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Leucine Supplementation

Daily L-Leucine Supplementation in Novice Trainees During a 12-Week Weight Training Program

Theocharis Ispoglou, Roderick F.G.J. King, Remco C.J. Polman, and Cathy Zanker

Purpose: To investigate the effects of daily oral L-leucine ingestion on strength, bone mineral-free lean tissue mass (LTM) and fat mass (FM) of free living humans during a 12-wk resistance-training program. Methods: Twenty-six initially untrained men (n = 13 per group) ingested either 4 g/d of L-leucine (leucine group: age 28.5 ± 8.2 y, body mass index 24.9 ± 4.2 kg/m²) or a corresponding amount of lactose (placebo group: age 28.2 ± 7.3 y, body mass index 24.9 ± 4.2 kg/m²). All participants trained under supervision twice per week following a prescribed resistance training program using eight standard exercise machines. Testing took place at baseline and at the end of the supplementation period. Strength on each exercise was assessed by five repetition maximum (5-RM), and body composition was assessed by dual energy X-ray absorptiometry (DXA). Results: The leucine group demonstrated significantly higher gains in total 5-RM strength (sum of 5-RM in eight exercises) and 5-RM strength in five out of the eight exercises (P < .05). The percentage total 5-RM strength gains were 40.8% (± 7.8) and 31.0% (± 4.6) for the leucine and placebo groups respectively. Significant differences did not exist between groups in either total percentage LTM gains or total percentage FM losses (LTM: 2.9% ± 2.5 vs 2.0% ± 2.1, FM: 1.6% ± 15.6 vs 1.1% ± 7.6). Conclusion: These results suggest that 4 g/d of L-leucine supplementation may be used as a nutritional supplement to enhance strength performance during a 12-week resistance training program of initially untrained male participants.

Keywords: ergogenic aids, resistance training, muscle growth, oral ingestion

Leucine is one of the essential amino acids that belongs to the group of branched-chain amino acids (BCAA) and may play a specific and pertinent role in the regulation of protein turnover. It has been demonstrated clearly that leucine may act as nutrient signal to stimulate protein synthesis in animal studies, thus having the potential to enhance anabolism and act as an ergogenic aid. In humans,
intravenous infusion of leucine may decrease protein degradation at rest, while oral ingestion of leucine and essential amino acids may increase muscle protein synthesis at rest in both young and elderly individuals. Addition of leucine to a carbohydrate/protein supplement postexercise has been shown to elicit greater responses in the rate of muscle protein synthesis in healthy male subjects than a carbohydrate and a carbohydrate/protein supplement following a resistance exercise bout. However, others have suggested that coinigestion of leucine and whey protein before exercise does not result in a greater anabolic response than whey protein alone. In terms of sports performance, leucine supplementation has been shown to increase time to exhaustion and the upper body power of rowers during 6 wk of supplementation to a greater extent than a placebo condition, while coinigestion of protein and leucine during 8 wk of resistance training has shown to elicit further gains in one repetition maximum strength (1-RM) than a placebo (carbohydrate) and a control condition.

Bearing in mind the results of the aforementioned studies, it is probable that supplementation of leucine during resistance exercise training may be beneficial in terms of strength gains and muscle growth. Surprisingly, the capacity of orally ingested leucine alone to enhance strength and muscle mass in humans, while following an identical prescribed resistance-training program has not been tested. It has also been suggested that the recommended dietary intake of leucine of 14 mg·kg⁻¹·BW·d⁻¹ (where BW = body weight) is inadequate in individuals who are physically active. At a protein intake of 1.26 g·kg⁻¹·BW·d⁻¹, the serum concentrations of amino acids are lowered considerably during training of athletes. Leucine supplementation of 50 mg·kg⁻¹·BW·d⁻¹ appears to prevent the decrease in the serum leucine concentration during training. Therefore, a dose of similar magnitude was utilized in the present study.

