

# Daily myofibrillar protein synthesis rates in response to low- and high-frequency resistance exercise training in healthy, young men

Shad, Brandon J; Thompson, Janice L; Mckendry, James; Holwerda, Andrew M; Elhassan, Yasir S; Breen, Leigh; van Loon, Luc J C; Wallis, Gareth A

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1 **Daily myofibrillar protein synthesis rates in response to low and high frequency**  
2 **resistance exercise training in healthy, young men**

3

4 Brandon J. Shad<sup>1</sup>, Janice L. Thompson<sup>1</sup>, James Mckendry<sup>1</sup>, Andrew M. Holwerda<sup>2</sup>, Yasir S.  
5 Elhassan<sup>3,4</sup>, Leigh Breen<sup>1,5,6</sup>, Luc J.C. van Loon<sup>2</sup> and Gareth A. Wallis<sup>1\*</sup>

6

7 <sup>1</sup>*School of Sport, Exercise and Rehabilitation Sciences, University of Birmingham,*  
8 *Birmingham, United Kingdom*

9 <sup>2</sup>*Department of Human Biology, NUTRIM School of Nutrition and Translational Research in*  
10 *Metabolism, Maastricht University Medical Centre, Maastricht, The Netherlands*

11 <sup>3</sup>*Institute of Metabolism and Systems Research, University of Birmingham, Birmingham,*  
12 *United Kingdom*

13 <sup>4</sup>*Centre for Endocrinology, Diabetes and Metabolism, Birmingham Health Partners,*  
14 *Birmingham, United Kingdom.*

15 <sup>5</sup>*NIHR Birmingham Biomedical Research Centre, United Kingdom.*

16 <sup>6</sup>*MRC- Versus Arthritis Research UK Centre for Musculoskeletal Ageing Research,*  
17 *University of Birmingham, United Kingdom.*

18

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20 **Running title:** Resistance exercise frequency and muscle protein synthesis

21

22 **\*Corresponding author:**

23 Dr Gareth A. Wallis

24 School of Sport, Exercise and Rehabilitation Sciences

25 University of Birmingham

26 Edgbaston, B15 2TT, UK.

27 Phone: +44(0) 121 414 4129

28 Email: g.a.wallis@bham.ac.uk

29 **ABSTRACT**

30 The impact of resistance exercise frequency on muscle protein synthesis rates remains  
31 unknown. The aim of this study was to compare daily myofibrillar protein synthesis rates  
32 over a seven-day period of low frequency versus high frequency resistance exercise training.  
33 Nine young men ( $21\pm 2$  y) completed a seven-day period of habitual physical activity  
34 (BASAL). This was followed by a seven-day exercise period of volume-matched, low  
35 frequency (10 x 10 repetitions at 70% 1RM, once per week; LF) or high frequency (2 x 10  
36 repetitions at  $\sim 70\%$  1RM, five times per week; HF) resistance exercise training. Participants  
37 had one leg randomly allocated to LF and the other to HF. Skeletal muscle biopsies and daily  
38 saliva samples were collected to determine myofibrillar protein synthesis rates using  $^2\text{H}_2\text{O}$ ,  
39 with intracellular signalling determined using Western blotting. Myofibrillar protein synthesis  
40 rates did not differ between LF ( $1.46\pm 0.26\ \%\cdot\text{d}^{-1}$ ) and HF ( $1.48\pm 0.33\ \%\cdot\text{d}^{-1}$ ) conditions over  
41 the seven-day exercise training period ( $P>0.05$ ). There were no significant differences  
42 between LF and HF conditions over the first two days ( $1.45\pm 0.41$  vs  $1.25\pm 0.46\ \%\cdot\text{d}^{-1}$ ) or last  
43 five days ( $1.47\pm 0.30$  vs  $1.50\pm 0.41\ \%\cdot\text{d}^{-1}$ ) of the exercise training period ( $P>0.05$ ). Daily  
44 myofibrillar protein synthesis rates were not different from BASAL at any time point during  
45 LF or HF ( $P>0.05$ ). The phosphorylation status and total protein content of selected proteins  
46 implicated in skeletal muscle ribosomal biogenesis were not different between conditions  
47 ( $P>0.05$ ). Under the conditions of the present study, resistance exercise training frequency  
48 did not modulate daily myofibrillar protein synthesis rates in young men.

49 **Key words:** Exercise frequency, muscle protein synthesis, skeletal muscle, deuterated water

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51

## 52 INTRODUCTION

53 The muscle hypertrophic response to resistance exercise training can be modulated by  
54 manipulating variables such as absolute load, total exercise volume, proximity to failure and  
55 rest interval between exercise sets (Burd et al., 2010b; Mitchell et al., 2012; Schoenfeld et al.,  
56 2016). Less clear is the impact of resistance exercise training frequency (i.e., the number of  
57 times a muscle group is exercised over a given period of time) on muscle hypertrophy.

58 Understanding the relative importance of exercise training frequency is necessary to optimize  
59 the skeletal muscle adaptive response to prolonged resistance exercise training.

60 Whilst some studies have shown muscle hypertrophy to be enhanced by a higher (i.e., two or  
61 more times per week) resistance exercise training frequency (Schoenfeld et al., 2015; Zaroni  
62 et al., 2018), most have shown no differences (Schoenfeld et al., 2018). However, most  
63 studies to date have examined the impact of resistance exercise training frequencies in the  
64 range of one-to-three times per week. It is possible that higher resistance exercise training  
65 frequencies (e.g., five times per week) are required to enhance muscle protein synthesis rates  
66 and subsequent muscle hypertrophy. The evidence currently available is equivocal, with one  
67 study (Zaroni et al., 2018) showing greater muscle hypertrophy with a relatively high (five  
68 times per week) resistance exercise training frequency whereas another study (Gomes et al.,  
69 2018) reported no differences. As such, the impact of high versus low resistance exercise  
70 training frequency on muscle hypertrophy remains unclear.

