The effect of short-term exercise prehabilitation on skeletal muscle protein synthesis and atrophy during bed rest in older men

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Abstract

Background  Poor recovery from periods of disuse accelerates age-related muscle loss, predisposing individuals to the development of secondary adverse health outcomes. Exercise prior to disuse (prehabilitation) may prevent muscle deterioration during subsequent unloading. The present study aimed to investigate the effect of short-term resistance exercise training (RET) prehabilitation on muscle morphology and regulatory mechanisms during 5 days of bed rest in older men.

Methods  Ten healthy older men aged 65–80 years underwent four bouts of high-volume unilateral leg RET over 7 days prior to 5 days of inpatient bed rest. Physical activity and step-count were monitored over the course of RET prehabilitation and bed rest, whilst dietary intake was recorded throughout. Prior to and following bed rest, quadriceps cross-sectional area (CSA), and hormone/lipid profiles were determined. Serial muscle biopsies and dual-stable isotope tracers were used to determine integrated myofibrillar protein synthesis (iMyoPS) over RET prehabilitation and bed rest phases, and acute postabsorptive and postprandial myofibrillar protein synthesis (aMyoPS) rates at the end of bed rest.

Results  During bed rest, daily step-count and light and moderate physical activity time decreased, whilst sedentary time increased when compared with habitual levels (P < 0.001 for all). Dietary protein and fibre intake during bed rest were lower than habitual values (P < 0.01 for both). iMyoPS rates were significantly greater in the exercised leg (EX) compared with the non-exercised control leg (CTL) over prehabilitation (1.76 ± 0.37%/day vs. 1.36 ± 0.18%/day, respectively; P = 0.007). iMyoPS rates decreased similarly in EX and CTL during bed rest (CTL, 1.07 ± 0.22%/day; EX, 1.30 ± 0.38%/day; P = 0.037 and 0.002, respectively). Postprandial aMyoPS rates increased above postabsorptive values in EX only (P = 0.018), with no difference in delta postprandial aMyoPS stimulation between legs. Quadriceps CSA at 40%, 60%, and 80% of muscle length decreased significantly in EX and CTL over bed rest (0.69%, 3.5%, and 2.8%, respectively; P < 0.01 for all), with no differences between legs. No differences in fibre-type CSA were observed between legs or with bed rest. Plasma insulin and serum lipids did not change with bed rest.

Conclusions  Short-term resistance exercise prehabilitation augmented iMyoPS rates in older men but did not offset the relative decline in iMyoPS and muscle mass during bed rest.

Keywords  Bed rest; Protein synthesis; Sarcopenia; Muscle

Received: 9 June 2020; Revised: 19 October 2020; Accepted: 24 November 2020

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Introduction

Sarcopenia is a condition characterized by skeletal muscle mass and strength loss with increased risk of frailty, falls, metabolic disease, and all-cause mortality in older individuals.\(^1\)\(^,-\)\(^2\) It is estimated that as many as 32 million older individuals across Europe could be diagnosed with sarcopenia by 2045.\(^3\) Currently, the annual cost to the National Health Service (NHS) of treating age-related muscle weakness is ~£2.5 billion (\(\sim 2\)–\(3\)\% of budget).\(^4\) with health care costs \(\sim 2\)–\(3\) times greater in those with muscle weakness.\(^5\) Thus, sarcopenia poses a major current and future predicted socio-economic threat.

Periods of disuse and inactivity, typical during illness and hospitalization, result in rapid muscle atrophy and impaired postprandial muscle protein synthesis (MPS) stimulation in older individuals.\(^6\)\(^–\)\(^9\) The impaired ability of older individuals to fully recover from these acute bouts of inactivity is thought to accumulate over time and contribute to the sarcopenic progression.\(^10\) This so-called catabolic crisis model is exemplified by the failure of >\(50\)% of hospitalized older individuals to regain pre-admission mobility levels 12 months after discharge.\(^11\) Alarmingly, low muscle mass/attenuation and poor physical function at discharge are associated with (i) greater risk of readmission, (ii) longer length of stay, (iii) greater reliance on external care after discharge, and (iv) greater mortality.\(^12\)\(^–\)\(^14\) Thus, mitigating disuse-induced muscle atrophy in older individuals could ultimately delay sarcopenia progression and improve quality of life, with important implications for health care expenditure.

Resistance exercise training (RET) stimulates MPS and can attenuate muscle atrophy when implemented during disuse.\(^15\)\(^,\)\(^16\) However, imposed disuse as a result of illness/co-morbidities or (planned) elective surgery may preclude older individuals from performing RET. Exercise interventions initiated prior to disuse (‘prehabilitation’) capitalize on a better physical and emotional condition of the patient as compared with peri-disuse or post-disuse RET and could potentially benefit a number of clinically relevant outcomes.\(^17\) Although multi-modal prehabilitation has been shown to offset the decline in function during disuse and enhance recovery during rehabilitation, findings are inconsistent,\(^18\)\(^–\)\(^22\) potentially because RET stimuli are insufficient (dose and/or time frame) to increase strength and muscle mass in older patients prior to disuse.\(^23\)\(^,\)\(^24\) Furthermore, the effects of RET prehabilitation on muscle morphology and regulatory mechanisms have yet to be explored. Whilst longer-term high-volume RET prehabilitation may be required to build a muscle mass and strength reserve in older patients, evidence suggests that several bouts of low-load high-volume RET, accumulated over a relatively short period, can augment postprandial MPS responsiveness.\(^16\)\(^,\)\(^25\) Given the possibility that short-term targeted RET prehabilitation could mitigate disuse-induced muscle anabolic resistance and, therefore, atrophy, it is imperative to investigate such concise, cost-saving interventions.

The primary aim of this study was to determine the effects of short-term unilateral leg RET prehabilitation on quadriceps muscle mass during 5 days of bed rest in older individuals. Using dual-stable isotope tracer and serial muscle biopsy sampling, we determined integrated myofibrillar protein synthesis (iMyoPS) rates over prehabilitation and bed rest phases, acute postabsorptive and postprandial myofibrillar protein synthesis (aMyoPS) rates, and gene/protein expression of targets known to modulate muscle mass. We hypothesized that RET prehabilitation would augment iMyoPS, gene expression, and anabolic signalling during prehabilitation and that this prior stimulus would offset the expected decline in iMyoPS and muscle mass during bed rest, through greater postprandial aMyoPS responsiveness.

Methods

Participants

Ten healthy older men (65–80 years) were recruited through local advertisements and deemed eligible for study participation if they had no history of structured RET within 10 years prior to study participation, were deemed healthy and free of sarcopenia diagnosis as assessed by a general health questionnaire, had a score of ≥9 on the Short Physical Performance Battery test, appendicular lean mass of ≥7.25 kg/m\(^2\), and a body mass index (BMI) < 30 kg/m\(^2\).\(^26\) Participants were excluded from study participation if they had a coagulation disorder, myocardial infarction, artery/vein disease, chronic/systemic illness, or (pre-)diabetes or underwent hormone replacement therapy. Furthermore, participants were excluded if they currently smoked and consumed any anticoagulant medication or medication that might affect muscle metabolism. Participants were asked to refrain from consuming any nutritional supplements that might affect muscle metabolism during the bed-rest phase. All participants were informed of the study purpose and procedures and provided their written informed consent. Ethical approval was obtained through the West Midlands—Black Country Research Ethics Committee (16/WM/0483) and was registered at clinicaltrials.gov (NCT04422665; RG_16-100). The study conformed to the
standards outlined by the Declaration of Helsinki (seventh edition).