The potential of leucine to stimulate primarily protein synthesis provides the basis and justification to ingest leucine during resistance exercise training in order to enhance muscle growth and ultimately improve strength and personal performance. An increase in muscle size is generally accompanied by an increase in strength. Strength gains may lead to improved performance in athletic disciplines and everyday life. Gains in lean mass and losses in fat mass may result in physiological adaptations that decrease the risk of diseases such as cardiovascular disease. Therefore, the main aim of the present study is to investigate the capacity of 4 g·d⁻¹ oral ingestion of leucine alone to enhance strength, lean mass and decrease fat mass during a 12-wk resistance-training program of initially untrained male participants. Our expectation was that supplementation with leucine would lead to further gains in muscle growth and consequently strength when compared with a placebo condition.

**Methods**

**Participants**

Forty healthy males were recruited in stages. Participants who failed to follow the resistance training program and did not comply with our participation requirements described below were removed from the study. Thus, the required number of 13 participants per group (determined by power calculation using Machin’s software [AUQ2] version 2) was met. The volunteers were assigned to a placebo group (P) (28.2 ± 7.3 y; 1.77 ± 0.10 m; 78.4 ± 16.8 kg; BMI 24.9 ± 4.2 kg·m⁻²) and a leucine supplemented group (L) (28.5 ± 8.2 y; 1.76 ± 0.05 m; 77.6 ± 7.9 kg; BMI
24.6 ± 3.2 kg·m⁻²). Attempts were made to match pairs of participants as closely as possible on age and BMI. Medical history and lifestyle evaluation questionnaires were collected before the assignment of the participants to one of the two conditions. The participants had never taken part in any structured resistance training program or any other type of exercise training program in the past. Volunteers who had participated in any type of training the 6 mo preceding the baseline measurements were also excluded. None of the participants had diabetes, high blood pressure, or symptomatic disease, including cardiovascular disease. All participants were free from medication and were not taking any other nutritional supplements during or before the intervention. The study was approved by the Leeds West Research Ethics Committee. All participants received information sheets and had the opportunity to ask questions about the potential benefits, risks and design of the study before informed consent was obtained.

Experimental Design and Strength Training Protocol

The design of the study was placebo-controlled and double blind. The participants agreed to take part exclusively in the assigned resistance training program. Both groups received identical prescribed resistance training over a period of 12 wk. The participants were advised to maintain the same lifestyle and physical activity levels before the commencement of the experimental period. The standard machines used during the strength assessment and strength training sessions were as follows: leg press, bench press, chest cross, pullover, overhead press, preacher curls, triceps press (All Nautilus, USA) and prone leg curl (Nautilus Nitro, USA). All sessions took place in a university gym. The participants trained twice per week (with 2–3 rest days between sessions). According to the position stand of the American College of Sports Medicine (ACSM) the initial resistance for novice subjects should be in the range of 8-12 repetition maximum (RM) strength, and the training frequency 2–3 d/wk. The duration of the training program was set at 12 weeks as muscle growth is more likely to become evident after 6–7 wk of resistance training. The five repetition maximum (5-RM) testing took place at baseline, at weeks 4, 8 and 12 of the experimental period. The participants warmed-up using a cycle ergometer (Startrac Pro, USA) for a period of 5 min followed by 5 min of dynamic arm and leg movements combined with gentle stretching of arms and legs. A 5-min rest was given between each trial to allow for adequate recovery. 5-RM was employed for assessment of strength in order to minimize the risk of injury because all subjects were untrained at baseline, and to determine rather than estimate the training loads during the strength workouts. Multiple RM testing between 3-RM to 6-RM has also been shown to be a valid index of assessment of strength. In addition to measuring 5-RM strength in all eight exercises, total 5-RM was determined by the sum of 5-RM strength on each individual exercise. One week before the first 5-RM testing all participants were familiarized with the training equipment using light resistance.

The training protocol combined two different types of workouts shown to increase strength and muscle size. The hypertrophy workout involved 3 sets of 10 repetitions with 1 min of rest between sets; the training loads were adjusted in order to allow the participants to perform just 10 repetitions during the last set. During the strength workouts the resistance used was the 5-RM achieved during the previous assessment; participants in this case performed 4 × 5-RM with 3 min of recovery between sets. If for any reason participants were unable to complete 5 repetitions
during their final set, resistance was adjusted in a subsequent session to allow them to perform 5 repetitions during the final set.

