in buffer consisting of (mM) 210 sucrose, 2 EGTA, 40 NaCl, 30 HEPES, 5 EDTA, and 2 PMSF (pH 7.4) and stored at −80°C. CS activity was assayed by a spectrophotometric method according to Srere (45). Changes in absorbance were recorded over 3 min at 412 nm and at 25°C.

Statistical Analysis

Values are means ± SD. One-way ANOVA, with repeated measures for time, was used to test for the effects of training on V̇O2max, the lactate threshold, MCT content, and muscle buffer capacity. Two-way ANOVA, with repeated measures for time, was used to test for the effects of training on muscle and blood metabolites. Least-squares linear regression analysis was used to calculate correlation coefficients between measured variables using Pearson’s product moment (r). Selected correlation coefficients were also reexamined by the use of semipartial correlations to adjust for the contribution of possible confounding variables. Sigma Stat software (Jandel Scientific, San Rafael, CA) was used for all statistical analyses, and the level of statistical significance was set at P < 0.05.

RESULTS

Aerobic Fitness

There were no significant changes in body mass following training (63.9 ± 7.8 vs. 64.0 ± 7.5 kg). There were significant improvements in V̇O2max (43.2 ± 4.9 vs. 47.4 ± 3.9 ml·kg⁻¹·min⁻¹, +9.7%, P < 0.05), power at V̇O2max (210 ± 20 vs. 231 ± 18 W, +10.0%, P < 0.05), and lactate threshold (140 ± 16 vs. 158 ± 19 W, +12.9%, P < 0.05). The increase in power at V̇O2max after training resulted in a decrease in the relative intensity of the CIT45 from 190% (before training) to 170% (after training) of the power at V̇O2max. There was no significant change in CS activity with training (35.1 ± 5.3 vs. 37.1 ± 7.1 μmol/min, P > 0.05).
**βm_{in vitro} and MCT Relative Abundance**

Individual changes in βm_{in vitro} (along with mean values) are shown in Fig. 2. Very high-intensity training was associated with a significant decrease (-11%) in βm_{in vitro} (from 145.7 ± 6.6 to 129.6 ± 7.7 mmol H⁺·kg dry wt⁻¹pH⁻¹, P < 0.05). After training, there was no significant change in relative abundance of MCT1 (96 ± 12%) or MCT4 (119 ± 21%), expressed relative to pretraining values. There was a significant correlation between CS activity and relative abundance of MCT1 (r = 0.75, P < 0.05), but not MCT4 (r = 0.28, P > 0.05).

**Muscle Metabolites**

At rest. Muscle metabolite data measured before and after the 45-s cycle test before and after training are summarized in Fig. 3. Resting muscle ATP, PCr, lactate, and H⁺ concentrations were not significantly different after training (P > 0.05).

After high-intensity exercise. There were significant changes in muscle ATP, PCr, lactate, and H⁺ from rest to after exercise before and after training. However, muscle lactate and H⁺ accumulation after exercise was lower after than before training (P < 0.05). A significant correlation was observed between muscle lactate and H⁺ accumulation after exercise when pre- and posttraining data were pooled (lactate = 0.90·[H⁺] + 81, r = 0.88, P < 0.05, n = 12). There was no significant change in postexercise ATP or PCr after training (P > 0.05) compared with before training. Muscle ATP and PCr measured 60 s after exercise were significantly higher after training than at the same time point before training, whereas muscle lactate and H⁺ accumulation was significantly lower.

After training, there was no significant change in the decrease in muscle lactate during the first 60 s of recovery (14.9 ± 6.4 vs. 14.8 ± 4.4 mmol/kg dry wt, P = 0.16). In contrast, there was a significantly smaller decrease in muscle H⁺ during the first 60 s of recovery following training (25.7 ± 5.2 vs. 17.8 ± 8.9 nmol/l, P < 0.05). This is equivalent to an increase in pH during the first 60 s of recovery of 0.09 ± 0.02 and 0.07 ± 0.02 pH unit before and after training, respectively. There was a significant correlation between the postexercise H⁺ and the decrease in H⁺ (r = 0.78, P < 0.05; Fig. 4). There was no significant correlation between the decrease in muscle

![Fig. 2](image-url)  
**Fig. 2.** Individual changes in titrated muscle buffer capacity (βm_{in vitro}) in resting skeletal muscle biopsies from untrained women (S1–S6) before and after 5 wk of very high-intensity interval training. Values are means ± SD. *P < 0.05 vs. pretraining mean.