**Experimental design**

After an initial screening visit and obtainment of study consent, participants visited the Wellcome Trust/National Institute for Health Research (NIHR) Clinical Research Facilities (CRF) at the Queen Elizabeth Hospital Birmingham for a preliminary testing visit (Day 1), prehabilitation phase (Days 2–7), mid-testing visit (Day 8), bed-rest phase (Days 8–13), and final testing day (Day 13). For each testing visit, participants reported to the CRF at 8 a.m. after an overnight fast (or were already present in the case of Day 13). An overview of the study timeline is depicted in Figure 1.

**Preliminary testing visit (Day 1)**

Following an overnight fast, participants provided a single saliva sample after which a baseline muscle biopsy from the vastus lateralis was obtained under 1% lidocaine using the Bergström needle technique, as described in our previous work. The leg to be biopsied first was randomly determined for the first participant and then alternated between subsequent participants. Participant height, body mass, compartmental body composition, estimated one-repetition maximum (1RM), and physical function were then determined (described subsequently). Participants were then provided with a bolus of deuterated water ($D_2O$) and daily top-up doses (described subsequently) for the measurement of iMyoPS rates. Prior to leaving the CRF, participants were given a 3 day weighed food diary to be completed during the subsequent prehabilitation phase, to determine habitual dietary intake. Participants were fitted with a hip-worn pedometer and wrist-worn accelerometer for the remainder of the study to monitor physical activity levels/intensity throughout prehabilitation and bed-rest phases.

**Prehabilitation phase (Days 2–7)**

Participants reported to the CRF to complete a bout of one-legged RET on Days 2, 4, 6, and 7. RET was performed by the strongest leg, as determined by the 1RM testing during the preliminary testing visit. Each RET bout consisted of two warm-up sets at 50% of 1RM, followed by six sets at 75% of 1RM for both the leg extension and leg curl. RET sets consisted of a target 12 repetitions and were separated by 2 min of passive rest. The exercise load was adjusted to maintain a subjective rating of perceived exertion of 8–9 on the modified Borg category-ratio scale (CR-10). RET sessions were conducted at a time convenient for the participant.

![Figure 1](image_url) Schematic overview of the longitudinal experimental design (Days 1–13; top) and the acute infusion trial conducted at the end of the bed-rest period (Day 13; bottom).
except for the last RET session, which was performed in the afternoon.

Mid-phase testing visit (Day 8)
On the morning of Day 8 and following an overnight fast, participants reported to the CRF. A skeletal muscle biopsy from the vastus lateralis was obtained from both the exercised (EX) and non-exercised control leg (CTL). Following this, repeat assessments of height, body mass, and body composition were taken. Finally, participants underwent a magnetic resonance imaging (MRI) scan to determine quadriceps muscle cross-sectional area (CSA) in both legs (described subsequently).

Bed-rest phase (Days 8–13)
To mimic the effects of a traditional inpatient hospital stay, participants underwent a 5-day period of strict bed rest. Once participants returned from the mid-testing MRI scan, the bed-rest period commenced, and participants remained in bed. During the day, participants were allowed to sit up in bed or in a recliner chair. Bathing and sanitary activities were performed in a wheelchair. Accelerometer and pedometer devices were briefly removed during showering. To prevent coagulation disorders and bed sores, participants were given light, non-weight-bearing exercises to be performed hourly each day (e.g. knee bends, rolling side to side, and ankle rotations). Participants also wore lower limb compression stockings and received a daily subcutaneous enoxaparin injection (20 mg). To mimic a typical inpatient stay, participants choose from a selection of meals/snacks provided by the CRF during bed rest. Dietary energy and macronutrient intake were not strictly controlled but were closely monitored and logged by CRF nursing staff.

Post-bed-rest experimental trial (Day 13)
Following an overnight fast, participants were woken up at 6 a.m. A 21 G cannula was inserted in an antecubital vein of both forearms. One cannula was used for serial blood sampling, whilst the other one was used to administer a stable amino acid isotope infusion. After a baseline blood sample was obtained, a primed-continuous infusion of L-[ring-13C6] phenylalanine was initiated (prime, 2 μmol/kg; infusion, 0.05 μmol/kg, Cambridge Isotope Laboratories, Andover, MA, USA). Blood samples were drawn from the contralateral arm at −185, −120, −60, and −5 min prior to consumption and 20, 30, 60, 90, 120, 180, and 240 min post-consumption of a milk protein drink (described subsequently). Blood samples were collected in serum separator and ethylenediamine-tetraacetic acid (EDTA)-treated vacutainers (BD Biosciences, Oxford, UK) and centrifuged at 3000 rpm at 4°C, with serum and plasma aliquots stored at −80°C for further analyses. After 150 min of infusion, a muscle biopsy was obtained from the vastus lateralis of both legs. Immediately after biopsy obtainment, participants consumed 18.75 g of milk protein isolate (MyProtein, Cheshire, UK), providing 15 g of milk protein, dissolved in 300 mL of water; 240 min after drink consumption, a second muscle biopsy from the vastus lateralis of each leg was obtained ~3 cm proximal to the first biopsy and indicated the end of the infusion trial. Participants were then fed a meal of their choice and transported in a wheelchair to the MRI scanner for post-bed-rest measurement of quadriceps muscle CSA. Obtention of the MRI scan indicated the end of the bed-rest phase. Participants walked back to the CRF for a final body composition assessment and consultation/assessment with a trained physiotherapist prior to discharge.

Experimental procedures

Body mass, height, and body composition
Participants’ body mass and height were determined in light clothing to the nearest 0.1 kg and 0.1 cm using electronic weighing scales and a stadiometer, respectively. Compartmenal body composition was determined using bioelectrical impedance analysis (TANITA BC-148), with participants holding an electrode in each hand whilst standing barefoot on two other electrodes. Participants were asked to consume 0.5 L of water 30 min before bioelectrical impedance assessments to standardize hydration status, having refrained from fluid consumption beforehand.

Maximal strength assessment
Participant knee extensor and flexor estimated 1RM strength was assessed for both legs separately. The leg reaching the highest estimated 1RM was assigned to the exercise intervention. Leg extension 1RM for both right and left leg was determined first, after which the protocol was repeated to determine leg flexor 1RM. Briefly, participants performed a one-set warm-up consisting of 12 repetitions at 10 kg. Thereafter, exercise load was gradually increased over subsequent sets until participants were unable to perform >10 repetitions. Increments in exercise load were based on subjective ratings of perceived exertion using the modified Borg category-ratio scale (CR-10). The Brzycki equation was used to estimate 1RM for both knee extensor and flexor strength.

D2O dosing protocol
The D2O dosing protocol consisted of a loading day and 12 maintenance days. On Day 1 of the trial and after providing a background saliva sample, participants consumed a loading dose of 70% D2O equalling three times their body mass in millilitres with the aim to label the body water pool to ~0.3 atom per cent excess (APE). The loading dose was split up in 50 mL doses and consumed every 30 min to avoid nausea and light-headedness. Body water enrichment was maintained in a pseudo steady state using daily top-ups based upon a 6% per day decay rate. Participants were instructed to provide a daily saliva sample upon waking, followed by consumption of the D2O maintenance dose. The D2O loading
and maintenance protocol was well tolerated, and no adverse effects were reported by the participants.

**Quadriiceps cross-sectional area**
Quadriiceps and vastus lateralis CSA was determined on Days 8 and 13 of the experimental trial using a 3 T MRI scanner (Phillips Achieva 3 T scanner) at the Birmingham University Imaging Centre (BUIC). Participants were placed on the scanner bed in a supine position and entered the scanner feet first. To ensure consistent positioning across pre-bed-rest and post-bed-rest MRI scans and participant comfort, participants’ feet were taped to internally rotate the toes, and sandbags were placed over the ankles. To aid alignment of pre-bed-rest and post-bed-rest MRI scans, cod liver oil tablets were taped every 5 cm on the lateral part of the limb starting at the fibula head running proximal up to the greater trochanter. Accurate placement of cod liver oil capsules was achieved by re-marking the sites of capsule placement from images obtained at 20% of the length between the top of the patella and the greater trochanter for each participant (OsiriX medical imaging software, OsiriX, Atlanta, USA).

