Skeletal muscle water and electrolytes following prolonged dehydrating exercise

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We studied if dehydrating exercise would reduce muscle water (H_{2}O_{\text{muscle}}) and affect muscle electrolyte concentrations. Vastus lateralis muscle biopsies were collected prior, immediately after, and 1 and 4 h after prolonged dehydrating exercise (150 min at 33 ± 1 °C, 25% ± 2% humidity) on nine endurance-trained cyclists (V_{\text{O}_{2}\max } = 54.4 ± 1.05 mL/kg/min). Plasma volume (PV) changes and fluid shifts between compartments (CI method) were measured. Exercise dehydrated subjects 4.7% ± 0.3% of body mass by losing 2.75 ± 0.15 L of water and reducing PV 18.4% ± 1% below pre-exercise values (P < 0.05). Right after exercise H_{2}O_{\text{muscle}} remained at pre-exercise values (i.e., 398 ± 6 mL/100 g dw muscle^{-1}) but declined 13% ± 2% (342 ± 12 mL/100 g dw muscle^{-1}; P < 0.05) after 1 h of supine rest. At that time, PV recovered toward pre-exercise levels. The Cl^{-} method corroborated the shift of fluid between extracellular and intracellular compartments. After 4 h of recovery, PV returned to pre-exercise values; however, H_{2}O_{\text{muscle}} remained reduced at the same level. Muscle Na^{+} and K^{+} increased (P < 0.05) in response to the H_{2}O_{\text{muscle}} reductions. Our findings suggest that active skeletal muscle does not show a net loss of H_{2}O during prolonged dehydrating exercise. However, during the first hour of recovery H_{2}O_{\text{muscle}} decreases seemingly to restore PV and thus cardiovascular stability.

During prolonged exercise in the heat, subjects lose large amounts of fluid by sweating. However, the contribution of each body fluid compartment (intracellular, extracellular, and vascular) to sweat loss is not well defined. With the transition from rest to exercise, plasma volume (PV) rapidly declines by ~8% and from there, there is a progressive loss of PV to around 13% when exercise lasts 120 min (i.e., 450 mL loss; Costill et al., 1976). However, PV accounts for a small percentage of all the fluid losses that take place during prolonged exercise. Necessarily, fluid from the interstitial and intracellular fluid (ICF) spaces ought to be contributing to water loss during prolonged exercise. However, it is unclear to what extent water from active muscles contributes to the sweat losses during or after prolonged dehydrating exercise. Water has an important metabolic role in contracting muscle and its deficit could trigger catabolic responses (Häussinger, 1996) and have negative consequences in pH regulation and gene expression (Schäfer et al., 2007).

Costill and associates studied muscle water changes during (Costill et al., 1981) and after prolonged dehydrating exercise (Costill et al., 1976) in active and inactive muscles. They obtained muscle biopsies and carefully measured the loss of tissue weight after thorough drying using high temperature. They found that if biopsies are obtained during exercise, active muscle does not decrease but rather increases its water content (Costill, 1977). Sjøgaard and Saltin (1982) also reported increases in active muscle water when sampling muscle right after three bouts of supramaximal cycling exercise. Likewise, increases in muscle water have also been reported after intense knee extension exercise (Sjøgaard et al., 1985; Ploutz-Snyder et al., 1995; Kristiansen et al., 2014). However, when exercise is prolonged, Costill et al. (1976) were able to measure progressive reductions in muscle water that reached 7% when body weight decreased 5.8% by dehydration. Surprisingly, in a follow-up study, Costill et al. (1981) reported no changes in total muscle water content after a 3.2% reduction in body weight by exercise inducing sweating. They speculated that the muscle water loss could have been compensated by the water released from the breakdown of glycogen, water formed during oxidative phosphorylation, or water drawn from the gut. In this study, muscle biopsies were taken 30 min after exercise discontinuation. It is also possible that 30 min was insufficient time to allow fluid to shift from muscle to the vasculature, explaining the unchanged muscle water content after 3.2% whole-body dehydration.

Neufer et al. (1991) reported an 8% reduction in active muscle water after 4.8% whole-body dehydration by exercise. With similar degree of dehydration, Hackney...
et al. (2012) report only 5% reduction in knee extensor water content [measured by magnetic resonance imaging (MRI)]. The exercise intensity was higher in the Neufer et al. study and the sampling time also differed between studies. It is possible that the time elapsed between ending the exercise and biopsy (or MRI) sampling plays a role in the muscle water deficit results obtained. After exercise, hydrostatic pressure rapidly declines to resting values allowing muscle water to shift to other fluid spaces replenishing their fluid losses. Lastly, the role of muscle electrolytes on preserving or allowing muscle water losses is undetermined.

In the present study, we examine the changes in water and electrolyte contents of active muscle after large sweat losses induced by prolonged exercise in the heat. Our hypothesis is that during recovery from dehydrating exercise, water shifts from muscle to the extracellular fluid (ECF) space to help in the recovery of PV. Although indirect measures (i.e., Cl⁻ method) suggest that fluid shifts occur after exercise to replenish PV (Nose et al., 1988; Sanders et al., 1999; Hamouti et al., 2013), the specific role of muscle water on this recovery is unclear. To test our hypothesis, we sampled and analyzed muscle water and electrolyte contents before, right after exercise, and after 1 and 4 h of recovery. This should allow us to describe the timing and participation of muscle water in post-exercise fluid and electrolyte shifts among compartments.

