Combined ingestion of protein and carbohydrate improves protein balance during ultra-endurance exercise


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Koopman, René, Daphne L. E. Pannemans, Asker E. Jeukendrup, Annemie P. Gijsen, Joan M. G. Senden, David Halliday, Wim H. M. Saris, Luc J. C. van Loon, and Anton J. M. Wagenmakers. Combined ingestion of protein and carbohydrate improves protein balance during ultra-endurance exercise. Am J Physiol Endocrinol Metab 287: E712–E720, 2004. First published May 27, 2004; 10.1152/ajpendo.00543.2003.—The aims of this study were to compare different tracer methods to assess whole body protein turnover during 6 h of prolonged endurance exercise when carbohydrate was ingested throughout the entire exercise period and to investigate whether addition of protein can improve protein balance. Eight endurance-trained athletes were studied on two different occasions at rest (4 h), during 6 h of exercise at 50% of maximal O2 uptake (in sequential order: 2.5 h of cycling, 1 h of running, and 2.5 h of cycling), and during subsequent recovery (4 h). Subjects ingested carbohydrate (CHO during 6 h of exercise at 50% of maximal O2 uptake (in sequential order: 2.5 h of cycling, 1 h of running, and 2.5 h of cycling), and during subsequent recovery (4 h). Subjects ingested carbohydrate (CHO trial; 0.7 g CHO·kg−1·h−1) or carbohydrate/protein beverages (CHO + PRO trial; 0.7 g CHO·kg−1·h−1 and 0.25 g PRO·kg−1·h−1) at 30-min intervals during the entire study. Whole body protein metabolism was determined by infusion of I-[1-13C]leucine, I-[2H5]phenylalanine, and [15N2]urea tracers with sampling of blood and expired breath. Leucine oxidation increased from rest to exercise (27 ± 2.5 vs. 74 ± 8.8 (CHO) and 85 ± 9.5 vs. 200 ± 16.3 mg protein·kg−1·h−1 (CHO + PRO), P < 0.05), whereas phenylalanine oxidation and urea production did not increase with exercise. Whole body protein balance during exercise with carbohydrate ingestion was negative (−74 ± 8.8, −17 ± 1.1, and −72 ± 5.7 mg protein·kg−1·h−1) when I-[1-13C]leucine, I-[2H5]phenylalanine, and [15N2]urea, respectively, were used as tracers. Addition of protein to the carbohydrate drinks resulted in a positive or less-negative protein balance (−32 ± 16.3, 165 ± 4.6, and 151 ± 13.4 mg protein·kg−1·h−1) when I-[1-13C]leucine, I-[2H5]phenylalanine, and [15N2]urea, respectively, were used as tracers. We conclude that, even during 6 h of exhaustive exercise in trained athletes using carbohydrate supplements, net protein oxidation does not increase compared with the resting state and/or postexercise recovery. Combined ingestion of protein and carbohydrate improves net protein balance at rest as well as during exercise and postexercise recovery.

protein metabolism; dietary supplements; protein intake

IN THE NINETEENTH CENTURY, skeletal muscle protein was thought to be the main fuel used to generate energy for muscle contractions (33). However, controlled nitrogen balance studies invalidated the proposed hypothesis, inasmuch as they showed no substantial increase in nitrogen loss during and/or after prolonged exercise (9, 14). Since then, various methods have been applied to investigate the effects of exercise on protein metabolism. This has resulted in many discrepant findings in the literature, which are due in part to differences in the methodology employed and in part to the conditions under which different exercise interventions have been performed.

Since the introduction of stable isotope tracers in metabolic research, various methods have been applied to study whole body protein metabolism. Most studies have used plasma I-[1-13C]leucine kinetics as a model for whole body protein metabolism. Whereas several studies have reported increases in whole body protein degradation during exercise (23, 25, 39, 40), others have failed to observe such changes (6, 26). Wolfe et al. (39) measured the rate of appearance of 13CO2 in the expired breath from infused 13C-labeled leucine and reported a threefold increase in leucine oxidation rates during exercise. The latter was shown to occur in the absence of a change in total leucine flux, which implies that the observed increase in protein breakdown was accompanied by a reduction in the rate of protein synthesis. Another tracer method that has been developed to determine whole body protein metabolism, without the necessity of analyzing breath gases, is the use of I-[2H5]phenylalanine according to the model of Clarke and Bier (10) and Thompson et al. (27). However, that method has not been applied to investigate the effect of prolonged endurance exercise. Studies applying [15N]urea to determine the rate of urea production as a measure of the amount of amino acids that are liberated by net protein degradation (degradation synthesis) and oxidized (with conversion of the amino group to urea) during exercise have not confirmed the catabolic nature of prolonged endurance exercise (7, 39).

