Anabolic resistance assessed by oral stable isotope ingestion following bed rest in young and older adult volunteers: Relationships with changes in muscle mass

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Summary

Background & aims: Aging and experimental bed rest are associated with muscle atrophy and resistance to post-prandial stimulation of protein synthesis or anabolic resistance (AR). We have used in young and older adult volunteers, during short-term bed rest, a quick and non-invasive method, based on a single oral bolus of the stable isotope L[ring-2H5]phenylalanine (D5Phe), to determine post-prandial AR, defined as ratio between irreversible hydroxylation and incorporation into body protein of ingested phenylalanine.

Methods: We compared in older (O, 59 ± 1 y) and young (Y, 23 ± 1 y) healthy male volunteers the effects of two-week bed rest on post-prandial protein kinetics, assessed during absorption of a standard ready-to-use oral nutritional supplement, through stable-labeled isotope amino acid D5Phe, diluted in water, given as single oral load. The metabolic fate of D5Phe is either utilization for protein synthesis or irreversible hydroxylation to L[ring-2H4]tyrosine (D4Tyr). AR was defined as ratio between irreversible hydroxylation and incorporation into body protein of ingested phenylalanine.

Results: At baseline, in pooled Y and O subjects, values of AR were inversely correlated with QMV (R = −0.75; p < 0.03). Following 2-weeks of inactivity, there were significant bed rest effects on AR (p < 0.01) and QMV (p < 0.03), as well as significant bed rest × group interaction for AR (p < 0.03; +9.2% in Y; +21.9% in O) and QMV (p < 0.05; −5.7% in Y; −7.3% in O). In pooled subjects, the percentage delta changes in AR and QMV, induced by bed rest, were inversely correlated (R = −0.57; p < 0.05).

Conclusion: Bed rest-induced AR is much greater in the older than in younger adults. We have developed a new, simple, non-invasive method for the assessment of AR. The results indicate that this metabolic abnormality is a key mechanism for sarcopenia of aging and inactivity.

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1. Introduction

Several physiological and pathological conditions are characterized by a reduced ability of meal proteins to stimulate post-prandial protein synthesis. This metabolic alteration, defined as
anabolic resistance, is evaluated through complex and invasive methodology based on intravenous infusion of stable isotopes of amino acids [1-4]. We, and others, have found that anabolic resistance (AR) is the key mechanism leading to muscle atrophy following prolonged physical inactivity [2,3,5]. These observations were obtained using the experimental bed rest model in young volunteers [5].

Evidence in healthy, active older subjects indicates that physiological aging is associated with AR [4]. Furthermore, aging is also characterized by sarcopenia [6]. These results support the latest recommendation for a higher protein requirement in older subjects in order to counteract AR and delay sarcopenia [7]. Nonetheless, the effects of experimental bed rest on the onset of AR in an aged population compared to young controls have never been investigated.

We recently completed a European funded project (PANGEA 2014, ClinicalTrials.gov Identifier: NCT02694471) to compare the response of skeletal muscle mass and function to a two-week period of bed rest in older adults and young men [8]. The results showed that the impact of inactivity on muscle was greater in the older than in the younger subjects. In the present study we propose a rapid and non-invasive method, tested within the frame of the PANGEA project, eight healthy older adults (age, 59 ± 1 y; body mass index, BMI, 27 ± 1 kg/m²) and seven young healthy adults (age, 23 ± 1 y; BMI 24 ± 1 kg/m²) participated to 14-day experimental bed rest. The study was performed at the Orthopaedic Valdoltra Hospital, University of Primorska (Ankaran-Capodistria, Slovenia). The protocol was approved by the Ethic Committee of the University of Ljubljana (Slovenia), and conformed to the standards set by the Declaration of Helsinki (2002) and its amendments. A written informed consent was obtained from each subject upon enrollment. Subjects were admitted at the hospital 3 days before bed rest in order to undergo basal measurements and get acquainted with the hospital environment (ambulatory period) and remained in the hospital for another 3 days (recovery phase) after the bed rest period. The whole study design is reported elsewhere [8]. Daily dietary intake was planned and monitored by an expert dietitian to maintain each subject in near-neutral eucaloric balance [10] and to provide a standard macronutrient intake, of around 60% carbohydrate, 25% fat, and 15% protein (about 1.0–1.1 g/kg/day). In spite of the careful monitoring of food intake by the dietitian, one of the older subjects was excluded from the data analysis because of a severe negative energy balance (body mass loss of 4.5 kg) developed during the bed rest period, since we have previously shown that a negative energy balance accelerates muscle loss during bed rest [3].

