A validation of the application of D₂O stable isotope tracer techniques for monitoring day-to-day changes in muscle protein subfraction synthesis in humans

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Wilkinson DJ, Franchi MV, Brook MS, Narici MV, Williams JP, Mitchell WK, Szweczyk NJ, Greenhaff PL, Atherton PJ, Smith K. A validation of the application of D₂O stable isotope tracer techniques for monitoring day-to-day changes in muscle protein subfraction synthesis in humans. Am J Physiol Endocrinol Metab 306: E571–E579, 2014. First published December 31, 2013; doi:10.1152/ajpendo.00650.2013.—Quantification of muscle protein synthesis (MPS) remains a cornerstone for understanding the control of muscle mass. Traditional [¹³C]amino acid tracer methodologies necessitate sustained bed rest and intravenous cannulation(s), restricting studies to acid tracer methodologies necessitate sustained bed rest and intravenous cannulation(s), restricting studies to monitoring day-to-day changes in muscle protein subfraction synthesis in humans.

Innovative Methodology

SKELETAL MUSCLE IS THE BODY’S LARGEST ORGAN and represents an important tissue for maintenance of metabolic health. Muscle not only provides locomotory function but also acts as a major glucose disposal sink and a fuel reservoir for other organs in times of fasting and stress (60). Thus, unsurprisingly, declines in muscle mass with aging (sarcopenia) or during acute or chronic illness [i.e., sepsis, chronic obstructive pulmonary disease, cancer cachexia, and organ failure (31, 52, 56, 59)] lead to frailty and loss of mobility and independence in addition to increased risk of cardiovascular and metabolic disease [e.g., type 2 diabetes; (54)] or, in severe cases, even death (2).

Over recent years, considerable effort has been focused on research into techniques and interventions for understanding the regulation of skeletal muscle mass and minimizing loss of muscle mass with aging and disease (26, 41). Despite this, effective treatments to mitigate muscle loss remain limited. Indeed, despite extensive investment in pharmaceutical interventions (6, 39), the current most effective and safe means by which to maintain or increase muscle mass remains resistance exercise training (RET) with accompanying protein nutrition (11, 19, 25, 29, 40). The hypertrophic response to RET is regulated via cumulative postexercise increases in muscle protein synthesis (MPS) (22, 30, 42), which after several weeks/months leads to muscle hypertrophy that is quantifiable using imaging techniques such as dual-energy X-ray absorptiometry (DEXA), ultrasound, MRI, or CT.

Traditionally, MPS has been measured in vivo using stable isotope tracers of amino acids (AA) with either heavy carbon (¹³C), deuterium (²H), or nitrogen (¹⁵N) motifs. The incorporation of these labeled AAs into muscle protein allows for the calculation of a fractional synthetic rate (FSR) of MPS (45, 61). Although they provide great utility, these methods are not without their inherent limitations. For instance, ¹³C-AA tracers permit measurements to be performed only over short durations (typically <8–12 h) and require preparation of sometimes costly sterile infusions, venous/arterial cannulation, and multiple biopsy collection, all within a controlled laboratory environment. Therefore, there is a need to develop less restrictive and longer-term means to determine MPS in “free-living” humans. It would clearly be advantageous were this method to be sensitive enough to detect changes in MPS and anabolic and/or catabolic scenarios over periods in which it is not plausible to measure changes in mass via imaging methods.

Recent advances in mass spectrometry, in particular the development of pyrolysis-isotope ratio mass spectrometry (IRMS) systems for high-precision measurement of hydrogen...
and oxygen stable isotopes (21, 47), have led to the recent re-introduction of the first stable isotope tracer used in metabolic research, deuterium oxide (D₂O, or “heavy water”) (48, 51). Upon oral ingestion, D₂O equilibrates within the body water pool (20 min in rats (7) and 2 h in humans (24)), and through exchange with hydrogen the deuterium can be incorporated into multiple metabolic pools and tissues. This negates the need for intravenous administration and allows subjects to administer tracer while performing normal activities. D₂O has been successfully integrated into the measurement of protein synthesis in both animal models (7, 9, 12a, 14, 18, 27, 37, 62) and humans (9, 16, 33, 43, 50). Nonetheless, to date few studies have concentrated on its application and validity in the arena of human muscle.