Methods

Participants

Nine endurance-trained male cyclists, who routinely cycled at least 2 h/day, 4–7 days/week during the last 3 years, participated in the study. Participants were 24 ± 9 years old, weighed 69 ± 7 kg, and their height and percent body fat [dual-energy X-ray absorptiometry (DXA) evaluated] were 1.73 ± 7.1 m, and 14.8% ± 1.7%, respectively [all mean ± standard deviation (SD)]. Their VO₂max was of 54.4 ± 3.1 mL/kg/min. Subjects were fully informed about the experimental procedures and the possible risks and discomforts associated with the experiment before they gave their written informed consent to participate. The study was approved by the Virgen de la Salud Hospital Research Ethics Committee in accordance with the latest version of the Declaration of Helsinki.

Preliminary testing

In their first visit to the laboratory, participants underwent a physical examination including rest and exercise electrocardiogram (Cosmed T12, Rome, Italy). They performed an incremental cycling test to volitional fatigue using an electromagnetically braked cycle ergometer (Cardiotest 100, Seca, Hamburg, Germany) to determine their maximal aerobic power (VO₂max). After a 5-min warm-up at 100 W, participants began cycling at 125 W with increments of 25 W each minute. Gas exchange data were collected using an automated breath-by-breath system (Quark b2, Cosmed) and averaged every 15 s. VO₂max was defined as the highest plateau (two successive maximal readings within 0.15 L/min) reached. Percent body fat and legs lean soft-tissue mass were determined by DXA (DXA Hologic Series Discovery Wi QDR, Bedford, Massachusetts, USA). Regional analysis of leg soft-tissue mass included the gluteal and psoas muscles by analyzing scans from the iliac crest to the foot.

Experimental protocol

Participants refrained from exercising the day before the trials and performed a standardized low-intensity workout 48 h before the trial. Subjects were instructed to follow a diet rich in carbohydrates (5–8 g/kg/day) and drink liberally in the 24 h before the trial. Upon awakening, 1 h before arriving to the laboratory, participants were required to eat a standardized breakfast [i.e., 330 mL of a fruit milkshake (168 kcal) and a pastry (456 kcal)] and drink 500 mL of water to promote euhydration. Upon arrival to the laboratory, subjects voided and their nude body weight was measured using a ±0.05 sensitive scale (Seca 764). A urine sample was analyzed for specific gravity (Usg) using a hand-held refractometer (Master-Sur/NT, Atago, Tokyo, Japan) to confirm euhydration (i.e., <1.020; Sawka et al., 2007). If subject’s Usg was above 1.020, the trial was aborted and the subject was re-scheduled. Then, subjects lay down on a stretcher and were catheterized (20-gauge Teflon®; BD Insite, Becton Dickinson, Madrid, Spain) in an antecubital vein. The catheter was kept patent by frequent flushing with 1 mL of 0.9% sterile saline (Grifols, Barcelona, Spain). After 20 min, a muscle biopsy was obtained from the vastus lateralis and a blood sample drawn to be used as baseline for PV changes calculations.

Immediately after, participants dressed up in shorts and cleated cycling shoes, and entered in the climatic chamber, set at 33 ± 1 °C with 25% ± 2% relative humidity (mean ± SD) and 2.5 m/s wind speed, where they sat quietly on the cycle ergometer for 15 min. A heart rate monitor was positioned around their chest (RS400, Polar, Kempele, Finland) to record heart rate during exercise. Then, a pre-exercise blood sample was drawn (time 0 in Fig. 1) and subjects started pedaling at 65% VO₂max during 120 min followed by 30 more minutes at 55% VO₂max to complete 150 min of exercise. During exercise blood samples were collected at 15, 60, 120, and 150 min of exercise. Customized adhesive non-allergenic sweat patches (10 × 12 cm, Tegaderm™ plus Pad, 3M, St. Paul, Minnesota, USA) were placed in the middle back after cleaning the skin with deionized water as previously described (Hamouti et al., 2011) and collected after 30 min of sweat collection at 60, 120, and 150 min of exercise. Immediately after, participants dressed up in shorts and cleated cycling shoes, and entered in the climatic chamber, set at 33 ± 1 °C with 25% ± 2% relative humidity (mean ± SD) and 2.5 m/s wind speed, where they sat quietly on the cycle ergometer for 15 min. A heart rate monitor was positioned around their chest (RS400, Polar, Kempele, Finland) to record heart rate during exercise. Then, a pre-exercise blood sample was drawn (time 0 in Fig. 1) and subjects started pedaling at 65% VO₂max during 120 min followed by 30 more minutes at 55% VO₂max to complete 150 min of exercise. Exercise blood samples were collected at 15, 60, 120, and 150 min of exercise. Customized adhesive non-allergenic sweat patches (10 × 12 cm, Tegaderm™ plus Pad, 3M, St. Paul, Minnesota, USA) were placed in the middle back after cleaning the skin with deionized water as previously described (Hamouti et al., 2011) and collected after 30 min of sweat collection at 60, 120, and 150 min of exercise. At these time points nude body weights were also measured to calculate sweat rate. VO₂ and VCO₂ were measured using indirect calorimetry for 3 min after 15, 55, 115, and 145 min of exercise. Upon completion of the dehydrating exercise, the last sweat patch and the heart rate chest band were removed and participants exited the hot chamber. Subjects moved to a stretcher and then a second muscle biopsy sample was obtained (immediately post). Participants’ toweled-dry before post-exercise nude body weight was collected and then subjects voided when possible. Participants remained on the stretcher for 4 h, whereas blood samples, muscle biopsies, and body weight were obtained 1 and 4 h post-exercise. In an attempt to maintain the same level of dehydration and knowing that during the 3 h of rest water will be loss by sweating and breathing and urine will be formed, we provided subjects with 330 mL of water right after the 1-h biopsy. Urine was collected at any time during the resting period that the subjects could urinate. Once the trial concluded, participants were provided with food and drink, and instructions to take care of the biopsy wounds.