The reports using urea as a tracer indicate that protein synthesis and degradation rates do not change during endurance-type activities. However, nitrogen balance data from field studies suggest that prolonged endurance exercise, leading to glycogen depletion, is accompanied by a substantial increase in net protein breakdown (13). Therefore, activation of the branched-chain α-keto acid dehydrogenase complex, the enzyme responsible for leucine oxidation in the muscle, has been shown to increase during exercise when glycogen stores are low (34, 35). During competition, many endurance athletes (e.g., cyclists, triathletes, and ultra-marathon runners) exercise for >5 h at relatively high workloads and deplete their glycogen stores to a large extent. However, whole body protein turnover measurements using multiple amino acid tracers have not been performed during such prolonged exhaustive exercise activity.
Endurance-trained athletes try to optimize carbohydrate availability through carbohydrate ingestion before, during, and after exercise. However, most studies measuring protein turnover during exercise have been performed after an overnight fast and with nutritional interventions generally applied during the postexercise recovery phase. Little information is available about the effects of carbohydrate or carbohydrate + protein supplementation on protein metabolism during exercise. The available literature indicates that ingestion of carbohydrate + protein in the postexercise recovery phase improves net protein balance (12, 22), at least after resistance exercise (2, 3, 24, 28, 29). Ingestion of such a substrate mixture during exercise could possibly also affect protein balance during and after prolonged endurance exercise.

In the present study, we determined whole body protein synthesis and degradation rates at rest, during prolonged exhaustive exercise, and during subsequent recovery. A multiple-tracer approach applying [1-13C]leucine, [1-3H3]phenylalanine, and [15N2]urea infusions was chosen to investigate whether there are differences in outcome, depending on the applied tracer model. These studies were performed in a laboratory setting that resembles real-life endurance exercise competition in elite triathletes, combining 6 h of exercise with regular carbohydrate supplementation. In addition, we investigated whether the addition of protein to carbohydrate ingestion can improve net protein balance during exercise and subsequent recovery compared with the ingestion of only carbohydrate.

METHODS

Subjects. Eight well-trained male subjects [age = 31 ± 3 yr, height = 1.84 ± 0.03 m, weight = 72.4 ± 2.3 kg, body mass index = 21.4 ± 0.3 kg/m², %body fat = 9.6 ± 0.8%, fat-free mass = 65.7 ± 1.3 kg, maximal workload capacity (Wmax) = 383 ± 13 W, maximal O2 uptake (V̇O2max) = 4.9 ± 0.2 l/min] participated in the study. Subjects trained ~15 h/wk (3 h of swimming, 7 h of cycling, and 5 h of running) and had a >5-yr training history. Subjects were informed about the nature and risks of the procedures before their written informed consent was obtained. The study was approved by the local medical ethical committee of the Academic Hospital Maastricht.

Pretesting. V̇O2max and Wmax were measured on an electronically braked cycle ergometer (Lode Excalibur, Groningen, The Netherlands) during an incremental exhaustive exercise test (17) 1 wk before the first trial to determine 45% Wmax (179 ± 8.6 W, ~50% V̇O2max), which was applied in these studies. Maximum running velocity was determined on a motor-driven treadmill with a 1% incline (to represent outdoor running) using an incremental exhaustive exercise test to determine the 45% maximal running velocity (11.0 ± 0.3 km/h, ~50% V̇O2max). After a 5-min warm-up at 10 km/h, speed was increased every 3 min by 2 km/h and from a heart rate of 85% of maximal heart rate with 1 km/h every 3 min until the subject was unable to continue.

Diet and activity before testing. All subjects were instructed not to consume any products with a high natural abundance of 13C (carbohydrates derived from C₄ plants: corn and sugar cane) 1 wk before the first trial and during the entire test period. This has been shown to minimize possible shifts in background enrichment due to changes in endogenous substrate utilization (36). All subjects were also instructed to refrain from heavy physical exercise and to maintain normal dietary habits for 3 days before each trial. In addition, we asked subjects to record their food intake for 48 h before the start of the first experimental trial and to consume exactly the same food 48 h before the start of the second test.

Studies. All subjects were studied on two different occasions. Each test day consisted of three parts, during which whole body protein turnover and whole body urea production rates were measured at rest, during prolonged exercise, and during subsequent recovery. Via a randomized crossover design, subjects received beverages containing carbohydrate (CHO) or carbohydrate with protein hydrolysate (CHO + PRO) during each trial. An outline of the study design is presented in Fig. 1.

