Blood $^{15}$N:$^{13}$C Enrichment Ratios Are Proportional to the Ingested Quantity of Protein with the Dual-Tracer Approach for Determining Amino Acid Bioavailability in Humans

Nikkie van der Wielen,1,2 Nadezda V Khodorova,3 Walter JJ Gerrits,2 Claire Gaudichon,3 Juliane Calvez,3 Daniel Tomé,3 and Marco Mensink1

1Division of Human Nutrition, Wageningen University and Research, Wageningen, Netherlands; 2Animal Nutrition Group, Wageningen University and Research, Wageningen, Netherlands; and 3UMR Physiologie de la nutrition et du comportement alimentaire, AgroParisTech, National Research Institute for Agriculture, Food and Environment, Université Paris-Saclay, Paris, France

ABSTRACT

Background: Assessment of amino acid bioavailability is of key importance for the evaluation of protein quality; however, measuring ileal digestibility of dietary proteins in humans is challenging. Therefore, a less-invasive dual stable isotope tracer approach was developed.

Objective: We aimed to test the assumption that the $^{15}$N:$^{13}$C enrichment ratio in the blood increases proportionally to the quantity ingested by applying different quantities of $^{15}$N test protein.

Methods: In a crossover design, 10 healthy adults were given a semi-liquid mixed meal containing 25 g (low protein) or 50 g (high protein) of $^{15}$N-labeled milk protein concentrate simultaneous with 0.4 g of highly $^{13}$C–enriched spirulina. The meal was distributed over multiple small portions, frequently provided every 20 min during a period of 160 min. For several amino acids, the blood $^{15}$N–related to $^{13}$C-isotopic enrichment ratio was determined at $t = 0, 30, 60, 90, 120, 180, 240, 300, and 360$ min and differences between the 2 meals were compared using paired analyses.

Results: No differences in $^{13}$C AUC for each of the measured amino acids in serum was observed when ingesting a low- or high-protein meal, whereas $^{15}$N AUC of amino acids was $\sim 2$ times larger on the high-protein meal ($P < 0.001$).

Doubling the intake of $^{15}$N-labeled amino acids increased the $^{15}$N:$^{13}$C ratio by a factor of 2.04 ± 0.445 for lysine and a factor between 1.8 and 2.2 for other analyzed amino acids, with only phenylalanine (2.28), methionine (2.48), and tryptophan (3.02) outside this range.

Conclusions: The amino acid $^{15}$N:$^{13}$C enrichment ratio in the peripheral circulation increased proportionally to the quantity of $^{15}$N-labeled milk protein ingested, especially for lysine, in healthy adults. However, when using $^{15}$N-labeled protein, correction for, e.g., $\alpha$-carbon $^{15}$N atom transamination is advised for determination of bioavailability of individual amino acids. This trial was registered at www.clinicaltrials.gov as NCT02996704. J Nutr 2020;150:2346–2352.

Keywords: protein digestibility, stable isotopes, dual-tracer approach, amino acid bioavailability, milk protein, spirulina

Introduction

A critical factor in protein quality assessment is amino acid (AA) bioavailability, which represents the fraction of AA available to the organism after digestion and absorption (1, 2). It is usually derived from ileal AA digestibility—that is, digestibility measured at the terminal ileum after ingestion of a protein. As it is assumed that AAs are absorbed exclusively in the small intestine and that the fraction entering the large intestine is not absorbed in nutritionally relevant quantities (3–5). A promising non-, or minimally, invasive approach for measuring AA bioavailability in humans is the dual stable isotope tracer approach (4, 6, 7).

This dual stable isotope tracer method has been applied for measuring the bioavailability of AAs from different intrinsically labeled proteins in humans (8, 9). The method relies on the ingestion of a meal containing a test protein of unknown AA bioavailability, intrinsically labeled with $^2$H or $^{15}$N, together with a tracer dose of a $^{13}$C-labeled protein or standard free...
AA mixture of known AA bioavailability. The comparison of the differential isotopic enrichment (\(^{15}\)N or \(^{2}\)H vs \(^{13}\)C) of each AA in the blood and in the meal reflects, when correcting for label transfer, for example, due to transamination, the relative absorption of each AA that originates from the 2 labeled proteins.

Even though the dual-tracer method has been applied in several studies (6, 8, 10) and its results were comparable with bibliographic true ileal or apparent fecal digestibility values in pigs and humans (9, 11), the approach has not been validated. The approach is based on several assumptions. One of these assumptions is that the \(^{15}\)N:\(^{13}\)C enrichment ratio for each AA in blood is identical to the ratio after absorption and that postabsorptive metabolism, including isotopic fractionation, does not influence this ratio. One can test this assumption by varying the quantity of bioavailable protein. Varying the protein quantity provided or its bioavailability should result in a proportional response of this ratio in peripheral blood.

In this study we applied the dual-tracer approach for AA bioavailability using \(^{15}\)N-milk protein concentrate together with \(^{13}\)C-spirulina as test and standard protein, respectively. These protein sources were chosen as their AA digestibility is known (8, 12). The aim was to test the critical assumption of the method that variation in bioavailability should be proportionally reflected in changes in isotope ratios. Therefore, different quantities of test protein were applied to validate whether the \(^{15}\)N:\(^{13}\)C enrichment ratio of AAs in the peripheral circulation increased proportionally to the quantity of \(^{15}\)N-labeled milk protein concentrate ingested while keeping the quantity of \(^{13}\)C spirulina identical.