2. Subjects and methods

In the framework of the PANGEA project, eight healthy older adults (age, 59 ± 1 y; body mass index, BMI, 27 ± 1 kg/m²) and seven young healthy adults (age, 23 ± 1 y; BMI 24 ± 1 kg/m²) participated to 14-day experimental bed rest. The study was performed at the Orthopaedic Valdoltra Hospital, University of Primorska (Ankaran-Capodistria, Slovenia). The protocol was approved by the Ethic Committee of the University of Ljubljana (Slovenia), and conformed to the standards set by the Declaration of Helsinki (2002) and its amendments. A written informed consent was obtained from each subject upon enrollment. Subjects were admitted at the hospital 3 days before bed rest in order to undergo basal measurements and get acquainted with the hospital environment (ambulatory period) and remained in the hospital for another 3 days (recovery phase) after the bed rest period. The whole study design is reported elsewhere [8]. Daily dietary intake was planned and monitored by an expert dietitian to maintain each subject in near-neutral eucaloric balance [10] and to provide a standard macronutrient intake, of around 60% carbohydrate, 25% fat, and 15% protein (about 1.0–1.1 g/kg/day). In spite of the careful monitoring of food intake by the dietitian, one of the older subjects was excluded from the data analysis because of a severe negative energy balance (body mass loss of 4.5 kg) developed during the bed rest period, since we have previously shown that a negative energy balance accelerates muscle loss during bed rest [3].

2.1. Metabolic test

Before the start (baseline) and at the last day of the bed rest protocol, we performed in each subject a metabolic test aimed at assessing the ratio between irreversible hydroxylation and incorporation into body protein of ingested phenylalanine, applying a new method based on bolus meal and oral stable isotope administration. The day of the study, at 7 AM, after an overnight fast, a polyethylene catheter was inserted into a forearm vein for blood collection. After a blood sampling, at time 0, for baseline data, a liquid oral meal and an oral bolus of D5Phe (0.3 g), dissolved in 150–200 mL of water, were administered to each subject to be consumed in 10 min. The meal was a ready-to-use standard product (Nutricomp®, B.Braun, 500 mL, 500 Kcal, vanilla flavor) composed by 15% protein (i.e., 18.75 g of proteins mainly derived from casein and soy), 30% fat and 55% carbohydrate. We have assessed in post-prandial conditions, changes in plasma D5Phe concentration, which is the net result of D5Phe absorption, utilization for protein synthesis and hydroxylation into D4Tyr (Fig. 1). The ratio between the AUC of both amino acids (i.e., AUC D4Tyr-to-AUC D5Phe ratio) expresses the amount of D5Phe, hydroxilated into D4Tyr and not incorporated into body protein (Fig. 2A and B). Blood was collected in EDTA tubes, immediately centrifuged at 3000 g at 4°C for 10 min and plasma was immediately stored at −80°C. In the same days, before the metabolic test, body fat mass has been assessed by bioimpedance analysis (BIA101, Akern, Florence, Italy), following manufacturer instructions, while muscle mass changes were assessed by continuous magnetic resonance imaging (MRI) (Magnemot Avanto; Siemens Medical Solution, Erlangen, Germany).

2.1.1. Anabolic resistance: analysis and calculations

Isotopic enrichments of plasma D5Phe and D4Tyr, derived by phenylalanine hydroxylation, were determined by gas chromatography-mass spectrometry (GC–MS) (HP 5890; Agilent Technologies, Santa Clara, CA) as t-butyldimethylsilyl derivatives [11]. Plasma concentrations of phenylalanine and tyrosine were assessed in all samples by GC–MS, using the internal standard technique, as previously described [11]. Known amounts of 13C-phenylalanine and 2H2-tyrosine (Cambridge Isotope Laboratories, Andover, MA) were added as internal standards. Isotopic enrichments were assessed considering the following mass-to-charge ratios (m/z): phenylalanine m/z 234–239; tyrosine m/z 146–470. Amino acid concentrations were assessed considering the following mass-to-charge ratios (m/z): phenylalanine m/z 336–337 and tyrosine m/z 466–468.
Plasma concentrations of unlabeled amino acid \([\text{aa}]\) were determined by the internal standard technique as follows:

\[
[\text{aa}] = \frac{a}{C_2} \times \text{TTR}_{\text{is}}
\]

where \(a\) is the concentration of the internal standard added to plasma samples and \(\text{TTR}_{\text{is}}\) is the isotopic enrichment of the internal standard.

