Effects of depletion exercise and light training on muscle glycogen supercompensation in men

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1Naval Health Research Center, San Diego, California 92186; 2Departments of Internal Medicine and 3Cellular and Molecular Physiology, and 4Howard Hughes Medical Institute, Yale University, School of Medicine, New Haven, Connecticut 06510; and 5GEO-CENTERS, Inc., Newton, Massachusetts 02459

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Goforth, Harold W., Jr., Didier Laurent, William K. Prusaczyk, Kevin E. Schneider, Kitt Falk Petersen, and Gerald I. Shulman. Effects of depletion exercise and light training on muscle glycogen supercompensation in men. Am J Physiol Endocrinol Metab 285: E1304–E1311, 2003.—Supercompensated muscle glycogen can be achieved by using several carbohydrate (CHO)-loading protocols. This study compared the effectiveness of two “modified” CHO-loading protocols. Additionally, we determined the effect of light cycle training on muscle glycogen. Subjects completed a depletion (D, n = 15) or nondepletion (ND, n = 10) CHO-loading protocol. After a 2-day adaptation period in a metabolic ward, the D group performed a 120-min cycle exercise at 65% peak oxygen uptake (V˙O2 peak) followed by 1-min sprints at 120% V˙O2 peak to exhaustion. The ND group performed only 20-min cycle exercise at 65% V˙O2 peak. For the next 6 days, both groups ate the same high-CHO diets and performed 20-min daily cycle exercise at 65% V˙O2 peak followed by a CHO beverage (105 g of CHO). Muscle glycogen concentrations of the vastus lateralis were measured daily with 13C magnetic resonance spectroscopy. On the morning of day 5, muscle glycogen concentrations had increased 1.45 (D) and 1.24 (ND) times baseline (P < 0.001) but did not differ significantly between groups. However, on day 7, muscle glycogen of the D group was significantly greater (p < 0.01) than that of the ND group (130 ± 7 vs. 104 ± 5 mmol/l). Daily cycle exercise decreased muscle glycogen by 10 ± 2 (D) and 14 ± 5 mmol/l (ND), but muscle glycogen was equal to or greater than preexercise values 24 h later. In conclusion, a CHO-loading protocol that begins with a glycogen-depleting exercise results in significantly greater muscle glycogen that persists longer than a CHO-loading protocol using only an exercise taper. Daily exercise at 65% V˙O2 peak for 20 min can be performed throughout the CHO-loading protocol without negatively affecting muscle glycogen supercompensation.

carbohydrate loading; detraining; 13C magnetic resonance spectroscopy

MANY ENDURANCE ATHLETES practice muscle glycogen supercompensation or carbohydrate (CHO) loading, even though the factors affecting glycogen kinetics and the underlying mechanisms are not completely under-

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of repetitive biopsies is limited to three to five per subject. Even this low number of biopsies, if taken from an area of muscle ≤7 cm², can negatively affect glycogen synthesis (7). Other limitations of many CHO-loading studies include a lack of rigorous control of diet and/or exercise throughout the study and measurement of muscle glycogen for only 3–4 days of CHO loading, even when a plateau has not been reached.

Since the validation of ¹³C magnetic resonance spectroscopy (MRS) (36), it has become possible to make frequent, noninvasive measures of muscle glycogen without traumatizing the muscle. It is now possible to profile the time course of muscle glycogen kinetics with good precision (±5 mmol/l muscle) and determine the relative effects of important variables of CHO-loading protocols (e.g., diet and exercise).

When CHO loading, competitive athletes generally prefer moderate training to rest, out of a concern that detraining may occur and offset the potential benefits of supercompensated muscle glycogen. Their concerns are supported by studies reporting that even 3–6 days of rest can significantly reduce several metabolic adaptations acquired through endurance training (25, 26). On the other hand, CHO-loading protocols that include a high-intensity depletion exercise (90–180 min) and/or daily long (45- to 60-min) training sessions can prevent muscle glycogen supercompensation (10, 34).

The present study was designed to determine 1) the relative importance of an exhaustive depletion exercise bout and a high-CHO diet on the time course of muscle glycogen supercompensation and 2) the effect of daily light training sessions [20 min at 65% peak oxygen uptake (VO₂peak)] on muscle glycogen concentrations during and after CHO loading.

METHODS

Subjects

Twenty-five male volunteers from US Navy and Marine Corps Special Operations commands gave written informed consent to participate in this study. Subjects were engaged in 60–120 min of daily training that included aerobic training (running, swimming, cycling), calisthenics, and resistance exercises (weight lifting). This study was approved by the Human Investigation Committee of Yale University School of Medicine (New Haven, CT) and the Committee for the Protection of Human Subjects at the Naval Health Research Center (San Diego, CA). Partial data from this study were previously presented in a study examining the effects of caffeine on the neuroendocrine axis and muscle glycogen utilization during a 2-h cycle exercise bout (24).