### Study population

The study population consisted of 10 healthy subjects (5 males, 5 females), with an age between 20 and 35 y. Habitual dietary intake was measured with a food-frequency questionnaire, and general health was assessed using a basic questionnaire. Height and body weight were measured. To be included in the study, subjects needed to have a BMI (kg/m\(^2\)) between 18.5 and 25 and a habitual dietary protein intake between 10% and 30% of total energy intake. Subjects were excluded when having a history of medical or surgical events or a disease that might influence study outcomes, using medical drugs except for an alcohol consumption that might influence study outcomes, using medical drugs except for physical activity for >5 h/wk, reporting weight loss or gain of >3 kg in the last month, reporting a slimming or medical diet, or being pregnant or breastfeeding. All subjects gave their written informed consent. The included subjects were 23.5 ± 3.2 y old, had a BMI of 22.2 ± 0.58, and habitual protein intake of 14.4% ± 2.5% of total energy intake.

### Study design

A randomized crossover design was applied with 2 different experimental meals, providing either 25 or 50 g of \(^{15}\)N-enriched milk proteins. These meals were provided on 2 different test days, which were separated by 1 wk. Order of the meals was randomized. The day prior to each test day subjects were asked to refrain from vigorous exercise and foods naturally enriched in \(^{13}\)C, like corn or pineapple. Furthermore, subjects were asked to copy their dietary behavior from the first test day to the second test day. For practical reasons, the study was executed in 2 parts: the first part, executed in August 2016, included 4 participants (2 males, 2 females) and the second part, executed in November 2016, included 6 participants (3 males, 3 females).

### Experimental meals

To produce \(^{15}\)N-labeled milk protein concentrate, 2 Holstein Friesian dairy cows were rumen-infused with \(^{15}\)N-ammonium sulfate (98% enriched; Cambridge Isotope Laboratories, Inc.) and milk protein was extracted. The protein content of the \(^{15}\)N-labeled milk protein concentrate was 67.9% (N × 6.38) and the total \(^{15}\)N enrichment was 2.27 atom percent (AP) or 1.90 atom percent excess (APE), as determined using an elemental analyzer coupled to isotope ratio mass spectrometer. The 2 experimental meals consisted of either 25 g or 50 g of \(^{15}\)N-labeled milk protein concentrate providing 18.2 g (low-protein meal) and 35.5 g (high-protein meal) of crude protein, respectively. Both meals contained 400 mg U\(^{13}\)C-labeled whole spirulina (>98% enriched (98%AP); Cambridge Isotope Laboratories, Inc.). The commercially obtained \(^{13}\)C-labeled whole spirulina (>98% enriched; Cambridge Isotope Laboratories, Inc.) consisted of 39.6% (N × 6.25) protein, 2.83% fat, 23.2% neutral detergent fiber, 4.94% ash, and 2.30% moisture. It was produced under quality standards: ISO Guide 34, ISO/IEC 17025, ISO 13485, current good manufacturing practice. The low-protein and high-protein meal also contained 55 g or 32 g dextrose-maltose (Fantomalt; Nutricia), 50 g concentrations of lemonade syrup (Karvan cevitam, Cassis), 5.5 g or 5.0 g sunflower oil, 12 g or 7 g locust bean gum (Johannesbroodpitmeel), and 338 g or 341 g water, respectively. Both mixed meals were similar in energy content (513 kcal for the low-protein meal and 514 kcal for high protein) and total weight; moreover, no visual differences in color or viscosity were noticed. Nutrient composition was different between the meals, with the high-protein meal containing 27.8% of energy (E%) protein and 61.9 E% carbohydrates and the low-protein meal contained 14.3 E% protein and 75.4 E% carbohydrates. Both meals contained 10.3 E% fat.

### Details of the test day

On the test days, subjects arrived at the human research facilities of Wageningen University after an overnight fast of ≥10 h. After measurement of body weight, a catheter was placed in the antecubital vein and baseline blood samples were taken. Subsequently, the subjects ingested the semi-liquid meal, divided in 9 portions, every 20 min (t = 0, 20, 40, 60, 80, 100, 120, 140, and 160 min). Postprandial blood samples were taken at t = 30, 60, 90, 120, 180, 240, 300, and 360 min after the first portion of the semi-liquid meal was consumed. Blood serum samples were used to determine isotopic enrichment of free AAs. At each time point, 16 mL of blood was collected in two 8-mL SST II Advance blood collection tubes (BD Biosciences). After 30–60 min of clotting, the tubes were centrifuged (1550 × g, 10 min, room temperature) and subsequently serum was placed into aliquots and stored at −80 °C until measurement.