Body water balance

Urine volume during the trial was measured with a graduated cylinder (Simax 1000 mL, Prague, Czech Republic) and 1.5 mL of aliquots stored at −80 °C until analysis. Sweat volume was calculated from the losses of body weight subtracting respiratory and
metabolic weight losses. Respiratory water loss (i.e., \( m_e \)) and weight loss associated with excess CO\(_2\) expired (i.e., \( m_r \)) were estimated according to the following equations (Mitchell et al., 1972):

\[
m_e = 0.019 \times \text{VO}_2 \times (44 - \text{Pa}) \times \text{exercise time (min)}
\]

\[
m_r = \text{VO}_2 \times R \times (\rho_{\text{CO}_2} - \rho_{\text{O}_2}) \times \text{exercise time (min)}
\]

where \( m_e \) is evaporative water loss from the respiratory tract, \( \text{VO}_2 \) is oxygen uptake in L/min at standard temperature pressure dry air (STPD), and \( \text{Pa} \) is the ambient vapor water pressure in mmHg (i.e., 9.77 mmHg in our hot environment). \( m_r \) is the metabolic weight loss, \( R \) is the respiratory exchange ratio (VCO\(_2\)/VO\(_2\)), and \( \rho_{\text{CO}_2} \) and \( \rho_{\text{O}_2} \) are the densities of carbon dioxide (1.96 g/L STPD) and oxygen (1.43 g/L STPD), respectively.

Metabolic water formation during exercise was calculated from substrate oxidation. Substrate oxidation rates (fat and carbohydrate) were calculated from non-protein RQ measured by indirect calorimetry (Jeukendrup & Wallis, 2005) after 15, 55, 115, and 145 min of exercise. It was assumed that the oxidation of 1 g of glucose results in the formation of 0.6 g of water and the oxidation of 1 g of fat in the formation of 1.13 g of water (Maughan et al., 2007).

Urine, blood, and sweat analysis

Blood samples were immediately analyzed for hemoglobin concentration (ABL-520, Radiometer, Brønshøj, Denmark) and hematocrit by microcentrifugation (Biocen, Alresa, Madrid, Spain) in triplicate. Relative changes in PV (\( \Delta \text{PV} \)) were calculated with the equations outlined by Dill and Costill (1974). Based on subjects' pre-exercise body mass, PV was calculated following Sawka et al. (1992):

\[
\text{PV (mL)} = [(0.026 \times \text{body mass in kg}) + 1.239] \times 1000
\]

Absolute changes in PV (in mL) were calculated from this baseline calculation.

Sweat collected in the patches was separated by centrifugation in sealed tubes and stored at \(-80^\circ \text{C} \) until analysis. Urine and sweat samples were thawed to room temperature and analyzed for Na\(^+\), K\(^+\), and Cl\(^-\) concentrations using an ion-selective analyzer (Easylyte Plus, Medica Corporation, Bedford, Massachusetts, USA). Blood samples were allowed to clot in serum tubes (Z Serum Sep Clot Activator Vacuette®, Greiner Bio-One GmbH, Kremsmünster, Austria) and followed by centrifugation at 4000 rpm for 10 min at a temperature of 4 °C (MPW-350R, Med. Instruments, Warsaw, Poland) to obtain serum that was stored at \(-80^\circ \text{C} \). Serum was analyzed for Na\(^+\), K\(^+\), and Cl\(^-\) concentrations using an ion-selective analyzer (Easylyte Plus, Medica Corporation). Cl\(^-\) concentration from sweat patches was extrapolated to whole body concentrations according to Patterson et al. (2000) using the specific equation for lower back sweat collection. Serum protein was analyzed using the refractive index in a hand-held refractometer (Master-Sur/\( \alpha \), Atago). Percent change in the concentration and content of serum constituents was calculated from pre-exercise values of corrected hematocrit and solute concentrations according to the equation proposed by van Beaumont et al. (1973).

