Accepted Manuscript

A systematic review and meta-analysis of the safety, feasibility and effect of exercise in women with stage II+ breast cancer.

Ben Singh, MSc, Rosalind R. Spence, Phd, Megan L. Steele, Phd, Carolina X. Sandler, Phd, Jonathan M. Peake, Phd, Sandra C. Hayes, Phd

PII: S0003-9993(18)30280-6

DOI: 10.1016/j.apmr.2018.03.026

Reference: YAPMR 57222

To appear in: ARCHIVES OF PHYSICAL MEDICINE AND REHABILITATION

- Received Date: 21 December 2017
- Revised Date: 24 February 2018

Accepted Date: 23 March 2018

Please cite this article as: Singh B, Spence RR, Steele ML, Sandler CX, Peake JM, Hayes SC, A systematic review and meta-analysis of the safety, feasibility and effect of exercise in women with stage II+ breast cancer., *ARCHIVES OF PHYSICAL MEDICINE AND REHABILITATION* (2018), doi: 10.1016/j.apmr.2018.03.026.

This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.



- 1 **Running head:** Exercise and stage II+ breast cancer.
- 2 Title: A systematic review and meta-analysis of the safety, feasibility and effect of exercise
- 3 in women with stage II+ breast cancer.
- 4 **Authors:** Ben Singh, MSc^{1, 2}, Rosalind R Spence, Phd^{1, 2}, Megan L Steele, Phd^{1, 2}, Carolina
- 5 X Sandler, Phd^{1, 2, 3}, Jonathan M Peake, Phd^{2, 4} and Sandra C Hayes, Phd^{1, 2}

6 Affiliations:

- ⁷ ¹ School of Public Health and Social Work, Queensland University of Technology, Victoria
- 8 Park Road, Brisbane, Queensland, Australia
- 9 ² Institute of Health and Biomedical Innovation, Queensland University of Technology, Musk
- 10 Avenue, Kelvin Grove, Queensland, Australia
- ³ Cancer Prevention Research Centre, School of Public Health, Faculty of Medicine, The
- 12 University of Queensland, Herston, Queensland, Australia
- ⁴ School of Biomedical Sciences, Queensland University of Technology, Victoria Park Road,
- 14 Kelvin Grove, Queensland, Australia
- **Suppliers:** RevMan software (version 5.3); R statistical software (version 3.4.1).
- 16 **Funding:** The authors received no specific funding for this work.
- 17 **Disclosure:** The authors declare no conflicts of interest.
- 18 Corresponding author:
- 19 Ben Singh
- 20 School of Public Health and Social Work, Queensland University of Technology, Victoria
- 21 Park Road, Kelvin Grove, Queensland, Australia
- 22 b6.singh@qut.edu.au

1 Abstract

Objective: To systematically evaluate the safety, feasibility and effect of exercise 2 among women with stage II+ breast cancer. Data Sources: CINAHL, Cochrane, Ebscohost, 3 MEDLINE, Pubmed, ProQuest Health and Medical Complete, ProQuest Nursing and Allied 4 5 Health Source, Science Direct and SPORTDiscus were searched for articles published prior 6 to March 1, 2017. Study selection: Randomised, controlled, exercise trials involving at least 50% of women diagnosed with stage II+ breast cancer were included. **Data Extraction:** Risk 7 8 of bias was assessed and adverse event severity was classified using the Common Terminology Criteria. Feasibility was evaluated by computing median (range) recruitment, 9 10 withdrawal and adherence rates. Meta-analyses were performed to evaluate exercise safety and effects on health outcomes only. The influence of intervention characteristics (mode, 11 supervision, duration and timing) on exercise outcomes were also explored. Data Synthesis: 12 13 There were no differences in adverse events between exercise and usual care (risk difference: <0.01 [95% CI: -0.01, 0.01]), p=0.38). Median recruitment rate was 56% (1%-96%), 14 withdrawal rate was 10% (0%-41%) and adherence rate was 82% (44%-99%). Safety and 15 16 feasibility outcomes were similar, irrespective of exercise mode, supervision, duration, or timing. Effects of exercise for quality of life, fitness, fatigue, strength, anxiety, depression, 17 body mass index and waist circumference compared with usual care were significant 18 (standardised mean difference range: 0.17–0.77, p<0.05). Conclusion: The findings support 19 20 the safety, feasibility and effects of exercise for those with stage II+ breast cancer, suggesting 21 that national and international exercise guidelines appear generalizable to women with local, regional and distant breast cancer. 22

23 Key words: breast neoplasm, aerobic exercise, resistance exercise, exercise oncology.

1

24 There is growing scientific and community support to incorporate exercise into standard breast cancer care.¹ Previous systematic reviews have demonstrated a low risk of 25 serious adverse events with exercise.²⁻⁴ Specifically, no serious adverse events have been 26 reported and over 80% of trials included in previous reviews have reported no exercise-27 related adverse events for individuals with cancer.²⁻⁴ Exercise is also considered feasible. 28 Previously reported recruitment rates have ranged between 20 to 70%⁵, withdrawal rates have 29 been low (<10%) and exercise adherence rates have been high (80-90%).²⁻⁵ The health 30 benefits of exercise both during and following treatment have also been well described in 31 systematic reviews and meta-analyses.^{3, 6, 7} Specifically, exercise improves fatigue, aerobic 32 fitness, muscular strength, anxiety, body image and self-esteem, cognitive health, 33 psychosocial distress and overall quality of life (QOL).^{1, 3, 4, 7-10} Observational evidence also 34 indicates that among women with breast cancer, exercise reduces the risk of subsequent 35 chronic disease (including diabetes, osteoporosis, cardiovascular disease), reduces the risk of 36 cancer recurrence and improves survival,^{7, 10-12} 37

Most studies included in systematic reviews on exercise and breast cancer have 38 comprised of a sample primarily with early-stage and localised breast cancer.^{13, 14} However, 39 population-based statistics suggest that approximately 50% of women with breast cancer are 40 diagnosed with regional or distant disease (Stage II+).^{13, 15} As such, it is plausible that women 41 with stage II+ disease are underrepresented in the body of evidence, which currently supports 42 exercise as being safe, feasible and effective during and following breast cancer treatment. 43 This is of note since breast cancer stage influences the types of treatment prescribed. More 44 invasive surgery and higher doses of adjuvant treatment are associated with more frequent 45 and severe treatment-related sequelae.¹⁶⁻²⁰ Further, five-year relative survival declines with 46 advancing stage (Stage I: 99%; Stage II: 93%, Stage III: 72%; Stage IV: 22%^{15, 21, 22}). As 47 such, compared with early stage breast cancer, the higher disease and treatment-related 48

burden associated with later stage breast cancer may also influence safety, feasibility andexercise outcomes.

The aim of this systematic review and meta-analysis was to assess the safety, feasibility 51 and effect of exercise in women with stage II, III or IV disease (i.e., II+). Specifically, this 52 review evaluated: 1) the number, type and severity of adverse events (safety): 2) study 53 recruitment, withdrawal and adherence rates (feasibility); and 3) effect of exercise (as 54 assessed immediately post-intervention) on survivorship outcomes including QOL, aerobic 55 56 fitness and fatigue. This analysis was performed by evaluating findings derived from randomised, controlled trials (RCTs) that involved samples with >50% of women with stage 57 II+ breast cancer. As a secondary objective, we also explored the relationship between safety, 58 feasibility, effect, and intervention characteristics, including exercise mode, degree of 59 intervention supervision, intervention duration and timing of intervention (during or 60 61 following treatment).

62

63 Methods

64 Search strategy and selection criteria

Eligibility criteria were established using the Participants, Intervention, Comparator, 65 and Outcome (PICO) framework²³ as follows: Participants: RCTs in which at least 50% of 66 the sample was diagnosed with Stage II+ breast cancer, either undergoing or completed 67 treatment. If a study involved multiple intervention arms, groups consisting of less than 50% 68 of participants with Stage II+ disease were excluded. Intervention: Exercise intervention trials 69 70 were eligible for inclusion. Exercise was defined as any form of planned, structured, and repetitive bodily movements performed in order to improve or maintain fitness, performance 71 or health.^{24, 25} Exercise mode was classified as aerobic, resistance or other. 'Other exercise' 72

73 was considered a form of exercise that: 1) was not specified as aerobic or resistance (e.g., yoga); and 2) did not constitute complete decongestive therapy-based exercise, or common 74 forms of lymphoedema treatment (e.g., stretching, passive, assistive, remedial or range of 75 76 motion exercise performed against no resistance). Trials were eligible regardless of the level of supervision provided, mode of intervention delivery, intervention duration or intensity. 77 Studies that involved multiple intervention groups consisting of different exercise intensities 78 or modes were eligible if they included a control group. Studies that involved exercise in 79 addition to other interventions such as dietary or other lifestyle interventions were excluded if 80 81 the outcomes of the exercise could not be isolated. Comparators: Studies were included if they involved a usual care or control group (i.e., any type of control group not involving 82 exercise therapy). 83

The following electronic databases were searched by one reviewer (BS): CINAHL, 84 Cochrane, Ebscohost, MEDLINE, Pubmed, ProQuest Health and Medical Complete, 85 ProQuest Nursing and Allied Health Source, Science Direct and SPORTDiscus. A faculty 86 liaison librarian was consulted in the development of search terms. Titles and abstracts were 87 searched for the following terms: 'breast neoplasm' or 'breast cancer' 'or 'breast' and 88 '(cancer or neoplasm)' and/or 'advanced' or 'metastatic' or 'stage II, III, IV' or 'late stage' or 89 'palliative' and 'physical activity' or 'aerobic' or 'exercise' or 'training' or 'fitness' or 90 'physical' or 'jogging' or 'walking' or 'running' or 'swim*' or 'bik*' or 'bicyc*' or 'cycl*' 91 or 'weight lifting' or 'aerobics' or '(strength or resistance)' or 'hydrotherapy' or 'water*' or 92 'yoga' and/or 'exercise' or 'movement' or 'exercise tolerance' or 'exercise therapy'. 93 Database searches were limited to peer-reviewed scholarly journal articles published in 94 English-language prior to March 1, 2017. There was no registered protocol for this review. 95

96

97 **Outcomes of interest**

4

Safety

98

99 Adverse events were defined as any undesirable medical or health-related event that occurred during study participation. They were classified as either non-exercise adverse 100 events (adverse events reported to have occurred during study participation, but considered 101 unrelated to exercise) or exercise-related adverse events (events which occurred during, or as 102 a direct result of exercise). Adverse events were categorised according to the Common 103 Terminology Criteria for Adverse Events, Version 4^{26} as grade 1: asymptomatic or mild 104 105 symptoms, clinical or diagnostic observations only and/or intervention not indicated; grade 2: moderate, minimal, local or non-invasive intervention required and/or limiting age-106 appropriate activities of daily living; grade 3: severe or medically significant but not 107 immediately life-threatening, hospitalisation and/or prolongation of hospitalisation indicated, 108 disabling and limiting self-care activities of daily living; grade 4: life-threatening 109 consequences and urgent intervention indicated, or; grade 5: death. Serious adverse events 110 were considered any "adverse medical event that required hospitalization, resulted in 111 significant disability, was life threatening or resulted in death".²⁷ The lack of reporting and 112 categorisation of health-related withdrawals as adverse events is common in exercise trials, 113 and is suggestive of under-reporting of adverse events.²⁸ Therefore, we considered any 114 withdrawal that occurred due to health-related reasons as an adverse event (e.g., illness or 115 cancer recurrence). However, if participants withdrew for reasons such as time constraints, 116 travel or family reasons, these were not considered adverse events (i.e., non-health-related 117 reasons). If the severity of an adverse event was not reported, and the event resulted in study 118 withdrawal, or if a participant withdrew from a trial due to unspecified health or medical 119 reasons, these events were categorised as grade 3. If a study did not report on the occurrence 120 121 of adverse events, and no health-related withdrawals occurred, it was considered that no adverse events had occurred. If a study involved more than one intervention group and did 122

not specify in which intervention group an adverse event occurred, the data were not includedin the meta-analysis.

125 *Feasibility*

Feasibility was determined by computing recruitment rate, withdrawal rate, reason for withdrawals, and exercise adherence rate. Recruitment rates were computed as the proportion of those who were eligible and consented to participate in the study. Withdrawal rates were calculated as the percentage of those enrolled who did not complete the study. Exercise adherence rates were calculated as a percentage of the scheduled number of exercise sessions that were completed by participants.

132 *Health outcomes*

Health outcomes that were reported in a minimum of two studies were included in a
meta-analysis. These included QOL, aerobic fitness, fatigue, upper-body strength, anxiety,
depression, body mass index, body fat percentage, body mass index and waist circumference.

136 **Data extraction and management**

The titles and abstracts of all articles identified through an electronic database search were screened for eligibility by one reviewer (BS). Reference lists of all eligible and original manuscripts, and reviews were manually checked to identify additional articles (BS). Relevant records were then retrieved in full-text and screened further against the eligibility criteria (BS). Study and participant characteristics, intervention features and outcomes assessed from included articles were extracted into tabular format using predefined data fields (BS).

The quality of methods used in each RCT was assessed independently by two investigators (CS and BS) using the Physiotherapy Evidence Database (PEDro scale). The PEDro scale is a valid and reliable tool for evaluating risk of bias and quality in RCTs.^{29, 30}

The scale consists of 11 items (eligibility criteria, random allocation, allocation concealment, baseline differences between groups, subject blinding, therapist blinding, assessor blinding, attrition, intention-to-treat analyses, between-group statistical comparisons and reporting of measures), with the total PEDro score ranging from 0 to 10 points (item 1 not contributing to the total score). RCTs with a score 6 or higher were considered high quality. RCTs receiving less than 6 were classified as low quality.^{29, 31} Discrepancies in ratings were resolved by discussion and consultation with a third reviewer (SH) when required.

154 Statistical analyses

155

Meta-analysis of adverse events

Adverse events were treated as a count variable for inclusion in the meta-analysis. The 156 number of adverse events that occurred in the exercise participants compared to the usual 157 care participants was pooled and analysed, using a Mantel-Haenszel random effects model. 158 The risk difference (RD) and 95% confidence interval was calculated as the effect measure. 159 The RD was considered most appropriate since there were studies included in this review that 160 reported no adverse events in either group. Performing a meta-analysis using risk ratio as the 161 effect measure would also exclude all studies with zero adverse events.^{32, 33} A negative value 162 for RD indicates a lower risk of an adverse event with exercise compared with usual care. 163 Meta-analysis was performed only for adverse events that were grade 3 or higher. This was 164 considered appropriate because evaluation of grade 3 or higher adverse events was more 165 166 likely to be consistent across the intervention versus usual care groups. Conversely, reporting of grade 1–2 events may not have been comprehensively evaluated for those in the usual care 167 groups due to reduced contact with study staff. Further, some grade 1-2 events may reflect 168 normal physiological responses to exercise (e.g., mild muscle stiffness or soreness) as 169

170 compared to potentially avoidable adverse events.^{34, 35} All adverse events (grade 1–5) were
171 also evaluated descriptively.

172 *Feasibility*

Feasibility was evaluated by calculating study recruitment rate, withdrawal rate and exercise adherence rate (all as a percentage); median, interquartile range, minimum and maximum rates were reported to ensure adequate description of data and to account for skewed data. We defined feasibility of exercise as achievement of a recruitment rate of $\geq 25\%^{36}$, a withdrawal rate of <25% (i.e., retention of $\geq 75\%^{37}$) and adherence of $\geq 75\%^{.37}$ These values were determined *a priori* as clinically relevant cut-offs to establish feasibility based on previous literature.^{38, 39}

180 *Meta-analysis of health outcomes*

All health outcomes of interest were analysed as continuous variables and involved 181 comparisons of post-intervention means and standard deviations (SDs) between exercise and 182 usual care participants. To allow comparison of data from different scales, pooled statistics 183 were calculated using standardised mean differences (SMDs) using RevMan software 184 (version 5.3). Forest plots were created using R statistical software (version 3.4.1). When 185 means and SDs were not available (n=9 studies), authors were contacted (two responded), or 186 187 means and/or SDs were calculated using reported data (e.g., using median, range and sample size) and recommended formulas.⁴⁰ If authors could not be contacted, and means or SDs 188 could not be calculated (because of insufficient data or data being reported in graph format 189 only), the study was not included in meta-analyses (n=7). When two or more methods of 190 assessing outcomes were used in a study, the method defined as being the gold standard or 191 the method/instrument with demonstrated validity and reliability was used. 192

193 Data were combined at the study level for each meta-analysis. Publication bias was assessed by plotting RDs or SMDs against corresponding standard errors and determining the 194 presence of asymmetries or missing sections within the funnel plot when ten or more studies 195 were available.⁴¹ Statistical heterogeneity was assessed using Cochran's Q test and the I^2 196 statistic to quantify the proportion of the overall outcome attributed to variability.^{42, 43} The 197 following values were used to determine level of heterogeneity: $l^2=0-25\%$: low 198 heterogeneity; $I^2 = >25-50\%$: moderate heterogeneity; $I^2 = >75-100\%$.^{3, 43} Planned subgroup 199 analyses were performed to assess the influence of: 1) exercise mode (aerobic, resistance, 200 combined and 'other' exercise); 2) degree of intervention supervision (supervised and 201 unsupervised); 3) intervention duration (12 weeks or less and greater than 12 weeks), and; 4) 202 timing of the intervention with respect to treatment status of participants (during treatment, 203 post-treatment and mixed [i.e., samples consisting of those currently receiving and completed 204 treatment]) on adverse events, recruitment, withdrawal and adherence rates, and effect of 205 exercise on health outcomes. Sensitivity analyses were also performed by repeating all meta-206 analyses with: 1) only trials rated as high quality using the PEDro scale, and; 2) only trials 207 with 100% of samples with stage II+ disease. Standardised classifications for the magnitude 208 of effect were used, with less than 0.20 representing a small effect; >0.20–0.50 representing a 209 moderate effect; and >0.50 representing a large effect.⁴⁴ A p-value of less than 0.05 was 210 considered statistically significant. 211

212

213 **Results**

214 *Literature search*

Following a search of databases, 2,391 articles were identified (Supplementary Material
1). After removal of duplicates and screening of titles and abstracts, 406 publications were

retrieved and examined. Of these, 345 were excluded (with 60% of exercise and breast cancer trials excluded as they comprised samples with <50% of participants with stage II+ breast cancer). After these exclusions, 61 trials were included in the systematic review (low quality, n=24, 39%; high quality, n=37, 61%, Supplementary Material 2).

221 *Participant characteristics*

Median sample size was 63 (range: 10–377), with a participant mean age of 53 years 222 (SD=3.6, see Online Supplementary Material 3). The period since breast cancer diagnosis 223 ranged between 8 months⁴⁵ and 6 years^{46, 47}; 41% (n=25) of trials involved participants who 224 225 were currently undergoing treatment, including neoadjuvant, adjuvant or palliative treatment. The median proportion of the samples with stage II+ disease was 72% (range: 50% (n=2 226 studies^{48, 49}) to 100% (n=10 studies, 50-59). Within the ten trials that involved only participants 227 with stage II+ disease⁵⁰⁻⁵⁹, one trial included only participants with stage II disease⁵⁴, seven 228 trials included only those with stage II or III disease^{50, 55-59}, and two trials included only 229 women with stage IV disease.^{51, 53} 230

231 *Intervention characteristics*

Details of intervention characteristics are shown in Table 1. Approximately one-third of 232 studies (n=20^{45, 47, 48, 50, 52-54, 59-71}) evaluated aerobic exercise only, whereas another third 233 (n=21^{9, 55, 57, 72-89}) evaluated combined aerobic and resistance exercise. The remaining studies 234 evaluated resistance exercise only (n=6 studies^{46, 49, 90-93}), or other modes of exercise (n=11, 51 , 235 ^{56, 58, 94-101}), and three trials involved separate aerobic and resistance exercise arms (n=3, 102-1)236 ¹⁰⁴). Home-based exercise was prescribed for approximately one-third of the interventions 237 (n=20 studies, 32%^{9, 50, 53, 61, 64, 65, 67-72, 75, 79, 81, 84, 88, 89, 103, 104}), while the other two-thirds 238 involved interventions conducted at a range of facilities including local gymnasiums, 239 hospital, clinical, university or rehabilitation settings. Approximately half of the interventions 240

241 involved supervised exercise sessions (i.e., over half of the exercise sessions involved faceto-face supervision, $n=31^{45-49, 52, 54, 57, 59, 60, 62, 73, 74, 76-78, 80, 82, 83, 85, 86, 90, 92-99, 102}$), with one trial 242 evaluating a supervised and an unsupervised intervention group.⁸⁷ Supervision in these trials 243 was provided by an accredited exercise physiologist (n=9, $^{46-48, 52, 60, 80, 85, 86, 102}$), other exercise 244 trainers with or without tertiary qualifications (n=10,^{49, 57, 76, 78, 82, 83, 90, 93, 95, 96}), or other allied 245 health professionals such as an occupational therapist or physical therapist (n=8, 45, 73, 78, 87, 92, 246 ⁹⁵⁻⁹⁷). The interventions in 29 trials were classified as unsupervised (i.e., less than half of the 247 prescribed exercise sessions involved face-to-face supervision: n=29,^{9, 50, 51, 53, 55, 56, 58, 61, 63-72,} 248 75, 79, 81, 84, 88, 89, 91, 100, 101, 103, 104). Of these 29 trials, nine involved predominantly unsupervised 249 exercise sessions, supplemented with some face-to-face contact or supervision, commonly 250 once per week.^{9, 53, 55, 56, 58, 66, 91, 100, 101} Eleven trials involved telephone contact with an 251 exercise specialist⁷⁵, research staff member^{61, 68-70, 81, 104}, accredited exercise physiologist⁷², 252 nurse⁶³ or a physical activity counsellor^{64, 65} throughout the intervention. The remaining nine 253 unsupervised trials involved other forms of intervention support such as provision of 254 guidebooks or print materials^{50, 67, 71, 89, 103}, emails with support from an e-counsellor exercise 255 physiologist⁷⁹, a website⁸⁸ or exercise instructional videos or CDs.^{51, 84} Intervention durations 256 ranged between 6 weeks and 1 year (median 12 weeks, Table 1). 257

258

Safety - summary of adverse events

259 *Adverse events in exercise participants*

From 61 studies included in this review, 41% (n=25) explicitly reported that no adverse events had occurred, while 34% (n=21) did not mention adverse events (see Online Supplementary Material 4). There were a total of 116 adverse events among participants allocated to exercise reported in 15 trials^{52, 55, 64, 65, 68, 70, 72, 73, 81, 87, 93, 95-97, 103} (grade 1: n=42 events; grade 2: n=20 events; grade 3: n=52 events; grade 4: n=0 events; grade 5: n=2 events, Table 2). The most common adverse events among exercise participants were unspecified

266 health or medical problems or illness leading to withdrawal (n=20 events, grade 3), discomfort or low-level muscle pain, stiffness or soreness after an exercise session (n=18 267 events, grade 1) and musculoskeletal injuries (e.g., sprains: n=8 events, grade 1). While 58% 268 (n=66) of reported adverse events were considered unrelated to exercise, 42% (n=50) were 269 exercise-related. Of these events, most (n=43, 88%) were classified as grade 1 or 2 (grade 1: 270 n=34 events; grade 2: n=9 events; grade 3: n=6 events). Of the six exercise-related adverse 271 events that were grade 3, five of these events resulted in participant withdrawal. These were 272 severe headaches (n=1 event), an unspecified physical accident (n=1 event), severe 273 discomfort (n=1 event), dizziness (n=1 event) and foot pain requiring surgery (n=1 event). 274

275

Adverse events in usual care participants

Seventeen studies^{45, 49, 52, 53, 58, 60, 62, 83, 86-88, 90-93, 100, 101} reported a total of 40 adverse events in those allocated to usual care (grade 1: n=2 events; grade 2: n=1 event; grade 3: n=34 events; grade 4: n=0; grade 5: n=3, Table 2). The most common adverse events among usual care participants were unspecified health or medical problems or illness leading to withdrawal (n=11 events, grade 3), infections, secondary suturing, seroma discharge or uncontrollable pain (not reported individually, n=8 events, grade 3) and breast cancer progression (n=4 events, grade 3).