Protocol. The subjects reported to the laboratory at 7 AM after an overnight fast. A Teflon catheter (Baxter, Utrecht, The Netherlands) was inserted into an antecubital vein of one arm for blood sampling; another catheter was inserted in the contralateral arm for isotope infusion. A resting blood sample was drawn, and expired breath samples were collected into Vacutainer tubes (Becton Dickinson, Meylan, France) to measure baseline enrichments. Thereafter, the bicarbonate pool was primed with a single intravenous dose of NaH13CO3 (5.9 μmol/kg), which was followed by priming with [1-15N2]urea (7.6 μmol/kg), [1-3H3]phenylalanine (3.0 μmol/kg), [15N2]urea (88 μmol/kg), and [1-13C]tyrosine (0.46 μmol/kg). Thereafter (time 0), continuous infusion of [1-15N2]leucine (7.6 μmol·kg⁻¹·h⁻¹), [1-3H3]phenylalanine (3 μmol·kg⁻¹·h⁻¹), and [15N2]urea (88 μmol·kg⁻¹·h⁻¹) was started via a calibrated pump (model 560, IVAC, San Diego, CA) and continued for 14 h. After the start of the tracer infusion, subjects rested for 4 h (resting period). Subsequently, subjects exercised for 6 h at a moderate intensity (~45% Wmax), starting with 2.5 h of cycling followed by 1 h of treadmill running and another 2.5 h of cycling exercise. Subjects were not allowed to rest between the different exercise bouts, except for the time necessary to change from ergometer to treadmill and back to ergometer. After cessation of exercise, subjects rested for 4 h (recovery). During the preexercise resting period and the postexercise recovery phase, breath and blood samples were obtained every 30 and 60 min, respectively. During cycling exercise periods, blood and breath samples were collected every 30 min, and O2 uptake and CO2 output were measured (Oxycon-øi, Mijnhardt, Bunnik, The Netherlands) at 30-min intervals for 5 min.

Beverages. Directly after the start of the stable isotope infusion and every 30 min throughout the entire trial (before, during, and after exercise), subjects received a beverage volume of 4 ml/kg to ensure a given dose of 0.7 g·kg⁻¹·h⁻¹ carbohydrates (maltodextrin) in the control trial (CHO) and 0.7 g·kg⁻¹·h⁻¹ carbohydrate and 0.25 g·kg⁻¹·h⁻¹ protein hydrolysate in the CHO + PRO trial. The drinks were prepared by Quest International (Naarden, The Netherlands). [1-3H3]phenylalanine and [1-15N2]leucine enrichment in the beverage

![Fig. 1. Schematic outline of study protocol. Every 30 min throughout the entire trial, subjects received a beverage volume of 4 ml/kg to ensure a given dose of 0.7 g·kg⁻¹·h⁻¹ carbohydrates (maltodextrin) in the control trial (CHO) and 0.7 g·kg⁻¹·h⁻¹ carbohydrate and 0.25 g·kg⁻¹·h⁻¹ protein hydrolysate in the CHO + PRO trial. V̇O2max, maximal O2 uptake.](http://ajpendo.physiology.org/doi/abs/10.220.32.247)
Breath 13 CO2 enrichment was significant in contrast to the CHO trial, with significant differences in protein synthesis, breakdown, oxidation, net balance, and whole body protein synthesis between the different interventions over time. A two-way repeated-measures ANOVA was used with subject and treatment as factors. Scheffe's post hoc test was applied in case of a significant F-ratio to locate specific differences. Student's t-tests for paired observations were used to compare differences in protein synthesis, breakdown, oxidation, net balance, and urea production between the CHO and PRO trial. Significance was set at the 0.05 level of confidence.

**RESULTS**

**Tracer kinetics.** Plasma t-[^2]H5phenylalanine and t-[^2]H3tyrosine enrichment, plasma [1-13C]α-KIC enrichment, breath CO2 enrichment, and plasma [15N2]urea enrichment are shown in Fig. 2. Plasma t-[^2]H5phenylalanine, t-[^2]H3tyrosine, and [1-13C]α-KIC reached a steady state after 2 h of infusion. Breath CO2 enrichment was significantly lower during exercise (P < 0.05) than at rest and during postexercise recovery. Plasma [15N2]urea enrichment slightly increased over time in the CHO trial, with significantly higher plasma [15N2]urea enrichment during recovery than during preexercise rest (P < 0.05). In contrast, in the CHO + PRO trial, plasma [15N2]urea enrichment remained constant throughout the trial. In the CHO + PRO trial, plasma t-[^2]H5phenylalanine, t-[^2]H3tyrosine, [1-13C]α-KIC, and [15N2]urea enrichments were significantly lower than in the CHO trial (P < 0.05). Breath CO2 enrichment was elevated in the CHO + PRO trial compared with the CHO trial (P < 0.05).

**Phenylalanine model.** Protein flux was similar at rest, during exercise, and during postexercise recovery in the CHO trial (176 ± 6, 170 ± 6, 172 ± 7, and 162 ± 15 mg protein-kg^{-1}h^{-1}, respectively) and in the CHO + PRO trial (303 ± 26, 294 ± 14, 264 ± 14, and 293 ± 21 mg protein-kg^{-1}h^{-1}, respectively). However, in the CHO + PRO trial, protein flux was increased ~70% at rest, during exercise, and in the postexercise recovery phase compared with the CHO trial (P < 0.01).

