1. Introduction

Aging is commonly associated with the progressive loss of skeletal muscle tissue and functional capacity. Illness and injury notwithstanding, sarcopenia is facilitated by factors secondary to the adoption of a more sedentary lifestyle and consumption of a less than optimal diet. While maintaining or increasing physical activity is clearly desirable, establishing a basic nutrition foundation focusing on adequate protein and energy consumption is necessary if other interventions are to succeed.

It is generally accepted that aging is associated with a blunted protein synthetic response to meals containing less than approximately 15–20 g of protein or the equivalent essential amino acid content.1–3 While this deficiency may be partially or wholly overcome by simply increasing protein intake, obstacles such as total energy consumption, palatability, cost, satiety and habitual practices must be considered.4,5 Consequently, the practical validity of prescribed supplementation regimens must also be critically examined. For example, in controlled metabolic studies it is easy to demonstrate acute increases in muscle protein synthesis or markers of anabolism following ingestion of a protein or amino acid supplement.6–8 However, in practice, there is a risk that acute responses will not translate into positive chronic adaptations in muscle mass or function. The reasons for a lack of success are varied, but we suspect they more often reflect compliance-related issues or consumer choice, rather than a temporally diminished physiological capacity.9,10 To this end, we propose that an effective supplement should produce a robust anabolic response, and be i) low-volume and easily incorporated into existing menu plans, ii) palatable, and iii) cost effective.

From a practical and mechanistic perspective, the branch chain amino acid leucine is an attractive supplement. Leucine has many well described effects on the regulatory mechanisms controlling translation initiation and muscle protein synthesis.9,10,11–18 Nevertheless, aging muscle appears to be less responsive to the stimulatory effects of normal/typical post-prandial concentrations of leucine.11 Several studies in both
animals and humans suggest that increasing the leucine content of regular mixed nutrient meals may normalize or even increase protein synthesis in older populations. Specifically, Rieu et al. reported that following a mixed meal, muscle protein synthesis in an older rat population was blunted compared to younger rats. However, the addition of supplemental leucine to the meal restored/normalized muscle protein synthesis in the older animals. In a follow-up study in older adults (70 ± 1 yr), muscle protein synthesis was acutely increased in response to a leucine supplemented meal (0.053 ± 0.009%/h vs. 0.083 ± 0.008%/h, p < 0.05). Of note, however, is the fact that for 4 days preceding the metabolic study, the older adults consumed a controlled diet that limited protein intake to 0.8 g/kg/day. As described below, we propose that habitual protein intake is a key determinant of the efficacy of leucine/protein supplementation regimens.

While the headline result from the leucine studies mentioned above are encouraging, an important emerging caveat is that leucine supplementation is not unconditionally effective. Specifically, ingestion of supplemental leucine appears to be of little value to: i) younger adults, and/or ii) individuals who habitually consume a protein/leucine-deficient diet. For example, in two diabetesogenic murine models, no change in skeletal muscle mass or strength was reported in a cohort of healthy well-designed studies, no change in skeletal muscle mass or strength was determined via Dual Energy X-Ray Absorptiometry (DEXA, Hologic, Inc., Natick, MA). On the morning of the metabolic study, following an overnight fast, polyethylene catheters were inserted into the antecubital vein of both arms for infusion of the stable isotope and arterialized venous blood sampling. Baseline blood samples were drawn for the analysis of hormones and background amino acid enrichment. A primed (2 μmol/kg), continuous infusion of L-[ring-13C6] phenylalanine (Phe) (0.08 μmol kg⁻¹ min⁻¹) was started (time 0) and continued uninterrupted until the conclusion of each metabolic study (Fig. 1). Blood samples were obtained at hours 2 and 3 (fasting) and every 20 min for the hour prior to and following ingestion of the simulated meal. To simulate a mixed meal, a standardized beverage containing 7 g of essential amino acids and 10 g carbohydrates (sucrose) (Table 1) dissolved in approximately 300 mL of diet soda was ingested at hour 4. 0.05 g of L-[ring-13C6] phe was added to maintain an isotopic steady state. The amino acid content of this meal was consistent with a 80–90 g serving of most animal proteins (e.g., beef, chicken, fish) and while the digestion/absorption kinetics of amino acids would certainly differ from protein-rich foods, this relatively small simulated meal has been previously shown to elicit only a modest anabolic response in older adults.

