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Effect of exercise on cardiometabolic risk factors in adults with chronic spinal cord injury

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Exercise and CMS risk in SCI

1 The effect of exercise on cardiometabolic risk factors in adults with chronic spinal cord 2 injury: A systematic review 3 Mr Matthew Farrow, MSci¹, Dr Thomas E Nightingale, PhD^{2,3}, Dr Jennifer Maher, PhD¹, Dr 4 Carly D McKay, PhD¹, Professor Dylan Thompson, PhD¹, Professor James Bilzon, PhD¹ 5 6 7 ¹Department for Health, University of Bath 8 ²International Collaboration on Repair Discoveries (ICORD), University of British Columbia 9 ³Faculty of Medicine, Division of Physical Medicine and Rehabilitation, University of British 10 Columbia 11 Conflict of Interest The authors declare no conflicts of interest 12 13 Funding This research did not receive any specific grant from funding agencies in the public, 14 commercial, or not-for-profit sectors **Corresponding author:** 15 16 Professor James Bilzon, Department for Health, University of Bath, BA2 7AY, UK 17 Email: J.Bilzon@bath.ac.uk 18 Tel: +44 (0)1225 383174 19 20 **Trial registration number** CRD4201815110

- 1 The effect of exercise on cardiometabolic risk factors in adults with chronic spinal cord
- 2 injury: A systematic review

ABSTRACT

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- 5 **Objective** To determine the effects of exercise on individual cardiometabolic syndrome (CMS)
- 6 risk factors in adults with chronic spinal cord injury (SCI).
- 7 **Design** Systematic review.
- 8 **Data sources** English language searches of PubMed, Web of Science, EMBASE, and Scopus
- 9 (01/01/1970 to 31/07/2019).
- 10 Eligibility criteria for selecting studies (1) original articles with statistical analysis, (2)
- participants were adults with a SCI sustained \geq 1-year ago, (3) exercise intervention duration
- ≥ 2 weeks, and (4) included any CMS risk factor as an outcome. The methodological quality
- of articles was assessed using the Downs and Black score.
- 14 **Results** Sixty-five studies were included for the final analysis, including nine studies classified
- as high quality (\geq 66%), 35 studies classified as fair quality (50-66%), and 21 studies classified
- as low quality (<50%). Improvements in waist circumference (4/6 studies) and markers of
- hepatic insulin sensitivity (4/5 studies) were reported following upper-body aerobic exercise
- training, but no improvements in fasting glucose (8/8 studies), lipid profile (6/8 studies),
- 19 systolic (8/9 studies) or diastolic blood pressure (9/9 studies) were observed. Improvements in
- 20 markers of peripheral insulin sensitivity (5/6 studies) were observed following functional
- 21 electrical stimulation (FES)-cycling. Improvements in lipid profile (4/5 studies) were observed
- following upper-body resistance training (RT) (with or without aerobic exercise). No consistent
- 23 improvements in CMS risk factors were observed following assisted ambulation, FES-hybrid,
- 24 FES-rowing, and FES-RT.

25	Conclusion Upper-body aerobic exercise training (>75% maximum heart rate) appears to
26	improve waist circumference and hepatic insulin sensitivity, but appears insufficient for
27	improving fasting glucose, lipid profile, or resting blood pressure. The addition of RT to
28	upper-body aerobic exercise may elicit favourable changes in the lipid profile. More high
29	quality studies are needed to confirm if FES-cycling is effective at improving peripheral
30	insulin sensitivity.
31	
32	Key Words spinal cord injuries, exercise therapy, metabolic diseases
33	
34	Abbreviations
35	CMS cardiometabolic syndrome
36	DBP diastolic blood pressure
37	ES effect size
38	FES functional electrical stimulation
39	HDL-C high-density lipoprotein-cholesterol
40	HOMA-IR homeostatic model assessment insulin resistance
41	HRR heart rate reserve
42	LDL-C low-density lipoprotein-cholesterol
43	RT resistance training
44	RCT randomised controlled trial
45	SBP systolic blood pressure
46	SCI spinal cord injury
47	TC total cholesterol
48	TG triglycerides

Persons with a spinal cord injury (SCI) are at an increased risk of cardiovascular disease and diabetes compared to able-bodied individuals [1, 2]. The risk of developing these chronic diseases is raised in individuals who present with a clustering of associated risk factors including: obesity, insulin resistance, dyslipidaemia, and hypertension, or as commonly referred to, cardiometabolic syndrome (CMS) [3]. The International Diabetes Federation defines CMS as central obesity (indicated by waist circumference), plus the presence (or treatment) of two of more of the following: hypertriglyceridemia ($\geq 1.7 \text{ mmol/L}$), reduced high-density lipoprotein-cholesterol (HDL-C) (< 1.03 mmol/L for men, < 1.29 mmol/L for women), hypertension (systolic blood pressure $\geq 130 \text{ mmHg}$, or diastolic blood pressure $\geq 85 \text{ mmHg}$), and raised fasting plasma glucose ($\geq 5.6 \text{ mmol/L}$, or diagnosed with type 2 diabetes) [4]. A waist circumference greater than 94 cm and/or a body mass index of greater than 22 kg/m² have been suggested as suitable cut-points to define central obesity in SCI [5, 6]. The prevalence of CMS in chronic SCI appears to be high; with the largest study to date (n=473) reporting a prevalence rate of 57.5% [7].

There is strong evidence that exercise is an effective countermeasure for the prevention of chronic disease and the treatment of CMS risk factors in the able-bodied population [8]. This has allowed national and global health organisations to produce guidelines regarding the total volume and intensity of physical activity (minimum of 150 min/week of moderate-intensity, or 75 minutes/week of vigorous-intensity) required to improve cardiometabolic health [9, 10]. However, as the most recent systematic review of the effect of exercise on health in SCI concluded, the evidence base for spinal cord injured persons "lags far behind" that for the general population [11]. This review formed the basis for the latest SCI-exercise guidelines, which recommend adults with a chronic SCI perform a minimum of 90 min/week of moderate-to-vigorous intensity aerobic exercise to improve cardiometabolic health [12]. Additional systematic reviews have also reported beneficial effects of exercise on specific CMS risk

factors, including systemic inflammation (C - reactive protein) and obesity (fat mass and waist circumference) in persons with chronic SCI [13, 14].

Since the last systematic search of the literature by van der Sheer and colleagues (search date: 1st Jan 2016), several randomised controlled trials assessing the effect of exercise training on CMS risk factors in SCI have been published. However, this systematic review did not address clinical thresholds for CMS risk factors at baseline, the magnitude of change following exercise training, and how different exercise modalities may impact specific individual CMS biomarkers. These questions are important for practitioners prescribing exercise to patients presenting with CMS risk factors, and researchers designing future studies in this field. A review which addresses these importance issues and focuses specifically on how different forms of exercise impacts on individual CMS risk factors in chronic SCI is therefore required. The aim of this systematic review is to determine the effect of different exercise modality interventions on CMS risk factors in adults with chronic SCI.

METHODS

The study inclusion criteria and planned analysis were specified in advance (PROSPERO: CRD42018105110) and the Preferred Reporting Items for Systematic Review and Meta-Analyses (PRISMA) guidelines were followed [15]. The databases of PubMed, Web of Science, EMBASE, and Scopus (Elsevier) were searched on 22nd August 2018, using a search strategy formulated based on a similar previous systematic review [11]. The search was repeated on 31st July 2019 to identify any additional articles prior to publication. The search strategy was piloted to ensure known articles were included and reviewed by two authors (MF & TN). The full search strategy for PubMed is presented in Supplement 1 as an exemplar. Briefly, the search was performed by combining key words associated with SCI (e.g., "paraplegia", "spinal cord lesion"), exercise, (e.g., "physical activity", "resistance training",

"functional electrical stimulation") and CMS risk factors (e.g., "glucose", "BMI", "blood pressure"). The reference list of included items and previous systematic reviews were checked, and hand-searching of relevant journals was performed to search for any additional studies (Journal of Spinal Cord Medicine (1982-2018) and Archives of Physical Medicine and Rehabilitation (1985-2018)).

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Titles and abstracts of retrieved articles were independently screened for relevance by two reviewers (MF & TN). The same two reviewers independently assessed the full text of relevant articles for eligibility. In the event of any disagreements in article selection, a third reviewer (JB) made the final decision. Articles were included if they met the criteria according to the PICOS structure: i) participants - \geq 50% of participants were aged \geq 18 years old, and had a chronic SCI (≥1 year post-injury), ii) *intervention* - included an exercise training programme (any, or combination of: voluntary upper-body exercise, lower-body functional electrical stimulation (FES), and assisted ambulation training) lasting ≥2 weeks, iii) *comparison* – studies comparing exercise intervention to a control group or pre-intervention data, iv) outcomes study included at least one CMS risk factor as an outcome variable (see Table 1) [4], and v) study design - study employed and reported quantitative statistical analysis to determine the impact of the exercise intervention on the relevant CMS risk outcome(s) (i.e. case reports and case-series were excluded), and was published in an English-language peer-reviewed journal (i.e. abstracts and conference proceedings were excluded) between 1st January 1970 and the final search date. Studies involving solely neuromuscular electrical stimulation (NMES) with no functional movement and passive cycling were excluded on the basis that the skeletal muscle contractions produced during these activities do not directly produce a functional movement, and therefore cannot be classed as exercise, per se. Studies assessing the impact of exercise on solely blood pressure amongst tetraplegics were excluded on the basis that the aim of the exercise intervention was to increase resting blood pressure, and therefore was not reflective of a CMS risk factor (i.e. hypertension).

Two articles did not identify participants' time since injury [16, 17]. The corresponding authors were contacted by email and asked to provide clarification and given two weeks to respond. Both articles were excluded as the corresponding authors were unable to provide this information.

Two reviewers (MF and JM) independently evaluated the quality of included studies using a modified Downs and Black scale [18]. In the modified version, the scoring for question 27 (relating to statistical power) is simplified to "Yes" (1) or "No" (0). In the event of any discrepancies in scoring, discussion between the reviewers was used to reach a consensus. The total Downs & Black score for each article was expressed as a percentage of the maximum score possible (28) to allow categorisation of study quality [19]. Articles were classified as high (\geq 66.7%), fair (between 50.0% and 66.6%), or low (<50.0%) quality [19].

An insufficient number of studies examined the same outcomes following similar exercise modalities, precluding a meta-analysis. Therefore, a coding system [19] was used to summarise the effect of different exercise training modalities on each CMS risk factor. If 0-33% of studies reported a statistically significant change in a specific CMS risk factor following exercise training, the result was categorised as 'no effect'. If 34-59% of studies reported a statistically significant change in a CMS risk factor following exercise training, the result was categorised as 'inconsistent'. If 60-100% of studies reported a statistically significant change in a CMS risk factor following exercise training, the result was categorised as 'positive'. If four or more studies reported the same effect, the result was highlighted in bold to indicate a consistent finding. The findings from one particular study [20] were counted as non-significant for summary coding, due to the significance being set at p<0.10, with actual p

values not reported. Data extraction was performed by MF, and later checked independently by TN, JM, and JB.

To aid interpretation of results, group average values at baseline for body mass index (≥22 kg/m²) [6], waist circumference (>94 cm) [5], triglycerides (TG) (≥1.7 mmol/L), total cholesterol (TC) (≥5 mmol/L), low-density lipoprotein (LDL-C) (>3 mmol/L), HDL-C (<1.03 mmol/L), fasting glucose (≥5.6 mmol/L), systolic blood pressure (SBP) (≥130 mmHg), and diastolic blood pressure (DBP) (≥85 mmHg) [4] were highlighted to indicate that they can be classified as clinically high, according to the International Diabetes Federation and SCI-specific guidelines (Tables 3-9).

RESULTS

The initial database search yielded a total of 2450 unique records, of which 2245 were excluded following title and abstract screening. An additional 10 articles were retrieved from; hand-searching of relevant journals (n=1), relevant systematic reviews (n=2), the associated reference list of an included paper (n=4), and the updated search (n=3). Therefore, the full-text of 215 studies were subsequently assessed, three papers [21-23] contained data presented in another article, and these were removed from all analysis, leaving 65 articles for final review. The study selection process is detailed in Figure 1.

There was substantial agreement between reviewer's for title and abstract screening (k=0.635, 95% CI: 0.581, 0.689), and almost perfect agreement for the full-text screening (k=0.880, 95% CI: 0.811, 0.949) [24].

We identified studies as pre-post designs (n=47), RCTs (n=15), non-randomised controlled trials (n=2), and a retrospective cohort study (n=1). Numerous studies utilised arm-cranking (n=9), wheelchair ergometry (n=3), wheelchair treadmill propulsion (n=2), or hand-

cycling (n=2). These 16 studies were grouped together for analysis as voluntary upper-body aerobic exercise (Table 3). Seven studies utilised upper-body resistance training (RT) (with or without upper-body aerobic exercise) (Table 4). The most common exercise modality was FEScycling (n=17) (Table 5). Six studies utilised FES-resistance training (FES-RT) exercise (in the form of non-isometric knee extensions), and three studies involved a combination of FEScycling and FES-RT (Table 6). Studies which involved hybrid functional electrical stimulation (FES)-cycling (n=4) or FES-rowing (n=4) were grouped together as they both involve lower-body FES combined with voluntary upper-body aerobic exercise (Table 7). Several studies utilised solely body weight supported treadmill training (n=6), FES-walking, exoskeletal body weight supported treadmill training (n=1), or robotic body weight supported treadmill training (n=1). These 10 studies were grouped together for analysis (Table 8). Studies that involved a combination of upper-body aerobic, upper-body RT and neuromuscular stimulation (n=1), or a combination of lower-body FES-RT, and BWSTT (n=1), were not grouped for qualitative analysis (Table 9).