Physical Characteristics

Anthropometric and VO₂peak measurements were made 1–2 wk before the study. Percent body fat was estimated from seven-site skinfold thickness (18). VO₂peak was determined by use of a multistage exercise test on a mechanically braked cycle ergometer (model 818-E, Monark). The test began with exercise at 60 W, which increased in a stepwise fashion by 60 W every 4 min until volitional fatigue or failure to maintain 60 rpm (19). Respiratory gases were measured over 15-s intervals with open-circuit spirometry (Horizon 4400, SensorMedics). Heart rate was monitored (Polar Vantage XL) during the final 15 s of each 4-min stage. Subjects were admitted to the Yale/New Haven Hospital General Clinical Research Center (GCRC), where they underwent medical screening that included a physical exam, medical history evaluation, and routine blood tests.

Experimental Design

Subjects completed a depletion (n = 15) or nondepletion (n = 10) modified CHO-loading protocol (Fig. 1). Subjects were assigned to treatment groups matched for age, VO₂peak, body weight, and body fat. The study began with a 2-day

![Fig. 1. Experimental design with time line for diet, exercise, muscle, and blood samples. D, depletion group; ND, nondepletion group; B, blood sampling; MG, muscle glycogen; CHO, carbohydrate; VO₂peak, peak oxygen uptake.](https://www.ajpendo.org/ AJP-Endocrinol Metab • VOL 285 • DECEMBER 2003 • www.ajpendo.org)
adaptation period during which subjects stayed in the GCRC and consumed a normal mixed diet. On the morning after the adaptation period, groups performed one of two exercise protocols. For the remainder of the study, both groups performed 20 min of daily cycle exercise and consumed the same diets. Under fasting conditions at the same time every day (0800–1200), muscle glycogen concentrations in the vastus lateralis were measured using $^{13}$C MRS. Glycogen measurements also were made before and after daily 20-min cycle exercise bouts (twice for the depletion group and once for the nondepletion group) during the loading and maintenance phases.

Urinary output, food and water intake, and body weights were measured daily. Fasting venous blood samples were collected from all subjects at 8:00 AM on days 1–6 and on the morning of day 7 (end of study). Blood samples also were collected from the depletion group before exercise, at 120 min, and after exhaustion in the depletion exercise bout. Samples were analyzed for selected hormones, metabolites, and standard blood chemistry.

**Exercise Protocols**

On the first day of the loading protocol, the depletion group performed a depletion exercise by cycling at 70 rpm for 120 min at 65% $V_{\text{O2 peak}}$, followed by repeated 1-min sprints at 90 rpm and 120% $V_{\text{O2 peak}}$, separated by 1-min rest periods (13). Exercise ended with volitional fatigue or failure to maintain a pedal rate of $\geq 90$ rpm. The nondepletion group performed 20 min of cycle exercise at 65% $V_{\text{O2 peak}}$ on the 1st day. On all other days, both groups performed 20 min of cycle exercise at 65% $V_{\text{O2 peak}}$ at 70 rpm.

**Dietary Protocols**

All meals were planned by a registered dietician and prepared in the metabolic kitchen of the GCRC. Diets for both treatment groups were designed to contain 195 kJ (46.65 kcal)-kg body wt $^{-1}$-day $^{-1}$; however, slight adjustments were made to maintain body weight over the course of the study. Subjects were encouraged to eat all of the food provided. Food and beverage consumption was carefully monitored, and any uneaten food was documented. During the 2-day adaptation period, subjects consumed a normal mixed diet (44% CHO, 38% fat, and 18% protein). During the 3-day loading phase (days 1–3), subjects consumed a high-CHO diet (~9 g CHO·kg$^{-1}$·day$^{-1}$, ~675–745 g CHO/day) composed of 80% CHO, 10% fat, and 10% protein. This diet included three bottles of a glucose-polymer beverage containing 105 g CHO/532 ml bottle (CarboForce, American Body-Building, Walterboro, SC). During the 3-day maintenance phase (days 4–6), subjects consumed a moderate-CHO diet (~6.5 g CHO·kg$^{-1}$·day$^{-1}$, or 480–530 g CHO/day) composed of 56% CHO, 26% fat, and 18% protein, and including two bottles of CarboForce. Each day of the loading and maintenance phases, one bottle of CarboForce (~1.4 g CHO/kg body wt) was consumed within 30 min after exercise.