Assessment of Body Composition

Total and regional (arms, legs and trunk) body composition was assessed at baseline and at the end of the supplementation period by dual energy X-ray absorptiometry (DXA). A DXA scanner (Prodigy Lunar, GE Medical Systems, serial No: 10395, USA) was used for the assessment of bone mineral-free lean tissue mass (LTM) and fat mass (FM). The testing took place in the Centre for Bone and Body Composition Research, Leeds University. DXA has been validated against other methods and has shown little bias based on age, fat distribution, activity levels, gender or race, while at the same time is a more sensitive method for assessing small and regional changes in body composition.18,19

Diet and Supplements

The participants were asked to maintain their habitual diet throughout the experimental period and were asked to keep diet diaries for a period of 4 d at the midpoint of the study to include two training, one weekend and one week nontraining days. The parameters assessed for dietary analysis were: carbohydrate, protein, fat and alcohol intake. The CompEat program 5.7 (CompEat version 5, Nutrition systems, Grantham, UK) was used for the analysis of diary records. The resting energy expenditure (REE) was calculated using the Harris-Benedict equation20 and the total energy requirements were estimated by multiplying REE with a factor of 1.35.21

Supplementation was continuous for the duration of the study. On nontraining days participants were asked to take the supplements in three equal doses during the day (morning, midday, evening). On training days, the supplements were ingested immediately following exercise. The doses were 4 g·d⁻¹ (approximately 50 mg·kg⁻¹·BW·d⁻¹) for leucine (L-leucine white crystalline powder, Ajinomoto Co. Inc., Tokyo, Japan) and a corresponding amount of lactose (99% lactose monohydrate, Dairy Crest Ingredients, Surrey, UK). The composition of the placebo was not revealed to the participants. The leucine dose (4 g·d⁻¹) resulted from the average weight for British men of 77.2 kg (Allied Dunbar National Fitness Survey); 22 50 mg·kg⁻¹·BW·d⁻¹ seems to be an adequate quantity to prevent a decrease in serum leucine concentration during intensive training of athletes.7 There was good reason to believe that postexercise supplementation in addition to regular supplementation through the day would be more likely to enhance any anabolic effect of leucine than daily supplementation alone. This is because previous research has shown that ingestion of protein and amino acid mixtures immediately after (ideally within the first hour) exercise can create a more anabolic environment conducive to muscle hypertrophy.23 Continuity of supplementation was also of significant importance as protein turnover may remain elevated for up to 48 h following the end of exercise.24 We expected that ingestion of leucine at different time points during nonexercising days (in combination with ingestion of nutrients through their normal diet) would create a sustained positive or a more positive protein balance conducive to muscle hypertrophy throughout the day.

The volume of each drink was approximately 150 mL and consisted of four parts water, one part squash (Kia-Ora, Coca Cola Enterprises Ltd) and the daily
supplement doses; the energy value excluding the supplements was approximately 16 kcal (11 kcal·100 mL⁻¹). The squash was added in the drinks in an attempt to disguise the taste of the supplements and in particular the bitterness of leucine. The energy value of 4 g·d⁻¹ of lactose was 16 kcal (400 kcal·100 g⁻¹), while the energy value of 4 g·d⁻¹ of leucine was 24.8 kcal (6.2 kcal·g⁻¹). Powders were dispensed in plastic food bags that were sealed, placed in opaque envelopes, which were labeled as A, or B and then distributed to the participants. The participants were instructed to shake well the mix in the supplied beakers (black color). As a result there were no residues of leucine or lactose left in the beakers. Furthermore, as the experimental groups were discreet groups, neither the participants in the leucine group nor the participants in the placebo group knew what the drink tasted like in the opposite group.

Ratings of Perceived Exertion and 7-d Physical Activity Recall

Ratings of perceived exertion (RPE) were recorded at the end of each exercise within each training bout to compute session RPE (average of RPE in eight exercises). The 6–20 Borg scale has been used in resistance training studies to monitor exercise intensity²⁵ and has been found to be a very useful tool to distinguish between different training intensities. In addition to RPE, a 7-d physical activity recall (7-d PAR) interview took place at the end of the supplementation period. The purpose of the 7-d PAR, a reliable index in assessing activity levels²⁶ was to explore the activity levels of the two groups toward the end of the experimental period.