Robinson et al. (50) reported that it was possible to measure MPS over extended periods (i.e., 6 wk) in groups of free-living adults, highlighting its unique suitability for determining longer-term MPS. Furthermore, Gasier et al. (16) showed stimulation of myofibrillar protein synthesis (MyoPS) but not mixed-muscle protein synthesis 24 h following a single exercise bout highlighting the importance of delineating fraction-specific differences. Nonetheless, the authors of that study did not measure responses over longer periods in an attempt to resolve the temporal and cumulative relationship between successive bouts of resistance exercise and MPS, the latter being the key underlying feature regulating muscle hypertrophy. Finally, both the Gasier et al. (16) and Robinson et al. (50) studies used relatively large doses of D₂O to be able to chart MPS due to sensitivity issues involved with gas chromatography-mass spectrometry (GC-MS) as an analytical tool. In comparison, IRMS adds ≥100-fold increases in sensitivity for measuring tissue isotopic enrichment compared with GC-MS, as demonstrated by MacDonald et al. (33), who showed that MyoPS could be measured over a period of 4–14 days following a small single bolus of 100 ml of 70 atom% D₂O in healthy adults under rested conditions.

Our goal was to assess the utility of D₂O for quantifying temporal and cumulative MPS under habitual and stimulated conditions (to assess its potential sensitivity to inform on anabolic/anticatabolic interventions). We employed unilateral exercise (28), where one leg is exercised and the contralateral serves as internal control, to determine the efficacy of D₂O for quantifying cumulative MPS at rest and in response to exercise. We hypothesized that D₂O alongside IRMS techniques could be used to distinguish the synthesis profiles of muscle protein fractions (myofibrillar, collagen, sarcoplasmic) over a period of short-term RET (4 bouts over 8 days). Our rationale for this was that 1) it is unfeasible to use AA tracers over this time, 2) it would inform about diurnal MPS in real-life settings, and 3) it is a period too short to quantify increases in muscle mass as a consequence of RET in healthy, young volunteers.

MATERIALS AND METHODS

Subject characteristics and ethics. Eight young healthy males (22 ± 3.5 yr, body mass index 23.5 ± 0.8 kg/m²) were recruited. All volunteers were screened by means of a medical questionnaire, physical examination, and resting electrocardiogram, with exclusions for metabolic, respiratory, and cardiovascular disorders or other symptoms of ill health. Subjects had normal blood chemistry, were normotensive (blood pressure <140/90), and were not prescribed any medications; all subjects performed activities of daily living and recreation but did not participate routinely in any formal exercise. All subjects gave their written informed consent to participate after all procedures and risks were explained. This study was approved by the University of Nottingham Ethics Committee and complied with the Declaration of Helsinki.

Study procedures. Volunteers were asked to refrain from heavy exercise for the 72 h before the start of the study and perform no exercise over the course of the study other than that prescribed as part of the protocol. One week before the first visit, strength was assessed for the training leg using an isokinetic dynamometer (Icom; Isokinetic Technologies, Eurokinetics, UK), with a one-repetition maximum (1RM) also assessed (Dominant leg; Technogym, Gambettola, Italy). In addition, muscle architecture was assessed by ultrasound (Mylab 70; Esaote Biomedica). The same measurements were performed 8 days later following the final training and biopsy session. Participants then completed a unilateral resistance exercise training program over a period of 8 days; training consisted of single-leg knee extension exercise (4 × 8 reps at 80% 1RM) performed on days 0, 2, 4, 6, and 8. Twenty grams of whey protein isolate (Pro-Isolate Tech; Muscletech) was provided postexercise to support the increased demands of exercise on MPS processes. Bilateral biopsies (nonexercised and exercised) of m. vastus lateralis were taken under sterile conditions, using the conchothermy biopsy technique (13) with 1% lidocaine (B. Braun Melsungen) as local anaesthetic, on days 0 (basal), 2, 4, and 8. A rested biopsy was taken prior to the exercise bout to avoid acute stimulation from the exercise and whey protein feed. Muscle was rapidly dissected free of fat and connective tissue, washed in ice-cold saline, and then frozen in liquid N₂ and stored at −80°C until further analysis. Single venous blood samples were collected on days 0 and 8 into lithium heparin-coated tubes; these were immediately cold centrifuged at 3,200 rpm, and the plasma fraction was then aliquoted and frozen at −80°C until analysis.