**Sample analyses**

**Plasma amino acids, plasma isotope enrichment, and body water $^2$H enrichment**
Plasma $[^{13}C_6]$ phenylalanine enrichment was determined by gas chromatography–mass spectrometry (Model 5973; Hewlett Packard, Palo Alto, CA, USA) by monitoring ion 234/240. Briefly, 100 μL of plasma was diluted 2:1 with acetic acid before being purified through cation-exchange columns and eluted using 2 M of NH₄OH. Eluents were dried down under nitrogen and converted to their N-tert-butylidimethylsilyl-N-methyltrifluoracetamide (MTBSTFA) derivative. Leucine and phenylalanine concentrations were measured using internal standards U-[13C₆] leucine (ions 302/308) and U-[13C₆,15N] phenylalanine (ions 336/346). Body water enrichment was measured as described by Wilkinson et al. Briefly, 50 μL of saliva was heated in an inverted auto-sampler vial for 4 h at 100°C and placed upright on ice afterwards to condense extracted body water. This was then transferred to a clean auto-sampler vial, and a total of 0.1 μL of body water was injected into a high-temperature elemental analyser (Thermo Finnigan; Thermo Scientific, Hemel Hempstead, UK) connected to an isotopic ratio mass spectrometer (Delta V Advantage; Thermo Scientific).

**Isolation of myofibrillar protein fractions and protein-bound alanine and $^{13}$C₆ phenylalanine enrichment**
Myofibrillar proteins were extracted by homogenizing 20–30 mg of muscle in ice-cold homogenization buffer [50 mM of Tris-HCl (pH 7.4), 50 mM of NaF, 10 mM of β-glycerophosphate disodium salt, 1 mM of EDTA, 1 mM of ethylene glycol tetracetic acid, and 1 mM of activated Na₂VO₄ and a complete protease inhibitor cocktail tablet (Roche, West Sussex, UK)] at 10 μL/μg tissue and shaken for 10 min. Homogenates were spun at 1000 g for 5 min at 4°C, and the supernatant was collected. The myofibrillar fraction of the pellet was solubilized for 30 min at 37°C in 0.3 M of NaOH and separated from the insoluble collagen fraction by centrifugation. Myofibrillar proteins were precipitated using 1 M of perchloric acid and spun down at 3200 g for 20 min at 4°C before being hydrolysed overnight in 1 mL of 0.1 M of HCl and 1 mL of Dowex H⁺ Resin. Proteins were eluted from the resin using 2 M of NH₄OH and dried down at 70°C under a constant nitrogen flow. Amino acids for $^{13}$C₆ phenylalanine enrichment were derivatized as their n-acetyl-n-propyl ester, and labelling was determined using a Thermo Delta V isotope ratio mass spectrometer with a Thermo GC ultra and PAL auto-sampler, Thermo GC Combustion III interface, and Conflow IV interface. Amino acids for incorporation of deuterium into muscle-bound alanine were derivatized as their N-methoxy carbonyl methyl esters. Labelling was determined using chromatography: pyrolysis:isotope ratio mass spectrometry (Delta V Advantage) and ran alongside a standard curve of known DL-alanine-2.3.3.3-d₄ enrichment to ensure measurement accuracy of the machine.

**Gene expression analysis**
Gene expression analysis was performed as previously described. Briefly, RNA was isolated from ~20 mg of frozen powdered muscle homogenized in 1 mL of TRI Reagent (Sigma Aldrich, Gillingham, UK) and 200 μL of chloroform added to achieve phase separation. The RNA containing supernatant was removed and purified using RNeasy spin columns (Promega, Madison, Wisconsin, USA). RNA concentration and purity (ratio of the absorbance at 260 and 280 nm) and was ≥1.85 for all samples was determined using a FLUOstar Omega microplate reader; 700 ng of total RNA was reverse transcribed to cDNA in 20 μL volumes using the nanoScript 2 RT kit in combination with oligo (dT) and random primers (Primerdesign, Southampton, UK). cDNA was diluted to 5 ng/μL prior to RT-qPCR analysis. All analyses were performed in triplicate using Primerdesign custom-made primer sequences or commercially available 18S, B2M, GAPDH, and ACTB; 5 and 20 ng of cDNA was added, respectively, for housekeeping and human genes of interest to a 20 μL reaction volume. Thermal cycling conditions consisted of 2 min at 95°C, followed by 40 cycles of 10 s at 95°C and 60 s at 60°C. A melt curve was performed (Applied Biosystems, Thermo Fisher, UK) post-qPCR to assure primer specificity. Results were analysed using Thermo Fisher Connect (Thermo Fisher) and expressed as fold change relative to baseline using the $2^{-\Delta\DeltaCT}$ method. Data were
normalized to the geometric mean of the three most stable housekeeping genes (GAPDH, 18S, and ACTB) to minimize variation of the individual housekeeping genes. All gene expression results are presented for variation of the individual housekeeping genes. All gene expression targets were measured in muscle biopsy tissue obtained in the postabsorptive state.

Protein expression analysis
Protein expression was measured by western blot analysis on the sarcoplasmic protein fraction obtained during myofibrillar protein isolation. Gels were loaded according to the sarcoplasmic protein concentration assessed by the DC protein assay (Bio-Rad, Hertfordshire, UK), before aliquots of 2 μg/mL were prepared in 4× Laemml sample buffer and 100% D2O. Samples were boiled for 5 min, and equal amounts of protein (30 μg) were loaded into Criterion™ TGX™ Precast Midi protein gels (Bio-Rad, Hertfordshire, UK) and separated by sodium dodecyl sulfate–polyacrylamide gel electrophoresis at a constant voltage (100 V for 10 min followed by 150 V for 1 h). Protein samples were transferred to a polyvinylidene difluoride (Whatman, Dassel, Germany) membrane at 100 V for 1 h. The membranes were then incubated overnight (4°C) with appropriate primary antibodies; muscle ring finger protein 1 (MuRF1; sc-398608), muscle atrophy f-box (MAFbx; AM-3141), and actin (ACTB) to minimize a constant voltage (200 V for 150 min). Muscle cross sections were washed three times in 1× phosphate-buffered saline (PBS) and incubated in their respective secondary antibodies with wheat germ agglutinin Igg for 90 min. Finally muscle cross sections were washed in 1× PBS, and slides were mounted with Pro-Long Gold anti-fade reagent (P36930, Invitrogen). Images were captured using an Eclipse E600 (Nikon, Badhoevedorp, the Netherlands) and a 20× zoom. All images were analysed using ImageJ Fiji software.

Plasma insulin, glucose, triglycerides, and non-esterified fatty acids
Plasma insulin concentrations were analysed using a commercially available enzyme-linked immunosorbent assay according to the manufacturer’s instructions (R&D Systems, MN, USA). Plasma glucose was measured using a Roche Cobas 8000 analyser (Roche Diagnostics, Basel Switzerland). Serum total cholesterol, high-density lipoprotein cholesterol (HDL-C), triglyceride (TG), and non-esterified fatty acid (NEFA) were incubated overnight in myosin heavy chain type I (CST), total ribosomal protein S6 (rpS6), and phospho-Akt (pAkt) were determined as percentage per hour and percentage per day for [13C6] phenylalanine and [1H] alanine, respectively, with the use of the precursor-product equation, as previously described. The use of tracer naïve participants allowed us to use the pre-infusion plasma [13C6] phenylalanine enrichment as a proxy for basal muscle protein enrichment for measurement of postabsorptive aMyoPS rates. This approach has been validated for use in older individuals.

