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Effects of high-intensity training on MCT1, MCT4, and NBC expressions in rat skeletal muscles: influence of chronic metabolic alkalosis

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This study investigated the effects of high-intensity training, with or without induced metabolic alkalosis, on lactate transporter (MCT1 and MCT4) and sodium bicarbonate cotransporter (NBC) content in rat skeletal muscles. Male Wistar rats performed high-intensity training on a treadmill 5 times/wk for 5 wk, receiving either sodium bicarbonate (ALK-T) or a placebo (PLA-T) prior to each training session, and were compared with a group of control rats (CON). MCT1, MCT4, and NBC content was measured by Western blotting in soleus and extensor digitorum longus (EDL) skeletal muscles. Citrate synthase (CS) and phosphofructokinase (PFK) activities and muscle buffer capacity (3 ± m) were also evaluated. Following training, CS and PFK activities were significantly higher in the soleus only (P < 0.05), whereas 3 ± m was significantly higher in both soleus and EDL (P < 0.05). MCT1 (PLA-T: 2%; ALK-T: 23%) and NBC contents (PLA-T: 85%; ALK-T: 60%) increased significantly only in the soleus following training (P < 0.01). MCT4 content in the soleus was significantly greater in ALK-T (115%) but not PLA-T compared with CON. There was no significant change in protein content in the EDL. Finally, NBC content was related only to MCT1 content in soleus (r = 0.50, P < 0.01). In conclusion, these results suggest that MCT1, MCT4, and NBC undergo fiber-specific adaptive changes in response to high-intensity training and that induced alkalosis has a positive effect on training-induced changes in MCT4 content. The correlation between MCT1 and NBC expression suggests that lactate transport may be facilitated by NBC in oxidative skeletal muscle, which may in turn favor better muscle pH regulation.

monocarboxylate transporter; sodium bicarbonate transporter; lactate transporter; muscle pH regulation; exercise

DURING HIGH-INTENSITY EXERCISE, contracting skeletal muscles produce and accumulate lactate and hydrogen ions. Lactate is either removed by oxidative processes or released into the blood and removed by other cells [i.e., the lactate shuttle (11)]. Muscle pH dramatically decreases during high-intensity exercise [pH = 7.4 = > 6.86 (34)] and then, consequently, in the blood. Efficient pH regulation depends on different mechanisms, including membrane transport proteins (for review, see Ref. 21), sarcolemmal carbonic anhydrase (16), and also the intracellular muscle buffer capacity (33).

The main transport proteins that regulate muscle pH are the sodium/proton exchanger [NHE (19, 24)], the monocarboxylate transporters [MCT (6)], and the sodium bicarbonate cotransporter [NBC (26)]. NHE plays an important role in pH regulation at rest, whereas MCTs are involved mainly during high-intensity exercise (19, 24). Different isoforms of MCT are expressed in skeletal muscle (8), with the most commonly described being the MCT1 and MCT4 isoforms. MCT1 has been found predominantly in oxidative muscles, and only in small amounts in glycolytic muscles, whereas MCT4 has been reported to exist with less fiber-type specificity (14, 35). Finally, NBC has been identified in different cells, and only recently in human (25, 26) and rat (26) skeletal muscle. NBC and MCT1 seem to functionally cooperate with Becker et al. (3) reporting increased lactate transport activity when MCT1 is coexpressed together with NBC in xenopus oocytes. Facilitation of MCT1 transport activity appears due mainly to the dissipation of the proton gradient with the transport of bicarbonate by NBC.

Although many studies (21) have investigated changes in MCT relative abundance following various forms of training, there are limited data regarding the training response of NHE and no data regarding training-induced changes in NBC abundance. The expression of MCT1, but not MCT4, is responsive to endurance training (14, 34), whereas the expression of both isoforms is responsive to strength (23) and sprint (5) training. Thus, the type of training does not seem of importance for the regulation of MCT1 content in skeletal muscle, although it appears important for the regulation of MCT4 content (23). In contrast, both muscle buffer capacity (15) and NHE (24) have been reported to undergo only adaptive changes in response to high-intensity training. These results suggest that a reduction in internal pH could be an important stimulus for training-induced adaptations in the MCT1 and MCT4 protein expression and possibly also NBC expression.