Immediately post-biopsy on day 0, participants provided a saliva sample (collected in sterile plastic tubes) and were asked to consume a single 150-ml oral bolus of D₂O (70 atom%; Sigma-Aldrich, Poole, UK), and this was performed with the aim to label the body water pool to ~0.2%. To monitor the body water enrichment throughout the study, each participant was asked to provide a single daily saliva sample collected at midday (time equal ± 30 min after their last meal or drink). These were collected in sterile plastic tubes and kept refrigerated, and participants were asked to bring these to each training session. Upon receipt of saliva samples, they were immediately cold centrifuged at 16,000 g to remove any debris that might have been present; they were then aliquoted into 2-ml glass vials and frozen at −20°C until analysis. A detailed schematic of the study protocol is provided in Fig. 1.

Body water enrichment. Body water enrichment was determined through direct liquid injection of saliva samples (0.1–µl volume) into a high-temperature conversion elemental analyzer (TC/EA; Thermo Finnigan, Thermo Scientific, Hemel Hempstead, UK) connected to an isotope ratio mass spectrometer (IRMS; Delta V Advantage, Thermo). To minimize the effect of carryover between samples, each was injected a minimum of four times. To validate the accuracy of the TC/EA for measuring body water enrichment from saliva, the analysis was repeated with two participants’ saliva sets using GC-Pyrolysis-IRMS (Trace GC isokin Delta V Advantage; Thermo Scientific) and a modification of the protocol by Mahsri et al. (34). Briefly, 100 µl of each saliva sample was incubated with 2 µl of 10 M NaOH and 1 µl of acetone for 24 h at room temperature; this high pH incubation leads to the exchange of deuterium from water with the hydrogen positions on the acetone. Following incubation the acetone was extracted into 200 µl of h-heptane, and 0.5 µl of the heptane phase was injected into the GC for analysis. A standard curve of known D₂O enrichment was run alongside the saliva samples for calculation of enrichment. Further validation was also provided via analysis of water extracted from day 0 and day 8 plasma. In addition, 50- to 100-µl aliquots of plasma were
placed in the cap of inverted autosampler vials; these were then placed on a heating block set at 90°C for 2 h. Water distillate was then collected by rapidly cooling the vials on ice for 10 min and transferred to fresh vials for direct liquid injection into the TC/EA, same as for the saliva.

Isolation and derivatization of myofibrillar, sarcoplasmic, and collagen protein fractions. For isolation of myofibrillar, sarcoplasmic, and collagen fractions, ~30–50 mg of muscle was used. The muscle was homogenized in ice-cold homogenization buffer [50 mM Tris-HCl (pH 7.4), 50 mM NaF, 10 mM β-glycerophosphate disodium salt, 1 mM EDTA, 1 mM EGTA, and 1 mM activated Na3VO4 (all from Sigma-Aldrich)] and a complete protease inhibitor cocktail tablet (Roche, West Sussex, UK) at 10 μg/μg tissue. Homogenates were rotated for 10 min, and the supernatant was collected by centrifugation at 13,000 g for 5 min at 4°C. The myofibrillar pellet was solubilized in 0.3 M NaOH and isolated from the insoluble collagen by centrifugation, and the myofibrillar protein was precipitated with 1 M perchloric acid. The sarcoplasmic proteins were precipitated from the initial homogenate supernatant fraction with 1 M perchloric acid and separated by centrifugation. The insoluble collagen fraction was washed sequentially with 0.3 M NaOH, 70% ethanol, 0.5 M acetic acid, and 0.5 M acetic acid and isolated by centrifugation. The protein-bound alanine was determined by gas chromatography-pyrol-