D5Phe and D4Tyr concentrations \([\text{aatracer}]\) were calculated as follows:

\[
[\text{aatracer}] = \frac{[\text{aa}]}{C_2} \times \text{TTR}_{\text{tracer}}
\]

where \([\text{aa}]\) is plasma concentration of the tracee (unlabeled phenylalanine or tyrosine) and \(\text{TTR}_{\text{tracer}}\) is the isotopic enrichment of D5Phe or D4Tyr.

The areas under plasma labeled amino acid concentrations versus time curves, were estimated using the linear trapezoidal method from time 0–6 h. AR was defined as ratio between the areas under the curves of D4Tyr-to-D5Phe plasma concentrations over 6 h meal absorption, thereby expressing the fraction of D5Phe, hydroxilated into D4Tyr, not incorporated into body protein.

### 2.1.2. Magnetic resonance images

Quadriceps muscle volume of the dominant leg was measured by turbo spin-echo, T1-weighted, MRI obtained with 1.5 T (Magnetom Avanto; Siemens Medical Solution, Erlangen, Germany). On each MRI slice, contours corresponding to the quadriceps muscles were delineated by an MRI imaging expert, using the image processing tools OsiriX (Pixmeo Sarl, v.4.1.2). The muscle volume was then derived by summing a series of evenly spaced truncated cones, each encompassed between two axial images. The process included an average of 25 images (range 23–28) and covered the entire length of the quadriceps [8].

### 2.2. Statistics

Data were expressed as mean ± SEM. Repeated measures ANOVA (or ANCOVA when appropriate) was applied to define the effects of bed rest and bed rest × group interaction on investigated parameters. Spearman’s test was used to assess the correlations among changes in anabolic resistance and QMV in the two groups or changes between indices of anabolic resistance at different timing. Data were log-transformed where appropriate. \(p < 0.05\) was considered statistically significant. Statistical analysis was performed using SPSS statistical software (v12; SPSS, Inc., Chicago, IL). Statistical significant main effects were confirmed at alpha 0.05 and interaction effect at alpha 0.10.

### 3. Results

Table 1 shows the effects of bed rest on body mass, fat mass and quadriceps muscle volume in young and older adults. In basal conditions, muscle mass, expressed as quadriceps muscle volume (QMV) was significantly greater in the young as compared to the older subjects. In contrast, body mass, fat mass and fat-free mass were not significantly different in the two groups. After two-weeks of experimental inactivity we observed a
significant effect of bed rest on body mass and fat-free mass with no bed rest × group interaction. This effect determined a significant reduction of body mass in both groups. As expected, because of the maintained isocaloric condition, we did not observe any significant effect of bed rest or bed rest × group interaction on body fat mass (FM). QMV was significantly affected by 14 days of bed rest with a significant bed rest × group interaction (−7.3 ± 0.4% in older and −5.7 ± 1.5% in young). Individual values of FFM in young and older adults significantly (p < 0.05) correlated with QMV in the basal ambulatory conditions (R = 0.54), but not during bed rest (R = 0.42).

Figure 2A and B shows the values of fasting and post-prandial plasma concentrations of labeled (D5Phe and D4Tyr), before (Baseline) and at the end of 14 days of bed rest in the young and older subjects. In the older volunteers, mean time to the point of maximum concentration of labeled amino acids tended to be delayed in the baseline condition as compared to the bed rest period. These differences, however, were not statistically significant.

At baseline, fasting plasma concentrations of phenylalanine and tyrosine were not significantly different in young (65 ± 5 and 73 ± 8 μmol/L, respectively) and older subjects (54 ± 4 and 78 ± 7 μmol/L, respectively). We did not observe significant effects of bed rest on fasting plasma concentrations of phenylalanine and tyrosine in young and older subjects. There was not significant bed rest × group interaction for fasting plasma concentrations of phenylalanine and tyrosine. The AUC of plasma concentrations of labeled and unlabeled tyrosine and phenylalanine are reported in Table 2.

