1	Nil Whey Protein Effect on Glycaemic Control after Intense Mixed-Mode Training in T2D
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#### Abstract 24

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While intense endurance and resistance exercise training and whey protein supplementation 25 have both been shown to independently improve glycaemic control, no known studies have 26 examined the effect of high-intensity mixed-mode interval training (MMIT) and whey 27 supplementation in adults with Type-2 diabetes (T2D). 28 Purpose: To determine if peri-training whey protein supplementation combined with MMIT 29 30 can improve glycaemic control. Methods: In a double-blind randomised controlled trial, 24 men (55.7±5.6 y) with T2D 31 32 performed MMIT with whey (20 grams) or placebo control for 10 weeks. Glycaemic control was assessed via glucose disposal rate (GDR) during a euglycaemic insulin clamp, fasting 33 blood glucose concentration (FBG), and HOMA-IR. Changes in peak oxygen consumption 34 35 (VO<sub>2peak</sub>), 1-repetition maximum strength (1RM), Vastus lateralis (VL) muscle and subcutaneous adipose thicknesses (SAT), and waist circumference (WC) were also assessed. 36 Results: 10-weeks of MMIT substantially improved GDR by 27.5% (90%CI 1.2%, 60.7%) 37 and 24.8% (-5.4%, 64.8%) in the whey and control groups, respectively. There were likely 38 and possible reductions in FBG by -17.4% (-30.6%, -1.6) and HOMA-IR by -14.1% (-25.3%, 39 1.08%) in the whey group, however, whey effects were not clearly beneficial to glycaemic 40 outcomes, relative to control. MMIT also clearly substantially improved 1RM by 20.6% 41 (16.3%, 24.9%) and 22.7% (18.4%, 27.2%), VO<sub>2peak</sub> by 22.6% (12.0%, 26.2%) and 18.5% 42 43 (10.5%, 27.4%), VL muscle thickness by 18.9% (12.0%, 26.2%) and 18.6% (10.5%, 27.4%) and possibly reduced WC by -2.1% (-3.1%, -1.0%) and -1.9% (-3.7%, -0.1%) in the control 44 and whey groups respectively, but the whey-control outcome was trivial or unclear.

46 Conclusion: A clinically-meaningful enhancement in glycaemic control following 1047 weeks of MMIT was not clearly advanced with peri-training whey protein supplementation in
48 middle-aged men with Type-2 diabetes.

Key Words: Milk-protein, exercise, diabetes, interval training, high-intensity, glucosedisposal.

#### 51 Introduction

A central pathology of type-2 diabetes (T2D) is impaired glycaemic control, a 52 53 condition characterised by a diminished capacity to restore postprandial blood glucose concentrations to homeostatic levels. Skeletal muscle is the major tissue of postprandial 54 glucose disposal (1), and a well-established site of dysfunction in T2D (2). It is well-55 documented that T2D skeletal muscle displays low expression of proteins contributing to 56 glucose uptake and metabolism, including: contractile, glucose transporter, and mitochondrial 57 proteins (3-5). Exercise has been shown to upregulate the expression of these proteins (6, 7), 58 and it is well-established that the improvements in aerobic capacity, lean mass and strength 59 60 that follow progressive aerobic or circuit resistance training are also associated with better glycaemic control (8, 9). High-intensity interval training has emerged as an effective low-61 volume and time-efficient exercise mode for rapidly improving glycaemic control. In middle-62 aged men with T2D, 2 weeks of high-intensity cycle interval training was shown to 63 significantly increase the expression of glucose transporter 4 (GLUT4) and mitochondrial 64 proteins in the vastus lateralis muscle and lower 24-hour blood glucose concentrations (6). 65 Milk protein supplementation has shown promise as a complementary therapeutic 66 agent to exercise for improving glycaemic control. Milk proteins are rich in amino acids that 67 stimulate protein synthesis in skeletal muscle (10), which may, like exercise training, lead to 68 This is a non-final version of an article published in final form in Gaffney, Kim A.; Lucero, Adam; Stoner, Lee; Faulkner, James; Whitfield, Patricia; Krebs, Jeremy; Rowlands, David S. (2017) "Nil Whey Protein Effect on Glycaemic Control after Intense Mixed-Mode Training in T2D." Medicine & Science in Sports & Exercise: Post Acceptance: 14 August 2017.