A total of four muscle biopsies (~100–200 mg) were taken from the midportion of the vastus lateralis muscle during each stable isotope study to capture data from the postabsorptive (biopsy 1 & 2) and postprandial (biopsy 3 & 4) periods (Fig. 1). Samples were snap frozen in liquid nitrogen to abruptly stop all enzymatic reactions and frozen at −80 °C for later analysis.

2. Materials and methods

2.1. Participants

Eight healthy but sedentary older adults participated in this project. The study was approved by the Institutional Review Board of the University of Texas Medical Branch. Written informed consent was obtained prior to participation. Volunteers were screened at the University Of Texas Medical Branch (UTMB) Institute for Translational Sciences Clinical Research Center (ITS-CRC) to determine eligibility. Inclusion criteria included a habitual protein intake (assessed via a 3-day dietary recall questionnaire) at or near the current RDA for protein (range: 0.75–0.85 g protein/kg/day). Exclusion criteria included: a history of regular physical exercise or training (~30 min, 3 times/wk) cardiac, liver, kidney or autoimmune disease; hypo- or hypercoagulation disorders; diabetes; cancer; uncontrolled hypertension, infectious disease, a BMI > 30; recent history of anabolic steroid or corticosteroid use; or recent participation in a weight loss diet. During the experimental period, subjects were instructed to continue all regular activities of daily living and maintain their usual diet.

2.2. Experimental protocol

Each subject completed two inpatient visits at the ITS-CRC (days 1 and 15) with an intervening two-week outpatient period involving leucine supplementation (4 g per meal; 3 meals per day; days 2–14) (Fig. 1). Subjects reported to the ITS-CRC at noon the day before each metabolic study at which time total body fat and lean mass were determined via Dual Energy X-Ray Absorptiometry (DEXA, Hologic, Inc., Natick, MA). On the morning of the metabolic study, following an overnight fast, polyethylene catheters were inserted into the antecubital vein of both arms for infusion of the stable isotope and arterialized venous blood sampling. Baseline blood samples were drawn for the analysis of hormones and background amino acid enrichment. A primed (2 μmol/kg), continuous infusion of L-[ring-13C6] phenylalanine (Phe) (0.08 μmol kg⁻¹ min⁻¹) was started (time 0) and continued uninterrupted until the conclusion of each metabolic study (Fig. 1). Blood samples were obtained at hours 2 and 3 (fasting) and every 20 min for the hour prior to and following ingestion of the simulated meal. To simulate a mixed meal, a standardized beverage containing 7 g of essential amino acids and 10 g carbohydrates (sucrose) (Table 1) dissolved in approximately 300 mL of diet soda was ingested at hour 4. 0.05 g of L-[ring-13C6] phe was added to maintain an isotopic steady state. The amino acid content of this meal was consistent with a 80–90 g serving of most animal proteins (e.g., beef, chicken, fish) and while the digestion/absorption kinetics of amino acids would certainly differ from protein-rich foods, this relatively small simulated meal has been previously shown to elicit only a modest anabolic response in older adults.

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2.3. Analytical methods

Dietary analysis from a 3-day diet recall period (pre-study screening) and daily diet logs (14 day supplementation period) were analyzed using Nutrition Data System for Research software version 2006, (Nutrition Coordinating Center Minneapolis, MN).

Phenylalanine enrichments in arterialized venous blood samples were determined after deproteinization with sulfosalicylic acid, extraction with cation exchange chromatography (Dowex AG 50W-8X, 100–200 mesh H⁺ form; BioRad Laboratories, Inc., Richmond, CA), dried under vacuum (Savant Speedvac, Thermo Fisher Scientific, Inc., Waltham, MA), and tert-butyldimethylsilyl (t-BDMS)
derivation using gas-chromatography mass-spectrometry (GCMS) in electron impact mode (GC HP 5890, MSD HP 5898, Hewlett Packard, Palo Alto, CA).26

Muscle samples were weighed and the proteins precipitated with 800 μl of 10% sulfosalicylic acid. Tissue homogenization and centrifugation were performed on two occasions, and the pooled supernatant was collected. Intracellular amino acids were purified using cation exchange chromatography (Dowex AG 50W-8X, 200–400 mesh H+ form; BioRad Laboratories, Inc., Richmond, CA) and t-BDMS derivatization using GCMS in electron impact mode.27