**MRS**

Natural abundance $^{13}$C MRS was performed daily to determine noninvasively the time course of muscle glycogen depletion and repletion during the CHO-loading and -post-loading periods. Baseline muscle glycogen concentration values were determined after the 2-day adaptation period in the GCRC. Muscle glycogen concentration was also measured before and after the exhausting (2-h) and moderate (20-min) cycle exercise bouts. Measurements were performed in a 2.1-T Bruker Biospec spectrometer with a 1-m-diameter magnet bore, as previously described (28). A mark was made on the subject’s quadriceps muscle to facilitate repeated measures of muscle glycogen from the same site.

During the measurements, subjects remained supine, with the observation radio frequency (RF) probe resting above the quadriceps muscle. The probe consisted of a 9-cm-diameter inner coil for $^{13}$C acquisition and a 13-cm outer butterfly coil for $^1$H acquisition, image-guided positioning, and decoupling. Proton water line widths were shimmed to $<50$ Hz. The probe was positioned by an image-guided localization routine that used a T$\text{$_1$}$-weighted gradient-echo image so that the observation volume was typically ~1 cm$^3$ into the vastus lateralis muscle. The reference standard consisted of a microsphere containing [1$^{13}$C]formic acid (99% $^{13}$C enriched) fixed at the center of the double-tuned RF coil. Calibration of RF pulse widths was performed by determining the 180° flip angle at the center of the observation coil from the microsphere standard. Then the RF pulse width was set so that the 90° pulse was sent to the center of the muscle to obtain maximum suppression of the lipid signal from the subcutaneous fat layer and optimized signal from the muscle. $^{13}$C spectra were obtained with a $^1$H-decoupled pulse-acquired sequence in 10-min blocks consisting of 5,500 scans with a 90° pulse at coil center and a repetition time (TR) of 120 ms. Decoupling at the $^1$H frequency at a power of 15 W was applied at the C1 proton resonance frequency during the 25.6-ms acquisition period with a power deposition $<$4 W/kg (29).

Intramuscular glycogen concentrations were determined by comparison with signal from an external standard solution (150 mmol/l glycogen in 50 mmol/l KCl). The KCl in the glycogen standard allows for the phantom to have properties similar to a human leg. $^{13}$C spectra were processed by methods that have previously been described (28). Briefly, Gaussian broadened spectra (30 Hz) were baseline corrected by ±300 Hz on either side of the [1,1$^{13}$C]glycogen resonance of both subject spectra and standard spectra. Peak areas were then assessed at ±150 Hz of the resonance. Greater precision in measuring the exercise-induced change in muscle glycogen concentration was gained by using differential spectral analysis. The $^{13}$C MRS glycogen signal was corrected for the proton volume of the $^{13}$C coil. A phantom phantom solution of glycogen was acquired using a dedicated proton coil of the same size as the $^{13}$C coil and a fully relaxed gradient-echo sequence with a 90° excitation pulse. The sensitive volume of that image was manually drawn and compared with the set of images previously recorded from each individual by means of the butterfly coil. The corresponding filling coefficient was calculated according to the ratio of the regions of interest defined from the leg muscles and the in vitro data set (1.18 in average). In addition, the $^{13}$C MRS signal was corrected for the load of the $^{13}$C coil. A fully relaxed spectrum (4 pulses, TR = 15 s, pulse length 100 μs) of the formic acid sphere was recorded both in vivo and in the presence of the phantom solution of glycogen. The ratio of the formate peak area obtained in both conditions was used as the loading correction factor (0.91 in average). The detection threshold for minimum change is ~5 mmol/l muscle, which is 7% of the baseline concentration. This method has a reported coefficient of variation of 4.3% in resting subjects and is lower than the one typically reported for the biopsy technique (36). The $^{13}$C MRS technique for assessing intramuscular glycogen concentrations has been validated in situ in frozen rabbit muscle (12) and by comparison with human gastrocnemius muscle biopsies (36).
Blood Analyses

Plasma glucose was measured by the glucose oxidase method (Beckman glucose analyzer, Fullerton, CA). Plasma immunoreactive insulin, glucagon, cortisol, and human growth hormone (hGH) concentrations were measured using commercially available double-antibody radioimmunoassay kits (insulin: Diagnostic System Laboratories, Webster, TX; glucagon: Linco Research, St. Charles, MO; cortisol: Diagnostic Products, Los Angeles, CA; hGH: Sanofi Diagnostics Pasteur, Chaska, MN). Plasma triglycerides were determined fluorometrically at 340 nm with a quantitative enzymatic kit (Triglyceride Assay Kit; Wako Chemicals, Richmond, VA). Plasma lactate was measured using a lactate dehydrogenase assay kit (Sigma Diagnostics, St. Louis, MO). Plasma fatty acids were measured using a microfluorimetric assay (27). Plasma catecholamine (epinephrine and norepinephrine) concentrations were determined using a three-step procedure involving adsorption onto alumina (pH = 8.6), elution with a dilute acid, and analysis by high-pressure liquid chromatography.