Whole body protein synthesis (mg protein-kg^{-1}h^{-1}) was measured using the phenylalanine model and the urea isotopomer model. No differences were found in protein synthesis rates during exercise compared with rest and the recovery phase. However, under all conditions, protein synthesis rates were higher in the CHO + PRO trial than in the CHO trial (P < 0.01).

Whole body protein degradation rates (mg protein-kg^{-1}h^{-1}) did not change during exercise (Fig. 3B). A marked decrease in protein breakdown was observed throughout the CHO + PRO trial compared with the CHO trial (P < 0.01).

During exercise, whole body protein balance (mg protein-kg^{-1}h^{-1}) was not different from resting conditions and postexercise recovery (Fig. 5). Protein balance was positive in the CHO + PRO trial and negative in the CHO trial.

**Leucine model.** Protein flux (mg protein-kg^{-1}h^{-1}) was not different during exercise compared with rest and during postexercise recovery. Protein flux was ~50% higher at rest, during exercise, and in the postexercise recovery phase (P < 0.01) in the CHO + PRO trial than in the CHO trial.

Whole body protein synthesis (mg protein-kg^{-1}h^{-1}) was shown in Fig. 4. Whole body protein oxidation was similar at rest, during exercise, and during postexercise recovery in the CHO trial (13 ± 1, 17 ± 1, 19 ± 2, and 15 ± 2 mg protein-kg^{-1}h^{-1}, respectively) and in the CHO + PRO trial (31 ± 4, 30 ± 3, 25 ± 2, and 27 ± 3 mg protein-kg^{-1}h^{-1}, respectively). In the CHO + PRO trial, protein oxidation rates were higher at rest, during exercise, and during recovery than in the CHO trial (P < 0.05).

**Table 1. Amino acid composition of the protein hydrolysate**

<table>
<thead>
<tr>
<th>Amino Acids (Arg)</th>
<th>Rice Hydrolysate</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-Arginine</td>
<td>5.6</td>
</tr>
<tr>
<td>L-Histidine</td>
<td>1.5</td>
</tr>
<tr>
<td>L-Asparagine</td>
<td>4.1</td>
</tr>
<tr>
<td>L-Glutamine (Gln)</td>
<td>7.7</td>
</tr>
<tr>
<td>L-Lysine</td>
<td>2.8</td>
</tr>
<tr>
<td>L-Glycine</td>
<td>2.7</td>
</tr>
<tr>
<td>L-Alanine</td>
<td>3.5</td>
</tr>
<tr>
<td>L-Tryptophan</td>
<td>0.6</td>
</tr>
<tr>
<td>L-Serine</td>
<td>3.4</td>
</tr>
<tr>
<td>L-Proline</td>
<td>2.4</td>
</tr>
<tr>
<td>L-Valine</td>
<td>3.9</td>
</tr>
<tr>
<td>L-Threonine</td>
<td>2.4</td>
</tr>
<tr>
<td>L-Cysteine</td>
<td>0.6</td>
</tr>
<tr>
<td>L-Isoleucine</td>
<td>2.8</td>
</tr>
<tr>
<td>L-Leucine</td>
<td>5.2</td>
</tr>
<tr>
<td>L-Aspartic Acid</td>
<td>2.4</td>
</tr>
<tr>
<td>L-Glutamate</td>
<td>4.5</td>
</tr>
<tr>
<td>L-Methionine</td>
<td>1.4</td>
</tr>
<tr>
<td>L-Phenylalanine</td>
<td>3.6</td>
</tr>
<tr>
<td>L-Tyrosine</td>
<td>3.3</td>
</tr>
</tbody>
</table>

Values are expressed in g/100 g dry product.
during exercise, and during recovery compared with the CHO trial ($P < 0.05$).

In the CHO trial, whole body protein balance (mg protein·kg$^{-1}$·h$^{-1}$) was negative at rest, during exercise, and during subsequent recovery. Protein balance was more negative during exercise than at rest and during recovery ($P < 0.01$; Fig. 5). In the CHO + PRO trial, protein balance was less negative at rest, during exercise, and during postexercise recovery than in the CHO trial ($P < 0.01$); however, protein balance during exercise remained negative.

Phenylalanine model vs. leucine model. The relative increase in protein flux in the CHO + PRO trial compared with the CHO trial was similar for both applied tracers: $+70 \pm 14\%$ for L-[$^3$H]phenylalanine and $+48 \pm 12\%$ for L-$[1-^{13}C]$leucine.

Protein synthesis was increased $69 \pm 14\%$ in the CHO + PRO trial compared with the CHO trial when phenylalanine was used as a tracer. The leucine model showed no differences in protein synthesis rates between studies.

The relative decrease in protein degradation in the CHO + PRO trial compared with the CHO trial was similar for both

Protein oxidation was increased with both tracer models in the CHO PRO trial compared with the CHO trial. The increase in protein oxidation in the CHO PRO trial compared with the CHO trial was higher when measured with L-[1-13C]leucine than with L-[2H5]phenylalanine: 190 ± 30% vs. 82 ± 22% \((P < 0.05)\).