283

284 Meta-analyses of adverse events

Adverse event data from one trial⁶⁶ (n=5: wheezing requiring physician evaluation for asthma, cholinergic urticarial, herpes zoster, sinusitis, and back pain related to a fall) were not included in the meta-analysis since group allocation was unclear. Further, adverse event data (n=2: shoulder tendonitis and foot tendonitis) from another trial¹⁰³ involving two exercise

intervention groups were excluded from subgroup analyses of exercise mode due to a lack ofclarity of intervention group allocation.

Pooled analyses of 60 RCTs involving 5,200 participants (exercise: n=2,621; usual 291 care: n=2,579) showed no difference in the risk of a grade 3-5 adverse event between 292 exercise and usual care (n=91 adverse events [exercise: n=54 events; usual care: n=37 293 events], RD: <0.01 [95% CI= -0.01, 0.01]; p=0.38; $I^2=0\%$: low heterogeneity, Figure 1). 294 Evaluation of funnel plots indicated there was no evidence of publication bias (data not 295 296 shown). The results of subgroup analyses suggested that results were similar irrespective of exercise mode (aerobic, resistance, combined and other exercise), intervention supervision 297 (supervised and unsupervised), intervention duration (12 weeks or less and longer than 12 298 weeks) and intervention timing (during and after treatment). The RD remained unchanged 299 following sensitivity analyses involving only high-quality trials and trials with 100% of 300 samples with stage II+ disease (Figure 1). 301

302 Feasibility outcomes: recruitment, withdrawals, and exercise adherence

303 Recruitment, withdrawal and adherence rates are shown in Table 3. Recruitment rates: Study recruitment rates were calculated for 48 studies (data from 13 studies were 304 unavailable). Median recruitment rate met the pre-defined criterion of >25%, with an overall 305 rate of 45%. Recruitment rates varied based on exercise mode, with aerobic exercise studies 306 showing the lowest rates (32%) and studies evaluating 'other' modes of exercise showing the 307 308 highest rates (65%). Withdrawals: Overall withdrawal rate was 11%, across a total of 69 intervention groups, with similar rates irrespective of subgroup (Table 3). Lower withdrawal 309 rates occurred in studies with a high-quality rating compared with low quality studies 310 311 (exercise groups: 18% [low-quality studies] versus 9% [high-quality studies]; usual care groups: 16% [low-quality studies] versus 11% [high-quality studies]). Health-related reasons 312

for withdrawal were similar between exercise and usual care groups. Unspecified health or medical reasons were the most common reason (see Online Supplementary Material 5 for all reasons for withdrawals). *Exercise adherence:* Overall median adherence to the scheduled number of exercise sessions was 82% (Table 3), and rates were similar irrespective of subgroup.

318

319 *Health Outcomes: assessment of outcomes.*

An overview of all instruments and methods used to assess specific health outcomes, including QOL, aerobic fitness, fatigue, upper-body strength, anxiety, depression and body composition, body mass index, body weight and waist circumference is shown in Supplementary Material 6.

324

325 Meta-analyses results of health outcomes: exercise versus usual care

Large effects in favour of exercise compared with usual care were observed for aerobic 326 fitness (SMD=0.62 [95% CI: 0.42, 0.81], p<0.01, I^2 =75%; moderate heterogeneity, n=31 327 trials, Figure 2), anxiety (SMD=0.77 [95% CI: 0.64, 0.91]; p<0.01, I^2 =89%; high 328 heterogeneity, n=14 trials, see Supplementary Content 7) and depression (SMD=0.66 [95% 329 CI: 0.52, 0.80]; p<0.01, I^2 =90%; high heterogeneity, n=14 trials, see Supplementary Content 330 8). Compared with usual care, there were moderate effects in favour of exercise for QOL 331 (SMD=0.40 [95% CI: 0.33, 0.47]; p<0.01, I^2 =78%; high heterogeneity, n=40 trials, Figure 3), 332 fatigue (SMD=0.30 [95% CI: 0.23, 0.38], p<0.01, I^2 =75%; moderate heterogeneity, n=31 333 trials, Figure 4), upper-body strength (SMD=0.43 [95% CI: 0.33, 0.53]; p<0.01, I^2 =49%; 334 moderate heterogeneity, n=22 trials, see Supplementary Content 9) and waist circumference 335 (SMD=0.22 [95% CI: 0.02, 0.43]; p=0.03, I^2 =0%; low heterogeneity, n=8 trials, see 336

Supplementary Content 10). Small effects from exercise were observed for body mass index (SMD=0.17 [95% CI: 0.01, 0.32]; p=0.03, I^2 =0%; low heterogeneity, n=13 trials, see Supplementary Content 11), body weight (SMD=0.08 [95% CI: -0.04, 0.20]; p=0.22, I^2 =0%; low heterogeneity, n=15 trials) and body fat (SMD=0.11 [95% CI: -0.02, 0.24]; p=0.11, I^2 =0%; low heterogeneity, n=13 trials), with effect on only body mass index also being supported statistically (see Supplementary Content 12–13).

The results of subgroup analyses showed that exercise mode significantly influenced 343 344 exercise effect on QOL (γ^2 =26.36, df=3, p<0.01), with evidence of small-to-moderate effects in favour of aerobic (SMD=0.22 [95% CI: 0.10, 0.33], p<0.01), resistance (SMD=0.29 [0.09, 345 0.49], p<0.01) and combined exercise (SMD=0.51 [95% CI: 0.39, 0.62] p<0.01), and large 346 effects in favour of 'other' exercise (SMD=0.75 [95% CI: 0.55, 0.95], p<0.01) compared with 347 usual care. Subgroup analysis suggested that exercise mode influenced the effect on aerobic 348 fitness (χ^2 =6.05, df=3, p=0.05), with aerobic (SMD=0.62 [95% CI: 0.43, 0.81], p<0.01) and 349 combined exercise (SMD=0.65 [95% CI: 0.26, 1.03]) having a large effect, and resistance 350 exercise having a small to moderate effect (clinically), although not supported statistically 351 (SMD=0.23 [95% CI: -0.07, 0.53], p=0.13). Exercise mode also influenced upper-body 352 strength (χ^2 =12.44, df=2, p<0.01), anxiety (χ^2 =40.91, df=3, p<0.01) and depression 353 $(\chi^2=40.54, df=3, p<0.01)$. For upper-body strength, a large effect was observed for resistance 354 exercise (SMD=0.68 [95%CI: 0.05, 0.85]; p<0.01). For anxiety and depression, large effects 355 were observed for combined exercise (anxiety: SMD=1.36 [95% CI: 1.10, 1.62]; p<0.01; 356 depression: SMD=0.62 [95% CI: 0.18, 1.06]; p<0.01) and 'other' exercise (anxiety: 357 SMD=0.83 [95% CI: 0.61, 1.06]; p<0.01 depression: SMD=1.16 [95% CI: 0.94, 1.38); 358 p<0.01) compared with small-to-moderate effects for aerobic exercise (anxiety: SMD=0.37 359 [95% CI: 0.09, 0.65]; p=0.01; depression: SMD=0.53 [95% CI: 0.24, 0.82]; p<0.01) and no 360

361 effect for resistance exercise (anxiety: SMD=0.08 [95% CI:-0.30, 0.45]; p=0.68; depression:
362 SMD=0.04 [95% CI: -0.23, 0.31]; p=0.79).

Intervention supervision influenced the effect of exercise on QOL (χ^2 =13.74, df=1, 363 p<0.01), fatigue (χ^2 =5.87, df=1, p=0.02), anxiety (χ^2 =5.26, df=1, p=0.02,) and depression 364 $(\chi^2=16.51, df=1, p<0.01)$. Supervised interventions had large effects on QOL (SMD=0.59) 365 [95% CI: 0.46, 0.71], p<0.01) and fatigue (SMD= 0.44 [95% CI: 0.30, 0.57]; p<0.01), while 366 small effects were observed for unsupervised interventions (OOL: SMD=0.30 [95% CI: 0.22, 367 368 0.39], p<0.01; fatigue: SMD= 0.24 [95% CI: 0.15, 0.33]; p<0.01). In contrast, large effects were observed during unsupervised interventions for anxiety and depression (anxiety: 369 SMD=0.93 [95% CI: 0.74, 1.13], p<0.01; depression: SMD=1.18 [95% CI: 0.89, 1.47], 370 p<0.01), while moderate-to-large effects were observed during supervised interventions 371 (anxiety: SMD=0.62 [95% CI: 0.43, 0.81], p<0.01; depression: SMD=0.50 [95% CI: 0.34, 372 0.66], p<0.01). Neither the timing of the interventions (i.e., during or following treatment) 373 nor the intervention duration influenced the effect on outcomes, except in the case of 374 depression. Intervention duration had an effect on depression (γ^2 =7.93, df=1, p<0.01), with 375 interventions lasting longer than 12 weeks producing a large effect (SMD=0.84 [95% CI: 376 0.65, 1.03]; p<0.01) and interventions lasting 12 weeks or less having a moderate effect 377 (SMD=0.44 [95% CI: 0.23, 0.65]; p<0.01). 378

379

Sensitivity analyses

High quality trials: Results remained unchanged after performing meta-analyses with only high-quality trials, except for body mass index and waist circumference, for which the effect of exercise became smaller compared with results from meta-analyses using all available data. That is, exercise had no effect on body mass index (SMD=0.12 [95% CI: – 0.86, 0.73], p=0.87, I^2 =0%: low heterogeneity) and waist circumference (SMD= -0.07 [95%

CI: -0.09, 0.33], p=0.27, I^2 =2%; low heterogeneity) when analysis was restricted to including data only from high quality trials. *Trials with 100% of samples with stage II+ disease:* Compared with results from meta-analyses using all available data, effect sizes of exercise tended to be larger in trials involving only women with stage II+ breast cancer for QOL (0.78 vs. 0.40), fatigue (0.41 vs. 0.30) and depression (0.80 vs. 0.66).

390

391 Discussion

These findings suggest that exercise is safe, feasible and effective for improving health 392 outcomes among women with stage II+ breast cancer. More specifically, adverse events 393 reported as a consequence of participating in exercise during or following treatment for stage 394 II+ breast cancer were uncommon (occurring in <5% of women, Table 2). When adverse 395 events were reported, they were typically mild in nature and represented acute and normal 396 physiological adaptations to exercise. These results are similar to findings reported in 397 previous reviews and meta-analyses, which had underrepresentation of women with regional 398 and advanced breast cancer.²⁻⁴ Nonetheless, caution and care with exercise prescription 399 remains relevant because about one-third of studies (n=21) provided no comment on the 400 occurrence (or lack thereof) of adverse events. Studies that did report adverse events, mostly 401 did not comprehensively describe monitoring and recording procedures. Similar to our 402 findings, Speck *et al.*³ reported in their review of mixed-cancer types that only 44% (n=36) of 403 404 studies documented the presence or absence of adverse events, with 81% (n=29) of these studies reporting no harm as a result of exercise. These findings highlight the need for 405 standardised recording of adverse events to be incorporated into the design of RCTs. While 406 only a minimal amount of events that occurred in the exercise intervention group (5%) were 407 classified as severe (grade 3), these results nonetheless suggest a need for a thorough health 408

and medical history evaluation prior to exercise prescription, as well as individualised
exercise approaches and patient education to ensure that individuals can take appropriate
action, should an adverse event occur.

The safety findings were similar irrespective of the mode of exercise evaluated, the 412 degree of supervision provided, intervention duration and whether the intervention was 413 conducted during or following breast cancer treatment. However, caution is advised when 414 interpreting these results. For example, the exercise intensity of unsupervised interventions 415 416 was generally less vigorous compared with supervised exercise interventions. This difference in intensity may have been intentional, or it may suggest that individuals are more cautious 417 when exercising unsupervised. Also, compared with aerobic interventions, which were 418 mostly home-based walking programs, resistance exercise interventions were more 419 commonly performed at a supervised facility, involving specialised equipment (e.g., pin-420 loaded machines), instruction of technique and monitoring and progression of intensity (e.g., 421 progressing from 50 to 80% of 1RM). As such, paying particular attention to the provision of 422 safety information when prescribing unsupervised resistance-based exercise is paramount to 423 maintaining safety in this setting. Low withdrawal rates (approximately 11%) and high 424 adherence (approximately 80%) identified in this review suggest that exercise during and 425 following treatment for stage II+ breast cancer is highly feasible. These findings may in part 426 reflect recruitment bias (e.g., exercise readiness tends to be higher in those who agree to 427 participate in exercise trials compared with those who do not⁵). Alternately, the findings may 428 reflect the perceived or real physical and psychosocial benefit achieved through exercise 429 during the breast cancer survivorship period.¹ Specifically, the outcomes from this meta-430 analysis also demonstrated that for women with stage II+ breast cancer, exercise during and 431 following treatment led to improvements in QOL (SMD=0.4), fatigue (SMD=0.3), aerobic 432

433 fitness (SMD=0.6), upper-body strength (SMD=0.4), anxiety (SMD=0.8), depression
434 (SMD=0.7), waist circumference (SMD=0.2) and body mass index (SMD=0.2).

The magnitude of the effects reported here is similar to those reported in previous 435 reviews that likely overrepresented women with early-stage disease.^{2-4, 105-111} However, 436 greater effects of exercise were observed for depression and anxiety in this review. When 437 analyses were restricted to include data only from those studies involving all participants with 438 stage II+ disease, the effect was also higher for OOL, fatigue and depression. In contrast to 439 440 previous findings that showed larger effects of exercise when conducted following compared with during adjuvant treatment³, our findings showed similar effects irrespective of 441 intervention timing. It seems plausible that these differences are influenced by capacity for 442 change. That is, compared with those women with early-stage breast cancer, those with stage 443 II+ disease experience poorer health and greater morbidity (e.g., higher rates of depression 444 and anxiety are observed in women with more advanced disease compared with local disease 445 during and after treatment¹¹²). Women with more advanced breast cancer may therefore 446 experience greater benefits of exercise for improving their mental health and wellbeing. 447 Irrespective, the consistent message from findings reported here and that of others previously, 448 is that exercise is effective for preventing treatment-related morbidity and health declines, 449 and can be used to facilitate recovery post-treatment.¹¹³ 450

Exercise, irrespective of intervention characteristics, led to favourable effects, yet there was some evidence to suggest that the magnitude of effect differed for some outcomes depending on exercise mode, degree of supervision, timing (during versus following treatment) and duration of the intervention. For example, stronger effects for QOL were evident for supervised compared with unsupervised exercise, and when the intervention involved more than one exercise mode compared with only one mode. In contrast, greater benefits in psychological outcomes (anxiety and depression) occurred during unsupervised

458 interventions, compared with supervised interventions. Resistance exercise was more effective for improving strength compared with other modes of exercise, whereas 459 interventions that included aerobic exercise were more effective at improving fitness, anxiety 460 and depression. Finally, interventions lasting longer than 12 weeks produced larger effects on 461 depression than shorter interventions. This provides support for the important role of exercise 462 in longer term management of psychosocial wellbeing post-diagnosis. These findings also 463 support the notion that best clinical practice includes an exercise prescription that considers a 464 patient's physical and psychosocial needs, as well as their personal interests and preferences. 465

466 *Limitations*

Key limitations of this review include the poor reporting of adverse events by over 60% 467 of included studies, and the likelihood of a response bias. The mean age of the study 468 participants was 53 years, whereas the international average age of breast cancer diagnosis is 469 between 56 and 62 years.¹¹⁴ The samples included in this review were also likely healthier 470 compared with the wider breast cancer population. Most (79%; n=48) of the trials excluded 471 participants with various comorbidities, yet 90% of women with breast cancer report at least 472 one comorbidity.¹¹⁵ Consenting women were also likely to live in more urban environments 473 with easier access to care, and have a history of exercise participation. In contrast, 474 approximately 60% of the wider breast cancer population is sedentary or insufficiently active 475 at time of breast cancer diagnosis.¹¹⁶ Considering these limitations, we advise caution against 476 over-interpreting the results of this review. Another potential limitation of this review is the 477 inclusion of studies that involved women with early-stage, local disease. However, these 478 women represented less than 50% of the data. Further, findings from the sensitivity analyses 479 (which involved only RCTs with 100% of the sample being women with stage II+ disease) 480 were consistent with those findings when all studies were included. Finally, exercise effects 481 were examined based on immediately post-intervention results and the longest intervention 482

length was 1 year. As such, the longer term effects of exercise among women with stage II+
breast cancer remain unknown.

485 **Conclusions**

This review highlights the need for improved and standardised recording and 486 monitoring of adverse events, which is relevant in both clinical and research settings. Further, 487 demonstrating exercise that exercise is safe, feasible and effective in women with stage II+ 488 489 disease represents an important contribution to the literature. Future research will lead to greater understanding of the role of exercise with respect to survival outcomes, and will help 490 491 to refine optimal exercise prescription and the diagnosis, treatment, personal and behavioural characteristics that influence exercise safety, feasibility and effectiveness. Until this 492 information is available, the findings reported here indicate that most individuals with Stage 493 II+ breast cancer should be able to participate safely in exercise, according to established 494 general guidelines that are available and promoted to women with breast cancer. Specifically, 495 exercise should include mixed exercise modes (including aerobic- and resistance-based 496 exercise), and should be performed at moderate or higher intensities, three to five times per 497 week, for a total of at least 150 minutes per week of exercise.^{1, 117} 498

- 499 Acknowledgments: Nil.
- 500 **Funding:** The authors received no specific funding for this work.
- 501 **Disclosure:** The authors declare no conflicts of interest.

502 **Suppliers:** RevMan software (version 5.3); R statistical software (version 3.4.1).

503 List of Tables, Figures and Supplementary Files

504

Table 1: Summary of exercise intervention characteristics separated by exercise mode (n=61).

21

Table 2. Adverse events by grade of severity described for those in the exercise and usualcare groups.

Table 3. Study recruitment rate, withdrawal rate and exercise adherence by exercise mode,

509 treatment status, intervention supervision and intervention duration.

510

511 Figure 1. Meta-analysis of all grade 3 to 5 adverse events in exercise compared to usual care

512 presented as overall and separated by exercise mode, treatment status, intervention duration

513 and degree of supervision.

514 Figure 2. Meta-analyses results of aerobic fitness with subgroup analyses for exercise mode,

515 intervention supervision, timing and duration, and sensitivity analyses (positive SMD values516 favour exercise).

517 Figure 3. Meta-analyses results of quality of life with subgroup analyses for exercise mode,

518 intervention supervision, timing and duration and sensitivity analyses (positive SMD values

519 favour exercise).

520 Figure 4. Meta-analyses results of fatigue with subgroup analyses for exercise mode,

521 intervention supervision, timing and duration and sensitivity analyses (positive SMD values

522 favour exercise).

523

524 Supplementary Content 1: Systematic review flow diagram.

Supplementary Content 2: Ratings of all studies included in systematic review using thePEDro scale (n=61).

- 527 Supplementary Digital 3: Overview of samples and exercise details of included studies
 528 (n=61)
- 529 Supplementary Digital 4: Summary of study recruitment, retention, adherence, reasons for
- 530 withdrawal, intervention settings and supervision and exercise related events (n=61)
- 531 Supplementary Content 5. Overview reasons for withdrawals across all trials (n=61).
- 532 Supplementary Content 6. Overview of health outcomes and methods of assessment across533 all trials (n=61).
- 534 Supplementary Content 7. Meta-analyses results of anxiety with subgroup analyses for
- 535 exercise mode, intervention supervision, timing and duration and sensitivity analyses
- 536 (positive SMD values favour exercise).
- 537 Supplementary Content 8. Meta-analyses results of depression with subgroup analyses for
- 538 exercise mode, intervention supervision, timing and duration and sensitivity analyses
- 539 (positive SMD values favour exercise).
- 540 Supplementary Content 9. Meta-analyses results of upper-body strength with subgroup
- 541 analyses for exercise mode, intervention supervision, timing and duration and sensitivity
- 542 analyses (positive SMD values favour exercise).
- 543 Supplementary Digital 10. Meta-analyses results of waist circumference with subgroup
- analyses for exercise mode, intervention supervision, timing and duration and sensitivity
- 545 analyses (positive SMD values favour exercise).
- 546 Supplementary Digital 11. Meta-analyses results of body weight with subgroup analyses for
- 547 exercise mode, intervention supervision, timing and duration and sensitivity analyses
- 548 (positive SMD values favour exercise).

- 549 Supplementary Digital 12. Meta-analyses results of body mass index with subgroup analyses
- 550 for exercise mode, intervention supervision, timing and duration and sensitivity analyses
- 551 (positive SMD values favour exercise).
- 552 Supplementary Digital 13. Meta-analyses results of body fat with subgroup analyses for
- exercise mode, intervention supervision, timing and duration and sensitivity analyses
- 554 (positive SMD values favour exercise).

ANA ANA

556

557	1. Hayes SC, Spence RR, Galvão DA, Newton RU. Australian Association for Exercise and
558	Sport Science position stand: optimising cancer outcomes through exercise. J Sci Med Sport
559	2009;12(4):428-434.
560	2. Cheema B, Gaul CA, Lane K, Fiatarone Singh MA. Progressive resistance training in
561	breast cancer: a systematic review of clinical trials. Breast Cancer Res Treat 2008; 109(1):9-
562	26.
563	3. Speck RM, Courneya KS, Mâsse LC, Duval S, Schmitz KH. An update of controlled
564	physical activity trials in cancer survivors: a systematic review and meta-analysis. J Cancer
565	Surviv 2010;4(2):87-100.
566	4. Tian L, Lu HJ, Lin L, Hu Y. Effects of aerobic exercise on cancer-related fatigue: a meta-
567	analysis of randomized controlled trials. Support Care Cancer 2016. 24(2): p. 969-83.
568	5. Spence RR, Heesch KC, Brown WJ. Exercise and cancer rehabilitation: a systematic
569	review. Cancer Treat Rev 2010;36(2):185-94.
570	6. Schmitz KH, Holtzman J, Courneya KS, Mâsse LC, Duval S, Kane R. Controlled Physical
571	Activity Trials in Cancer Survivors: A Systematic Review and Meta-analysis. Cancer
572	Epidemiol Biomarkers Prev 2005;14(7):1588-1595.
573	7. Cormie P, Zopf EM, Zhang X, Schmitz KH. The Impact of Exercise on Cancer Mortality,
574	Recurrence, and Treatment-Related Adverse Effects. <i>Epidemiol Rev</i> 2017:39(1):1-22.
575	8. Baumann, FT, Physical exercise programs following cancer treatment. Eur Rev Aging Phys
576	Act 2013;10(1):57-59.

- 577 9. Hayes SC, Rye S, Disipio T, et al. Exercise for health: a randomized, controlled trial
- 578 evaluating the impact of a pragmatic, translational exercise intervention on the quality of life,
- 579 function and treatment-related side effects following breast cancer. Breast Cancer Res Treat

580 2013;137(1):175-186.