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69 better glycaemic control. Independently, whey supplementation was shown to improve glucose tolerance and FBG after 8 weeks in insulin resistant rats (11, 12) and HOMA-IR after 70 12 weeks in overweight and obese adults (13). As an adjunct therapy to exercise and 71 compared to carbohydrate consumption alone, milk-protein supplementation for 6 weeks was 72 reported to improve VO<sub>2max</sub> in treadmill trained sedentary men (14) and lean mass and 1RM 73 bench press strength after 8 weeks in mixed-mode trained female college basketball players 74 75 (15). As each of those outcomes has been previously associated with improved glycaemic control (9, 16, 17) combined treatments may also provide better therapeutic outcomes than 76 77 exercise alone in populations with T2D.

78 The aim of this study was to determine whether whey supplementation for 10 weeks

vould improve glycaemic control in a population with T2D performing high-intensity mixed-

80 mode interval training. We hypothesised that whey supplementation would enhance

81 glycaemic control to a greater extent than exercise alone. If effective, this may provide a

82 practical adjunct therapy to exercise for improving T2D rehabilitation outcomes.

83

#### 84 Methods

85 Participants

86 Men with T2D (n=24) were recruited from local medical centres in Wellington, NZ.

87 Inclusion characteristics were age 40-65 y, BMI<40, not requiring insulin therapy, and not

meeting the ACSM guidelines for exercise for T2D (18). Ethics was approved by the

89 Northern B Health and Disability Ethics Committee, Ministry of Health, Wellington NZ

90 (13/NTB/69). Participants provided written informed consent.

91

#### 92 *Experimental Design*

93 The design was a double blind, randomized (Research Randomizer, Version 4.0, http://www.randomizer.org), placebo controlled trial (http://www.anzctr.org.au/, Registration 94 number ACTRN12613000340730). At early stages of data collection, the original intended 95 third group: whey without MMIT, was removed from the study design because recruited 96 eligible participants declined to participate if not randomised to an exercise group creating 97 sampling bias. In the two-group design, participants consumed a whey-protein beverage or 98 99 carbohydrate placebo before and after 45 early-morning MMIT sessions over 10 weeks. Participants were encouraged to maintain dietary and medication habits throughout the 100 101 experimental period and not to participate in strenuous activity within 2 days of testing sessions. Participants were familiarised with all testing procedures except the euglycaemic 102 insulin clamp prior to baseline testing. Cardiac screening via ECG was performed at 103 104 familiarisation during a VO<sub>2peak</sub> cycling test. Baseline testing occurred 5-10 days prior to commencement of the intervention with post-testing 2 days after 45 exercise sessions. The 105 post glucose clamp was performed 48 hours after maximal cycling and strength tests to 106 provide a washout period that would allow for the bulk of the acute effects of intense exercise 107 on glycaemia to return to pre-exercise levels without inducing a period of deconditioning (19, 108 20). 109

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# 111 Exercise Protocol

Participants completed 27 cycling and 18 resistance training sessions (4-5 sessions each
week). Sessions included a 5-minute warm-up at low intensity on a cycle ergometer or
rowing machine followed by 20 minutes of 1-minute interval style cycling or resistance
exercise. Pre-programmed cycling sessions (VeloTron Racer Mate, Seattle, WA) included 10

intervals at 70%-90% (increased 5% every 2 weeks) of the participants' peak oxygen 116 consumption volume (VO<sub>2peak</sub>) obtained from baseline and fortnightly cycle testing 117 (SensorMedics Vmax, YorbaLinda, CA), with 1-minute active recovery intervals at 40% of 118 peak workload. Resistance training included 5 sets of 30 repetitions of each exercise (Day 1: 119 bench press and seated rows), and (Day 2: lateral pulldowns and barbell upright rows) with 1 120 minute of crunches on a fitball (Hart Sport, Auckland, NZ) as active recovery. Intensity was 121 122 set at 20% of 1-repetition maximum (1RM) during weeks 1-2 and increased to 25% of 1RM to elicit a high-intensity workload, for the remainder of the intervention based upon baseline 123 124 and fortnightly testing. If participants were unable to maintain a set cycling or strength workload for a full minute the subsequent interval was reduced by 10%. All exercise and 125 testing sessions were supervised by the researchers. 126