Blood Samples

Blood samples were drawn by venipuncture after a minimum of 10-h overnight fast at the beginning and at the end of the supplementation period to assess liver function (liver enzymes, bilirubin and albumin) and blood profile (standard full blood count). Subjects remained seated in a reclined position for 5-min before the sample was drawn from one of the brachial, medial cubital or radial veins. The samples were drawn in succession into a gold 3.5-mL (13 × 75 mm) BD SST II Vacutainer (BD Vacutainer Systems Preanalytical Solutions, UK) and a purple EDTA 4-mL Vacuette (Vacuette, Greiner Bio-One, Austria). The samples were transferred to the department of Clinical Biochemistry and Immunology, Leeds General Infirmary) for subsequent analysis. No other tests were conducted on blood sampling days.

Statistical Analysis

Data are presented as means (± standard deviations). Independent t test were employed to test for significant differences between groups in percentage strength and percentage LTM and FM differences (the difference between the absolute values expressed as percentage of the baseline value). In addition, effect sizes were computed by using the method of Cohen. Independent t test were also employed for analyzing 7-d PAR data and mean differences of blood data (postexercise absolute values minus baseline absolute values). Comparison of RPE data (average for each training session) were made using a 2-way ANOVA (condition × training sessions) with repeated measures across the training period for both the hypertrophy and strength training workouts. The alpha level of statistical significance was set at P <
Results

Strength Variables
Table 1 provides the descriptive statistics at baseline and end of the supplementation period. The percentage gains from baseline until the end of the experimental period in 5-RM strength for both groups ranged between 19% and 60% with a gain of approximately 30-40% in the majority of exercises (Table 1). On average the percentage 5-RM strength gains for the L group were approximately 10% higher than the P group. Statistical analysis on the mean percentage gains across the 12-week experimental period showed that the L group had significantly greater percentage mean gains than the P group in 5 out of 8 exercises and total strength ($P < .05$) (Table 1). The statistical analysis showed significant differences between groups for leg press ($t_{24} = –2.79; P = .010$), bench press ($t_{24} = –2.45; P = .02$), pullover ($t_{24} = –2.33; P = .03$), preacher curls ($t_{24} = –3.17; P = .004$), triceps press ($t_{24} = –3.47; P = .002$) and total strength ($t_{24} = –4.06; P < .001$). No significant differences between conditions were observed for leg curls ($P = .19$), chest cross ($P = .08$) and overhead press ($P = .35$).

Body Composition
Table 2 reports the body composition variables at baseline and week 12. The mean gains in total bone mineral-free lean tissue mass (LTM) and losses in total fat mass (FM) were 1.08 kg ($±1.1$), 1.53 kg ($±1.3$) and 0.41 kg ($±1.4$), 0.93 kg ($±3.0$) on average for P and L groups respectively. Independent $t$ test showed that these percentage changes were not significantly different between groups (LTM: $t_{24} = –0.94; P = .36$ and FM: $t_{24} = 0.10; P = .92$). Similarly, significant differences did not exist between groups in LTM or FM of arm, leg and trunk regions (Table 2).

Dietary Intake
The daily percentage contribution of different nutrients and alcohol to total energy as well as the total energy intakes were similar for both groups (Table 3). One participant from the P and two participants from the L group failed to submit their diet diaries. The P group met 98.2% ($±24.6$) and the L group 89.8% ($±18.2$) of the estimated energy requirements. The dietary grams of protein per kilogram of body weight were 0.88 g ($±0.28$) and 0.90 g ($±0.15$) for P and L groups respectively. There were no significant differences between the two groups in either absolute or percentage terms.
RPE
Ratings of perceived exertion (RPE) during each hypertrophy workout (session RPE) ranged from 16.2 (±1.9) to 17.6 (±1.2) and 16.1 (±1.9) to 17.8 (±1.3) for P and L groups respectively. During the strength workouts session RPE ranged from 16.2 (±1.4) to 18.3 (±0.8) and 16.3 (±1.6) to 18.1 (±1.1) for P and L groups respectively. Significant differences did not exist between groups. There was neither a condition main effect ($P = .499$ for hypertrophy and $P = .865$ for strength workouts) nor an interaction main effect ($P = .175$ for hypertrophy and $P = .540$ for strength workouts).