- 581 10. Courneya KS. Physical activity and cancer survivorship: a simple framework for a
- 582 complex field. *Exerc Sport Sci Rev* 2014;42(3):102-109.
- 11. Irwin ML. Physical activity interventions for cancer survivors. *Br J Sports Med*2009;43(1):32-38.
- 585 12. Ibrahim EM, Al-Homaidh A, Physical activity and survival after breast cancer diagnosis:
- meta-analysis of published studies. *Medical Oncology* 2011;28(3):753-765.
- 587 13. Spence RR, Di Sipio T, Schmitz K and Hayes SC. Is unsupervised exercise following
- 588 breast cancer safe for all women? *Int J Phys Med Rehabil* 2014;2(197):1-8.
- 589 14. Maddocks M, Mockett S, Wilcock A. Is exercise an acceptable and practical therapy for
- people with or cured of cancer? A systematic review. *Cancer Treat Rev* 2009;35(4):383-90.
- 591 15. Australian Institute of Health and Welfare and National Breast and Ovarian Cancer
- 592 Centre. Breast cancer in Australia: an overview. Canberra. Australian Institute of Health and593 Welfare; 2009.
- 16. American Cancer Society. Cancer Treatment and Survivorship Facts & Figures 20142015. Atlanta: American Cancer Society; 2014.
- 596 17. Oldervoll LM, Loge JH, Paltiel H, Asp MB, Vidvei U, Hjermstad MJ, Kaasa S. Are
- 597 palliative cancer patients willing and able to participate in a physical exercise program?

598 *Palliat Support Care* 2005;3(4):281-287.

18. Dixon JM., ABC of breast diseases. 4th ed. Chichester, West Sussex: Blackwell; 2012.

- 19. Brown P, Clark MM, Atherton P, et al. Will improvement in quality of life (QOL) impact
- 601 fatigue in patients receiving radiation therapy for advanced cancer? *Am J Clin Oncol*
- 602 2006;29(1):52-58.
- 603 20. Cheville AL, Troxel AB, Basford JR, Kornblith AB. Prevalence and treatment patterns of
- 604 physical impairments in patients with metastatic breast cancer. J Clin Oncol
- 605 2008;26(16):2621-9.
- 606 21. Cancer Australia. Report to the nation breast cancer 2012. Surry Hills, NSW:
- 607 Australian Government; 2012.
- 608 22. American Cancer Society. Breast Cancer Survival Rates. Atlanta: American Cancer
- 609 Society; 2014.
- 610 23. Schardt C, Adams MB, Owens T, Keitz S, Fontelo P. Utilization of the PICO framework
- 611 to improve searching PubMed for clinical questions. BMC Med Inform Decis Mak

612 2007;7(1):16.

- 613 24. Howley ET. Type of activity: Resistance, aerobic and leisure versus occupational
- 614 physical activity. *Med Sci Sports Exerc* 2001;33(6):S364-S369.
- 615 25. McNeely ML, Campbell KL, Rowe BH, Klassen TP, Mackey JR, Courneya KS. Effects
- of exercise on breast cancer patients and survivors: a systematic review and meta-analysis.
- 617 *CMAJ* 2006;175(1):34-41.
- 618 26. National Cancer Institute. *Common Terminology Criteria for Adverse Events (CTCAE)*
- 619 *version 4.0.* Bethesda: Maryland: National Institutes of Health; 2009.
- 620 27. Australian Government Department of Health and Ageing. *The Australian Clinical Trial*
- 621 Handbook. A simple, practical guide to the conduct of clinical trials to International

- 622 standards of Good Clinical Practice (GCP) in the Australian context.
- 623 *Canberra:Commonwealth of Australia;* 2006.
- 624 28. Wayne PM, Berkowitz DL, Litrownik DE, Buring JE, Yeh GY. What do we really know
- about the safety of tai chi?: A systematic review of adverse event reports in randomized trials.
- 626 *Arch Phys Med Rehabil* 2014; 95(12):2470-83.
- 627 29. Maher CG, Sherrington C, Herbert RD, Moseley AM, Elkins M. Reliability of the PEDro
- 628 Scale for Rating Quality of Randomized Controlled Trials. *Phys Ther* 2003;83(8):713-721.
- 629 30. Sherrington C, Herbert RD, Maher CG, Moseley AM. PEDro. A database of randomized
- trials and systematic reviews in physiotherapy. *Man Ther* 2000;5(4):223-226.
- 631 31. Armijo-Olivo S. da Costa BR, Cummings GG, Ha C, Fuentes J, Saltaji H, Egger M.
- 632 PEDro or Cochrane to Assess the Quality of Clinical Trials? A Meta-Epidemiological Study.
- 633 *PLOS ONE* 2015;10(7):e0132634.
- 634 32. Harrison RW, Hasselblad V, Mehta RH, Levin R, Harrington RA, Alexander JH. Effect
- of levosimendan on survival and adverse events after cardiac surgery: a meta-analysis. J *Cardiothorac Vasc Anesth* 2013;27(6):1224-32.
- 637 33. Hasselblad V, Mosteller F, Littenberg B, et al. A survey of current problems in meta-
- analysis. Discussion from the Agency for Health Care Policy and Research inter-PORT Work
- 639 Group on Literature Review/Meta-Analysis. *Med Care* 1995;33(2):202-20.
- 640 34. Lewis PB, Ruby D, Bush-Joseph CA, Muscle Soreness and Delayed-Onset Muscle
- 641 Soreness. *Clin Sports Med* 2012;31(2):255-262.
- 642 35. Riebe D, Ehrman JK, Liguori G, Meir M. ACSM's guidelines for exercise testing and
- 643 prescription. 10th ed. Philadelphia, PA: Wolters Kluwer Health; 2018.

- 36. De Jesus S, Fitzgeorge L, Unsworth K, et al. Feasibility of an exercise intervention for
 fatigued breast cancer patients at a community-based cardiac rehabilitation program. *Cancer Manag Res* 2017;9:29-39.
- 647 37. Newton MJ, Hayes SC, Janda M, et al. Safety, feasibility and effects of an individualised
- 648 walking intervention for women undergoing chemotherapy for ovarian cancer: A pilot study.
- 649 *BMC Cancer* 2011;11(1):389-389.
- 650 38. Valkenet K, Trappenburg JC, Schippers CC, et al. Feasibility of Exercise Training in
- 651 Cancer Patients Scheduled for Elective Gastrointestinal Surgery. *Dig Surg* 201633(5):439-
- **652** 447.
- 653 39. Coats V, Maltais F, Simard S, et al. Feasibility and effectiveness of a home-based
- exercise training program before lung resection surgery. *Can Respir J* 2013;20(2):e10-e16.
- 40. Higgins JPT, Deeks JJ, Altman DG. Special Topics in Statistics, in Cochrane Handbook
- 656 for Systematic Reviews of Interventions. John Wiley & Sons, Ltd; 2008: 481-529.
- 41. Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a
 simple, graphical test. *BMJ* 1997;315(7109):629-34.
- 42. Thompson G, Higgins JP, How should meta-regression analyses be undertaken and
 interpreted? *Stat Med* 2002;21(11):1559-73.
- 43. Deeks JJ, Higgins JPT, Altman DG. Analysing Data and Undertaking Meta-Analyses, in
 Cochrane Handbook for Systematic Reviews of Interventions. John Wiley & Sons, Ltd; 2008:
 243-296.
- 44. Cohen J. Statistical Power Analysis for the Behavioral Sciences. 2nd ed. Hillsdale (NJ):
- 665 Lawrence Erlbaum Associates; 1988.

- 45. Guinan EM, Connolly EM, Hussey J, Exercise training in breast cancer survivors: a
- review of trials examining anthropometric and obesity-related biomarkers of breast cancer
- risk. *Phys Ther Rev* 2013;18(2):79-89.46. Cormie, P., et al., *Is it safe and efficacious for*
- 669 women with lymphedema secondary to breast cancer to lift heavy weights during exercise: a
- 670 *randomised controlled trial.* Journal of cancer survivorship : research and practice, 2013.
- 671 **7**(3): p. 413-424.
- 46. Cormie P, Pumpa K, Galvão DA, Turner E, Spry N, Saunders C, Zissiadis Y, Newton

673 RU. Is it safe and efficacious for women with lymphedema secondary to breast cancer to lift

heavy weights during exercise: a randomised controlled trial. J Cancer Surviv 2013;7(3):413-

675 424.

- 47. Dolan LB, Campbell K, Gelmon K, Neil-Sztramko S, Holmes D, McKenzie DC, Interval
 versus continuous aerobic exercise training in breast cancer survivors--a pilot RCT. *Support Care Cancer* 2016;24(1):119-27.
- 48. Kim CJ, Kang DH, Smith BA, Landers KA. Cardiopulmonary responses and adherence to
 exercise in women newly diagnosed with breast cancer undergoing adjuvant therapy. *Cancer Nurs* 2006; 29(2):156-65.
- 49. Winters-Stone KM, Dobek J, Nail L, et al. Strength training stops bone loss and builds
 muscle in postmenopausal breast cancer survivors: a randomized, controlled trial. *Breast Cancer Res Treat* 2011;127(2):447-56.
- 50. Gokal K. Wallis D, Ahmed S, Boiangiu I, Kancherla K, Munir F. Effects of a self-
- 686 managed home-based walking intervention on psychosocial health outcomes for breast cancer
- 687 patients receiving chemotherapy: a randomised controlled trial. *Support Care Cancer*
- 688 2016;24(3):1139-66.

689	51. Headley, J.A., K.K. Ownby, and L.D. John, The effect of seated exercise on fatigue and
690	quality of life in women with advanced breast cancer. Oncol Nurs Forum 2004;31(5):977-
691	983.

- 692 52. Hornsby WE, Douglas PS, West MJ, et al. Safety and efficacy of aerobic training in
- 693 operable breast cancer patients receiving neoadjuvant chemotherapy: a phase II randomized
- 694 trial. *Acta Oncol* 2014;53(1):65-74.
- 53. Ligibel JA, Giobbie-Hurder A, Shockro L, et al. Randomized trial of a physical activity
- intervention in women with metastatic breast cancer. *Cancer* 2016;122(8):1169-1177.
- 697 54. Macvicar MG, Winningham ML, Nickel JL. Effects of Aerobic Interval Training on
- 698 Cancer Patients' Functional Capacity. *Nursing Research* 1989;38(6): 348-353.
- 699 55. Portela ALM, Santaella CL, Gómez CC, Burch A. Feasibility of an Exercise Program for
- Puerto Rican Women who are Breast Cancer Survivors. *Rehabil Oncol* 2008;26(2):20-31.
- 56. Raghavendra RM, Nagarathna R, Nagendra HR, et al. Effects of an integrated yoga
- 702 programme on chemotherapy-induced nausea and emesis in breast cancer patients. *Eur J*
- 703 *Cancer Care* 2007;16(6):462-74.
- 57. Rao R, Cruz V, Peng Y, et al. Bootcamp during neoadjuvant chemotherapy for breast
 cancer: a randomized pilot trial. *Breast Cancer* 2012;6:39-46.
- 58. Rao RM, Raghuram N, Nagendra HR, et al. Effects of an integrated Yoga Program on
- 707 Self-reported Depression Scores in Breast Cancer Patients Undergoing Conventional
- Treatment: A Randomized Controlled Trial. *Indian J Palliat Care* 2015;21(2):174-181.
- 59. Campbell KL, Kam JW, Neil-Sztramko SE, et al. Effect of aerobic exercise on cancer-
- associated cognitive impairment: A proof-of-concept RCT. *Psychooncology* 2017:1-8.

- 711 60. Courneya KS. Mackey JR, Bell GJ, Jones LW, Field CJ, Fairey AS. Randomized
- 712 controlled trial of exercise training in postmenopausal breast cancer survivors:
- cardiopulmonary and quality of life outcomes. *J Clin Oncol* 2003;21(9):1660-8.
- 61. Drouin JS, Young TJ, Beeler J, et al. Effects of Aerobic Exercise Training on Peak
- 715 Aerobic Capacity, Fatigue, and Psychological Factors During Radiation for Breast Cancer.
- 716 *Rehabil Oncol* 2005;23(1):11-17.
- 62. Murtezani A, Ibraimi Z, Bakalli A, Krasniqi S, Disha ED, Kurtishi I. The effect of
- 718 aerobic exercise on quality of life among breast cancer survivors: a randomized controlled
- 719 trial. J Cancer Res Ther 2014;10(3):658-64.
- 720 63. Naraphong W. Exercise intervention for fatigue-related symptoms in Thai women with
- 721 breast cancer: A pilot study., Exercise intervention for fatigue-related symptoms in Thai
- women with breast cancer: A pilot study. *Nurs Health Sci* 2014;17(1):33-41.
- 723 64. Pinto BM, Papandonatos GD, Goldstein MG. A randomized trial to promote physical
- activity among breast cancer patients. *Health Psychol* 2013;32(6):616-26.
- 725 65. Pinto BM, Stein K, Dunsiger S. Peers Promoting Physical Activity Among Breast Cancer
- 726 Survivors: A Randomized Controlled Trial. *Health Psychol.* 2015;34(5):463-472.
- 727 66. Rogers LQ, Hopkins-Price P, Vicari S, et al. A randomized trial to increase physical
- activity in breast cancer survivors. *Med Sci Sports Exerc* 2009;41(4):935-46.
- 729 67. Vallance JK, Courneya KS, Plotnikoff RC, Yasui Y, Mackey JR. Randomized Controlled
- 730 Trial of the Effects of Print Materials and Step Pedometers on Physical Activity and Quality
- of Life in Breast Cancer Survivors. *J Clin Oncol* 2007;25(17):2352-2359.

- 68. Wang YJ, Boehmke M, Wu YW, Dickerson SS, Fisher N. Effects of a 6-week walking
- program on Taiwanese women newly diagnosed with early-stage breast cancer. *Cancer Nurs*2011;34(2):E1-13.
- 69. Yang CY, Tsai JC, Huang YC, Lin CC. Effects of a home-based walking program on
- 736 perceived symptom and mood status in postoperative breast cancer women receiving
- 737 adjuvant chemotherapy. *J Adv Nurs* 2011;67(1):158-68.
- 738 70. Pinto BM, Frierson GM, Rabin C, Trunzo JJ, Marcus BH. Home-based physical activity
 739 intervention for breast cancer patients. *J Clin Oncol* 2005;23(15):3577-87.
- 740 71. Vallance JK, Friedenreich CM, Lavallee CM, et al. Exploring the Feasibility of a Broad-
- 741 Reach Physical Activity Behavior Change Intervention for Women Receiving Chemotherapy
- for Breast Cancer: A Randomized Trial. *Cancer Epidemiol Biomarkers Prev* 2016;25(2):391-

743 8.

- 744 72. Eakin EG, Lawler SP, Winkler EA, Hayes SC. A randomized trial of a telephone-
- 745 delivered exercise intervention for non-urban dwelling women newly diagnosed with breast
- cancer: exercise for health. *Ann Behav Med* 2012;43(2):229-38.
- 747 73. Anderson RT, Kimmick GG, McCoy TP, et al. A randomized trial of exercise on well748 being and function following breast cancer surgery: the RESTORE trial. *J Cancer Surviv*749 2012;6(2):172-181.
- 74. Cantarero-Villanueva I, Fernández-Lao C, Del Moral-Avila R, Fernández-de-Las-Peñas
 C, Feriche-Fernández-Castanys MB, Arroyo-Morales M. Effectiveness of core stability
 exercises and recovery myofascial release massage on fatigue in breast cancer survivors: a
- randomized controlled clinical trial. *Evid Based Complement Alternat Med* 2012:620619.

- 754 75. Cornette T, Vincent F, Mandigout S, et al. Effects of home-based exercise training on
- VO2 in breast cancer patients under adjuvant or neoadjuvant chemotherapy (SAPA): a
- randomized controlled trial. *Eur J Phys Rehabil Med* 2016;52(2):223-32.
- 757 76. De Luca V, Minganti C, Borrione P, et al. Effects of concurrent aerobic and strength
- training on breast cancer survivors: a pilot study. *Public Health* 2016;136:126-32.
- 759 77. Dethlefsen C, Lillelund C, Midtgaard J, et al. Exercise regulates breast cancer cell
- viability: systemic training adaptations versus acute exercise responses. Breast Cancer Res
- 761 *Treat* 2016;159(3):469-79.
- 762 78. Fernández-Lao C, Cantarero-Villanueva I, Ariza-Garcia A, et al. Water versus land-based
- 763 multimodal exercise program effects on body composition in breast cancer survivors: a
- real controlled clinical trial. *Support Care Cancer* 2013;21(2):521-530.
- 765 79. Hatchett A, Hallam JS, Ford MA. Evaluation of a social cognitive theory-based email
- intervention designed to influence the physical activity of survivors of breast cancer.
- 767 *Psychooncology* 2013;22(4):829-836.
- 768 80. Herrero F, San Juan AF, Fleck SJ, et al. Combined aerobic and resistance training in
- 769 breast cancer survivors: A randomized, controlled pilot trial. Int J Sports Med
- 770 2006;27(7):573-80.
- 81. Husebø AML, Dyrstad SM, Mjaaland I, Søreide JA, Bru E. Effects of Scheduled Exercise
- on Cancer-Related Fatigue in Women with Early Breast Cancer. *Sci World J* 2014:9.
- 82. Hutnick NA, Williams NI, Kraemer WJ, et al. Exercise and lymphocyte activation
- following chemotherapy for breast cancer. *Med Sci Sports Exerc*. 2005;37(11):1827-35.

- 83. Ligibel JA. Campbell N, Partridge A, et al. Impact of a mixed strength and endurance
 exercise intervention on insulin levels in breast cancer survivors. *J Clin Oncol*2008;26(6):907-12.
- 84. Maryam A, Fazlollah A, Eesa M, Ebrahim H, Abbas VF. The effect of designed exercise
- programme on quality of life in women with breast cancer receiving chemotherapy. *Scand J Caring Sci* 2010;24(2):251-8.
- 781 85. Milne HM, Wallman KE, Gordon S, Courneya KS. Effects of a combined aerobic and

resistance exercise program in breast cancer survivors: a randomized controlled trial. Breast

783 *Cancer Res Treat* 2008; 108(2):279-88.

784 86. Naumann F. Munro A, Martin E, et al. An individual-based versus group-based exercise

and counselling intervention for improving quality of life in breast cancer survivors. A

feasibility and efficacy study. *Psychooncology* 2012;21(10):1136-9.

- 787 87. van Waart H, Stuiver MM, van Harten WH, et al. Effect of Low-Intensity Physical
- 788 Activity and Moderate- to High-Intensity Physical Exercise During Adjuvant Chemotherapy
- on Physical Fitness, Fatigue, and Chemotherapy Completion Rates: Results of the PACES

Randomized Clinical Trial. J Clin Oncol 2015;33(17):1918-1927.

- 88. Galiano-Castillo N, Cantarero-Villanueva I, Fernández-Lao C, et al. Telehealth system: A
 randomized controlled trial evaluating the impact of an internet-based exercise intervention
 on quality of life, pain, muscle strength, and fatigue in breast cancer survivors. *Cancer*2016;122(20):3166-3174.
- 89. Short CE, James EL, Girgis A, D'Souza MI, Plotnikoff RC. Main outcomes of the Move
- 796 More for Life Trial: a randomised controlled trial examining the effects of tailored-print and
- targeted-print materials for promoting physical activity among post-treatment breast cancer

798 survivors. *Psychooncology* 2015;24(7):771-778.

- 90. Ahmed RL, Thomas W, Yee D, Schmitz KH. Randomized controlled trial of weight
- training and lymphedema in breast cancer survivors. *J Clin Oncol* 2006;24(18):2765-2772.
- 801 91. Kilbreath SL, Refshauge KM, Beith JM, et al. Upper limb progressive resistance training
- and stretching exercises following surgery for early breast cancer: a randomized controlled
- 803 trial. Breast Cancer Res Treat 2012;133(2):667-676.
- 804 92. Schmidt ME, Wiskemann J, Armbrust P, et al. Effects of resistance exercise on fatigue
- and quality of life in breast cancer patients undergoing adjuvant chemotherapy: A
- randomized controlled trial. *Int J Cancer* 2015;137(2):471-80.
- 93. Winters-Stone KM, Dobek J, Nail LM, Bennett JA, Leo MC, Torgrimson-Ojerio B, Luoh
- 808 SW, Schwartz A. Impact + resistance training improves bone health and body composition in
- prematurely menopausal breast cancer survivors: a randomized controlled trial. *Osteoporos Int* 2013;24(5):1637-46.
- 811 94. Banerjee B, Vadiraj HS, Ram A, et al. Effects of an integrated yoga program in
- 812 modulating psychological stress and radiation-induced genotoxic stress in breast cancer
- patients undergoing radiotherapy. *Integr Cancer Ther* 2007;6(3):242-50.
- 814 95. Cantarero-Villanueva I, Fernández-Lao C, Caro-Morán E, et al. Aquatic exercise in a
- 815 chest-high pool for hormone therapy-induced arthralgia in breast cancer survivors: a
- 816 pragmatic controlled trial. *Clin Rehabil* 2013;27(2):123-32.
- 817 96. Cantarero-Villanueva I, Fernández-Lao C, Cuesta-Vargas AI, et al. The effectiveness of a
- 818 deep water aquatic exercise program in cancer-related fatigue in breast cancer survivors: a
- 819 randomized controlled trial. *Arch Phys Med Rehabil* 2013;94(2):221-30.
- 820 97. Cantarero-Villanueva I, Fernández-Lao C, Fernández-de-Las-Peñas C, et al. Effectiveness
- of water physical therapy on pain, pressure pain sensitivity, and myofascial trigger points in

- breast cancer survivors: a randomized, controlled clinical trial. *Pain Med* 2012;13(11):150919.
- 824 98. Chandwani KD, Perkins G, Nagendra HR, et al. Randomized, Controlled Trial of Yoga in
- 825 Women with Breast Cancer Undergoing Radiotherapy. *J Clin Oncol* 2014;32(10):1058-1065.
- 826 99. Danhauer SC, Mihalko SL, Russell GB, et al. Restorative yoga for women with breast
- 827 cancer: findings from a randomized pilot study. *Psychooncology* 2009;18(4):360-8.
- 828 100. Loudon A, Barnett T, Piller N, Immink MA, Williams AD. Yoga management of breast

829 cancer-related lymphoedema: a randomised controlled pilot-trial. *BMC Complement Altern*

- *Med* 2013;14(1):214-214.
- 101. Moadel AB, Shah C, Wylie-Rosett J, et al. Randomized controlled trial of yoga among a

832 multiethnic sample of breast cancer patients: effects on quality of life. *J Clin Oncol*

833 2007;25(28):4387-95.

102. Courneya KS, Segal RJ, Mackey JR, et al. Effects of aerobic and resistance exercise in

breast cancer patients receiving adjuvant chemotherapy: a multicenter randomized controlled
trial. *J Clin Oncol* 2007;25(28):4396-404.

- 837 103. Musanti R. A study of exercise modality and physical self-esteem in breast cancer
 838 survivors. *Med Sci Sports Exerc* 2012;44(2):352-61.
- 839 104. Schwartz AL, Winters-Stone K, Gallucci B. Exercise effects on bone mineral density in
- 840 women with breast cancer receiving adjuvant chemotherapy. Oncol Nurs Forum
- 841 2007;34(3):627-33.
- 105. Zeng Y, Huang M, Cheng AS, Zhou Y, So WK. Meta-analysis of the effects of exercise
- intervention on quality of life in breast cancer survivors. *Breast Cancer* 2014;21(3):262-74.