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#### 128 Supplement

129 Participants appeared each morning to the exercise laboratory in a fasted state. A chocolate

130 flavoured whey protein isolate (WPI-895, Fonterra, Auckland, New Zealand) beverage (20

131 grams protein/10 grams carbohydrate/3 grams milk-fat) or an identically-flavoured but non-

132 protein formulated isocaloric beverage (30 grams carbohydrate/3 grams milk-fat) was

133 consumed immediately before and after each exercise session. Each drink contained 175

134 calories (731 kilojoules). To reduce hunger and provide opportunity for a clear peri-training

- 135 whey compared to carbohydrate consumption effect to be observed, each participant
- 136 consumed a low-protein snack bar (Nature Valley, General Mills, Auckland, NZ) 1 hour after

137 exercise and resumed normal eating habits after 2 hours.

138

#### 139 *Glycaemic Measures*

140 Glucose disposal rate (GDR) for each individual was determined via a modified euglycaemic insulin clamp as described previously (21). Briefly, participants appeared for testing between 141 7 and 9 am after an overnight fast and at least 48 hours after the last exercise testing session. 142 A catheter was placed at the antecubital vein for insulin and glucose infusion, and dorsally at 143 the hand for blood draws. Arterialised blood was obtained by placing the hand in a heater box 144 at 50 °C. Participants received priming insulin doses of 160 mU·m<sup>2</sup>·min<sup>-1</sup> for 4 minutes and 145 80 mU·m<sup>2</sup>·min<sup>-1</sup> for 3 minutes, after which the dosage was reduced to 40 mU·m<sup>2</sup>·min<sup>-1</sup> for 146 the remainder of the clamp. A 25% glucose infusion was initiated at 15 minutes or sooner if 147 fasting blood glucose levels were below 6.5 mmol·L<sup>-1</sup> and adjusted after 5-minute blood 148 glucose readings until stabilised at 5 mmol $\cdot$ L<sup>-1</sup>. As this method elevated blood insulin 149 concentrations within a physiological rather than a supraphysiological range, the time to 150 151 stabilisation was variable between participants. GDR was calculated from the average rate of glucose infused (mg·kg<sup>-1</sup>·min<sup>-1</sup>) during a 60 minute stabilisation phase. Fasting blood 152 samples were obtained to determine FBG and HOMA-IR. 153

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# 155 *Physical Exercise Capacity*

Participants completed a continuous ramp protocol to volitional exhaustion on a cycle ergometer commencing at 40 Watts for 3 minutes and increasing 1 Watt every 4 seconds. Participants were encouraged to maintain a cadence of 70 rpm during the ramp. Peak oxygen consumption (mL·kg<sup>-1·</sup>min<sup>-1</sup>) was measured as the average of the highest 30-second consumption rate during the test. Acceptance of a maximal effort was dependent upon the participant achieving a maximal Borg Scale (1-20) rating and/or an RER>1.15. Estimated 1 repetition-maximum (1RM) tests were completed at baseline and every 2 weeks for smith

machine bench press, lateral pulldown, seated row and barbell upright row during a maximal-effort of 3-6 repetitions and predicted via the Brzycki Formula (22).

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166 *Body Composition* 

Body composition measures were taken in a fasted state prior to exercise testing. VL 167 thickness and subcutaneous adipose tissue (SAT) were measured after lying supine for 15 168 169 minutes via B-mode ultrasound (Terason T32000, Teratech Corp., Burlington, MA) using previously validated protocols (23, 24) modified to include measurement of SAT at the 170 171 biceps and cross-section diameter at the VL muscle. Measurements were taken in a supine position after participants had been lying relaxed for 15 minutes and then analysed using 172 ImageJ software (National Institute of Health, Bethesda, Maryland). SAT was determined 173 174 from the sum of adipose thickness at 4 standard calliper sites: thigh, calf, biceps, and triceps and VL thickness from the maximal cross-sectional diameter measured at 1/3 the distance 175 from the centre of the patella to the tubercle of the anterior superior iliac spine. 176

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178 Statistical Methods

Sample size estimation was based upon the primary outcome GDR using the test-179 retest values reported by Defronzo et al (25) in a healthy adult population and upon sample 180 size estimations for magnitude based clinical inference (26, 27). The typical error of 181 182 measurement was doubled to allow for uncertainty in variability in a T2D population and n was increased by 10% to allow for potential dropouts, which brought the required 183 sample to 24. The threshold for smallest worthwhile clinical change in GDR was 5.4% based 184 185 upon the effect of 3 months of hypoglycaemic therapy (Metformin) on naïve Type-2 diabetics (28).186