Statistical Analyses

Blood metabolites, hormones, and muscle glycogen concentrations are presented as means ± SE in the text, Figs. 1–3, and Tables 1–4, unless noted otherwise. Statistical comparisons were made using a two-way repeated-measures analysis of variance (ANOVA). For within-subject comparisons, paired t-tests were used. When significant interactions were observed, unpaired t-tests were performed for time-point comparisons. When both multiple paired and unpaired t-tests were used, the alpha level was adjusted using the Bonferroni procedure. Significant differences between values and groups were accepted at P < 0.05.

RESULTS

Subject Characteristics

There were no significant differences between the depletion and nondepletion groups for any anthropomorphic or physiological characteristic. The depletion and nondepletion groups had mean ages (25.7 ± 1.1 and 27.1 ± 1.8 yr), heights (177.6 ± 1.6 and 175.3 ± 2.6 cm), weights (80.7 ± 2.0 and 75.1 ± 3.4 kg), body fat values (9.3 ± 0.7 and 9.2 ± 0.9%), and VO₂peak values (49.1 ± 1.4 and 47.4 ± 1.1 ml·kg⁻¹·min⁻¹), respectively.

Dietary Intake

There were no significant differences in the diets consumed by the two treatment groups in total kilocalories and absolute or percent intake of macronutrients (i.e., CHO, fat, and protein) during any phase of the study. Daily CHO intake for depletion and nondepletion groups averaged 9.2 and 9.0 g/kg total body mass (675–745 g CHO/day) during the loading phase and 6.6 and 6.4 g/kg (480–530 g CHO/day) during the maintenance phase, respectively. Daily caloric intake for both groups averaged 185 kJ/kg during the adaptation phase, 193 and 189 kJ/kg during the loading phase, and 197 and 191 kJ/kg during the maintenance phase for depletion and nondepletion groups, respectively. These daily intake values approximated the target value of 195 kJ/kg.

Muscle Glycogen Utilization

Effects of depletion exercise (2 h at 65% VO₂peak) and sprints. Baseline (preexercise) muscle glycogen concentrations for the depletion (89 ± 4 mmol/l) and the nondepletion (91 ± 5 mmol/l) groups did not differ significantly (Fig. 2). After the depletion exercise, the muscle glycogen of the depletion group decreased to 38 ± 6 mmol/l (i.e., 40 ± 19% of the preexercise value). Effects of moderate exercise (20 min at 65% VO₂peak). The muscle glycogen used by the depletion group during the 20 min of cycle exercise on day 3 of CHO loading and on day 5 (2nd day of maintenance diet) averaged 11 ± 3 and 9 ± 3 mmol/l, respectively (Fig. 3). The muscle glycogen of the nondepletion group during the 20-min exercise bout on day 4 (1st day of maintenance diet) decreased by 14 ± 5 mmol/l. The mean decrease in muscle glycogen during the 20-min cycle exercise bout under CHO-loaded conditions did not differ significantly between the two groups and had a pooled average of 11 ± 2 mmol/l for the 20-min bout.

Muscle Glycogen Repletion during CHO Loading

High-CHO diet/loading phase (days 1–3). During the first 24 h after the depletion exercise, muscle glycogen
of the depletion group increased significantly from 40 to 92 mmol/l muscle (Fig. 2). In contrast, after 24 h on the same diet, muscle glycogen of the nondepletion group remained approximately the same (85–92 mmol/l muscle). After 3 days of loading, the muscle glycogen of the depletion and nondepletion groups had increased to levels that were significantly (P < 0.01) greater (138 and 124%, respectively) than those at baseline. We found no significant correlation between the relative or absolute level of muscle glycogen depletion of individuals and their initial rate of repletion or peak muscle glycogen value.