ECF and ICF shift

We determined extra- and intracellular water after 60, 120, and 150 min of exercise as well as 1 and 4 h into recovery. This determination was based on the chloride method (Costill et al., 1976). Changes in body water loss were estimated from body mass loss. Chloride losses were calculated by multiplying the volume of urine and sweat excreted by the respective concentrations ([Cl\(^-\)\(_\text{urine} \), Cl\(^-\)\(_\text{sweat} \)]. Changes in extracellular fluid, ECF, and ICF were calculated using Donnan factor-specific formulae for exercise conditions (Sanders et al., 2001).

Muscle biopsy

Muscle biopsy samples were taken from the vastus lateralis using the suction-modified Bergstrom technique (Tarnopolsky et al., 2011). Skin was prepared with povidone-iodine (Betadine, MEDA, Bordeaux, France), followed by injection of 2% lidocaine without epinephrine (Braun 2%, Braun Medical, Rubi, Spain). Then, the skin and underlying tissues were surgically opened (scalpel blade number 10, Braun, Tuttingen, Germany) and muscle tissue obtained using a 4-mm internal diameter Bergstrom biopsy needle. Upon collection, muscle samples were immediately cleaned of
connective tissue, divided into two pieces, and rapidly frozen in liquid nitrogen for subsequent analysis of water and electrolyte contents. The incision was closed using adhesive strips (Steri-Strip™, 3M) covered with an adhesive dressing pad (Tegaderm™+ Pad, 3M) and compressive dressing (IcoVenda, Novico Medica, Barcelona, Spain). All four biopsies were taken from the right leg using one incision for each biopsy. Incisions were spaced ~5 cm in between and were taken starting in the mid leg and advancing distally (Costill et al., 1988). Two incisions were made before exercise: one to obtain the pre-exercise biopsy and the other to save time in the immediately post-exercise biopsy procedure.

Measurement of muscle water and electrolytes

All the samples from a given subject were analyzed in the same assay batch. Frozen samples were weighed on an electronic balance with a sensitivity of 0.1 μg (XB220A, Precisa, Dietikon, Switzerland). Elapsed time from sample removal from the freezer until weighing was recorded to permit correction for tissue water evaporation. Samples were freeze dried in a thermodryelectric freeze dryer (Cryodos-50, Telstar, Madrid, Spain) for 12 h at −50 °C and at a vacuum of 10-2 Torr. In brief, this apparatus freezes the liquid in the sample to ~−50 °C then sublimates it with a potent vacuum pump at a high flux rate (83 L/min). Samples were then reweighed in the same precision scale, and the difference in weight (i.e., water) was expressed per 100 g of dry tissue. Pilot data in our laboratory in fresh pig leg muscle indicated that $H_2O_{\text{muscle}}$ measurement was highly reproducible (i.e., 6% CV). Data in one subject who underwent 10 resting biopsies within 8 weeks in a euhydrated state (Usg < 1.020 and body weight ± 0.25 kg) confirmed the high reproducibility of this technique in our laboratory in human muscle (i.e., 5% CV).

Segmental body DXA scan permitted calculation of active legs lean soft-tissue mass (i.e., muscle and connective tissue), which contain most of the limb water. Soft-tissue mass DXA analysis focused in the body segment from the iliac crests to the ankles. Combining those measures with the losses of water in muscle from the biopsies, and assuming a uniform loss of water throughout the leg musculature, we estimated leg muscle water losses after exercise as follows:

$$\text{Legs} \quad H_2O \text{loss} \quad (L) = \text{Biopsy} \quad H_2O \text{loss} \quad (L/\text{kg wt muscle}) \times \text{DXA Legs lean soft tissue (kg)}$$

Muscle electrolytes were measured according to the methods described in Costill et al. (1981). Muscle samples were submerged overnight in petroleum ether to extract neutral fat. Then, the pieces were reweighed to determine fat-free solid (FFS) weight. The dried specimen was subsequently treated with 100 μL of 2N nitric acid overnight to extract electrolytes. Muscle sodium ($Na^{+}_{\text{muscle}}$) and potassium ($K^{+}_{\text{muscle}}$) were determined in that extract using flame photometry (Clinical PFP7; Jenway, Stone, UK).

Statistical analysis

Data are presented as mean ± standard error of the mean unless otherwise stated. When the Shapiro–Wilk test revealed non-normally distributed data ($P < 0.05$), we used Friedman’s non-parametric statistical test. Normally distributed data were analyzed using one-way analysis of variance with repeated measures. After a significant $F$-value (Greenhouse–Geisser adjustment for sphericity), pairwise differences were identified using Tukey’s (honest significant difference) post-hoc procedure for normally distributed data, and Mann-Whitney $U$-test with correction for multiplicity for non-normally distributed data. Level of significant was set at $P < 0.05$. Data analysis was performed using SPSS software for windows (v.18, SPSS Inc., Chicago, Illinois, USA).