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The effect of treatment and time on all dependent variables was estimated from mixed 187 models (Proc Mixed, SAS Version 9.1; SAS Institute, Cary, NC). Data were log transformed 188 prior to analysis. Total 1RM strength was expressed as the back log-transformed average of 4 189 log-transformed lift scores. Uncertainty was presented as 90% confidence limits or P value. 190 Magnitude-based inference was employed to infer clinical and mechanistic outcome effects 191 (27, 29). The probability that a contrast was at least greater than the clinical threshold or 192 smallest Cohen's d standardized difference ( $0.2 \times$  baseline SD) was: 25-75% possible, 75-193 95% likely, 95-99.5% very likely, >99.5% almost certain (27). In the case where the majority 194 (>50%) of the CI lay between the thresholds for positive and negative substantiveness, the 195 effect was qualified trivial (negligible) with the respective probabilities as above (30). The 196 terms *benefit*, *trivial* (*negligible*), and *harm* refer to the most likely directional outcome, 197 relative to the smallest effect threshold. The terms *unclear*, *inconclusive* refers to outcomes 198 199 where the likelihood of both benefit and harm exceeded 5%. The likelihood of a clinical benefit of intervention was expressed as the benefit:harm odds ratio, with 66:1 the smallest 200 adoption threshold (27). Pre- and post-intervention scores are presented in figures as raw 201 202 means and standard deviations.

203

# 204 **Results**

Twenty-four men with T2D were recruited to the study (Figure 1). There were no clear differences between group characteristics at baseline (Table 1). All participants completed the 45 exercise sessions within the 10-week period. The glycaemic control outcomes and statistical summary for all parameter measures are in Figure 2 and Table 2, respectively. Ten weeks of MMIT produced a clinically meaningful enhancement in GDR in

210 the whey and control groups, respectively, relative to the smallest threshold change (5.4%); however, the whey-control difference was unclear. The secondary outcome measures of 211 212 glycaemic control (FBG and HOMA-IR) showed a likely and possible benefit of whey supplementation on FBG, and possible and unclear benefits on HOMA-IR in the whey and 213 control groups respectively, reaching the adoption threshold (OR>66:1) only in the Whey 214 group; however, there was also no clear difference in the Whey-Control contrast. Very likely 215 216 and almost certain improvements in VO<sub>2peak</sub>, 1RM strength and VL muscle thickness in response to 10-weeks of MMIT in both the Whey and Control groups were observed (Figure 217 218 3), but the whey-control differences were also negligible and unclear. There was a possible decrease in WC in both groups and a possible decrease in SAT in the whey group only (Table 219 220 2).

221

#### 222 Discussion

The current study showed that consumption of 20 grams of whey protein before and 223 after MMIT for 10 weeks did not enhance glycaemic control in a T2D population assessed 224 via measures of glucose disposal rate, fasting blood glucose, and HOMA-IR. Similarly, whey 225 supplementation did not enhance any of the exercise performance adaptations accruing in 226 227 response to MMIT, including VO<sub>2peak</sub>, 1RM strength, and muscle thickness. While previous evidence indicates that whey supplementation and high-intensity interval training 228 independently improve glycaemic control (6, 13), no clear benefit of combined therapies was 229 230 observed. Previously, consumption of 10 grams of whey protein hydrolysate before and after 231

resistance training for 10 weeks was shown to significantly increase quadriceps cross-

sectional area in healthy trained men (31). In addition, consumption of a single-dose of a 233 mixed milk-protein (20 grams) carbohydrate beverage after treadmill training for 6 weeks 234 235 significantly increased VO<sub>2max</sub> in sedentary middle-aged men compared to an isocaloric carbohydrate control (14). As both the increase in mid-thigh muscle cross-sectional area and 236 VO<sub>2peak</sub> following exercise intervention have been previously associated with improved 237 HbA1c in populations with Type-2 diabetics (9, 17), we predicted that peri-training whey 238 239 supplementation for 10 weeks would lead to better glycaemic control than the MMIT alone. Our observation that whey supplementation did not clearly increase muscle thickness at the 240 241 VL or VO<sub>2peak</sub> suggests that adaptive responses previously seen in exercising healthy populations may be lost with the development of T2D and may explain why we saw no effect 242 of whey protein on GDR, FBG or HOMA-IR. 243