\(^{15}\)N and \(^{13}\)C AA enrichments

The meal samples were hydrolyzed (6 M HCl at 110°C for 24 h) and filtered (0.22 μm) prior to the extraction. To obtain a proper estimate for meal enrichment, the meals of all study days were analyzed in duplicate. Serum samples were acidified using 1 M HCl and directly mixed with the resin. Serum free AAs were extracted using cation-exchange Dowex AG50×8 resin conditioned under H\(^{+}\) form and eluted with 6 M NH\(_4\)OH. The purified AAs were analyzed by Gas chromatography combustion isotope ratio mass spectrometry.
as N(O)-ethoxycarbonyl ethyl ester derivatives. Briefly, 3.2 mL of an ethanol:pyridine solution (80:20, vol:vol) were added to 4 mL of the aqueous sample, the resulting mixture was vortexed before the addition of 400 μL ethyl chloroformate, and the solution gently shaken several times until no bubbles were formed. Then, 2 mL of dichloromethane:hexane (50:50, vol:vol) with 1% ethyl chloroformate (vol:vol) was added and the solution was vigorously vortexed. The upper organic phase was collected and evaporated under stream N2. The obtained residue was diluted in 50 μL of ethylacetate and placed in chromatographic vials, and the samples were stored at −20°C until injection. The 15N and 13C enrichments of derivatized AAs were determined using an Agilent 7890B gas chromatograph (Agilent Technologies) coupled to an Isoprime isotope ratio mass spectrometer (Isoprime; GV Instrument) via the GC5 Isoprime interface. The temperature in the interface was regulated at 350°C and combustion oven was maintained at 930°C and 850°C for 15N and 13C analysis, respectively. A 30-m Rxi-17 capillary column (0.25 mm i.d. and 0.5-μm film thickness; Restek) was used in a constant flow mode, and the high-purity helium was used as the carrier gas at a flow rate of 1.2 mL/min. The inlet temperature was set at 270°C. Samples (2 μL) were injected in splitless mode for 15N analysis and in split mode (10:1) for 13C analysis. The temperature program started at 150°C, increased to 200°C by 4°C/min, and then to 270°C by 25°C/min. The temperature was maintained for 20 min at 270°C. The stable isotopic compositions of nitrogen and carbon were measured using the conventional delta notation: the 15N and δ13C values were expressed relative to the international standards (AIR-N2 and PDB, respectively). In serum, the enrichment (AP) of 15N and 13C was corrected for the baseline (t = 0) enrichment of each individual AA to obtain APE. To obtain APE, in the meal the enrichment of 15N and 13C was corrected for overall background protein 15N and 13C abundance determined using an elemental analyzer coupled to an isotope ratio mass spectrometer and measured in the unlabeled milk collected from cows before the start of label infusion.

**Calculations and statistical analyses**

The postprandial response in plasma AA 15N and 13C enrichment (APE) is given for each of the analyzed AAs. To quantify the postprandial response, the AUC of plasma AA enrichment over the entire study day (t = 0–360 minutes) was calculated using the trapezoid rule. The AUC was chosen as the main response parameter in the absence of a clear steady state in our experiment.

The blood-to-meal AA 15N- to 13C-enrichment ratio (R1) between the 2 protein sources was calculated according to the following formulas below:

\[
R_{\text{meal},i} = \frac{15\text{NAA}_{i}/13\text{CAA}_{i}}{\text{R1}}
\]  
\[
R_{\text{meal},i}\text{ is the meal enrichment ratio of the } i \text{ AA, with } 15\text{NAA} \text{ and } 13\text{CAA} \text{ the } 15\text{N}-\text{AA} \text{ and } 13\text{C}-\text{AA} \text{ enrichments of the } i \text{ AA (in APE).}
\]

\[
R_{\text{blood},i} = \frac{15\text{NAAUC}_{i}/13\text{CAAUC}_{i}}{\text{R2}}
\]  
\[
R_{\text{blood},i}\text{ is the blood enrichment ratio of the } i \text{ AA, with } 15\text{NAAUC} \text{ AA and } 13\text{CAAUC} \text{ AA the AUC values of isotopic enrichment of a specific AA (in APE). Finally, } R_{1}\text{ is the blood-to-meal } 15\text{N-} \text{to } 13\text{C}-\text{enrichment ratio of the } i \text{ specific AA.}
\]

\[
R_{1,i} = R_{\text{blood},i}/R_{\text{meal},i}
\]  
\[
R_{1,i}\text{ depends for each AA on the ratio between AA digestibility of milk protein and spirulina.}
\]

Because of a potential label transfer (e.g., the exchange of 15N, but not of 13C) between indispensable AAs through transamination of the α-carbon 15N, digestibility can either be over- or underestimated using R1. Therefore, the R1 value for each AA was compared with R2 as follows:

\[
R_{3,i} = \text{AA digestibility milk protein/AA digestibility spirulina}
\]  

R2 is the relative digestibility of the i AA between milk protein and spirulina of the i AA as obtained from the literature (8, 12).

Comparing R1 to R2 allows estimation of a transamination correction factor (TcF):

\[
\text{TcF} = \frac{R_{1}}{R_{2}}
\]

When TcF = 1 there is no label transfer due to, for example, transamination; with TcF < 1 there is label transfer with a gain in 15N compared with 13C; and with TcF > 1 there is label transfer with a reduction in 15N compared with 13C.

All data are expressed as means ± SDs. Differences in enrichment of the 2 meals were tested using a t test with differences being significant when P < 0.05. Within-subject differences in AUC and Rblood values between the 2 test meals for each label and each AA were compared using a paired t test. The 95% CI was used to check if the difference in Rblood between the high- and low-protein meal was 2-fold.