71 Muscle hypertrophy following prolonged resistance exercise training is the product of  
72 sustained elevations in muscle protein synthesis that exceed muscle protein breakdown. It has  
73 recently been posited that relatively high resistance exercise training frequency is required to  
74 maximize muscle hypertrophy by regularly stimulating the acute myofibrillar protein  
75 synthetic response to a single bout of resistance exercise (Dankel et al., 2017). Following an

76 acute bout of resistance exercise, myofibrillar protein synthesis rates remain elevated for  
77 approximately twenty-four hours before returning to basal levels (Burd et al., 2011; Damas et  
78 al., 2016). Furthermore, a relatively low volume (~three sets) of resistance exercise appears  
79 to maximize post-exercise myofibrillar protein synthesis rates, at least in young men (Burd et  
80 al., 2010a; Kumar et al., 2012). On this basis, it has been speculated that more frequent, low-  
81 volume, resistance exercise could induce more frequent elevations in myofibrillar protein  
82 synthesis rates which in the long-term would lead to greater muscle hypertrophy (Dankel et  
83 al., 2017). Whilst plausible, this hypothesis has yet to be tested.

84 The purpose of the present study was to compare daily myofibrillar protein synthesis rates,  
85 measured using deuterated water ( $^2\text{H}_2\text{O}$ ) under free-living conditions, in young men over a  
86 seven-day period while performing low (once per week; LF) versus high (five times per  
87 week; HF) frequency resistance exercise training. As muscle protein synthesis rates are  
88 facilitated by transcriptional capacity (Figueiredo & McCarthy, 2019), we also aimed to  
89 assess whether resistance exercise training frequency impacts the phosphorylation status and  
90 total protein content of selected proteins implicated in ribosomal biogenesis.

## 91 **METHODS**

### 92 *Participants and ethical approval*

93 Nine young men participated in the present study between February 2018 and August 2018.  
94 Participant characteristics are presented in **Table 1**. Prior to providing written consent, each  
95 volunteer was informed of the experimental procedures and potential risks. Participants were  
96 screened prior to inclusion and deemed healthy based on their responses to a general health  
97 questionnaire. Inclusion criteria included being male, aged 18-35 years, a BMI between 18.5-  
98 29.99 kg/m<sup>2</sup>, being recreationally active and untrained (i.e., performing activities of daily  
99 living and recreation but no regular lower body resistance exercise in the last year), and being

100 willing and able to comply with all procedures. Exclusion criteria included having a lidocaine  
101 allergy, hypertension ( $\geq 140/90$  mmHg) or bleeding disorders, current participation in another  
102 study, being a current/recent smoker, vegetarian/vegan or a history of substance abuse and/or  
103 taking prescription or non-prescription medication or supplements that may influence normal  
104 metabolic responses. The study was approved by the National Research Ethics Service  
105 Committee West Midlands, Edgbaston, UK (Reference: 17/WM/0430) and conformed to  
106 standards for the use of human participants in research as outlined in the Declaration of  
107 Helsinki. The intervention was registered at [clinicaltrials.gov](http://clinicaltrials.gov) prior to data collection  
108 (Identifier: NCT03275779).

### 109 *Pretesting*

110 During the initial screening visit, participants underwent maximal strength testing and a  
111 familiarization session. First, participants completed a 5 minute warm-up of self-paced  
112 cycling. Maximal leg strength was then determined for each leg on a plate loaded  $45^\circ$  leg  
113 press. This process was then repeated on a weight-stacked leg extension machine.  
114 Participants first performed a submaximal warm-up set of eight-to-ten repetitions and had  
115 their lifting form critiqued and corrected when necessary. This was followed by sets at  
116 progressively increasing loads until only one valid repetition could be completed. The load  
117 for each set was chosen based on the participant's rating of perceived exertion following the  
118 previous set. A three-minute rest interval was provided between each set. Once completed,  
119 the corresponding load ( $\sim 70\%$  1RM) to be used during the subsequent familiarization session  
120 and resistance exercise sessions was calculated.

121 To familiarize participants with the exercise volume to be completed during the experimental  
122 trials, and to minimize muscle damage associated with an unfamiliar bout of resistance  
123 exercise (Damas et al., 2016; Nosaka et al., 2001), participants completed five sets of

124 bilateral leg press followed by five sets of bilateral leg extension at ~70% 1RM, with two  
125 minutes rest between each set. Total exercise volume completed during the familiarization  
126 (12121±2206 kg) was similar to that completed in total by both legs during the experimental  
127 resistance exercise sessions (11952±2700 kg). Pretesting and the first experimental trial (day  
128 0) were separated by  $\geq$  seven days.

### 129 *Study overview*

130 A study overview is presented in **Figure 1**. The study was designed to assess whether  
131 resistance exercise frequency impacts daily myofibrillar protein synthesis rates measured  
132 under free-living conditions. Participants arrived at ~08:00 in a fasted state on day 0 and had  
133 a muscle biopsy collected. All muscle biopsies were collected from the *vastus lateralis* using  
134 the Bergström needle with manual suction under local anaesthesia (1% lidocaine).  
135 Participants then completed a seven-day basal period (BASAL) where they were instructed to  
136 maintain habitual physical activity (i.e., activities of daily living and recreation without  
137 structured physical activity). Participants returned on day 7 and had a second muscle biopsy  
138 collected from the alternate leg. Following this, participants had each leg randomly allocated  
139 to one of low frequency (LF) or high frequency (HF) resistance exercise (see *Resistance*  
140 *exercise sessions* section below). A bout of LF and HF was completed on day 7.  
141 Approximately forty-eight hours later (day 9), participants returned and had one muscle  
142 biopsy collected from each leg. This was followed by the second bout of HF. Additional  
143 bouts of HF were completed on days 10, 11 and 12. Participants returned on day 14 (~48  
144 hours after the final HF bout) and had the final muscle biopsies collected from each leg,  
145 signifying the end of the study. A pedometer was worn throughout and weighed food diaries  
146 were completed to assess daily step count and dietary intake, respectively, across the study.

### 147 *<sup>2</sup>H<sub>2</sub>O dosing protocol*

148 The  $^2\text{H}_2\text{O}$  dosing protocol consisted of one dosing day and sixteen maintenance days (Shad et  
149 al., 2019). The  $^2\text{H}_2\text{O}$  protocol was well tolerated with none of the participants reporting any  
150 adverse effects.