The efflux of H+ from the muscle can be enhanced by an elevated extracellular buffer concentration (20). The ingestion of a buffering agent [i.e., sodium bicarbonate (NaHCO3) or sodium citrate] prior to exercise has subsequently been shown to reduce the accumulation of H+ in skeletal muscle, interstitium, and blood during repeated, intense muscle contractions (27, 38). Although previously used as an ergogenic aid, ingestion of a blood-buffering agent provides a novel approach to alter the H+ accumulation during training and to investigate
the role of H⁺ accumulation as a stimulus for changes in transport proteins.

The main purpose of this study, therefore, was to determine the effects of high-intensity exercise with or without the ingestion of NaHCO₃ prior to each training session on MCT1, MCT4, and NBC protein relative abundance in rats. Due to the possible ergogenic effects of NaHCO₃ ingestion, the experimental training groups were matched for work (supervised training program for 5 wk). We hypothesized that high-intensity exercise training would increase MCT1, MCT4, and NBC protein relative abundance in rat skeletal muscle. Furthermore, if intracellular pH is an important stimulus for changes in these proteins, then induced metabolic alkalosis, via the chronic ingestion of bicarbonate prior to each training session, should differently influence training-induced changes in their relative abundance. Finally, due to their functional cooperation (3) and their expression in skeletal muscle (26), we further posited that NBC and MCT1 relative abundance would be correlated.

METHODS

Animals

Experiments were conducted on 28 male Wistar rats (Charles River, St. Germain sur l’Arbresle, France) according to the guidelines of the National Research Council for the care and use of laboratory animals and French law on animal handling and protection. The protocol was approved by the local ethics committee. The rats were individually housed in a temperature-controlled room and maintained with food and drink ad libitum in a 12:12-light-dark cycle (lights on at 7 PM), thus allowing exercise during their active phase. All exercise tests were conducted in the daytime, but a dark cover was placed over the treadmill to ensure that the rats exercised in darkness. The body mass of the rats was monitored throughout the experimental period.

Experimental Design

All animals were familiarized with a motor-driven treadmill for 3–4 days, 5 min/day, on a 10% grade. The running speed was initially 15 m/min and was gradually increased to 25 m/min within 2 days. Following the familiarization, all animals performed an incremental test to exhaustion on a treadmill inclined to 15° [starting at 10 m/min with increments of 3 m/min every 2 min. (18)]. Exhaustion was defined as the inability of the rats to run on the treadmill despite small electric shocks and an inability to upright themselves when placed on their backs.

Rats were randomly assigned to one of the following three groups: control (CON, n = 10), high-intensity training supplemented with a placebo (PLA-T, n = 9), or high-intensity training supplemented with NaHCO₃ (ALK-T, n = 9). Forty-eight hours after the last exercise, all rats were killed by cervical dislocation. Portions of soleus and extensor digitorum longus (EDL) were quickly removed, frozen in liquid nitrogen, and stored at -80°C until biochemical analysis and Western blotting.

Training Intervention

As shown in Table 1, training consisted of six (1st week) to 12 (5th week) 2-min intervals (interspersed with 1 min of rest) performed 5 times/wk for 5 wk. The intensity of the intervals was initially set at 80% of the peak speed reached for each rat during the incremental test and was increased by 10% each week.

Supplementation

Thirty minutes prior to each training session, PLA-T received water, whereas ALK-T received NaHCO₃ (0.05 mg/kg body mass) administered by esophageal catheter. The timing was based on our pilot work, which showed that there was a significant increase in blood pH 15–30 min after the administration of NaHCO₃, which was maintained to a steady state for 45 min (Fig. 1). The dose was based on previous work in rats (36) that showed that this dose was able to prevent a significant drop in blood bicarbonate concentration following 60 min of exhaustive exercise. In addition, our further pilot work demonstrated that the decrease in muscle pH immediately following a typical training session (6 X 2-min intervals interspersed with 1 min of rest) was significantly less following the administration of NaHCO₃ than without (6.81 ± 0.04 vs. 6.96 ± 0.04 pH units, P < 0.05).