- 844 106. Duijts SF, Faber MM, Oldenburg HS, van Beurden M, Aaronson NK. Effectiveness of
- behavioral techniques and physical exercise on psychosocial functioning and health-related
- quality of life in breast cancer patients and survivors--a meta-analysis. *Psychooncology*

847 2011;20(2):115-26.

- 848 107. Lipsett A, Barrett S, Haruna F, Mustian K, O'Donovan A. The impact of exercise during
- 849 adjuvant radiotherapy for breast cancer on fatigue and quality of life: A systematic review
- and meta-analysis. *Breast* 2017; 32:144-155.
- 108. Carayol M, Bernard P, Boiché J, et al. Psychological effect of exercise in women with
- breast cancer receiving adjuvant therapy: what is the optimal dose needed? Ann Oncol
- 853 2013;24(2):291-300.
- 109. Juvet LK, Thune I, Elvsaas IKØ, et al. The effect of exercise on fatigue and physical
- 855 functioning in breast cancer patients during and after treatment and at 6 months follow-up: A
- 856 meta-analysis. Breast 2017;33:166-177.
- 110. van Vulpen JK, Peeters PH, Velthuis MJ, van der Wall E, May AM. Effects of physical
- 858 exercise during adjuvant breast cancer treatment on physical and psychosocial dimensions of
- 859 cancer-related fatigue: A meta-analysis. *Maturitas* 2016;85:104-11.
- 860 111. Kim C, Kang D, Park J, A meta-analysis of aerobic exercise interventions for women
 861 with breast cancer. *West J Nurs Res* 2009;31(4):437-461.
- 862 112. Massie MJ, Prevalence of depression in patients with cancer. *J Natl Cancer Inst Monogr*863 2004;32: 57-71.
- 864 113. Mishra SI, Scherer RW, Snyder C, et al. Exercise interventions on health-related quality
- of life for people with cancer during active treatment. *Cochrane Database Syst Rev*,
- 866 2012;8:Cd008465.

- 867 114. McCaskill-Stevens W, Abrams JS. Comorbidities in the aging breast cancer population:
- 868 are current assessments leading to improved outcomes? J Natl Cancer Inst
- 869 2011;103(14):1072-3.
- 870 115. Schmitz KH, Speck RM, Rye SA, DiSipio T, Hayes SC. Prevalence of breast cancer
- treatment sequelae over 6 years of follow-up: The Pulling Through Study. Cancer

872 2012;118(8):2217-25.

- 873 116. Branstrom, R, Petersson LM, Saboonchi F, et al. Physical activity following a breast
- 874 cancer diagnosis: Implications for self-rated health and cancer-related symptoms. *Eur J*
- 875 *Oncol Nurs* 2015:19(6):680-5.
- 876 117. Schmitz KH, Courneya KS, Matthews C, et al. American College of Sports Medicine
- 877 roundtable on exercise guidelines for cancer survivors. *Med Sci Sports Exerc*
- 878 2010;42(7):1409-1426.

879

Table 1: Summary of exercise intervention characteristics separated by exercise mode (n=61).

		0
	Intervention details	
erobic exercise studies (
Mode	Continuous and interval training: cycle ergometer, outdoor cycling, elliptical trainer, treadmill, brisk walking	, jogging, rowing ergometer, stair-climbing machine (stair-master).
Intensity	40-85 HR _{max} ; 60-80% of age-adjusted HR _{max} ; 35-85% HRR; up to 90% of the HR reached in the 6MWT; 60	-80% VO _{2max} ; 55–100% VO _{2peak} (Interval training: <2 min intervals at
~	>80 % VO _{2peak}); 12–14 RPE (6–20 scale); 4–6 RPE (0–10 Scale, "breathing hard but able to talk"); \leq 3–6 ME	TS.
Session duration	15–60 minutes' overall duration (5–10 min warm-up and cool-down).	
Frequency	2–7 sessions per week (range 6 weeks to 1 year).	
Supervision	Supervised interventions ² : 2–3 supervised sessions per week.	
	Unsupervised interventions ³ : 2–5 unsupervised sessions per week; weekly supervised exercise sessions and/o guidebook; weekly to fortnightly in-person and telephone physical activity counselling sessions; tailored-phy	
esistance exercise studie		sical activity print-materials, physical activity booklet.
Mode	Resistance machines, free weights (dumbbells and barbells), weighted vests, resistance-bands. Exercises inclu	ided upper and lower body exercises targeting all major muscle groups
Widde	(e.g., squat, lunge, leg extension, leg curl, leg press, calf raises, chest press, seated row, triceps extension, bic	
	raise, shoulder flexion, hip flexion, hip extension, abdominal crunches, lower back hyperextensions and 2 for	
Intensity	50-85% of 1RM; 15 RPE (6–20 scale), $3-8$ RPE (0–10 scale); $0-10%$ of body weight (weighted vests).	ted jumps with weighted (ests).
2	6–20 repetitions per set.	
	1–3 sets per exercise.	
Session duration	30–60 min overall session duration (5–10 min warm-up and cool-down).	
Frequency	1–4 sessions per week (range 4 weeks to 12 months).	
Supervision	Supervised interventions ² : In-person supervision 2–3 sessions per week; 1 unsupervised session per week.	
	Unsupervised interventions ³ : 2-4 unsupervised sessions per week; 1 supervised session per week; exercise in	struction guidebook.
ombined exercise studie		
Mode	Aerobic-based, resistance-based, circuit training (including pump class and boot camp style training), stretchi	
	Pilates, hydrotherapy and patient-specific rehabilitation performed either on separate days or in combination session).	
	Aerobic exercise: Same as aerobic exercise studies plus aerobics classes, running, hiking, Nordic walking, flo	
	boat rowing, mini-trampoline, step-up blocks, antigravity treadmill, floor-based aerobic exercise to music, jun	
	Resistance exercise: Same as resistance exercise studies plus Flexband exercises, water-based resistance exer	
	exercises using steps and balls, small soft ball, mats, fit-balls and bodyweight exercises. Upper- and lower bo	dy exercises targeting all major muscle groups including lower back and
Intensity	abdominals. Exercises targeting all major muscle groups. Aerobic exercise:	Resistance exercise:
intensity	$12-16 \text{ RPE} (6-20 \text{ scale}), 7-8 \text{ RPE} (0-10 \text{ RPE scale}); 40-75\% \text{ VO}_{2\text{max}}; 55-75\% \text{ VO}_{2\text{peak}}; 55-85\% \text{ HR}_{\text{max}}$	40–90% 1RM; 13–15 RPE (6–20 scale), 4–7 RPE (0–10 scale).
	(intervals ranged from 30 secs [100% HR _{max})] to 6 mins [90%–95% of HR _{max}]; 40–65% HRR; 3-6 METS;	6-20 repetitions per set.
	60 RPM (cycle ergometer); \geq 50% bodyweight (antigravity treadmill).	1–4 sets per exercise.
Session duration	15–90 min overall session duration (5–10 min warm-up and cool-down).	
Frequency	1 to 7 sessions per week (range 6 weeks to 12 months).	
Supervision	Supervised interventions ² : In-person supervised 1–3 sessions per week; unsupervised sessions 2–3 times per	week.
	Unsupervised interventions ³ : 1–7 unsupervised sessions per week; weekly to monthly supervised sessions; w	eekly to monthly telephone calls, instructional exercise videos; weekly
	email messages; internet-based support (telehealth).	
ther exercise (n=11)		
Mode	Yoga (n=5): Stretching and isometric floor-based and standing exercises, whole-body postures, breathing exer	rcises, meditation and relaxation techniques and post-operative shoulder
	mobility exercises involving use of a mat, bolsters, chairs, blankets, blocks, and $a \le 1$ kg hand weights.	mining of stratching throughing maning and an ability and it.
	Pilates (n=1): Whole-body, standing, seated and floor based-exercises (including pelvic floor exercises) compression and shared and standing movements.	brising of stretching, breathing, running and mobility exercises using
	resistance bands, foam rollers, \leq 1kg hand weights; floor-based, seated and standing movements.	

	Hydrotherapy (n=3): Various whole body aerobic-based, strength-based, mobility and stretching movements targeting all major muscle groups including running in water and
	swimming, forward and backward jogging with arms pushing, pulling and pressing, leaps, leg crossovers and movements using pool noodles, swimming boards, and swimming belts.
	Seated exercise (n=1): Stretching and repeated flexion and extension of the arms, head, upper-torso, and legs while seated.
	Nia exercise (n=1): Aerobic-based and whole-body conditioning program that integrates strength, flexibility, mobility, agility, and stability exercises incorporating martial arts, dance
	and yoga style movements.
Intensity	Yoga: Low-moderate, low-impact and gentle stretching and postures; moderate (<12 RPE); Individual poses were held from 20 seconds to 5 minute.
	Pilates: Low-moderate.
	Hydrotherapy: Aerobic-based components performed at 60% HR _{max} and strength-based components performed for 2–3 sets of 8–12 repetitions.
	Seated exercise: Low-moderate.
Session duration	15–120 min overall session duration (including 5-10 min warm up, cool down and stretching).
Frequency	1–7 sessions per week for 4 weeks up to 24 weeks.
Supervision	Supervised interventions ² : In-person supervision 1–3 sessions per week.
•	Unsupervised interventions ³ : 1–7 sessions per week unsupervised; weekly supervised sessions; instructional exercise videos and audiotapes.
Studies involving separate	e aerobic and resistance exercise arms (n=3)
* *	See aerobic and resistance exercise studies for details

¹ N=3 additional trials involved separate aerobic and resistance exercise arms,

² Supervised were interventions whereby 50% or more of prescribed exercise was supervised in-person.

³ Unsupervised were interventions whereby less than 50% of prescribed exercise was supervised in-person. HR: Heart rate; HR_{max}: Heart rate maximum; HRR: Heart rate reserve; METS: Metabolic equivalents; RPE: Rating of perceived exertion; RPM: Revolutions per minute; VO_{2max}: Maximal oxygen consumption;

VO_{2peak}: Peak oxygen consumption; 6MWT: 6-minute walk test.

CER

Table 2. Adverse events by grade of severity described for those in the exercise and usual care groups.

Adverse	Exerc	rise group	Usual ca	re group		
event	(116 adverse even	nts, 2621 participants)	(40 adverse events, 2579 participants)			
grade ¹	Total number of adverse event	s ² / exercise-related adverse events	Total number of adverse events ²	/ exercise-related adverse events		
Grade 1	Grade 1 adve	erse events: 42/34	Grade 1 adve	rse events: 2/0		
	Low-severity musculoskeletal symptoms (pain/stiffness/soreness/tendonitis) (18/18) Lymphoedema onset or worsening (8/3) Increase in fatigue (4/4) Mild cardiac symptoms or angina (7/6)	Unspecified minor injuries (3/2) Acute illness (1/0) Vertigo (1/1)	Acute illness (1/0) Lymphoedema onset (1/0)			
Grade 2		erse events: 20/10	Grade 2 adve	rse events: 1/0		
	Musculoskeletal injuries (mild fractures, strains, tendinitis) (9/6) High blood pressure (>140/90 mmHg) (4/3) Gynaecologic complication or urinary tract infection (2/0)	Influenza or upper respiration tract infection (2/0) Hypoglycaemia (1/1) Haemorrhoids (1/0) Diabetes mellitus (1/0)	Shingles secondary to varicella zoster infecti	on (1/0)		
Grade 3 ³	Grade 3 adv	erse events: 52/6	Grade 3 adverse events: 34/0			
	Unspecified health/medical problems or illness leading to withdrawal (20/0) Infections/secondary suturing/seroma, discharge/uncontrollable pain (2/0) ⁴ Breast cancer progression (3/0) Breast cancer recurrence (2/0) Cancer ⁵ or developed other cancer (n=5) Hospitalisation (4/0) Lymphoedema (3/0) Musculoskeletal symptoms or injuries leading to withdrawal (2/0) Discomfort with exercise (1/1)	Dizziness and dyspnoea (1/1) Unspecified physical accident (2/1) Foot pain requiring surgery (1/1) Mild chest pain during exercise (1/1) Gastrointestinal complication (1/0) Deep vein thrombosis and pulmonary embolism (1/0) Diverticulosis (1/0) Syncope (1/1)	Unspecified health/medical problems or illness leading to withdrawal (12/0) Infections/secondary/suturing/seroma/ discharge/uncontrollable pain (8/0) Breast cancer progression (4/0) Breast cancer recurrence (3/0) Uncontrolled cardiac disease and hypertension leading to withdrawal (2/0)	Gynaecologic problems (1/0) Anaemia leading to withdrawal (1/0) Foot fracture (1/0) Chemotherapy-induced severe discomfort leading to withdrawal (1/0) Bronchitis (1/0)		
Grade 4	Grade 4 adv	verse events: 0/0	Grade 4 adve	rse events: 0/0		
	Nil		Nil			
Grade 5	Grade 5 adv	verse events: 2/0	Grade 5 adve	rse events: 3/0		
	Death (2/0)		Death (3/0)			

¹ Adverse events were classified using the Common Terminology Criteria²⁶ as; grade 1: asymptomatic or mild symptoms; grade 2: moderate, minimal, local or non-invasive intervention indicated and limiting age-appropriate instrumental activities of daily living; grade 3: severe or medically significant but not immediately life-threatening; grade 4: life-threatening consequences and urgent intervention indicated, or; grade 5: death.

² Includes all adverse events (both exercise- and non-exercise related).

³Adverse events in which the severity was not reported were considered Grade 3 or higher if the event led to study withdrawal.

⁴Not reported individually.

⁵ Reported as "cancer" with no further detail provided on whether the withdrawals were due to cancer progression, recurrence or development of other cancer.

Table 3. Study recruitment rate, withdrawal rate and exercise adherence by exercise mode, treatment status, intervention supervision and intervention duration.

	Recruitment rate (%)	Withdrawa	Adherence rate (%)		
	Median	Median (minimum	, maximum [IQR])	Median ²	
	(minimum, maximum [IQR])	Exercise	Usual care	(minimum, maximum [IQR])	
Overall	45 (1, 96 [40]), n=48	11 (0, 41 [15.5]), n=69	12 (0, 49 [13]), n=69	81 (44, 99 [21]), n=52	
Exercise mode					
Aerobic exercise	32 (1, 96 [45]), n=22	11 (0, 41 [14]), n=26	12 (0, 49 [20]), n=26	86 (71, 99 [18]), n=16	
Resistance exercise	40 (28, 83 [50]), n=6	7 (0, 34 [19]), n=10	11 (4, 43 [15]), n=10	84 (44, 96 [32]), n=9	
Combined exercise	49 (33, 95 [23]), n=13	11 (0, 32 [24]), n=23	7 (0, 32 [16]), n=23	79 (55, 93.9 [20]), n=17	
Other exercise	65 (15, 85 [32]), n=7	17 (0, 41 [19]), n=10	15 (0, 36 [19]), n=10	81 (58, 92 [34]), n=10	
Treatment status			1		
During treatment	48 (14, 96 [35]), n=17	9 (0, 41 [14]), n=29	14 (0, 49 [16]), n=29	80 (58, 99 [15]), n=19	
Post treatment	46 (1, 95 [41]), n=27	12 (0, 41 [20]), n=36	11 (0, 43 [17]), n=36	84 (44, 98 [20]), n=30	
Mixed	26 (13, 47 [- ⁴]), n=4	12 (5, 32 [23]), n=4	16.5 (7, 25 [14.75]), n=4	84 (71, 92 [- ⁴]), n=3	
Supervision					
Supervised	45 (14, 83 [41]), n=24	7 (0, 41 [19]), n=33	14 (0, 49 [IQR]), n=33	79 (44, 98 [20]), n=31	
Unsupervised	46 (1, 96 [48]), n=24	12 (0, 38 [38]), n=36	10 (0, 36 [IQR]), n=36	84 (55, 99 [14]), n=21	
Intervention duration					
<12 weeks	45 (1, 95 [47]), n=27	12 (0, 41 [15]), n=37	14 (0, 49 [14]), n=37	85 (58, 99 [16]), n=27	
>12 weeks	47 (14, 96 [46]), n=21	9 (0, 38 [21]), n=32	11 (0, 43 [16]), n=32	79 (44, 98 [17]), n=25	
Study quality rating ³					
Low	61 (15, 81 [26]) n=14	17 (0, 41 [30]), n=27	16 (0, 49 [25]), n=27	79 (55, 98 [20]), n=20	
High	38 (1, 96 [36]), n=34	9 (0, 34 [12]), n=42	11 (0, 43 [10]), n=42	84 (44, 99 [20]), n=32	
¹ n= values represent nur	nber of studies.)			
² n= values represent nur	nber of groups. Withdrawal and adh	erence rates reported by intervent	ion groups because n=7 studies in	volved multiple intervention group	
³ Low quality: PEDro sca	ale score of less than 6; high quality	: PEDro scale score of 6 or higher	·.		
⁴ Interquartile range not c		-			
IOR · Interquartile range					

IQR: Interquartile range.

Figure 1. Meta-analysis of all grade 3 to 5 adverse events in exercise compared to usual care presented as overall and separated by exercise mode, treatment status, intervention duration and degree of supervision.

Subgroup	Studies (n=)	Participants (n=) Intervention vs. Control	Total adverse events (n=) Intervention vs. Control	RD (95% CI)	P-value	
Exercise mode						
Aerobic exercise	22	960 vs. 938	22 vs. 11	0.00 (-0.01, 0.02)	0.45	-
Resistance exercise	9	389 vs. 383	3 vs. 9	-0.01 (-0.02, 0.01)	0.33	H
Combined exercise	27	846 vs. 871	22 vs. 7	0.01 (-0.01, 0.02)	0.16	-
Other exercise	11	426 vs. 387	6 vs. 10	0.00 (-0.02, 0.02)	0.89	H
Supervision [1]						
Supervised interventions	42	1136 vs. 1125	29 vs. 14	0.00 (-0.01, 0.01)	0.35	•
Unsupervised interventions	27	1485 vs. 1454	25 vs. 23	0.00 (-0.01, 0.01)	0.79	
Intervention timing						
During treatment	29	1128 vs. 1145	24 vs. 17	0.00 (-0.01, 0.01)	0.85	
Post-treatment	36	1273 vs. 1213	25 vs. 16	0.01 (-0.01, 0.02)	0.23	-
Mixed	4	220 vs. 221	5 vs. 4	0.00 (-0.03, 0.02)	0.79	H
Duration						
<12 week interventions	36	1578 vs. 1469	20 vs. 19	0.00 (-0.01, 0.01)	0.98	
>12 week interventions	33	1043 vs. 1110	34 vs. 18	0.01 (-0.01, 0.01)	0.25	
Sensitivity analyses [2]						
High quality studies only	42	1862 vs. 1888	40 vs. 31	0.00 (-0.00, 0.01)	0.48	•
Stage II+ only	10	257 vs. 258	16 vs. 10	0.02 (-0.03, 0.06)	0.40	⊢
Overall	69	2621 vs. 2579	54 vs. 37	0.00 (-0.01, 0.01)	0.38	-0.1 0 0.1

-0.1 0 0.1

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Figure 2. Meta-analyses results of aerobic fitness with subgroup analyses for exercise mode, intervention supervision, timing and duration, and sensitivity analyses (positive SMD values favour exercise).

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value		
Exercise mode		Intervention vs. Control					
Aerobic exercise	15	395 vs. 355	31%	0.62 (0.43, 0.81)	<0.01		H
Resistance exercise	3	108 vs. 76	0%	0.23 (-0.07, 0.53)	0.13	ŀ	
Combined exercise	13	487 vs. 498	87%	0.65 (0.26, 1.03)	<0.01		
Other exercise	0	-	-	-	-		
Supervision [1]							
Supervised interventions	15	491 vs. 413	84%	0.66 (0.29, 1.03)	<0.01		
Unsupervised interventions	16	499 vs. 516	43%%	0.53 (0.35, 0.70)	<0.01		H
Intervention timing							
During treatment	14	611 vs. 546	68%	0.58 (0.36, 0.81)	<0.01		H
Post-treatment	15	279 vs. 278	0%	0.50 (0.33, 0.67)	<0.01		H
Mixed	2	100 vs. 105	98%	1.47 (-1.01, 3.95)	0.25	<	
Duration							
<12 week interventions	13	255 vs. 250	34%	0.59 (0.36, 0.83)	<0.01		⊢∎⊣
>12 week interventions	18	724 vs. 642	83%	0.62 (0.35, 0.90)	<0.01		⊢ ∎
Sensitivity analyses [2]							
High quality studies only	18	714 vs. 663	84%	0.60 (0.32, 0.89)	<0.01		⊢∎⊣
Stage II+ only	4	83 vs. 81	34%	0.57 (0.15, 0.99)	<0.01		⊢ I
Overall	31	990 vs. 929	75%	0.62 (0.42, 0.81)	<0.01		•
						-1 (0 1

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision. [2] Sensitivity analyses were performed only on i) high quality studies, and ii) studies with 100% of samples with stage II+ disease.

Figure 3. Meta-analyses results of quality of life with subgroup analyses for exercise mode, intervention supervision, timing and duration and sensitivity analyses (positive SMD values favour exercise).

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs Control				
Aerobic exercise	9	653 vs. 645	80%	0.38 (0.24, 0.53)	<0.01	⊢∎→
Resistance exercise	5	247 vs. 171	3%	0.29 (0.09, 0.49)	<0.01	⊢∎1
Combined exercise	20	646 vs. 609	82%	0.35 (0.25, 0.44)	<0.01	HEH
Other exercise	6	231 vs. 202	75%	0.75 (0.55, 0.95)	<0.01	⊢
Supervision [1]						
Supervised interventions	19	643 vs. 521	75%	0.59 (0.46, 0.71)	<0.01	⊢∎⊣
Unsupervised interventions	21	1134 vs. 1076	79%	0.30 (0.22, 0.39)	<0.01	H
Intervention timing						
During treatment	16	849 vs. 714	79%	0.43 (0.33, 0.54)	<0.01	HEH
Post-treatment	21	776 vs. 727	81%	0.35 (0.25, 0.45)	<0.01	HEH
Mixed	3	152 vs. 156	27%	0.45 (0.22, 0.68)	<0.01	⊢ −■−−1
Duration						
<12 week interventions	24	1053 vs. 955	82%	0.42 (0.33, 0.51)	<0.01	H
>12 week interventions	16	724 vs. 642	71%	0.37 (0.26, 0.47)	<0.01	HEH
Sensitivity analyses [2]						
High quality studies only	24	1345 vs. 1202	80%	0.31 (0.23, 0.39)	<0.01	H
Stage II+ only	6	128 vs. 144	77%	0.78 <mark>(</mark> 0.55, 1.01)	<0.01	⊢ −■−−1
Overall	40	1777 vs. 1597	78%	0.40 (0.33, 0.47)	<0.01	•

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Figure 4. Meta-analyses results of fatigue with subgroup analyses for exercise mode, intervention supervision, timing and duration and sensitivity analyses (positive SMD values favour exercise).