#### 151 *Dietary intake and physical activity*

152 The evening prior to each experimental visit involving muscle biopsies, participants received  
153 the same standardized meal (~689 kcal, providing ~55 energy% (En%) carbohydrate, ~20  
154 En% protein, and ~25 En% fat). A weighed four-day food diary was completed over the first  
155 seven-day period of habitual physical activity (BASAL) and over the second seven-day  
156 period of LF and HF resistance exercise to assess energy and macronutrient intake.

157 Participants were required to include two week-days and both weekend days in their  
158 recordings. Dietary records were analysed using Dietplan software (Forestfield Software Ltd.,  
159 v6.70.67). Participants were instructed to refrain from structured physical activity throughout  
160 the study other than the prescribed resistance exercise completed as part of the study.

161 Participants were also provided with a hip-worn pedometer (Yamax Digi-Walker SW-200) to  
162 wear throughout the study to assess daily step count.

#### 163 *Resistance exercise sessions*

164 Using a within-subject design, participants had one leg randomized to complete LF and the  
165 other to HF. Prior to all resistance exercise sessions, participants completed a five-minute  
166 warm-up of self-paced cycling at ~100 W. On day 7, a single bout of unilateral high volume  
167 LF was completed. This consisted of five sets of ten repetitions at ~70% 1RM on the 45° leg  
168 press machine followed by five sets of ten repetitions at ~70% 1RM on the weight-stacked  
169 leg extension machine. A single bout of unilateral low volume HF was also completed on day  
170 7 using the opposite leg. This consisted of one set of ten repetitions at ~70% 1RM on the 45°  
171 leg press machine followed by one set of ten repetitions at ~70% 1RM on the weight-stacked



172 leg extension machine. A further four bouts of unilateral low volume HF was completed on  
173 days 9, 10, 11 and 12. This design ensured that total exercise volume and the number of sets  
174 completed were matched between the LF and HF conditions. Total exercise volume was  
175 intentionally matched as exercise volume has been shown, at least when comparing low  
176 volumes of resistance exercise, to modulate the magnitude of the myofibrillar protein  
177 synthetic response to resistance exercise (Burd et al., 2010a). Two minutes of rest was  
178 allowed between all sets, and five minutes of rest was allowed between the bouts of LF and  
179 HF on day 7. Following all resistance exercise sessions, participants ingested 25 g of whey  
180 protein powder (Impact Whey Protein; Myprotein), containing 21 g of protein (equating to  
181 ~0.29 g/kg), dissolved in water.

#### 182 *Body water <sup>2</sup>H enrichment*

183 Body water <sup>2</sup>H enrichment was analysed from daily saliva samples collected throughout the  
184 study as previously described (Holwerda et al., 2018; Shad et al., 2019).

#### 185 *Myofibrillar bound <sup>2</sup>H-alanine enrichment*

186 <sup>2</sup>H-alanine enrichment in the myofibrillar fraction of muscle biopsy samples was measured as  
187 previously described (Shad et al., 2019).

#### 188 *Western blotting*

189 Western blot analyses were performed on the sarcoplasmic fraction obtained during  
190 myofibrillar protein extraction as previously described (McKendry et al., 2019). The  
191 following primary antibodies were used ((1:1000) in 2.5% bovine serum albumin (BSA)):  
192 total eukaryotic translation initiation factor 4E (eIF4E) (ab33766), phospho-eIF4E Ser209  
193 (ab76256), total cyclin D1 (ab16663) and total upstream binding factor (UBF) (ab244287) all  
194 purchased from Abcam (Abcam, Cambridge, U.K). Imaging was undertaken using a G:Box

195 Chemi-XR5 (Syngene, Cambridge, UK) and bands were quantified using Image Studio Lite  
196 (Li-Cor, Lincoln, Nebraska, U.S).

### 197 *Calculations*

198 Myofibrillar protein fractional synthetic rate (FSR) was determined using the incorporation of  
199 <sup>2</sup>H-alanine into myofibrillar protein and the mean <sup>2</sup>H enrichment in body water between  
200 sequential biopsies, corrected by a factor of 3.7, as the surrogate precursor based upon <sup>2</sup>H  
201 labelling during *de novo* alanine synthesis (Belloto et al., 2007). The standard precursor-  
202 product method was used to calculate FSR:

$$203 \quad FSR (\% \cdot \text{day}^{-1}) = \left( \frac{E_{m2} - E_{m1}}{E_{\text{precursor}} \times t} \right) \times 100$$

204 where  $E_{m1}$  and  $E_{m2}$  are the myofibrillar protein-bound <sup>2</sup>H-alanine enrichments between  
205 sequential muscle biopsies.  $E_{\text{precursor}}$  represents the mean body water <sup>2</sup>H enrichment between  
206 sequential biopsies corrected by a factor of 3.7 based upon the <sup>2</sup>H labelling of alanine during  
207 *de novo* synthesis (Belloto et al., 2007).  $t$  represents the time between sequential biopsies in  
208 days.

### 209 *Statistics*

210 Based on the hypothesis that high frequency resistance exercise training would result in more  
211 frequent elevations in myofibrillar protein synthesis rates compared to low frequency  
212 resistance exercise training, and previous research (Holwerda et al., 2018; Wilkinson et al.,  
213 2014), an effect size of 1.1 was estimated. Sample size calculations showed that n=9 would  
214 be sufficient to detect a difference in daily myofibrillar protein synthesis rates between LF  
215 and HF conditions over the seven-day exercise training period using a two-tailed paired  
216 samples t test (80% power,  $\alpha$ -level of 0.05, G\*power). All statistical analyses were performed