**Isokinetic dynamometer.** Iso-exercise testing was performed on an isokinetic dynamometer (Isocom; Isokinetic Technologies, Euronet Health). The visible portion of the fascicle length was assessed from the ultrasound scans using Image J 1.42q (National Institutes of Health). The number of fascicles and the number of deuterium moieties incorporated per alanine, 3.7) as the surrogate precursor labeling between subsequent biopsies. In brief, the standard equation used was:

\[
\text{FSR} = \frac{A P E_{\text{Ala}} - A P E_{\text{Ala}}}{A P E_{\text{Ala}}} = \text{deuterium enrichment of protein-bound alanine, }
\]

where \( A P E_{\text{Ala}} = \text{deuterium enrichment of protein-bound alanine} \)

and \( A P E_{\text{Ala}} = \text{mean precursor enrichment over the time period, and } t \) is the time between biopsies.

**Muscle architecture and strength assessment.** We also analyzed changes in muscle architecture, as these represent a technique with sufficient resolution to detect subtle changes in muscle architecture (albeit not muscle mass) after only short periods of exercise training (1). Exercised leg vastus lateralis ultrasonographic images were obtained at rest using B-mode ultrasonography (MyLab 70; Esaote Biomedica) with a 100 mm, 10- to 15-MHz, linear array probe. Ultrasound images were taken at a specific point at full knee extension when the participant was lying supine. The transducer was aligned in the fascicle plane to capture an optimal portion of fascicles. The muscle architecture parameters fascicle length and pennation angle were quantified by the same unblinded investigator (M. V. Franchi) from the ultrasound scans using Image J 1.42q (National Institutes of Health). The visible portion of the fascicle length was assessed directly using this software. Pennation angle was measured as the intersection between fascicles and the deep tendon aponerose. Isometric muscle strength was measured in a sitting position using an isokinetic dynamometer (Isocom; Isokinetic Technologies, Eurokinetics) throughout a range of six knee joint angles from 90° to 40°, with full extension corresponding to 0°. Subjects were seated in the dynamometer chair and secured into position using straps across the chest. Contractions lasted for 4 s, with a rest period of 45 s between contractions and a rest period of 90 s between knee joint angles.

**Plasma (20 μl) proteins were precipitated with 100% ethanol, the supernatant evaporated to dryness and was reconstituted in 0.5 M HCl, and the lipid fraction was removed using ethyl acetate extraction; alanine was then converted to its MCME derivative, as described above. All samples were run alongside an t-alanine-2,3,3,3-d4 standard curve. Enrichment (mole %excess) of alanine was then determined using gas chromatography-mass spectrometry (GC-MS: MD800, Fison, UK) and single ion monitoring of m/z 102, 103, 104, 105, and 106.

**GC-pyrolysis IRMS deuteration analyses.** Prior to each analysis, the IRMS system was calibrated and tested for measurement accuracy and was not used unless it fell within the manufacturer’s technical specifications. For GC-pyrolysis-IRMS analysis of acetone and MCME alanine, samples were separated on a DB-wax column (30 m x 0.32 mm × 0.25 μm; Agilent J & W) following splitless injection. The oven temperature program for acetone was started at 50°C and held for 2 min and then ramp at 30°C/min to 240°C and hold for 2 min, and for alanine it was started at 70°C and held for 3 min and then ramp at 10°C/min to 240°C and hold for 15 min. The separated samples were then passed through a high temperature (1,420°C) conversion reactor, where the analytes were converted to H2 gas before being directed to the IRMS, where the δ2H/1H ratio was determined. For TC/EA analysis of body water enrichment, following direct liquid injection into the TC/EA, where samples were immediately converted to H2 gas, sample gases were directed to the IRMS, where the δ2H/1H ratio was determined. The deuterium isotopic enrichment provided as δ2H was converted to atom% using the following equation:

\[
\text{Atom}% = \frac{100 \times \text{AR} \times (\delta^2 H \times 0.001 + 1)}{1 + \text{AR}(\delta^2 H \times 0.001 + 1)}
\]

where AR represents the absolute ratio constant for deuterium based on the VSMOW standard and equates to 0.00015595. This was then converted to atom percent excess (APE) by correcting for baseline sample, i.e., background enrichment. If any sample fell outside the dynamic range of the instrument, these were re-injected or re-prepared.