Intervention durations ranged from four to 52 weeks, with the most common length of 12 weeks (n=14). Training frequency ranged from 1 to 7 sessions per week, with three times per week the most common frequency of exercise performed (n=35). No serious adverse events were reported in any of the included studies.

Sample sizes ranged from four to 48. Only seven studies reported a-priori sample size calculations, and four of these met their target sample size (Table 10). There was a total of 872 participants (658 men, 110 women, 104 NR) (Table 10). There were nine studies classified as high quality, 35 studies classified as fair quality, and 21 studies classified as low quality. The most commonly assessed outcome measures for obesity, glycaemic control, dyslipidaemia, inflammation, vascular dysregulation, and thrombotic state were body mass (n=28), interleukin-6 (n=7), HDL-C (n=23), fasting glucose (n=18), PAI-1 (n=3), and systolic blood

- pressure (n=22), respectively. No studies reported outcome measures of hip circumference,
- liver fat content, apolipoprotein B, or proinsulin.

DISCUSSION

There are consistent findings that voluntary upper-body aerobic exercise (>75% HR_{MAX}) is effective in reducing waist circumference, and improving hepatic insulin sensitivity (i.e. fasting insulin concentration and HOMA-IR), however it does not appear to improve fasting glucose concentrations, lipid profile or resting blood pressure in persons with chronic SCI. The addition of upper-body RT appears to have an inconsistent effect on lipid profiles, but given the limited number of high-quality studies on combined exercise modalities, more research is needed in this area. FES-cycling may improve outcomes relating to peripheral insulin sensitivity (i.e. ability of the skeletal muscle to dispose of glucose), but more high-quality studies are required to strengthen the available evidence. There is insufficient evidence to conclude if FES-resistance training, FES-hybrid, FES-rowing, or assisted ambulation training improves any of these CMS risk factors.

Four [27, 25, 34, 33] of the six studies utilising upper-body aerobic exercise reported a reduction in supine waist circumference (-1.9 to -3.7 cm, ES: 0.26-2.67), indicating that this form of exercise is effective for reducing central obesity. A reduction in waist circumference (-2.5 cm) was achieved with as few as 64 min/week of exercise at 65-75% HRR [25], though this reduction did not translate to any change in android fat mass [25]. There was also no change in visceral adipose tissue [26] following 180 min/week at 60-65% VÔ2peak of upper-body aerobic exercise. Future studies should combine both surrogate and gold-standard measures (i.e. DEXA/CT derived) of central obesity/adiposity to further elucidate changes in body composition. Given the relatively small skeletal muscle mass involved in upper-body aerobic exercise, it is perhaps unsurprising that there were consistent findings that body mass and BMI were unchanged, as reported in a previous systematic review [14]. Whilst not part of the search strategy, only one study in this category measured free-living energy intake and expenditure

during the exercise intervention [26]. In order to better understand the isolated impact of prescribed exercise interventions on energy balance and body composition, future studies should also attempt to estimate total energy intake and total energy expenditure. This would account for any compensatory changes in diet or exercise behaviours, providing a better understanding of the overall impact of exercise interventions on energy balance in SCI [90]. Guidelines for measuring these variables in persons with chronic SCI have been published elsewhere [91].

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Four [25, 28, 26, 33] of the five studies that measured fasting insulin resistance by HOMA-IR and/or fasting insulin concentrations reported a reduction (22-40%, ES: 1.07-1.78) following upper-body aerobic exercise, suggesting that this form of exercise is effective at improving hepatic insulin sensitivity (i.e. ability of the liver to dispose of glucose). The single study [31] to find no statistically significant change in fasting insulin concentration following upper-body aerobic exercise, reported that all five participants had a lower insulin concentration (22-76%, ES: 0.41) post-training, indicating that the study simply lacked the statistical power to demonstrate an effect. Despite the improvement in hepatic insulin sensitivity [92] observed following upper-body aerobic exercise, the three studies [26, 28, 31] that measured outcomes relating to peripheral insulin sensitivity [93] found no changes following training. This is likely as a result of the limited skeletal muscle mass involved (i.e. limited sink for glucose disposal). Furthermore, the upper-body skeletal musculature is usually already well-conditioned from habitual wheelchair propulsion, meaning that moderateintensity upper-body exercise is likely an insufficient stimulus to substantially promote molecular adaptations (e.g. GLUT4 translocation, mitochondrial biogenesis) associated with improved peripheral insulin sensitivity [94]. One high quality study reported no improvement in glucose or insulin area under the curve despite 180 min/week of exercise at 60-65% VO₂peak [26]. This suggests that even large volumes of upper-body aerobic exercise above the

recommended guidelines of 90 min/week [12] may be insufficient to improve markers of peripheral insulin sensitivity.

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There are also numerous studies indicating that upper-body aerobic exercise alone does not improve fasting glucose, resting blood pressure (SBP, DBP), or lipid profiles (TC, HDL-C, LDL-C, and TG). All eight studies [25, 26, 28, 31-35] measuring fasting glucose reported no change following upper-body aerobic exercise. However, only one study [34] reported a clinically elevated group mean glucose concentration at baseline (≥5.6 mmol/L). Nine studies [29, 35, 38, 39, 25, 26, 34, 32, 31] measured changes in resting blood pressure following upperbody aerobic exercise. The only study [34] where participants presented with clinically elevated systolic blood pressure (≥130 mmHg) at baseline reported a reduction (3 mmHg, ES: 0.66) following 10 weeks of exercise training (4 sessions/week 50-70% HRR, 60 min). Thus, a basement effect may explain the lack of significant changes in fasting glucose and resting blood pressure in participants presenting with healthy values at baseline. Eight studies measured TG, TC, HDL-C, or LDL-C [25, 26, 28, 32-35, 20] following upper-body aerobic exercise, including four with clinically high mean concentrations at baseline. Only two studies reported a significant reduction in any variable. One study [34] reported a 25% reduction (ES: 0.31) in TG in participants with a clinically elevated mean concentrations at baseline (≥1.7 mmol/L). One study reported improvements in HDL-C, LDL-C, TC: HDL-C and TG following 60 mins/week at 70-80% HRR, however the threshold for significance was set at p<0.10 [40]. It therefore appears that upper-body aerobic exercise may not be an adequate stimulus to improve blood lipid profile irrespective of baseline values. This is likely due to the low energy expenditure achieved through upper-body exercise, which appears to drive changes in the lipid profile [95].

Upper-body RT (with or without aerobic exercise) appears to reduce central obesity, with three [42-44] out of four studies reporting a reduction in waist circumference (-

1.0 to -2.6 cm) or waist to hip ratio (-0.02). These changes were accompanied by a decrease in whole-body fat mass and visceral adipose tissue following 120 min/week of training (3 x 10 of 50-70% 1RM, 20 min at 3-6 RPE) [42]. Upper-body RT (with or without aerobic exercise) may elicit improvements in lipid profile, with four [43-45, 40] out of the five retrieved studies reporting a beneficial effect of at least one marker (TC, HDL-C, LDL-C, TC: HDL-C, and TG). However, more studies are needed to determine this, particularly given the high-quality study reporting no change in the lipid profile following 16-weeks of twice-weekly combined training [42].

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Five [50, 54, 58, 60, 62] of the six studies to measure outcomes relating to peripheral insulin sensitivity reported a significant improvement following FES-cycling. The largest of these studies (n=18) [54] reported a significant reduction in glucose and insulin at multiple time-points during a 2-h oral glucose tolerance test following 10 weeks of exercise (2-3 sessions/week, 30 min). However, four of these studies were rated as low quality, and therefore more high-quality studies are needed to confirm if FES-cycling can improve peripheral insulin sensitivity, which upper-body exercise appears unable to achieve. Surprisingly, we identified no RCT's assessing the efficacy of FES-cycling compared to a true control group (i.e. passive cycling or stretching), which should be addressed in future research. Four studies reported no change in body mass following FES-hybrid or FES-rowing training. There was a distinct lack of training studies with sufficient breadth of outcomes to make any other meaningful conclusions on the effect of FES-RT, FES-hybrid, FES-rowing and assisted ambulation on CMS risk factors. Nonetheless, given that hybrid training (2 sessions/week, 18-32 min, 65-75% HRR) [25] improved a multitude of CMS risk factors (waist circumference, android fat percentage, TG, DBP), and that different exercise modalities appear to offer specific benefits to CMS risk factors, other rigorously conducted prospective studies assessing multimodal (e.g.

FES-cycling combined with upper-body aerobic and resistance exercise) interventions should be conducted in this area of promise.

This review has highlighted the lack of research assessing novel markers of CMS risk, including outcomes relating to inflammation, DEXA/CT derived measured of central adiposity, and endothelial function. It is clear that many studies in the area recruit a convenience sample of relatively active and lean individuals, who are not reflective of the wider, chronic SCI population (i.e. poor metabolic health), which should be considered when interpreting results. For example, individuals with SCI have a significantly lower HDL-C compared to able-bodied controls (1.06 vs 1.28 mmol/L) [96], however only five of the 23 studies to measure HDL-C had a clinically low mean concentration at baseline (<1.03 mmol/L). As is widely acknowledged, this review has also confirmed the existing evidence base of exercise and CMS risk in SCI lacks sufficiently powered (four in total identified), high-quality studies (eight in total identified). However, this review identified 16 additional studies, published since the previous systematic review by van der Scheer and colleagues [11] that were all categorised as fair or high quality, including eight RCT's.

Study Limitations

The major limitation of this systematic review is the use of summary coding to draw conclusions regarding the effect of each exercise modality on specific CMS risk factors. Due to the variability in CMS risk factors measured, exercise modes and training parameters (i.e. exercise intensity and volume), and participant characteristics (i.e. paraplegic vs. tetraplegic), a meta-analysis was not possible. Whilst the coding system provides a useful assessment of the consistency of findings in the field, it uses arbitrary classifications and does not distinguish studies of differing quality. However, when studies rated as 'low-quality' were removed from this analysis (Supplement 2), the conclusions remained unchanged, with the exception of

potential of FES-cycling to improve peripheral insulin sensitivity. Further, given that the vast majority of included studies lacked sufficient statistical power, there is a risk of a type II error in the conclusions formed. Finally, this review did not include acute SCI as van der Scheer and colleagues [11] determined there was an "absence of high-quality, consistent evidence" in this area, a view which still appears to be true.

CONCLUSIONS

In summary, this systematic review has provided evidence that in adults with chronic SCI, upper-body aerobic exercise improves outcomes relating to central obesity and hepatic insulin sensitivity, but is not sufficient to improve fasting glucose, lipid profiles, or resting blood pressure. Practitioners should consider prescribing moderate-to-vigorous intensity (>75% HR_{MAX}) upper-body aerobic exercise to improve fasting glycaemic control and central obesity. To elicit improvements in lipid profile, this should be combined with upper-body resistance training. More high-quality randomised controlled trials assessing novel markers of CMS and responses to combined exercise interventions (e.g. aerobic exercise with resistance training), high-intensity exercise interventions, and FES-based exercise are needed to inform and refine evidence-based exercise guidelines for the prevention and management of CMS in this population.

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Figure 1. PRISMA flow diagram

- 1 The effect of exercise on cardiometabolic risk factors in adults with chronic spinal cord
- 2 injury: A systematic review

ABSTRACT

3

- 5 **Objective** To determine the effects of exercise on individual cardiometabolic syndrome (CMS)
- 6 risk factors in adults with chronic spinal cord injury (SCI).
- 7 **Design** Systematic review.
- 8 **Data sources** English language searches of PubMed, Web of Science, EMBASE, and Scopus
- 9 (01/01/1970 to 31/07/2019).
- 10 Eligibility criteria for selecting studies (1) original articles with statistical analysis, (2)
- participants were adults with a SCI sustained \geq 1-year ago, (3) exercise intervention duration
- 12 \geq 2 weeks, and (4) included any CMS risk factor as an outcome. The methodological quality
- of articles was assessed using the Downs and Black score.
- 14 **Results** Sixty-five studies were included for the final analysis, including nine studies classified
- as high quality (\geq 66%), 35 studies classified as fair quality (50-66%), and 21 studies classified
- as low quality (<50%). Improvements in waist circumference (4/6 studies) and markers of
- hepatic insulin sensitivity (4/5 studies) were reported following upper-body aerobic exercise
- training, but no improvements in fasting glucose (8/8 studies), lipid profile (6/8 studies),
- systolic (8/9 studies) or diastolic blood pressure (9/9 studies) were observed. Improvements in
- 20 <u>markers of peripheral insulin sensitivity (5/6 studies)</u> were observed following functional
- 21 electrical stimulation (FES)-cycling. Improvements in lipid profile (4/5 studies) were observed
- following upper-body resistance training (RT) (with or without aerobic exercise). No consistent
- 23 improvements in CMS risk factors were observed following assisted ambulation, FES-hybrid,
- 24 FES-rowing, and FES-RT.