<table>
<thead>
<tr>
<th>Table 1. Body water loss and formation and metabolic weight loss during 150 min of dehydrating exercise in a hot environment</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Liter (%BW loss)</strong></td>
</tr>
<tr>
<td><strong>Body weight loss (kg)</strong></td>
</tr>
<tr>
<td><strong>Sweat loss (L)</strong></td>
</tr>
<tr>
<td><strong>Breath $H_2O$ loss (L)</strong></td>
</tr>
<tr>
<td><strong>Urine void (L)</strong></td>
</tr>
<tr>
<td><strong>$CO_2$ loss (kg)</strong></td>
</tr>
<tr>
<td><strong>Metabolic $H_2O$ formation (L)</strong></td>
</tr>
<tr>
<td><strong>Total $H_2O$ loss (L)</strong></td>
</tr>
<tr>
<td><strong>%Body weight loss from $H_2O$</strong></td>
</tr>
</tbody>
</table>

Data are means ± standard error of the mean for nine endurance-trained subjects.

Results

Percent dehydration and sweat rate

After 150 min of exercise subjects dehydrated by 4.7% ± 0.3% of their initial body weight, losing 2.75 ± 0.15 L of water by sweating at a rate of 1.06 ± 0.05 L/h. Total water lost explained 86% ± 1% of body weight losses (Table 1) even when metabolic water formation was accounted. Weight losses associated with excess $CO_2$ accounted for another 5% with a total of 92% ± 1% of the weight losses accounted in our calculations (Table 1).

PV changes

The transition from supine rest to standing rest in the cycle-ergometer reduced PV to −4.6% ± 0.6%. After 15 min of exercise PV was further reduced to −13% ± 0.6% (0.400 ± 0.02 L; Fig. 1; $P < 0.05$). The progressive dehydration reduced PV to −18.4% ± 1% after 120 min of exercise. That reduction was maintained during the last 30 min of exercise (i.e., loss of 0.545 ± 0.04 L after 150 min; $P < 0.05$). One hour after exercise, PV returned halfway toward the resting initial values (−10% ± 1%; Fig. 1), although it was still lower than pre-exercise resting values. After 4 h of supine recovery, PV was fully recovered with values not different from rest (−4.4% ± 2%; Fig. 1).

Changes in muscle water content

Prior to exercise vastus lateralis contained 395 ± 6 mL of water per each 100 g dw muscle$^{-1}$. $H_2O_{\text{muscle}}$ remained at pre-exercise values immediately after exercise (398 ± 5; Fig. 1). However, 1 h after exercise $H_2O_{\text{muscle}}$ declined by 13% ± 2% to 342 ± 12 mL/100 g dw muscle$^{-1}$ (Fig. 1; $P < 0.05$). This reduction of $H_2O_{\text{muscle}}$ was maintained for another 3 h with levels after 4 h of exercise of 346 ± 11 mL/100 g dw muscle$^{-1}$ (Fig. 1; $P < 0.05$). Leg water loss was calculated combining biopsy vastus lateralis $H_2O_{\text{muscle}}$ data with legs DXA scan of lean soft-tissue mass, assuming that this tissue contains most of the leg water. We detected a loss of $−0.529 ± 0.10$ L of...
Fluid losses in exercising skeletal muscle

Table 2. Water losses from resting values after 150 min of exercise in a hot environment (33 ± 1 °C, 25% ± 2% humidity)

<table>
<thead>
<tr>
<th></th>
<th>Immediately post-exercise</th>
<th>1 h post-exercise</th>
<th>4 h post-exercise</th>
</tr>
</thead>
<tbody>
<tr>
<td>Whole-body H_2O loss (L)</td>
<td>−2.750 ± 0.15</td>
<td>−2.750 ± 0.15</td>
<td>−2.590 ± 0.12</td>
</tr>
<tr>
<td>Legs H_2O loss (L)</td>
<td>0.012 ± 0.02</td>
<td>−0.529 ± 0.10*</td>
<td>−0.476 ± 0.10*</td>
</tr>
<tr>
<td>Plasma H_2O loss (L)</td>
<td>−0.545 ± 0.05</td>
<td>−0.307 ± 0.04*</td>
<td>−0.132 ± 0.05 †</td>
</tr>
<tr>
<td>Legs + plasma H_2O loss (L)</td>
<td>−0.533 ± 0.03</td>
<td>−0.836 ± 0.12*</td>
<td>−0.608 ± 0.12 †</td>
</tr>
<tr>
<td>% from whole-body H_2O loss (%)</td>
<td>19% ± 1%</td>
<td>30% ± 4%</td>
<td>23% ± 2%</td>
</tr>
</tbody>
</table>

Data are divided into whole-body, plasma, and previously active muscle (i.e., both legs). Data are means ± standard error of the mean for nine endurance-trained subjects.
*Significantly different from immediately post-exercise.
†Significantly different from 1 h post-exercise.

Fig. 2. Changes in muscle sodium and potassium during 150 min of dehydrating exercise and subsequent 4-h recovery. Data are means ± standard error of the mean for nine endurance-trained subjects. *Significantly different from rest values (P < 0.05).