We did not observe either significant bed rest effect or a bed rest × group interaction on tyrosine AUC, however phenylalanine AUC was significantly affected by 14 days of bed rest in both groups with no significant bed rest × group interaction. We did not observe significant bed rest effect or bed rest × group interaction on AUC D5Phe, index of tracer absorption, whereas AUC D3Tyr, derived from the hydroxylation of D5Phe, was significantly affected by bed rest with a strong tendency on the bed rest × group interaction evidencing a greater increment of AUC D3Tyr in the older (+33 ± 16%) volunteers than in the young (+11 ± 14%) subjects.

We have defined the ratio between the AUC of D3Tyr and the AUC of D5Phe as an index of anabolic resistance to ingested protein. The baseline ratio between the AUC of D3Tyr and the AUC of D5Phe was greater in the older (0.23 ± 0.01), as compared to the younger volunteers (0.19 ± 0.02), without achieving statistical significance (unpaired t-test, p = 0.07). After 14 days of experimental inactivity, we observed a significant bed rest effect on the anabolic resistance index in both groups (older: 0.28 ± 0.02; young: 0.20 ± 0.01) (Fig. 3), with a significant bed rest × group interaction. There was an increment of 22 ± 6% and 9 ± 7% in the older and young volunteers, respectively.

At baseline, individual values of anabolic resistance (i.e., AUC D3Tyr-to-AUC D5Phe ratio) and quadriceps muscle volume were inversely correlated in young and older subjects (R = −0.56, p < 0.03; Fig. 4). Following 14 days of bed rest, percentage changes from baseline of individual values of AR (i.e., AUC D3Tyr-to-AUC D5Phe ratio) and quadriceps muscle volume were inversely correlated in young and older subjects (R = −0.75, p < 0.03; Fig. 5).

In postprandial conditions, changes in plasma L[ring-2H5] phenylalanine (D5Phe) concentration depend on D5Phe absorption, degradation and utilization for protein synthesis, as well as on the appearance of L[ring-2H4]tyrosine, produced by D5Phe hydroxylation. We administered a single oral bolus of D5Phe stable isotope to determine the anabolic resistance to dietary protein stimuli for whole body protein synthesis.

Plasma concentrations of D5Phe (continuous line) and D3Tyr (dotted line), after meal test, before (black) and at the end (grey) of 14-d bed rest, in healthy young (A) and older (B) adults. Values are expressed as mean ± SEM. Time is given as minutes from the ingestion of the ready-to-use liquid oral meal plus the D5Phe oral bolus. D5Phe, L[ring-2H5]phenylalanine; D3Tyr, L[ring-2H4]tyrosine.

Table 2

<table>
<thead>
<tr>
<th>Group</th>
<th>Basal</th>
<th>Bed rest</th>
<th>Bed rest × group interaction</th>
<th>p-value&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>AUC Phe (μmol × h/L)</td>
<td>Young adults</td>
<td>413 ± 28</td>
<td>430 ± 19</td>
<td>0.03</td>
</tr>
<tr>
<td></td>
<td>Older adults</td>
<td>401 ± 12</td>
<td>416 ± 16</td>
<td>0.68</td>
</tr>
<tr>
<td>AUC Tyr (μmol × h/L)</td>
<td>Young adults</td>
<td>516 ± 59</td>
<td>502 ± 45</td>
<td>0.34</td>
</tr>
<tr>
<td></td>
<td>Older adults</td>
<td>600 ± 44</td>
<td>615 ± 52</td>
<td>0.23</td>
</tr>
<tr>
<td>AUC D5Phe (μmol × h/L)</td>
<td>Young adults</td>
<td>16.7 ± 1.6</td>
<td>16.4 ± 1.4</td>
<td>0.21</td>
</tr>
<tr>
<td></td>
<td>Older adults</td>
<td>16.1 ± 1.6</td>
<td>18.0 ± 2.3</td>
<td>0.06</td>
</tr>
<tr>
<td>AUC D3Tyr (μmol × h/L)</td>
<td>Young adults</td>
<td>3.3 ± 0.6</td>
<td>3.3 ± 0.4</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td></td>
<td>Older adults</td>
<td>3.8 ± 0.4</td>
<td>4.9 ± 0.7</td>
<td>0.06</td>
</tr>
</tbody>
</table>

<sup>a</sup> ANOVA or ANCOVA (basal values as covariates) were used when appropriate. Values were log-transformed when appropriate. AUC, areas under the curve; Phe, phenylalanine; Tyr, Tyrosine; D5Phe, L[ring-2H5]phenylalanine; D3Tyr, L[ring-2H4]tyrosine.
assessed after 6-h meal test. Values are expressed as mean ± SEM.