217 using SPSS 25.0 (SPSS, USA). Differences between the seven-day basal period and seven-  
218 day exercise period (i.e., BASAL vs. LF/HF) for daily step count and dietary intake were  
219 compared using paired sample t-tests. Differences between exercise conditions (LF vs. HF)  
220 for exercise variables (i.e., maximal strength and total exercise volume) were compared using  
221 a paired sample t-test. Body water <sup>2</sup>H enrichment was analysed using a one-factor repeated  
222 measures ANOVA with time as the within-subjects factor. Myofibrillar protein FSR over the  
223 seven-day resistance exercise training period was compared between LF and HF conditions  
224 using a paired samples t-test (n=9). All other comparisons over time and between conditions  
225 for myofibrillar protein FSR were analysed using two-factor repeated measures ANOVAs  
226 (condition x time) with condition (BASAL vs. LF vs. HF) and time (days 0-7, 7-9, 9-14 and  
227 7-14) as within-subjects factors. Intracellular signalling was analysed using a two-factor  
228 repeated measures ANOVA (condition x time) with condition (BASAL vs. LF vs. HF) and  
229 time (days 7, 9 and 14) as within-subjects factors. A biopsy sample for one participant could  
230 not be collected on day 9, and thus myofibrillar protein FSR data for days 7-9 and 9-14 and  
231 all intracellular signalling data were analysed on n=8. When a significant main effect or  
232 interaction was found, t-tests with Bonferroni correction for multiple comparisons were  
233 performed. All data are presented as mean±SD.

234

## 235 RESULTS

### 236 *Exercise variables*

237 Maximal strength values at baseline were not different between the LF and HF conditions for  
238 the leg press ( $P=0.397$ ) and leg extension ( $P=0.650$ ) exercises (**Table 1**). By design, total  
239 exercise volume completed was not different between the LF ( $5933\pm 1357$  kg) and HF  
240 ( $6019\pm 1347$  kg) conditions ( $P=0.121$ ).

### 241 *Daily step count and dietary intake*

242 Daily step count and dietary intake are presented in **Table 2**. Daily step count was not  
243 different between BASAL and the seven-day period of resistance exercise ( $P=0.167$ ). The  
244 relative contribution of dietary fat to overall energy intake significantly decreased during the  
245 period of resistance exercise ( $P=0.041$ ). There was also a trend for daily protein intake  
246 ( $P=0.061$ ) and protein intake relative to body weight ( $P=0.089$ ) to increase during the period  
247 of resistance exercise. All other dietary variables were unchanged across the study.

### 248 *Body water $^2\text{H}$ enrichment*

249 **Figure 2A** presents the mean body water  $^2\text{H}$  enrichment. Following the loading phase on day  
250 -2 and a single maintenance day on day -1, body water  $^2\text{H}$  enrichment reached  $0.55\pm 0.05\%$   
251 (day 0). Body water  $^2\text{H}$  enrichment did not change significantly over the duration of the  
252 study, with an average body water  $^2\text{H}$  enrichment of  $0.58\pm 0.08\%$  during BASAL and  
253  $0.62\pm 0.13\%$  during the period of resistance exercise ( $P=0.107$ ).

### 254 *Myofibrillar protein synthesis*

255 Daily myofibrillar protein synthesis rates were not different between LF ( $1.46\pm 0.26\% \cdot \text{d}^{-1}$ )  
256 and HF ( $1.48\pm 0.33\% \cdot \text{d}^{-1}$ ) conditions over the entire seven-day exercise period (**Figure 2B**;

257 P=0.801). Moreover, there were no significant differences between LF and HF conditions  
258 over the first two days (days 7-9) ( $1.45\pm 0.41$  vs.  $1.25\pm 0.46$  %·d<sup>-1</sup>; **Figure 3**; P=0.342) or over  
259 the last five days (days 9-14) of the exercise period ( $1.47\pm 0.30$  vs.  $1.50\pm 0.41$  %·d<sup>-1</sup>; **Figure**  
260 **3**; P=0.342). Daily myofibrillar protein synthesis rates were not different from BASAL at any  
261 time point during LF or HF (**Figures 2B and 3**; P=0.591).

### 262 *Intracellular signalling*

263 A main effect of time was observed for eIF4E total protein content (**Figure 4A**; P=0.029).  
264 Following correction for multiple comparisons, pairwise comparisons showed a tendency  
265 (P=0.056) for greater total protein content 48 hours (i.e., day 9) following the initial LF and  
266 HF resistance exercise bouts compared to day 7. A main effect of time was also observed for  
267 cyclin D1 total protein content (**Figure 4C**; P=0.046). However, following correction for  
268 multiple comparisons, pairwise comparisons showed no significant difference between time  
269 points. There were no significant changes over time (P=0.407) or differences between LF and  
270 HF conditions (P=0.345) for phosphorylation of eIF4E at Ser209 (**Figure 4B**). There were  
271 no significant changes over time (P=0.217) or differences between LF and HF conditions  
272 (P=0.891) for UBF total protein content (**Figure 4D**).

273

## 274 **DISCUSSION**

275 The present study is the first to determine the impact that resistance exercise training  
276 frequency may have on myofibrillar protein synthesis rates. The major finding was that daily  
277 myofibrillar protein synthesis rates did not differ between volume-matched low and high  
278 frequency resistance exercise training performed over a seven-day period in young men. In  
279 line with these findings, resistance exercise training frequency did not modulate the  
280 phosphorylation status and total protein content of selected proteins implicated in skeletal  
281 muscle ribosomal biogenesis.

282 Manipulation of resistance exercise training frequency (i.e., the number of times a muscle  
283 group is exercised over a given period of time) has been proposed as a key factor determining  
284 exercise training induced muscle hypertrophy (Dankel et al., 2017; Schoenfeld et al., 2018).  
285 This is based on the premise that high resistance exercise training frequency induces greater  
286 overall myofibrillar protein synthesis rates and thus results in a greater amount of time spent  
287 in a greater net positive protein balance (Dankel et al., 2017). In the present study, a unilateral  
288 exercise model was utilized where each participant had one leg assigned to complete  
289 resistance exercise training once per week (i.e., low frequency; LF) and the other leg to  
290 complete resistance exercise training five times per week (i.e., high frequency; HF). This  
291 experimental design ensured that factors known to influence day-to-day muscle protein  
292 synthesis rates (e.g., sleep (Saner et al., 2020), protein intake (Witard et al., 2014), dietary  
293 composition (van Vliet et al., 2017) and habitual physical activity (Shad et al., 2019)) were  
294 identical between conditions, thereby allowing the impact of different resistance exercise  
295 training frequency on myofibrillar protein synthesis rates to be assessed in isolation. In  
296 contrast to the aforementioned hypothesis, the findings of the present study demonstrate that  
297 under volume-matched conditions, a high resistance exercise training frequency did not result  
298 in greater daily myofibrillar protein synthesis rates. These findings lend support to the

299 preponderance of evidence showing that resistance exercise training frequency has little  
300 impact on muscle hypertrophy (Barcelos et al., 2018; Schoenfeld et al., 2018).