Skeletal Muscle Analysis

Citrate synthase and 6-phosphofructokinase activities. Homogenates for citrate synthase (CS) activity were prepared in buffer [nm: sucrose 210, EGTA 2, NaCl 40, HEPE 30, EDTA 5, and phenylmethylsulfonyl fluoride 2 (pH 7.4)] and stored at -80°C. CS activity was assayed by a spectrophotometric method according to Srere (37). Changes in absorbance were recorded for 3 min at 412 nm and at 25°C. Phosphofructokinase (PFK) activity was determined on fresh soleus and EDL homogenates according to the method of Opie and Newsholme (32) and followed the rate of decrease in NADH at 340 nm and at 25°C. Muscle protein concentrations were determined in duplicate by Bradford assay with BSA as standard (Bio-Rad, Ivy-sur-Seine, France).

Muscle buffer capacity. Resting muscle samples (45–50 mg) were homogenized on ice for 2 min in a solution containing sodium fluoride (10 mM) at a dilution of 25 µl of homogenizing solution per mg wet muscle (see Ref. 29). The muscle homogenate was then placed in a circulating water bath at 37°C for 5 min prior to and during the measurement of pH. The pH measurements were made with a microelectrode (MI-415; Microelectrodes, Bedford, NH) connected to a pH meter (SA 520; Orion Research, Cambridge, MA). After initial pH measurement, muscle homogenates were adjusted to a pH of 7.2 with a sodium hydroxide solution (0.1 M) and then titrated to a pH of 6.2 by the serial addition of 10 µl of hydrochloric acid (0.01 M). From the fitted titration trendline, the number of moles of H⁺ (per g muscle) required to change the pH from 7.1 to 6.5 was interpolated. This value was then normalized to the whole pH unit for final display as micromoles H⁺ per gram wet muscle per unit pH (µmol H⁺·g muscle wet wt·pH⁻¹) and determined as the rat’s 3 mH⁺/vitrō.

Transport Protein Expression

Sample preparation for Western blotting. Proteins were isolated from muscles for Western blotting by a method previously described by McCullagh et al. (30) and previously used in our laboratory (39). Muscle protein concentrations were determined in duplicate by bicinchoninic acid assay (Pierce, Interchim, Montluçon, France) with the use of BSA as a standard.

Western blotting of MCT1, MCT4, and NBC. MCT and NBC contents were determined using previously described methods.

<table>
<thead>
<tr>
<th>Training Weeks</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of intervals/training session</td>
<td>6</td>
<td>7</td>
<td>9</td>
<td>10</td>
<td>12</td>
</tr>
<tr>
<td>Mean speed/wk, m/min</td>
<td>33</td>
<td>35</td>
<td>39</td>
<td>44</td>
<td>48</td>
</tr>
<tr>
<td>Minimal speed at beginning of training session, m/min</td>
<td>48</td>
<td>28</td>
<td>31</td>
<td>31</td>
<td>37</td>
</tr>
<tr>
<td>Maximal speed reached during training session, m/min</td>
<td>48</td>
<td>40</td>
<td>43</td>
<td>49</td>
<td>52</td>
</tr>
<tr>
<td>Mean running distance/training session, m</td>
<td>435</td>
<td>489</td>
<td>731</td>
<td>962</td>
<td>1,202</td>
</tr>
</tbody>
</table>