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	11	671 vs. 628	80%	0.31 (0.17, 0.45)	<0.01	H
Resistance exercise	2	128 vs. 90	0%	0.25 (-0.02, 0.53)	0.07	⊢
Combined exercise	12	470 vs. 481	64%	0.33 (0.23, 0.44)	<0.01	HEH
Other exercise	6	225 vs. 185	87%	0.21 (0.00, 0.41)	0.04	
Supervision [1]						
Supervised interventions	14	534 vs. 451	79%	0.44 (0.30, 0.57)	<0.01	H
Unsupervised interventions	17	960 vs. 933	70%	0.24 (0.15, 0.33)	<0.01	H - H
Intervention timing						
During treatment	17	829 vs. 737	79%	0.29 (0.19, 0.39)	<0.01	H
Post-treatment	13	618 vs. 596	74%	0.33 (0.22, 0.45)	<0.01	H
Mixed	1	47 vs. 51	-	0.11 (-0.28, 0.51)	0.25	⊢
Duration						
<12 week interventions	18	843 vs. 798	84%	0.34 (0.24, 0.44)	<0.01	H
>12 week interventions	13	651 vs. 586	13%	0.26 (0.14, 0.37)	<0.01	HEH
Sensitivity analyses [2]						
High quality studies only	21	407 vs. 422	67%	0.30 (0.22, 0.38)	<0.01	
Stage II+ only	4	92 vs. 95	73%	0.46 (0.16, 0.75)	<0.01	
Overall	31	1494 vs. 1384	75%	0.30 (0.23, 0.38)	<0.01	•

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Supplementary Content 7. Meta-analyses results of anxiety with subgroup analyses for exercise mode, intervention supervision, timing and duration and sensitivity analyses (positive SMD values favour exercise).

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	4	84 vs. 127	0%	0.37 (0.09, 0.65)	0.01	⊨■
Resistance exercise	1	41 vs. 82	-	0.08 (-0.30, 0.45)	0.68	—
Combined exercise	4	143 vs. 143	59%	1.36 (1.10, 1.62)	<0.01	⊢
Other exercise	5	168 vs. 217	94%	0.83 (0.61, 1.06)	<0.01	⊢-■1
Supervision [1]						
Supervised interventions	7	213 vs. 299	92%	0.62 (0.43, 0.81)	<0.01	⊢ ∎1
Unsupervised interventions	7	223 vs. 270	82%	0.93 (0.74, 1.13)	<0.01	⊢∎→
Intervention timing						
During treatment	10	328 vs. 465	92%	0.75 (0.60, 0.91)	<0.01	H -1
Post-treatment	4	108 vs. 104	14%	0.84 (0.56, 1.13)	<0.01	⊢
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	7	198 vs. 254	91%	0.83 (0.62, 1.03)	<0.01	⊢-■
>12 week interventions	7	238 vs. 315	89%	0.73 (0.55, 0.91)	<0.01	+∎
Sensitivity analyses [2]						
High quality studies only	10	341 vs. 433	91%	0.81 (0.65, 0.96)	<0.01	H
Stage II+ only	4	102 vs. 98	84%	0.80 (0.51, 1.09)	<0.01	⊢
Overall	14	436 vs. 569	89%	0.77 (0.64, 0.91)	<0.01	•

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Supplementary Content 10. Meta-analyses results of waist circumference with subgroup analyses for exercise mode, intervention supervision,

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs Control				
Aerobic exercise	3	30 vs. 39	0%	0.04 (-0.44, 0.51)	0.89	⊢
Resistance exercise	0	0 vs. 0	-	-	-	
Combined exercise	4	120 vs. 152	0%	0.32 (0.08, 0.57)	<0.01	⊢
Other exercise	1	20 vs. 20	-	-0.11 (-0.73, 0.51)	0.72	⊢−−−− (
Supervision [1]						
Supervised interventions	8	170 vs. 211	0%	0.22 (0.02, 0.43)	0.03	H-
Unsupervised interventions	0	-	-	-	-	
Intervention timing						
During treatment	0	-	-	-	-	
Post-treatment	8	170 vs. 211	0%	0.22 (0.02, 0.43)	0.03	⊨ _
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	6	84 vs. 123	0%	0.22 (-0.06, 0.51)	0.12	⊢
>12 week interventions	2	170 vs. 211	0%	0.22 (-0.08, 0.52)	0.14	⊢
Sensitivity analyses [2]						
High quality studies only	1	10 vs. 16	-	-0.07 (-0.86, 0.73)	0.87	·
Stage II+ only	0	-	-	-	-	
Overall	8	170 vs. 211	0%	0.22 (0.02, 0.43)	0.03	<u> </u>
						-1 -0.5 0 0.5

< Favours Control Favours Exercise >

timing and duration and sensitivity analyses (positive SMD values favour exercise).

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised exercise involved face-to-face supervised exercise involved face-to-face supervision and unsupervised exercise involved face-to-face supervised exercise involved face-to-face supervised exercise involved exercise involved exercise involved exercise involved exercise exercise involved exercise exerc face supervision.

Supplementary Content 11. Meta-analyses results of body weight with subgroup analyses for exercise mode, intervention supervision, timing

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	6	172 vs. 164	0%	0.11 (-0.11, 0.32)	0.32	⊢
Resistance exercise	3	172 vs. 169	0%	0.00 (-0.21, 0.21)	0.99	⊢
Combined exercise	5	162 vs. 160	0%	0.14 (-0.08, 0.36)	0.22	⊢
Other exercise	1	20 vs. 20	-	-0.03 (-0.65, 0.59)	0.92	$\longleftrightarrow \bullet \bullet$
Supervision [1]						
Supervised interventions	15	526 vs. 513	0%	0.08 (-0.04, 0.20)	0.22	⊢
Unsupervised interventions	0	-	-	-	-	
Intervention timing						
During treatment	2	164 vs. 160	0%	0.18 (-0.04, 0.40)	0.11	
Post-treatment	13	362 vs. 353	0%	0.03 (-0.12, 0.18)	0.67	⊢
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	7	148 vs. 145	0%	0.05 (-0.18, 0.28)	0.65	⊢
>12 week interventions	8	378 vs. 368	20%	0.09 (-0.06, 0.23)	0.24	⊢
Sensitivity analyses [2]						
High quality studies only	8	373 vs. 361	0%	0.08 (-0.07, 0.22)	0.29	⊢
Stage II+ only	0	-	-	-	-	
Overall	15	526 vs. 513	0%	0.08 (-0.04, 0.20)	0.22	•
						-0.5 0 0.5

< Favours Control Favours Exercise >

and duration and sensitivity analyses (positive SMD values favour exercise).

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Supplementary Content 12. Meta-analyses results of body mass index with subgroup analyses for exercise mode, intervention supervision,

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	4	123 vs. 119	0%	0.14 (-0.12, 0.39)	0.29	⊢
Resistance exercise	0	-	-	-	-	
Combined exercise	8	165 vs. 204	5%	0.22 (0.01, 0.43)	0.04	⊢_ ∎(
Other exercise	1	20 vs. 20	-	-0.06 (-0.68, 0.56)	0.86	F
Supervision [1]						
Supervised interventions	8	205 vs. 232	0%	0.10 (-0.09, 0.29)	0.31	⊨∎→
Unsupervised interventions	5	103 vs. 111	0%	0.31 (0.04, 0.58)	0.03	⊢ −−■−−−1
Intervention timing						
During treatment	2	27 vs. 27	83%	0.28 (-0.28, 0.84)	0.33	· • •
Post-treatment	11	281 vs. 316	0%	0.16 (-0.00, 0.32)	0.05	
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	6	149 vs. 178	0%	0.13 (-0.09, 0.35)	0.24	⊢ ∎1
>12 week interventions	7	159 vs. 165	20%	0.21 (-0.01, 0.43)	0.07	⊢_ ∎1
Sensitivity analyses [2]						
High quality studies only	6	177 vs. 175	2%	0.12 (-0.09, 0.33)	0.27	⊢ -∎1
Stage II+ only	0	-	-	-	-	
Overall	13	308 vs. 343	0%	0.17 (0.01, 0.32)	0.03	-1 -0.5 0 0.5 1

< Favours Control Favours Exercise >

timing and duration and sensitivity analyses (positive SMD values favour exercise).

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involved face-to-face supervision and unsupervised exercise involved face-to-face supervised exercise involved face-to-face supervision and unsupervised exercise involved face-to-face supervised exercise involved face-to-face supervised exercise involved exercise involved exercise involved exercise involved exercise exercise involved exercise exerc face supervision.

Supplementary Content 13. Meta-analyses results of body fat with subgroup analyses for exercise mode, intervention supervision, timing and

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs Control				
Aerobic exercise	5	185 vs. 177	0%	0.10 (-0.10, 0.31)	0.33	F
Resistance exercise	4	150 vs. 150	0%	0.03 (-0.19, 0.26)	0.78	⊢
Combined exercise	4	104 vs. 106	0%	0.23 (-0.04, 0.51)	0.09	⊢_
Other exercise	0	-	-	-	-	
Supervision [1]						
Supervised interventions	9	352 vs. 350	0%	0.12 (-0.03, 0.26)	0.13	+■-
Unsupervised interventions	4	87 vs. 83	0%	0.09 (-0.22, 0.39)	0.57	⊢
Intervention timing						
During treatment	2	164 vs. 160	0%	0.14 (-0.07, 0.36)	0.20	
Post-treatment	11	275 vs. 273	0%	0.09 (-0.08, 0.26)	0.29	⊢∎→
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	5	95 vs. 91	0%	0.08 (-0.21, 0.37)	0.59	⊢
>12 week interventions	8	344 vs. 342	20%	0.12 (-0.03, 0.27)	0.12	+-■
Sensitivity analyses [2]						
High quality studies only	10	384 vs. 378	0%	0.09 (-0.05, 0.23)	0.21	H <mark>a</mark> H
Stage II+ only	0	-	-	-	-	
Overall	13	439 vs. 433	0%	0.11 (-0.02, 0.24)	0.11	•
						-1 -0.5 0 0.5

< Favours Control Favours Exercise >

1

duration and sensitivity analyses (positive SMD values favour exercise).

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

Supplementary Content 3: Overview of samples and exercise details of included studies (n=61)
--

	Sample	Exercise	Intervention setting	Supervision ¹
Ahmed 2006	N=46: 0–III BC with lymphoedema	Type: Resistance exercise	Recreation centre	Supervised: ACSM certified fitness
	Exercise group:	Frequency: 2/week for 6 months		professional.
	DCIS: 1 (4.4%);	Intensity: 3 sets per exercise of 8–12		
	Stage I: 6 (26.1%);	repetitions		
	Stage II: 13 (56.5%)	Time: ~60 minutes		
	Stage III: 3 (13.0%)			
Anderson 2012	N=104 stage I-III BC	Type: Combined aerobic and resistance	University health and exercise research	Supervised: Occupational or physical
	Exercise group	Frequency: 2/week for 12 months	centre	therapist.
	I: n=25 (48%); II: n=19 (37%); III: n=8 (15%);	Intensity: Resistance exercise:		-
	N/A: n=0.	>50%1RM, 12 repetitions 14–16 RPE		
	Usual care	Time: ~60 min total session (30min		
	I: n=26 (50%); II: n=21 (40%); III: n=4 (8%); N/A:	continuous walking)		
	n=1 (2%).	· · · · · · · · · · · · · · · · · · ·		
Banerjee 2007	N=68 undergoing RT; Stage II–III	Type: Other (Yoga)	Hospital outpatient setting and home	Supervised: Yoga instructors
5	Exercise group (n=23):	Frequency: 6 weeks		1 0
	II: 17 (48%)	Intensity: Low		
	III: 18 (52%)	Time: 90 min		
Campbell 2017	N=19 completed chemotherapy for stages I-IIIA	Type: Aerobic	Research gym and home	Supervised: Not reported by whom
•	BC	Frequency: 4/week for 24 weeks		
	I: 0	Intensity: 60–80% of HRR		Unsupervised
	II: 10 (100%)	Time: 30–45 min		*
	III: 0	Y		
Cantarero-Villanueva	N=40 BC; Stage I–IIIA currently receiving HT.	Frequency: 3/week for 8 weeks	University medical centre outpatient	Supervised: Exercise trainer specialist
2012a	Exercise group n=20	Intensity: Not specified	clinic and swimming pool	and physiotherapists
	I: 6 (30)	Time: 60 min (5 minutes of warm-up,		
	II: 8 (40)	15-20 min of aerobic exercise, 15 min		
	IIA: 6 (30)	of mobility exercise and 20 min of		
		recovery techniques.		
		Type: Other, hydrotherapy		
Cantarero-Villanueva	n=78 BC survivors; stage I–IIIA	Frequency: 3/week for 8 weeks	University medical centre outpatient	Supervised: not reported
2012b	Exercise group n=32:	Intensity: Aerobic exercise[]: Resistance	clinic and swimming pool	1 1
	I: 4 (12.5)	exercise: 75% maximum load, 2–3 sets		
	II: 23 (71.9)	of 10–15 repetitions,		
	IIIA: 5 (15.6)	Time: 90 min		
		Type: Aerobic, resistance & mobility		
		exercise (Core Stability Exercises; all		
		major muscle groups; small soft ball,		
		mats, fit-ball & resistance bands).		
Cantarero-Villanueva	N=66; stage I–IIIA	Frequency: 3/week for 8 weeks	University medical centre outpatient	Supervised: Physical therapist
2012c;	Exercise group (n=33)	Intensity: Low-intensity	clinic and swimming pool	
,	I: 16 (48)	Time: 60 min (10-min warm-up, 35 min,	Summing Poor	
	II: 10 (40) II: 10 (30)	15 min cool-down)		
	IIIA: 7 (22)	Type: Hydrotherapy		
		i jpe. iijulouleupj		1

2013	hormone therapy Exercise group: n=32 I: 4 (12.5)	Intensity: Moderate; 2–3 sets of 8–12 repetitions Time: 60 min (10 min warm-up, 40 min	clinic and swimming pool	physical therapists
	I. 4 (12.5) II: 23 (72)	aerobic & endurance exercises, 10 min		
	IIIA: 5 (15.5)	cool-down).		
		Type: Hydrotherapy		
Chandwani 2014	N=109 stage 0-III breast undergoing RT	Frequency: 3/week for 6 weeks	Cancer treatment centre	Supervised: certified yoga instructor
	Exercise group:	Intensity: Low		
	0: 5 10	Time: 60 min		
	I: 16 30	Type: Yoga		
	II: 15 28			
	III: 17 32			
Cormie 2013	N=63; stage I–III BC;	Frequency: 2/wk for 12 weeks	Hospital, health clinic	Supervised: Accredited exercise
	High-load exercise:	Intensity: 12–16 RPE; High load: 75–85		physiologist
	I: 2 (9.1)	% of 1RM using 10-6 RM Low load:		
	II: 18 (81.8)	55-65 % of 1RM using 20-15 RM		
	III: 2 (9.1)	Time: 60 min (inc. 10 min warm-up & 5		
	Low-load exercise:	min cool-down)		
	I: 5 (23.8)	Type: Resistance exercise		
	II: 10 (47.6)			
	III: 6 (28.6)			
Cornette 2016	N=44; stage I-IIIB during CT	Frequency: ≥ 3 /week for 1 year	Home-based	Unsupervised: Weekly telephone
	Exercise group (n=20)	Intensity: Aerobic: 60RPM; 70-80%		contact from an exercise specialist
	I: 3 (15%)	HRmax; 3-6 METS; Resistance: 2 sets		
	IIA: 8 (40%)	per exercise of 8–12RM		
	IIB: 3 (15%)	Time: Aerobic: 20–40 min (+5 min		
	IIIA: 5 (25%)	warm-up and 5 min cool down);		
	IIIB: 1 (5%)	Resistance: Not specified.		
		Type: Resistance- and aerobic-based.		
Courneya 2003	N=53; stage I-IIIa; completed treatment	Frequency: 3/week for 15 weeks	University Cancer institute	Supervised: Accredited exercise
	Exercise group:	Intensity: 70–75% VO ₂ max		physiologist
	I: 10 (42%)	Time: 15–35 mins (+5 min warm-up &		
	IIa: 6 (25%)	5 min cool down)		
	IIb: 6 (25%)	Type: Aerobic; cycle ergometer		
	IIIa: 2 (8%)			
Courneya 2007	N=242; stage I-IIIa; BC initiating adjuvant CT	Frequency: 3/week for 15 weeks	University Cancer institute	Supervised: Accredited exercise
	Aerobic exercise group:	Intensity: Aerobic exercise group: 60-		physiologist
	I: 18 (23.1%)	80% VO ₂ max Resistance exercise group:		
	IIa: 33 (42.3%)	2 sets of 8–12 repetitions at 60-70%		
	IIb: 17 (21.8%)	1RM		
	IIIa: 10 (12.8%)	Time: Aerobic exercise group: 15–45		
	Resistance exercise group:	min (+5 min warm-up and 5 min cool		
	I: 22 (26.8%)	down) Resistance exercise group: Not		
	IIa: 36 (43.9%)	specified.		
	IIb: 9 (11.0%)	Type: Aerobic exercise (cycle		
	IIIa: 15 (18.3%)	ergometer, treadmill, or elliptical) or		
		resistance exercise		

Danhauer 2009	N=44; DCIS- stage VI Exercise group: DCIS: 13.6 (3%) I: 22.7 (5%) II: 45.5 (10%)	Type: Other, yoga Frequency: 1/week for 10 weeks Intensity: Low Time: 75 mins	Yoga studio	Supervised: Certified yoga instructor
	III: 13.6 (3%) IV: 4.6 (1%)			
De Luca 2016	N=20; stage I–III completed treatment Exercise group n=10: I: 4 (x%) II: 5 (x%) III: 2 (x%)	Frequency: 2/ week for 24 weeks Intensity: Aerobic exercise: 70–80% HRmax; Resistance exercise: 2–4 sets per exercise, 6–10 repetitions at 40–60% IRM. Time: 90 min (10 min warm-up, 40 min resistance exercise, 30 min aerobic exercise, 10 min cool down); Type: Aerobic and resistance exercise	University gymnasium	Supervised: Fitness professional and physician
Dethlefsen 2016	N= 74 diagnosed with operable (stage I–III); <6 months since completing CT Exercise group (n=37) I: 4% II: 60% III: 36%	Type: Aerobic and resistance exercise Frequency: 1/week for 6 months (supervised) Intensity: Aerobic exercise: Intervals ranged from 30 s (maximum intensity) to 6 min (90%–95% of HRmax); Resistance exercise: 3 sets of 8–12 repetitions at 70–90% of 1RM Time: 90 min (supervised), >3 h/week	Hospital	Supervised: not reported
Dolan 2016	N=33; stage 0-III; postmenopausal; Interval exercise group: 0: 0 I: 3 II: 5 III: 3 Other: 1 Continuous intensity exercise group: 0: 1 I: 2 II: 2 III: 5 Other: 1	Frequency: 3/week for 6 weeks Intensity: Interval exercise: 3.22–4.02 km; ≤2 min bouts at ≥80 % VO ₂ peak; Continuous exercise: 3.22–4.02 km at 55–70 % VO ₂ peak Time: Interval exercise: Continuous exercise: Type: Aerobic (treadmill)	Location not specified	Supervised: Accredited exercise physiologist
Drouin 2005	N=20 stage 0–IIIb; during RT Exercise group (n=13): 0: 3 I: 2 II: 6 III: 2	Frequency: 3–5/week for 7 weeks Intensity: 50–70% HRmax Time: 20–45 min Type: Aerobic, walking	Home-based	Unsupervised: Weekly contact (face-to- face or by telephone) with researcher
Eakin 2012	N=143; invasive BC, 6 weeks post-surgery Telephone group: 0–I: 26 (35.6%) II+: 38 (52.1%)	Frequency: ≥4/week for 8 months Intensity: Low-high Time: 20–45+ min Type: Aerobic & resistance exercise	Home-based	Unsupervised: Weekly to monthly telephone contact with an Accredited exercise physiologist

		(telephone-delivered exercise)		
Fernández-Lao 2013	N=98; stage I–IIIA) completed treatment (excluding HT) Land-based exercise group n=31: I: 5 II: 21	Frequency: 3/ week for 8 weeks Intensity: 60% HRmax (aerobic exercise) and 2–3 sets of 8–12 repetitions (resistance exercise) Time: 60 min (inc. 10 min warm-up and	A gymnastic hall and heated swimming pool	Supervised: Fitness specialist and physical therapist
	IIIa: 5 Water-based exercise group n=33: I: 13 II: 13 IIIa: 7	10 min cool-down) Type: Aerobic and resistance exercise; land- or water-based	S.	
Galiano-Castillo 2017	N=81 completed adjuvant therapy (except hormone treatment) for stage I to IIIA breast cancer Exercise group n=40 I: 14 (35%) II: 18 (45%) IIA: 8 (20%)	Type: Combined aerobic and resistance Frequency: 3/week for 8 weeks Intensity: Moderate Time: 90 min	Home-based	Unsupervised: Internet-based (Tele- rehabilitation)
Gokal 2015	N=50; stage I–III BC N=25 exercise; n=25 control I: 0 0 II: 5 20 III: 20 80	Frequency: 5/week for 12 weeks Intensity: 12–14 RPE Time: 10–30 mins Type: Aerobic, Walking	Home-based	Unsupervised: Physical activity booklet
Guinan 2013	N=26 BC survivors Stage I–III Exercise group n=16: I: 3 (18.8) II: 10 (62.6) III: 3 (18.8)	Frequency: 2/week for 8 weeks Intensity: 35–65% HRR Time: 21–42 min Type: Aerobic (stationary bike, treadmill, rowing ergometer).	Centre- (unspecified) and home-based	Supervised: Physiotherapist and a research assistant
Hatchett 2013	N=85 stage I-IV BC completed treatment Intervention group (n=36) I: 10 II: 17 III: 6 IV: 3	Frequency: 3-7/week for 12 weeks Intensity: 12-14 RPE Time: 10-60 min (150 min/week total) Type: Aerobic and resistance exercise	Home-based	Unsupervised: email delivered intervention (e-counselor exercise physiologist)
Hayes 2012	N=194 Stage 0-III; 6-weeks post- surgery Face-to-face exercise (n=67) 0: 2 (3.0) I: 23 (34.3) II-III: 38 (56.7) Unknown: 4 (6.0) Telephone exercise (n=67) 0: 3 (4.5) I: 18 (26.9) II-III: 45 (67.2) Unknown: 1 (1.5)	Frequency: ≥4/week for 8 months Intensity: Low-high Time: 20–45+ min Type: Aerobic & resistance exercise	Home-based	Supervised and unsupervised: Accredited exercise physiologist
Headley 2004	n=32; stage IV BC. Exercise group (n=16)	Frequency ⁴ : 3/week for 12 weeks Intensity: "Low-to-moderate" (RPE not	Home-based	Unsupervised: exercise DVD