25	Conclusion Upper-body aerobic exercise training (>75% maximum heart rate) appears to
26	improve waist circumference and hepatic insulin sensitivity, but appears insufficient for
27	improving fasting glucose, lipid profile, or resting blood pressure. The addition of RT to
28	upper-body aerobic exercise may elicit favourable changes in the lipid profile. More high-
29	quality studies are needed to confirm if FES-cycling is effective at improving peripheral
30	insulin sensitivity.
31	
32	Key Words spinal cord injuries, exercise therapy, metabolic diseases
33	
34	Abbreviations
35	CMS cardiometabolic syndrome
36	DBP diastolic blood pressure
37	ES effect size
38	FES functional electrical stimulation
39	HDL-C high-density lipoprotein-cholesterol
40	HOMA-IR homeostatic model assessment insulin resistance
41	HRR heart rate reserve
42	LDL-C low-density lipoprotein-cholesterol
43	RT resistance training
44	RCT randomised controlled trial
45	SBP systolic blood pressure
46	SCI spinal cord injury
47	TC total cholesterol
48	TG triglycerides

Persons with a spinal cord injury (SCI) are at an increased risk of cardiovascular disease and diabetes compared to able-bodied individuals [1, 2]. The risk of developing these chronic diseases is raised in individuals who present with a clustering of associated risk factors including: obesity, insulin resistance, dyslipidaemia, and hypertension, or as commonly referred to, cardiometabolic syndrome (CMS) [3]. The International Diabetes Federation defines CMS as central obesity (indicated by waist circumference), plus the presence (or treatment) of two of more of the following: hypertriglyceridemia ($\geq 1.7 \text{ mmol/L}$), reduced high-density lipoprotein-cholesterol (HDL-C) (< 1.03 mmol/L for men, < 1.29 mmol/L for women), hypertension (systolic blood pressure $\geq 130 \text{ mmHg}$, or diastolic blood pressure $\geq 85 \text{ mmHg}$), and raised fasting plasma glucose ($\geq 5.6 \text{ mmol/L}$, or diagnosed with type 2 diabetes) [4]. A waist circumference greater than 94 cm and/or a body mass index of greater than 22 kg/m² have been suggested as suitable cut-points to define central obesity in SCI [5, 6]. The prevalence of CMS in chronic SCI appears to be high; with the largest study to date (n=473) reporting a prevalence rate of 57.5% [7].

There is strong evidence that exercise is an effective countermeasure for the prevention of chronic disease and the treatment of CMS risk factors in the able-bodied population [8]. This has allowed national and global health organisations to produce guidelines regarding the total volume and intensity of physical activity (minimum of 150 min/week of moderate-intensity, or 75 minutes/week of vigorous-intensity) required to improve cardiometabolic health [9, 10]. However, as the most recent systematic review of the effect of exercise on health in SCI concluded, the evidence base for spinal cord injured persons "lags far behind" that for the general population [11]. This review formed the basis for the latest SCI-exercise guidelines, which recommend adults with a chronic SCI perform a minimum of 90 min/week of moderate-to-vigorous intensity aerobic exercise to improve cardiometabolic health [12]. Additional systematic reviews have also reported beneficial effects of exercise on specific CMS risk

factors, including systemic inflammation (C - reactive protein) and obesity (fat mass and waist circumference) in persons with chronic SCI [13, 14].

Since the last systematic search of the literature by van der Sheer and colleagues (search date: 1st Jan 2016), several randomised controlled trials assessing the effect of exercise training on CMS risk factors in SCI have been published. However, this systematic review did not address clinical thresholds for CMS risk factors at baseline, the magnitude of change following exercise training, and how different exercise modalities may impact specific individual CMS biomarkers. These questions are important for practitioners prescribing exercise to patients presenting with CMS risk factors, and researchers designing future studies in this field. A review which addresses these importance issues and focuses specifically on how different forms of exercise impacts on individual CMS risk factors in chronic SCI is therefore required. The aim of this systematic review is to determine the effect of different exercise modality interventions on CMS risk factors in adults with chronic SCI.

METHODS

The study inclusion criteria and planned analysis were specified in advance (PROSPERO: CRD42018105110) and the Preferred Reporting Items for Systematic Review and Meta-Analyses (PRISMA) guidelines were followed [15]. The databases of PubMed, Web of Science, EMBASE, and Scopus (Elsevier) were searched on 22nd August 2018, using a search strategy formulated based on a similar previous systematic review [11]. The search was repeated on 31st July 2019 to identify any additional articles prior to publication. The search strategy was piloted to ensure known articles were included and reviewed by two authors (MF & TN). The full search strategy for PubMed is presented in Supplement 1 as an exemplar. Briefly, the search was performed by combining key words associated with SCI (e.g., "paraplegia", "spinal cord lesion"), exercise, (e.g., "physical activity", "resistance training",

"functional electrical stimulation") and CMS risk factors (e.g., "glucose", "BMI", "blood pressure"). The reference list of included items and previous systematic reviews were checked, and hand-searching of relevant journals was performed to search for any additional studies (Journal of Spinal Cord Medicine (1982-2018) and Archives of Physical Medicine and Rehabilitation (1985-2018)).

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Titles and abstracts of retrieved articles were independently screened for relevance by two reviewers (MF & TN). The same two reviewers independently assessed the full text of relevant articles for eligibility. In the event of any disagreements in article selection, a third reviewer (JB) made the final decision. Articles were included if they met the criteria according to the PICOS structure: i) participants - ≥50% of participants were aged ≥18 years old, and had a chronic SCI (≥1 year post-injury), ii) *intervention* - included an exercise training programme (any, or combination of: voluntary upper-body exercise, lower-body functional electrical stimulation (FES), and assisted ambulation training) lasting ≥2 weeks, iii) *comparison* – studies comparing exercise intervention to a control group or pre-intervention data, iv) outcomes study included at least one CMS risk factor as an outcome variable (see Table 1) [4], and v) study design - study employed and reported quantitative statistical analysis to determine the impact of the exercise intervention on the relevant CMS risk outcome(s) (i.e. case reports and case-series were excluded), and was published in an English-language peer-reviewed journal (i.e. abstracts and conference proceedings were excluded) between 1st January 1970 and the final search date. Studies involving solely neuromuscular electrical stimulation (NMES) with no functional movement and passive cycling were excluded on the basis that the skeletal muscle contractions produced during these activities do not directly produce a functional movement, and therefore cannot be classed as exercise, per se. Studies assessing the impact of exercise on solely blood pressure amongst tetraplegics were excluded on the basis that the aim

of the exercise intervention was to increase resting blood pressure, and therefore was not reflective of a CMS risk factor (i.e. hypertension).

Two articles did not identify participants' time since injury [16, 17]. The corresponding authors were contacted by email and asked to provide clarification and given two weeks to respond. Both articles were excluded as the corresponding authors were unable to provide this information.

Two reviewers (MF and JM) independently evaluated the quality of included studies using a modified Downs and Black scale [18]. In the modified version, the scoring for question 27 (relating to statistical power) is simplified to "Yes" (1) or "No" (0). In the event of any discrepancies in scoring, discussion between the reviewers was used to reach a consensus. The total Downs & Black score for each article was expressed as a percentage of the maximum score possible (28) to allow categorisation of study quality [19]. Articles were classified as high (≥66.7%), fair (between 50.0% and 66.6%), or low (<50.0%) quality [19].

An insufficient number of studies examined the same outcomes following similar exercise modalities, precluding a meta-analysis. Therefore, a coding system [19] was used to summarise the effect of different exercise training modalities on each CMS risk factor. If 0-33% of studies reported a statistically significant change in a specific CMS risk factor following exercise training, the result was categorised as 'no effect'. If 34-59% of studies reported a statistically significant change in a CMS risk factor following exercise training, the result was categorised as 'inconsistent'. If 60-100% of studies reported a statistically significant change in a CMS risk factor following exercise training, the result was categorised as 'positive'. If four or more studies reported the same effect, the result was highlighted in bold to indicate a consistent finding. The findings from one particular study [20] were counted as non-significant for summary coding, due to the significance being set at p<0.10, with actual p

values not reported. Data extraction was performed by MF, and later checked independently by TN, JM, and JB.

To aid interpretation of results, group average values at baseline for body mass index (\geq 22 kg/m²) [6], waist circumference (>94 cm) [5], triglycerides (TG) (\geq 1.7 mmol/L), total cholesterol (TC) (\geq 5 mmol/L), low-density lipoprotein (LDL-C) (>3 mmol/L), HDL-C (<1.03 mmol/L), fasting glucose (\geq 5.6 mmol/L), systolic blood pressure (SBP) (\geq 130 mmHg), and diastolic blood pressure (DBP) (\geq 85 mmHg) [4] were highlighted to indicate that they can be classified as clinically high, according to the International Diabetes Federation and SCI-specific guidelines (Tables 3-9).

RESULTS

The initial database search yielded a total of 2450 unique records, of which 2245 were excluded following title and abstract screening. An additional 10 articles were retrieved from; hand-searching of relevant journals (n=1), relevant systematic reviews (n=2), the associated reference list of an included paper (n=4), and the updated search (n=3). Therefore, the full-text of 215 studies were subsequently assessed, three papers [21-23] contained data presented in another article, and these were removed from all analysis, leaving 65 articles for final review. The study selection process is detailed in Figure 1.

There was substantial agreement between reviewer's for title and abstract screening (k=0.635, 95% CI: 0.581, 0.689), and almost perfect agreement for the full-text screening (k=0.880, 95% CI: 0.811, 0.949) [24].

We identified studies as pre-post designs (n=47), RCTs (n=15), non-randomised controlled trials (n=2), and a retrospective cohort study (n=1). Numerous studies utilised arm-cranking (n=9), wheelchair ergometry (n=3), wheelchair treadmill propulsion (n=2), or hand-

cycling (n=2). These 16 studies were grouped together for analysis as voluntary upper-body aerobic exercise (Table 3). Seven studies utilised upper-body resistance training (RT) (with or without upper-body aerobic exercise) (Table 4). The most common exercise modality was FEScycling (n=17) (Table 5). Six studies utilised FES-resistance training (FES-RT) exercise (in the form of non-isometric knee extensions), and three studies involved a combination of FEScycling and FES-RT (Table 6). Studies which involved hybrid functional electrical stimulation (FES)-cycling (n=4) or FES-rowing (n=4) were grouped together as they both involve lower-body FES combined with voluntary upper-body aerobic exercise (Table 7). Several studies utilised solely body weight supported treadmill training (n=6), FES-walking, exoskeletal body weight supported treadmill training (n=1), or robotic body weight supported treadmill training (n=1). These 10 studies were grouped together for analysis (Table 8). Studies that involved a combination of upper-body aerobic, upper-body RT and neuromuscular stimulation (n=1), or a combination of lower-body FES-RT, and BWSTT (n=1), were not grouped for qualitative analysis (Table 9).

Intervention durations ranged from four to 52 weeks, with the most common length of 12 weeks (n=14). Training frequency ranged from 1 to 7 sessions per week, with three times per week the most common frequency of exercise performed (n=35). No serious adverse events were reported in any of the included studies.

Sample sizes ranged from four to 48. Only seven studies reported a-priori sample size calculations, and four of these met their target sample size (Table 10). There was a total of 872 participants (658 men, 110 women, 104 NR) (Table 10). There were nine studies classified as high quality, 35 studies classified as fair quality, and 21 studies classified as low quality. The most commonly assessed outcome measures for obesity, glycaemic control, dyslipidaemia, inflammation, vascular dysregulation, and thrombotic state were body mass (n=28), interleukin-6 (n=7), HDL-C (n=23), fasting glucose (n=18), PAI-1 (n=3), and systolic blood

- pressure (n=22), respectively. No studies reported outcome measures of hip circumference,
- liver fat content, apolipoprotein B, or proinsulin.

DISCUSSION

There are consistent findings that voluntary upper-body aerobic exercise (>75% HR_{MAX}) is effective in reducing waist circumference, and improving hepatic insulin sensitivity (i.e. fasting insulin concentration and HOMA-IR), however it does not appear to improve fasting glucose concentrations, lipid profile or resting blood pressure in persons with chronic SCI. The addition of upper-body RT appears to have an inconsistent effect on lipid profiles, but given the limited number of high-quality studies on combined exercise modalities, more research is needed in this area. FES-cycling may improve outcomes relating to peripheral insulin sensitivity (i.e. ability of the skeletal muscle to dispose of glucose), but more high-quality studies are required to strengthen the available evidence. There is insufficient evidence to conclude if FES-resistance training, FES-hybrid, FES-rowing, or assisted ambulation training improves any of these CMS risk factors.

Four [27, 25, 34, 33] of the six studies utilising upper-body aerobic exercise reported a reduction in supine waist circumference (-1.9 to -3.7 cm, ES: 0.26-2.67), indicating that this form of exercise is effective for reducing central obesity. A reduction in waist circumference (-2.5 cm) was achieved with as few as 64 min/week of exercise at 65-75% HRR [25], though this reduction did not translate to any change in android fat mass [25]. There was also no change in visceral adipose tissue [26] following 180 min/week at 60-65% VÔ2peak of upper-body aerobic exercise. Future studies should combine both surrogate and gold-standard measures (i.e. DEXA/CT derived) of central obesity/adiposity to further elucidate changes in body composition. Given the relatively small skeletal muscle mass involved in upper-body aerobic exercise, it is perhaps unsurprising that there were consistent findings that body mass and BMI were unchanged, as reported in a previous systematic review [14]. Whilst not part of the search strategy, only one study in this category measured free-living energy intake and expenditure

during the exercise intervention [26]. In order to better understand the isolated impact of prescribed exercise interventions on energy balance and body composition, future studies should also attempt to estimate total energy intake and total energy expenditure. This would account for any compensatory changes in diet or exercise behaviours, providing a better understanding of the overall impact of exercise interventions on energy balance in SCI [90]. Guidelines for measuring these variables in persons with chronic SCI have been published elsewhere [91].

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Four [25, 28, 26, 33] of the five studies that measured fasting insulin resistance by HOMA-IR and/or fasting insulin concentrations reported a reduction (22-40%, ES: 1.07-1.78) following upper-body aerobic exercise, suggesting that this form of exercise is effective at improving hepatic insulin sensitivity (i.e. ability of the liver to dispose of glucose). The single study [31] to find no statistically significant change in fasting insulin concentration following upper-body aerobic exercise, reported that all five participants had a lower insulin concentration (22-76%, ES: 0.41) post-training, indicating that the study simply lacked the statistical power to demonstrate an effect. Despite the improvement in hepatic insulin sensitivity [92] observed following upper-body aerobic exercise, the three studies [26, 28, 31] that measured outcomes relating to peripheral insulin sensitivity [93] found no changes following training. This is likely as a result of the limited skeletal muscle mass involved (i.e. limited sink for glucose disposal). Furthermore, the upper-body skeletal musculature is usually already well-conditioned from habitual wheelchair propulsion, meaning that moderateintensity upper-body exercise is likely an insufficient stimulus to substantially promote molecular adaptations (e.g. GLUT4 translocation, mitochondrial biogenesis) associated with improved peripheral insulin sensitivity [94]. A high quality study reported no improvement in glucose or insulin area under the curve despite 180 min/week of exercise at 60-65% VO₂peak [26]. This suggests that even large volumes of upper-body aerobic exercise above the recommended guidelines of 90 min/week [12] may be insufficient to improve markers of peripheral insulin sensitivity.