**Moderate CHO diet/maintenance phase (days 4–6).** During this phase, muscle glycogen concentrations of the depletion group continued to increase (nonsignificantly) on days 4 and 5 and peaked on day 6 at 147% of baseline (Fig. 2). This level of supercompensation persisted until the final measurement on the morning of day 7. In contrast, muscle glycogen levels of the nondepletion group peaked on day 4 (at 124% of baseline), and then decreased back to baseline concentration on day 7. On day 7, the muscle glycogen of the nondepletion group (104 ± 5 mmol/l) was ~25% lower than that of the depletion group (130 ± 7 mmol/l; Fig. 2). This approximates the average amount of muscle glycogen (i.e., 26 mmol/l) used per hour during the 2-h depletion exercise performed by the depletion group.

**Metabolite and Hormonal Data**

**Fasted and rested conditions.** Throughout the study, resting plasma triglyceride, fatty acid, glucose, and lactate concentrations were within normal limits and did not differ between groups, with one minor exception (Table 1). On the morning after the depletion exercise, fatty acid values were within normal values but significantly greater in the depletion group (346 ± 41 μM) than in the nondepletion group (226 ± 40 μM). Throughout the study period, there were no significant differences between the depletion and nondepletion groups in fasting plasma insulin, glucagon, epinephrine, norepinephrine, and growth hormone concentrations (Table 2). Fasting plasma cortisol concentrations remained within normal limits at all times but were significantly greater in the depletion group than in the nondepletion group (24 ± 1 vs. 16 ± 3 μg/dl, P < 0.05) on the morning of day 7.

**Postexercise condition.** DEPLETION BOUT (2 H AT 65% \( \dot{V} \text{O}_{2\text{PEAK}} \)) AND SPRINTS. After the 2-h depletion exercise, plasma lactate, free fatty acids, epinephrine, norepinephrine, glucagon, cortisol, and growth hormone had increased significantly, whereas insulin was significantly reduced (Tables 1 and 2). Blood glucose concentrations also decreased from 89 ± 2 to 81 ± 3 mg/dl after 2 h of cycle exercise but returned to preexercise values (91 ± 7 mg/dl) immediately after the last sprint. After 2 h of cycle exercise, lactate levels increased from 0.9 ± 0.2 to 1.8 ± 0.2 mM, but after the last sprint (volitional exhaustion), they had increased (P < 0.01) to 6.5 ± 0.6 mM. Plasma norepinephrine and cortisol also increased significantly (P < 0.05) after the last sprint. Plasma glucose, free fatty acids, triglycerides, insulin, glucagon, growth hormone, and epinephrine were unchanged after the last sprint (Tables 1 and 2).

**MODERATE EXERCISE (20 MIN AT 65% \( \dot{V} \text{O}_{2\text{PEAK}} \)).** During the adaptation period (normal mixed diet), 20 min of cycle exercise had no effect on plasma glucose and triglycerides except for plasma lactate concentrations, which increased in both groups to ~2.0 mM (Table 3). Additionally, the plasma fatty acids decreased significantly in the depletion and nondepletion groups from 272 ± 19 to 194 ± 19 μM and from 224 ± 25 to 155 ± 14 μM, respectively. The hormonal response to the cycle exercise during this period did not differ between groups: insulin decreased significantly; glucagon, epinephrine, and norepinephrine increased significantly; and plasma growth hormone and cortisol were unchanged (Table 4).

The same exercise performed on day 3 of the high-CHO diet produced no change in the plasma glucose, triglycerides, or free fatty acids, but it increased

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**Table 1. Plasma metabolite concentrations in response to glycogen depletion exercise bout**

<table>
<thead>
<tr>
<th>Time</th>
<th>Glycogen, mmol/l</th>
<th>Glucose, mg/dl</th>
<th>Lactate, mM</th>
<th>FFA, μM</th>
<th>Triglycerides, mg/dl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>89 ± 4</td>
<td>89 ± 9</td>
<td>0.8 ± 0.2</td>
<td>237 ± 21</td>
<td>104 ± 9</td>
</tr>
<tr>
<td>After 2 h of exercise</td>
<td>ND</td>
<td>81 ± 3*</td>
<td>1.8 ± 0.2†</td>
<td>1,465 ± 157†</td>
<td>125 ± 10*</td>
</tr>
<tr>
<td>After 5–21 sprints</td>
<td>38 ± 6*</td>
<td>91 ± 7</td>
<td>6.5 ± 0.6†</td>
<td>1,295 ± 157†</td>
<td>121 ± 9*</td>
</tr>
</tbody>
</table>

Values are means ± SE of 15 subjects. FFA, free fatty acids. *P < 0.05 vs. baseline; †P < 0.01 vs. baseline; ‡P < 0.01 vs. after sprints.