Changes in muscle and blood serum electrolytes

Prior to exercise muscle Na⁺ and K⁺ were 9.5 ± 0.3 and 46.3 ± 0.5 mEq/100 g of FFS, respectively. After exercise muscle Na⁺ and K⁺ remained at pre-exercise levels according to the lack of changes in H₂O_muscle. However, after 1 and 4 h of supine rest, values increased above basal values to 11.6 ± 0.3 and 47.5 ± 0.2 mEq/100 g of FFS, respectively (Fig. 2; P < 0.05). These increases in muscle electrolytes coincided with the decreases in muscle water content. Changes from pre-exercise in concentration and contents of protein and electrolytes are depicted in Fig. 3. Serum K⁺, Na⁺, chloride (Cl⁻), and total protein concentration increased during exercise (32% ± 4%, 4% ± 1%, 6% ± 1%, 15% ± 3%, respectively; P < 0.05). These increases in concentration were due to increased contents as Na⁺, Cl⁻, and total protein contents were reduced. This was not the case with serum K⁺ content that increased during exercise (10% ± 2%; P < 0.05). During the recovery after exercise, serum concentrations of electrolytes and protein declined while the contents returned to pre-exercise levels. Plasma protein contents actually increased during recovery above its pre-exercise levels (9% ± 2%).

Body fluid shift

During dehydrating exercise PV declined and as a consequence plasma Cl⁻ increased (Table 3; P < 0.05). Average sweat chloride concentration ranged from 51 to 55 mEq/L during exercise. Urine chloride concentration varied from 124 to 160 mEq/L as an average (Table 3). During exercise ECF declined to −2.79 ± 0.08 L (P < 0.05) while ICF increased from basal levels (Table 3). Lowering the work load during the last 30 min of exercise partly recovered ECF returning ICF to basal levels. One hour after exercise ECF returned to pre-exercise values (Table 3) while ICF was at the lower level. Four hours after exercise ECF and ICF remained at the levels observed after 1 h of exercise.
During prolonged exercise (150 min) in a hot environment (33 °C; 25% ± 2% humidity), body water is lost mainly through sweating (Table 1). In this situation, intravascular fluid (i.e., blood plasma) decreases due to its participation in sweat formation. However, the losses of PV (0.545 ± 0.05 L; Fig. 1 and Table 2) only explain 19% of all the fluid lost during exercise (Table 2). Thus, 

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**Figure 3.** Changes in plasma concentration (a) and contents (b) of protein, sodium, potassium, and chloride during 150 min of dehydrating exercise and subsequent 4-h recovery. Data are means ± standard error of the mean for nine endurance-trained subjects. *Significantly different from rest values (P < 0.05).

**Table 3.** Chloride concentration in serum, sweat, and urine prior, during, and after exercise and calculated extracellular (ECF) and intracellular fluid (ICF) following the chloride method

<table>
<thead>
<tr>
<th></th>
<th>Rest</th>
<th>60 min</th>
<th>120 min</th>
<th>150 min</th>
<th>Post 1 h</th>
<th>Post 4 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cl⁻ serum blood (mEq/L)</td>
<td>103 ± 0.6</td>
<td>106 ± 0.6</td>
<td>107 ± 0.5</td>
<td>109 ± 0.4*</td>
<td>107 ± 1.7</td>
<td>105 ± 0.9</td>
</tr>
<tr>
<td>Cl⁻ sweat (mEq/L)</td>
<td>–</td>
<td>51 ± 3</td>
<td>55 ± 4</td>
<td>51 ± 3</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Cl⁻ urine (mEq/L)</td>
<td>124 ± 16</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>141 ± 17</td>
<td>160 ± 9</td>
</tr>
<tr>
<td>ECF (L)</td>
<td>0.0 ± 0.0</td>
<td>–2.2 ± 0.8*</td>
<td>–2.8 ± 0.8*</td>
<td>–1.6 ± 0.6*</td>
<td>0.0 ± 0.1</td>
<td>0.0 ± 0.2</td>
</tr>
<tr>
<td>ICF (L)</td>
<td>0.0 ± 0.0</td>
<td>1.2 ± 0.8</td>
<td>0.5 ± 0.8</td>
<td>–1.8 ± 0.5*</td>
<td>–3.4 ± 0.2*</td>
<td>–2.9 ± 0.3*</td>
</tr>
</tbody>
</table>

Data are means ± standard error of the mean for nine endurance-trained subjects. *Significantly different from rest values.

**Discussion**

During prolonged exercise (150 min) in a hot environment (33 °C; 25% ± 2% humidity), body water is lost mainly through sweating (Table 1). In this situation,
participation of interstitial and intracellular fluid on sweat formation for heat dissipation was initially hypothesized. Three decades ago, David Costill’s group measured muscle water (H2O_{muscle}) changes after dehydrating exercise using muscle biopsies. Although in one study they showed a parallel reduction of H2O_{muscle} along with whole-body dehydration (Costill et al., 1976), they were not able to show reductions in H2O_{muscle} with similar dehydration in a subsequent study (Costill et al., 1981). Our data help to shed light into this apparent controversy. No net H2O_{muscle} loss was detected early after exercise (Costill et al., 1981) but after 1 h of supine rest a 13% reduction H2O_{muscle} appeared (Costill et al., 1976). Thus, a fluid equilibration time of 1 h is needed to measure the effects of whole-body dehydration on the previously active musculature.