	IV: 16	reported) Time ⁵ : 30 min Type: Other; Stretching and repeated flexion and extension of the arms, head, upper torso, and legs while seated.		
Herrero 2005	N=16; stage I–II ductal breast carcinoma Exercise group; n=8 I: 3 II: 5	Frequency: 3/week for 8 weeks Intensity: Aerobic: 70–80% HRmax; Resistance: 1–3 sets of 8–20 repetitions; Time: 90 min (inc. 10 min warm-up & 10 min cool down) Type: Aerobic (cycle ergometer) & resistance exercise	Community fitness centre	Supervised: Exercise physiologists
Hornsby 2014	n=20; stage IIB-IIIC BC n=10 exercise; n=10 control;	Frequency ⁴ : 3/week for 12 weeks Intensity: 60–100% VO ₂ peak Time ⁵ : 15–45 mins Type: Aerobic exercise (cycle ergometer; continuous and interval training)	Cancer institute	Supervised: Exercise physiologist
Husebø 2014	N=67 stage I–III BC during CT Exercise group: I: 7 (24.2) II: 19 (65.5) III: 3 (10.3)	Frequency: Aerobic: Daily; Resistance exercise: 3/week for 6 months Intensity: Moderate Time: Aerobic exercise: 30 mins; Resistance exercise: Not specified Type: Aerobic & resistance exercise	Home-based	Unsupervised: fortnightly telephone calls from the research team
Hutnick 2005	N=49 stage I–III Exercise group: I: 6 (21.4%) II: 19 (67.9%) III: 2 (7.1%) Unknown: 1 (3.6%)	Frequency: 3/week for 6 months Intensity: Aerobic: 60–75% functional capacity; Resistance; 3 sets of 8-12 repetitions Time: 40–90 min Type: Aerobic & resistance exercise	University clinical setting and home	Supervised: exercise trainer
Kilbreath 2012	N=160 stage I–III BC Exercise group n=81 I: 17% II: 44% III: 38%	Type: Resistance exercise & stretching Frequency: ≥1/week for 8 weeks Intensity: Resistance exercises: 2 sets per exercise for 8–15 repetitions, 15 Borg RPE; Stretching: hold each stretch for 5–15 min Time: Not specified	Centre-based (unspecified) and home- based	Unsupervised and supervised: Not reported
Kim 2006	N=41 stage 0-III undergoing adjuvant therapy Exercise group: 0: 1 (4.5%) I: 10 (45.5%) II: 8 (36.4%) III: 3 (13.6%)	Type: Aerobic exercise Frequency: ≥3/week for 8 weeks Intensity: 60–70% HRR and/or VO ₂ peak Time: 30 min (+5 min warm-up and 5 min cool-down)	University exercise facility	Supervised: Exercise physiologists
Ligibel 2008	N=101 I-III Exercise group: I: 22 43 II: 22 43	Type: Aerobic & resistance exercise Frequency: >2/week for 16 weeks Intensity: Aerobic: 55–80% HRmax; Resistance: 80% 1RM, 2–4 sets per	Centre- (unspecified) and home-based,	Supervised: Personal trainer

	III: 6 12	muscle group Time: Aerobic exercise: 90 min; Resistance exercise: 50 min		
Ligibel 2016	n=101; metastatic BC. n=48 exercise; n=53 control;	Frequency ⁴ : \geq 150 min/week for 16 weeks Intensity: 55–80% HRmax Time ⁹ : \geq 150 min/week Type: Aerobic exercise	Home-based, supervised and unsupervised: Exercise physiologist	Unsupervised and supervised: Face-to- face and telephone contact with an exercise physiologist
Loudon 2014	N=28 stage 0-III BC Exercise group (n=15) 0: 0 I: 3 (25%) II: 6 (50%) III: 3 (25%)	Type: Yoga Frequency: 7/week for 8 weeks Intensity: Not specified Time: 40–90 min	Centre- (unspecified) and home-based,	Unsupervised and supervised: Yoga instructor and at home DVD
Macvicar 1989	n=45; stage II BC. n=18 exercise; age=45±10. n=11 stretching exercises); age=46±10. n=16 control; age=43±9	Type: Aerobic exercise (Interval cycle ergometry) Frequency ⁴ : 3/week for 10 weeks Intensity: 60–85% HRR Time ⁵ : Not reported	Centre-based (unspecified)	Supervised: Not reported
Maryam 2010	N=56 women with BC receiving CT stage I–III Exercise group I: 3 (10.7%) II: 20 (71.4%) III: 5 (17.9%)	Type: Aerobic & resistance Frequency: 3–5/week for 9 weeks Intensity: Light Time: 20–30 min	Home-based	Unsupervised: CD
Milne 2008	N = 58 within 2 years of completing adjuvant therapy stage I–IIIa Exercise group n=29: I: 15 (25.9%) IIa: 25 (43.1%) IIb: 16 (27.6%) IIIa: 2 (3.4%)	Type: Aerobic & resistance exercise Frequency: 3/week for 12 weeks Intensity: Aerobic exercise: Not specified; Resistance exercise: 12 exercises, 2 sets of 10–15 repetitions Time: Aerobic: 25 min (inc. 5 min cool- down).	Rehabilitation clinic	Supervised: Exercise physiologists
Moadel 2007	N=128 stages I to IV) BC Exercise group (n=84): I: 42 II: 36 III: 17 IV: 5	Frequency: 1/week for 12 weeks Intensity: Low Time: 90 min Type: Yoga	Cancer centre	Unsupervised and supervised: Oncologist and Yoga instructor
Mohan Rao 2015	N=98 stage II–III BC undergoing surgery followed by adjuvant RT and/or CT Exercise group n=45 II: 17 (54.83%) III: 16 (42.1%)	Frequency: 3/week for 24 weeks Intensity: Low Time: 60 min Type: Yoga	Hospital	Unsupervised and supervised: Yoga instructor
Mulero Portela 2008	N=44 Stage I–IV BC completed treatment Home exercise group: I: 3 II: 5	Home exercise group: Type: Aerobic & resistance exercise Frequency: 5/week for 26 weeks Intensity: Aerobic exercise: 12–16 RPE	Gymnasium or Home-based	Supervised and unsupervised: physical therapists

	III: 2 IV: 0 Unknown: 3 Gym exercise group: I: 0 II: 3 III: 6 IV: 0 Unknown: 3	(6–20 scale); Resistance exercise: 13–15 RPE (6–20 scale); 2–3 sets per exercise, 10–15 repetitions per set. Time: Aerobic exercise: 30 min; Resistance exercise: Not specified Gym exercise group: Type: Aerobic & resistance exercise Frequency: 5/week for 26 weeks Intensity: Aerobic exercise: 60–80% of HRmax; Resistance exercise: 13–15 RPE (6–20 scale); 2–3 sets per exercise, 10–15 repetitions per set. Time: Aerobic exercise: 30 min; Resistance exercise: Not specified	CRR S	
Murtezani 2014	N=62 completed surgery, RT, and/or CT with or without current HT use stage I–IIIa. Exercise group (n=30) I: 10 (33%) IIa: 11 (37%) IIb: 6 (20%) IIIa: 3 (10%)	Type: Aerobic exercise Frequency: 3/week for 10 weeks Intensity: 50–75% HRR Time: 25–45 min	University clinical rehabilitation centre,	Supervised: Not reported
Musanti 2012	N=42 stage I–IIIB BC who had completed adjuvant chemotherapy Aerobic group: I: 5 II: 5 III: 2 Resistance group: I: 5 II: 10 III: 2	Aerobic group: Type: Aerobic exercise Frequency: 3/week for 12 weeks Intensity: 40–85 HRmax Time: 15–30 min Resistance group: Type: Resistance exercise Frequency: 3/week for 12 weeks Intensity: 3–5 RPE (0–10 scale, up to 8 RPE at the completion of 12 repetitions); 1 set of 10–12 repetitions Time: Not specified	Home-based	Unsupervised: Exercise booklet
Naraphong 2015	N=23 with postoperative stage I–IIIa breast cancer, scheduled to receive CT Exercise n=11: I: 1 (9.09%) II: 7 (63.64%) IIIA: 3 (27.27%)	Type: Aerobic exercise Frequency: 3–7/week for 10 weeks Intensity: 12–14 RPE (6–20 scale), 40– 60% of HRmax; <3-6 METS Time: 20–30 min (plus 5 min warm-up & 5 min cool-down)	Home- and community-based	Unsupervised: Weekly telephone contact with a nurse
Naumann 2012	N=36 BC survivors stage I–III breast cancer, within 12 months of treatment completion Group-based exercise group (n=14) Stage (mean±SD): 2.0 ± 0.6	Type: Aerobic and resistance exercise Frequency: 3/week for 9 weeks Intensity: Moderate Time: 45–60 min	Gymnasium	Supervised: Accredited exercise physiologist
Pinto 2005	N=86: stage 0–II BC, completed treatment	Type: Aerobic	Home-based	Unsupervised: telephone support from

	0: 8 (18.6%) I: 17 (39.5%) II: 18 (41.9%)	Frequency: ≥5/week for 12 weeks Intensity: 55–65% HR _{max} Time: 30 min		research staff and pedometer
Pinto 2013	N=192: stage 0–IV BC currently undergoing treatment Exercise: 0: 12 (11%) I: 41 (39%) II: 44 (42%) III/IV: 9 (8%)	Type: Aerobic Frequency: ≥5/week for 12 weeks Intensity: 55–65% HR _{max} Time: 30 min	Home-based	Unsupervised: telephone counselling from a physical activity counsellor and pedometer
Pinto 2015	N=76 stage 0–III BC completed treatment 0: 3 (7.69%) I: 16 (41.03%) II: 16 (41.03%) III: 4 (10.26%)	Type: Aerobic (walking) Frequency: 5–7/week for 12 weeks Intensity: Moderate Time: ≥30 min	Home-based	Unsupervised: telephone counselling from a physical activity counsellor, pedometer and heart rate monitor
Raghavendra 2007	N=62 stage II–III BC on chemotherapy Exercise group n=28 II: 16 (57.1%) III: 12 (42.9%)	Frequency: 6/wk during the course of chemotherapy Intensity: Low Time: 60 min Type: Yoga	Hospital	Unsupervised and supervised: yoga instructor and home exercise video
Rao 2012	n=10; stage II-III BC. n=5 exercise; n=5 control;	Frequency: 3/week for 16 weeks Intensity: Not specified Time: 60 min Type: Combined resistance- and aerobic exercise (involving bouts of jumping jacks, running in place, arm and leg exercises with exercise balls, bands and weights)	Home- and community-based	Supervised: personal trainer
Rogers 2009	N=41 stage I-IIIA BC Intervention n=21: I: 6 (29) II: 11 (52) III: 4 (19)	Frequency: 3–5/week for 12 weeks Intensity: Moderate Time: 150 min/week Type: Walking	Centre- (unspecified) and home-based	Unsupervised and supervised: ACSM exercise specialist and/or certified exercise physiologist
Schmidt 2015	N=101 Stage I-IV BC starting CT Exercise group (n=49): I: 37 (38.9%) II: 41 (43.2%) III: 15 (15.8%) IV: 2 (2.1%)	Frequency: 2/week for 12 weeks Intensity: 3 sets, 8–12 repetitions at 60– 80% of 1RM Time: 60 min Type: Resistance exercise (8 different machine exercises)	Hospital	Supervised: Physical therapists
Schwartz 2007	N=66 stage I–III BC beginning adjuvant CT Aerobic group n=22: I: 4 (18%) II: 13 (59%) III: 5 (12%) Resistance group n=21: I: 6 (28) II: 11 (52%) III: 4 (19%)	Aerobic group: Frequency: 4/week for 6 months Intensity: Moderate intensity ("breathing hard but able to talk") Time: 15–30 min Type: Self-selected (e.g., walking or jogging) Resistance group: Frequency: 4/wk for 6 months	Home-based	Unsupervised: telephone contact from research staff

		Intensity: 2 sets of 8–10 repetitions Time: ~30 min Type: Thera-Band [™] exercises (4 upper body & 4 lower body exercises).		
Short 2015	Tailored group n=109 0: 3 (2.9%) I: 27 (26.5%) II: 32 (31.4%) III: 23 (22.6%) IV: 2 (1.9%) Unknown: 15 (14.7%) Targeted group n=110 0: 3 (2.8%) II: 22 (20.8%) II: 45 (42.5%) III: 20 (18.8%) IV: 1 (0.9%) Unknown: 15 (14.5%)	Frequency: 4–7/week for 12 weeks Intensity: Moderate Time: ≥30 min Type: Aerobic and resistance (Tailored- print intervention: three computer- tailored physical activity newsletters; targeted-print group: physical activity booklet)	Home-based	Unsupervised; tailored-physical activity print materials or targeted physical activity booklet
Vallance 2007	N=377 stage I–III BC completed treatment Pedometer group (n=94) I: 38 (40.4%) II: 50 (53.2%) III: 6 (6.4%)	Frequency: ≥5/week for 12 weeks Intensity: Moderate Time: ≥30 min Type: Aerobic	Home-based	Unsupervised.
Vallance 2015	N=95 stage I–III BC receiving adjuvant chemotherapy Intervention group (n=49) I: 10 (20%) II: 31 (63%) III: 8 (16%)	Frequency: ≥5/week for 12 weeks Intensity: Moderate Time: ≥30 min Type: Aerobic	Home-based	Unsupervised: tailored print materials and pedometer
Van Waart 2016	N=230 stage I-III undergoing adjuvant chemotherapy OnTrack (n=76) I: 5 (7) II: 32 (42) III: 39 (51) Onco-Move (n=77) I: 2 (3) II: 40 (52) III: 35 (45)	Onco-Move (n=77) Frequency: 5/week for duration of chemotherapy Intensity: 12–14 RPE Time: >30 min Type: Combined resistance and aerobic OnTrack (n=76) Frequency: 5/week for duration of chemotherapy Intensity: Aerobic: 50% to 80% of the maximal workload; Resistance: 6 exercises, 2 sets, 8 repetitions 80% of IRM Time: 50 min Type: Combined resistance and aerobic	Home-based, or Centre-based (unspecified)	Unsupervised, or Supervised: physical therapists
Wang 2011	N=72 Stage I–II undergoing CT Exercise group (n=35) I: 9 (25.7%) II: 26 (74.3%)	Type: Combined resistance and acrobic Type: Aerobic exercise, walking Frequency: 3–5/week for 6 weeks Intensity: 40–60% HR _{max} or Time: 30 min	Home-based	Unsupervised: weekly telephone calls

Winters Stone 2011	N=106 >1 year post-RT and/or CT	Type: Resistance exercise (+impact	University setting and home-based	Unsupervised and supervised: certified
	0: 7.7%	training)		exercise instructors
	I: 38.5%	Frequency: 2–3/week for 12 months		
	II: 48.1%	Intensity: 60–70% of 1-RM for 1–3 sets		
	IIIa: 1.9%	of 8–12 repetitions		
		Time: 45–60 min		
Winters-Stone 2013	71 BCS Stage I–IIIa: prematurely menopausal	Frequency: 3/week for 12 months (2	University setting	Supervised: certified exercise instructors
	Impact + resistance group (N=35)	supervised + 1 unsupervised)		
	I: 22.9 %	Intensity: 8–15 repetition maximum		
	II: 65.7 %	Time: 30-60 min/session		
	III: 11.4 %	Type: Free weights (e.g., dumbbells,		
		barbells, resistance bands, and weighted		
		vests Jump: 0-10% BW, 3-10 sets, 10		
		repetitions; Upper and lower-body RE:		
		2–3 sets per exercise, 6–14 repetitions		
		(6-14RM; upper body RE); 0-10% BW		
		(lower body RE)		
Yang 2010	N=40 stage I-II BC receiving adjuvant	Type: Aerobic	Home-based	Unsupervised: weekly telephone calls
-	chemotherapy	Frequency: 3/week for 12 weeks		
	I: 9 (47.4%)	Intensity: 60–80% of age-adjusted		
	II: 10 (52.6%)	maximal heart rate		
		Time: 30 min (plus 5 min warm-up and		
		5 min cool-down)		

¹Interventions were considered supervised if 50% or more of the prescribed exercise involved face-to-face supervision.

CERTEN

Supplementary Content 4: Summary of study recruitment, retention, adherence, reasons for withdrawal, intervention settings and supervision and exercise related events (n=61)

Study	Recruitment (eligible / total number screened)	Retention	Reasons for withdrawal	Adherence (to scheduled exercise sessions)	Location and supervision	Adverse events
Ahmed 2006	85/238 eligible & consented ² 64%	Exercise: 100% Baseline: n=23 Completed: n=23 Control: 96% Baseline: n=23 Completed: n=22	Exercise: n=0 Control n=1 Breast cancer recurrence n=1	92%	Recreation centre, supervised: ACSM certified fitness professional.	Exercise: Not reported Control: Not reported
Anderson 2012	104/625 eligible & consented 16%	Exercise: 83% Baseline: n=52 Completed: n=43 Control: 75% Baseline: n=52 Completed: n=39	Not reported by group: Feeling overwhelmed or a lack of time to participate (38%), lost to follow-up (19%), lack of interest (10%), family issues (10%), death (n=2, 10%), and other reasons (10%).	71.2%	University health and exercise research centre, supervised: Occupational or physical therapist.	Exercise: N=2 exercise related adverse events (n=1 pectoral muscle pain; n=1 stress fracture in foot). Control: Not reported
Banerjee 2007	Not reported	Exercise: 100% Baseline: n=35 Completed: n=35 Control: 70% Baseline: n=33 Completed: n=23	Exercise: n=0 Control: n=10 Reason not reported	Not reported.	Hospital outpatient, supervised: Yoga instructors + Home, unsupervised	Exercise: Not reported Control: Not reported
Campbell 2017	19/102 eligible and consented 18.6%	Exercise: 100% Baseline: n=10 Completed: n=10 Control: 100% Baseline: 9 Completed: n=9	Exercise: n=0 Control: n=0	Overall: 87.5% Supervised gym: 88% Unsupervised home: 87%	Research gym, supervised: not reported by whom + Home, unsupervised	Exercise: No adverse events occurred. Control: No adverse events occurred.
Cantaro- Villanueva 2012a	40/62 eligible & consented 65%	Exercise: 100% Baseline: n=20 Completed: n=20 Control: 100% Baseline: n=20 Completed: n=20	Exercise: n=0 Control: n=0	79%	University medical centre outpatient clinic and swimming pool, supervised: Exercise trainer specialist and physiotherapists	Exercise: N=4 in the hydrotherapy group showed a temporal (1–3 days) increase of pain after one session, but this event did not stop them continuing the programme. Control: No further adverse events were reported.
CantarerVillanueva 2012b	78/238 eligible and consented 33%	Exercise: 84% Baseline: n=38 Completed: n=32 Control: 88% Baseline: n=40 Completed: n=35	Exercise: n=6 (health problems n=1; family problems n=1; never started program n=2; too busy n=2) Control: n=5 (not contactable n=1; absent from test n=4)	Overall: 83.5% Completed treatment ≤6 months: 79.6% Completed treatment >6 months:	University medical centre outpatient clinic and swimming pool, supervised: not reported	Exercise: No exercise-related adverse events Control: No exercise related adverse events

				87.4%		
CantareVillanueva 2012c;	66/95 eligible and consented 69%	Exercise: 97% Baseline: n=33 Completed: n=32 Control: 100% Baseline: n=33 Completed: n=33	Exercise: n=1 (breast cancer recurrence) Control: n=0	>85%	University medical centre outpatient clinic and swimming pool, supervised: Physical therapist	Exercise: N=3: transient increase of edema, N=4: increase in fatigue immediately after the beginning of the first session, which improved in the next few days. Control: None
CantareVillanueva 2013	68/163 eligible and consented 42%	Exercise: 94% Baseline: n=34 Completed: n=32 Control: 85% Baseline: n=34 Completed: n=29	Exercise: n=2 (did not commence n=1; too busy n=1) Control: n=5 (not contactable n=1; absent from test n=4)	84%	University medical centre outpatient clinic and swimming pool, supervised: Exercise specialist and physical therapists	Exercise N=3: discomfort or low-intensity pain/stiffnes after an exercise session (nevertheless, they continued the program.) Control: None
Chandwani 2014	178/294 eligible and consented 61%	Exercise: 81% Baseline: n=53 Completed: n=43 Control (WL): 86% Baseline: n=54 Completed: n=46	Exercise: n=10 reasons not reported Control: n=8 reasons not reported	78%	Cancer treatment centre, supervised: certified yoga instructor	Exercise: Not reported. Control: None reported
Cormie 2013	62/135 eligible & consented 46%	High-load: 86% Baseline: n=22 Completed: n=19 Low-load: 100% Baseline: n=21 Completed: n=21 Control: 89% Baseline: n=19 Completed: n=17	High-load exercise (n=3): Unrelated medical condition n=1; time constraints n=2 Control (n=2): Unrelated medical condition n=1; time constraints n= 1	High-load exercise: 96% Low-load exercise: 96%	Hospital/health clinic, supervised: Accredited exercise physiologist	No exercise-related adverse events.
Cornette 2016	44/89 eligible and consented 49%	Exercise: 68% Baseline: n=22 Completed: n=15 Control: 68% Baseline: n=22 Completed: n=15	Exercise (n=7): N=2 excluded (n=1 did not complete baseline CTEP; n=1 using beta-blockers); n=5 no reason Control (n=7): n=7 no reason	88%	Home-based, unsupervised	No exercise-related adverse events.
Courneya 2003	53/370 eligible and consented 14%	Exercise: 96% Baseline: n=25 Completed: n=24 Control: 93% Baseline: n=28 Completed: n=26	Exercise: N=1 gastrointestinal complication Control: n=2 N=1 orthopaedic complication; n=1 Bronchitis	98%	Cancer institute and University, supervised: Accredited exercise physiologist	Five participants (20.8%) in the exercise group experienced an adverse event compared with two participants (7.1%) in the control group. The adverse events in the exercise group were lymphedema (n= 3), gynecologic complication (n=1), and influenza (n=1). Control: The control group's events were foot fracture (n =1) and bronchitis (n=1).
Courneya 2007	242/1468 eligible and	Aerobic: 95% Baseline: n=78	Aerobic n=4 Reasons not reported	Aerobic 72.0%	Cancer institute and University, supervised: Accredited exercise	Exercise N=2 after baseline maximal treadmill testing (n=1 light

	consented 16%	Completed: n=74 Resistance: 94%	Resistance n=5	Resistance 68%	physiologist	headedness, hypotensive, and moderately nauseous; n=1dizziness
	10%	Baseline: n=82	Reasons not reported	0870		weakness,
		Completed: n=77	-			and mild diarrhoea).
		Control: 89%	Control n=9			Control: none
		Baseline: n=82	Reasons not reported			
D 6 1 - 001 6	014/1400	Completed: n=73	TY 11 . 1 1. • 1	6604		XY 1.11
Defelson 2016	214/1400 eligible and	Exercise: 100% Baseline: n=37	Unable to be determined.	66%	Hospital, supervised: not reported.	No exercise-related adverse events.
	consented	Completed: n=37				
	15.2%	Control: 100%				
	1012/0	Baseline: n=37				
		Completed: n=37				
Danhauer 2009	44/299	Exercise: 59%	Exercise (n=9):	60%.	Yoga studio, supervised: Certified	No exercise-related adverse events.
	responded,	Baseline: n=22	N=9 did not return		yoga instructor	
	eligible and	Completed: n=13	questionnaire (lost to follow			
	consented	Control: 64%	up)			
	15%	Baseline: n=22	Control (n=8):			
		Completed: n=14	N=7 did not return			
			questionnaire (lost to follow			
			up); N=1 dropped out of study	1		
					N .	
De Luca 2016	Not	Exercise: 100%	Exercise n=0	Not reported	University gymnasium, supervised:	No exercise-related adverse events.
	reported.	Baseline: n=10			Fitness professional and physician	
		Completed: n=10	Control n=0			
		Control: 100%				
		Baseline: n=10 Completed: n=10				
Dolan 2016	36/59	Interval: 100%	Interval: n=0	Interval exercise:	Location not specified, supervised:	No exercise-related adverse events.
2010	eligible and	Baseline: n=12	interval n=0	98%	Accredited exercise physiologist	The exercise related adverse events.
	consented	Completed: n=12	Continuous: n=1	Continuous	r j. g.	
		Continuous: 92%	Reason not reported	exercise:		
	61%	Baseline: n=12		98%		
		Completed: n=11	Control: n=2			
		Control: 83%	Reason not reported			
		Baseline: n=12				
Drouin 2005	23/39	Completed: n=10 Exercise: 100%	Exercise: N=0	Mean $= 3.6$	Home-based, unsupervised	Not reported.
D10u111 2005	eligible and	Baseline: n=13	Excicise. IN-0	days/week that	nome-based, unsupervised	The reported.
	consented	Completed: n=13	Control: N=2	aerobic exercise		
	59%	Control: 80%	personal commitments	was performed.		
		Baseline: n=10				
		Completed: n=8	r			
Eakin 2012	143/383	Exercise: 93%	Exercise (n=5):	88%	Home-based, telephone delivered:	N=3: muscle soreness (n=2);
	eligible and	Baseline: n=73	n=4 health concerns; n=1 no		Accredited exercise physiologist	musculoskeletal injury (n=1).
	consented	Completed: n=68	longer has cancer			
	37%	Control: 99%				

		Baseline: n=70	Control (n=1):			
Galiano-Castillo 2017	81/99 eligible and consented 82%	Completed: n=69 Exercise: 87.8% Baseline: n=41 Completed: n=36 Control: 87.8% Baseline: n=41 Completed: n=36	n=1 health concerns Exercise n=5 Busy n=1 Health problems n=3 Not reported n=1 Control n=5 Busy n=3 Personal problems n=1 Death n=1	93.9%	Home-based, internet-based (tele- rehabilitation): unsupervised	Exercise: no intervention-related adverse events. Control: no intervention-related adverse events.
Gokal 2015	63/164 eligible and consented 38%	Exercise: 84% Baseline: n=25 Completed: n=21 Control: 100% Baseline: n=25 Completed: n=25	Exercise: n=5: Hospitalisation n=4; Medical difficulties n=1 Control: n=0	80%	Home-based, unsupervised	Not reported
Guinan 2013	26/32 eligible & consented 81%	Exercise: 88% Baseline: n=16 Completed: n=14 Control: 80% Baseline: n=10 Completed: n=8	Exercise group: N=2: N=2 time constraints Control group: N=2: N=2 illness unrelated to their breast cancer.	Not reported	Location not specified, supervised: Physiotherapist and a research assistant + Home-based, unsupervised	Not reported.
Fernández-Lao 2013	98/132 eligible and consented 74%	Land-based exercise Baseline: n=31 ^{CC} Water-based exercise Baseline: n=33 ^{CC} Control Baseline: n=34 ^{CC}	Not reported	Land-based: 85% Water-based: 92%	A gymnastic hall and heated swimming pool, supervised: Fitness specialist and physical therapists	Not reported.
Hatchett 2013	85/200 eligible & consented 42.5%	Exercise: 88% Baseline: n=43 Completed: n=38 Control: 86% Baseline: n=42 Completed: n=36	Exercise n=5: Discontinued participation (n=5) Control n=6: Discontinued participation (n=6)	Not reported	Home-based, unsupervised email delivered intervention: e-counselor exercise physiologist	Not reported
Hayes 2012	194/402 eligible & consented 48%	Exercise: 91% Baseline: n=67 Completed: n=61 Telephone: 94% Baseline: n=67 Completed: n=63 Control: 93% Baseline: n=60 Completed: n=56	N=14 ^{LL99} (Reasons: too busy (n=4); unhappy with allocation (n=2); not coping with treatment (n=2); unknown (n=2); unable to contact/passive withdrawal (n=2); Reasons: no longer interested (n=2))	Exercise: 88% Telephone: 81%	Home-based, supervised and unsupervised: face-to-face or telephone contact with accredited exercise physiologist	No exercise-related adverse events.