Calculations
The iMyoPS and aMyoPS fractional synthetic rate (FSR) was determined as percentage per hour and percentage per day for [13C6] phenylalanine and [1H] alanine, respectively, with the use of the precursor-product equation, as previously described. The use of tracer naïve participants allowed us to use the pre-infusion plasma [13C6] phenylalanine enrichment as a proxy for basal muscle protein enrichment for measurement of postabsorptive aMyoPS rates. This approach has been validated for use in older individuals.

Statistical analyses
Anthropometric, physical activity, dietary characteristics, and blood hormone/analytes were analysed using a paired-samples t-test. Quadriceps CSA, fibre-type morphology, gene expression, and iMyoPS and aMyoPS were analysed using a two-way repeated-measures ANOVA (condition × time) with condition (CTRL vs. EX) and time (pre-bed rest vs. post-bed
rest). Body water $^2$H enrichment, plasma amino acid, and $^{13}$C$_6$ phenylalanine were analysed using a one-way repeated-measures ANOVA with time as the within-subject factor. Delta and percentage change in quadriceps CSA, iMyoPS, and aMyoPS for CTL and EX were analysed using a Student’s t-test. Bonferroni post-hoc tests were performed to correct for multiple comparisons when a significant condition × time interaction was identified. All analyses were performed using SPSS 26 (SPSS, Chicago, IL, USA). Significance was set at $P \leq 0.05$. All data are presented as mean ± SD unless otherwise indicated.

## Results

### Anthropometric, physical activity, and dietary characteristics

Participant body mass and BMI did not change from pre-bed rest to post-bed rest. However, following bed rest, a decrease in relative fat-free mass (~5%; $P = 0.02$) and appendicular lean mass (~6%; $P = 0.002$) and an increase in relative fat mass (~13%; $P = 0.02$) were observed. Average daily step-count and percentage of daily time spent performing light-intensity and moderate-intensity activities were significantly lower during bed rest compared with habitual levels ($P < 0.001$ for all). Percentage of daily sedentary time increased significantly during bed rest (~27%; $P < 0.001$), whilst light and moderate activity decreased (both $P < 0.001$). Percentage of daily vigorous activity did not significantly change during bed rest. During bed rest, total energy ($P = 0.025$), dietary protein ($P = 0.01$), and fibre ($P = 0.001$) intake significantly decreased from habitual levels. Anthropometric, physical activity, and dietary characteristics are presented in Table 1.

### Maximal strength and resistance training parameters

Estimated 1RM strength was 64.7 ± 14.8 and 58.6 ± 7.7 for the leg extension and leg curl machines, respectively. The average total training volume over the four sessions of prehabilitation RET was 22 407 ± 3340 kg, performed at an average Borg CR-10 rating of 8.1 ± 1.0. RET prehabilitation data are presented in Table 2.

### Quadriceps cross-sectional area and fibre-type morphology

Quadriceps CSA significantly decreased from pre-to-post bed rest at 40%, 60%, and 80% ($P < 0.01$ for all), but not at 20% of the distance measured between the top of the patella and greater trochanter in CTL and EX. Vastus lateralis CSA significantly decreased from pre-to-post bed rest at 40% and 60% ($P < 0.05$ for all), but not at 20% and 80% of the distance measured between the top of the patella and greater trochanter in CTL and EX. No differences for quadriceps or vastus lateralis CSA between EX and CTL were found before or after bed rest. Furthermore, there was no significant difference between EX and CTL in the relative change in quadriceps CSA or vastus lateralis CSA from pre-to-post bed rest at any anatomical muscle length. Quadriceps and vastus lateralis CSA are presented in Table 3 and Figure 2. To determine

### Table 1 Participant anthropometric, activity, and dietary characteristics before and during/after bed rest

| Characteristic         | Pre-bed rest          | Peri-to-post-bed rest | P-value |
|------------------------|-----------------------|-----------------------|---------|
| Age (years)            | 71.5 ± 4.0            | 79.0 ± 8.4            | 0.212   |
| Height (m)             | 1.77 ± 0.07           | 25.3 ± 2.7            | 0.207   |
| Weight (kg)            | 79.6 ± 9.0            | 27.1 ± 4.0*           | 0.021   |
| BMI (kg/m$^2$)         | 25.5 ± 2.8            | 72.9 ± 4.0*           | 0.022   |
| Body Fat (%)           | 23.3 ± 4.9            | 7.45 ± 0.85           | 0.002   |
| Fat-free mass (%)      | 76.7 ± 5.0            | 7.90 ± 0.96           |         |
| ALM (kg/m$^2$)         | 7.90 ± 0.96           | 7.45 ± 0.85           |         |
| SPPB                   | 12 ± 1.1              | 134 ± 161*            | <0.001  |
| Daily step-count       | 10 177 ± 3695         | 89.40 ± 6.87*         | <0.001  |
| Sedentary activity (%) | 70.67 ± 9.38          | 4.83 ± 2.30*          | <0.001  |
| Light activity (%)     | 12.32 ± 3.85          | 5.70 ± 4.98*          | <0.001  |
| Moderate activity (%)  | 16.47 ± 6.85          | 0.08 ± 0.11           | 0.014   |
| Vigorous activity (%)  | 0.67 ± 1.21           | 1990 ± 369*           | 0.025   |
| Total energy intake (kcal) | 2451 ± 750         | 0.97 ± 0.25*          | 0.004   |
| Protein intake (g/kg/day) | 1.37 ± 0.47          | 2.73 ± 0.54           | 0.932   |
| CHO intake (g/kg/day)  | 2.71 ± 1.08           | 1.12 ± 0.24           | 0.521   |
| Fat intake (g/kg/day)  | 1.19 ± 0.49           | 0.10 ± 0.06*          | <0.001  |
| Fibre intake (g/kg/day)| 0.39 ± 0.19           | 0.00 ± 0.00           | 0.085   |

Step-count, physical activity, and dietary intake data are daily averages obtained over 5 days of bed rest. Values are means ± SD for $n = 10$ participants.

ALM, appendicular lean mass; BMI, body mass index; SPPB, Short Physical Performance Battery.

*Significantly different from corresponding pre-bed-rest value ($P < 0.05$).

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Journal of Cachexia, Sarcopenia and Muscle 2021; 12: 52–69
DOI: 10.1002/jcsm.12661

B. Smeuninx et al.
Table 2: Leg strength and resistance training parameters for leg extension and leg curl exercises

| Parameter               | Leg extension | Leg curl  |
|-------------------------|---------------|-----------|
| Estimated 1RM (kg)      | 64.7 ± 14.8   | 58.6 ± 7.7|
| Avg load per set (kg)   | 41.3 ± 8.7    | 36.7 ± 4.9|
| Total load (kg)         | 986 ± 203     | 880 ± 118 |
| Avg repetitions per set | 12.0 ± 0.1    | 12.0 ± 0.0|
| Total repetitions       | 287 ± 1       | 288 ± 1   |
| Avg volume per set (kg) | 494 ± 106     | 440 ± 58  |
| Total volume (kg)       | 11 857 ± 2541 | 11 500 ± 1390|
| T-U-T total (s)         | 551.8 ± 3.5   | 23.0 ± 2.9|
| T-U-T per set (s)       | 23.0 ± 3.5    | 23.0 ± 2.9|
| Avg Borg CR-10          | 8.2 ± 1.0     | 8.0 ± 0.9 |

Values are means ± SD for n = 10 participants. 1RM, one-repetition maximum strength; T-U-T, time under tension.

Blood hormones and analytes

No significant differences between fasting pre-bed-rest and post-bed-rest values were found for total cholesterol to HDL ratio and concentrations of plasma insulin, glucose, serum total cholesterol, serum HDL, serum non-HDL, serum NEFA, and serum TG. Homeostatic Model Assessment of Insulin Resistance (HOMA-IR), derived from fasting glucose and insulin, did not differ between pre-bed-rest and post-bed-rest. Blood hormone and analyte data are presented in Table 4.