Protein balance was improved with both tracer models in the CHO PRO trial compared with the CHO trial \((P < 0.05)\). The increase in protein balance in the CHO PRO trial compared with the CHO trial was higher when measured with L-[2H5]phenylalanine than with L-[1-13C]leucine \((P < 0.01)\).

**Urea model and comparison with other tracers.** Urea kinetics were calculated using a \([^{15}N_2]\)urea tracer, and results are illustrated in Fig. 6. In the CHO trial, urea production decreased during exercise and recovery compared with resting values, whereas in the CHO + PRO trial, urea production remained constant. Urea production was higher in the CHO + PRO trial than in the CHO trial \((P < 0.05)\).

Net protein balance calculated from urea production rates was less negative during exercise and recovery than at rest in the CHO trial \((P < 0.05)\). No changes in net protein balance were observed in the CHO PRO trial. Calculated net protein balance was positive in the CHO PRO trial, whereas in the CHO trial net protein balance was negative.

In the CHO trial, whole body protein oxidation did not change in time with L-[2H5]phenylalanine applied as a tracer, increased during exercise with L-[1-13C]leucine used as a tracer to return to resting values in the recovery period, and gradually decreased in time with \([^{15}N_2]\)urea as tracer. In the CHO PRO trial, whole body protein oxidation did not change in time with L-[2H5]phenylalanine and \([^{15}N_2]\)urea as tracers and increased during exercise with L-[1-13C]leucine as tracer to return to resting values in the recovery period.

Protein balance was improved with all tracer models applied in the CHO PRO trial compared with the CHO trial. The phenylalanine model and the urea model indicate that the net protein balance is positive in the CHO PRO trial at rest, during exercise, and during recovery.

**DISCUSSION**

In the present study, we show that whole body estimates of protein metabolism during exercise are highly dependent on the specific method and tracer that are applied. When L-[1-13C]leucine was used as a tracer to study the effects of exercise on protein metabolism, protein oxidation was increased two- to threefold during exercise. However, when L-[2H5]phenylalanine was used as a tracer, protein oxidation, synthesis, and degradation rates were not affected by exercise. In addition, urea production, measured using \([^{15}N_2]\)urea, decreased in time in the CHO trial. Thus two of the three methods
indicate that prolonged exhaustive exercise does not lead to increased protein oxidation and/or a more negative protein balance. Although the application of different tracer methods results in different effects of exercise on protein metabolism, each method shows that protein balance is negative when only carbohydrates are ingested. The addition of protein to carbohydrate ingestion improves whole body protein balance when measured using the phenylalanine and leucine models used in the present study.

Urea production in subjects not ingesting any protein is generally considered a reflection of the amount of amino acids that are liberated by net protein degradation (degradation – synthesis) and subsequently oxidized with conversion of the amino group to urea. Studies on the effect of exercise on urea production have consistently found no significant change from the basal rate (7, 39, 40), leading to the conclusion that the rate of net protein catabolism (degradation – synthesis) and amino acid oxidation was not changed. However, data obtained using another approach to determine whole body protein turnover, involving the infusion of \(^1\text{H}\)phenylalanine and the determination of leucine turnover and \(^{13}\text{CO}_2\) appearance in the expired air, indicated that leucine oxidation was markedly increased during exercise (4, 5, 39, 40). Because of this discrepancy between methods, we applied both tracer methods in the same study. In addition, we also applied a third tracer, \(^1\text{H}^2\)phenylalanine, which has been successfully applied at rest (10, 27), but not during exercise conditions. In contrast to most studies, we used an experimental setup that included exercise and the continuous ingestion of carbohydrate (CHO) or carbohydrate and protein supplements (CHO + PRO), inasmuch as athletes always ingest carbohydrates during exercise (training or competition) of this duration and intensity. With leucine as tracer, protein oxidation was two- to threefold higher during exercise than at rest or during recovery (Fig. 4). The widely used \(^1\text{H}^2\)phenylalanine model for the calculation of whole body protein synthesis and breakdown, therefore, suggests that exercise inhibits protein synthesis (Fig. 3A). However, with phenylalanine used as tracer, protein oxidation, synthesis, and degradation were similar at rest, during exercise, and during recovery (Figs. 4, 3A, and 3B, respectively). Moreover, urea production and the amount of protein oxidation calculated from it also did not increase during exercise (Fig. 6). Thus, the phenylalanine model and urea data show that prolonged moderate-intensity exercise does not lead to an imbalance between protein synthesis and degradation and subsequent oxidation of the liberated amino acids. This is consistent with tracer incorporation studies (8, 30), which reported no change in muscle protein synthesis during moderate endurance exercise and recovery. However, results from the phenylalanine and leucine models in the present study show that whole body protein balance is negative at rest, during exercise, and during postexercise recovery when only carbohydrates are ingested (Fig. 5).