![Fig. 3](image-url)  
**Fig. 3.** Muscle ATP (A), lactate (B), phosphocreatine (PCr, C), and H⁺ (D) at rest, immediately after exercise (after-ex), and after 60 s of passive recovery (+60 s) from CIT45 before and after 5 wk of training. Values are means ± SE (n = 6, except for +60 s lactate, where n = 5). *Significantly different from pretraining (P < 0.05). †Significantly different from after-ex (P < 0.05). ‡Significantly different from rest (P < 0.05).
H⁺ and βm\textsubscript{in vitro} (r = 0.48, P > 0.05). Furthermore, the relationship between the postexercise H⁺ and the decrease in muscle H⁺ remained significant (r = 0.70, P < 0.05), even when the influence of βm\textsubscript{in vitro} on the decrease in muscle H⁺ during recovery was separated out.

**Plasma Lactate Concentration and pH**

Plasma lactate and pH data measured before and after the 45-s cycle test before and after training are summarized in Fig. 5. After training, there was a significant decrease in plasma lactate concentration immediately after exercise (P = 0.04) and 3 min after exercise (P = 0.03), but not 5 min after exercise (P = 0.08); there was no significant change in plasma pH at any time point (P = 0.07–0.13).

**DISCUSSION**

**Main Findings**

This is the first study to investigate training-induced changes in βm\textsubscript{in vitro}, MCT1, MCT4, and muscle lactate/H⁺ accumulation after high-intensity exercise in untrained women. The major finding was that high-intensity interval training, interspersed with short recovery periods, was associated with a significant decrease in βm\textsubscript{in vitro} without a significant change in MCT1 or MCT4 relative abundance in resting muscle. After training, reductions in muscle lactate and H⁺ accumulation were observed in response to the same absolute workload. There was also a significantly smaller decrease in muscle H⁺, but not muscle lactate, in the first 60 s after exercise following training. Thus these findings allow a discussion about the importance of the local formation of H⁺ and/or lactate within the skeletal muscle during exercise as an important stimulus to induce adaptations in βm\textsubscript{in vitro}, MCT1, and MCT4 and changes in muscle metabolites during recovery from intense exercise. These results have implications for the design of exercise interventions to modify these characteristics.

**Effects of Training on βm\textsubscript{in vitro} and MCT Relative Abundance**

Consistent with our previous research (15), high-intensity interval training (6–12, 2-min intervals at ~100% VO\textsubscript{2max}), with short rest periods, resulted in a significant decrease in βm\textsubscript{in vitro}, which was observed in all subjects (Fig. 2). This result is likely to be due to a decrease in the intracellular buffers that are measured using the in vitro titration technique (i.e., phosphate, protein, and dipeptides within the muscle) (33). Our previous research (15) indicates that a decrease in muscle carnosine content is not the cause of the decrease in βm\textsubscript{in vitro}. Furthermore, although we did not measure the content of other phosphate-bound compounds or the total adenine nucleotide concentration, the absence of changes in resting ATP or PCr after training suggests that the decrease in βm\textsubscript{in vitro} was not due to changes in intracellular phosphate. Therefore, the decrease in βm\textsubscript{in vitro} may be related to a decrease in intracellular protein buffering as a result of the large acidic load during training (pH <6.8) (9, 10) and/or the cumulative effect of repeated, transient decreases in intracellular protein buffering during the high-intensity training sessions (3).

Although H⁺ can be buffered intracellularly, sarcolemmal lactate transporters also have an important role in the regulation of intracellular pH during high-intensity exercise (29). In the present study, there was no significant change in resting MCT1 or MCT4 relative abundance after training. The lack of change in MCT4 is consistent with the observation that MCT4 has typically been shown to be less sensitive to training than MCT1. Furthermore, significant changes in MCT4 in response to high-intensity training were not observed in two previous studies (31, 35). However, the absence of a significant change in MCT1 is in contrast to previous research that has reported significant increases after low-intensity (30), moderate-intensity (6, 11), and high-intensity (7, 31) training.