Plasma amino acid, $^{13}$C$_6$ phenylalanine, and $^2$H body water enrichment

Plasma leucine concentrations were significantly elevated above postabsorptive values at 20, 40, 60, and 90 min post-drink consumption (Figure 3(A)), whilst plasma phenylalanine concentrations were significantly elevated above postabsorptive values at 20, 40, 60, and 90 min post-drink consumption (Figure 3(B)). Plasma $^{13}$C$_6$ phenylalanine enrichment significantly increased above basal values 180 min after initiation of the stable isotope tracer infusion and remained elevated for the duration of the trial ($P < 0.001$; Figure 3(C)). Plasma $^{13}$C$_6$ phenylalanine enrichment at 180 min post-drink consumption was significantly different to values at 20, 40, 60, and 90 min post-drink consumption. Nonetheless, linear regression analysis revealed that the $^{13}$C$_6$ phenylalanine enrichment slopes in both groups were not significantly different from zero. Body water $^2$H enrichment, assessed via saliva, was 0.31 ± 0.02 APE at 24 h after the first bolus D$_2$O dose (Figure 3(D)). Body water $^2$H enrichment averaged 0.29 ± 0.02 APE over the duration of the study. Linear regression analysis indicated that the slope of the body water enrichment curve was significantly different from zero ($r^2 = 0.41; P = 0.025$).

Integrated myofibrillar protein synthesis

A significant main effect for time was found ($P = 0.001$), whilst no interaction effect was observed. Rates of iMyoPS were significantly elevated in EX compared with CTL during the prehabilitation phase ($1.36 ± 0.18%/day vs. 1.76 ± 0.37%/day; P = 0.007$). Both CTL and EX iMyoPS rates significantly decreased during bed rest compared with prehabilitation to 1.07 ± 0.22%/day ($P = 0.037$) and 1.30 ± 0.38%/day ($P = 0.002$), respectively. iMyoPS rates did not differ between EX and CTL over the bed-rest phase. The decrease in iMyoPS from prehabilitation to bed-rest values was not significantly different between EX and CTL (Figure 4(A) and 4(B)).

Acute myofibrillar protein synthesis

A significant main effect for time ($P = 0.005$) was found with no apparent interaction effect. Whilst not significant, there

Table 3: Quadriceps and fibre cross-sectional area for non-exercised control and exercised legs measured prior to and following 5 days of bed rest

| Parameter               | Pre-bed rest | Post-bed rest | P-value | Pre-bed rest | Post-bed rest | P-value |
|-------------------------|--------------|---------------|---------|--------------|---------------|---------|
| Quadriceps CSA 20% (mm$^2$) | 4770 ± 649   | 4760 ± 641    | 0.542   | 4787 ± 600   | 4774 ± 620    | 0.465   |
| Quadriceps CSA 40% (mm$^2$) | 6823 ± 677   | 6776 ± 663*   | 0.004   | 6855 ± 692   | 6809 ± 671*   | 0.005   |
| Quadriceps CSA 60% (mm$^2$) | 7168 ± 826   | 6917 ± 717*   | 0.001   | 7260 ± 688   | 7040 ± 783*   | 0.002   |
| Quadriceps CSA 80% (mm$^2$) | 5086 ± 759   | 4963 ± 733*   | <0.001  | 5148 ± 679   | 5027 ± 679*   | <0.001  |
| VL CSA 20% (mm$^2$)       | 1198 ± 145   | 1186 ± 119    | 0.429   | 1203 ± 116   | 1199 ± 138    | 0.560   |
| VL CSA 40% (mm$^2$)       | 1889 ± 193   | 1860 ± 157*   | 0.021   | 1919 ± 184   | 1892 ± 193*   | 0.018   |
| VL CSA 60% (mm$^2$)       | 2246 ± 338   | 2158 ± 325*   | <0.001  | 2316 ± 300   | 2221 ± 357*   | 0.007   |
| VL CSA 80% (mm$^2$)       | 1290 ± 266   | 1271 ± 185    | 0.083   | 1338 ± 202   | 1315 ± 174    | 0.064   |
| Type I fibre CSA (µm$^2$) | 6162 ± 2100  | 5638 ± 1175   | 0.397   | 6222 ± 1689  | 5703 ± 2281   | 0.466   |
| Type II fibre CSA (µm$^2$) | 6089 ± 2343  | 5743 ± 1156   | 0.505   | 5863 ± 1689  | 5577 ± 1530   | 0.633   |

Values are means ± SD for n = 10 participants. CSA, cross-sectional area; VL, vastus lateralis. *Significantly different from corresponding pre-bed-rest value ($P < 0.05$).
was a trend for greater aMyoPS rates in EX vs. CTL in the postabsorptive (0.022 ± 0.012%/h vs. 0.032 ± 0.009%/h; \( P = 0.054 \)) and postprandial (0.026 ± 0.012%/h vs. 0.038 ± 0.012%/h; \( P = 0.052 \)) states. aMyoPS rates significantly increased from the postabsorptive to postprandial state in EX only (\( P = 0.018 \)). The postprandial change in aMyoPS from the postabsorptive state did not differ between EX and CTL (Figure 4(C) and 4(D)).

**Gene expression**

Both and p70S6K1 (Figure 5(A)) and mTOR (Figure 5(B)) gene expression was significantly increased after bed rest compared with prehabilitation in CTL only (\( P = 0.026 \) and \( P = 0.047 \), respectively). Whilst no differences between EX and CTL were found for p70S6K1, mTOR gene expression was significantly higher in CTL vs. EX after bed rest (\( P = 0.002 \)). Myostatin mRNA expression was significantly increased in both CTL (\( P = 0.001 \)) and EX (\( P = 0.023 \)) following bed rest, with no difference between EX and CTL (Figure 5(C)). In regard to genes associated with proteolysis, only MAFbx (Figure 5(D)) was significantly higher in CTL compared with EX following prehabilitation (\( P = 0.004 \)). There was a significant main effect of time on MAFbx expression from prehabilitation to bed rest (\( P = 0.016 \)). No significant group, time, or interaction effects were found for MuRF1 mRNA expression (Figure 5(E)).

**Figure 2** Percentage change in quadriceps cross-sectional area during 5 days of bed rest in healthy older men in a leg that had undergone resistance exercise prehabilitation over the preceding 7 days (EX) or the contralateral non-exercised control leg (CTL). Magnetic resonance imaging was obtained at 20%, 40%, 60%, and 80% of the length between the top of the patella and the greater trochanter (distal to proximal). Between-leg differences were analysed using a Student’s paired t-test. Boxes represent the 25th to 75th percentiles, error bars represent SEM, and horizontal lines and crosses within boxes represent median and mean values, respectively (\( n = 9 \)). Significance was set at \( P < 0.05 \). A significant reduction in quadriceps cross-sectional area (CSA) was noted at 40%, 60%, and 80% of muscle length (*\( P < 0.01 \), **\( P < 0.001 \)), with no between-group differences observed at any length.

**Table 4** Fasting blood hormone and analyte concentrations prior to and following 5 days of bed rest

|                      | Pre-bed rest | Post-bed rest | \( P \)-value |
|----------------------|--------------|---------------|---------------|
| Plasma insulin (pmol/L) | 38.8 ± 19.4  | 46.4 ± 22.6   | 0.094         |
| Plasma glucose (mmol/L) | 5.29 ± 0.72  | 5.45 ± 0.82   | 0.116         |
| HOMA-IR              | 1.54 ± 0.90  | 1.92 ± 1.08   | 0.070         |
| Serum total cholesterol (mmol/L) | 5.11 ± 1.24  | 4.50 ± 0.77   | 0.073         |
| Serum HDL-C (mmol/L) | 1.65 ± 0.44  | 1.44 ± 0.38   | 0.588         |
| Non-HDL-C (mmol/L) | 3.46 ± 1.12  | 3.06 ± 0.68   | 0.124         |
| Total cholesterol:HDL-C ratio | 3.20 ± 0.73  | 3.26 ± 0.78   | 0.588         |
| Serum NEFA (mmol/L) | 0.49 ± 0.14  | 0.46 ± 0.17   | 0.390         |
| Serum triglycerides (mmol/L) | 1.07 ± 0.36  | 1.11 ± 0.46   | 0.602         |

Values are means ± SD for \( n = 10 \) participants.