A severalfold increase in leucine oxidation without changes in urea production during exercise has been previously observed (39, 40). Moreover, other studies also showed no effect of exercise on plasma urea concentration (26), production (7), or urinary excretion (6, 18, 23). An explanation for the discrepancy between the leucine and urea data is that leucine metabolism during exercise is not representative of the fate of the other amino acids that are present in proteins. The stochastic leucine model, therefore, cannot be used to investigate whole body protein metabolism during exercise. A difference in time response of leucine and urea kinetics theoretically also leads to an apparent absence of an increase in urea production during high rates of leucine oxidation. The urea pool is large and turns over with a half-life of 8–10 h (16, 19, 37). This implies that any increase in urea production due to an accelerated protein oxidation rate during exercise should have become apparent during or after the 6-h exercise period. However, in the CHO trial, we observed a gradual decrease, instead of an increase, in urea production over time. Such a decrease in urea production in time has previously been reported in studies at rest after an overnight fast (for references see Ref. 37). The latter can be attributed to the delayed response of the urea production after changes in protein intake (16, 37).

In the present study, we also investigated phenylalanine metabolism. We determined the rate of conversion (hydroxylation) of \(^1\text{H}^2\)phenylalanine to \(^1\text{H}^2\)tyrosine at rest, during exercise, and during recovery. This is the first step of phenylalanine oxidation, and we found that its rate did not change during exercise and/or recovery compared with the resting state. This is consistent with findings reported by Wolfe et al. (40) using a lysine tracer. In the latter study, the increase in lysine oxidation during exercise was small compared with the increase in leucine oxidation. In addition, a discrepancy existed between the \(^{15}\text{N}\)- and \(^{13}\text{C}\)lysine flux data and the \(^{13}\text{C}\)- and \(^{15}\text{N}\)leucine flux data that was similar to the discrepancy between phenylalanine and leucine observed in the present study.

One of the likely causes of the discrepancy between the lysine and phenylalanine tracer and the leucine tracer methods is a severalfold increase in uptake (1, 32) and oxidation (32, 34, 35) of branched-chain amino acids (leucine, isoleucine, and valine) during exercise in contracting muscle. As such, the leucine tracer does not seem to be representative of the fate of the other essential amino acids during exercise. The latter means that the stochastic leucine model of whole body protein metabolism (20) becomes invalid during exercise and cannot be used to estimate qualitative or quantitative changes in whole body protein synthesis and/or degradation.

Most studies measuring protein turnover during exercise were performed in the morning after an overnight fast, which does not represent everyday practice in competitive athletes. Athletes optimize carbohydrate availability before and during exercise to enhance endurance performance capacity. During subsequent recovery, they ingest mixed meals or carbohydrate/protein recovery drinks. The present study demonstrates that net protein degradation does not increase during moderate-intensity exercise when carbohydrate is ingested. This implies that net protein breakdown and oxidation are minimal during athletic events such as (ultra)marathons and prolonged cycling races when carbohydrate is ingested. Although protein degradation and synthesis are unaltered by exercise when carbohydrate (0.7 g/min) is ingested, whole body protein balance measured with \(^1\text{H}^2\)phenylalanine was slightly negative during exercise and during recovery in the present study. However, the imbalance between protein synthesis and degradation and the amount of protein oxidized was not larger than in the resting state (Fig. 5).

It has previously been shown that infusion or ingestion of amino acids alone or in combination with carbohydrate increases protein synthesis and reduces protein degradation at a
whole body level and in skeletal muscle tissue under resting conditions (12, 15, 22) and after resistance exercise (2, 3, 24, 28, 29), resulting in a positive protein balance. We aimed to investigate whether the addition of protein to a carbohydrate supplement can improve whole body protein balance during and after prolonged endurance exercise leading to a (more) positive protein balance, as previously observed in the resting state. Combined ingestion of protein and carbohydrate resulted in an increased whole body protein synthesis rate at rest, during exercise, and during recovery compared with carbohydrate supplementation only. Although protein oxidation was increased almost twofold at rest, during exercise, and during recovery with the protein supplement, whole body protein degradation decreased by ~60%. Thus the present study shows that the combined ingestion of protein and carbohydrate improves net protein balance at rest, during prolonged moderate-intensity exercise, and during subsequent recovery in elite endurance athletes. With phenylalanine and urea as tracers, a positive net protein balance was achieved, whereas ingestion of only carbohydrate resulted in a negative protein balance.

The L-[2H5]phenylalanine tracer method indicates that protein flux and oxidation rates are increased as expected when protein is orally ingested. In addition, coingestion of protein and carbohydrate results in a decreased rate of protein degradation and improved protein balance. The [15N2]urea tracer provides an effect of exercise on net protein balance similar to that when L-[2H5]phenylalanine is used as tracer. Urea production slightly increases when protein is ingested (Fig. 6), because more protein is ingested than is used for protein synthesis. *Significantly different between studies (P < 0.05). †Significantly different from rest (P < 0.01). ‡Significantly different from recovery (P < 0.01).