244 It is possible that adults with T2D require a larger dose of milk-protein to induce clinically meaningful outcomes. 20 grams of protein has been reported to be the optimal 245 dosage for improving protein synthetic responses in the skeletal muscle of healthy young men 246 (32). While we also provided a total of 40 g of protein as 20 g before and 20 g after MITT, in 247 another study in healthy elderly individuals (71±4 y), 40 grams of whey protein increased 248 muscle protein synthesis after resistance training compared to a 20 gram dose (33). The 249 cohort in the current study was middle-aged (55.6±5.7 y), however, T2D skeletal muscle has 250 been shown to display characteristics of aged tissue, including: accelerated muscle wasting 251 252 (34); lower contractile strength to muscle volume (35), and decreased mitochondrial density (36). Future investigations should test dosage effects on muscle protein synthetic responses in 253 a population with T2D. 254

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While we saw no clear benefits of whey supplementation on glycaemic control in this study, there was some evidence that the protein exposure produced a more pronounced effect 256 This is a non-final version of an article published in final form in Gaffney, Kim A.; Lucero, Adam; Stoner, Lee; Faulkner, James; Whitfield, Patricia; Krebs, Jeremy; Rowlands, David S. (2017) "Nil Whey Protein Effect on Glycaemic Control after Intense Mixed-Mode Training in T2D." Medicine & Science in Sports & Exercise: Post Acceptance: 14 August 2017. https://doi.org/10.1249/MSS.000000000001404

on each of the glycaemic measures compared to exercise alone, as suggested through the 257 observation of a substantially larger clinically-beneficial odds ratios for GDR, FBG, and 258 HOMA-IR in the whey compared to the control group. We also observed that the adoption 259 threshold (odds ratio >66) was reached for FBG, HOMA-IR and SAT only in the whey 260 group, suggesting that the magnitude of the improvements in those secondary outcomes was 261 sufficient to justify treatment use only when therapies were combined. It is important to 262 263 acknowledge, however, that the full placebo-control adjusted outcome (whey-control), which takes into account the on-study effect, left a statistically unclear whey-protein effect. We 264 265 suggest that a longer intervention or a larger cohort (to increase study power) may have clarified whether whey supplementation was enhancing the pattern for improvement in 266 clinical outcomes. 267

268 An inherent potential confounder of investigations with control of energy intake was that the control group was consuming substantially more carbohydrate each training morning 269 than the whey group (60 compared to 20 grams). We reasoned that while there was potential 270 for the control group to be consuming more carbohydrate than their normal dietary intake, 271 which could be deleterious to glycaemic control, we expected that the metabolic demands of 272 20 minutes of MMIT would obviate any effect on post-exercise blood glucose concentration 273 in a previously sedentary population. In addition, 6 x 20 minute sessions of high-intensity 274 interval cycling was previously shown to significantly improve postprandial and 24-hour 275 276 blood glucose regulation in middle-aged adults with T2D (6). Our findings confirm that chronic intense interval training is effective for improving glycaemic control in populations 277 with T2D. We also found that the 5-days per week, mixed-mode training regime was well-278 279 adhered to by a previously sedentary, middle-aged T2D population, improved glucose disposal rates by a 4-5-fold greater magnitude than an equivalent duration of 280 This is a non-final version of an article published in final form in Gaffney, Kim A.; Lucero, Adam; Stoner, Lee; Faulkner, James; Whitfield, Patricia; Krebs, Jeremy; Rowlands, David S. (2017) "Nil Whey Protein Effect on Glycaemic Control after Intense Mixed-Mode Training in T2D." Medicine & Science in Sports & Exercise: Post Acceptance: 14 August 2017. https://doi.org/10.1249/MSS.000000000001404