7-d Physical Activity Recall
The physical activity recall data showed no differences in the number or intensity in work related and/or other activity levels between both groups.

Blood
All blood variables were within the normal range (as provided by the Department of Clinical Biochemistry and Immunology, Leeds General Infirmary). No significant differences between groups were found.

Discussion
The resistance training program for the novice weight trainers employed in the current study was associated with positive adaptations in strength and LTM for both the P and L groups. The magnitude of gains for both groups was in the expected range of approximately 20–40% for moderately trained and untrained individuals. Daily ingestion of leucine during 12 wk of resistance training was accompanied by significantly greater gains in 5-RM strength in five out of eight resistance exercises and 5-RM total strength compared with similar training with a placebo.

The gains observed in strength in both groups in the present study are most likely the result of the prescribed supervised resistance training program. Participants were clearly instructed to refrain from any lifestyle or activity modifications (other than the weight training sessions) during the experimental period. Hence, differences between the two conditions are likely to be attributed to the ingestion of leucine supplementation. The groups engaged in exactly the same type and volume of supervised resistance exercises and received the same training stimulus during these sessions as indexed by similar RPE ratings for both the hypertrophy and strength workouts. In addition, participants did not differ in activity or exercise levels toward the end of the intervention program nor did they differ in their dietary intake. Of course, differences in genetics or muscle fiber distribution might result in interindividual differences in progress. Because genotype has a bearing on the magnitude of gains in strength and/or FFM on its own right, recruitment of identical twins might be the best way to test the effectiveness of leucine as an ergogenic aid. However, the control mechanisms in place in the present study provide support for the notion that leucine supplementation might act as an ergogenic aid in novice weight trainers.

Our hypothesis that leucine supplementation would lead to greater gains in strength due to additional gains in muscle growth was supported by the strength data but only weakly supported by the LTM data (Table 2). The leucine group had
slightly higher percentage gains and losses in total LTM and FM respectively than the P group but none of these differences were significant. However, the small effect size (Table 2) in trunk and total LTM suggests that a study with a longer duration and a higher training frequency may have resulted in significant differences between groups in LTM. The current study design was implemented to improve adherence to the study protocol. Despite the fact that participants were only required to engage in training twice per week only 26 out of 40 participants completed the required number of training sessions. This resulted in a dropout of 35%. An aim of future studies would be to improve adherence to training programs.

The dietary intakes of the participants were under the estimated energy requirements but all within the 15% expected underestimation when food diaries are used. Ingestion of extra amino acids through the diet, which may have favored one of the groups in terms of muscle protein synthesis, was not the case since the protein intake per kilogram of body weight was identical in both groups. Thus, the most likely explanation of the marginally higher gains in LTM of the L group, supported by a small effect size in LTM and accompanied by significantly greater gains in 5-RM strength, was leucine supplementation. We acknowledge that four days of diet records may not be adequate to estimate actual intakes across the twelve weeks. However, it has been shown that as dietary record duration increases, the accuracy and recording declines.

We can only speculate about the mechanisms of action since the data collected is not appropriate for this purpose. Coingestion of leucine and protein has shown to elicit a greater anabolic response in untrained individuals than carbohydrate alone or carbohydrate/protein, while removal of leucine in the presence of all other amino acids can decrease protein synthesis by 40%. Thus, it is probable that leucine supplementation further enhanced the rates of muscle protein synthesis in our L group when compared with diet alone and to some extent this is supported by the small effect size in LTM. Others have suggested that addition of leucine to a whey protein supplement before exercise does not result in greater anabolic responses than whey protein alone. However, this needs to be further investigated since neither a whey alone nor a leucine alone group were included in their study. The role of amino acids, obtained from either endogenous or exogenous sources, in enhancing the rate of protein synthesis cannot be ignored; as in some conditions leucine enhances protein synthesis only in the presence of these other amino acids. However, it is unlikely our participants were depleted of any amino acids since protein intake was adequate and similar in both groups and supplements were coingested with food during the nonexercising days. It is also unlikely the significant gains in 5-RM strength of L group was the result of amelioration of central fatigue since leucine ingestion elevates plasma leucine and BCAA concentrations without affecting plasma free-tryptophan and the ratio of free-tryptophan to BCAA ratio.