We are not the first to report muscle water reductions induced by exercise in a hot environment. Using MRI technique, Hackney et al. (2012) reported a 1.2% reduction in leg volume for each 1% decrease in body mass. Costill et al. (1976) also reported a similar ratio between losses of muscle water and whole-body weight. This ratio is interesting to define as it could allow predictions of muscle water losses by only measuring changes in body weight after exercise. However, the ratio between whole-body and muscle dehydration seems to depend on the intensity of exercise. Both investigators (Costill et al., 1976; Hackney et al., 2012) used very hot environments (37–39 °C) and low workload (97 ± 39 W; Hackney et al., 2012) to achieve up to 5% dehydration. In contrast, Neufer et al. (1991) using intense intermittent cycling and a warm environment reported an 8% reduction in active muscle water after 4.8% whole-body dehydration by exercise.

Our moderately intense pedaling produced a ratio between whole-body and muscle dehydration similar to Neufer’s (i.e., 2.5% per each 1% loss of body weight). This suggests that exercise intensity influences the rate of muscle water loss in contracting skeletal muscle, with a higher rate indicating a higher workload. At low workloads, glycogenolysis rate is low (Horowitz et al., 1999) and less water may be liberated and available to be shifted away from the muscles. Prior studies (Costill et al., 1976; Neufer et al., 1991; Hackney et al., 2012) established that muscle water participates in the losses of fluid that take place during prolonged exercise in the heat. Our study helps to define the factors that influence the rate of muscle water loss during prolonged dehydrating exercise (e.g., exercise intensity). Furthermore, our study is unique in that it explores the connection between plasma and water muscle changes during the acute and prolonged phases of recovery (i.e., immediately, 1 and 4 h after exercise) after exercise.

During recovery from exercise, we observed that PV returned toward basal levels at the same time that H2O_{muscle} sharply declined 1-h post-exercise (Fig. 1). Our interpretation is that H2O_{muscle} reduction after exercise reflects the contribution of muscle fluid to the initial stages of PV recovery. Nose et al. (1988) have shown that two-thirds of PV are recovered 1 h after dehydrating exercise (2.3% weight losses). We have recently shown (Hamouti et al., 2013) that, even if no rehydration is allowed, one-half of PV is recovered overnight after 3% dehydration is induced by exercise. At that time and based on indirect calculations (chloride method), we speculated that interstitial and intracellular water participated in the partial restoration of blood volume after exercise (Hamouti et al., 2013). Now, in addition to the indirect chloride method, we detected a post-exercise reduction in H2O_{muscle} coinciding with a recovery of PV (Fig. 1). Our data suggest that to restore PV after dehydration, exercise water from previously active skeletal muscle is likely transferred to the vasculature. This restoration of PV helps to regain cardiovascular stability.

Using DXA regional analysis we assessed lower body soft-tissue mass. We extrapolated the water loss in the biopsy tissue after exercise to all the lower body soft-tissue mass measured in each individual (e.g., DXA; Table 2). To perform this calculation, we assumed that most leg lean soft tissue is muscle and that water was lost evenly throughout the leg musculature. We may have incurred into a slight overestimation as we chose to biopsy vastus lateralis which together with vastus medialis are the most heavily recruited muscles during cycling (Farina et al., 2004) and likely more prone to lose water (Hackney et al., 2012). Despite this overestimation, muscle and plasma water losses together could not account for the water lost during exercise as sweat, breath, and urinary water (Table 2). Thus, other tissues must have participated in body water losses. Data from Costill et al. (1981) indicate that inactive muscle groups (i.e., deltoids during cycling) do not lose water during dehydrating pedaling in the heat. Other candidate tissues to provide water for thermoregulation during prolonged exercise are the interstitial space, the skin, and the gastrointestinal system.

During exercise or immediately after exercise, something prevented H2O_{muscle} from mitigating PV reductions. During exercise the increased muscle perfusion pressure (i.e., blood pressure) forces plasma into the interstitial space of active muscle (Friedman et al., 1982). In addition, the increased metabolite concentration during muscle contraction raises osmotic forces that draw fluid out of the plasma into the muscle (Lundvall, 1972). However, upon exercise discontinuation, blood pressure drops back to resting values or lower (Senay et al., 1980) allowing fluid to be reincorporated into plasma. In addition, during recovery, the metabolites accumulated during exercise are gradually removed or oxidized and osmotic forces re-equilibrate between blood and muscle. Still, right after exercise, H2O_{muscle} content did not drop below basal levels. It is possible that during exercise H2O_{muscle} increased above basal levels, and that there was an actual drop of H2O_{muscle} returning to pre-exercise levels, explaining the lack of net changes at the end of prolonged exercise. Short-term intense exercise increases active
muscle water content (Sjøgaard & Saltin, 1982; Ploutz-Snyder et al., 1995; Kristiansen et al., 2014). However, it is unclear if that acute increase in exercising muscle water is maintained during prolonged dehydrating exercise.