**Calculation of the FSR.** The FSR of MyoPS, sarcoplasmic synthesis (SPS), collagen synthesis (CPS), and protein synthesis was determined from the incorporation of deuterium-labeled alanine into protein, using the enrichment of body water (corrected for the mean number of deuterium moieties incorporated per alanine, 3.7) as the surrogate precursor labeling between subsequent biopsies. In brief, the standard equation used was:

\[
\text{FSR} = \frac{A P E_{\text{Ala}} - A P E_{\text{Ala}}}{A P E_{\text{Ala}}} = \text{deuterium enrichment of protein-bound alanine, APE}_{\text{Ala}} = \text{mean precursor enrichment over the time period, and } t \text{ is the time between biopsies.}
\]
A single bolus of 150 ml of deuterium oxide (D2O; 70 atom%). Body water enrichment over the 8-day period can be observed in Fig. 2A.

**RESULTS**

**Body water and plasma alanine enrichment.** Mean body water enrichment over the 8-day period can be observed in Fig. 2A. A single bolus of 150 ml of 70% D2O led to a body water enrichment of 0.202 ± 0.009% 24 h postingestition (range: 0.162–0.237%). Body water enrichment followed an exponential decay pattern (Fig. 2A), decaying slowly and significantly over the 8-day period at a rate of ~0.009/day. Raw APE values were natural logarithm converted to determine the mean decay constant (see Fig. 2B), and from this an estimation of mean half-life for body water elimination can be made; this was calculated as 11 ± 0.9 days.

Saliva samples from two subjects were prepared via a modification of a method described by Mahsut et al. (34), where deuterium from water is transferred onto the hydrogen positions of acetone under high pH incubation. These samples were then analyzed via GC-pyrolysis-IRMS. This was performed to validate that there was no dilution in enrichment with saliva due to contaminants that might have been present.

**Table 1.** Comparison between MS techniques for measuring BW enrichment following D2O ingestion and the use of saliva as a surrogate for plasma

<table>
<thead>
<tr>
<th>Participant 1 BW (APE)</th>
<th>Participant 2 BW (APE)</th>
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<tr>
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<td>TC/EA</td>
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<td>0.124</td>
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MS, mass spectrometry; D2O, deuterium oxide; BW, body water; APE, atom %excess; TC/EA, high-temperature conversion elemental analyzer; GC-Pyr-IRMS, gas chromatography-pyrolysis-isotope ratio mass spectrometry.
factor is used in the precursor-product equation to calculate FSR when body water enrichment is being used as the surrogate precursor. To validate this for our own calculations, the alanine enrichment in 8-day plasma samples was determined by GC-MS, and the mean plasma alanine/saliva body water ratio was calculated as 3.67 (Fig. 2A), therefore confirming the utility of 3.7 as a correction factor for use in the calculation of FSR in this study.

FSRs of MyoPS, SPS, and CPS. Figure 3, A and B, shows the mean $\delta^2$H in protein-bound alanine over the course of the 8-day training period. An exponential nonlinear pattern of incorporation was observed. Breaking this down into sections of time, we identify linear kinetics as being maintained for 0–2 days (Fig. 3A), after which it switches to nonlinear from 4 to 8 days (Fig. 3B). Therefore, the standard precursor-product calculation (as detailed in MATERIALS AND METHODS) was only valid for FSR 0–4 days, after which a nonlinear model was applied to the data whereby