**Results**

15N- and 13C-AA meal enrichment and Rmeal calculation

15N and 13C enrichment (APE) of AAs in the meal and the ratio 15N:13C enrichment of the AAs in the meal (Rmeal) are given in Table 1. The 15N enrichment was numerically similar in both meals and did not differ significantly for all AAs except for tryptophan and phenylalanine (P < 0.05; Table 1). The 13C enrichment was lower on the high-protein meal compared with the low-protein meal for all AAs, except for lysine (P = 0.057) and methionine (P = 0.128), due to dilution of the 13C-labeled AAs by carbons from the milk protein AAs.

Appearance of 15N- and 13C-labeled AA in blood and Rblood calculation

The appearance of 13C and 15N enrichment in the blood AA pool (Figure 1) showed that 15N enrichment was numerically lower after ingestion of 25 g compared with 50 g 15N-labeled milk protein concentrate, whereas no visual difference was observed on 13C enrichment. Enrichment of both 13C and 15N AAs steadily increased during consumption of the multiple small portions and peaked at 180 min. In the period after the ingestion of the multiple small portions, towards the end of the study day (t = 240–360 min), a decrease in labeled AAs in blood was observed for most AAs. The AUC 13C and 15N enrichment and Rblood of the different circulating AAs are reported in Table 2. Comparing both meals, no significant differences in the AUC for 13C enrichment for each of the AAs was observed, whereas the AUC of 15N AAs in serum was significantly larger on the high-protein meal compared with the low-protein treatment for all AAs except for tryptophan. Doubling the intake of 15N-labeled AAs increased the 15N:13C ratio by a factor varying between 1.8 and 2.2 for the analyzed AAs, with only phenylalanine (2.26), methionine (2.48), and tryptophan (3.02) outside this range. For lysine, the fold-difference between the meals was numerically 2. Other AAs tended to deviate numerically from the expected 2-fold difference, even though the factor 2 was in the CI of each AA due to high variation between individuals.

**Blood-to-meal 15N:13C enrichment ratio (R1) and estimation of 15N transamination for each AA (TcF)**

R1 for AAs was in the range from 0.544 to 3.44 (Table 3). For the different AAs, R1 depends on the difference in the
TABLE 1  Enrichment of individual amino acids in the semi-liquid mixed meals with 25 g (low protein) or 50 g (high protein) of
$^{15}$N-labeled milk protein concentrate with 0.4 g of $^{13}$C-labeled spirulina$^1$

| Amo acid | $^{15}$N enrichment (APE) | $^{13}$C enrichment (APE) | R$_{meal} = ^{15}$N/$^{13}$C |
|----------|---------------------------|---------------------------|---------------------|
| Ala      | 1.66 ± 0.091              | 1.83 ± 0.058              | >0.05               |
| Gly      | 1.45 ± 0.084              | 1.58 ± 0.133              | >0.05               |
| Val      | 1.61 ± 0.050              | 1.65 ± 0.087              | >0.05               |
| Leu      | 1.53 ± 0.028              | 1.58 ± 0.024              | >0.05               |
| Ile      | 1.71 ± 0.118              | 1.90 ± 0.060              | >0.05               |
| Thr      | 1.28 ± 0.063              | 1.80 ± 0.232              | >0.05               |
| Phe      | 1.53 ± 0.029              | 1.61 ± 0.014              | <0.05               |
| Lys      | 2.03 ± 0.071              | 2.07 ± 0.022              | >0.05               |
| Met      | 1.54 ± 0.083              | 1.76 ± 0.141              | >0.05               |
| Trp      | 0.87 ± 0.027              | 0.94 ± 0.034              | <0.05               |
|          | Low protein               | High protein              |                     |
|          | 1.33 ± 0.120              | 0.79 ± 0.062              | <0.001              |
|          | 1.53 ± 0.123              | 0.97 ± 0.061              | <0.001              |
|          | 0.59 ± 0.050              | 0.33 ± 0.022              | <0.001              |
|          | 0.53 ± 0.049              | 0.30 ± 0.019              | <0.001              |
|          | 0.62 ± 0.054              | 0.34 ± 0.020              | <0.001              |
|          | 0.31 ± 0.031              | >0.05                     |
|          | 0.40 ± 0.064              | 0.22 ± 0.026              | <0.01               |

$^{1}$The table shows the mean ± SD of meals provided on different test days, $n = 4$ for $^{13}$C and $n = 3$ for $^{15}$N. APE, atom percent excess.

Discussion

The dual-tracer approach has been proposed to measure food protein AA bioavailability by using a labeled ($^{15}$N or $^2$H) test protein together with a tracer dose of labeled ($^{13}$C) standard protein or AA mixture in a test meal (4). This approach was shown to be a minimal invasive technique to measure dietary AA digestibility in adults, children, and cystic fibrosis patients (6, 10, 13). This dual-tracer approach was applied using $^2$H-labeled test protein in combination with $^{13}$C-labeled standard spirulina (8, 9, 11) as well as with $^{15}$N-labeled spirulina protein in combination with $^2$H-labeled phenylalanine (6). In this context, the present study evaluated the potential of using $^{15}$N-labeling of milk protein with $^{13}$C spirulina...
for measuring dietary protein AA bioavailability in humans. Variation in bioavailability should be proportionally reflected in changes in isotope ratios. The validity of this approach is difficult to test with protein sources differing in bioavailability as, by definition, variations in bioavailability between such sources and methodology (dual isotope technique vs. true ileal AA disappearance) have to be combined. To circumvent this problem, we tested this crucial assumption by doubling the provision of a highly digestible protein source.