Table 1. Summary of the training performed by both training groups during the experimental period
Affinity polyclonal antibodies directed against the carboxy terminus of rat MCT1 and MCT4 were produced with the synthetic peptide C-Ahx-PLQNSSGDAEESPV-OH (aa 478–494) for MCT1 and C-Ahx-LREVEFLKAEPKNG-OH (aa 440–455) for MCT4, like the sequences used by Butz et al. (12), McCullagh et al. (30), and Wilson et al. (41). Such peptides were linked with their cystein residues at KLH EDC-activated carrier protein according to a protocol recommended by the manufacturer (Pierce) and the resulting solutions injected as specific antigens for immunization of New Zealand white rabbits. Western blots were probed with these antibodies and Chemicon International (Temecula, CA) antibodies (rabbit anti-MCT1: AB3540P; rabbit anti-MCT4: AB3114P). Polyclonal antibodies yielded a single band on a Western blot that corresponded to 43 kDa, consistent with the molecular mass reported earlier (30). Antibody specificities were confirmed in preliminary experiments in which the peptides blocked the detection of MCT1 and MCT4 (39). To determine NBC expression, we voluntarily chose an NBC antibody against the NBC isoform, which seemed to be related to the kidney/pancreas/heart isoform NBC1 (150 kDa). Indeed, Kristensen et al. (26) have pointed out that this isoform was clearly present in different rat skeletal muscles, but without any correlation to fiber type. In addition, an NBC construct derived from the kidney corresponding to this isoform was also used by Becker and colleagues (3, 4) in their in vitro studies to demonstrate the functional cooperation between NBC and MCT1. The NBC content was quantified by isoform-nonspecific NBC antibodies as previously used by Juel et al. (25) and Kristensen et al. (26) in skeletal muscle (NBC antibody, AF04017; Chemicon International 3212). These NBC antibodies were produced to a sequence (rat kidney NBC amino acids 338–391) with a high degree of homology to most NBC isoforms that have been cloned. Preliminary experiments also confirmed the specificity, and the band was detected at 140–150 kDa.

MCT1, MCT4, and NBC expression were detected by ECL (Biomax MR films; Kodak, Reuil-Malmaison, France). Films were developed and processed using a Hyperprocessor, RNP 1700 (Amer sham, Les Ulis, France). MCT1 and MCT4 protein band densities were determined by scanning the blots on a scanner (AGFA Duo Scan T1200) and then analyzing with Scion Image software (Scion, Frederick, MD). Results were expressed in arbitrary optical density units as previously used (39).

**Statistical Analysis**

Descriptive statistics are expressed as means ± SE One-way ANOVA was performed to compare the effects of high-intensity training (comparison with CON group) and the effects of NaHCO3 administration (PLA-T vs. ALK-T). Individual relationships between variables were studied by means of linear regressions. Statview software was used for all tests, and the level of significance was set at \( P < 0.05 \).

**RESULTS**

**Body Mass**

The mean body mass of the three groups of animals before training was 235 ± 17 g. After 5 wk of treadmill training, the body masses of the PLA-T and ALK-T rats were 380 ± 8 and 367 ± 8 g, respectively, with no significant difference between groups. After 5 wk of caged living, the control rats weighed significantly more than both training groups (428 ± 7 g, \( P < 0.05 \)).

**Effects of Training on Changes to CS and PFK Enzyme Activities and Buffer Capacity**

**Muscle enzyme activities.** As shown in Table 2, compared with CON, the activity of CS in the trained groups was +16% higher for PLA-T (\( P = 0.18 \)) and +23% higher for ALK-T (\( P < 0.05 \)) in the soleus, with no significant change in the EDL. Furthermore, CS activity was significantly higher in ALK-T than in PLA-T in the soleus (\( P < 0.05 \)). There were also significant changes in PFK activity (\( P < 0.05 \)) in both trained groups (+48% for PLA-T and +45% for ALK-T) only in the soleus, without change in the EDL after training.

**Muscle buffer capacity.** As reported in Table 3, muscle buffer capacity significantly increased after both training conditions (placebo and alkalosis) in both the soleus (\( P < 0.01 \), +41% for PLA-T and +40% for ALK-T) and in the EDL (\( P < 0.05 \), +27% for PLA-T and +26% for ALK-T).

**Membrane Transport Proteins of Rat Skeletal Muscles**

**Skeletal muscle and membrane protein contents.** As reported in Table 4, total skeletal muscle and membrane protein contents did not increase after training in PLA-T and ALK-T compared with CON (\( P > 0.05 \)).

**MCT1 relative abundance.** After training, MCT1 relative abundance in the soleus was significantly greater in the PLA-T (+30%) and ALK-T (+23%) groups compared with CON (\( P < 0.05 \)), with no significant difference between the two training groups (Fig. 2A). There were no significant differences between the three groups for MCT1 relative abundance in the EDL (Fig. 2B).