The remaining pellet containing bound mixed-muscle proteins was repeatedly washed, dried at 50 °C overnight and hydrolyzed in 3 ml of 6 N HCl at 110 °C for 24 h. Amino acids in the hydrolysate were extracted and derivatized, as previously described, by monitoring the ions 238 and 240 and using the external standard curve approach.28

2.4. Calculations

Mixed muscle fractional synthesis rate (FSR) was calculated by directly measuring the incorporation of L-[ring-13C6]-phenylalanine into muscle protein using the precursor-product model:

\[
FSR = \frac{(E_{BP2} - E_{BP1})}{(E_p \times t)} \times 60 \times 100
\]

where \(E_{BP1}\) and \(E_{BP2}\) are the enrichments of bound L-[ring-13C6]-phenylalanine in sequential muscle biopsies, \(t\) is the time interval between biopsies, and \(E_p\) is the mean L-[ring-13C6]-phenylalanine enrichment in the muscle intracellular or plasma precursor pools.

Muscle phenylalanine concentrations \((C)\) (μmol/L) in total muscle tissue fluid was determined using a chloride correction29 and calculated as:

\[
C = \left[\frac{Q_{IS}}{V \times E_{IS}}\right]
\]

where \(Q_{IS}\) (μmol) is the amount of internal standard added to the sample, \(V\) is the volume of muscle tissue fluid and \(E_{IS}\) is the tracer-to-tracee ratio of internal standard in total muscle water. Measured values of intracellular phe concentrations relative to total tissue water were corrected using the chloride correction method.29 The chloride method is based on the premise that chloride is freely diffusible across the skeletal muscle fiber membrane and is distributed according to Nernst’s equation and assuming a normal resting membrane potential of muscle in man to be −872 mV.

2.5. Immunoblot analysis

Biopsy samples from the postabsorptive (biopsy 1) and postprandial (biopsy 3 and 4) periods were homogenized in tissue lysis buffer, and protein concentrations were determined. A total of 80 μg of protein was loaded per lane and separated by SDS-PAGE. Following SDS-PAGE, proteins were transferred to polyvinylidene difluoride membranes (PVDF) (Hybond-P, Amersham Biosciences, Piscataway, NJ) and blocked for 1 h. Blots were serially washed and incubated with primary antibody overnight at 4 °C with constant agitation. The next day, the blots were washed and incubated with secondary antibody for 1 h at room temperature with constant agitation. Blots were serially washed and incubated for 5 min with enhanced chemiluminescence reagent (ECL Advanced Western Blotting Detection System, Amersham Biosciences, Piscataway, NJ). Optical density measurements were obtained with a ChemiDoc XRS imaging system (BioRad, Hercules, CA). Densitometric analysis was performed by using Quantity One 1-D analysis software (Ver 4.5.2) (BioRad, Hercules, CA). Data are expressed as the phosphorylation divided by total protein content (in arbitrary units) normalized to a rodent standard.

2.6. Antibodies

Primary antibodies were purchased from Cell Signaling (Beverly, MA): phosphor-mTOR (Ser2448), phosphor-p70S6K1 (Thr389), phospho–4E-BP1 (Thr37/46), total mTOR, total p70S6K1 and total 4E-BP1. Anti-rabbit IgG horseradish peroxidase-conjugated secondary antibody was purchased from Amersham Bioscience (Piscataway, NJ).

2.7. Amino acid analysis

Amino acids were analyzed by a solid phase extraction followed by derivatization and a liquid/liquid extraction (EZ:faast Amino Acid Analysis Kit, Phenomenex, Inc., Torrance, CA). Briefly, calibration standards were prepared at concentration of 50, 100 and 200 nmol/ml from 2 standard amino acid solutions. Standard 1 contained a mixture of AAA (α-aminoacidic acid), ABA (α-amino-butyric acid), (allo-isoleucine), Ala, Asp, βAB (β-isobutyric acid), C–C, Glu, Gly, His, Hyp (4-hydroxyproline), Ile, Leu, Lys, Met, Orn, Phe, Pro, Sar, Ser, Thr, Tyr, and Val. Standard 2 contained Asn, Gln and Trp, which were unstable in acid solution. Calibration standards and serum samples were passed through a sorbent packed tip, binding the amino acids while allowing interfering compounds to flow through. The amino acids on sorbent were then extruded and quickly derivatized in an aqueous solution to facilitate concomitant migrate of the amino acids to the organic layer for additional separation from interfering compounds. The organic layer was then removed, evaporated, and re-suspended in redissolution solvent and analyzed using gas-chromatography mass-spectrometry (GCMS: GC HP 5890, MSD HP 5898, Hewlett Packard, Palo Alto, CA).