We evaluated the effect of the quantity of 15N-labeled test milk protein when applying the dual-tracer method on blood isotopic 15N and 13C-AA enrichment by providing 25 g or 50 g of 15N-labeled milk protein concentrate with 0.4 g of 13C-labeled spirulina. Doubling the quantity of milk protein provided was intended to double the quantity absorbed. Increasing the quantity of protein in a meal does not have the measured blood isotopic 15N:13C ratio is not affected by postabsorptive metabolism.

No difference was observed in the 13C enrichment of circulating AAs between the low- and high-protein meal, which both contained an equal quantity of 13C-labeled spirulina. As expected, the 15N AUC was higher after ingestion of the high-protein meal compared with the low-protein meal. Most importantly, the 2-fold difference for Rblood between these meals for lysine confirmed our idea that the 15N:13C ratio in the blood is proportional to the quantity of AAs absorbed. Furthermore, for all other AAs no significant deviation from 2 was observed, although this might be related to the high variation, as phenylalanine, methionine, and tryptophan are numerically very deviating. Thereby the results validate a crucial assumption in the application of the dual-isotope approach that the isotopic appearance of AAs is proportional to the absorbed quantities, especially for lysine. Thus, potential differences in postabsorptive metabolism for 15N- and 13C-labeled AAs are not affected by the quantity absorbed (7). This observation is also critical for the future application of the dual-tracer approach as it was unclear whether the relative digestibility estimated with the dual-tracer approach would be affected by a quantity of protein provided.

In previous dual-tracer studies, similar protein quantities (21–24 g vs 18.2 g in the present study for the low-protein meal)

### TABLE 2

|  | 15N AUC | 13C AUC | Rblood = 15N AUC/13C AUC | Lower and upper 95% CI (high/low protein) |
|---|---|---|---|---|
| | Low protein | High protein | P | Low protein | High protein | | |
| Ala | 44.2 ± 7.65 | 83.8 ± 13.1 | <0.001 | 19.8 ± 2.87 | 21.3 ± 4.48 | >0.05 | 2.28 ± 0.47 | 4.00 ± 0.69 | <0.001 | 1.81 ± 0.44 | 1.55 ± 2.09 |
| Gly | 37.5 ± 8.70 | 69.1 ± 14.7 | <0.001 | 18.1 ± 4.37 | 17.3 ± 5.18 | >0.05 | 2.10 ± 0.41 | 4.33 ± 0.66 | <0.001 | 2.10 ± 0.58 | 1.70 ± 2.50 |
| Val | 82.9 ± 14.1 | 151.6 ± 25.0 | <0.001 | 37.9 ± 6.75 | 33.3 ± 5.38 | >0.05 | 2.21 ± 0.27 | 4.57 ± 0.43 | <0.001 | 2.11 ± 0.44 | 1.84 ± 2.38 |
| Leu | 62.9 ± 11.5 | 125.0 ± 18.9 | <0.001 | 32.4 ± 4.51 | 29.8 ± 4.83 | >0.05 | 1.94 ± 0.27 | 4.22 ± 0.31 | <0.001 | 2.21 ± 0.39 | 1.97 ± 2.46 |
| Ile | 73.4 ± 15.9 | 142.3 ± 25.1 | <0.001 | 48.3 ± 7.91 | 42.7 ± 5.94 | >0.05 | 1.51 ± 0.19 | 3.33 ± 0.38 | <0.001 | 2.19 ± 0.35 | 1.95 ± 2.42 |
| Thr | 42.9 ± 10.7 | 89.6 ± 20.5 | <0.001 | 27.1 ± 5.53 | 26.6 ± 5.16 | >0.05 | 1.75 ± 0.45 | 3.46 ± 0.95 | <0.001 | 1.92 ± 0.58 | 1.45 ± 2.39 |
| Phe | 63.6 ± 12.4 | 125.7 ± 27.8 | <0.001 | 28.6 ± 4.84 | 25.8 ± 5.27 | >0.05 | 2.23 ± 0.36 | 4.91 ± 0.78 | <0.001 | 2.28 ± 0.59 | 1.89 ± 2.64 |
| Lys | 108 ± 15.9 | 190 ± 17.1 | <0.001 | 19.9 ± 2.66 | 17.8 ± 4.05 | >0.05 | 5.33 ± 0.30 | 11.0 ± 2.02 | <0.001 | 2.04 ± 0.44 | 1.76 ± 2.33 |
| Met | 72.6 ± 20.2 | 150 ± 36.2 | <0.001 | 9.3 ± 7.80 | 14.6 ± 16.0 | >0.05 | 10.0 ± 4.69 | 19.5 ± 12.3 | <0.05 | 2.48 ± 1.85 | 1.10 ± 3.85 |
| Trp | 31.6 ± 7.39 | 62.1 ± 3.55 | <0.001 | 7.91 ± 7.50 | 7.68 ± 4.50 | >0.05 | 5.26 ± 3.35 | 10.0 ± 4.51 | >0.05 | 3.02 ± 3.02 | 0.376 ± 0.56 |

1Values are means ± SDs unless otherwise indicated, n = 10. The meal was frequently distributed in 9 discrete portions given every 20 min. Rblood for both meals was determined in 9 discrete portions given every 20 min. Rblood for both meals was determined in 9 discrete portions given every 20 min.