The most likely explanation for our contrasting results is our novel training protocol: this is the first study to use a training program consisting of relatively long, high-intensity intervals (120 s at 100% VO\textsubscript{2max}) interspersed with short (60-s) rest
Muscle Lactate and $H^+$

Despite no change in MCT relative abundance, we observed the typical reductions in muscle lactate and $H^+$ accumulation that have been observed in previous studies after submaximal (6, 31, 32) and supramaximal (22) exercise training. These changes have predominantly been attributed to reductions in lactate production and/or increases in lactate removal (37). The improvements in aerobic fitness (10–13% increase in $V_{\text{O}}_{2\text{max}}$ and lactate threshold) suggest a decrease in anaerobic metabolism related to an improved potential for aerobic metabolism (24). Similar improvements in aerobic fitness have previously been associated with an increased aerobic contribution to supramaximal exercise (12, 13, 49). In addition, attenuated muscle glycogenolysis has been reported after endurance training (21, 22). A decrease in anaerobic glycogenolysis (nonmitochondrial ATP turnover) will result in a decrease in $H^+$ and lactate production, as increased lactate accumulation coincides with cellular acidosis (39, 42), as verified by the results of the present study ($r = 0.88, P < 0.05$). Thus an increased aerobic contribution and a decreased anaerobic contribution are likely to have contributed to the decreased postexercise lactate and $H^+$ values after training in the present study.

After training, there was no significant change in the decrease in muscle lactate content during the first 60 s after exercise. This contrasts with the results of previous interval training studies that have reported greater posttraining decreases in muscle lactate during the first 3–5 min of recovery after exercise (35, 43). However, it is difficult to interpret these findings, as changes in muscle lactate were examined while the subjects performed more work (and produced more lactate) after training. It is well established that lactate-$H^+$ cotransport is driven by the lactate gradient across the sarcolemma (29). Furthermore, it appears that most of the greater muscle lactate decrease reported by Mohr et al. (35) can be explained by the increase in postexercise lactate after training (Fig. 6). Thus our results suggest that training that induces large changes in muscle lactate does not necessarily induce adaptations that lead to a greater decrease in muscle lactate after intense exercise.

Consistent with previous research, the decrease in muscle $H^+$ during recovery was greater than the decrease in muscle lactate ($H^+$-to-lactate ratio of 1.9 to 1.2 pre- to posttraining). These values are close to the $H^+$-to-lactate ratio of 1.6 obtained with the NMR technique and human muscle (36). The greater
postexercise decrease in H\(^+\) than in lactate can be explained by the release of H\(^+\) from muscle fibers, by not only the lactate-H\(^+\) transporter, but also by the Na/H\(^+\) exchanger and HCO\(_3\)-dependent systems (28). However, a novel finding of the present study was the significantly smaller postexercise decrease in muscle H\(^+\) after training (25.7 vs. 17.8 mmol/l, \(P < 0.05\)). Our results suggest that this is not related to the decrease in \(\beta_{\text{in vitro}}\) as we found no correlation between \(\beta_{\text{in vitro}}\) and the decrease in muscle H\(^+\). Instead, the smaller postexercise decrease in muscle H\(^+\) was correlated with the lower H\(^+\) accumulation during exercise (Fig. 4).

In addition, the greater rate of PCr resynthesis after training (and subsequent release of H\(^+\)) would have contributed to the smaller decrease in muscle H\(^+\) after training. As \(pK_a\) values for PCr and P\(_i\) are 4.5 and 6.8, respectively, it can be calculated that, at a muscle pH of 6.9, \(~40\%\) of the smaller decrease in muscle H\(^+\) in the present study can be attributed to the increased rate of PCr resynthesis (25). Thus our results indicate that training that results in a decrease in H\(^+\) accumulation and an increase in the rate of PCr resynthesis can actually lower the decrease in muscle H\(^+\) following supramaximal exercise.

**Perspectives and Significance**

The local formation of H\(^+\) and/or lactate within the skeletal muscle during exercise has been proposed to be an important stimulus for adaptations of the muscle pH-regulating systems (13, 31, 50). However, an important finding of the present study is that changes in \(\beta_{\text{in vitro}}\) and MCT relative abundance do not appear to be linked to the degree of muscle lactate and H\(^+\) accumulation during training. Further research is therefore required to understand the complex intramuscular signals that are responsible for training-induced changes in the pH-regulating systems of the muscle; this has important implications for the design of exercise interventions to modify these characteristics. The smaller decrease in muscle H\(^+\) during the recovery from the CIT45 is also a novel finding and suggests that training that results in a decrease in H\(^+\) accumulation and an increase in the rate of PCr resynthesis can actually reduce the decrease in muscle H\(^+\) following supramaximal exercise.

**ACKNOWLEDGMENTS**

We thank the subjects for their commitment to the challenging training program and for their maximal effort during all tests. We also thank Prof. Dominique Mornet and Gerald Hugon for the production of MCT1 and MCT4 antibodies.

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