HDL-C, high-density lipoprotein cholesterol; HOMA-IR: Homeostatic Model Assessment of Insulin Resistance; NEFA, non-esterified fatty acids.
Protein expression

Following bed rest, there was a significant main effect of time on Akt$^{473}$ (Figure 6(A)) and mTOR$^{2448}$ expression (Figure 6(C)), which were greater than CTL values after prehabilitation ($P < 0.01$ and $P < 0.001$, respectively), with no difference between legs. Following bed rest, there was a significant main effect of time on 4E-BP1$^{137/46}$ expression, which was lower than CTL values after prehabilitation ($P = 0.04$; Figure 6(E)), with no difference between legs. A significant main effect of leg was observed with rpS6$^{240/244}$ expression following exercise prehabilitation and bed rest (Figure 6(G), $P = 0.004$) and between the postabsorptive and postprandial states following bed rest (Figure 6(H), $P = 0.033$). Post-hoc analysis revealed that, following exercise prehabilitation and bed rest, rpS6$^{240/244}$ expression (Figure 6(G)) was significantly higher in EX vs. CTL ($P = 0.011$ and 0.043, respectively). Further, following bed rest, rpS6$^{240/244}$ expression (Figure 6(H)) was significantly higher in EX vs. CTL in the postabsorptive and postprandial states ($P = 0.043$ and 0.013, respectively). Following bed rest, Akt$^{473}$ expression was significantly lower than the postabsorptive state in EX only ($P = 0.015$; Figure 6(B)). No significant group, time, or interaction effects were found for MuRF1 or MAFbx protein expression (Figure 6(I)–(L), all $P > 0.05$). Representative western blot images are presented in Figure S2.

Discussion

Age-related skeletal muscle loss (sarcopenia) is driven by impairments in muscle protein turnover that are accelerated during periods of disuse (i.e. hospitalization). Muscle mass/attenuation is associated with length of stay in hospital, risk of readmission, recovery of function, and mortality. Given the treatment costs associated with sarcopenia, and the increasing prevalence of this condition, attenuating...
muscle mass loss in older individuals during an in-hospital stay is crucial to maintain functional and metabolic health and lower the strain on health services (e.g. decreased bed occupancy and external care provision). Compared with peri-disuse or post-disuse interventions, prehabilitation capitalizes on better patient health and could potentially benefit a number of clinically relevant outcomes. Here, we report that short-term RET prehabilitation augmented iMyoPS over the week prior to 5 days of in-hospital bed rest in older individuals. However, the relative decline in iMyoPS with bed rest was not offset by our prehabilitation protocol, and quadriceps muscle atrophy was not abated. In support of these findings, although postabsorptive and postprandial aMyoPS rates tended to be higher with RET prehabilitation, the net postprandial aMyoPS response was not enhanced/maintained.

The removal of muscle contractile activity in older individuals undergoing bed rest results in blunted rates of postprandial and postabsorptive aMyoPS, which drives the reduction in iMyoPS rates (i.e. incorporating postabsorptive and postprandial aMyoPS over a longer-term) reported during disuse events. RET is a potent stimulus for aMyoPS that, when undertaken routinely, augments iMyoPS for muscle protein accrual. A single bout of RET enhances postprandial aMyoPS for at least 24 h afterwards, whilst the accumulation of six bouts of low-load high-volume RET over 14 days augments postprandial aMyoPS in older individuals.

Figure 4 Integrated myofibrillar protein fractional synthesis rates over the course of 7 days of resistance exercise prehabilitation and 5 days of bed rest in exercised (EX) and non-exercised (CTL) legs in healthy older men (A). Delta change in integrated myofibrillar protein fractional synthesis rates from prehabilitation to bed rest in EX and CTL (B). Acutely measured myofibrillar protein synthesis rates in the postabsorptive and postprandial state, after ingestion of 15 g of milk protein, in EX and CTL (C). Delta change in acute myofibrillar protein fractional synthesis rates from postabsorptive to postprandial state in EX and CTL (D). Values in (A) and (C) are means ± SEM and individual participant data (n = 10). (B) and (D) The boxes represent the 25th to 75th percentiles, error bars represent SEM, and horizontal lines and crosses within boxes represent median and mean values, respectively (n = 10). Data in (A) and (C) were analysed using a two-way repeated-measures ANOVA (condition × time) with condition (CTL vs. EX) and time (prehabilitation vs. post-bed rest). Data in (B) and (D) were analysed using a Student’s t-test. Bonferroni post-hoc tests were performed to correct for multiple comparisons when a significant condition × time interaction was identified. Significance was set at P < 0.05. Integrated myofibrillar protein synthesis (iMyoPS) was higher in EX vs. CTL during prehabilitation (#P < 0.01) and was lower during bed rest vs. prehabilitation in EX (**P < 0.01) and CTL (*P < 0.05), with no difference between groups. Acute postabsorptive and postprandial myofibrillar protein synthesis (aMyoPS) was stimulated above postabsorptive values in EX (*P < 0.05), with no differences between groups. † indicates a significant time effect of bed rest vs. prehabilitation (P < 0.05).
individuals, when measured 72 h after the final bout. Thus, the present study sought to determine whether four sessions of moderate-load RET prehabilitation over 7 days (higher total volume and a shorter time frame than our earlier work) would rescue iMyoPS, and consequently muscle mass, through repeatedly elevating postprandial aMyoPS in older individuals during bed rest. As expected, RET increased iMyoPS by 23% over the prehabilitation phase compared with the non-exercised CTL, analogous to the increase in iMyoPS reported in younger adults with several bouts of RET. Interestingly, the gene expression of p70S6K and mTOR increased above basal values during bed rest in CTL only (*P < 0.05 for both). There expression of mTOR after bed rest was significantly lower in EX vs. CTL (#P < 0.05). Myostatin expression was greater after bed rest compared with prehabilitation in EX and CTL (*P < 0.05) with no difference between legs. MAFbx expression was greater in CTL vs. EX after prehabilitation (#P < 0.05). There was a significant main effect of time on MAFbx expression from prehabilitation to bed rest (P = 0.016).

Figure 5 Changes in mRNA expression of p70S6K (A), mTOR (B), myostatin (C), MAFbx (D), and MuRF1 (E) after 7 days of resistance exercise prehabilitation and a subsequent 5 days of bed rest in older men. Data are expressed as the fold change from levels measured in non-exercised control leg (CTL) after prehabilitation, which was normalized to a value of 1. All targets were measured in muscle biopsy tissue obtained in the postabsorptive state. Boxes represent the 25th to 75th percentiles, error bars represent SEM, and horizontal lines and crosses within boxes represent median and mean values, respectively [n = 9 for all targets except p70S6K (n = 8)]. Significance was set at P < 0.05. There was a significant increase in p70S6K and mTOR expression after bed rest compared with prehabilitation for CTL only (*P < 0.05 for both). There expression of mTOR after bed rest was significantly lower in EX vs. CTL (#P < 0.05). Myostatin expression was greater after bed rest compared with prehabilitation in EX and CTL (*P < 0.05) with no difference between legs. MAFbx expression was greater in CTL vs. EX after prehabilitation (#P < 0.05). There was a significant main effect of time on MAFbx expression from prehabilitation to bed rest (P = 0.016).
4E-BP1 with bed rest did not differ between EX and CTL. Collectively, the similar relative changes to iMyoPS and anabolic signalling expression over the bed rest period in CTL and EX might partially explain the similar degree of quadriceps muscle loss as measured by MRI. Interestingly, rpS6 phosphorylation was greater in EX vs. CTL after prehabilitation and bed rest, but this clearly did not impact on the extent of disuse atrophy.