### TABLE 3

| Digestibility, % | R1 = Rblood/Rmeal | R2 | TCF = R2/R1 |
|---|---|---|---|
| Milk8 | Spirulina8 | Low protein | High protein | Low protein | High protein |
| Ala | 95.9 | — | — | 1.84 ± 0.377 | 1.72 ± 0.273 | — | — |
| Gly | 99.3 | — | — | 2.21 ± 0.434 | 2.66 ± 0.403 | — | — |
| Val | 93.4 | 87.1 | 1.07 | 0.82 ± 0.101 | 0.92 ± 0.087 | 1.32 ± 0.155 | 1.17 ± 0.100 |
| Leu | 95.0 | 86.0 | 1.10 | 0.68 ± 0.094 | 0.79 ± 0.060 | 1.65 ± 0.235 | 1.40 ± 0.100 |
| Ile | 95.0 | 84.2 | 1.13 | 0.54 ± 0.070 | 0.61 ± 0.069 | 2.11 ± 0.321 | 1.89 ± 0.204 |
| Thr | 95.4 | 82.5 | 1.16 | 1.11 ± 0.287 | 1.05 ± 0.291 | 1.10 ± 0.273 | 1.18 ± 0.313 |
| Phe | 94.9 | 95.3 | 1.00 | 0.79 ± 0.131 | 0.93 ± 0.148 | 1.29 ± 0.224 | 1.10 ± 0.159 |
| Lys | 95.6 | 77.5 | 1.23 | 1.28 ± 0.072 | 1.39 ± 0.254 | 0.96 ± 0.053 | 0.91 ± 0.162 |
| Met | 91.6 | 84.1 | 1.09 | 2.97 ± 1.390 | 3.44 ± 2.170 | 0.42 ± 0.145 | 0.57 ± 0.544 |
| Trp | — | — | — | 2.39 ± 1.521 | 2.37 ± 1.074 | — | — |

1Values are means ± SDs unless otherwise indicated, n = 10. The blood-to-meal 15N/13C ratios (R1) obtained in healthy participants who received 25 g (low protein) or 50 g (high protein) of 15N-labeled milk protein concentrate with 0.4 g of 13C-labeled spirulina are shown. Literature data on milk protein and spirulina amino acid digestibility were used to calculate R1 and subsequently TCF. TCF, transamination correction factor.

2Data from reference 12

3Data from reference 8.
were provided in a meal with higher energy content (714–834 kcal vs this study’s meal of 513 kcal) (9, 11). Therefore, our study’s protein-to-energy ratio was only slightly higher for the low-protein meal (14.3 E% protein) but markedly higher for the present high-protein meal (27.8 E% protein) compared with other dual-tracer studies. As it is well known that the amount of carbohydrates in a protein meal can influence the retention of AAAs in the body (17), this could theoretically have influenced postabsorptive metabolism of labeled AAAs and, in the case of isotopic fractionation, result in deviating labeled AA fluxes (18). However, here we showed that postabsorptive metabolism of labeled AAAs does not influence the proportional appearance of labeled lysine and potentially other AAAs.

Unlike earlier work using the dual-tracer approach (8) we used the combination of $^{15}$N test protein and $^{13}$C reference protein to test the critical assumption of proportional isotope ratios. Depending on the protein source, labeling with $^{15}$N is more convenient and efficient than $^2$H labeling, although it comes with additional challenges. In the dual-isotope approach with a $^{13}$C-labeled reference protein, there is for most AAAs no exchange of $^{13}$C for the $^{12}$C-AAAs absorbed from the reference $^{12}$C-labeled protein, and the $^{13}$C enrichment of circulating AAAs is only related to its digestibility and absorption, except for methionine. In contrast, for the $^{15}$N-labeled test protein, the $^{15}$N enrichment of circulating AAAs is proportional to the digestibility and absorption of the AA from the test protein but also to the level of $\alpha$-carbon $^{15}$N atom transamination of the AAAs. Transamination is a complex process of AA metabolism that is active in different tissues and differently affects AAAs (19–21). Assuming that transamination is the main source of discrepancy between predicted (i.e., $R_2$) and observed relative digestibility (i.e., $R_1$) between our 2 protein sources, TcFs could be estimated. The TcF was calculated by comparing the relative digestibility measured in this study ($R_1$) with that calculated from data in the literature ($R_2$), although these literature values were determined with different methods and in different populations (8, 12). The values of the transamination factors ranged from 0.42 to 2.11 for $^{15}$N atom transamination. It is important to realize that these TcFs are much higher than those calculated for loss of deuterium ($^2$H) from indispensable AAAs that ranged from 1.002 to 1.081 (8). This is unavoidable because only 1 hydrogen atom out of multiple hydrogen atoms on the AA molecule is lost during transamination in comparison with the loss (or gain) of the only nitrogen atom in many AAAs. The estimated TcFs >1 indicate that for the branched-chain AAAs valine, leucine, and isoleucine $^{15}$N compared with $^{13}$C is lost. This is substantiated by the fact that, even though the transamination of branched-chain AAAs is reversible, the likeliness of re-amination of the $\alpha$-keto-acid with labeled nitrogen is virtually nil because of the rate of the process and the large amino-nitrogen pool size (18, 22). In contrast, for methionine, a transamination factor $<1$ was estimated from our data, suggesting an increased $^{15}$N relative to $^{13}$C, which may be caused by homocysteine remethylation (23). From our results it is clear that measuring the AA bioavailability of a test protein using $^{15}$N-labeling of the test protein requires a specific correction factor for most AAAs. The rate of label transfer due to metabolism such as transamination and the appropriate correction factors for future studies need to be quantified before the dual-tracer approach can be routinely applied using $^{15}$N-labeled proteins. However, for AAAs that do not undergo transamination, lysine and threonine, a correction factor for transamination may be less critical for the right determination of AA bioavailability than for other indispensable AAAs such as the branched-chain AAAs.