**Table 2. Plasma hormone concentrations in response to glycogen depletion exercise bout**

<table>
<thead>
<tr>
<th>Time</th>
<th>Insulin, μU/ml</th>
<th>Glucagon, pg/ml</th>
<th>Epinephrine, pg/ml</th>
<th>Norepinephrine, pg/ml</th>
<th>Cortisol, μg/dl</th>
<th>hGH, μg/ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>9.2 ± 0.8</td>
<td>60 ± 5</td>
<td>21 ± 5</td>
<td>208 ± 29</td>
<td>22.0 ± 1.6</td>
<td>0.9 ± 0.1</td>
</tr>
<tr>
<td>After 2 h of exercise</td>
<td>5.3 ± 0.8**</td>
<td>126 ± 17**</td>
<td>217 ± 34**</td>
<td>857 ± 120*</td>
<td>32.0 ± 2.1*</td>
<td>8.8 ± 1.0*</td>
</tr>
<tr>
<td>After 5–21 sprints</td>
<td>4.5 ± 0.7**</td>
<td>119 ± 16**</td>
<td>336 ± 83**</td>
<td>1,372 ± 159†</td>
<td>39.0 ± 1.9†</td>
<td>12.7 ± 1.4†</td>
</tr>
</tbody>
</table>

Values are means ± SE of 15 subjects. hGH, human growth hormone. *P < 0.05 vs. baseline; †P < 0.01 vs. baseline; ‡P < 0.01 vs. after sprints.
plasma lactate significantly to $\sim 4.0 \pm 0.5$ mM in both groups. Plasma lactate returned to preexercise values (0.7–2.4 mM) at 60 min postexercise for both groups under both dietary conditions. As expected, this relatively short exercise again produced no significant increase in plasma cortisol or growth hormone concentrations for either group. Plasma glucagon showed a nonsignificant increase in both groups. However, plasma epinephrine increased significantly only in the depletion group, and norepinephrine increased significantly in both groups (Table 4).

**DISCUSSION**

The classical CHO-loading protocol (3, 20, 35) used by endurance athletes in the 1960s and 1970s has been largely replaced by less demanding modified protocols (24, 31, 35). Modified protocols include a training taper and may or may not begin with a depletion bout of exercise. Using $^{13}$C MRS to follow the time course of muscle glycogen supercompensation, we demonstrated that moderately trained males achieve greater muscle glycogen concentration and maintain it longer when exhaustive cycle exercise was performed and the group that performed only 20 min of cycle exercise (decreasing muscle glycogen to 90% of baseline) and ate the same high-CHO diet showed no change in muscle glycogen at 24 h. Muscle glycogen concentration of the nondepletion group peaked at 72 h (124% of baseline) and was not different from baseline on day 7.

These results agree with earlier findings (2, 31, 38) that muscle glycogen depletion (i.e., exhaustive exercise) affects the initial rate of muscle glycogen synthesis and the level of repletion (4, 7).

Several studies have suggested that when muscle glycogen is severely depleted, glycogen resynthesis is markedly activated (6, 16, 29). Muscle biopsy studies (6, 16) have found that when glycogen concentration in the vastus lateralis is decreased to 66–70 mmol/kg wet wt, glycogen synthase activity increases rapidly (6, 16).

More recent studies using MRS reported an increase in the resynthesis rate when muscle glycogen was decreased to 30–40 mmol/l or 25% of baseline (29). These findings agree with the rapid glycogen synthesis observed in our depletion group, whose mean postexercise muscle glycogen was 38 $\pm$ 6 mmol/l.

Glycogen synthase activity is increased by conversion of its D form to the active I form (8, 22). Glycogen repletion after exercise is biphasic (21) and is controlled by the rates of glucose transport and disposal. During the early rapid phase (0–6 h postexercise), glucose transport across the muscle membrane is maximally stimulated and insulin independent (29). Orig-

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**Table 3. Plasma metabolite concentrations in response to 20 min of cycle exercise at 65% $\dot{V}O_{2peak}$ during CHO loading**