pharmacotherapy (Metformin) alone (28), and negated the potentially deleterious impact of 281 consuming  $2 \times 30$  grams of a carbohydrate beverage each morning. Therefore, the MMIT 282 mode of exercise training may prove to be highly effective for improving T2D health 283 outcomes in long-term rehabilitation programs where high intensity exercise is appropriate. 284 285 In conclusion, consumption of 20 grams of whey protein before and after highintensity mixed-mode interval training for 10 weeks, compared to isocaloric non-protein 286 control, did not clearly enhance glycaemic control, VO<sub>2peak</sub>, 1RM strength, or VL muscle 287 cross-section diameter in middle-aged men with T2D. These findings suggest that over short-288 term interventions, populations with T2D may be resistant to nutritional stimulation of this 289 nature. However, recent dose response data, and patterns for greater gains in some clinical 290 parameters in the whey group support further investigation of the nutritional intervention, 291 possibly increasing the supplement dose or the intervention period. 292

293

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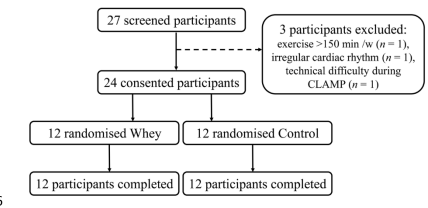
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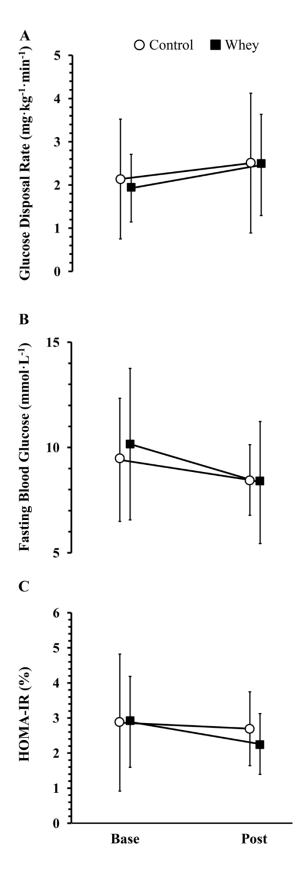
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406

407 **Figure 1.** Recruitment flowchart.



408

- 409 Figure 2. Effect of 10 weeks of peri-training whey supplementation on: A) glucose disposal
- 410 rate; B) fasting blood glucose concentration; and, C) HOMA-IR. Data are raw means and SD
- 411 for the Pre (baseline) and Post testing time points.

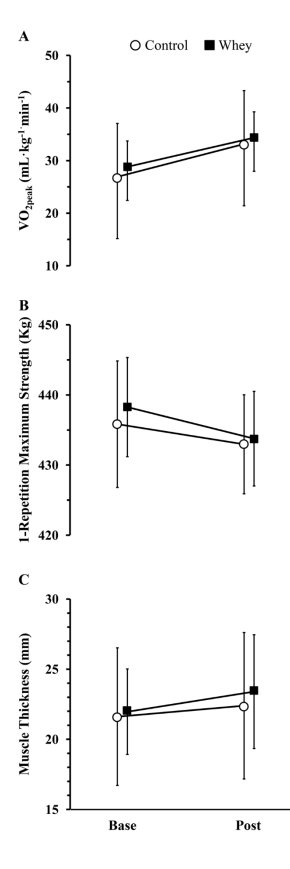




Figure 3. Effect of 10 weeks of peri-training whey supplementation on: A) VO<sub>2peak</sub>; B) 1RM
strength (the back log-transformed average of 4 log-transformed lift scores); and, C) *vastus lateralis* muscle thickness. Data are raw means and SD for the Pre (baseline) and Post testing
time points.

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	<b>Control</b> <i>n</i> =12	<b>Whey</b> <i>n</i> =12
Parameter	Mean SD	Mean SD
Age (y)	$57.8\pm5.2$	$53.5\pm5.6$
Height (cm)	$174.6\pm7.1$	$177.1\pm8.7$
Weight (kg)	$91.9 \pm 15.5$	$92.8 \pm 11.0$
BMI (kg⋅m <sup>2</sup> )	$30.1\pm4.9$	$29.6\pm2.7$
VO <sub>2peak</sub> (mL·kg <sup>-1</sup> ·min <sup>-1</sup> )	$26.9\pm10.2$	$28.7\pm4.9$
FBG (mmol·L <sup>-1</sup> )	$9.4\pm2.9$	$10.2\pm3.6$
GDR (mg·kg <sup>-1</sup> ·L <sup>-1</sup> )	$2.11 \pm 1.4$	$1.93\pm0.8$
Time to euglycaemia (min)	$106.3\pm67.2$	$106.7\pm53.9$

**Table 1.** Baseline characteristics of theControl and Whey groups.

Data are presented as means and standard deviations.