The repeated daily stimulation of postprandial aMyoPS over time is considered the main locus of muscle mass regulation, which is further enhanced by RET. Prehabilitation could, therefore, provide a stimulus to bolster/retain muscle anabolic sensitivity to subsequent feeding occasions during bed rest. The ability of prehabilitation to chronically augment postprandial aMyoPS was investigated after the final day of bed rest. Ingestion of 15 g of milk protein isolate (~0.19 g/kg/BW) significantly increased aMyoPS in EX only, whilst no increase was observed in CTL. The latter aligns with observations of muscle anabolic resistance in older individuals after 5 days of bed rest without prior RET. On the other hand, although a trend for higher postabsorptive (~32%; $P = 0.054$) and postprandial (~31%; $P = 0.052$) aMyoPS rates was observed in EX vs. CTL, the relative increase in aMyoPS from prehabilitation to bed rest did not differ between legs. Furthermore, with the exception of Akt, there was no postprandial change in anabolic signalling.

Figure 6 Changes in protein expression of Akt$^{473}$, mTOR$^{2448}$, 4E-BP1$^{137/46}$ and rpS6$^{2420/244}$, MuRF1, and MAFbx following 7 days of resistance exercise prehabilitation (EX) and a subsequent 5 days of bed rest in older men. Akt$^{473}$, mTOR$^{2448}$, 4E-BP1$^{137/46}$, and rpS6$^{2420/244}$ are expressed relative to respective total protein expression. MuRF1 and MAFbx are expressed relative to ponceau loading control. (A), (C), (E), (G), (I), and (K) show protein expression in EX and CTL after exercise prehabilitation and after bed rest and are expressed as the fold change from levels measured in control (CTL) after prehabilitation, which was normalized to a value of 1 (all targets were measured in a postabsorptive state). (B), (D), (F), (H), (L), and (J) show protein expression in EX and CTL after bed rest in the postabsorptive (BR-PA) or 4 h postprandial state after ingestion of 15 g of milk protein (BR-PP), expressed as the fold change from levels measured in CTL after bed rest, which was normalized to a value of 1. For all panels, boxes represent the 25th to 75th percentiles, error bars represent SEM, and horizontal lines and crosses within boxes represent median and mean values, respectively ($n = 9$ for all targets except Akt, mTOR, rpS6 ($n = 8$)). Significance was set at $P < 0.05$. Following bed rest, there was a significant main effect of time on Akt$^{473}$ and mTOR$^{2448}$ expression, which was greater than CTL values after exercise prehabilitation ($^{*}P < 0.05$ for both, (A)/A and (C), respectively), with no difference between legs. Following bed rest, there was a significant main effect of time on 4E-BP1$^{137/46}$ expression, which was significantly lower than CTL values after exercise prehabilitation ($^{*}P < 0.05$, (E)), with no difference between legs. Following exercise prehabilitation and bed rest, rpS6$^{2420/244}$ expression was significantly higher in EX vs. CTL, $^{*}P < 0.05$ for both, (I)). Following bed rest, rpS6$^{2420/244}$ expression was significantly higher in EX vs. CTL in the postabsorptive and postprandial state ($^{*}P < 0.05$ for both, (H)). Following bed rest, Akt$^{473}$ expression was significantly lower than the postabsorptive state in EX only ($^{*}P < 0.05$, (B)).
phosphorylation in EX or CTL, and no differences in postpran- 
dial anabolic signalling between legs. Therefore, it appears 
that moderate-load high-volume RET prehabilitation, under-
taken over 7 days, was insufficient to favourably influence 
postprandial aMyoPS stimulation or anabolic signalling over 
a subsequent 5 day bed-rest period in older individuals. We 
acknowledge that any potential effect of RET prehabilitation 
on postprandial aMyoPS and anabolic signalling responsive-
ness may have dissipated over the course of bed rest and 
been absent by the time of assessment (6 days after the final 
bout). This opens the possibility that short-term RET 
prehabilitation could alleviate muscle anabolic resistance 
and atrophy in older individuals during very short periods of 
disuse.\textsuperscript{16,40}

The possibility that net muscle protein deposition during 
RET prehabilitation could counteract muscle loss during sub-
sequent disuse is based on the assumption that all synthe-
sized proteins are incorporated into bound myofibrillar 
protein and that muscle protein breakdown (MPB) remains 
unchanged. If this holds true, the absolute increase in 
iMyoPS over 7 days of prehabilitation in the present study 
(2.8% or 0.4% daily) would have buffered against muscle 
mass loss during bed rest (~0.7%, 3.3% and 2.4% at 40%, 
60%, and 80% of muscle length, respectively). Unfortunately, 
we were unable to perform MRI scans at baseline to better 
understand the influence of short-term RET prehabilitation 
on quadriceps CSA. A more viable explanation for the disso-
ciation between elevated iMyoPS in EX and the observed 
muscle loss is the redirection of newly synthesized proteins 
toward RET-induced muscle damage repair rather than mus-
cle hypertrophy per se, as has been reported in younger 
men in response to a similar RET stimulus as that used 
herein.\textsuperscript{41} Furthermore, given that a single bout of 
high-volume RET elevates MPB over 24 h post-exercise,\textsuperscript{32,43} 
a cumulative MPB response over the 7 day prehabilitation 
phase in the present study would detract from overall net 
muscle protein balance. Nonetheless, we did not observe 
any difference in the protein expression of the ubiquitin li-
gases, MAFbx or MuRF1, between EX and CTL after 
prehabilitation. Paradoxically, MAFbx gene expression was 
lower after prehabilitation in EX vs. CTL, suggestive of atten-
umated MPB. However, in light of evidence suggesting that 
MAFbx expression may increase in the immediate hours af-
after RET, before falling below basal levels at 24 and 72 h 
post-exercise,\textsuperscript{44} this finding may be more indicative of active 
remodelling processes in EX vs. CTL. Additional biopsy and 
blood sampling would have allowed us to better reconcile 
the role of muscle damage and proteolytic signalling in re-
modelling processes during short-term exercise 
prehabilitation and bed rest. Notwithstanding, it is unlikely 
that our short-term RET prehabilitation programme was suf-
icient for the level of net muscle protein accretion that 
would effectively cancel out the subsequent muscle loss dur-
ing bed rest in older individuals.

Whilst impaired aMyoPS is regarded as the primary mech-
anistic driver of disuse-induced muscle atrophy, evidence of a 
transient up-regulation in genes associated with proteolysis 
in the first several days of disuse in older\textsuperscript{8,40} implicates a role 
of elevated MPB in disuse atrophy. Given the effect of RET on 
MPB,\textsuperscript{42,43} it is possible that short-term RET prehabilitation 
may have influenced the MPB response to bed rest in older 
individuals in the present study. In contrast to Tanner \textit{et al.},\textsuperscript{8} 
we did not observe an increase in the gene or protein expres-
sion of MAFBx and MuRF1, after 5 days of bed rest in older 
individuals, despite comparable levels of muscle loss. How-
ever, we did observe an increase in myostatin, a negative reg-
ulator of muscle mass associated with MPB,\textsuperscript{46} in EX and CTL. 
Based on these findings, it is not possible to determine the 
time course of change in MPB over 5 days of bed rest in older 
individuals, or the contribution of MPB to the observed 
muscle loss. Regardless, there is no evidence to suggest that MPB 
differed between legs. Given that the relative change in 
iMyoPS and aMyoPS during bed rest was similar between 
EX and CTL, any difference in MPB between legs would have 
influenced overall net protein balance and muscle atrophy.