ANCOVA analysis was performed using washout values as covariates. AUC, area under the curve; D5Phe, L[ring-2H5]phenylalanine; D4Tyr, L[ring-2H4]tyrosine.

4. Discussion

Aging is associated with reduced whole-body and muscle protein synthesis after protein intake. This metabolic response, termed anabolic resistance contributes to sarcopenia [4,15]. In this study we assessed in young and older adults the effects of two weeks of experimental bed rest on the ability of a standardized meal to promote incorporation of dietary amino acids into body proteins. Changes in AR, defined as ratio between irreversible hydroxylation and incorporation into body proteins of ingested phenylalanine and quantified by a simple and non-invasive method based on stable isotope ingestion, were compared with effects of bed rest on muscle mass measured by MRI [8]. We found that bed rest increased AR four-times more in the older than in the younger volunteers. Changes in AR inversely correlated with changes in muscle volume.

Table 3 reports selected studies describing the effects of inactivity and exercise on estimation of anabolic resistance in elderly and young subjects by different methods using stable isotopes [1,2,4,13,16,17]. Results and methods are compared with the present investigation.

The previous methodologies involved administration of stable isotopes of amino acids to follow their incorporation into body proteins. In order to apply traditional tracer dilution or incorporation equations, a metabolic steady-state is required, i.e., substrate concentrations in body fluids should not change over time. Thus, anabolic resistance has been often evaluated in non-physiological conditions, as during constant nutrient administration, either parenterally or enterally [1-4]. In addition, the assessment of AR in skeletal muscle requires multiple biopsies or vascular catheters, in femoral artery and vein, to measure fractional protein synthesis and monitor tracer disappearance into leg muscle protein, respectively [4,12,16]. At the whole-body level, two methods are currently available to determine the rates of amino acid utilization for protein synthesis in steady state condition. With the 13C-leucine method, irreversible amino acid oxidation is determined through the assessment of the 13CO2 rate of appearance in the breath by indirect calorimetry combined with isotope-ratio mass-spectrometry [3]. The second method is based on continuous infusion of two independent tracers of phenylalanine and tyrosine. Irreversible phenylalanine hydroxylation is determined by monitoring transformation of labeled phenylalanine into the correspondent tyrosine tracer (e.g., D5-phenylalanine into D4-tyrosine) [9]. With both methods, the rate of amino acid incorporation into body protein is derived from the difference between rate of amino acid appearance and irreversible loss (e.g., oxidation for leucine or hydroxylation for phenylalanine). These methods require tracer infusions for several hours to achieve steady state conditions in both the postabsorptive and postprandial states.

Thus, several limitations are associated with current methodologies to assess AR in vivo, in humans. First, intravenous tracer infusion requires preparation of sterile and apyrogenic solutions. Second, application of steady state equations to determine protein synthesis requires intravenous tracer infusions for several hours and constant (non-physiological) nutrient administration to simulate experimental post-prandial state [3,9,11]. Third, muscle biopsies or femoral arterial and venous catheters are required to assess AR in muscle proteins [4,12].
In the present study, we have described a new methodology to assess AR in humans. An oral bolus of D5Phe was administered in parallel with a standard ready-to-use oral nutritional supplement containing 500 kcal, 15% protein, 30% fat and 55% carbohydrate. Plasma concentrations of D5Phe and D4Tyr (derived from D5Phe through irreversible hydroxylation) were monitored for 6 h after meal ingestion. Tracer concentrations, calculated as tracer-to-tracee-ratio, multiplied by unlabeled amino acid concentration, were virtually zero after 6 h from meal ingestion. AUC of D5Phe and meal ingestion. Tracer concentrations, calculated as tracer-to-

parallel with a standard ready-to-use oral nutritional supplement

D4Tyr concentrations were then calculated from time 0
delay in enteral phenylalanine absorption, as long as the entire
versible hydroxylation to tyrosine. The results are not affected by
of phenylalanine is incorporation into protein synthesis or irre-
utilized for body protein synthesis, i.e. AR. In fact, the only fate

Application of the new methodology

The new methodology has some limitations, which have to be
pointed out. First, only the metabolic fate of ingested phenylalanine
(i.e., irreversible hydroxylation or incorporation into body proteins)
has been traced by the oral isotope. Postprandial changes of synthesis
and degradation of endogenous body proteins could not be assessed
in our study. Constant intravenous isotope infusion and steady state
conditions were required to trace endogenous body protein kinetics.