Table 2. CS and PFK activity

<table>
<thead>
<tr>
<th></th>
<th>CON</th>
<th>PLA-T</th>
<th>ALK-T</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS activity, μmol/min</td>
<td>42.9±1.6</td>
<td>49.7±3.4</td>
<td>52.9±3.2†</td>
</tr>
<tr>
<td>EDL</td>
<td>34.1±3.2</td>
<td>32.4±3.3</td>
<td>38.6±3.5</td>
</tr>
<tr>
<td>PFK activity, μmol·min⁻¹·mg protein⁻¹</td>
<td>9.4±1.1</td>
<td>13.9±1.0*</td>
<td>13.6±1.5*</td>
</tr>
<tr>
<td>SOL</td>
<td>45.0±4.3</td>
<td>49.2±7.4</td>
<td>47.2±7.4</td>
</tr>
</tbody>
</table>

Mean values ± SE for citrate synthase and phosphofructokinase activities in soleus (SOL) and extensor digitorum longus (EDL) muscles in control rats (CON) and trained rats, following the ingestion of either NaHCO3 (ALK-T) or a placebo (PLA-T). CS, citrate synthase; PFK, phosphofructokinase. *\( P < 0.05 \), significantly different from CON; †\( P < 0.05 \), significantly different from PLA-T.
Table 3. Muscle buffer capacity

<table>
<thead>
<tr>
<th></th>
<th>CON</th>
<th>PLA-T</th>
<th>ALK-T</th>
</tr>
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<tbody>
<tr>
<td>Muscle buffer capacity, µmol·g⁻¹·pH⁻¹</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SOL</td>
<td>32.9±1.0</td>
<td>46.4±2.8*</td>
<td>46.1±2.0*</td>
</tr>
<tr>
<td>EDL</td>
<td>44.0±1.4</td>
<td>56.1±3.1*</td>
<td>55.5±3.9*</td>
</tr>
</tbody>
</table>

Mean values ± SE for muscle buffer capacity in SOL and EDL muscles in CON and trained rats following ALK-T or PLA-T. *P < 0.05, significantly different from CON.

*MCT4 relative abundance.* MCT4 relative abundance in the soleus was significantly greater in ALK-T (+115%, P < 0.01) than both PLA-T and CON (Fig. 2A). Although a trend was apparent, there was no significant difference between PLA-T (+35%) and CON (P = 0.30). There were no significant differences between the three groups for MCT4 relative abundance in the EDL (Fig. 2B).

*NBC relative abundance.* NBC relative abundance in the soleus was significantly greater in both groups (+85% for PLA-T and +60% for ALK-T) compared with CON, with no significant difference between the two training groups (Fig. 2A). There were no significant differences between the three groups for NBC relative abundance in the EDL (Fig. 2B).

*Relationships between parameters.* A positive relationship was observed between MCT1 and NBC (r = 0.50, P < 0.05; Fig. 3), whereas there was no significant relationship between MCT4 and NBC (r = 0.31, P > 0.05). Furthermore, MCT1 and CS activity were positively correlated in the soleus (r = 0.57, P < 0.05).

**DISCUSSION**

We have observed typical increases in MCT1 and MCT4 relative abundance in response to high-intensity training. This is the first report, however, to demonstrate that NBC relative abundance can also be increased by training. Changes in MCT1, MCT4, and NBC were observed in the soleus only (i.e., a highly oxidative muscle) and not in the EDL (i.e., a highly glycolytic muscle). In contrast, high-intensity training significantly altered the in vitro muscle buffer capacity in both types of muscle. A second novel finding of the present study was that inducing metabolic alkalosis during training was associated with a significantly greater increase in MCT4 relative abundance in the soleus but did not affect other parameters in either muscle. Although there was no relationship between MCT4 and NBC relative abundance, MCT1 and NBC relative abundance were positively related in oxidative, but not glycolytic, muscle.