Headley 2004	Not reported	84% Baseline: n=38 Completed: n=32	n= 6; disease progression.	75%	Cancer centre outpatient clinic, supervised: Oncology nurse	No exercise-related adverse events.
Herrero 2005	20/37 eligible & consented 54%	Exercise: 80% Baseline: n=10 Completed: n=8 Control: 80% Baseline: n=10 Completed: n=8	Exercise: N=2; Reasons not reported Control: n=2; Reasons not reported	91%	Community fitness centre, supervised: Exercise physiologists	No exercise-related adverse events.
Hornsby 2014	20/1445 eligible & consented 1%	Exercise: 90% Baseline: n=10 Completed: n=9 Control: 100% Baseline: n=10 Completed: n=10	Exercise: n=1; and DVT and PE Control: n=0	82%	Cancer institute, supervised: Accredited exercise physiologist	Exercise: n=1 (unexplained leg pain that quickly resolved following exercise cessation); n=3 during exercise testing (n=1 exercise-induced oxygen desaturation, SpO ² 84%), n=1 anxiety attack, n=1 dizziness). Exercise:N=7 events (persistent tachycardia n=1, diverticulosis n=1, urinary tract infection (UTI) n=1, diabetes mellitus n=1, upper respiration tract infection n=1, hemorrhoids n=1; and DVT and PE n=1 (more than one event was observed in the same patient) Control:N=1 shingles to secondary to varicella zoster infection
Husebø 2014	67/93 eligible & consented 72%	Exercise: 76% Baseline: n=33 Completed: n=25 Control: 85% Baseline: n=34 Completed: n=28	Exercise: n=8 (n=7 no reason reported; n=1 syncope due to a comorbid condition) Control: n=6 (no reason reported)	58%	Home-based, unsupervised	Exercise N=1 reported knee discomfort (remained in trial); n=1 syncope during the walking exercise (related to a secondary chronic condition, withdrew from trial). Control N=0
Hutnick 2005	Not reported.	Exercise: 75% Baseline: n=28 Completed: n=21 Control: 71% Baseline: n=21 Completed: n=15	Exercise: n=7 (reasons not reported) Control: n=6 (reasons not reported)	Overall: 79% Months 1-3: 82.2% Months 4-6: 75.9%	University clinical setting, supervised: exercise trainer And/or Home-based, unsupervised (periodic contact with exercise trainer)	Not reported.
Kilbreath 2012	160/457 eligible & consented 35%	Exercise: 95% Baseline: n=81 Completed: n=77 Control: 93% Baseline: n=79 Completed: n=74	Exercise: N=4 time constraints Control: N=5 (n=3 time constraints, n=1 developed metastases, n=1 unable to contact).	Overall: 84% Supervised: 78% Unsupervised: 90%	Location and supervision not specified + Home-based, supervision not specified	Not reported.
Kim 2006	Not reported.	Exercise: 59% Baseline: n=37 Completed: n=22 Control: 51% Baseline: n=37	Exercise: N=5 intervention withdrew Control: N=6 control withdrew Personal problems $(n = 2)$, problems at home $(n = 2)$,	78%	University exercise facility, supervised: Exercise physiologists	No exercise-related adverse events. Exercise: Not reported. Control: None Baseline testing: N=2 ECG abnormality or hypertensive episodes during baseline graded exercise testing.

			1	1	l	
		Completed: n=19	problems related to chemotherapy ($n = 3$), thrombophlebitis in the lower leg ($n = 2$), non-exercise- related injuries ($n = 1$), or death ($n = 1$). N=12 control missed either a pre- or post-intervention graded exercise test. N=10 intervention missed either a pre- or post- intervention graded exercise test.		R.	
Ligibel 2008	101/199 eligible & consented 51%	Exercise: 78% Baseline: n=51 Completed: n=40 Control: 84% Baseline: n=49 Completed: n=42	Exercise: $n=11$ Lost to follow-up ($n = 2$)Family emergency ($n=1$); too much of a time commitment ($n=3$); too ill for final measurements ($n=1$); disease recurrence ($n=1$), developed unrelated cancer ($n=1$), withdrew consent ($n=1$), need for unrelated surgery ($n=1$)	73%	Location and supervision not specified + Home-based, unsupervised	Not reported.
			Control: n=7 Lost to follow-up (n = 3) Disease recurrence (n=2), withdrew upon assignment to control group (n=1), family problems (n=1)	P		
Ligibel 2016	Not reported.	Exercise: 68% Baseline: n=48 Completed: n=33 Control Baseline: n=53 Completed: n=43 81%	Exercise group (n=15): n=4 stopped attending and unreachable by study team; n=4 time and travel reasons; n=3 disease progression; n=1 deceased due to disease; n=2 moved during intervention n=1 no reason Control group (n=10): n=5 unreachable by study team; n=2 disease progression; n=1 time and travel reasons; n=1 no reason; n=1 ineligible due to active brain metastases	Not reported	Home-based, supervised and unsupervised: Exercise physiologist	No exercise-related adverse events.
Loudon 2014	28/59 eligible & consented 47%	Exercise: 80% Baseline: n=15 Completed: n=12 Control: 85%	Exercise n=3 Surgery n=1; broken hip n=1; acute illness n=1 Control n=2	Overall: 92% Home-practice: 86% Group yoga	Location not specified, supervised: certified yoga instructor + Home-based, unsupervised	No exercise-related adverse events.

		Baseline: n=13 Completed: n=11	Family reasons n=1; acute illness n=1	sessions: 97%		
Macvicar 1989	Not reported	72% Baseline: n=62 Completed: n=45	n=9 disease progression; n=1 transportation problems; n=2 commenced cardio-toxic medications; n=2 extreme chemotherapy associated side effects; n=3 equipment failure.	Not reported.	Location not specified, supervision not specified	Not reported.
Maryam 2010	Not reported	Exercise: 100% Baseline: n=28 Completed: n=28 Control: 100% Baseline: n=28 Completed: n=28	Exercise: n=0 Control: n=0	Not reported.	Home-based, unsupervised	Not reported.
Milne 2008	58/131 eligible & consented 44%	Exercise: 100% Baseline: n=29 Completed: n=29 Control: 100% Baseline: n=29 Completed: n=29	Exercise: n=0 Control: n=0	60%	Rehabilitation clinic, supervised: Exercise physiologists	Not reported.
Moadel 2007	164/193 eligible & consented 85%	Exercise: 78% Baseline: n=108 Completed: n=84 Control: 73% Baseline: n=56 Completed: n=44	Exercise n=24 Loss to follow up: 16; Refused: 5; Change in health status: 3 Control; n=12 Loss to follow up: 8; Refused: 3; Change in health status: 1	58%	Cancer centre, supervised: Oncologist and certified yoga instructor	Not reported.
Mohan Rao 2015	Not reported	Exercise: 73% Baseline: n=45 Completed: n=33 Control: 68% Baseline: n=53 Completed: n=36	Exercise n=12 Reason not reported Control n=17 Reason not reported	Not reported.	Hospital, supervised: yoga instructor	Exercise: N=2 (infections, secondary suturing, seroma, discharge , uncontrollable pain) Control: N=8 (infections, secondary suturing, seroma, discharge, uncontrollable pain) *2008 Rao paper included:
Mulero Portela 2008	Not reported	Gym exercise: 75% Baseline: n=16 Completed: n=12 Home exercise: 68% Baseline: n=19 Completed: n=13 Control: 100% Baseline: n=9 Completed: n=9	Gym exercise: N=4; moved to the United States (n=1); developed eye cancer (n=1); developed headaches with referral or MRI(n=1); foot surgery (n=1) Home exercise: N=6; developed uterine cancer (n=1); no show with no reason given (n=1); asthma complications and non-clearance from	Gym exercise Overall: 55% Aerobic exercise: 47% Resistance exercise: 63% Home exercise Overall: 79% Aerobic exercise: 71% Resistance exercise: 86%	Gymnasium, supervised and unsupervised: physical therapists or Home-based, supervised and unsupervised: physical therapists	Gym exercise; n=1 hypoglycaemia while at the gym during an exercise; n=1 high blood pressure (>140/90 mmHg) during their participation in the exercise programs; n=1 severe headache at during post-intervention exercise testing; n=1 foot pain which worsening during the first exercise session leading to study withdrawal and surgery. Home exercise N=1 asthma episode during the 12- minute walk

			physician to continue (n=1); personal problems (n=1); high blood pressure with referral for stress test (n=1); discontent with schedule (n=1). Control: N=0		Å	test at baseline (leading to withdrawal prior to commencing intervention); n=2 high blood pressure (>140/90 mm Hg) during their participation in the exercise programs
Musanti 2012	55/314 eligible & consented 18%	Overall 76% Baseline: n=55 Completed: n=42 Aerobic Baseline: ? Completed: n=12 Resistance Baseline: ? Completed: n=17	N=7 difficulty fitting the exercise into their lives because of work and/or family responsibilities; n=1 breast reconstruction surgery rescheduled; n=1 one did not give a reason; n=1 could not complete the initial fitness testing because of an elevated HR; n=1 wanted more supervised exercise;N=1 appendicitis	Aerobic exercise: 81% Resistance exercise: 91%	Home-based, unsupervised	Exercise: N=2 tendonitis (n=1 shoulder, n=1 foot) Control: none
Murtezani 2014	73/241 eligible & consented 30%	Exercise: 81% Baseline: n=37 Completed: n=30 Control: 89% Baseline: n=36 Completed: n=32	Exercise n=7 Transportation difficulties n=3; lymphoedema n=3; low back pain n=1 Control n=4 Gynaecologic problems n=1; unreachable n=2; personal reason n=1	85%	University clinical rehabilitation centre, supervised: not reported by whom	Not reported.
Naraphong 2015	26/177 eligible & consented 15%	Exercise: 81% Baseline: n=11 Completed: n=9 Control: 100% Baseline: n=12 Completed: n=12	Exercise n=2: Moved & withdrew from care at the site at week 7 (n=1); Too busy for exercising at week 10 (n = 1) Control n=0	Not reported	Home- and community-based, unsupervised (weekly contact with a nurse)	Not reported.
Naumann 2012	40/48 eligible & consented 83%	Exercise: 93% Baseline: n=15 Completed: n=14 Control: 83% Baseline: n=12 Completed: n=10	Group exercise: N=1 (n=1 unrelated injury) Control: N=2 (n=1 unrelated injury; failed to commence participation n=1)	74%	Gymnasium, supervised: Accredited exercise physiologist	No exercise-related adverse events.
Pinto 2005	86/424 eligible & consented 20%	Exercise: 90.7% Baseline: n=43 Completed: n=39 Control: 100% Baseline: n=43 Completed: n=43	Exercise (n=4): n=1; could not be contacted to determine reasons, n=2; and participation terminated, n=1; the study team terminated one woman's participation because of symptoms of chest pain during exercise and her refusal to have these symptoms	Not reported	Home-based, unsupervised: telephone support from research staff	Not reported

			evaluated by her physician).			
			Control (n=0)			
Pinto 2013	192/351 eligible & consented 55%	Exercise: 79% Baseline: n=106 Completed: n=89 Control: 91% Baseline: n=86 Completed: n=84	Exercise (n=17): Lost contact=8, family issues=4, cancer=2, no interest=2, too busy=1 Control (n=2): Lost contact=2	Not reported	Home-based, unsupervised: telephone counselling: physical activity counsellors	N=1 sustained minor injuries related to falling off a treadmill, n=1 died during the trial for reasons unrelated to study participation.
Pinto 2015	76/595 eligible & consented 13%	Intervention: 92% Baseline: n=39 Completed: n=36 Control: 86% Baseline: n=37 Completed: n=32	Intervention (n=3): nonresponsive (n = 2), health issues (n = 1) Control (n=5): nonresponsive (n=2), too busy (n=2), physical health issues (n=1)	92%	Home-based, unsupervised: telephone counselling; physical activity counsellors	Intervention: chest pain and shortness of breath during exercise (n=6), vertigo (n=1), and ankle injury (n=4) Control: none
Raghavendra 2007	98/174 eligible & consented 56%	Exercise: 62% Baseline: n=45 Completed: n=28 Control: 64% Baseline: n=53 Completed: n=34	Intervention (n=17): Reason not reported Control (n=19): Reason not reported	Not reported.	Hospital, supervised: yoga instructor	Not reported.
Rao 2012	Not reported	100% Exercise: 100% Baseline: n=5 Completed: n=5 Control: 100% Baseline: n=5 Completed: n=5	Intervention (n=0): Control (n=0):	80%	Home- and community-based, supervised: personal trainer	Not reported.
Rogers 2009	41/119 eligible and consented 34%	Exercise: 95% Baseline: n=21 Completed: n=20 Control: 95% Baseline: n=20 Completed: n=19	Exercise: N=1 due to unrelated medical problems Control: N=1 due to travel distance	Overall: 99% Individual sessions: 100% Group sessions: 98%	Location not specified, supervised: ACSM exercise specialist and/or certified exercise physiologist + Home-based, unsupervised	No exercise-related adverse events; The following, non- exercise related events were recorded: wheezing requiring physician evaluation for asthma, cholinergic urticaria, herpes zoster, sinusitis, back pain related to falling, and elective cosmetic reconstructive surgery-non reported by group
Schmidt 2015	101/121 eligible & consented 83%	Exercise group: 98% Baseline: n=52 Completed: n=51 ^{GG} Control group: 94% Baseline: n=49 Completed: n=46	Exercise group (n=1): N=1 psychological problems Control group (n=3): N=1 disliked intervention; n=1 time constraints; n=1 death	71%	Hospital, supervised: physical therapists	No exercise-related adverse events.

Schwartz 2007	72/75	Aerobic group:	N=6 Too busy $(n = 4)$ or the	Not reported.	Home-based, unsupervised	Not reported.
	eligible & consented	92% Baseline: n=24	location was not convenient (n $= 2$).			
	96%	Completed: n=22				
		Resistance: 91% Baseline: n=23	N=4 exercise N=2 control			
		Completed: n=21	*Unable to determine whether			
		Control: 92%	withdrew prior or post-		\mathcal{R}	
		Baseline: n=25	randominisation			
Short 2015	330/349	Completed: n=23 Tailored-print:	Tailored-print (n=11):	NA	Home-based, unsupervised; tailored-	
511011 2015	eligible	89%	1 poor health; 2 no reason	INA	physical activity print materials or	
	95%	Baseline: n=109	given; 8 non responders		targeted physical activity booklet	
		Completed: n=98				
		Targeted-	Targeted-booklet (n=12):		S	
		booklet: 88% Baseline: n=110	8 non responders; 1 deceased; 9 non responders			
		Completed: n=97	non responders			
		Control group:	Control group (n=7):			
		93%	7 non responders			
		Baseline: n=111		· · · · · · · · · · · · · · · · · · ·		
		Completed: n=104			X'	
Vallance 2007	377/1590	Pedometer	Exercise (n=6):	Not reported	Home-based, unsupervised.	Not reported.
	eligible &	group: 94%	n=6 loss to follow-up		1 I	1
	consented	Baseline: n=94				
	24%	Completed: n=88	Control (n=11):			
		Control group: 89%	n=1 hadn't kept up with program; n=10 loss to follow-			
		Baseline: n=96	up.			
		Completed: n=85	-			
Vallance 2015	95/123	Intervention	Intervention (n=8):	95%	Home-based, unsupervised: tailored	Not reported.
	eligible & consented	group: 83.67% Baseline: n=49	No response (n=8)		print materials and pedometer	
	77%	Completed: n=49	Control (n=9):	r		
	/ / /0	Control group:	No response $(n=8)$; Passed			
		80.4%	away (n=1)			
		Baseline: n=46				
Van Waart 2016	230/536	Completed: n=37	High intensity n=5	High intensity:	Location not apositized sumarized	Iliah intensity oversion
van waart 2016	eligible &	High-intensity: 93%	High-intensity: n=5 n=2 felt to ill, n=1 physical	High-intensity: 71%.	Location not specified, supervised: supervised by specially trained	High-intensity exercise n=1 unspecified physical accident related to trial
	consented	Baseline: n=76	accident unrelated to trial, n=1	Low-intensity	physical therapists	Low-intensity exercise
	43%	Completed: n=71	physical accident related to	Attendance of	or	Not reported
		Low-intensity:	trial, n=1 unwilling	planned sessions:	Home-based, unsupervised	Control
		89%	I and internation of Q	N/A.		Not reported
		Baseline: n=77 Completed: n=69	Low-intensity: n=8 N=1 neuropathy, n=1			
		Control: 86%				
			emigrated, n=6 unwilling			

		Baseline: n=77 Completed: n=66	Control: n=11 n=2 felt to ill, n=7 unwilling, n=2 unknown.			
Wang 2011	72/160 eligible and consented 45%	Exercise: 86% Baseline: n=35 Completed: n=30 Control: 86% Baseline: n=37 Completed: n=32	Exercise n=5; Discomfort with exercise n=1; dizziness n=1; Dyspnoea n=1; Too busy n=1; No family support n=1 Control n=5: Anaemia n=1; Moved n=1; Prolonged treatment n=1; Progressed to metastatic disease n=1; Holiday n=1	93%	Home-based, unsupervised	Anemia n=1- control Dizziness with dyspnea n=1- exercise
Winters Stone 2011	106/359 eligible and consented 30%	Exercise: 69% Baseline: n=52 Completed: n=36 Control: 57% Baseline: n=54 Completed: n=31	Exercise: n=16 Lost to follow-up n=8; Too busy: n=5; Poor health: n=1; Dislike: n=1; Moved: n=1 Control: n=23 Lost to follow-up n=12; Too busy: n=4; Poor health: n=4; Dislike: n=3	57%	University setting, supervised: certified exercise instructors + Home-based, unsupervised	No exercise-related adverse events.
Winters-Stone 2013	71/258 eligible & consented 28%	Exercise: 66% Baseline: n=35 Completed: n=23 Control: 69% Baseline: n=36 Completed: n=25	Exercise: n=12 Reasons: Too busy (n=6); Poor health (n=1); Disinterested (n=1); Lost to follow-up (n=4) Control: n=11 Reasons: Too busy (n=5); Inconvenient (n=1); Cancer recurrence (1); Pregnancy (n=1); Lost to follow-up (n=3)	Overall: 44% Supervised sessions: 64% Home-based sessions: 26 %	University setting, supervised: certified exercise instructors	POWIR stopped increasing vest weight at month 6 due to back (N=2) or knee (N=1) pain, and one participant stopped lower body exercises at month 5 due to pain,
Yang 2010	Not reported	Intervention: 100% Baseline: n=19 Completed: n=19 Control: 100% Baseline: n=21 Completed: n=21	Intervention (n=0) Control (n=0)	77%	Home-based, unsupervised	No exercise-related adverse events.