$$FSR = -\ln\left(1 - \frac{(APE_{Ala})}{(APE_p)}\right)$$

Using this equation, in the rested leg, MyoPS was 1.45 ± 0.10, 1.47 ± 0.06, and 1.35 ± 0.07%/day at 0–2, 0–4, and 0–8 days, respectively, which when calculated as percent/hour represented between 0.052 and 0.061%/h; this rate is identical to the rates that are observed using traditional AA stable isotope tracers (53). MyoPS was increased significantly in the exercised leg at 0–2 (1.97 ± 0.13%/day), 0–4 (1.96 ± 0.15%/day, P < 0.01; Fig. 3A), and 0–8 days (1.79 ± 0.12%/day, P < 0.05). Figure 3D illustrates the temporal pattern for stimulation of MyoPS over the 8-day training period. FSR in the exercised leg is significantly increased above that of the rested leg at 2–4 days (P < 0.001), after which it rapidly falls by ~40% from 4 to 8 days (P < 0.01). CPS was, as expected, slower overall than MyoPS but followed a similar pattern over the 8 days, with the exercised leg showing a trend for greater FSR (2 days: 1.14 ± 0.13 vs. 1.45 ± 0.15%/day; 4 days: 1.13 ± 0.07 vs. 1.47 ± 0.18%/day; 8 days: 1.03 ± 0.09 vs. 1.40 ± 0.11%/day; Fig. 4A). However, due to a problem with sample recovery during collagen sample extraction, there were a number of missing data points; therefore, it was not possible to run statistical analyses on this data set. SPS, as with CPS, was found to be increased in the exercised leg at 0–2, 0–4, and 0–8 days (1.93 ± 0.15 vs. 1.48 ± 0.17, 1.67 ± 0.11 vs. 1.22 ± 0.07, and 1.88 ± 0.23 vs. 1.39 ± 0.23%/day, respectively; Fig. 4B); however, this was not statistically significant.

DISCUSSION

The purpose of the present study was to determine the efficacy of using the D2O tracer to determine the temporal

![Figure 3](https://example.com/figure3.jpg)

**Fig. 3. A:** linear incorporation of deuterium into protein bound alanine over the 1st 4 days. **B:** illustration of nonlinear incorporation of deuterium into protein-bound alanine beyond 4 days. **C:** myofibrillar protein synthesis rates in exercised (solid line) and nonexercised (dashed line) legs over 8-day training period. **D:** temporal pattern of myofibrillar protein synthesis in exercised and nonexercised legs. **Significantly different from rested leg at the same time point, P < 0.001; **Significantly different from previous time point in same leg P < 0.05. FSR, fractional synthetic rate.
MUSCLE PROTEIN SYNTHESIS USING D₂O

Innovative Methodology

Fig. 4. Muscle collagen (A) and sarcoplasmic protein synthesis (B) in exercised (solid line) and nonexercised (dashed line) legs over 8-day training period.

In the present data, we further validate the use of D₂O in measuring turnover in skeletal muscle (16, 18, 33, 50) by highlighting for the first time the efficacy of the D₂O tracer for monitoring the cumulative and temporal responses of protein synthesis to an anabolic stimulus (RET). Over 8 days of RET, MyoPS peaked between 2 and 4 days. Following this initial stimulation there was a drop in FSR of ~40% from 4 to 8 days, which could be suggestive of a “muscle full” effect (3, 4, 8, 38) where the muscle has achieved a new set point of accommodation, requiring greater stimulation to sustain growth. Finally, although not significant, there was a slight drop in MyoPS in the nonexercised leg over the 8 days (Fig. 3D), and this may have been influenced by the free-living nature of the design; for example, no strict controls on diet and exercise were imposed on the participants. We conclude that future studies of this kind would thus benefit from diet and activity monitoring.