To overcome this methodological limitation, postprandial steady state conditions have been artificially created by constant oral (sip feeding) or intravenous
nutrient administration. By these methods anabolic resistance has
been demonstrated in bed resting young subjects and in ambulatory elderly. This experimental approach should be used in future studies
to assess the response of endogenous protein kinetics to constant nutrient administration in young and elderly subjects in ambulatory conditions and during bed rest. **Second**, the ratio between irreversible hydroxylation and incorporation into body proteins can be affected by protein intake level. We could not perform multiple studies in our subjects to create a dose–response curve. Protein content of the isotope-enriched experimental meals (i.e., 18.75 g) was equivalent to
that provided in daily with each main meal (three main meals per day administered throughout the study). Total daily protein intake was
within a narrow range in the young and older subjects (i.e., about
10.1–11 g/kg/day) because body weight and fat-free mass were not significantly different in the two groups (Table 1). **Third**, the ability to utilize ingested amino acids for body protein synthesis can be affected by individual muscle mass. Sarcopenic conditions could be characterized by anabolic resistance simply because of reduced
mass. This mechanism may in part contribute to our results because bed rest induced muscle loss was greater in the older than in
the young volunteers. We have normalized the amount of ingested protein utilized for body protein synthesis (i.e., 18.75 g multiplied by
(1 - AUC D4Tyr/AUC D5Phe)) by kg of FFM in the young and older volunteers before and after bed rest. There was no significant effect of
bed rest on this figure, however we found a significant bed rest > group interaction (p = 0.025, ANCOVA). Therefore, the amount of ingested protein utilized for body protein synthesis normalized by FFM was 14% greater (p = 0.03, unpaired t-test with Bonferroni correction) in the young as compared to the older volunteers during bed rest. This value was not significantly different in the ambulatory condition. These results suggest that bed rest induced anabolic resistance directly contributed to the greater disuse atrophy observed in the quadriceps muscle of the elderly. In a previous 2-week bed rest study in young subject, we have shown that the development of anabolic resistance preceded detectable changes in lean body mass as assessed by DEXA, thereby confirming our data [3].