Although the dose used in the current study (4 g·d⁻¹) seems to be an adequate amount to effect positive adaptations during 12 wk of combined resistance training and supplementation in novice weight trainers, different dosages of leucine (higher or lower) should also be tested in future studies. Older individuals seem to require higher doses of leucine for stimulation of protein synthesis. However, more evidence-based research to test the efficacy and safety of different dosages is needed, before prescription of higher dosages. In our study, there was no evidence to suggest that 4 g·d⁻¹ leucine ingestion for a period of up to 12 wk during resistance
training negatively affected the health of the male participants as indicated by tests of liver function and full blood counts.

Practical Applications
The additional strength gains in the L group support the use of leucine as a diet supplement to improve strength and ultimately performance in everyday life and athletic disciplines. In cases where supplementation is deemed necessary (eg, high intensity, high volume exercise), supplementation could be considered and complement an appropriate diet. An investigation of special significance would be to test the hypothesis that leucine supplementation may be of benefit in diseases in which lean body mass is compromised by atrophy such as type II diabetes based on supporting evidence that leucine can attenuate body wasting.2

Conclusion
In conclusion, the resistance training program employed in the current study increased strength of initially untrained male participants regardless of whether they ingested leucine or a placebo. However, those participants ingesting leucine had the greatest increase in strength. The data therefore suggest that regular ingestion of leucine while undertaking a resistance training program may accentuate strength gains. However, given the small effect size in LTM, more research is needed before any concrete conclusions can be made regarding the efficacy of leucine as an ergogenic aid.

Acknowledgments
The research was funded by the Greek National Foundation Scholarships (IKY). Special thanks to the men who participated in this research study and the laboratory technician Brian Oldroyd (Centre for Bone and Body Composition, University of Leeds) for his assistance with the research.

References


Table 1  Mean strength expressed as five repetition maximum strength (5-RM) at baseline and at the end of the supplementation period in eight resistance exercises and total strength

<table>
<thead>
<tr>
<th>Exercise</th>
<th>Placebo (n = 13)</th>
<th>Leucine (n = 13)</th>
<th>Effect Size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Week 12</td>
<td>% Mean Gain</td>
</tr>
<tr>
<td>Leg press (kg)</td>
<td>171.1 ± 28.4</td>
<td>216.9 ± 35.1</td>
<td>27.0 ± 7.8</td>
</tr>
<tr>
<td>Bench press (kg)</td>
<td>71.3 ± 12.0</td>
<td>93.6 ± 12.9</td>
<td>31.9 ± 8.7</td>
</tr>
<tr>
<td>Leg curls (kg)</td>
<td>58.1 ± 12.4</td>
<td>68.3 ± 12.0</td>
<td>18.6 ± 7.5</td>
</tr>
<tr>
<td>Chest cross (kg)</td>
<td>56.5 ± 12.8</td>
<td>82.6 ± 14.4</td>
<td>48.2 ± 10.7</td>
</tr>
<tr>
<td>Pullover (kg)</td>
<td>71.1 ± 14.3</td>
<td>89.1 ± 15.4</td>
<td>26.3 ± 7.7</td>
</tr>
<tr>
<td>Overhead press (kg)</td>
<td>36.1 ± 6.9</td>
<td>53.0 ± 8.8</td>
<td>48.0 ± 14.7</td>
</tr>
<tr>
<td>Preacher curls (kg)</td>
<td>47.3 ± 9.0</td>
<td>60.4 ± 9.1</td>
<td>29.0 ± 9.0</td>
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<tr>
<td>Triceps press (kg)</td>
<td>75.3 ± 11.1</td>
<td>102.6 ± 12.4</td>
<td>36.9 ± 7.2</td>
</tr>
<tr>
<td>Total strength (kg)</td>
<td>587.1 ± 89.6</td>
<td>766.6 ± 103.7</td>
<td>31.0 ± 4.6</td>
</tr>
</tbody>
</table>