301 The present data are in line with evidence showing no differences in muscle hypertrophy with  
302 a resistance exercise frequency of one versus five times per week (Gomes et al., 2018), but  
303 are inconsistent with findings showing greater muscle hypertrophy under similar conditions  
304 (Zaroni et al., 2018). It is important to note that the total exercise volume completed in the  
305 study by Zaroni et al. (2018) was significantly higher in the group with a resistance exercise  
306 training frequency of five times per week. In contrast, in the present study, total exercise  
307 volume was intentionally matched between the low and high frequency exercise training  
308 conditions, which likely explains the lack of agreement between findings. Indeed, a recent  
309 meta-analysis, published whilst the present study was being undertaken, suggests that  
310 resistance exercise training frequency does not significantly impact muscle hypertrophy when  
311 conducted under volume-matched conditions (Schoenfeld et al., 2018). Taken together, it  
312 would appear that resistance exercise training frequency *per se* (i.e., under volume matched  
313 conditions) does not impact daily myofibrillar protein synthesis rates or subsequent muscle  
314 hypertrophy in young individuals.

315 In contrast to most (Brook et al., 2016; Damas et al., 2016; Wilkinson et al., 2014), although  
316 not all (Davies et al., 2020) previous studies, resistance exercise training failed to induce a  
317 detectable increase in daily myofibrillar protein synthesis rates (**Figure 3**). The volume of  
318 resistance exercise completed in the high volume, low frequency exercise bout would have  
319 been expected to increase daily myofibrillar protein synthesis rates, given that resistance  
320 exercise of a similar volume and relative intensity has previously been shown to increase  
321 muscle protein synthesis rates in young men (Wilkinson et al., 2014). As such, there appears  
322 to be no obvious explanation for the absence of a measurable increase in daily myofibrillar  
323 protein synthesis rates following resistance exercise training. A possible explanation is that

324 the impact of resistance exercise training on myofibrillar protein synthesis was ‘diluted’ over  
325 the measurement period, as  $^2\text{H}_2\text{O}$  measures myofibrillar protein synthesis rates continuously  
326 capturing all free-living activities including diet, sleep and inactivity. Whilst more  
327 representative of long-term muscle hypertrophy and remodelling (Damas et al., 2016), the  
328 free-living nature of the  $^2\text{H}_2\text{O}$  measurement may have masked the well-established increase  
329 in myofibrillar protein synthesis in the hours following resistance exercise (Burd et al.,  
330 2010a; Kumar et al., 2012).

331 An alternative explanation could be related to familiarizing participants with resistance  
332 exercise prior to the study. During the screening visit, participants completed a high volume  
333 familiarization bout. Given that Damas and colleagues demonstrated that the 48-hour  
334 myofibrillar protein synthetic response following resistance exercise is no longer different  
335 from resting values once participants have been familiarized with resistance exercise, this  
336 may explain the undetectable increase in daily myofibrillar protein synthesis rates in the  
337 present study (Damas et al., 2016). A final possibility is that factors known to influence  
338 muscle protein synthesis rates (e.g., sleep (Saner et al., 2020) and energy balance (Areta et  
339 al., 2014)) could have differed during the basal period and the exercise period and thus could,  
340 in part, explain the lack of an exercise effect. It must be acknowledged that the inability to  
341 detect an increase in daily myofibrillar protein synthesis rates in response to resistance  
342 exercise training may also have precluded differences from being detected between low  
343 frequency and high frequency resistance exercise training.

344 As muscle protein synthesis is partly regulated by translational capacity (i.e., ribosomal  
345 biogenesis) (Figueiredo & McCarthy, 2019), a secondary aim was to assess whether  
346 resistance exercise training frequency impacts the phosphorylation status and total protein  
347 content of selected proteins implicated in skeletal muscle ribosomal biogenesis (**Figure 4**).  
348 Transcription of ribosomal DNA (rDNA) requires the activation of eIF4E and cyclin D1



349 which can subsequently activate a number of transcription factors including UBF which  
350 forms part of the pre-initiation complex (Figueiredo & McCarthy, 2019). In line with  
351 previous findings (Figueiredo et al., 2016), there was a tendency ( $P=0.056$ ) for total eIF4E  
352 protein content (**Figure 4A**) to increase 48 hours following the initial bouts of LF and HF  
353 resistance exercise training. Consistent with the finding that resistance exercise training  
354 frequency had no impact on daily myofibrillar protein synthesis rates, no differences were  
355 observed at any time point for any marker of skeletal ribosomal biogenesis between LF and  
356 HF resistance exercise training (**Figure 4**). However, it should be acknowledged that skeletal  
357 muscle ribosomal biogenesis is activated at multiple time points following resistance exercise  
358 (Figueiredo et al., 2016) and thus it is possible that biopsy timing, primarily intended to  
359 assess myofibrillar protein synthesis rates, missed differences that may have occurred at  
360 earlier time points.

361 Although total exercise volume was intentionally matched to isolate the impact of resistance  
362 exercise training frequency *per se* on daily myofibrillar protein synthesis rates, it should be  
363 considered that higher resistance exercise training frequencies can be used effectively to  
364 increase overall exercise volume for a given muscle group (Barcelos et al., 2018). Indeed,  
365 under non-volume equated conditions, higher resistance exercise training frequencies have  
366 been associated with greater gains in muscle mass (Schoenfeld et al., 2018) and strength  
367 (Grgic et al., 2018). From a practical standpoint, high resistance exercise training frequency  
368 may be considered a useful means of achieving a given exercise training volume, particularly  
369 when time is a limiting factor.