Our results also indicate other aspects that are essential for future applications of the dual-tracer approach. The $R_{\text{meal}}$ was expected to be 2 times larger for the high-protein meal compared with the low-protein meal. The enrichments measured in our meal showed a high level of variation, especially for $^{13}$C, most likely due to flaws in the homogeneity of the meal and subsampling portions for analysis, as mixing small quantities of highly labeled spirulina within a semi-liquid meal does not necessarily lead to a homogeneous distribution. Moreover, the values differed slightly from our expectations; however, this could also be due to uncertainty in AA enrichment of spirulina and the unknown AA content and enrichment of the thickening agent. As the primary aim of the current study was not to calculate digestibility of the test protein, this uncertainty in our meal data did not influence our observation that the $^{15}$N:$^{13}$C enrichment ratio in the blood increased proportionally to the quantity of $^{15}$N-labeled protein ingested. However, it affected the size of the blood-to-meal ratio ($R_1$) and the TcF values; therefore, no quantitative comparisons between low- and high-protein meals in $R_1$ of TcF were made.

The pattern of frequently feeding small meals prevented the potential effect of large differences in the intestinal kinetics (24, 25) between spirulina and milk protein and led to a parallel kinetic response of isotope enrichment in blood AAAs. Accordingly, the pattern of $^{15}$C and $^{13}$N enrichment of circulating AAAs over time showed quite similar profiles, although with a slightly faster decrease in $^{15}$N enrichment after the last meal. However, we did not obtain an isotopic plateau, probably because of a too-short meal feeding period or a lack of an initial priming bolus as applied by Devi et al. (8). Due to a lack of plateau, AUC was chosen for calculations. However, it is uncertain whether this issue, analytical variations for the chosen isotopic labels, or day-to-day subject variation resulted in the relatively high variation that was observed in Table 2. Moreover, the high variation for some AAAs may have caused a lack in statistical power to show differences due to the quantity of test protein. As a result, the estimated TcFs also show high variability, especially compared with the previously reported $<$1% variation in $^2$HTcFs (8).

The calculations with the dual-tracer approach rely on multiple ratios; therefore, small deviations in isotopic enrichment in blood samples may have large effects on the final result of the digestibility calculation. For example, a 5% increase or decrease in $^{13}$N or $^{15}$N enrichment in the blood alone could result in an $\sim$10% increase or $\sim$10% decrease in $R_1$, and with similar variations in the meal enrichment this will be even larger. A slightly higher SD in mean indispensable AA digestibility value was also shown by Kashyap et al. (9), where digestibility values obtained with the dual-tracer approach were compared with bibliographic true ileal or apparent fecal digestibility values in pigs and humans. This potential variation should be taken into account when designing studies.

In conclusion, this study demonstrated that the $^{15}$N:$^{13}$C enrichment ratio of AAAs in the peripheral circulation increased proportionally to the quantity of $^{15}$N-labeled milk protein concentrate ingested, especially for lysine, indicating that postabsorptive metabolism is similar for an AA with either a $^{13}$C or $^{15}$N label, and that the ratio of labels in the blood is proportional and thereby representative for absorption. This illustrates the validity of the dual-isotope method to measure relevant variation in the bioavailability.

However, applying this dual-tracer approach with $^{15}$N to determine bioavailability of AAAs from food proteins requires
correction factors that are especially critical for AAs with high transamination rates and less essential for AAs that do not undergo transamination like threonine and lysine. The correction factors estimated in the current study need further quantification and validation before application.

Acknowledgments
The authors’ responsibilities were as follows—DT, MM, CG, and WJJG: designed the research; NvdW, NVK, and JC: conducted the research; NvdW and MM: analyzed data; NvdW, MM, and DT: wrote the manuscript; and all authors: read and approved the final manuscript.