### Table 3

<table>
<thead>
<tr>
<th>Reference</th>
<th>Methods</th>
<th>Experimental fed state</th>
<th>Conditions</th>
<th>Results</th>
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<tbody>
<tr>
<td>Biolo et al., 2004</td>
<td>Whole-body protein synthesis, degradation and balance by constant infusion of L-[1-13C]leucine</td>
<td>Constant IV amino acid infusion (130 mg/h/kg LBM)</td>
<td>Two weeks of experimental bed rest in young subjects</td>
<td>Net leucine deposition into body protein was ~8% lower during bed rest than during the ambulatory phase Muscles myofofrillar fractional synthetic rate in non-immobilized leg was greater by ~54% and ~68% with low and high dose AA infusion, respectively. Muscle fractional synthesis rate and net leg muscle protein balance were ~44% and ~120% greater after exercise than at rest, respectively. Lag muscle protein balance was 61% lower in the elderly than in the young adults.</td>
</tr>
<tr>
<td>Glover et al., 2008</td>
<td>Muscle myofofrillar fractional synthetic rate by constant infusion of L-[ring-13C6]phenylalanine</td>
<td>Constant IV amino acid infusion at low and high rates (43 and 261 mg/h/kg);</td>
<td>Two weeks of unilateral leg immobilization in young subjects</td>
<td>Muscle myofofrillar fractional synthetic rate in non-immobilized leg was greater by ~54% and ~68% with low and high dose AA infusion, respectively. Muscle fractional synthesis rate and net leg muscle protein balance were ~44% and ~120% greater after exercise than at rest, respectively.</td>
</tr>
<tr>
<td>Biolo et al., 1997</td>
<td>Muscle fractional synthesis rate and net leg muscle protein balance by constant infusion of L-[ring-13C6]phenylalanine</td>
<td>Constant IV amino acid infusion (150 mg/h/kg LBM)</td>
<td>Young subject following exercise</td>
<td>Young subject following exercise</td>
</tr>
<tr>
<td>Katsanos et al., 2005</td>
<td>Leg muscle arteriovenous phenylalanine balance by constant infusion of L-[ring-2H3]phenylalanine</td>
<td>Bolus ingestion of 15 g essential amino acids</td>
<td>Elderly and young subjects in ambulatory conditions</td>
<td>Elderly and young subjects in ambulatory conditions</td>
</tr>
<tr>
<td>Durham et al., 2010</td>
<td>Fractional muscle protein synthesis by primed continuous infusion of L-[ring-13C6]phenylalanine</td>
<td>Constant IV amino acid infusion (135 mg/h/kg)</td>
<td>Elderly and young subjects following endurance exercise</td>
<td>Efficiency of stimulation of muscle protein synthesis was reduced by ~40% in the elderly as compared to the young adults. The ratio between irreversible hydroxylation and incorporation into body protein of ingested phenylalanine increased by ~22% and ~9% following bed rest in the elderly and young adults, respectively.</td>
</tr>
<tr>
<td>Present study</td>
<td>Ratio between irreversible hydroxylation and incorporation into body protein of phenylalanine ingested by bolus oral administration of L-[ring-2H3]phenylalanine</td>
<td>Bolus ingestion of mixed meal (18.8 g of protein)</td>
<td>Two weeks of experimental bed rest in young and older subjects</td>
<td>Two weeks of experimental bed rest in young and older subjects</td>
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</tbody>
</table>

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In full agreement with previous studies [2,3,13], we found an effect of bed rest on post-prandial AR in the two groups of young and older adults. However, the impact of unloading on AR was significantly greater in the older, as compared to the younger volunteers. These results suggest that older adults are more susceptible to the negative effects of unloading on post-prandial protein anabolism, as compared to younger individuals.

Human studies clearly indicate that decreased protein synthesis in the postprandial state is the main protein catabolic mechanism associated with muscle unloading and aging [2,14]. Sarcopenia is defined as a progressive decline in skeletal muscle mass and function, strongly related to the decrease in autonomy and quality of life in the older population [6]. Inactivity is a key mechanism of sarcopenia of aging. In our study, the effects of bed rest in the young and in the older adults on skeletal muscle mass were assessed as quadriceps volume by MRI. As expected [18], bed rest significantly decreased QMV in both young and older adults, however percent decreases in muscle mass were greater in the old than in the young group. In pooled young and older subjects, changes induced by bed rest in AR and in QMV were inversely correlated (Fig. 5). In addition, at baseline, AR was inversely correlated with quadriceps muscle volume (Fig. 4). These results clearly suggest that AR is a key mechanism for sarcopenia of aging and inactivity.

We have previously shown that both underfeeding and overfeeding can accelerate muscle loss during physical inactivity [3,10]. In the present study, energy intake was carefully tailored to energy requirement of individual subjects and to the level of physical activity. Energy balance was maintained throughout the experimental periods as shown by the fact that fat mass of young and older adults were not modified during bed rest (Table 1). Thus, changes in AR may be muscle mass were not affected by changes in energy balance.

In conclusions, we have developed a new, simple, non-invasive methodology to quantify AR in humans. Results suggest cause-effect relationships between inactivity-induced AR and muscle loss in both young and older subjects. AR is associated with increased protein requirement, as in the older and during inactivity [7]. Our new method allows to closely monitor changes in AR in different conditions.

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Authors’ contributions to manuscript

GB, designed research, analyzed data and performed statistical analysis, wrote paper and had primary responsibility for final content; RP, designed research, conducted research; SM, conducted research, analyzed data and performed statistical analysis; FGDC, conducted research, analyzed data and performed statistical analysis, wrote paper; SL, conducted research; BG, designed research, conducted research; CR, designed research, conducted research; AP, designed research, conducted research; JR, designed research, conducted research; MC, conducted research; BS, designed research, analyzed data; MN, designed research, conducted research, analyzed data. All authors have read and approved the final manuscript. Additional contribution: we thank all participants of the study.

Conflict of interest

The authors declare no competing financial interest.

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