*Training Adaptations*

In the present study, MCT1 and MCT4 relative abundance were higher after training than in the control condition in oxidative muscle. These findings are in accordance with observations in other studies that have demonstrated that a period of training can increase the density of MCT1 (14, 24, 34) and MCT4 (34) proteins in skeletal muscle. Despite not reaching statistical significance, the percentage increase in MCT4 was similar to that reported in previous studies (24, 31, 34). The lack of significant increase in the present study can partly be attributed to the greater variability in the MCT4 response (i.e., low statistical power), which is again typical of previous research (9, 14).

In the present study, we confirm, via Western blotting, that NBC is present in the membranes of both oxidative and glycolytic skeletal muscle (25, 26). For the first time, however,
we have also shown that, like MCT1 and MCT4, NBC relative abundance can be increased by exercise training. Similar to previous results for MCT1 (2), we report that NBC also responds with a fiber-type specificity to high-intensity training (i.e., changes were observed in the soleus but not the EDL). Thus, we have shown that, like other membrane transport proteins that are involved in muscle pH regulation [i.e., MCT1 (14, 24), MCT4 (24, 31), and NHE (24, 31)], NBC is sensitive to chronic contractile activity in rat skeletal muscle.

The lack of change in either MCT1, MCT4, or NBC relative abundance in the EDL suggests that these proteins respond in a fiber-specific manner. One can suggest that potential increases in any of the proteins that we measured could have been underestimated due to an increase in total protein and/or total membrane protein contents. However, no change in protein contents in EDL and soleus was observed after training (Table 4), confirming the fiber-type specific response of these proteins. Furthermore, our results are consistent with previous research (2) that has reported greater training-induced increases in MCT1 in red than in white muscle in rats. Furthermore, Bonen (7) has demonstrated the same fiber-specific response of MCT1 to chronic muscle stimulation. These muscle-specific changes are unlikely to be due to less recruitment of the EDL during treadmill running exercise, as our pilot work demonstrated significant decreases in muscle pH in the EDL muscle after a typical exercise session. We also reported a significant increase in [3min \text{vitrō} in both the EDL and the soleus (Table 3).

Regulation of NBC, MCT1, and MCT4 Expression

Because MCT1, MCT4, and NBC all contribute to pH regulation, it might be predicted that H⁺ accumulation during training would be an important factor stimulating the synthesis of these proteins. Support for this idea is provided by previous research (13, 24, 31) that has shown that increases in NHE content (another important H⁺ transporter) tend to be greater in response to training that leads to a significant decrease in pH. Therefore, to investigate the role of H⁺ accumulation on MCT1, MCT4, and NBC adaptations to training in rats, we used sodium bicarbonate to experimentally manipulate the intracellular H⁺ concentration during high-intensity training.

As demonstrated by our pilot work, muscle proton accumulation differed in our two training groups (PLA-T > ALK-T).

In contrast to our hypothesis, the present results indicate that performing the same training with different H⁺ accumulation did not alter the training-induced improvements in MCT1 (+23% for ALK-T vs. +30% for PLA-T, $P > 0.05$). This finding is supported by the results of a recent study (31) demonstrating that increases in MCT1 in humans were not significantly different between two different training programs that resulted in significantly different changes in muscle pH (7.06 vs. 6.98). The observation that increases in NBC in the present study were also not significantly different between conditions (+60% for ALK-T vs. +85% for PLA-T, $P > 0.05$) may be related to the functional cooperation that has been identified between MCT1 and NBC (3).

Because an adequate contraction stimulus appears to be required to increase MCT1 density (2), we cannot exclude the possibility that a minimum accumulation of H⁺ is required to stimulate MCT1 and NBC synthesis. However, it appears that the degree of H⁺ accumulation is not the main factor stimulating further synthesis of these proteins. Recently, Hashimoto et al. (17) have demonstrated that lactate anion could be a signaling molecule to regulate MCT1 expression, but not MCT4, in L6 cells.