2.8. Insulin and glucose

Serum insulin concentrations were measured by using an IMMULITE 8000 (Siemens Healthcare Diagnostics, Deerfield, IL, USA). Plasma glucose concentrations were measured by using an automated glucose analyzer (YSI, Yellow Springs, OH, USA). Four blood draws were taken prior to ingestion of a test meal, and these samples were averaged and reported as the basal value. All post-prandial blood draws were analyzed and reported individually.

2.9. Statistical analysis

Each subject served as his or her own control. All values are expressed as means ± SEM. Comparisons were performed by using ANOVA with repeated measures, the effects being time and day. For post hoc testing we used Bonferroni when appropriate. Significance was set at \(P \leq 0.05\). Descriptive data are presented as means ± SEM.
Three weeks of leucine supplementation increased post-absorptive muscle protein synthesis (Day 1: 0.063 ± 0.004 vs. Day 15: 0.074 ± 0.007%/h; p = 0.004). The response to the simulated meal was also increased by leucine supplementation (Day 1: 0.075 ± 0.006 vs. Day 15: 0.10 ± 0.007%/h; p = 0.01), (Fig. 3). A similar pattern was observed using plasma phenylalanine as the precursor (see calculations) for postabsorptive (Day 1: 0.040 ± 0.004 vs. Day 15: 0.048 ± 0.004%/h; p = 0.02) and postprandial (Day 1: 0.060 ± 0.002 vs. Day 15: 0.077 ± 0.008%/h; p = 0.001) muscle protein synthesis (Fig. 4).

3.4. Nutrient anabolic signaling

The simulated meal increased the phosphorylation of mTOR, p70S6K1, and 4E-BP1 (p < 0.05; Fig. 5). Two weeks of leucine supplementation increased mTOR (Day 1: 1.03 ± 0.07 vs. Day 15: 1.23 ± 0.11 AU; p = 0.01) and p70S6K1 (Day 1: 0.89 ± 0.06 vs. Day 15: 1.01 ± 0.07 AU) phosphorylation in response to the simulated meal; (p ≤ 0.05). We also noted an increase in postabsorptive 4E-BP1 phosphorylation (Day 1: 0.30 ± 0.02 vs. Day 15: 0.37 ± 0.03 AU; p = 0.03). Total protein did not significantly change in response to the simulated meal or leucine supplementation.

3.5. Amino acid analysis

Postabsorptive plasma concentrations of essential and non-essential amino acids did not change following 14 days of leucine supplementation (Table 3).

3.6. Serum insulin and glucose concentrations

No changes in postabsorptive serum insulin concentrations following 2-weeks of leucine supplementation were noted (P < 0.05). However, on Day 15, subjects experienced a 33% lower insulin response 20 min following the simulated meal (Day 1: 18.0 ± 3.7 vs. Day 15: 12.0 ± 2.6 mg/dl; p = 0.004). There was no change in the concentration of plasma glucose following the simulated meal or leucine supplementation.
4. Discussion

Our data suggest that leucine supplementation may be an energetically efficient and practical means of chronically improving muscle protein synthesis in response to a low protein meal in older adults who habitually consume close to the RDA for protein. In the context of preventing or slowing sarcopenic muscle loss, the rationale for the use of a dietary supplement should take into account one or more of the following assumptions: i) supplementation will improve net muscle protein anabolism above that afforded by regular meals alone, ii) the supplement will not negate the rationale for the use of a dietary supplement, and iii) use of the supplement will not be compromised or restricted by a lack of compliance, complex preparation, high cost or poor palatability. To this end, we demonstrated that supplementing regular daily meals with a relatively small amount of leucine improves both mixed-muscle protein synthesis and anabolic signaling in older adults. Further, while we did not conduct a specific assessment of practical issues such as longer duration compliance, ease of use or palatability, our low-volume approach is intuitively less likely to be burdensome than most traditional higher-volume amino acid supplement regimens.