<table>
<thead>
<tr>
<th></th>
<th>Glucose, mg/dl</th>
<th>Lactate, mM</th>
<th>FFA, $\mu$M</th>
<th>Triglycerides, mg/dl</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasted baseline</td>
<td>84.4 $\pm$ 2.4</td>
<td>1.2 $\pm$ 0.1</td>
<td>261 $\pm$ 25</td>
<td>111 $\pm$ 15</td>
</tr>
<tr>
<td>Preexercise</td>
<td>86.8 $\pm$ 4.7</td>
<td>2.0 $\pm$ 0.2</td>
<td>152 $\pm$ 14*</td>
<td>130 $\pm$ 14</td>
</tr>
<tr>
<td>Postexercise</td>
<td>71.4 $\pm$ 7.1</td>
<td>4.2 $\pm$ 0.6†</td>
<td>165 $\pm$ 11*</td>
<td>168 $\pm$ 18</td>
</tr>
<tr>
<td>ND</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasted baseline</td>
<td>89.4 $\pm$ 2.3</td>
<td>1.1 $\pm$ 0.1</td>
<td>235 $\pm$ 19</td>
<td>145 $\pm$ 24</td>
</tr>
<tr>
<td>Preexercise</td>
<td>101.4 $\pm$ 9.5</td>
<td>2.2 $\pm$ 0.2</td>
<td>174 $\pm$ 18*</td>
<td>156 $\pm$ 25</td>
</tr>
<tr>
<td>Postexercise</td>
<td>81.6 $\pm$ 2.6</td>
<td>4.0 $\pm$ 0.5†</td>
<td>152 $\pm$ 23*</td>
<td>179 $\pm$ 26</td>
</tr>
</tbody>
</table>

Values are means $\pm$ SE of 15 subjects in depletion (D) protocol and 10 subjects in nondepletion (ND) protocol. $\dot{V}O_{2peak}$, peak oxygen uptake; CHO, carbohydrate. *$P < 0.05$, †$P < 0.01$ vs. preexercise.

**Table 4. Plasma hormone concentrations in response to 20 min of cycle exercise at 65% $\dot{V}O_{2peak}$ during CHO loading**

<table>
<thead>
<tr>
<th></th>
<th>Insulin, µU/ml</th>
<th>Glucagon, pg/ml</th>
<th>Epinephrine, pg/ml</th>
<th>Norepinephrine, pg/ml</th>
<th>Cortisol, µg/dl</th>
</tr>
</thead>
<tbody>
<tr>
<td>D</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasted baseline</td>
<td>10.2 $\pm$ 1.2</td>
<td>54.7 $\pm$ 4.6</td>
<td>27 $\pm$ 5.4</td>
<td>314 $\pm$ 48</td>
<td>19.9 $\pm$ 1.8</td>
</tr>
<tr>
<td>Preexercise</td>
<td>58.2 $\pm$ 12.1†</td>
<td>58.0 $\pm$ 5.4</td>
<td>19 $\pm$ 2.4</td>
<td>316 $\pm$ 55</td>
<td>16.5 $\pm$ 1.0</td>
</tr>
<tr>
<td>Postexercise</td>
<td>14.3 $\pm$ 2.1‡</td>
<td>78.1 $\pm$ 7.2</td>
<td>60 $\pm$ 9.6‡</td>
<td>838 $\pm$ 140‡</td>
<td>16.1 $\pm$ 0.6*</td>
</tr>
<tr>
<td>ND</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fasted baseline</td>
<td>8.5 $\pm$ 0.6</td>
<td>49.4 $\pm$ 4.7</td>
<td>46 $\pm$ 3.5</td>
<td>287 $\pm$ 56</td>
<td>17.6 $\pm$ 2.5</td>
</tr>
<tr>
<td>Preexercise</td>
<td>76.0 $\pm$ 18.3†</td>
<td>52.6 $\pm$ 4.6</td>
<td>43 $\pm$ 5.9</td>
<td>283 $\pm$ 42</td>
<td>15.5 $\pm$ 2.1</td>
</tr>
<tr>
<td>Postexercise</td>
<td>14.4 $\pm$ 2.7†</td>
<td>78.2 $\pm$ 8.5</td>
<td>49 $\pm$ 6.5</td>
<td>687 $\pm$ 234</td>
<td>17.7 $\pm$ 2.3</td>
</tr>
</tbody>
</table>

Values are means $\pm$ SE of 15 D group and 10 ND group subjects. *$P < 0.05$ vs. baseline; †$P < 0.01$ vs. baseline; ‡$P < 0.01$ vs. preexercise.
originally attributed to a "local factor" present in the muscle after exercise (3), it is now known to result from the translocation of an intracellular pool of the GLUT4 isoform of glucose transporter proteins (9), possibly secondary to activation of AMP-activated protein kinase (1, 37). Ren et al. (30) reported a rapid increase in the number of GLUT4 glucose transport receptors in rats in response to prolonged exercise. Kua et al. (23) reported that this increase in GLUT4 protein is controlled by both pretranslational and posttranslational mechanisms. This may explain why the depletion group achieved and maintained significantly greater muscle glycogen than the nondepletion group. When provided sufficient glucose, muscle glycogen synthesis continues during the slow phase (6–72 h) so that pre-exercise levels of glycogen can be reached by 24 h. Muscle glycogen can exceed normal levels by 72 h if a high-CHO diet is consumed and exercise is limited. It is well established that, during the slow phase, muscle glycogen can reach 1.5–2.0 times resting levels (16, 33); however, only a few studies have previously monitored muscle glycogen content longer than 72 h of CHO loading (11, 24).