In summary, this study shows that the L-[2H5]phenylalanine, L-[1-13C]leucine, and [15N2]urea tracer models reveal the same qualitative effect of the combined ingestion of protein and carbohydrate on whole body protein balance at rest and during recovery. However, the L-[1-13C]leucine model for measurements of whole body protein turnover overestimates protein oxidation rates and underestimates protein synthesis rates during exercise. We conclude that prolonged exhaustive moderate-intensity exercise does not result in an increase in protein degradation and/or a decrease in protein synthesis compared with resting situations in endurance-trained athletes. Whereas whole body protein balance remains negative when only carbohydrates are ingested, coingestion with protein improves protein balance by increasing synthesis and decreasing breakdown, resulting in a positive net protein balance at rest, during prolonged moderate-intensity exercise, and during recovery.

APPENDIX

Calculations. The rate of protein turnover was calculated as described previously (20). Briefly, leucine turnover (flux, \(\dot{Q}\)) was measured from the dilution of L-[1-13C]leucine infusion in plasma α-KIC at isotopic steady state

\[
\dot{Q} = i \cdot \left( \frac{E_s - 1}{E_p} \right)
\]

where i is the [1-13C]leucine infusion rate (\(\mu\text{mol} \cdot \text{kg body wt}^{-1} \cdot \text{h}^{-1}\)), \(E_s\) is the enrichment of the [1-13C]leucine infused [expressed in atom percent excess (APE)], and \(E_p\) is the [1-13C]α-KIC enrichment in the plasma at steady state (APE). The background enrichment of expired \(^1\text{CO}_2\) was measured before the start of tracer infusions. This background enrichment was subtracted from the isotopic plateau value for the calculation of leucine oxidation. Leucine oxidation rates were also corrected for bicarbonate retention. In endurance-trained subjects, bicarbonate retention factors were 83.1% for rest and 98.9% for exercise (11). The rate of leucine oxidation (O) is calculated as follows

Table 2. Protein metabolism at rest, during exercise, and during subsequent recovery

<table>
<thead>
<tr>
<th>Phenylalanine</th>
<th>Protein Oxidation, mg protein kg(^{-1}) h(^{-1})</th>
<th>Protein Synthesis, mg protein kg(^{-1}) h(^{-1})</th>
<th>Protein Degradation, mg protein kg(^{-1}) h(^{-1})</th>
<th>Protein Balance, mg protein kg(^{-1}) h(^{-1})</th>
<th>Urea Production, µmol kg(^{-1}) h(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>CHO</td>
<td>CHO + PRO</td>
<td>CHO</td>
<td>CHO + PRO</td>
<td>CHO</td>
<td>CHO + PRO</td>
</tr>
<tr>
<td>Rest</td>
<td>13±0.7</td>
<td>31±3.5*</td>
<td>163±6.0</td>
<td>272±22.3*</td>
<td>176±6.0</td>
</tr>
<tr>
<td>Exercise</td>
<td>17±1.1</td>
<td>30±3.2*</td>
<td>153±4.6</td>
<td>264±20.9*</td>
<td>170±5.7</td>
</tr>
<tr>
<td>Exercise</td>
<td>19±1.8</td>
<td>25±1.8*</td>
<td>154±5.3</td>
<td>238±13.1*</td>
<td>172±6.7</td>
</tr>
<tr>
<td>Recovery</td>
<td>15±2.1</td>
<td>27±2.8*</td>
<td>147±12.7</td>
<td>266±19.1*</td>
<td>162±14.5</td>
</tr>
<tr>
<td>Leucine</td>
<td>27±2.5</td>
<td>85±9.5*</td>
<td>178±8.1</td>
<td>206±14.5</td>
<td>205±8.1</td>
</tr>
<tr>
<td>Exercise</td>
<td>74±8.8††</td>
<td>200±16.3††</td>
<td>122±8.1††</td>
<td>102±15.6††</td>
<td>196±6.4</td>
</tr>
<tr>
<td>Exercise</td>
<td>71±6.7††</td>
<td>179±8.1††</td>
<td>125±7.1††</td>
<td>116±16.6††</td>
<td>196±4.6</td>
</tr>
<tr>
<td>Recovery</td>
<td>28±2.1</td>
<td>90±7.4*</td>
<td>164±14.1</td>
<td>187±13.4</td>
<td>191±14.8</td>
</tr>
<tr>
<td>Urea</td>
<td>Rest</td>
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<td>NA</td>
<td>NA</td>
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<tr>
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<td>NA</td>
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<td>NA</td>
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<td>Exercise</td>
<td>2</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
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</tr>
<tr>
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<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Values are means ± SE (n = 8). CHO, carbohydrate trial; CHO + PRO, carbohydrate and protein trial; NA, not applicable. *Significantly different between studies (P < 0.05). †Significantly different from rest (P < 0.01). ‡Significantly different from recovery (P < 0.01).
\[ O = F^{13}CO_2 \cdot \left[ \left( \frac{1}{E_i} \right) - \left( \frac{1}{E_p} \right) \right] \cdot 100 \]  