A key finding of this study was the postabsorptive and postprandial improvement in muscle protein synthesis after 2-weeks of leucine supplementation. Speed of digestion and absorption notwithstanding, the study meal containing 7 g of essential amino acids and 10 g carbohydrates was designed with two goals in mind. First, we wanted it to mimic the amino acid and carbohydrate content of a typical small meal or snack consumed by older adults (e.g., approximately a half of a chicken breast and a half cup of cooked rice). Second, we wanted to provide a meal that would initially produce a marginal protein synthetic response; a characteristic of meals ingested by older adults at risk of, or experiencing sarcopenic muscle loss. Data from Day 1 clearly show that the simulated meal had a minimal effect on muscle protein synthesis. However, the response to the same meal after 2-weeks of leucine supplementation was quite robust, despite the fact that leucine supplementation was not consumed during the metabolic study days (Day 1 and 15). Conceptually, this suggests that leucine supplementation is able to chronically up regulate the potential for muscle protein synthesis thereby making an otherwise insufficient or marginal quantity of protein more biologically available and efficient. Notably, we saw no differences in plasma amino acid concentrations during the metabolic studies on Day 1 and 15. In some instances, elevated plasma leucine concentrations have been associated with a reduction in the plasma or intracellular concentrations of other essential amino acids, particularly isoleucine and valine.39

The anabolic potential of leucine has been attributed to its capacity to stimulate translation initiation both dependently and independently of the mTOR signaling pathway.13,31 As noted, aging has been associated with impairment of nutrient-sensing signaling pathways.1 However, our data show that the postprandial phosphorylation of mTOR, and its downstream effector p70S6K, was more pronounced following chronic leucine supplementation.13,15,31 Further, leucine supplementation increased post-absorptive 4E-BP1 phosphorylation, suggesting that leucine supplementation effects muscle protein synthesis through increased elf4E availability and constitutive hyperphosphorylation of 4E-BP1.

In addition to its effects on muscle protein synthesis, there is some evidence suggesting that leucine augments glucose-induced
Compared with the status quo, leucine supplementation was more effective than the status quo. Further, there is no mechanism or previous data to suggest that an iso-

effect of chronic leucine supplementation on markers of nutrient signaling.

Fig. 5: Effects of chronic leucine supplementation on markers of nutrient signaling. Comparisons of the phosphorylation state of mTOR (A), p70S6K (B), and 4E-BP1 (C) obtained on Day 1 (□) and Day 15 (●). Protein phosphorylation and total protein were determined before and again at 30 and 180 min after ingestion of the simulated meal. A representative blot of the phosphorylation on Day 1 and Day 15 is shown above each graph. Values are expressed as means ± SEM. * significantly different from basal and # significantly different from Day 1; P < 0.05.

Table 3
Mean serum amino acid concentrations before and after 14 days of leucine supplementation.

<table>
<thead>
<tr>
<th>Essential amino acids</th>
<th>Day 1</th>
<th>Day 15</th>
</tr>
</thead>
<tbody>
<tr>
<td>Leucine</td>
<td>127 ± 9</td>
<td>132 ± 10</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>69 ± 7</td>
<td>66 ± 7</td>
</tr>
<tr>
<td>Valine</td>
<td>278 ± 19</td>
<td>250 ± 19</td>
</tr>
<tr>
<td>Histidine</td>
<td>87 ± 7</td>
<td>80 ± 5</td>
</tr>
<tr>
<td>Lysine</td>
<td>182 ± 10</td>
<td>196 ± 15</td>
</tr>
<tr>
<td>Methionine</td>
<td>21 ± 2</td>
<td>24 ± 2</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>71 ± 3</td>
<td>82 ± 6</td>
</tr>
<tr>
<td>Threonine</td>
<td>144 ± 14</td>
<td>140 ± 13</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Non-essential amino acids</th>
<th>Day 1 (μM)</th>
<th>Day 15 (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alanine</td>
<td>512 ± 51</td>
<td>483 ± 35</td>
</tr>
<tr>
<td>Asparagine</td>
<td>49 ± 5</td>
<td>56 ± 6</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>27 ± 5</td>
<td>28 ± 4</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>115 ± 13</td>
<td>119 ± 16</td>
</tr>
<tr>
<td>Glutamine</td>
<td>728 ± 58</td>
<td>645 ± 43</td>
</tr>
<tr>
<td>Glycine</td>
<td>336 ± 49</td>
<td>330 ± 44</td>
</tr>
<tr>
<td>Proline</td>
<td>305 ± 43</td>
<td>278 ± 34</td>
</tr>
<tr>
<td>Serine</td>
<td>115 ± 14</td>
<td>114 ± 12</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>68 ± 7</td>
<td>76 ± 4</td>
</tr>
<tr>
<td>Tryptophan</td>
<td>62 ± 4</td>
<td>67 ± 4</td>
</tr>
<tr>
<td>Sarcosine</td>
<td>631 ± 63</td>
<td>596 ± 43</td>
</tr>
<tr>
<td>a-Aminobutyric acid</td>
<td>14 ± 2</td>
<td>16 ± 23</td>
</tr>
<tr>
<td>b-Aminoisobutyric acid</td>
<td>89 ± 0.3</td>
<td>89 ± 0.3</td>
</tr>
<tr>
<td>Hydroxyproline</td>
<td>19 ± 2</td>
<td>22 ± 2</td>
</tr>
<tr>
<td>a-Aminoadipic acid</td>
<td>7 ± 0.4</td>
<td>5 ± 0.9</td>
</tr>
<tr>
<td>Ornithine</td>
<td>97 ± 7</td>
<td>96 ± 8</td>
</tr>
<tr>
<td>Cystine</td>
<td>51 ± 2</td>
<td>52 ± 3</td>
</tr>
</tbody>
</table>