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**Table 2.** The effect of 10-weeks peri-training whey-protein supplementation on established clinical measures of glycaemic control, exercise performance, and body composition.

Contrast <sup>a</sup>	% Change	Upper CI	Lower CI	Likelihood (%) benefit/trivial/harm <sup>b</sup>	Qualitative <sup>b</sup>	Benefit odds <sup>b</sup>		
Glucose Disposal Rate								
Control	24.8	64.8	-5.4	90.1/7.1/2.8	Benefit likely	318		
Whey	27.5	60.7	1.2	95.6/3.5/0.9	Benefit very likely	2424		
Whey-Control	2.2	44.8	-28.0	42.6/24.4/33.0	Unclear	2		
Fasting Blood Glucose								
Control	-8.1	10.7	-23.7	50.4/45.8/3.8	Benefit possible	26		
Whey	-17.4	-1.6	-30.6	88.8/11.0/0.2	Benefit likely	3291		
Whey-Control	-10	15.3	-29.8	57.3/35.9/6.8	Unclear	19		
HOMA-IR								
Control	-5.3	28.3	-30.1	23.7/68.8/7.6	Unclear	4		
Whey	-14.1	1.08	-25.3	42.0/58/0.0	Benefit possible	3331		

When Control	0.2	25.4	24.0	25 0/50 1/6 0	I in alarm	0			
Whey-Control	9.2	25.4	-34.2	35.0/59.1/6.0	Unclear	8			
VO <sub>2peak</sub>									
Control	22.6	26.2	12.0	99.8/0.2/0.0	Benefit almost certain	5.05E+07			
Whey	18.5	27.4	10.5	99.1/0.9/0.0	Benefit very likely	2.81E+06			
Whey-Control	-3.3	9.07	-8.75	4.4/69.1/26.5	Trivial possible	0			
1-Repetition Maximum Strength <sup>c</sup>									
Control	20.6	24.9	16.3	100/0.0/0.0	Benefit almost certain	3.29E+31			
Whey	22.7	27.2	18.4	100/0.0/0.0	Benefit almost certain	7.80E+35			
Whey-Control	1.8	7.1	-3.2	0.1/99.8/0.0	Trivial almost certain	11			
Muscle Thickness									
Control	18.9	26.2	12.0	100/0.0/0.0	Benefit almost certain	1.78E+09			
Whey	18.6	27.4	10.5	99.89/0.02/0.0	Benefit almost certain	6.62E+07			
Whey-Control	-0.2	9.1	-8.8	13.6/70.6/15.9	Unclear	1			
Waist Circumference									
Control	-2.1	-1.0	-3.1	41.0/59.1/0.0	Benefit possible	7.44E+05			
Whey	-1.9	-0.1	-3.7	28.6/71.4/0.0	Benefit possible	2888			
Whey-Control	0.1	2.1	-1.8	0.3/99.5/0.2	Trivial very likely	2			
Subcutaneous Adipose Tissue <sup>d</sup>									
Control	-1	6.9	-8.3	6.7/90.8/2.5	Trivial likely	3			
Whey	-6.9	3.5	-16.2	43.7/55.8/0.5	Benefit possible	151			
Whey-Control	-6.0	6.7	-17.1	40.1/57.9/2.0	Benefit possible	32			

<sup>a</sup> Data for each contrast are post-pre. <sup>b</sup> The threshold for smallest clinical effect for glucose disposal rate was 5.4% (28); and for all other measures the smallest standardised difference (0.2xSD). The likelihood that a contrast was at least greater than the clinical threshold was: 25-75% possible, 75-95% likely, 95-99.5% very likely, >99.5% almost certain. Unclear refers to outcomes where the likelihood of both benefit and harm exceeded 5%. The clinical adoption threshold was expressed as a benefit: harm odds ratio >66:1. <sup>c</sup> Total 1-Repetition Maximum strength was expressed as the back log-transformed average of 4 log-transformed lift scores. <sup>d</sup> Subcutaneous Adipose Tissue was expressed as the sum of 4 sites.

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