Our second finding has practical applications for competitive endurance athletes who may prefer exercise to rest while CHO loading. Previous research suggests there may be trade-offs associated with continuing training while attempting to achieve and maintain glycogen supercompensation. For example, because the rate of muscle glycogenolysis is most rapid during the early minutes of exercise, even 20 min of moderate-intensity (60–75% \( V_{\text{O}_2 \text{peak}} \)) exercise can significantly decrease muscle glycogen by 30–58% (1, 14). Rapid rates of glycogenolysis also occur with high exercise intensity and high initial muscle glycogen concentrations. In a study comparing two modified CHO-loading protocols (5), the muscle glycogen of well-trained endurance runners did not differ when they performed 40 min of "easy" daily running instead of resting while CHO loading. Because these researchers did not report the intensity of the exercise or the amount or timing of the postexercise CHO intake, it is difficult to directly compare our findings.

In our study, subjects used only 10–15% of their muscle glycogen during 20 min of daily cycle exercise at 65% \( V_{\text{O}_2 \text{peak}} \). The rate of glycogen resynthesis is maximal during the first 1–2 h postexercise (29), and maximum resynthesis occurs when 1.5 g glucose/kg body wt is consumed during this period (17). By consuming a CHO supplement (1.4 g glucose/kg body wt) immediately after exercise, within 24 h our subjects had replaced all of the muscle glycogen used during daily exercise. This demonstrates that an athlete can exercise daily during CHO loading without negatively affecting muscle glycogen supercompensation.

Metabolic and hormonal responses of our subjects were typical for the exercise stress and dietary conditions. The 2-h depletion exercise significantly increased plasma levels of fatty acids and triglycerides consistent with increased lipolysis. Elevated plasma fatty acid levels at 2 h typically occur with endurance exercise combined with glycogen sparing and the absence of lactate accumulation (32). Plasma lactate levels <2 mM at 2 h indicate that subjects exercising at 65% \( V_{\text{O}_2 \text{peak}} \) were below their "lactate threshold." However, the postexercise plasma lactate concentrations indicate a significant involvement of anaerobic glycolysis. During the first 2 h of cycle exercise, insulin decreased and glucagon increased significantly to maintain glucose availability and was unchanged after the sprints. The hormonal responses of our subjects (e.g., increased plasma epinephrine, norepinephrine, cortisol, and growth hormone) after cycling 2 h at 65% \( V_{\text{O}_2 \text{peak}} \) and after the series of intense sprints are consistent with changes typically seen after prolonged exhaustive exercise (32). There were no significant differences between treatment groups in insulin sensitivity as estimated by homeostasis model assessment-estimated insulin resistance at any point in the study.

CHO-loading studies generally do not measure muscle glycogen in every subject each day, nor do they study the maintenance of supercompensated muscle glycogen. We previously reported that supercompensated muscle glycogen can persist for ≥3 days after completion of a classical CHO-loading protocol, if subjects abstain from exercise (11). The current study is an extension of that work and profiled muscle glycogen during and after subjects completed two modified CHO-loading protocols, which included 20 min of daily cycle exercise.

The potential military application of CHO loading involves possible conditions that would not occur in sports. The most obvious is that missions can be delayed for a variety of reasons and the CHO-loaded personnel may be required to wait several days before deploying. Athletic endurance events, however, are postponed for only minutes, not days. Thus, unlike the sports athlete, military special operations personnel may need to maintain glycogen supercompensation and avoid physical detraining during the period of postponement. The findings of this study may, however, have application to the recreational athlete who could benefit by knowing the time profile of peak muscle glycogen and the effect of daily exercises on supercompensated muscle glycogen.

In conclusion, we have demonstrated that modified CHO-loading protocols that begin with an exhaustive depletion exercise achieve greater muscle glycogen concentrations that will persist longer than nondepletion protocols (e.g., training taper). These findings agree with earlier studies. However, equally important, we demonstrated that 20 min of moderate cycle exercise can be performed daily during all phases of CHO loading. These exercise bouts do not alter the profile muscle glycogen concentrations if a CHO supplement (~1.4 g CHO/kg) is consumed within 30 min after exercise. This suggests that endurance athletes can continue moderate daily exercise while CHO loading and still achieve and maintain muscle glycogen supercompensation.
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DISCLOSURES

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REFERENCES