The importance of studying distinct muscle protein subfractions can be seen via the preferential acute increase in myofibrillar MPS in response to resistance vs. mitochondrial MPS after endurance exercise, both of which are prophetic of ensuing adaptations to each exercise mode (58). As with MyoPS, both CPS and SPS have been shown to be stimulated in a similar fashion immediately post-resistance exercise, highlighting the coordinated response of muscle proteins to stimulation by exercise (36). In the present study, we have seen similar coordinated effects. SPS was greater, albeit not quite significantly (P = 0.14, 0.07, and 0.16 for 0–2, 0–4, and 0–8 days respectively), over the 8-day training period in the exercised compared with the nonexercised leg. This lack of significant findings may be influenced by the diverse nature of the sarcoplasmic protein pool, and dramatic variance in turnover rates within this pool will also reflect the variance observed within the present data. Our present data on CPS support other published data (5, 36) demonstrating that muscle collagen turns over at a slower rate than the myofibrillar and sarcoplasmic fractions (~25% lower than myofibrillar in the nonexercised state). As expected, CPS rates showed a pattern of change over the 8-day period that was similar to the myofibrillar fraction; however, because of sample processing problems, we were unable to produce complete data sets for all subjects, such that statistical representation of the data is limited. Despite this, we have shown that this approach can be used for the quantification of multiple muscle fractions in a single study, and further refinements to the protocol, sample processing, and the use of synthesis of muscle protein subfractions at “rest” and in response to short-term RET in the form of successive exercise bouts. Additionally, we hoped to demonstrate the utility of this method for providing short-term resolution of the regulation of muscle mass under conditions where traditional AA tracer approaches are untenable and where changes in mass due to RET are not yet quantifiable with standard image measurement techniques (ultrasound, MRI, DEXA). Indeed, although we observed increases in fascicle length and pennation angle at the end of 8 days, which is suggestive of increased hypertrophic response, this change was very small, increasing by 1.08 and 1.8%, respectively. Furthermore, we aimed to validate the use of IRMS for quantifying these responses rapidly and robustly. Here, we show for the first time that, using a single bolus of D₂O, we are able to detect synthesis rates of multiple protein fractions (MyoPS, CPS, and SPS) at rest and also increases in synthesis in response to an anabolic stimulus (exercise) vs. an internal nonexercised contralateral leg. This difference was detectable within 48 h of the first exercise bout, gradually declining from 2 to 8 days. This highlights the applicability of D₂O for quantifying cumulative MPS over periods when detection of an increase in muscle mass (via imaging techniques) is not feasible.

RET remains the most feasible tool for increasing or maintaining muscle mass in both healthy young and old individuals (20, 44) in addition to frail elderly and clinical populations (10, 32, 55). Hypertrophy associated with resistance exercise is caused primarily by cumulative increases in MPS, leading to net protein accretion (4). Using traditional AA tracer techniques, resistance exercise has been shown to induce increases in MPS following a single bout (22, 30, 42), which can be augmented by the addition of a protein feed postexercise (57) and with adequate nutrition can be sustained for up to and beyond 24 h (12, 36, 42). It was our aim to measure participants in a “free-living” normal environment such that these studies were not controlled for feeding, with the exception of the protein taken postexercise, to provide a “maximal stimulus” to ensure that the study was not confounded by individuals consuming insufficient protein with which to sustain MPS. Indeed, following the first bout of RET we observed that MyoPS was increased by ~36% compared with the nonexercised leg after 48 h (Fig. 3C). These data support the findings of the acute AA tracer work (acute infusions on different study days), which substantiates that with adequate nutrition MPS can be maintained for greater than 24 h postexercise (12, 36).

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MyoPS peaked between 2 and 4 days.
tandem MS will permit measurement of the turnover of additional protein pools, e.g., mitochondrial protein fractions.

It is also necessary to compare our FSRs to those captured using traditional $^{13}$C methodologies to qualify their quantitative utility. We obtained "rested" FSR for MyoPS of $1.25-1.47\%$/day, which equates to $0.05-0.06\%/h$, representative of that usually achieved with the use of $^{13}$C-AA tracers (53). This was also observed with rested SPS rates that were $0.051-0.062\%/h$ in the present study, similar to rates quoted within the literature using $^{13}$C tracers (3, 36). Although CPS rates were slightly higher than some values quoted in previous literature [$0.042-0.047\%/h$ here compared with $0.018-0.025\%/h$ (35, 36)], CPS rates of $0.06\%/h$ have been reported (22). Therefore, using the present approaches, we arrive at values approximating those expected.