There are alternative explanations for the lack of $H_2O_{\text{muscle}}$ loss right after exercise. Costill (1977) speculated that the unchanged $H_2O_{\text{muscle}}$ after exercise may be due to the compensatory role of water formation during oxidation and/or from water liberated from glycogen. These sources of water could have matched the $H_2O_{\text{muscle}}$ lost resulting in unchanged levels after exercise. Substrate oxidation was calculated based on non-protein respiratory exchange ratio, and water from oxidizing fat and carbohydrates was based on Maughan et al. ‘s (2007) publication. Water from oxidation mounted to 0.27 ± 0.01 L that could have been produced during the 150 min of exercise (Table 1).

Although the amount of water bound to glycogen that is liberated during exercise is unknown (Sherman et al., 1982), based on associations between water and glycogen storage after exercise (Olsson & Saltin, 1970), a ratio of 3 to 1 has been proposed. Based on a previous study (Horowitz et al., 1999), we calculated an average glycogen use of 1.8 g/min during the exercise intensity held in our trial. Thus, 0.81 L of water bound to glycogen could have been liberated in the exercising muscle in our study. Even taking into account these sources of water formation, it is possible that active leg muscles had contributed up to 1.1 L of the sweat losses, without incurring in any measurable $H_2O_{\text{muscle}}$ net deficit. Further studies with metabolic tracing of water may confirm or correct this prediction.

Our data suggest that muscle participates in the recovery of PV although it may be a limitation to the water that can be lost from muscle. This is evidenced by the lack of loss in $H_2O_{\text{muscle}}$ from 1 to 4 h of recovery despite progressive PV recovery during this time. Also, body fluid shift calculations corroborated biopsy measurements in that no recovery of ICF was present in the 4 h of supine post-exercise. It is possible that osmotic forces may have re-equilibrated between plasma and muscle 1 h after exercise. This may have halted the driving force for fluid exchange between these two compartments. Corroborating this osmotic equilibration, we reported that muscle electrolytes (Na⁺ and K⁺) remained at high levels from 1 to 4 h after exercise (Fig. 2). Despite the unchanged $H_2O_{\text{muscle}}$, PV kept recovering during the last 3 h of the 4-h post-exercise recovery period, which indicated that other tissues apart from muscle participated in PV restoration. Thus, although an important contributor, $H_2O_{\text{muscle}}$ is not the only fluid involved in the restoration of PV after exercise. Other fluid compartments (i.e., interstitial) are likely involved in this response.

The redistribution of water across the muscle cell membrane during dehydration is dependent on the osmotic gradient and the activity of the ionic pumps (Sawka, 1992). Osmotic forces that draw fluid into plasma are primarily determined by serum proteins (osmotic pressure) and electrolytes. However, measuring the concentration of these metabolites do not fully inform of the fluid shift but rather the end result of those shifts. To gain insight into the nature of the fluid shift during and after prolonged dehydrating exercise, we calculated plasma contents of protein and electrolytes (Fig. 3). During exercise, total protein concentration was increased while protein content decreased. This indicates that total protein concentration was primarily a result of the loss of fluid from plasma. However, after exercise, total protein concentration declined while content in plasma increased. Influx of protein into plasma during recovery has been previously reported (Senay et al., 1980; Gillen et al., 1991). During exercise, plasma contents of Na⁺ and Cl⁻ decreased due to their excretion in sweat, while their concentration increased (van Beaumont et al., 1973) due to the hypotonic nature of sweat inducing larger losses of plasma than electrolytes. However, plasma K⁺ content increased during exercise likely from muscle sources (Kilburn, 1966). In summary, the increase in plasma Na⁺, Cl⁻, and protein concentration with exercise are the results of hemococoncentration and not induced by a release of proteins from extravascular sources. The hemodilution observed during recovery from prolonged dehydrating exercise occurs despite release of proteins and electrolytes into the plasma.

In summary, we found that (a) right after exercise muscle $H_2O$ is not reduced from pre-exercise values suggesting that either muscle does not contribute to water for thermoregulation (i.e., sweat) or that water is formed (metabolic or glycogen bound) at the same rate that it is lost from the muscle; (b) however, 1 h after exercise $H_2O_{\text{muscle}}$ declines likely to restore PV; and (c) finally, 4 h after exercise PV keeps recovering without apparent contribution from muscle $H_2O$. Our data suggest that muscle water contributes to the first stages of PV recovery after prolonged dehydrating exercise. However, this contribution is not evident right after exercise and requires 60 min of supine rest to emerge.

**Perspectives**

Insufficient fluid replacement during physical activities that entail profuse sweating results in whole-body fluid deficit. However, how that deficit is shared by the different body fluid compartments is unclear, however relevant. Water has important metabolic roles in human skeletal muscle and its deficit may delay the recovery of muscle energy substrates and the elimination of built-up metabolites, and could even stimulate catabolism in muscle (Häussinger, 1996). Our data show that after large whole-body dehydration (4.7%) plasma fluid is fully recovered at the expense of fluid from skeletal muscle during the first hour of recovery (Fig. 1). In this study, we describe the magnitude and timing of this muscle water deficit after exercise. The metabolic consequences of this post-exercise skeletal muscle dehydration are still to be described.
**Key words:** Exercise, plasma volume, body water, water loss, intracellular fluid, extracellular fluid.

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