where \( F^{13}CO_2 \) is the rate of \(^{13}\)CO\(_2\) production and release in the expired breath (\(\mu\text{mol}^{13}\text{CO}_2\cdot\text{kg body wt}^{-1}\cdot\text{h}^{-1}\)). At isotopic steady state, protein \(Q\) equals the sum of protein synthesis (\(S\)) and \(O\) as well as the sum of the rate of appearance of meal protein from the gut (\(I\)) and protein breakdown (\(B\)); whole body protein synthesis rate was calculated as flux – oxidation

\[ Q = S + O = B + I \]  

\[ S = Q - O \]

The leucine parameters mentioned above were converted to corresponding estimates of whole body protein turnover by multiplying the leucine values by the constant \(1/(590 \text{ g protein per kilogram per hour})\). The \(590 \text{ g protein per kilogram protein} \) corresponds to the protein leucine content of 7.8% and was derived from averaged values for leucine content of protein of human and other mammalian muscles (20).

In addition, we used the phenylalanine balance model as described by Thompson et al. (27). In this model, phenylalanine \(Q\) (\(Q_p\)) can be calculated by isotope dilution using Eq. 1, where \(i\) is the \(l\-[ring-2H_5]\text{phenylalanine}\) infusion rate (\(\mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{h}^{-1}\)), \(E_i\) is the enrichment of the \(l\-[ring-2H_5]\text{phenylalanine}\) infused (expressed in APE), and \(E_p\) is the \(l\-[ring-2H_5]\text{phenylalanine}\) in the plasma at steady state (APE). At isotopic steady state, whole body phenylalanine oxidation can be determined from the conversion (hydroxylation) of \(l\-[ring-2H_5]\text{phenylalanine}\) to \(l\-[ring-2H_5]\text{tyrosine}\). The rate of hydroxylation (\(Q_{\text{pt}}\)) was calculated (27) using the formula

\[ Q_{\text{pt}} = \frac{P_i \cdot Q_p}{P_p \cdot \frac{E_i}{E_p} - 1 - (i_p - Q_p)} \]  

where \(P_i/P_p\) (0.73) is the molar ratio of tyrosine \(Q\) to \(Q_p\), arising from protein catabolism (27), \(E_i\) is the \(l\-[ring-2H_5]\text{tyrosine}\) enrichment, and \(i_p\) is the \(l\-[ring-2H_5]\text{phenylalanine}\) infusion rate (\(\mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{h}^{-1}\)). Protein synthesis-and-degradation using this approach was calculated as follows

\[ Q = S + Q_{\text{pt}} = B + I \]

Rates of protein synthesis and degradation (\(g\) protein\(\cdot\)kg body weight\(^{-1}\cdot\text{h}^{-1}\)) are calculated using the phenylalanine content of protein of 280 \(\mu\text{mol/g protein}\) (27). The 280 \(\mu\text{mol phenylalanine/g protein}\) factor was derived from protein composition measurements in animals (21).

The net protein balance (Table 2) was calculated from the difference between whole body protein synthesis and degradation determined using the leucine and phenylalanine model as described above.

The rate of urea production was calculated using equations described previously (38). Briefly, at isotopic steady state during the constant infusion of \(^{15}\text{N}_2\text{urea}\), two enrichments (expressed in tracer-to-tracer ratio) were measured, \(A\) and \(B\)

\[ A = \frac{\text{singly labeled urea}}{\text{unlabeled urea}} \]  

\[ B = \frac{\text{doubly labeled urea}}{\text{unlabeled urea}} \]

The rate of urea production was calculated using

\[ \dot{Q} = \frac{2F}{A + 2B} \]

where \(F\) is the \(^{15}\text{N}_2\text{urea}\) infusion rate (\(\mu\text{mol} \cdot \text{kg}^{-1} \cdot \text{h}^{-1}\)).

The net balance of the amount (\(g\)\(\cdot\)kg\(^{-1}\cdot\text{h}^{-1}\)) of protein that was broken down in the CHO trial and converted to urea was calculated by converting urea production rates to milligrams of nitrogen produced per kilogram per hour and multiplying by 6.25 (1 g of protein contains 160 mg of nitrogen). The positive protein net balance (protein synthesis – protein degradation) reported in Table 2 was calculated in this case by again converting the observed urea production rate to protein oxidation rates and then taking the difference between protein ingestion and protein oxidation (both in \(g\) protein\(\cdot\)kg\(^{-1}\cdot\text{h}^{-1}\)).

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