370 It is also important to note that any change in muscle mass is ultimately determined by the  
371 overall protein balance between muscle protein synthesis and breakdown. Whilst the absence  
372 of a measure of muscle protein breakdown may be considered a limitation of the present  
373 investigation, the myofibrillar protein synthesis measurements made in the present study

374 align well with the general finding that volume-matched resistance exercise training  
375 frequency has no impact on muscle hypertrophy (Schoenfeld et al., 2018). Finally, this study  
376 was conducted in individuals unaccustomed to regular lower limb resistance exercise, but it is  
377 possible that higher resistance exercise frequencies could be of greater benefit to more  
378 resistance-trained individuals as has been suggested previously (Dankel et al., 2017).

379 In conclusion, under the conditions of the present study, resistance exercise training  
380 frequency does not modulate daily myofibrillar protein synthesis rates or the phosphorylation  
381 status and total protein content of selected proteins implicated in skeletal muscle ribosomal  
382 biogenesis in young men. These findings suggest that for a given exercise volume, resistance  
383 exercise training frequency has little impact on skeletal muscle hypertrophy.

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390 **AUTHOR CONTRIBUTIONS**

391 B.J.S., J.L.T., A.M.H., L.J.C.v.L., and G.A.W. conception and design of research; B.J.S.,  
392 J.M., Y.S.E., L.B., and G.A.W. performed experiments; B.J.S., J.M., and A.M.H. analysed  
393 samples; B.J.S., and G.A.W. prepared figures and drafted manuscript; B.J.S., J.L.T., J.M.,  
394 A.M.H., Y.S.E., L.B., L.J.C.v.L., and G.A.W. edited and revised manuscript; B.J.S., J.L.T.,  
395 J.M., A.M.H., Y.S.E., L.B., L.J.C.v.L., and G.A.W. approved final version of manuscript.

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399 **CONFLICTS OF INTEREST**

400 None of the authors have any conflicts of interest or financial disclosures to declare.

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529 **Table 1.** Participants' characteristics at baseline

Variable	Value
Age (y)	21.0 ± 2.3
Height (m)	1.79 ± 0.07
Body mass (kg)	72.4 ± 7.1
BMI (kg·m <sup>-2</sup> )	22.7 ± 2.6
LF leg press 1RM (kg)	104 ± 22
HF leg press 1RM (kg)	106 ± 22
LF leg extension 1RM (kg)	82 ± 11
HF leg extension 1RM (kg)	81 ± 12

530 Values are mean±SD. n=9. BMI, body mass index; 1RM, one repetition maximum; LF, low

531 frequency; HF, high frequency.

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541 **Table 2.** Daily step count and dietary intake during the seven-day period of habitual physical  
 542 activity (BASAL) and seven-day period of low frequency (LF) and high frequency (HF)  
 543 resistance exercise

Variable	BASAL	LF/HF	<i>P</i> Value
Daily step count	10000 ± 2420	11458 ± 1871	0.167
Energy intake (kcal·d <sup>-1</sup> )	2253 ± 316	2336 ± 208	0.477
Protein (g·kg <sup>-1</sup> ·d <sup>-1</sup> )	1.3 ± 0.4	1.5 ± 0.2	0.089
Protein intake (g·d <sup>-1</sup> )	93 ± 25	104 ± 15	0.061
Carbohydrate intake (g·d <sup>-1</sup> )	278 ± 53	280 ± 43	0.931
Fat intake (g·d <sup>-1</sup> )	82 ± 12	82 ± 8	0.906
Protein (En%)	16 ± 5	18 ± 2	0.402
Carbohydrate (En%)	51 ± 7	52 ± 4	0.602
Fat (En%)	32 ± 3	30 ± 4*	0.041

544 Values are mean±SD. n=9. \*(*P*<0.05) indicates a significant difference between BASAL and  
 545 LF/HF conditions.

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553 **FIGURE LEGENDS**554 **Figure 1.** Study overview.

555 **Figure 2.** Body water  $^2\text{H}$  enrichment and daily myofibrillar protein fractional synthesis rates  
556 (FSR) during a seven-day period of habitual physical activity (BASAL) and a seven-day  
557 period of low frequency (LF) and high frequency (HF) resistance exercise (n=9). Data are  
558 displayed as mean $\pm$ SD with participants' individual FSR

559 **Figure 3.** Daily myofibrillar protein fractional synthesis rates (FSR) during a seven-day  
560 period of habitual physical activity (BASAL) and a seven-day period of low frequency (LF)  
561 and high frequency (HF) resistance exercise (n=8). Data are displayed as mean $\pm$ SD with  
562 participants' individual FSR

563 **Figure 4.** Impact of low frequency (LF) and high frequency (HF) resistance exercise on total  
564 protein content of eukaryotic translation initiation factor 4E (eIF4E; A), phosphorylation of  
565 eIF4E at Ser209 (B), total protein content of cyclin D1 (C) and total protein content of  
566 upstream binding factor (UBF; D) (n=8). Data are mean $\pm$ SD.