* Denotes significant difference in percentage gain between the placebo and leucine condition (* P < .05; **P < .01).
Table 2  Bone mineral-free lean tissue mass (LTM), fat mass (FM) and soft tissue mass (STM = LTM + FM) by DXA at baseline and at the end of the experimental period

<table>
<thead>
<tr>
<th>Body Composition Variable</th>
<th>Placebo (n = 13)</th>
<th>Leucine (n = 13)</th>
<th>Effect Size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Week 12</td>
<td>% Change</td>
</tr>
<tr>
<td>Arm FM (kg)</td>
<td>1.6 ±1.0</td>
<td>1.5± 1.0</td>
<td>–3.1 ± 12.7</td>
</tr>
<tr>
<td>Leg FM (kg)</td>
<td>6.8 ± 3.0</td>
<td>6.5 ± 2.8</td>
<td>–4.4 ± 7.1</td>
</tr>
<tr>
<td>Trunk FM (kg)</td>
<td>11.3 ± 6.2</td>
<td>11.2 ± 5.7</td>
<td>–1.3 ± 8.6</td>
</tr>
<tr>
<td>Arm LTM (kg)</td>
<td>6.6 ± 0.9</td>
<td>7.0 ± 0.9</td>
<td>6.1 ± 5.6</td>
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<tr>
<td>Leg LTM (kg)</td>
<td>20.0 ± 2.7</td>
<td>20.4 ± 2.8</td>
<td>1.8 ± 2.5</td>
</tr>
<tr>
<td>Trunk LTM (kg)</td>
<td>27.6 ± 5.0</td>
<td>27.9 ± 4.9</td>
<td>1.3 ± 3.1</td>
</tr>
<tr>
<td>Total FM (Arm+Leg+Trunk) (kg)</td>
<td>19.7 ± 9.9</td>
<td>19.3 ± 9.4</td>
<td>–1.1 ± 7.7</td>
</tr>
<tr>
<td>Total LTM (Arm+Leg+Trunk) (kg)</td>
<td>54.2 ± 7.9</td>
<td>55.2 ± 8.0</td>
<td>2.0 ± 2.1</td>
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<tr>
<td>Total STM</td>
<td>76.7 ± 16.7</td>
<td>77.4 ± 16.1</td>
<td>1.1 ± 2.2</td>
</tr>
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</table>
## Table 3  Average daily dietary intakes over a period of 4 (two training days, one weekend day, and one nontraining weekday) in the middle of the training period

<table>
<thead>
<tr>
<th>Dietary Variable</th>
<th>Placebo (n = 12)</th>
<th>Leucine (n = 11)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbohydrate intake (kcal)</td>
<td>1180.7 ± 344.9</td>
<td>1012.2 ± 210.6</td>
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<tr>
<td>Protein intake (kcal)</td>
<td>393.7 ± 108.6</td>
<td>355.0 ± 67.2</td>
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<tr>
<td>Fat intake (kcal)</td>
<td>853.5 ± 215.3</td>
<td>718.2 ± 165.7</td>
</tr>
<tr>
<td>Alcohol intake (kcal)</td>
<td>96.6 ± 163.8</td>
<td>182.7 ± 290.2</td>
</tr>
<tr>
<td>Carbohydrate (% of TEI)</td>
<td>48.2 ± 7.4</td>
<td>46.3 ± 7.5</td>
</tr>
<tr>
<td>Protein (% of TEI)</td>
<td>16.0 ± 2.5</td>
<td>16.5 ± 3.5</td>
</tr>
<tr>
<td>Fat (% of TEI)</td>
<td>34.8 ± 4.9</td>
<td>33.2 ± 8.0</td>
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<tr>
<td>Alcohol (% of TEI)</td>
<td>4.0 ± 6.6</td>
<td>6.9 ± 10.1</td>
</tr>
<tr>
<td>TEI (kcal)</td>
<td>2450.6 ± 554.0</td>
<td>2204.8 ± 444.8</td>
</tr>
</tbody>
</table>

*Note.* TEI = total energy intake.
Author Queries

[AUQ1] Please verify and/or supply the departmental affiliations.

[AUQ2] “Machin” --Please verify the spelling of this proper noun or how the 18th-century John Machin (?) and software are connected.