References

1. Tome D, Jahoor F, Kurpad A, Michaelensen KF, Pencharz P, Slater C, Weisell R. Current issues in determining dietary protein quality and metabolic utilization. Eur J Clin Nutr 2014;68:537–8.
2. Tome D. Digestibility issues of vegetable versus animal proteins: protein and amino acid requirements—functional aspects. Food Nutr Bull 2013;34:272–4.
3. Fuller MF, Tomé D. In vivo determination of amino acid bioavailability in humans and model animals. J AOAC Int 2005;88:923–34.
4. Lee WT, Weisell R, Albert J, Tome D, Kurpad AV, Uauy R. Research approaches and methods for evaluating the protein quality of human foods proposed by an FAO Expert Working Group in 2014. J Nutr 2016;146:929–32.
5. van der Wielen N, Moughan PJ, Mensink M. Amino acid absorption in the large intestine of humans and porcine models. J Nutr 2017;147(8):1493–8.
6. Engelen MP, Com G, Anderson PJ, Deutz NE. New stable isotope method to measure protein digestibility and response to pancreatic enzyme intake in cystic fibrosis. Clin Nutr 2014;33:1024–32.
7. Tome D. Editorial on “Measurement of protein digestibility in humans by a dual tracer method”—a key limiting factor of protein quality. Am J Clin Nutr 2018;107:855–6.
8. Devi S, Varkey A, Sheshishayee MS, Preston T, Kurpad AV. Measurement of protein digestibility in humans by a dual-tracer method. Am J Clin Nutr 2018;107:984–91.
9. Kashyap S, Shivakumar N, Varkey A, Duraisamy R, Thomas T, Preston T, Devi S, Kurpad AV. Ileal digestibility of intrinsically labeled hen’s egg and meat protein determined with the dual stable isotope tracer method in Indian adults. Am J Clin Nutr 2018;108:980–7.
10. Kashyap S, Shivakumar N, Varkey A, Preston T, Devi S, Kurpad AV. Co-ingestion of black tea reduces the indispensable amino acid digestibility of hens’ egg in Indian adults. J Nutr 2019;149:1363–8.
11. Kashyap S, Varkey A, Shivakumar N, Devi S, Reddy BHR, Thomas T, Preston T, Sreeman S, Kurpad AV. True ileal digestibility of legumes determined by dual-isotope tracer method in Indian adults. Am J Clin Nutr 2019;110(4):873–82.
12. Gaudichon C, Bos C, Morens C, Petzke KJ, Mariotti F, Everwand J, Benamouzig R, Dare S, Tome D, Metges CC. Ileal losses of nitrogen and amino acids in humans and their importance to the assessment of amino acid requirements. Gastroenterology 2002;123:50–9.
13. Shivakumar N, Kashyap S, Kishore S, Thomas T, Varkey A, Devi S, Preston T, Jahoor F, Sheshishayee MS, Kurpad AV. Protein-quality evaluation of complementary foods in Indian children. Am J Clin Nutr 2019;109:1319–27.
14. Donkoh A, Moughan PJ. The effect of dietary crude protein content on apparent and true ileal nitrogen and amino acid digestibilities. Br J Nutr 1994;72:59–68.
15. Bax ML, Buffiere C, Hafnaoui N, Gaudichon C, Savary-Auzeloux I, Dardevet D, Sante-Lhoutellier V, Remond D. Effects of meat cooking, and of ingested amount, on protein digestion speed and entry of residual proteins into the colon: a study in minipigs. PLoS One 2013;8:e61252.
16. Fan MZ, Sauer WC, McBurney MI. Estimation by regression analysis of endogenous amino acid levels in digesta collected from the distal ileum of pigs. J Anim Sci 1995;73:2319–28.
17. Deutz NE, Ten Have GA, Soeters PB, Moughan PJ. Increased intestinal amino-acid retention from the addition of carbohydrates to a meal. Clin Nutr 1995;14:334–64.
18. Waterlow JC. Protein turnover. Wallingford (England): CABI Publishing; 2006.
19. Braun A, Vikari A, Windisch W, Auerswald K. Transamination governs nitrogen isotope heterogeneity of amino acids in rats. J Agric Food Chem 2014;62:8008–13.
20. Macko SA, Estep MLF, Engel MH, Hare P. Kinetic fractionation of stable nitrogen isotopes during amino acid transamination. Geochim Cosmochim Acta 1986;50:2143–6.
21. Jungas RL, Halperin ML, Brosnan JT. Quantitative-analysis of amino-acid oxidation and related gluconeogenesis in humans. Physiol Rev 1992;72:419–48.
22. Matthews DE, Bier DM, Rennie MJ, Edwards RH, Halliday D, Millward DJ, Clugston GA. Regulation of leucine metabolism in man: a stable isotope study. Science 1981;214:1129–31.
23. Blom HJ, Smulders Y. Overview of homocysteine and folate metabolism: with special references to cardiovascular disease and neural tube defects. J Inherit Metab Dis 2011;34:75–81.
24. Boirie Y, Dangin M, Gachon P, Vasson MP, Maubois JL, Beaufreire B. Slow and fast dietary proteins differently modulate postprandial protein accretion. Proc Natl Acad Sci USA 1997;94:14930–5.
25. Lacroix M, Leonil J, Bos C, Henry G, Airinei G, Faquiant J, Tome D, Gaudichon C. Heat markers and quality indexes of industrially heat-treated [15N] milk protein measured in rats. J Agric Food Chem 2006;54:1508–17.