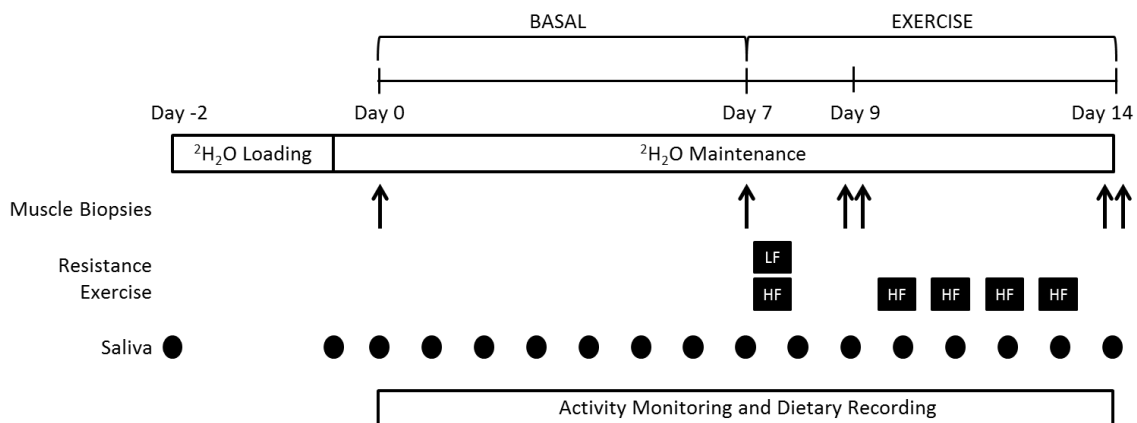
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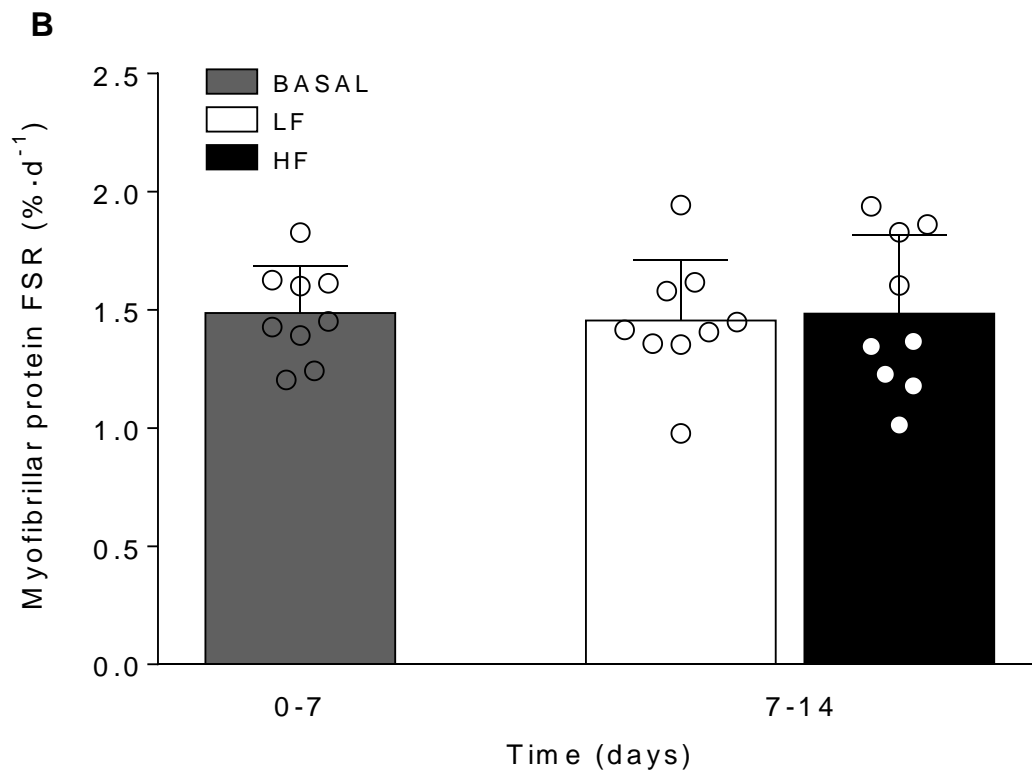
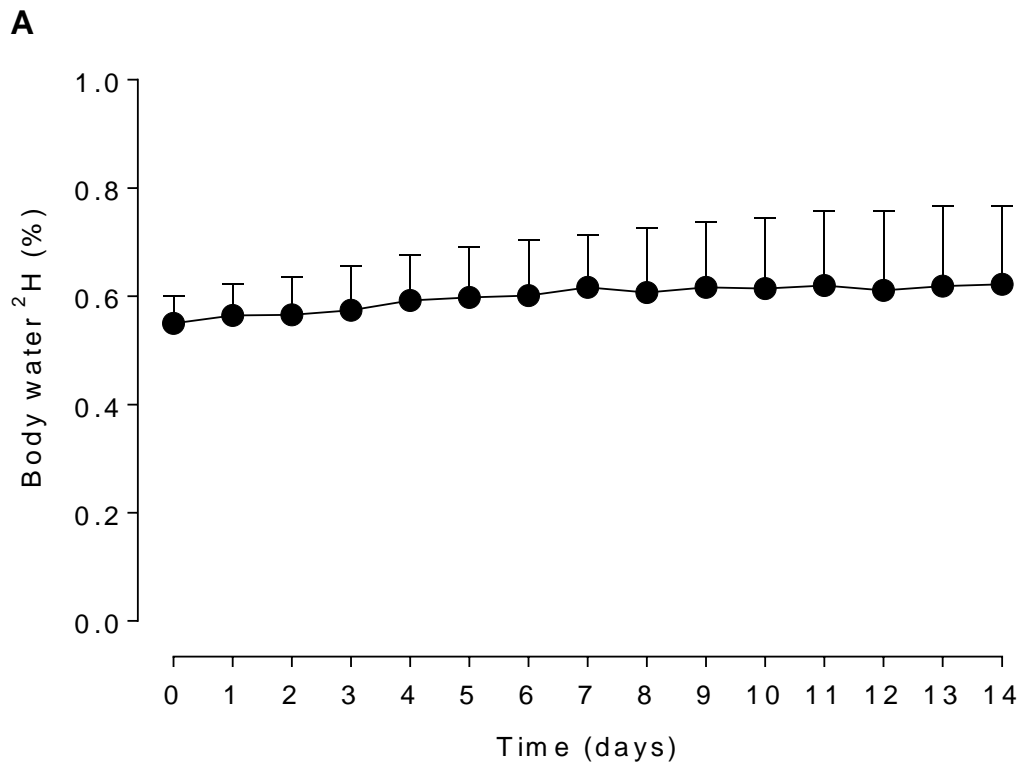
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