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There are also numerous studies indicating that upper-body aerobic exercise alone does not improve fasting glucose, resting blood pressure (SBP, DBP), or lipid profiles (TC, HDL-C, LDL-C, and TG). All eight studies [25, 26, 28, 31-35] measuring fasting glucose reported no change following upper-body aerobic exercise. However, only one study [34] reported a clinically elevated group mean glucose concentration at baseline (≥5.6 mmol/L). Nine studies [29, 35, 38, 39, 25, 26, 34, 32, 31] measured changes in resting blood pressure following upperbody aerobic exercise. The only study [34] where participants presented with clinically elevated systolic blood pressure (≥130 mmHg) at baseline reported a reduction (3 mmHg, ES: 0.66) following 10 weeks of exercise training (4 sessions/week 50-70% HRR, 60 min). Thus, a basement effect may explain the lack of significant changes in fasting glucose and resting blood pressure in participants presenting with healthy values at baseline. Eight studies measured TG, TC, HDL-C, or LDL-C [25, 26, 28, 32-35, 20] following upper-body aerobic exercise, including four with clinically high mean concentrations at baseline. Only two studies reported a significant reduction in any variable. One study [34] reported a 25% reduction (ES: 0.31) in TG in participants with a clinically elevated mean concentrations at baseline (≥1.7 mmol/L). One study reported improvements in HDL-C, LDL-C, TC: HDL-C and TG following 60 mins/week at 70-80% HRR, however the threshold for significance was set at p<0.10 [40]. It therefore appears that upper-body aerobic exercise may not be an adequate stimulus to improve blood lipid profile irrespective of baseline values. This is likely due to the low energy expenditure achieved through upper-body exercise, which appears to drive changes in the lipid profile [95].

Upper-body RT (with or without aerobic exercise) appears to reduce central obesity, with three [42-44] out of four studies reporting a reduction in waist circumference (-

1.0 to -2.6 cm) or waist to hip ratio (-0.02). These changes were accompanied by a decrease in whole-body fat mass and visceral adipose tissue following 120 min/week of training (3 x 10 of 50-70% 1RM, 20 min at 3-6 RPE) [42]. Upper-body RT (with or without aerobic exercise) may elicit improvements in lipid profile, with four [43-45, 40] out of the five retrieved studies reporting a beneficial effect of at least one marker (TC, HDL-C, LDL-C, TC: HDL-C, and TG). However, more studies are needed to determine this, particularly given the high-quality study reporting no change in the lipid profile following 16-weeks of twice-weekly combined training [42].

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Five [50, 54, 58, 60, 62] of the six studies to measure outcomes relating to peripheral insulin sensitivity reported a significant improvement following FES-cycling. The largest of these studies (n=18) [54] reported a significant reduction in glucose and insulin at multiple time-points during a 2-h oral glucose tolerance test following 10 weeks of exercise (2-3 sessions/week, 30 min). However, four of these studies were rated as low quality, and therefore more high-quality studies are needed to confirm if FES-cycling can improve peripheral insulin sensitivity, which upper-body exercise appears unable to achieve. Surprisingly, we identified no RCT's assessing the efficacy of FES-cycling compared to a true control group (i.e. passive cycling or stretching), which should addressed in future research. Four studies reported no change in body mass following FES-hybrid or FES-rowing training. There was a distinct lack of training studies with sufficient breadth of outcomes to make any other meaningful conclusions on the effect of FES-RT, FES-hybrid, FES-rowing and assisted ambulation on CMS risk factors. Nonetheless, given that hybrid training (2 sessions/week, 18-32 min, 65-75%) HRR) [25] improved a multitude of CMS risk factors (waist circumference, android fat percentage, TG, DBP), and that different exercise modalities appear to offer specific benefits to CMS risk factors, other rigorously conducted prospective studies assessing multimodal (e.g.

FES-cycling combined with upper-body aerobic and resistance exercise) interventions_should be conducted in this area of promise.

This review has highlighted the lack of research assessing novel markers of CMS risk, including outcomes relating to inflammation, DEXA/CT derived measured of central adiposity, and endothelial function. It is clear that many studies in the area recruit a convenience sample of relatively active and lean individuals, who are not reflective of the wider, chronic SCI population (i.e. poor metabolic health), which should be considered when interpreting results. For example, individuals with SCI have a significantly lower HDL-C compared to able-bodied controls (1.06 vs 1.28 mmol/L) [96], however only five of the 23 studies to measure HDL-C had a clinically low mean concentration at baseline (<1.03 mmol/L). As is widely acknowledged, this review has also confirmed the existing evidence base of exercise and CMS risk in SCI lacks sufficiently powered (four in total identified), high-quality studies (eight in total identified). However, this review identified 16 additional studies, published since the previous systematic review by van der Scheer and colleagues [11] that were all categorised as fair or high quality, including eight RCT's.

Study Limitations

The major limitation of this systematic review is the use of summary coding to draw conclusions regarding the effect of each exercise modality on specific CMS risk factors. Due to the variability in CMS risk factors measured, exercise modes and training parameters (i.e. exercise intensity and volume), and participant characteristics (i.e. paraplegic vs. tetraplegic), a meta-analysis was not possible. Whilst the coding system provides a useful assessment of the consistency of findings in the field, it uses arbitrary classifications and does not distinguish studies of differing quality. However, when studies rated as 'low-quality' were removed from this analysis (Supplement 3), the conclusions remained unchanged, with the exception of

potential of FES-cycling to improve peripheral insulin sensitivity. Further, given that the vast
majority of included studies lacked sufficient statistical power, there is a risk of a type II error
in the conclusions formed. Finally, this review did not include acute SCI as van der Scheer and
colleagues [11] determined there was an "absence of high-quality, consistent evidence" in this
area, a view which still appears to be true.

CONCLUSIONS

In summary, this systematic review has provided evidence that in adults with chronic SCI, upper-body aerobic exercise improves outcomes relating to central obesity and hepatic insulin sensitivity, but is not sufficient to improve fasting glucose, lipid profiles, or resting blood pressure. Practitioners should consider prescribing moderate-to-vigorous intensity (>75% HR_{MAX}) upper-body aerobic exercise to improve fasting glycaemic control and central obesity. To elicit improvements in lipid profile, this should be combined with upper-body resistance training. More high-quality randomised controlled trials assessing novel markers of CMS and responses to combined exercise interventions (e.g. aerobic exercise with resistance training), high-intensity exercise interventions, and FES-based exercise are needed to inform and refine evidence-based exercise guidelines for the prevention and management of CMS in this population.

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Figure 1. PRISMA flow diagram

Central	Body Mass Index (BMI)	Formatted: Pattern: Clear (Yellow)
Adiposity/Obesity	Body Mass (BM)	
	Waist Circumference (Waist)	
	Hip Circumference Waist to Hip Ratio (WHR)	Farmattada Dattara Class (Valland)
	Body Fat Percentage (BF%) (assessed via DEXA/CT) Fat Mass (FM) (assessed via DEXA/CT)	Formatted: Pattern: Clear (Yellow)
	Android Fat Mass	
	Visceral Adipose Tissue (VAT)	
	Liver Fat Content	
	Leptin	
Glycaemic Control	Fasting insulin and glucose	
	Glucose to insulin ratio Fasting proinsulin	
	Glycosylated haemoglobin (HbA1c)	
	Fasting/postprandial insulin sensitivity measures	
	C-peptide	
Dyslipidaemia	Triglycerides (TG)	Formatted: Pattern: Clear (Yellow)
	Low-density lipoprotein-cholesterol (LDL-C)	
	High-density lipoprotein-cholesterol (HDL-C) Total cholesterol (TC)	
	DL, HDL, TC, TC: HDL-C	
	Non-esterified fatty acids (NEFA)	
	Free-fatty acids (FFA)	
	Apolipoprotein B	
Inflammation	C-reactive Protein (CRP)	
	Interleukin-6 (IL-6)	
	Tumour necrosis factor-alpha (TNF-α) Adiponectin	
	•	
Vascular Dysregulation	Systolic Blood Pressure (SBP) Diastolic Blood Pressure (DBP)	Formatted: Pattern: Clear (Yellow)
	Pulse wave velocity (PWV)	
	Flow-mediated dilation (FMD)	
	Microalbuminuria	
Thrombotic State	Fibrinogen	
	Plasminogen activator inhibitor-1 (PAI-1)	
hrombotic State		_

Table 2. Summary coding of studies examining the effect of exercise on CMS outcome measures.

		Aerobic	Aerobic + RT	Ambulation	Hybrid and Rowing	FES-cycling	FES- RT/Combined
	BM	1/9 (11%)	1/2 (50%)	1/3 (33%)	0/5 (0%)	1/4 (25%)	0/4 (0%)
	BMI	1/4 (25%)	1/4 (25%)	1/1 (100%)	0/1 (0%)	0/2 (0%)	1/3 (33%)*
	Waist	4/6 (66%)	2/3 (67%)	=	1/2 (50%)	=	-
	WHR	-	1/1 (100%)	=	-	=	-
Central	BF%	0/2 (0%)	-	2/2 (100%)	0/2 (0%)	1/2 (50%)	0/2 (0%)
Adiposity/Obesity	FM	0/3 (0%)	1/2 (50%)	0/2 (0%)	-	1/2 (50%)	0/2 (0%)
	Android FM	0/1 (0%)	-	-	0/1 (0%)	-	-
	Abdominal AT	-	-	-		0/1 (0%)	-
	VAT	0/1 (0%)	1/1 (100%)	-		-	0/2 (0%)
	Leptin	1/1 (100%)	0/1 (0%)	-	1/1 (100%)	-	-
	CRP	0/1 (0%)		1/1 (100%)	0/1 (0%)	1/2 (50%)	0/1 (0%)
Inflammation	IL-6	1/2 (50%)	0/1 (0%)	-	0/1 (0%)	1/2 (50%)	0/1 (0%)
	TNF-α	1/1 (100%)	0/1 (0%)	-	-	1/2 (50%)	0/1 (0%)
	Adiponectin	0/1 (0%)	0/1 (0%)	-	-	-	1/1 (100%)
	TG	1/6 (17%)	2/4 (50%)	0/2 (0%)	1/1 (100%)	1/3 (33%)	1/3 (33%)
	FFA	-	-	-	-	0/1 (0%)	0/1 (0%)
	NEFA	0/1 (0%)	-	-	-	-	-
Dyslipidaemia	TC	1/6 (17%)	2/5 (40%)	1/2 (50%)	0/1 (0%)	0/2 (0%)	1/3 (33%)
	HDL-C	0/7 (0%)	1/5 (20%)	0/2 (0%)	0/2 (0%)	1/3 (33%)	1/3 (33%)
	LDL-C	0/5 (0%)	2/5 (40%)	1/2 (50%)	0/1 (0%)	1/3 (33%)	0/3 (0%)
	TC: HDL-C	0/1 (0%)	1/2 (50%)	1/1 (100%)	-	1/1 (100%)	1/2 (50%)
	Fasting Glucose	0/8 (0%)	0/3 (0%)	0/1 (0%)	1/2 (50%)	0/1 (0%)	0/2 (0%)
	Fasting Insulin	4/5 (80%)	1/3 (33%)	-	0/2 (0%)	0/3 (0%)	0/1 (0%)
	HbA1c	0/1 (0%)	0/1 (0%)	-	-	-	-
	HOMA-IR	4/4 (100%)	2/2 (100%)	=	0/2 (0%)	=	0/2 (0%)
	HOMA-%S	1/1 (100%)	-	=	-	=	0/1 (0%)
Glycaemic Control	НОМА-%β	0/2 (0%)	-	=	-	=	0/1 (0%)
	ISI-Matsuda	0/2 (0%)	-	-	-	-	-
	Glucose OGTT	0/2 (0%)	-	1/1 (100%)	0/1 (0%)	2/3 (66%)	0/3 (0%)
	Insulin OGTT	0/2 (0%)	-	1/1 (100%)	-	1/3 (33%)	0/2 (0%)
	IVGTT Si	0/1 (0%)	-	-	-	0/2 (0%)	0/1 (0%)
	Cederholm Index	-	-	-	-	1/1 (100%)	-
	HEC Si	-	-	-	-	1/1 (100%)	-
	HEC Glucose	_	-	-	-	1/1 (100%)	-

Thrombotic State	PAI-1	1/2 (50%)	0/1 (0%)	-	-	-	-
	Fibrinogen	0/1 (0%)	-	-	-	0/1 (0%)	-
	SBP	1/9 (11%)	0/3 (0%)	0/3 (0%)	0/2 (0%)	1/4 (25%)	0/1 (0%)
	DBP	0/9 (0%)	0/3 (0%)	0/3 (0%)	1/2 (50%)	1/3 (33%)	0/1 (0%)
Vascular	FMD	=	0/1 (0%)	-	1/2 (50%)	-	1/1 (100%)
Dysregulation	PWV	-	0/1 (0%)	=	-	0/1 (0%)	=
	Albumin	-	-	-	-	-	0/1 (0%)

Red: 0-33% of studies reported significant differences; yellow: 34-59% of studies reported significance differences; green: 60-100% of studies demonstrated positive significance differences, bold writing: ≥4 studies demonstrate the same effect. *one study reported a significant increase in BMI. NA; not applicable

HOMA-IR; homeostatic model assessment insulin resistance, HOMA-%S; insulin sensitivity; HOMA-%β; beta cell function, ISI-Matsuda; insulin sensitivity index-Matsuda. OGTT; oral glucose tolerance test, IVGTT Si; intravenous glucose tolerance test insulin sensitivity, HEC Si; hypereuglycaemic clamp insulin sensitivity.