Mobility during hospitalization is a central modifiable fac-
tor in preventing in-hospital functional decline and 
post-discharge adverse outcome in older adults.\textsuperscript{47} Average 
daily step-count in hospitalized older patients is ~600–1000 
during a 5 to 6 day inpatient stay,\textsuperscript{48,49} whilst >80% of time 
is typically spent lying and only ~3% standing or walking.\textsuperscript{50} 
In the present study, step-count in older individuals during 
5 days of bed rest was ~98% lower than habitual levels, and 
sedentary time increased at the expense of a reduction in ac-
tivity levels. These very low step-count and activity data were 
achieved by introducing a wheelchair to replace walking for 
most basic tasks and, hence, are more reflective of 
in-hospital values for frail older individuals with functional 
limitations.\textsuperscript{51} The dramatic alterations in activity during bed 
rest resulted in a loss of quadriceps CSA at 40%, 60%, and 
80% of muscle length (~0.7%, 3.5%, and 2.8%, respectively), 
which is broadly comparable with reports of an ~4% reduc-
tion in dual X-ray absorptiometry-derived whole-leg lean 
mass following 5 days of bed rest in older individuals.\textsuperscript{8} Con-
gruent with the present data, greater quadriceps atrophy at 
the proximal end of the muscle has been reported toward 
younger individuals during prolonged bed rest.\textsuperscript{52} A possible 
 explanation for the heterogeneity in quadriceps and vastus 
lateralis atrophy at various muscle lengths may relate to dif-
fferences in the function of certain muscle sub-regions in daily 
life and their consequent change in activity during bed rest. 
However, the relationship between quadriceps atrophy and 
anatomical length may depend on age status and the model 
of disuse, as others have observed distal quadriceps atrophy 
at 20% of muscle length in younger individuals after 7 days 
of leg immobilization.\textsuperscript{40} The decrease in quadriceps CSA was 
not confirmed at the microscopic level, which may have been 
due to the relatively small sample size. Indeed, similar
observations of a reduction in quadriceps CSA, but not fibre CSA, with short-term bed rest have been reported. Another possible explanation could be that biopsy samples were obtained in a distal to proximal orientation (~2–3 cm between biopsies). As fibre CSA is not constant throughout the muscle length, it is imperative for future studies to sample skeletal muscle from the same location in order to track changes in fibre CSA with disuse.

Dietary protein intake stimulates aMyoPS and, therefore, plays a pivotal role in muscle mass maintenance in ageing and disuse. Higher protein intakes are associated with greater long-term lean mass retention in older individuals, whereas a substantial increase in protein intake, on top of a diet providing the protein recommended dietary allowance or lower (<0.8 g/kg/BW/day), attenuates muscle loss in older individuals during bed rest. To mimic a typical inpatient stay in the present study, we provided participants with a choice of typical hospital meals/snacks. Protein intake during bed rest averaged 1.0 g/kg/BW/day, which was ~23% lower than habitual levels but greater than a real-world hospital setting, where older patients are often protein malnourished. The reduction in total energy and dietary protein intake during bed rest in the present study may be reflective of reduced energy requirements and/or lower nutritional quality of hospital meals. In support of the latter, dietary fibre intake was lower than habitual level during bed rest. Given that an increase in dietary protein during short-term bed rest, on top of a diet containing >1.0 g/kg/BW/day, does not attenuate muscle disuse atrophy in healthy older men, we contend that the reduction in dietary protein intake from habitual levels during bed rest would not have accelerated muscle loss. Because the majority of food intake during bed rest came from pre-prepared mixed-ingredient meals, we were unable to determine potential changes in dietary protein quality. This point is noteworthy, as recent evidence suggests that improving dietary protein quality during short-term bed rest offsets the decline in muscle mass and function in older individuals.

Although daily energy intake (EI) during bed rest was lower than habitual levels, the dramatic reduction in physical activity (and energy expenditure) may have left participants in a hypercaloric state, which may explain the observed increase in body fat. Thus, we cannot discount the notion that a hypercaloric state and increased fat mass could have influenced muscle atrophy. However, recent work showed that high-fat overfeeding exacerbated muscle amino acid balance but did not aggravate muscle atrophy during short-term forearm immobilization in young individuals. Furthermore, despite the increase in body fat after bed rest in the current study, lipid profiles remained unchanged. This is important, as elevated circulating lipids may underlie the postprandial muscle anabolic resistance reported in obese older adults. Thus, we speculate that the alterations in macronutrient/EI during bed rest had minimal impact on the observed muscle atrophy, although further clarification is required. The disconnect between body fat and lipid profile changes with bed rest may be explained by the limitations of BIA for body composition assessment. Pre-bed-rest and post-bed-rest BIA scans were conducted in the morning and afternoon, respectively (owing to logistical issues in the testing schedule), which could have led to slight discrepancies in fluid status and an overestimation in fat mass. Irrespective, any influence of alterations in macronutrient/EI during bed rest on muscle loss would have impacted EX and CTL similarly. Taken together, the alterations in macronutrient/EI observed during bed rest in our healthy older cohort reinforce the need to develop appropriate nutritional practices to support whole-body and muscle health during an in-hospital stay.

Although short-term RET prehabilitation did not offset disuse-induced muscle atrophy, there may be potential benefits for metabolic health. Bed rest and inactivity induce rapid insulin resistance and glucose dysregulation, which may underpin the progressive decline in metabolic health with advancing age. In contrast, plasma insulin, glucose, or HOMA-IR did not significantly change from pre-bed-rest to post-bed rest in the present study, although there was a trend for an increase in HOMA-IR. Thus, it is possible that short-term RET prehabilitation might offer some protection against disuse-induced declines in peripheral insulin sensitivity and glucose regulation, a process that usually occurs after the first few days of inactivity. The potential for unilateral RET prehabilitation to support whole-body insulin sensitivity in older adults during short-term disuse has important clinical implications and requires further exploration.

In conclusion, the present study is the first to investigate the influence of short-term RET prehabilitation on muscle morphology and associated regulatory mechanisms in older individuals during a period of disuse. Four bouts of single-leg high-volume RET, performed over 7 days prior to 5 days of bed rest, augmented iMyoPS as compared with the non-exercised leg. However, the relative reduction in iMyoPS during bed rest was similar between legs, as was the loss of quadriceps muscle mass. RET prehabilitation had no clear influence on relative postprandial aMyoPS stimulation compared with the non-exercised leg, when measured at the end of bed rest. The findings suggest that prior RET of sufficiently high-volume and duration may be required to build a muscle mass reserve to buffer against the atrophic response to a typical inpatient hospital stay. Irrespective, the appreciable loss of muscle mass in older individuals undergoing bed rest, reinforces the need to implement strategies during disuse events to protect muscle mass, function, and whole-body metabolic health. Whether short-term RET prehabilitation can protect postprandial muscle anabolism and offset muscle atrophy in older individuals during a very short bed-rest period (i.e., ≤3 days) remains to be seen, as do the potential benefits of RET prehabilitation on disuse-induced insulin resistance.
Acknowledgements

The authors would like to thank the research participants for their time and effort in completing the study. We extend our appreciation to the CRF nursing staff for their support throughout the trial.

Conflict of interest

None of the authors have any conflicts of interest to disclose.

Funding

This work was supported by an award from the Biotechnology and Biological Sciences Research Council to L.B. (BB/N018214/1) and the NIHR Clinical Research Facility in University Hospitals Birmingham NHS Foundation Trust, Birmingham. The authors of this manuscript certify that they comply with the ethical guidelines for authorship and publishing in the Journal of Cachexia, Sarcopenia and Muscle.

Data Availability Statement

The datasets generated during and/or analysed during the current study are not publicly available but are available from the corresponding author on reasonable request.

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