Herein, we have validated a highly sensitive stable isotope tracer approach for measuring human MPS using $D_2O$. In one of the only other human studies of its kind, Gasier et al. (16) showed that, by providing a 300-ml bolus of 70% $D_2O$, they were able to measure increased rates of MyoPS following a single bout of resistance exercise 24 h postexercise using the same unilateral model as the present study. To do this, these authors utilized GC-MS for measuring both bound protein and body water deuterium enrichment, reporting that because of low levels of enrichment in the body water ($<0.5\%$), accurate data quantification was difficult due to limitations of the GC-MS approach they used. Here, we show that by giving a single 150-ml bolus of $D_2O$ and using a combination of TC/EA-IRMS and GC-pyrolysis-IRMS, we are able to sensitively and accurately measure body water and bound protein alanine enrichment to a precision of 1–3 $\delta^2H$. The substantial increase in measurement sensitivity provided by IRMS allows us to uncover subtle temporal differences in response to exercise not possible with standard GC-MS. Indeed, $\delta^2H$ changes over 2 days in the present study ranged from 116 to 176 $\delta^2H$ for such large changes suggest that measurements of MyoPS may be possible over even shorter periods (e.g., several hours, given the appropriate $D_2O$ dosing protocol), with similar levels of accuracy using IRMS. Per hour, current $\delta^2H$ changes would be equivalent to 2–3 $\delta^2H$; however, by increasing the $D_2O$ dosing threefold, with appropriate timing of doses to avoid side effects, we believe it is likely that rates of MyoPS could be measured over periods of hours, enhancing the utility of this approach and potentially replacing the need for intravenous AA tracers for measurement of protein turnover in certain situations.

To ease the burden on the participants within the present study, we chose to provide only a single small bolus of $D_2O$ (150 ml) at the beginning of the study based on the experimental design provided by MacDonald et al. (33), where a single 100-ml $D_2O$ (70 atom%) bolus was applied to measure MyoPS over a period of 4–14 days. Many published uses of $D_2O$ for monitoring metabolic turnover in both animals and humans usually provide regular daily doses of $D_2O$ following an initial large bolus, and this is implemented in an attempt to mimic that of AA tracers, whereby a large primed and continuous infusion is provided to maintain an isotopic steady state. However, it is possible to measure protein synthesis in the non-steady-state conditions (15, 46). Because of the slow turnover of water in the body, half-life is reported as $~7–10$ days (17, 33) and was estimated in the present study as a mean of $11 \pm 0.9$ days. MacDonald et al. (33) has shown that $D_2O$ elimination follows a slow exponential decay after bolus ingestion, as was observed here (Fig. 2A). This slow decay maintains body water enrichment within adequate levels for continuous incorporation into muscle protein and measurement of MPS over periods of 1 wk and potentially longer. The use of IRMS, with its high measurement sensitivity for isotopic abundance, ensures that body water enrichment needs to be raised by only a small amount ($~0.2\%$) compared with the 1–2% quoted by other researchers (50). Furthermore, this also minimizes the potential for the onset of the reported side effects from $D_2O$, such as dizziness and nausea. This function of slow $D_2O$ elimination combined with high measurement precision makes this technique ideally suited for use within populations where regular $D_2O$ administration or high doses of $D_2O$ may be problematic (e.g., frail elderly) or contraindicated (e.g., children or critical care patient populations), highlighting the wide-ranging applicability of this tracer technique.

To summarize, using only a single 150-ml bolus of $D_2O$, we show for the first time that it is possible to measure the temporal response of FSR within multiple muscle protein subfractions over a short period and also validate the approach in response to an anabolic stimuli (RET), from which we could determine the temporal and cumulative anabolic responses to exercise (compared with the contralateral nonexercise control leg). The increased sensitivity provided by GC-pyrolysis-IRMS allows accurate measurement of MyoPS, SPS, and CPS over as little as 2 days while also being able to robustly detect a stimulation of protein synthesis. Therefore, we reaffirm that $D_2O$ is a valid tracer approach for measuring MPS and muscle anabolism in extended free-living situations and will have wide application to assess the efficacy of clinical/nutritional/exercise interventions for maximizing mass/attenuating atrophy. Finally, despite holding much promise, the extent to which $D_2O$ can be utilized as a tool to “predict” chronic anabolic outcomes remains to be determined.

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AUTHOR CONTRIBUTIONS

P.J.A. and K.S. approved the final version of the manuscript; M.V.F., M.S.B., J.P.W., and W.K.M. performed the experiments.

REFERENCES


Innovative Methodology

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