Table 3. Detailed findings from voluntary upper-body aerobic exercise studies included in this review.

Study Design D&B Quality	n	Intervention	CMS Outcome	Group Baseline Intervention (Control) Mean ± SD	Change Intervention (Control)	p value*	ES
[25]	10	Hand-cycle	Waist (cm)	89.7 ± 3.5	-2.5	0.03	0.75
Pre-post†		16 weeks	Android Fat Mass (kg)	2.6 ± 0.4	0.0	0.85	0.00
20		2 sessions/week	Android Fat (%)	38.6 ± 3.7	-1.3	0.26	0.40
High		65-75% HRR	TG (mmol/L)	1.2 ± 0.2	-0.1	0.67	0.63
6		18-32 mins	HDL-C (mmol/L)	1.4 ± 0.2	0.0	0.94	0.00
		10 32 111113	Fasting Glucose (mmol/L)	5.3 ± 0.2	-0.2	0.30	1.00
			Fasting Insulin (pmol/L)	54.6 ± 8.5	-14.3	0.01	1.78
			HOMA-IR	1.9 ± 0.3	-0.5	0.02	2.35
			SBP (mmHg)	119 ± 4	+4	0.30	1.13
			DBP (mmHg)	72 ± 3	-3	0.34	0.57
			CRP (mg/L)	2.86 ± 1.36	-0.39	0.34	0.37
20.63	2.1	A GE	IL-6 (pg/mL)	2.40 ± 0.57	-0.64	0.10	0.56
[26]	21	ACE	Body Mass (kg)	$76.8 \pm 13.3 \ (76.8 \pm 11.3)$	-1.1 (-0.7)	NS	-
RCT		6 weeks	Fat Mass (kg)	$27.6 \pm 10.0 \ (25.5 \pm 6.6)$	-0.6 (0.0)	NS	-
19		4 sessions/week	VAT (cm ²)	$181 \pm 85 \ (186 \pm 47)$	-22 (-3)	NS	-
High		60-65% VO _{2PEAK}	TG (mmol/L)	$1.2 \pm 0.5 \ (1.3 \pm 0.5)$	-0.1 (+0.5)	NS	1.02
		45 mins	TC (mmol/L)	$4.9 \pm 1.0 \ (5.1 \pm 0.9)$	-0.1 (+0.1)	NS	0.17
			HDL-C (mmol/L)	$1.1 \pm 0.3 \ (1.0 \pm 0.2)$	+0.1 (0.0)	NS	0.07
			LDL-C (mmol/L)	$3.2 \pm 0.9 \ (3.5 \pm 0.8)$	0.0 (-0.2)	NS	0.05
			NEFA (mmol/L)	$0.6 \pm 0.3 \ (0.7 \pm 0.6)$	+0.3 (-0.1)	NS	0.40
			Fasting Glucose (mmol/L)	$5.3 \pm 0.5 (5.7 \pm 1.3)$	0.0 (0.0)	NS	_
			Fasting Insulin (pmol/L)	$54.8 \pm 30.1 (41.3 \pm 18.1)$	-12.7 (+3.1)	0.03	0.54
			HOMA2-IR	$1.03 \pm 0.57 \ (0.80 \pm 0.35)$	-0.24 (+0.06)	0.04	0.49
			HOMA2-%ß (%)	87 ± 31 (66 ± 23)	-14 (+1)	NS	0.58
			ISI-Matsuda	$4.8 \pm 2.2 \ (6.4 \pm 3.1)$	+0.3 (-0.7)	NS NS	0.56
				$4.8 \pm 2.2 \ (0.4 \pm 3.1)$		NS NS	-
			Glucose OGTT (%)	-	+8 (-9)		-
			Insulin OGTT (%)	-	-8 (+6)	NS	-
			SBP (mmHg)	$128 \pm 23 \ (128 \pm 15)$	-3 (-2)	NS	-
			DBP (mmHg)	$77 \pm 15 (81 \pm 13)$	-1 (-4)	NS	-
[27]	17	ACE	BMI (kg/m²)	$27.6 \pm 4.1 \ (27.8 \pm 4.4)$	-0.2 (NR)	0.72	-
RCT		12 weeks	Waist (cm)	$98.1 \pm 6.6 (98.4 \pm 6.7)$	-3.7 (NR)	0.05	-
19		3 sessions/week	Leptin (ng/mL)	$9.6 \pm 2.7 \ (9.8 \pm 2.8)$	-2.1 (+0.1)	< 0.05	0.71
High		50-65% HRR	PAI-1 (ng/mL)	$29.8 \pm 6.2 \ (30.2 \pm 6.1)$	-0.7 (-0.1)	NS	0.09
		20-30 mins	IL-6 (pg/mL)	$6.7 \pm 2.2 \ (6.9 \pm 2.3)$	-2.6 (+0.1)	< 0.05	1.08
			TNF-α (pg/mL)	$23.3 \pm 5.6 (23.6 \pm 5.5)$	-2.7 (-0.1)	< 0.05	0.47
			Adiponectin (ng/mL)	$18.8 \pm 4.1 \ (18.5 \pm 4.2)$	+0.6 (+0.1)	NS	0.11
28]	10	ACE	BF (%)	34.9 ± 34.9	0.0	0.35	0.01
Pre-post		10 weeks	Fat Mass (kg)	25.1 ± 11.9	-0.3	0.75	0.02
7		3 sessions/week	TC (mmol/L)	4.50 ± 0.58	+0.04	0.75	0.08
air		70% VO _{2PEAK}	HDL-C (mmol/L)	0.94 ± 0.16	-0.06	0.07	0.22
an		30 mins	LDL-C (mmol/L)	2.71 ± 0.39	+0.31	0.12	0.72
		50 milis	Fasting Glucose (mmol/L)	5.54 ± 0.82	-0.05	0.12	0.72
			Fasting Glucose (fillio/L) Fasting Insulin (pmol/L)	3.34 ± 0.82 84.9 ± 38.8	-31.8	0.92	1.07
			Glucose: Insulin	9.77 ± 4.49	+3.92	0.03	1.00
			Glucose OGTT (AUC)	-	+6%	0.25	0.29
			Insulin OGTT (AUC)	-	+5%	0.92	0.13
			HOMA-IR	1.6 ± 0.7	-0.6	0.05	1.11
			HOMA-%ß (%)	111.4 ± 48.7	-29.0	0.12	0.78
			HOMA%S (%)	73.3 ± 31.6	+32.3	0.05	1.10
			ISI-Matsuda	3.4 ± 1.6	+0.2	0.35	0.16
29]	5	ACE	Body Mass (kg)	65.6 ± 6.6	+2.3	0.18	0.33
re-post		12 weeks	BMI (kg/m ²)	23.5 ± 3.4	+0.8	0.18	0.22
.7		3 sessions/week	SBP (mmHg)	110 ± 25	+1	0.13	0.04
air		Anaerobic	DBP (mmHg)	66 ± 12	+2	0.80	0.11
		Threshold 30 mins					
[30]	14	ACE	Body Mass (kg)	69.2	-2	NS	
	14		Dody Mass (kg)	09.2	- <u>∠</u>	149	1 -
Pre-post		10 weeks					
17 ∃air		3 sessions/week 25-35 mins					
		1 15 75 manna	1		i .		

		60% W _{PEAK}					
[31]	4	ACE	Body Mass (kg)	80 ± 12	0	NS	0.00
Pre-post†	1	16 weeks	BMI (kg/m ²)	28 ± 4	0	NS	0.00
16 post-		5 sessions/week	BF (%)	40 ± 3.7	-2	NS	0.52
Fair		75% HR _{MAX}	Fat Mass (kg)	31 ± 7	-2	NS	0.32
ran				5.27 ± 0.50	-0.06		0.31
		40 mins	Fasting Glucose (mmol/L)			0.9	
			Fasting Insulin (pmol/L)	76.4 ± 62.5	-23.6	NS	0.41
			IVGTT Insulin Sensitivity	-	+62.5%	NS	0.64
			IVGTT Glucose Effectiveness	-	+35%	NS	0.70
			SBP (mmHg)	119 ± 13	-1	NS	0.08
			DBP (mmHg)	75 ± 5	+2	NS	0.36
[32]	33	ACE	Waist (cm)	86.5 (94.5)	+4.75 (+1.5)	NS	-
RCT		12 weeks	TG (mmol/L)	1.50 (1.38)	+0.06 (+0.29)	NS	_
16		3 sessions/week	TC (mmol/L)	4.57 (4.60)	+0.26 (+0.05)	NS	_
Fair		50-70% VO _{2PEAK}	HDL-C (mmol/L)	0.96 (1.05)	0.0 (+0.14)	NS	_
1 411		30 mins	LDL-C (mmol/L)	2.87 (2.91)	0.0 (0.09)	NS	_
		30 mms	Fasting Glucose (mmol/L)	4.44 (4.47)	-0.19 (+0.14)	NS	
					0 (0)	NS NS	-
			SBP (mmHg)	100 (100)			-
			DBP (mmHg)	60 (60)	0 (0)	NS	-
[33]	16	Hand-cycle	BMI (kg/m ²)	$22.0 \pm 3.7 \ (20.8 \pm 2.7)$	-0.2 (+0.3)	<0.01	1.58
RCT		6 weeks	Waist (cm)	$88.3 \pm 13.1 \ (81.7 \pm 9.0)$	-2.6 (+0.8)	< 0.01	2.67
15		3 sessions/week	TG (mmol/L)	$1.16 \pm 0.47 \ (1.09 \pm 0.56)$	-0.01 (-0.12)	0.95	0.25
Fair		70-80% HR _{PEAK}	TC (mmol/L)	$4.56 \pm 0.92 \ (4.73 \pm 0.55)$	+0.03 (-0.09)	0.81	0.25
		44 mins	HDL-C (mmol/L)	$1.10 \pm 0.30 (1.17 \pm 0.18)$	+0.09 (-0.01)	0.29	0.82
			LDL-C (mmol/L)	$2.93 \pm 0.67 \ (3.07 \pm 0.62)$	-0.06 (-0.03)	0.99	0.09
			Fasting Glucose (mmol/L)	$4.36 \pm 0.46 \ (4.92 \pm 0.60)$	-0.09 (+0.04)	0.32	0.39
			Fasting Insulin (pmol/L)	$37.5 \pm 16.7 (34.0 \pm 20.1)$	-13.9 (+11.8)	<0.01	1.57
			HOMA-IR		-0.4 (0.4)	<0.01	1.40
F2.41	9	ACE		$1.0 \pm 0.6 \ (1.1 \pm 0.8)$		_	
[34]	9	ACE	Body Mass (kg)	61.0 ± 7.0	-1.9	<0.05	0.26
Pre-post		10 weeks	Waist (cm)	85.5 ± 6.2	-1.9	< 0.05	0.26
14		4 sessions/week	TG (mmol/L)	1.74 ± 0.78	-0.43	< 0.05	0.31
Fair		50-70% HRR	TC (mmol/L)	5.25 ± 0.88	-0.18	NS	0.14
		60 mins	HDL-C (mmol/L)	1.45 ± 0.18	+0.05	NS	0.20
			LDL-C (mmol/L)	2.95 ± 0.62	-0.10	NS	0.15
			Fasting Glucose (mmol/L)	5.66 ± 1.39	-0.17	NS	0.10
			HbA1c (%)	4.9 ± 0.6	-0.10	NS	0.14
			PAI-1 (g/L)	5.2 ± 1.1	-1.4	< 0.05	1.22
			Fibrinogen (g/L)	2.97 ± 5.7	-0.7	NS	0.14
			SBP (mmHg)	136 ± 5	-3	<0.05	0.66
					-3	NS	0.30
50.51	10	WICE	DBP (mmHg)	75 ± 8			_
[35]	12	WCE	Body Mass (kg)	74 ± 10	+2.0	NS	0.20
Pre-post		10 weeks	TG (mmol/L)	1.32 ± 0.59	-0.08	NS	0.12
14		2-3 sessions/week		4.78 ± 1.09	-0.39	0.04	0.40
Fair		Intensity NR	HDL-C (mmol/L)	1.24 ± 0.26	0.0	NS	0.00
		20-30 mins	TC: HDL-C	4 ± 1	-0.2	NS	0.20
			Fasting Glucose (mmol/L)	4.77 ± 1.94	-1.0	NS	0.03
			SBP (mmHg)	124 ± 10	0	NS	0.00
			DBP (mmHg)	85 ± 7	-3	NS	0.35
[36]	12	WCT	Body Mass (kg)	41.8 ± 5.8	0.0	NS	0.00
Pre-post	12	12 weeks	Dody Mass (Ng)	71.0 ± 3.0	0.0	140	0.00
14		14 sessions/week					
Fair	1	60-70% HR _{PEAK}			 	 	0.5-
[37]	9	WCT	Body Mass (kg)	82.1 ± 14.6	+1.2	NS	0.09
Pre-post		7 weeks	Waist (cm)	109.6 ± 12.2	+4.1	NS	0.28
13		5 sessions/week					
Low		Intensity NR					
		Duration NR					
[38]	11	WCE	SBP (mmHg)	126 ± 12	-2	NS	0.16
	11	5 weeks	DBP (mmHg)	82 ± 6	-2	NS NS	0.10
	1	2 sessions/week	DDI (IIIIIIIg)	02 ± 0	- <u>-</u>	140	0.29
Pre-post		⊥ ∠ sessions/week					
12							
		<80% HRPEAK					
12 Low		<80% HR _{PEAK} 30 mins					
12 Low	14	<80% HR _{PEAK} 30 mins ACE	SBP (mmHg)	122 ± 5 (114 ± 6)	+4 (+18)	NS	-
12 Low	14	<80% HR _{PEAK} 30 mins	SBP (mmHg) DBP (mmHg)	122 ± 5 (114 ± 6) 78 ± 5 (81 ± 4)	+4 (+18) -2 (+6)	NS NS	

controlled trial		50 or 70% VO _{2PEAK} 20 or 40 mins					
Low		20 01 10 111115					
[40] Pre-post 11 Low	11	WCE 8 weeks 3 sessions/week 70-80% HRR (or 50-60% HRR) 20 mins	TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) TC: HDL-C	$ \begin{array}{c} 1.08 \pm 0.32 \ (0.88 \pm 0.26) \\ 5.04 \pm 0.91 \ (4.81 \pm 0.70) \\ 1.01 \pm 0.28 \ (1.27 \pm 0.28) \\ 3.54 \pm 0.67 \ (3.15 \pm 0.44) \\ 5 \pm 0.9 \ (4 \pm 0.7) \end{array} $	-0.20 (-0.04) -0.41 (+0.16) +0.21 (-0.18) -0.54 (0.16) -1 (+1)	<0.1 (NS) NS (NS) <0.1 (NS) <0.1 (NS) <0.1 (NS)	0.76 (0.15) 0.63 (0.28) 0.83 (0.46) 1.12 (0.37) 1.37 (0.67)

Red font clinically high group average, bold font significant difference following intervention reported, ES effect size.