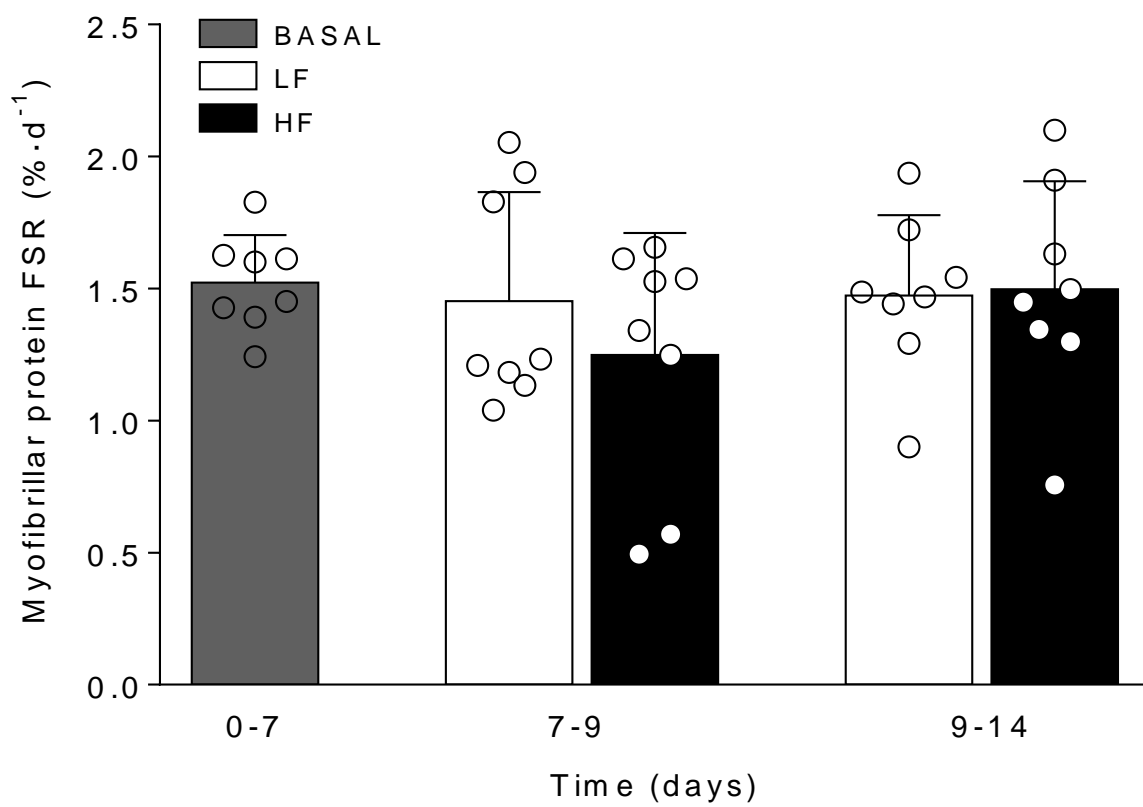
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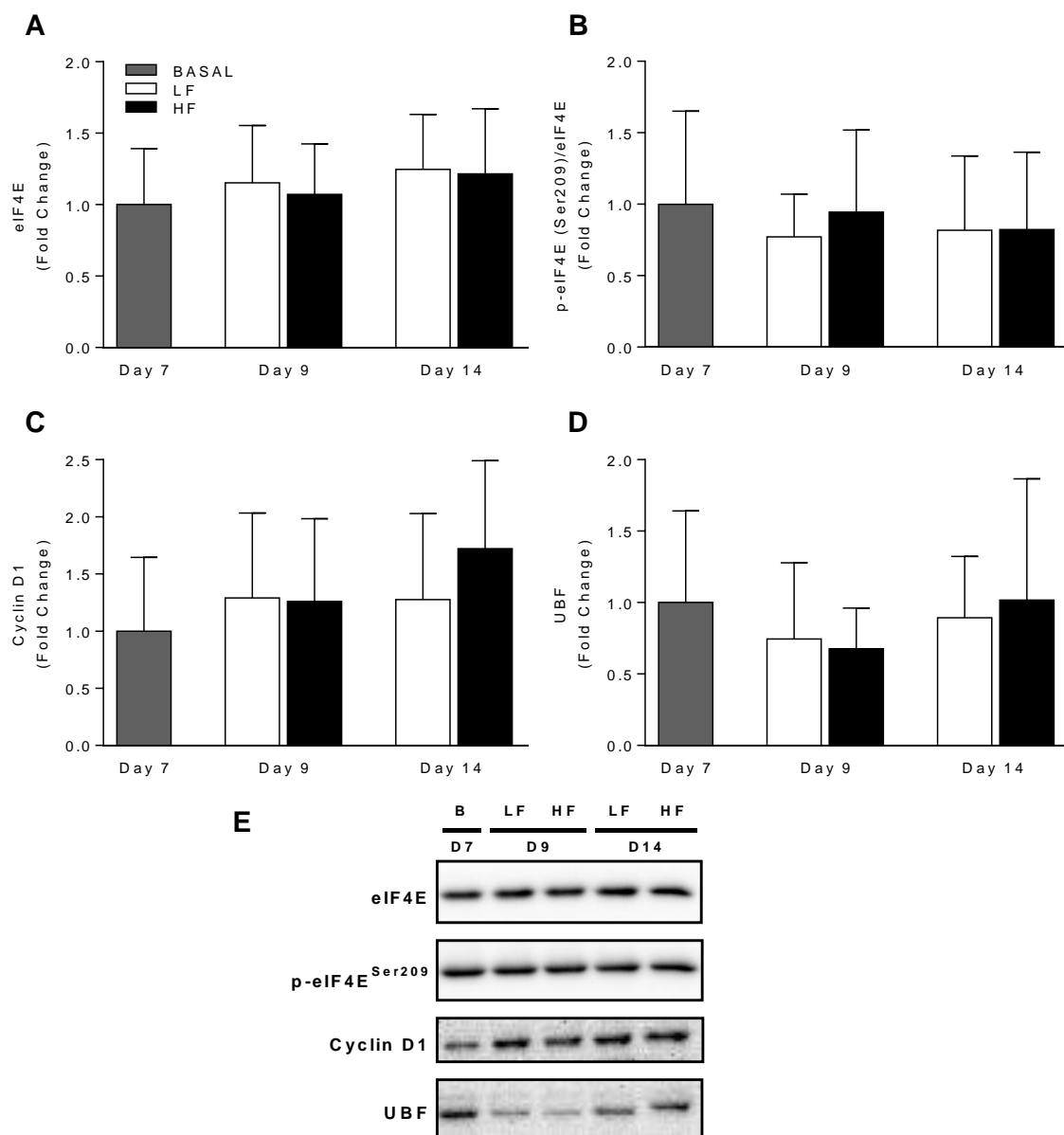
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