Values are expressed in μM as means ± S.E.M.

Insulin secretion. In animal models, chronic leucine supplementation improved insulin sensitivity despite the consumption of a high-fat diet. However, recently Verhoeven et al. reported no such benefit in healthy older individuals. Furthermore, in a group of type 2 diabetic older men no changes were observed in glycemic control with leucine supplementation. Consistent with these findings, basal glucose and insulin concentrations in the present study did not change. However, we did detect a decrease in insulin concentrations immediately after the consumption of the simulated meal. This subtle improvement suggests a contribution of leucine to the enhancement of insulin-mediated glucose disposal in normal healthy aging.

While our results are encouraging, there are clearly limitations both in the design of this study and its generalizability. This was a small, non-randomized study with a limited cohort and no direct measure of muscle protein breakdown. Participants were not overtly sarcopenic, but rather representative of a population at increased risk of muscle loss. Despite the lack of a traditional control group, our pre-post-test design demonstrated that leucine supplementation was more effective than the status quo further, there is no mechanism or previous data to suggest that an iso-
energetic control could provide a comparable result. Similarly, it is possible, but unlikely that an alternate, low-volume isonitrogenous supplement (e.g., 4 g BCAA or EAA/meal) would have produced a similar result. Enthusiasm for an amino acid cocktail is further diminished by palatability, cost and compliance issues.

Given the right circumstances it seems clear that leucine supplementation can acutely increase muscle protein synthesis. Unfortunately however, there is too often a disconnect between positive acute metabolic changes and longer-term improvement in outcomes that really matter (e.g. muscle mass, strength). In the current study, 2-weeks is obviously a very short period of time to observe a measurable change in muscle mass. However, the magnitude of the increase we observed in basal and postabsorptive muscle protein synthesis during this period should have mathematically increased muscle mass by approximately 4% – conservatively.
Assuming no change in muscle protein breakdown. Moving forward, it is clear that additional steps must be taken to clarify and define groups of individuals who will, and importantly, will not benefit from longer-term leucine supplementation.

In conclusion, we have shown that leucine supplementation increases the potential for muscle protein synthesis in older adults and may make an otherwise insufficient or marginal quantity of meal-derived protein more biologically available for muscle tissue growth and repair.

**Statement of authorship**

All authors read and approved the final manuscript. SLC carried out the studies, performed sample and data analysis, and drafted the manuscript. MSM participated in the design of the study. SJH performed analysis of the 3-day diet recall period and daily diet logs. DPJ conceived the study, participated in its design and coordination, and assisted with drafting the manuscript.

**Conflict of interest**

DPJ has participated on Scientific Advisory Panels for the National Dairy Council, the American Egg Board, the National Cattlemen’s Beef Association and Abbott Nutrition.

**Acknowledgments**

This research was supported by a pilot grant (DPJ) from the UTMB Claude D. Pepper Older Americans Independence Center # P30 AG024832. Additional supports was provided by NIH/NCI grant 5R01 CA127971 (MSM), NIH grant T32HD007539, the National Cattlemen’s Beef Association and the National Space Biomedical Research Institute grant NNJ08SA002N (DPJ). Studies were conducted in the Institute for Translational Sciences — Clinical Research Center at UTMB funded by grant 1UL1RR029876-01 from the National Center for Research Resources, National Institutes of Health.

**References**