ACE arm-crank ergometry, WCE wheelchair ergometer, WCT wheelchair treadmill ergometry, HRR heart rate reserve, VO2PEAK peak oxygen uptake, WPEAK peak power output, HRPEAK peak heart rate, HRMAX age-predicted maximum heart rate, BF body fat, HOMA-IR homeostatic model assessment of insulin resistance, OGTT oral glucose tolerance test, AUC area under the curve, IVGTT intravenous glucose tolerance test, NS non-significant, NR not reported

*Group x time interaction for RCT and non-randomised controlled trial, or pre-post change for pre-post study designs.

† True study design is RCT, presented as pre-post due to two different exercise modalities being tested.

Table 4. Detailed findings from upper-body RT (with or without aerobic training) studies included in this review.

Study Design D&B Quality	n	Intervention	CMS Outcome	Group Baseline Intervention (Control) Mean ± SD	Change Intervention (Control)	p value *	ES
[41] Pre-post† 23 High	17	16 weeks 3 sessions/week RT: 20-25 mins, 2- 3 sets at 12-15 repetition max resistance Aerobic: 20-25 mins, 3-5 RPE	Fat Mass (kg)	23.2 ± 10.8	-0.2	NS	0.02
[42] RCT 19 High	23	16 weeks 2 sessions/week RT: 3 x 10, 50-70% 1RM Aerobic: >20 mins, 3-6 RPE	Body Mass (kg) BMI (kg/m²) Waist (cm) Fat Mass (kg) VAT (kg) Leptin (ng/mL) TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) TC: HDL-C Fasting Insulin (pmol/L) HbA1c (mmol/L) PAI-1 (ng/mL) SBP (mmHg) DBP (mmHg) Brachial FMD Femoral FMD PWV - Central IL-6 (pg/mL) TNF-\(\alpha\) (pg/mL) Adiponectin (\(\pmu\)	83.4 ± 18.9 (78.6 ± 15.7) 27.3 ± 5.2 (25.7 ± 4.9) 96.2 ± 14.9 (89.6 ± 11.7) - (-) - (-) 10.12 ± 13.25 (10.2 ± 12.8) 1.3 ± 0.6 (1.1 ± 0.7) 4.5 ± 0.9 (4.1 ± 0.9) 1.01 ± 0.2 (1.13 ± 0.2) 2.9 ± 0.9 (2.5 ± 0.7) 4.6 ± 0.9 (3.8 ± 1.1) 39.2 ± 29.5 (68.2 ± 77.9) 1.01 ± 0.2 (1.13 ± 0.3) 30.4 ± 17.7 (31.1 ± 22.7) 116 ± 18 (118 ± 18) 68 ± 9 (74 ± 13) - - 2.5 ± 2.2 (3.7 ± 2.1) 4.7 ± 1.8 (4.1 ± 2.2) 76.7 ± 64.0 (82.02 ± 38.28)	↓ -0.3 (+0.9) -1.0 (+3.5) ↓ ↓ +1.0 (+4.1) +0.1 (-0.1) -0.2 (0.0) 0.0 (+0.04) -0.2 (-0.1) -0.2 (-0.2) +9.5 (+10.3) +0.9 (-0.2) +11.6 (+15.5) 0 (-2) -1 (-2) - -1.0 (+1.8) -0.3 (-0.1) +13.4 (+35.67)	0.03 0.02 0.03 0.04 0.04 NS NS NS NS NS NS NS NS NS NS NS NS NS	1.07 1.14 1.02 1.00 1.02 - - - - - - - - - - - - -
[43] RCT 17 Fair	20	8 weeks 3 sessions/week RT: 60-80% 1RM, 5 exercises.	BMI (kg/m²) Waist: Hip TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) Fasting Glucose (mmol/L) Fasting Insulin (pmol/L) HOMA-IR	$25.3 \pm 1.4 (24.9 \pm 1.0)$ $0.83 \pm 0.02 (0.83 \pm 0.14)$ $1.77 \pm 0.07 (1.80 \pm 0.11)$ $4.66 \pm 0.18 (4.78 \pm 0.10)$ $1.12 \pm 0.06 (1.15 \pm 0.11)$ $2.81 \pm 0.10 (2.82 \pm 0.12)$ $5.46 \pm 1.34 (5.45 \pm 1.42)$ $110.6 \pm 19.5 (116.7 \pm 24.9)$ $6.92 \pm 1.27 (7.27 \pm 2.09)$	-0.6 (+0.2) -0.02 (+0.01) -0.27 (+0.02) -0.38 (+0.04) +0.12 (+0.01) -0.12 (+0.05) -0.38 (-0.01) -2.4 (-3.5) -0.62 (-0.25)	NS 0.03 0.001 0.001 NS 0.001 NS NS 0.03	
[44] RCT 17 Fair	17	6 weeks 3 sessions/week RT: 1-3 x 10-20 Aerobic: 10-20 mins, 4-8 RPE or 65-85% HRMAX	BMI (kg/m²) Waist (cm) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) Fasting Glucose (mmol/L) Fasting Insulin (pmol/L) HOMA-IR	$21.8 \pm 2.9 (20.8 \pm 1.9)$ $84.1 \pm 11.9 (79.4 \pm 6.6)$ $4.20 \pm 0.88 (1.96 \pm 0.09)$ $1.26 \pm 0.55 (1.32 \pm 0.27)$ $2.42 \pm 0.81 (3.25 \pm 0.76)$ $4.50 \pm 0.30 (4.20 \pm 0.20)$ $52.1 \pm 32.6 (20.1 \pm 7.6)$ $1.5 \pm 1.0 (0.5 \pm 0.2)$	-0.4 (-0.1) -2.6 (-0.2) -0.04 (+0.05) +0.14 (-0.04) -0.12 (+0.36) -0.09 (+0.10) -20.1 (+2.1) -0.6 (+0.06)	0.08 0.02 0.46 0.05 0.12 0.23 0.05 0.05	1.17 1.94 0.40 1.24 0.85 0.62 1.24 1.33
[45] Pre-post 15 Fair	16	12 weeks 3 sessions/week RT: 2 x 8 to 3 x 12. Aerobic: 60-75% HRR 20-60 mins	Body Mass (kg) BMI (kg/m²) Waist (cm) TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) Fasting Glucose (mmol/L) SBP (mmHg) DBP (mmHg)	74.9 ± 7.2 26.0 ± 2.6 104.1 ± 7.9 1.41 ± 0.93 5.66 ± 1.32 1.26 ± 0.40 4.20 ± 1.15 5.81 ± 0.05 118 ± 20 80 ± 11	-2.9 -1.0 +1.3 - 0.30 - 0.68 +0.02 -0.19 -0.74 -5	NS NS NS <0.05 <0.05 NS NS NS NS	1.19 0.33 0.17 0.35 0.54 0.05 0.17 1.64 0.26 0.27
[46] RCT	34	36 weeks 2 sessions/week	SBP (mmHg)* DBP (mmHg)*	125 ± 23 (133 ± 20) 72 ± 16 (85 ± 14)	+2 (-2) +3 (-4)	NS NS	-

15 Fair		RT: 70-80% 1RM, Aerobic: 15-30 mins, 70% HR _{MAX} or 3-4 RPE.	*Paraplegics only				
[47]	5	12 weeks	TG (mmol/L)	2.29 ± 1.35	-0.14	0.63	0.12
Pre-post		3 sessions/week	TC (mmol/L)	4.73 ± 0.67	-0.42	0.20	0.56
12		Circuit Training:	HDL-C (mmol/L)	1.05 ± 0.14	+0.11	0.10	0.49
Low		50-60% 1RM	LDL-C (mmol/L)	3.06 ± 0.57	-0.79	0.05	1.17
		40-45 mins	TC: HDL-C	5.0 ± 1.1	-1.1	0.05	1.19

1RM *one-rep maximum*, RPE *rating of perceived exertion*. *Group x time interaction for RCT and non-randomised controlled trial, or pre-post change for pre-post study designs. †True study design is RCT, presented as pre-post due to two different exercise modalities being tested

 Table 5. Detailed findings of FES-cycling studies included in this review.

Study Design D&B Quality	n		CMS Outcome	Group Baseline Intervention (Control) Mean ± SD	Change Intervention (Control)	p value *	ES
[48] Pre-post 16 Fair	1 0	FES-cycling 12 weeks 3 sessions/week 90-95% of max tolerance 1-45 mins	TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) CRP (pg/mL) IL-6 (pg/mL) TNF-α (pg/mL)	0.37 ± 0.19 1.99 ± 0.46 0.48 ± 0.13 1.13 ± 0.33 12.59 ± 14.06 6.29 ± 4.65 25.62 ± 49.64	-0.01 +0.07 0.0 +0.07 -5.81 +0.61 +4.27	NS NS NS NS NS NS	0.06 0.15 0.00 0.22 0.55 0.13 0.07
[49] Retrospective cohort study 16 Fair	5	FES-cycling 3-168 weeks 3 sessions/week Intensity NR 45-60 mins	TG HDL-C LDL-C TC: HDL-C	NR NR NR 4.1 ± 1.0 (5.3 ± 1.9)	- - -	<0.05 NS <0.05 0.03	- - - 0.79
[31]† Pre-post 16 Fair	9	FES-cycling 16 weeks 5 sessions/week 75% HR _{MAX} 40 mins	Body Mass (kg) BMI (kg/m²) BF (%) Fat Mass (kg) Fasting Glucose (mmol/L) Fasting Insulin (pmol/L) IVGTT Insulin Sensitivity (%) IVGTT Glucose Effectiveness (%) SBP (mmHg) DBP (mmHg)	79 ± 12 26 ± 5 38 ± 5.7 29 ± 8.6 5.00 ± 0.11 97.2 ± 118.1 123 ± 8 79 ± 5	+6 +3 0 0 +0.33 -59.0 +129 +4 +4	NS NS NS NS 0.4 0.8 NS NS >0.5 >0.5	0.59 0.82 0.00 0.00 0.65 0.70 0.69 0.19 0.44 0.36
[50] Pre-post 14 Fair	7	FES-cycling 8 weeks 3 sessions/week Max load to finish 30 min 30 min	2-h Glucose OGTT (mmol/L) 2-h Insulin OGTT (pmol/L)	7.77 ± 0.89 822 ± 296	-0.98 -215	0.01 NS	2.13 1.00
[51] Pre-post 14 Fair	9	FES-cycling 6 weeks 3 sessions/week Max load to finish 30 min 30 min	SBP (mmHg)	131 ± 20	+6	NS	0.40
[52] Pre-post 14 Fair	1 8	FES-cycling 8 weeks 3 sessions/week Intensity NR 30 mins	Body Mass (kg) BMI (kg/m²)	73.8 ± 13.9 25.4 ± 3.9	+1.2 +0.3	0.06 NS	0.09 0.08
[53] Pre-post 13 Low	1 3	FES-cycling 12 weeks 3 sessions/week Max load to finish 30 min 30 min	SBP (mmHg) DBP (mmHg) *paraplegics only	-	<u></u>	<0.05 <0.05	-
[54] Pre-post 13 Low	1 8	FES-cycling 10 weeks 2-3 sessions/week Max load to finish 30 min or fatigue	Body Mass (kg) Fat Mass (kg) TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) 2-h Glucose OGTT 2-h Insulin OGTT CRP IL-6 TNF-α	69.6 ± 4.2 22.9 ± 2.3 1.18 ± 0.30 4.08 ± 0.16 0.88 ± 0.05 2.65 ± 0.16 $-$ 15.92 ± 1.57 4.91 ± 1.10 11.82 ± 0.63	-2.1 +0.6 -0.04 -0.04 -0.10 +0.07 ↓ -2.98 -1.12 -0.51	<0.05 <0.05 NS NS <0.05 NS <0.05 <0.05 <0.05 <0.05	0.12 0.06 0.04 0.06 0.43 0.12 - 0.57 0.31 0.19
[55] Pre-post 13 Low	8	FES-cycling 6 weeks 3 sessions/week Intensity NR 30 mins	SBP (mmHg) DBP (mmHg)	112 ± 6 77 ± 4	-3 -4	NS NS	0.63 1.00

		T	Γ		T		T
[56]	5		BF (%)	29.7 ± 2.6	-1.9	<0.05	0.80
Pre-post		8 weeks	Fasting Insulin	NR	NR	NS	-
12		7 sessions/week					
Low		Max load to finish 30					
		min					
		30 mins					
[57]	1	FES-cycling	Fibrinogen (mg/dL)	410 ± 78	+29	NS	0.17
Pre-post	2						
12		2 sessions/week					
Low		Intensity NR					
		30 mins					
[58]	5	FES-cycling	HEC Glucose Uptake (%)	-	+33	< 0.05	0.95
Pre-post		8 weeks	2				
11		7 sessions/week					
Low		Max load to finish 30					
		min					
		30 mins					
[59]	8	FES-cycling	Hyperaemic Flow	_	\leftrightarrow	NS	-
Pre-post		8 weeks					
11 1		2-3 sessions/week					
Low		Max load to finish 30					
		min					
		30 mins					
[60]	1	FES-cycling	FFA (mmol/L)	0.68 ± 0.08	-0.03	NS	0.13
Pre-post	0		Fasting Insulin (pmol/L)	83 ± 35	-28	NS	0.33
11		3 sessions/week	Glucose OGTT (AUC)	_	\leftrightarrow	NS	-
Low		Intensity NR	Insulin OGTT (AUC)	_	\leftrightarrow	NS	-
		30 mins	HEC SSGIR Step 1 (%)	_	+28	< 0.05	0.74
			HEC SSGIR Step 2 (%)	_	+17	NS	0.63
[61]	1	FES-cycling	Body Mass	NR	\leftrightarrow	NS	-
Pre-post	5		Abdominal Adipose Tissue	NR	\leftrightarrow	NS	-
10		3 sessions/week	1				
Low		Max load to finish 30					
		min					
		30 mins					
[62]	5		Cederholm Index	-	↑	< 0.05	-
Pre-post		8 weeks			'		
9		3 sessions/week					
Low		Intensity NR					
		30 mins					
			I .	L	L		l

^{*}Group x time interaction for RCT and non-randomised controlled trial, or pre-post change for pre-post study designs. \dagger True study design is RCT, presented as pre-post due to two different interventions (vs. high-protein diet).