A novel finding of the present study was that the group that ingested sodium bicarbonate prior to training had a significantly greater increase in MCT4 relative abundance in the soleus than the placebo training group (+115 vs. +35%, $P < 0.01$; Fig. 2). Because control rats did not receive bicarbonate ingestion, we cannot exclude the possible effects of bicarbonate ingestion alone. However, since MCT4 content tended to increase in PLA-T, this suggests that the greater improvements in the ALK-T were due to the combined effects of training and sodium bicarbonate ingestion, possibly mediated via a reduced H⁺ accumulation during training. Although sodium bicarbonate ingestion has also been reported to increase muscle lactate accumulation (27), this appears unlikely to explain the differences in MCT4, as Mohr et al. (31) have reported no significant differences in MCT4 following two training programs that elicited very different muscle lactate values.

The mechanisms responsible for the greater improvements in MCT4 in ALK-T, when compared with PLA-T, cannot be determined from the present study. However, flux through a transport system during high-intensity exercise may provide a crucial stimulus for adaptation (24), and hydrogen ion efflux out of muscle cells has been reported to be enhanced by a greater extracellular buffer concentration (28). Therefore, it is possible that, in the present study, alkalosis induced a greater release of lactate and protons from working muscle (27) and that this stimulated greater MCT4 expression. It is difficult to speculate as to why this effect was seen only in the soleus (a highly oxidative muscle). However, it has been reported (7) that MCT4 content is greater in glycolytic fibers, and this may make it difficult to further elevate MCT4 content in such fibers (e.g., the EDL). Although sodium bicarbonate ingestion did not influence training-induced adaptations in MCT1 (or NBC), this is in accordance with previous research (10, 17, 24) demonstrating that transporters are regulated in an isoform-specific manner.
Relationship Between MCT1 and NBC

A further novel finding was the moderate, positive relationship between MCT1 and NBC relative abundance in oxidative, but not glycolytic, muscle. A greater MCT1 content has previously been shown (30) to be related to an increase in the transport of lactate (and H+) into the muscle. It may, therefore, be important that increases in MCT1 are accompanied by increases in NBC, as observed in the soleus in the present study, to help suppress the buildup of extracellular H+ and to maintain the pH gradient that is required for continued MCT1 activity (22). This hypothesis is supported by previous in vitro research that has shown that, when MCT1 is coexpressed with NBC1 in oocytes, lactate transport is enhanced (3). In this way, MCT1 and NBC could cooperate to facilitate lactate uptake in oxidative skeletal muscle during high-intensity exercise and recovery.

The moderate correlation coefficient between NBC and MCT1 relative abundance (r = 0.50, P < 0.05), however, suggests that other parameters are also likely to be involved in the regulation of lactate transport during high-intensity exercise and recovery. Of crucial importance to this process is the distribution and activity of carbonic anhydrase (CA), both in skeletal muscle and in the extracellular space, which help to establish a fast equilibrium between carbonic acid and CO2, HCO3-, and H+ (16). In addition, it has been reported that carbonic anhydrase IV, which is the sarcomemal CA (sCA) isofrom (16), has an essential interaction for full NBC1 activity (1). Recently, sCA has been shown to facilitate lactate transport in rat skeletal muscles (40). In consequence, although we have not demonstrated the colocalization of MCT1 and NBC, our results and those of the literature are in favor of a combination of NBC, MCT1, and sCA that may help to make the lactate shuttle more efficient between blood and muscle and to improve intracellular pH regulation, particularly after training. Surprisingly, no data are currently available concerning the relationship between MCT4 and either NBC or sCA. We did not, however, find a significant relationship between MCT4 and NBC relative abundance in the present study.

Conclusions

In conclusion, the present study has shown that MCT1, MCT4, and NBC undergo fiber-specific adaptive changes in response to high-intensity training and that NBC relative abundance is moderately related to MCT1 expression in oxidative, but not glycolytic, muscle. In consequence, we can speculate that lactate transport may be facilitated by NBC in oxidative skeletal muscle. Furthermore, although MCT4 content tended to increase with high-intensity training, alkalosis induced prior to each training session resulted in a significantly greater increase in MCT4 relative abundance in the soleus muscle, without any effect on MCT1 and NBC content.

ACKNOWLEDGMENTS

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REFERENCES