Table 6. Detailed findings of FES-RT and combined (FES-cycling and FES-RT) studies included in this review.

Study Design D&B Quality	n	Intervention	CMS Outcome	Group Baseline Intervention (Control) Mean ± SD	Change Intervention (Control)	p value *	ES
[63] RCT 21 High	22	FES-knee extensions (with testosterone replacement therapy) 16 weeks 2 sessions/week 4 x 10 ~1 kg increments every 2 sessions	Body Mass (kg) BMI (kg/m²) BF (%) Fat Mass (kg) VAT (cm²) TG FFA TC HDL-C LDL-C IVGTT Insulin Sensitivity (%) IVGTT Glucose Effectiveness (%) CRP IL-6 (pg/mL) TNF-α Adiponectin (ng/mL)	$80.5 \pm 16 (77.5 \pm 9.0)$ $25 \pm 4.5 (24.4 \pm 3.6)$ $32 \pm 11 (33.4 \pm 9)$ $26.7 \pm 12.5 (26.1 \pm 8.0)$ $101 \pm 71 (91.5 \pm 49.5)$ NR NR NR NR NR NR NR NR NR NR 5.5 ± 5.6 (5.9 ± 6.0) NR	+2.6 (+0.2) +1.6 (-0.4) -1.3 (-1.4) 0.0 (-1.0) -13 (-7.0) \leftrightarrow \leftrightarrow \leftrightarrow \leftrightarrow 0.0 (0.0) 31.5 (28.6) \leftrightarrow -2.6 (-2.0) \leftrightarrow -624 (+1291)	NS 0.004 NS NS NS NS NS NS NS NS NS NS	-
[64] RCT 16 Fair	9	FES knee-extensions 12 weeks 2 sessions/week 4 x 10 Increased by ~1kg every 2 sessions	Body Mass (kg) BMI (kg/m²) BF (%) Fat Mass (kg) Trunk VAT CSA (cm²) TG (mmol/L) FFA (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) TC: HDL-C HOMA-IR (Log10) Glucose OGTT (AUC) (%) Insulin OGTT (AUC) (%)	$74 \pm 14 \ (76 \pm 8)$ $21 \pm 5 \ (23 \pm 3)$ $30 \pm 8 \ (29 \pm 3)$ $23.3 \pm 9 \ (22 \pm 2)$ $103 \pm 80 \ (106 \pm 32)$ $1.58 \pm 1.38 \ (1.25 \pm 0.28)$ $0.58 \pm 0.1 \ (0.53 \pm 0.1)$ $4.19 \pm 1.27 \ (3.93 \pm 0.70)$ $0.78 \pm 0.08 \ (0.83 \pm 0.16)$ $2.72 \pm 0.93 \ (2.53 \pm 0.67)$ $5.6 \pm 2 \ (5 \pm 1)$ $0.44 \pm 0.27 \ (0.33 \pm 0.17)$	+1 (-1) 0 (0) -1 (-1) -0.7 (1) -9 (-14) -0.60 (+0.16) -0.14 (-0.11) +0.05 (+0.2) +0.08 (-0.03) +0.21 (+0.16) -0.8 (+0.2) -0.03 (+0.06) -6.5 (-8.5) -33.9 (+22.0)	NS NS NS NS NS 0.05 0.3 0.1 0.07 0.5 0.02 NS NS	-
[65] Pre-post 14 Fair	12	FES knee-extensions 12 weeks 3 sessions/week 2 x 30 (25% Max), 1 x 60 (12.5% Max) Increased by 0.5 kg per session	Body Mass (kg)	67.6	-0.7	NS	-
[66] Pre-post 14 Fair	14	FES knee-extensions 16 weeks 2 sessions/week 4 x 10 Increased by 0.9 kg evert 2 successful sessions	BMI (kg/m²) TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) TC: HDL-C Fasting Glucose (mmol/L) 2-h Glucose OGTT (mmol/L) HOMA-IR HOMA%S HOMA%β	26.7 ± 4.7 1.55 ± 0.94 4.76 ± 1.03 1.09 ± 0.40 2.95 ± 0.94 4.8 ± 1.8 4.94 ± 1.05 6.62 ± 4.30 1.6 ± 1.4 136.0 ± 112.0 125.0 ± 68.0	-0.3 -0.18 +0.09 -0.21 -0.6 +0.22 +0.85 -0.1 +7.0 -14.0	0.70 0.36 0.05 0.02 0.11 0.43 0.16 0.41 0.73 0.65 0.17	0.07 0.16 0.16 0.24 0.21 0.33 0.07 0.19 0.06 0.07
[67] Pre-post 14 Fair	5	FES knee extensions 18 weeks 2 sessions/week 4 x 10 Increased by 0.9-1.8 kg every 2 sessions	Posterior Tibial FMD (when adjusted for resting diameter)	-	+3.9%	0.03	-
[68] Pre-post 13 Low	19	Combined 10-32 weeks 3 sessions/week	Albumin	NR	\leftrightarrow	NS	-

		Max load to fatigue or 45 reps (FES knee- extensions) 30 mins (FES-cycling)					
[69]	11	Combined	SBP (mmHg)	114 ± 4	-16	NS	1.21
Pre-post		13-28 weeks	DBP (mmHg)	71 ± 3	-4	NS	0.40
12		3 sessions/week					
Low		Max load to fatigue or					
		45 reps (FES knee-					
		extensions)					
		Duration NR					
[70]	5	FES knee-extensions	Fasting Glucose (mmol/L)	4.87 ± 0.58	0.0	NS	0.00
Pre-post		12 weeks	Fasting Insulin (mmol/L)	NR	\leftrightarrow	NS	-
11		2 sessions/week	2-h Glucose OGTT (mmol/L)	5.98 ± 1.44	-0.47	NS	0.24
Low		4 x 10	2-h Insulin OGTT	NR	\leftrightarrow	NS	-
		Increased by 0.9-1.8					
		kg every 2 sessions					
[71]	4	Combined	Body Mass (kg)	67.9 ± 5.2	+4.9	NS	0.65
Pre-post		4-12 weeks					
9		5 sessions/week					
Low		Intensity NR					
		15 mins each					

Table 7. Hybrid and FES-rowing studies included in this review.

Study Design	n	Intervention	CMS Outcome	Group Baseline Intervention (Control)	Change Intervention	p value	ES
D&B Quality				Mean ± SD	(Control)		
[25]	9	Hybrid	Waist (cm)	91.8 ± 4.7	-3.9	0.02	0.92
20		16 weeks	Android Fat Mass (kg)	2.0 ± 0.4	-0.1	0.34	0.25
Pre-post†		2 sessions/week	Android Fat (%)	33.4 ± 2.9	-2.1	0.02	0.76
High		65-75% HRR	TG (mmol/L)	1.7 ± 0.2	-0.3	0.01	1.50
		18-32 mins	HDL-C (mmol/L)	1.1 ± 0.1	+0.1	0.22	1.00
			Fasting Glucose (mmol/L)	5.7 ± 0.3	+0.1	0.38	0.28
			Fasting Insulin (pmol/L)	72.7 ± 10.6	-18.9	0.11	1.66
			HOMA-IR	2.8 ± 0.5	-0.6	0.16	1.09
			SBP (mmHg)	112 ± 6	+5	0.39	0.65
			DBP (mmHg)	69 ± 3	-6	0.04	1.70
			CRP (mg/L)	3.91 ± 1.75	-0.71	0.08	0.41
			IL-6 (pg/mL)	2.51 ± 0.91	-0.63	0.20	0.83
[72]	9	Hybrid	Body Mass (kg)	74 ± 18	+1	0.52	0.06
Pre-post		6 weeks	Relative Brachial FMD (%)	-	-	0.28	-
16		2 sessions/week	Relative Femoral FMD (%)	-	-	0.002	-
Fair		Intensity NR 30 mins					
[73]	12	FES-rowing	BMI (kg/m ²)	23.4 ± 3.7	-0.4	0.06	0.11
Pre-post	12	6 weeks	Waist (cm)	84.1 ± 10.3	-2.1	0.06	0.11
15 Fair		5 sessions/week >70% HR _{MAX} 42.5 mins	waist (ciii)	04.1 ± 10.5	-2.1	0.00	0.21
[74]	12	FES-rowing	Body Mass (kg)	72.5 ± 3.9	+0.8	NS	0.20
Pre-post		26 weeks					
14		1.8 ± 2					
Fair		sessions/week 75-85% HR _{PEAK} 30 mins					
[75]	10	Hybrid	Body Mass (kg)	73 ± 10	0	0.77	0.00
Pre-post		4 weeks	SBP (mmHg)	123 ± 18	-4	0.17	0.23
14		2-3 sessions/week	DBP (mmHg)	73 ± 14	-5	0.23	0.38
Fair		Intensity NR	Absolute Brachial FMD (mm)			0.48	-
		30 mins	Relative Brachial FMD (%)			0.68	-
			Absolute Femoral FMD (mm)			0.06	-
			Relative Femoral FMD (%)			0.10	-
[76]	10	FES-rowing	Body Mass (kg)	85.1 ± 19.6	0.0	0.18	0.00
Pre-post		6 weeks	BF (%)	36.9 ± 5.9	-0.2	0.64	0.03
14		3 sessions/week					
Fair		86 ± 8% HR _{PEAK} 30 mins					
[77]	7	FES-rowing	Body Mass (kg)	72.1 ± 3.6	-1.1	NS 0.07	0.14
Pre-post		12 weeks	BF (%)	25.5 ± 1.8	-1.1	0.07	0.26
14		3-4 sessions/week	Leptin (ng/mL)	6.9 ± 1.7	-2.2	0.05	0.60
Fair		80% VO _{2PEAK}	Fasting Glucose (mmol/L)	5.73 ± 0.09	-0.12	<0.05	0.73
		200 kcal/session	Fasting Insulin (pmol/L)	95.1 ± 14.6	-16.7	NS	0.49
F703		TT 1 ' 1	HOMA-IR	3.6 ± 0.8	-0.8	NS	0.65
[78]	8	Hybrid	TC	NR	NR	NS	-
Pre-post		6 weeks	HDL-C	NR	NR	NS	-
7		2 or 3 sessions/week	LDL-C	NR	NR	NS	-
Low		80-90% HR _{MAX}	Glucose OGTT	NR	NR	NS	-

HRPEAK peak heart rate, HRMAX age-predicted maximum heart rate, HOMA-IR homeostatic model assessment of insulin resistance, OGTT oral glucose tolerance test, NS non-significant, NR not reported †True study design is RCT, presented as pre-post due to two different exercise modalities being tested.

Table 8. Ambulation studies included in this review.

Study	n	Intervention	CMS Outcome	Group Baseline	Change	p	ES
Design D&B Quality				Intervention (Control) Mean ± SD	Intervention (Control)	value*	
[41] Pre- post†	17	FES-walking 16 weeks 3 sessions/week	Fat Mass (kg)	25.4	-1.1	NS	0.12
23 High		Max load without knee buckling 45 mins					
[79] RCT 19	18	Robotic BWSTT 12 weeks 3 sessions/week	Body Mass (kg) BF (%)	80.8 ± 14.6 (94.3 ± 25.0) 33.6 ± 7.9 (34.2 ± 6.9)	-1.0 (-2) -1.2 (-0.9)	0.72 0.20	-
High		80-85% HRR 20-45 mins					
[80] Pre-post 19 High	10	BWSTT 16 weeks 3 sessions/week Max speed without loss of gait	SBP (mmHg) DBP (mmHg)	114 ±19 66 ± 11	-1 -2	0.90 0.62	0.05 0.19
[81] Pre-post 18 Fair	8	60 mins BWSTT 26 weeks 3 sessions/week Max load and speed without knee bucking or loss of gait 60 mins	SBP (mmHg) DBP (mmHg)	117 ± 20 73 ± 11	-2 -1	NS NS	0.12 0.15
[82] Pre-post 17 Fair	14	BWSTT 6 weeks 5 sessions/week Intensity NR 45 mins	TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) Fasting Glucose (mmol/L) CRP (NR) SBP (mmHg) DBP (mmHg)	1.36 ± 0.17 4.67 ± 0.54 1.46 ± 0.31 2.61 ± 0.37 5.12 ± 0.67 NR 127 ± 10 75 ± 5	-0.20 -0.14 +0.07 -2.9 -0.19 - 0.15 -3	NS NS NS NS O.002 NS	0.33 0.28 0.26 0.21 0.54 - 0.21 0.49
[83] Pre-post 16 Fair	13	BWSTT 52 weeks 3 sessions/week Minimal load and max speed without knee buckling, losing proper weight shifting, and upright torso Up to 3 x 5-15 min bouts	Fat Mass (kg)	23.6 ± 11.0	+0.4	NS	0.04
[84] Pre-post 16 Fair	5	Robotic Exoskeleton Walking 60-70% HRR 6 weeks 3 sessions/week Up to 60 mins	Body Mass (kg) BMI (kg/m²) BF (%)	79.7 ± 12.5 24.5 ± 1.7 35.4 ± 7.1	+2.0 +0.6 -1.3	0.04 0.04 0.04	0.15 0.32 0.23
[85] Pre-post 15 Fair	9	BWSTT 26 weeks 3 sessions/week Intensity NR Until self-reported fatigue	TG (mmol/L) TC (mmol/L) HDL-C (mmol/L) LDL-C (mmol/L) TC: HDL	1.51 ± 0.20 4.91 ± 0.19 1.29 ± 0.19 3.25 ± 0.22 3.83 ± 0.33	-0.19 -0.55 +0.14 -0.42 -0.76	0.17 0.02 0.19 0.05 0.04	0.33 1.15 0.20 0.54 0.95
[86] Pre-post 14 Fair	9	BWSTT 24 weeks 3 sessions/week Based on self-reported fatigue Until self-reported fatigue	Glucose OGTT (AUC) Insulin OGTT (AUC)	-	-15% -33%	<0.05 <0.05	-
[87] Pre-post 13 Low	16	FES-walking 11 weeks 3 sessions/week Comfortable intensity Up to 3 sets	Body Mass (kg)	66.0	+1.3	0.06	-

BSWTT body-weight supported treadmill training, HRR heart rate reserve, AUC area under the curve † True study design is RCT, presented as pre-post due to two different exercise modalities being tested. *Group x time interaction for RCT and non-randomised controlled trial, or pre-post change for pre-post study designs.

Table 9. Overview of other exercise studies included in review but not grouped for qualitative analysis.

Study	n	Intervention	CMS Outcome	Group Baseline	Change	p	ES
Design				Intervention (Control)	Intervention	value*	
D&B				Mean \pm SD	(Control)		
Quality							
[88]	48	Lower body RT and BSWTT	Body Mass (kg)	$89.4 \pm 20.3 (75.7 \pm 21.0)$	-0.20 (+5.03)	0.31	0.45
RCT		or FES	BMI (kg/m ²)	$27.1 \pm 6.4 (24.8 \pm 6.6)$	0.0 (+0.7)	0.29	0.41
19		24 weeks	QUICKI	$0.35 \pm 0.04 (0.38 \pm 0.06)$	-0.002 (-0.012)	0.92	0.06
High		3 sessions					
Iligii		Intensity NR					
		Up to 180 mins					
[89]	6	Combined RT, ACE, and	Body Mass (kg)	87.7 ± 15.0	\leftrightarrow	NS	-
Pre-post†		FES	Fat Mass (kg)	-	\leftrightarrow	NS	-
18		8 weeks	Android Fat Mass (kg)	-	\leftrightarrow	NS	-
Fair		3 sessions/week	TG (mmol/L)	1.36 ± 0.66	+0.39	0.47	0.45
1 an		ACE: 80-90% VO _{2PEAK} , 15 x	TC (mmol/L)	4.44 ± 0.99	-0.21	0.94	0.25
		1 mins	HDL-C (mmol/L)	1.09 ± 0.16	-0.05	0.96	0.27
		Upper-body RT: 3 x 12	LDL-C (mmol/L)	2.73 ± 0.80	-0.34	0.75	0.48
		FES-knee extensions: 40	Fasting Glucose (mmol/L)	6.12 ± 1.14	-0.54	0.04	0.56
		reps, increased by ~0.5-1 kg	Fasting Insulin (pmol/L)	115.3 ± 127.1	-25.7	0.91	0.24
		every 2 weeks	Glucose OGTT (AUC)	-	+4%	0.87	0.14
			Insulin OGTT (AUC)	-	-27%	0.34	0.28
			HOMA-IR	4.6 ± 5.1	-1.3	0.83	0.31
			ISI-Matsuda	3.3 ± 2.0	+1.3	0.98	0.43
			IL-6 (pg/mL)	1.7 ± 1.0	-0.7	0.20	0.95
			TNF-α (pg/mL)	2.2 ± 0.4	-0.8	0.27	0.97

[|] TNF-α (pg/mL) | 2.2 ± 0.4 | -0.8 |
†True study design is RCT, presented as pre-post due to two different exercise modalities being tested *Group x time interaction for RCT and non-randomised controlled trial, or pre-post change for pre-post study design

Table 10. Participant characteristics, statistical power, and control group (if applicable) of included studies.

Study	Control Type	Statistical Power		Age (y)	TSI (y)	LOI	ASIA
[32]		10001			I: 1.3 (0.2-12), C: 1.3		
[32]	General Exercises	NR	33 (29/4)	I:33 (15-42), C:37 (19-62)	(0.3-10)	C7-L3	A-D
[48]	N/A	NR	10 (9/1)	39±10 (26-55)	9±9 (1-21)	C4-T11	A-C
[25]	1,11	1,12	10 (5/1)	Hybrid: 49±3 (31-64), Hand	Hybrid: 21±3 (13-34),	0.111	11.0
[]	N/A	No	19 (18/1)	cycle: 47±3 (30-63)	Hand cycle: 16±2 (9-21)	C2-L2	A-D
[28]	N/A	NR	10 (8/2)	37±13 (23-55)	12±14 (1-34)	C7-T5	A-B
[62]	N/A	NR	5 (4/1)	31-50	3-25	C5-T8	A
[45]	N/A	NR	16 (16/0)	45±12	12±10	Thoracic	A-C
[39]	No exercise intervention	NR	14 (14/0)	I: 30±3, C: 29±3	I: 19±3, C: 9±3	NR	NR
[42]	Instructed to maintain PA levels	NR	23 (21/2)	I: 39±11, C: 42±13	I: 15±10, C: 9±10	C1-T11	A-D
[81]	N/A	NR	8 (6/2)	28±5 (20-34)	10±8 (2-24)	C4-C5	B-C
[80]	N/A	NR	6 (4/2)	38±15	8±9	C4-T12	A-B
[53]	N/A	NR	13 (12/1)	31±5 (21-41)	8±4 (3-16)	C4-T10	A-D
[37]	N/A	NR	9 (NR)	35±11 (25-50)	12±5 (5-18)	C5-T4	NR
[51]	N/A	NR	9 (9/0)	39±11 (28-44)	11±10 (1-27)	C5-T8	A-C
[41]	N/A	NR	34 (26/8)	FES: 57±14, RT: 54±17	FES: 9±10, RT: 10±11	C2-T12	C-D
[83]	N/A	NR	14 (11/3)	29±8 (20-53)	8±7 (1-24)	C4-T12	NR
[63]	Testosterone replacement therapy only	Yes	22 (22/0)	I: 37±12, C: 35±8	I: 10±9; C: 7±6	C5-T11	A-B (ISNCSCI)
[31]				ACE: 41±13 (30-61); FES-	ACE: 11±9 (2-26); FES-		
	N/A	NR	9 (9/0)	Cycling: 37±7 (29-45)	Cycling: 7±5 (4-14)	C8-T10	A-B
[64]	Standardised diet with no exercise						
	intervention	NR	9 (9/0)	35±9 (21-47)	13±9 (2-26)	C5-T11	A-B
[79]				I: 52±12 (28-66), C: 52±15			
	Stretching (3 days/week for 20-25 mins)	NR	18 (NR)	(30-72)	NR	NR	C-D
[54]	N/A	NR	18 (13/5)	40±2 (25-57)	11±3	C4-T7	NR
[29]	N/A	NR	5 (5/0)	40±7	13.9±5.0	C4-L1	A-D
[78]	N/A	NR	8 (NR)	NR	NR	NR	NR
[46]				I: 37±11 (19-65); C: 43±9	I: 8±6 (1-22); C: 12±7 (3-		
	No exercise intervention	NR	34 (NR)	(29-63)	24)	C4-S1	A-D
[56]	N/A	NR	5 (5/0)	35±3 (28-44)	10±3 (4-23)	C5-C7	A-B
[58]	N/A	NR	5 (5/0)	35±3 (28-44)	10±3 (4-23)	C5-C7	A-B
[40]	N/A	NR	11 (6/5)	31±4 (23-36)	12±7 (2-19)	C5-T9	NR
[34]	N/A	NR	9 (9/0)	38±10	16±7	T8-L1	A-B
[77]	N/A	NR	6 (6/0)	46±5 (24-56)	NR	T4-T10	A-B
[50]	N/A	NR	7 (5/2)	45±8 (30-53)	20±14 (3-40)	C5-T10	NR
[88]	No exercise intervention	Yes	48 (30/11)	I: 42±13; C: 34±12	I: 7±10; C: 6±7	NR	C-D
[57]						C4-C8 and T1-	
	N/A	NR	12 (NR)	NR	>1	T10	NR

[84]	N/A	NR	5 (4/1)	60±6	8±5	C7-T10	NR
[33]	No exercise intervention	NR	15 (9/6)	33±6 (22-46)	7±4 (2-16)	C5-T11	A-B
[44]	Standard Care	NR	17 (11/6)	37±7 (23-53)	10±7 (2-27)	C4-L1	A-C
[73]	N/A	NR	12 (10/2)	36±12 (16-45)	11±6 (5-24)	C6-L1	A-C
[87]	N/A	NR	16 (13/3)	28±7 (21-45)	4±3 (0.7-9)	T4-T11	NR
[59]	N/A	NR	8 (8/0)	39±3	>4	C5-T11	A-B
[52]	N/A	NR	18 (16/2)	40±11 (26-61)	3±2 (1-9)	C3-L1	B-D
[89]	N/A	NR	6 (6/0)	50±8 (36-58)	24±8 (10-30)	C6-T6	A-B
[70]	N/A	NR	5 (5/0)	36±5	13±7	C5-T10	A
[30]	N/A	NR	14 (NR)	Supine: 34±12; Sitting: 33±7	Supine: 9±13; Sitting: 14±6	CT-T1	NR
[35]			12 (11/1) (2 non-				
	N/A	NR	SCI)	38±10 (22-58)	15±7 (4-29)	C6-L3	NR
[43]	No exercise intervention	NR	20 (20/0)	I: 25±3; C: 26±3	I: 10±4; C: 9±4	T9-T12	A
[60]	N/A	NR	10 (8/2)	35 (27-45)	12 (3-23)	C6 and T4	NR
[36]	N/A	NR	12 (12/0)	31±9 (19-45)	2±1 (1-3)	<t10< td=""><td>NR</td></t10<>	NR
[47]	N/A	NR	5 (5/0)	38±4 (34-43)	5±1 (1-7)	T6-T12	NR
[26]	No exercise intervention	Yes	21 (15/6)	I: 46±6, C: 48±10	I: 20±10; C: 14±11	T4-L3	A-D
[71]	N/A	NR	4 (4/0)	20-35	4±3 (1-8)	T4-T6	NR
[86]	N/A	NR	9 (8/1)	31±3	8±3	C4-T12	С
[69]	N/A	NR	11 (7/4)	29±15 (18-54)	6±3 (0.5-11)	C4-T6	NR
[68]	N/A	NR	19 (16/3)	19-47	2-17	C4-T10	NR
[55]	N/A	NR	8 (7/1)	32±2 (23-41)	12±2 (5-24)	C7-L1	NR
[65]	N/A	No	12 (9/3)	38±13 (19-63)	6±6 (1-17)	C4-T10	NR
[27]	No exercise intervention	NR	17 (17/0)	30±4 (I & C)	5±0	≤T5	NR
[66]	N/A	No	14 (11/3)	27±5 (28-57)	8±7 (2-22)	C4-T7	A-B
[49]	Standard Care	NR	45 (38/7)	I: 37±12; C: 35±12	I: 8 (1.5-43), C: 6 (1-27)	C1-L5	A-C
[74]	N/A	Yes	12 (11/1)	33±4 (22-60)	8±3 (0-33)	C4-T2	NR
[61]	No exercise intervention	NR	15 (15/0)	33 (21-48)	9 (1-21)	NR	A-B
[85]	N/A	NR	9 (8/1)	31±3	8±3	C4-T12	С
[67]	N/A	NR	5 (5/0)	36±5	13±7	C5-T10	A
[72]	N/A	NR	9 (8/1)	39±3 (25-52)	11±3 (1-25)	C5-T12	A, C
[75]	N/A	NR	10 (9/1)	39±9 (23-53)	11±6 (1-20)	T1-T12	A, C
[82]	N/A	NR	14 (10/4)	51±17	2-10	NR	Motor Incomplete
[76]	N/A	NR	10 (8/2)	47±18	18±14 (2-39)	T4-T12	A-C
[38]	N/A	NR	11 (11/0)	31±8 (20-49)	2±1 (0.5-4)	T8-T12	A

TSI time since injury, LOI level of injury, ASIA American Spinal Injury Association Impairment Scale, NR not reported, ISNCSCI International Standards for Neurological Classification of Spinal Cord Injury, ROM range of motion; I Intervention, C Control.