Supplementary Content 5. Overview reasons for withdrawals across all trials (n=61).

		n intervention group	Withdrawals from usual care group				
		out of total 2621 participants)	n=256 (9% withdrawals of				
	≤ 12 week interventions n=167	>12 week interventions n=144	≤ 12 week interventions n=124	>12 week interventions n=132			
Reason for	Health-related reasons n=33:	Health-related reasons n=27:	Health-related reasons n=15:	Health-related reasons n=17:			
withdrawals	Unspecified health or medical problems or	Unspecified health or medical problems or	Medical condition or illness unrelated to	Unspecified health or medical problem			
	deterioration of health n=10	deterioration of health n=9	breast cancer n=3	or deterioration of health n=8			
	Hospitalisation n=4	Breast cancer progression n=3	Unspecified health or medical problems or	Disease recurrence n=4			
	Lymphoedema n=3	Breast cancer recurrence n=2	deterioration of health n=1	Disease progression n=2			
	Unrelated medical condition n=2	No longer has cancer n=1	Developed metastases n=2	Orthopaedic complication n=1			
	Cancer $n=2^1$	Syncope due to a comorbid condition n=1	Acute illness n=1	Bronchitis n=1			
	Breast cancer recurrence n=1	Gastrointestinal complication n=1	Unrelated (unspecified) injury n=1	Became ineligible due to active brain			
	Deep vein thrombosis with pulmonary	Need for unrelated surgery n=1	Gynaecologic problems n=1	metastases n=1			
	embolism n=1	Developed other cancer $n=3^2$	Physical health issues n=1				
	Surgery n=1	Developed headaches with referral for MRI n=1	Anaemia n=1	Non-health-related reasons or other			
	Broken hip n=1	Foot surgery n=1	Prolonged treatment n=1	n=115:			
	Acute illness n=1	Asthma complications and non-clearance from	Death n=3	No reason for withdrawal or reason no			
	Low back pain n=1	physician to continue n=1		reported n=65			
	Unrelated (unspecified) injury n=1	High blood pressure with stress test referral n=1	Non-health-related reasons or other	Lost to follow-up n=18			
	Psychological problems n=1	Neuropathy n=1	n=109:	Too busy n=9			
	Symptoms of chest pain during exercise and	Death n=1	No reason for withdrawal or reason not	Unwilling to continue n=7			
	refusal to have these symptoms evaluated by		reported n=32	Uncontactable or non-responder n=5			
	her physician n=1	Non-health-related reasons or other n=117:	Did not return questionnaire or lost to	Dislike n=3			
	Discomfort with exercise n=1	No reason for withdrawal or reason not reported	follow up n=25	Time and travel reasons n=2			
	Dizziness n=1	n=62	Uncontactable or non-responder n=24	Did not complete baseline testing for			
	Dyspnoea n=1	Lost to follow-up n=14	Absent from test n=8	unspecified reason n=1			
	Death n=1	Too busy n=11	Time constraints or too busy n=10	Using beta-blockers n=1			
		Time and travel reasons n=7	Family or personal reasons n=5	Unhappy with group assignment n=1			
	Non-health-related reasons or other n=134:	Unwilling to continue for unspecified reason	Failed to commence participation for	Family problems n=1			
	Uncontactable or non-responder n=44	n=7	unspecified reason n=1	Inconvenient n=1			
	Did not return questionnaire (lost to follow up)	Moved during intervention n=5	Disliked intervention n=1	Pregnancy n=1			
	N=31	Uncontactable or non-responder n=4	Hadn't kept up with program n=1				
	No reason for withdrawal or reason not	Family or personal reasons n=2	Moved n=1				
	reported n=26	Discontent with schedule n=1	Holiday n=1				
	Time constraints n=8	Unspecified physical accident n=2					
	Too busy n=7	Dislike or disinterested n=2					
	Refused to continue for unspecified reason n=5						
	Family reasons n=5						
	Transportation difficulties n=3						
	Disinterested n=2						
	Moved & withdrew from care at the site n=1						
	No family support n=1						

¹ No further specification reported (i.e., unable to determine whether events were a cancer recurrence, cancer progression or development of new cancer) ² Other cancers were uterine (n=1); eye (n=1); and unspecified (n=1).

Supplementary Content 6. Overview of health outcomes and methods of assessment across all trials (n=61).

Outcome	Instrument/methods and number of studies
Quality of life (n=32)	FACT-B or FACT-B+4, n=14 ¹⁻¹⁴
	EORTC QLRC30, n=8 ¹⁵⁻²²
	Medical Outcomes Study 36-item short-form survey, n=3 ²³⁻²⁵
	FACIT-F, n=2 ^{26, 27}
	Functional Living Index of Cancer, n=2 ^{28, 29}
	FACT–Anemia scale, n=1 ³⁰
	FACT-G, $n=1^{31}$
	Lymphoedema QOL scale, n=1 ³²
	QOL-BC, $n=1^{33}$
Aerobic fitness (n=25)	VO ₂ peak testing using a modified Bruce treadmill protocol, $n=4^{19,34\cdot36}$
Refoble filless (II=25)	6-minute walk test, $n=3^{1,31,37}$
	12-minute walk test, $n=3^{8, 10, 38}$
	VO ₂ max or VO ₂ peak assessed using a cycle ergometer, $n=8^{2,5,15,17,26,39,41}$
	VO ₂ max or VO ₂ peak testing on a treadmill, $n=3^{30,42,43}$
	Submaximal treadmill test using the Naughton protocol with the end point of 85% of predicted HRmax, $n=1^{11}$
	Heart rate on completion of 3-minute step test, $n=1^4$
	Steep Ramp Test: maximal short exercise capacity, $n=1^{21}$
	Submaximal aerobic power cycle test, $n=1^6$
T	Rockport 1-mile test, n=1 ⁴⁴ FACT-F, n=12 ^{3-5, 7, 12-14, 19, 26, 27, 31, 45}
Fatigue (n=28)	FAC1+F, n=12 ⁻⁵⁵ , tiefer 17, et al. (31, 15), to
	Piper Fatigue Scale, $n=6^{22, 34, 36, 46, 48}$
	Schwartz Cancer Fatigue Scale-6 (SCFS-6), n=2 ^{6,37}
	Profile of mood states fatigue scale, $n=1^{49}$
	Brief Fatigue Inventory, n=1 ²³
	Multidimensional fatigue inventory- MFI-20, n=1 ¹⁵
	13-item Fatigue Scale, n=1 ²
	Functional Assessment of Cancer Therapy–Anemia fatigue scale, n=1 ³⁰
	Medical Outcomes Survey Short Form (SF)–36 fatigue symptoms, n=1 ⁵⁰
	Visual analogue scales, $n=1^{32}$
	Fatigue Assessment Questionnaire (FAQ), n=1 ²⁰
	Multidimensional Fatigue Inventory, $n=1^{21}$
	MDASI-T, $n=1^{51}$
Upper-body strength	1RM chest press, n=4 ^{24, 39, 52, 53}
(n=17)	Handgrip dynamometer, n=2 ^{10, 11, 22}
	1RM bench press, $n=1^{54}$
	Dynamometer elbow flexion, $n=1^{21}$
	Overhead press 1RM, $n=1^{38}$
	Bicep curl 1RM, $n=1^{42}$
	6 RM chest press, $n=1^{36}$
	Chest press – method or RM not specified, $n=1^6$
	Shoulder muscle strength using hand-held dynamometer, n=1 ¹⁸
	Bench press dynamic muscle strength tests performing as many reps with $100-110\%$ BW, $n=1^{17}$
	Shoulder press – method or RM not specified, $n=1^4$
	Shoulder press 1RM estimate based multiple repetition procedure, $n=1^{26}$
	8RM on the horizontal bench press, $n=1^{30}$
Anxiety (n=16)	Profile of Mood States-Anxiety, $n=6^{7, 34, 41, 47, 49, 51}$
Tunxiety (ii=10)	Hospital Anxiety and Depression Scale, n=4 ^{15, 36, 45, 55}
	State Trait Anxiety Inventory, $n=2^{12,29}$
	Functional Assessment of Cancer Therapy–Anemia scale Anxiety subscale, $n=1^{30}$
	Greene Climacteric Scale, n=1 ⁴
	Spielberger State Anxiety Scale, n=1 ⁴³
	Social Physique Anxiety Scale, n=1 ⁶
	Social Physique Anixety Scale, n=1
D : (16)	Functional Living Index of Cancer, $n=1^{28}$
Depression (n=16)	Profile of mood states-depression, n=5 ^{34, 41, 47, 49, 51} The Hospital Anxiety and Depression Scale, n=4 ^{15, 36, 45, 55}
	The Hospital Anxiety and Depression Scale, n=4 ^{13, 30, 43, 33}
	Centres for Epidemiological Studies-Depression (CES-D) measures, n=5 ^{3, 20, 23, 30, 43}
	Centres for Epidemiological Studies-Depression (CES-D) measures, $n=5^{3, 20, 23, 30, 43}$ Greene Climacteric Scale, $n=1^4$
	Centres for Epidemiological Studies-Depression (CES-D) measures, n=5 ^{3, 20, 23, 30, 43} Greene Climacteric Scale, n=1 ⁴ Beck Depression Inventory, n=1 ²⁹
Body fat (n=14)	Centres for Epidemiological Studies-Depression (CES-D) measures, $n=5^{3, 20, 23, 30, 43}$ Greene Climacteric Scale, $n=1^4$

	Dual-energy X-ray absorptiometry, n=5 ^{11, 30, 42, 52, 53}
	Sum of skinfolds measures, $n=4^{2, 17, 39, 44}$
Body mass index (n=13)	$n=14^2$, 8, 10, 11, 15, 16, 39, 42-44, 46, 54, 57, 58
	$n=13^{2}, 8, 16, 17, 30, 39, 40, 42, 43, 46, 52, 53, 57$
Body weight (n=12)	n=15 $n=7^{11, 16, 39, 40, 46, 57, 59}$
Waist circumference (n=7)	
	Organization for Research and Treatment of Cancer Quality of Life Questionnaire
FACT: Functional Assessme	nct of Cancer Therapy nctional Assessment of Cancer Therapy Questionnaire for Breast Cancer
	nent of Cancer Therapy: Fatigue
	ent of Chronic Illness Therapy
	ment of Cancer Therapy - General
HRmax: maximum heart rate	
MDASI-T: MD Anderson Sy	
	strument - Breast Cancer Patient Version
RM: repetition maximum	
VO2max: maximal oxygen co	
VO2peak: peak oxygen consu	umption

References

- 1. Anderson RT, Kimmick GG, McCoy TP, et al. A randomized trial of exercise on well-being and function following breast cancer surgery: the RESTORE trial. *J Cancer Surviv* 2012;6(2):172-181.
- 2. Courneya KS. Mackey JR, Bell GJ, Jones LW, Field CJ, Fairey AS. Randomized controlled trial of exercise training in postmenopausal breast cancer survivors: cardiopulmonary and quality of life outcomes. *J Clin Oncol* 2003;21(9):1660-8.
- 3. Danhauer SC, Mihalko SL, Russell GB, et al. Restorative yoga for women with breast cancer: findings from a randomized pilot study. *Psychooncology* 2009;18(4):360-8.
- 4. Hayes SC, Rye S, Disipio T, et al. Exercise for health: a randomized, controlled trial evaluating the impact of a pragmatic, translational exercise intervention on the quality of life, function and treatment-related side effects following breast cancer. *Breast Cancer Res Treat* 2013;137(1):175-186.
- 5. Hornsby WE, Douglas PS, West MJ, et al. Safety and efficacy of aerobic training in operable breast cancer patients receiving neoadjuvant chemotherapy: a phase II randomized trial. *Acta Oncol* 2014;53(1):65-74.
- 6. Milne HM, Wallman KE, Gordon S, Courneya KS. Effects of a combined aerobic and resistance exercise program in breast cancer survivors: a randomized controlled trial. *Breast Cancer Res Treat* 2008; 108(2):279-88.
- 7. Moadel AB, Shah C, Wylie-Rosett J, et al. Randomized controlled trial of yoga among a multiethnic sample of breast cancer patients: effects on quality of life. *J Clin Oncol* 2007;25(28):4387-95.
- 8. Murtezani A, Ibraimi Z, Bakalli A, Krasniqi S, Disha ED, Kurtishi I. The effect of aerobic exercise on quality of life among breast cancer survivors: a randomized controlled trial. *J Cancer Res Ther* 2014;10(3):658-64.
- 9. Naumann F. Munro A, Martin E, et al. An individual-based versus group-based exercise and counselling intervention for improving quality of life in breast cancer survivors. A feasibility and efficacy study. *Psychooncology* 2012;21(10):1136-9.
- 10. Portela ALM, Santaella CL, Gómez CC, Burch A. Feasibility of an Exercise Program for Puerto Rican Women who are Breast Cancer Survivors. *Rehabil Oncol* 2008;26(2):20-31.
- 11. Rogers LQ, Hopkins-Price P, Vicari S, et al. A randomized trial to increase physical activity in breast cancer survivors. *Med Sci Sports Exerc* 2009;41(4):935-46.
- 12. Eakin EG, Lawler SP, Winkler EA, Hayes SC. A randomized trial of a telephonedelivered exercise intervention for non-urban dwelling women newly diagnosed with breast cancer: exercise for health. *Ann Behav Med* 2012;43(2):229-38.
- 13. Short CE, James EL, Girgis A, D'Souza MI, Plotnikoff RC. Main outcomes of the Move More for Life Trial: a randomised controlled trial examining the effects of tailored print and targeted print materials for promoting physical activity among post treatment breast cancer survivors. *Psychooncology* 2015;24(7):771-778.
- 14. Vallance JK, Courneya KS, Plotnikoff RC, Yasui Y, Mackey JR. Randomized Controlled Trial of the Effects of Print Materials and Step Pedometers on Physical Activity and Quality of Life in Breast Cancer Survivors. *J Clin Oncol* 2007;25(17):2352-2359.
- 15. Cornette T, Vincent F, Mandigout S,et al. Effects of home-based exercise training on VO2 in breast cancer patients under adjuvant or neoadjuvant chemotherapy (SAPA): a randomized controlled trial. *Eur J Phys Rehabil Med* 2016;52(2):223-32.

- 16. Fernández-Lao C, Cantarero-Villanueva I, Ariza-Garcia A, et al. Water versus landbased multimodal exercise program effects on body composition in breast cancer survivors: a controlled clinical trial. *Support Care Cancer* 2013;21(2):521-530.
- 17. Herrero F, San Juan AF, Fleck SJ, et al. Combined aerobic and resistance training in breast cancer survivors: A randomized, controlled pilot trial. *Int J Sports Med* 2006;27(7):573-80.
- 18. Kilbreath SL, Refshauge KM, Beith JM, et al. Upper limb progressive resistance training and stretching exercises following surgery for early breast cancer: a randomized controlled trial. *Breast Cancer Res Treat* 2012;133(2):667-676.
- 19. Ligibel JA, Giobbie-Hurder A, Shockro L, et al. Randomized trial of a physical activity intervention in women with metastatic breast cancer. *Cancer* 2016;122(8):1169-1177.
- 20. Schmidt ME, Wiskemann J, Armbrust P, et al. Effects of resistance exercise on fatigue and quality of life in breast cancer patients undergoing adjuvant chemotherapy: A randomized controlled trial. *Int J Cancer* 2015;137(2):471-80.
- 21. Van Waart H, Stuiver MM, van Harten WH, et al. Effect of Low-Intensity Physical Activity and Moderate- to High-Intensity Physical Exercise During Adjuvant Chemotherapy on Physical Fitness, Fatigue, and Chemotherapy Completion Rates: Results of the PACES Randomized Clinical Trial. *J Clin Oncol* 2015;33(17):1918-1927.
- 22. Galiano-Castillo N, Cantarero-Villanueva I, Fernández-Lao C, et al. Telehealth system: A randomized controlled trial evaluating the impact of an internet-based exercise intervention on quality of life, pain, muscle strength, and fatigue in breast cancer survivors. *Cancer* 2016;122(20):3166-3174.
- 23. Chandwani KD, Perkins G, Nagendra HR, et al. Randomized, Controlled Trial of Yoga in Women with Breast Cancer Undergoing Radiotherapy. *J Clin Oncol* 2014;32(10):1058-1065.
- 24. Cormie P, Pumpa K, Galvão DA, Turner E, Spry N, Saunders C, Zissiadis Y, Newton RU. Is it safe and efficacious for women with lymphedema secondary to breast cancer to lift heavy weights during exercise: a randomised controlled trial. *J Cancer Surviv* 2013;7(3):413-424.
- 25. Pinto BM, Papandonatos GD, Goldstein MG. A randomized trial to promote physical activity among breast cancer patients. *Health Psychol* 2013;32(6):616-26.
- 26. De Luca V, Minganti C, Borrione P, et al. Effects of concurrent aerobic and strength training on breast cancer survivors: a pilot study. *Public Health* 2016;136:126-32.
- 27. Headley, J.A., K.K. Ownby, and L.D. John, The effect of seated exercise on fatigue and quality of life in women with advanced breast cancer. *Oncol Nurs Forum* 2004;31(5):977-983.
- 28. Raghavendra RM, Nagarathna R, Nagendra HR, et al. Effects of an integrated yoga programme on chemotherapy-induced nausea and emesis in breast cancer patients. *Eur J Cancer Care* 2007;16(6):462-74.
- 29. Rao RM, Raghuram N, Nagendra HR, et al. Effects of an integrated Yoga Program on Self-reported Depression Scores in Breast Cancer Patients Undergoing Conventional Treatment: A Randomized Controlled Trial. *Indian J Palliat Care* 2015;21(2):174-181.
- 30. Courneya KS, Segal RJ, Mackey JR, et al. Effects of aerobic and resistance exercise in breast cancer patients receiving adjuvant chemotherapy: a multicenter randomized controlled trial. *J Clin Oncol* 2007;25(28):4396-404.

- 31. Wang YJ, Boehmke M, Wu YW, Dickerson SS, Fisher N. Effects of a 6-week walking program on Taiwanese women newly diagnosed with early-stage breast cancer. *Cancer Nurs* 2011;34(2):E1-13.
- 32. Loudon A, Barnett T, Piller N, Immink MA, Williams AD. Yoga management of breast cancer-related lymphoedema: a randomised controlled pilot-trial. *BMC Complement Altern Med* 2013;14(1):214-214.
- 33. Maryam A, Fazlollah A, Eesa M, Ebrahim H, Abbas VF. The effect of designed exercise programme on quality of life in women with breast cancer receiving chemotherapy. *Scand J Caring Sci* 2010;24(2):251-8.
- 34. Drouin JS, Young TJ, Beeler J, et al. Effects of Aerobic Exercise Training on Peak Aerobic Capacity, Fatigue, and Psychological Factors During Radiation for Breast Cancer. *Rehabil Oncol* 2005;23(1):11-17.
- 35. Kim CJ, Kang DH, Smith BA, Landers KA. Cardiopulmonary responses and adherence to exercise in women newly diagnosed with breast cancer undergoing adjuvant therapy. *Cancer Nurs* 2006; 29(2):156-65.
- 36. Musanti R. A study of exercise modality and physical self-esteem in breast cancer survivors. *Med Sci Sports Exerc* 2012;44(2):352-61.
- 37. Husebø AML, Dyrstad SM, Mjaaland I, Søreide JA, Bru E. Effects of Scheduled Exercise on Cancer-Related Fatigue in Women with Early Breast Cancer. *Sci World J* 2014:9.
- 38. Schwartz AL, Winters-Stone K, Gallucci B. Exercise effects on bone mineral density in women with breast cancer receiving adjuvant chemotherapy. *Oncol Nurs Forum* 2007;34(3):627-33.
- 39. Dethlefsen C, Lillelund C, Midtgaard J, et al. Exercise regulates breast cancer cell viability: systemic training adaptations versus acute exercise responses. *Breast Cancer Res Treat* 2016;159(3):469-79.
- 40. Dolan LB, Campbell K, Gelmon K, Neil-Sztramko S, Holmes D, McKenzie DC, Interval versus continuous aerobic exercise training in breast cancer survivors--a pilot RCT. *Support Care Cancer* 2016;24(1):119-27.
- 41. Macvicar MG, Winningham ML, Nickel JL. Effects of Aerobic Interval Training on Cancer Patients' Functional Capacity. *Nursing Research* 1989;38(6): 348-353.
- 42. Hutnick NA, Williams NI, Kraemer WJ, et al. Exercise and lymphocyte activation following chemotherapy for breast cancer. *Med Sci Sports Exerc*. 2005;37(11):1827-35.
- 43. Campbell KL, Kam JW, Neil-Sztramko SE, et al . Effect of aerobic exercise on cancer-associated cognitive impairment: A proof-of-concept RCT. *Psychooncology* 2017:1-8.
- 44. Pinto BM, Frierson GM, Rabin C, Trunzo JJ, Marcus BH. Home-based physical activity intervention for breast cancer patients. *J Clin Oncol* 2005;23(15):3577-87.
- 45. Gokal K. Wallis D, Ahmed S, Boiangiu I, Kancherla K, Munir F. Effects of a selfmanaged home-based walking intervention on psychosocial health outcomes for breast cancer patients receiving chemotherapy: a randomised controlled trial. *Support Care Cancer* 2016;24(3):1139-66.
- 46. Cantarero-Villanueva I, Fernández-Lao C, Caro-Morán E, et al. Aquatic exercise in a chest-high pool for hormone therapy-induced arthralgia in breast cancer survivors: a pragmatic controlled trial. *Clin Rehabil* 2013;27(2):123-32.
- 47. Cantarero-Villanueva I, Fernández-Lao C, Fernández-de-Las-Peñas C, et al. Effectiveness of water physical therapy on pain, pressure pain sensitivity, and myofascial trigger points in breast cancer survivors: a randomized, controlled clinical trial. *Pain Med* 2012;13(11):1509-19.

- 48. Naraphong W. Exercise intervention for fatigue-related symptoms in Thai women with breast cancer: A pilot study., Exercise intervention for fatigue-related symptoms in Thai women with breast cancer: A pilot study. *Nurs Health Sci* 2014;17(1):33-41
- 49. Cantarero-Villanueva I, Fernández-Lao C, Del Moral-Avila R, Fernández-de-Las-Peñas C, Feriche-Fernández-Castanys MB, Arroyo-Morales M. Effectiveness of core stability exercises and recovery myofascial release massage on fatigue in breast cancer survivors: a randomized controlled clinical trial. *Evid Based Complement Alternat Med* 2012:620619.
- 50. Vincent F, Labourey JL, Leobon S, et al. Effects of a home-based walking training program on cardiorespiratory fitness in breast cancer patients receiving adjuvant chemotherapy: a pilot study. *Eur J Phys Rehabil Med* 2013;49(3):319-29.
- 51. Yang CY, Tsai JC, Huang YC, Lin CC. Effects of a home-based walking program on perceived symptom and mood status in postoperative breast cancer women receiving adjuvant chemotherapy. *J Adv Nurs* 2011;67(1):158-68.
- 52. Winters-Stone KM, Dobek J, Nail LM, Bennett JA, Leo MC, Torgrimson-Ojerio B, Luoh SW, Schwartz A. Impact + resistance training improves bone health and body composition in prematurely menopausal breast cancer survivors: a randomized controlled trial. *Osteoporos Int* 2013;24(5):1637-46.
- 53. Winters-Stone KM, Dobek J, Nail L, et al. Strength training stops bone loss and builds muscle in postmenopausal breast cancer survivors: a randomized, controlled trial. *Breast Cancer Res Treat* 2011;127(2):447-56.
- 54. Ahmed RL, Thomas W, Yee D, Schmitz KH. Randomized controlled trial of weight training and lymphedema in breast cancer survivors. *J Clin Oncol* 2006;24(18):2765-2772.
- 55. Banerjee B, Vadiraj HS, Ram A, et al. Effects of an integrated yoga program in modulating psychological stress and radiation-induced genotoxic stress in breast cancer patients undergoing radiotherapy. *Integr Cancer Ther* 2007;6(3):242-50.
- 56. Kim JJ, Shin YA, Suk MH. Effect of a 12-week walking exercise program on body composition and immune cell count in patients with breast cancer who are undergoing chemotherapy. *Journal of Exercise Nutrition & Biochemistry* 2015;**19**(3):255-262.
- 57. Ligibel JA. Campbell N, Partridge A, et al. Impact of a mixed strength and endurance exercise intervention on insulin levels in breast cancer survivors. *J Clin Oncol* 2008;26(6):907-12.
- 58. Rao R, Cruz V, Peng Y, et al. Bootcamp during neoadjuvant chemotherapy for breast cancer: a randomized pilot trial. *Breast Cancer* 2012;6:39-46.
- 59. Guinan EM, Connolly EM, Hussey J, Exercise training in breast cancer survivors: a review of trials examining anthropometric and obesity-related biomarkers of breast cancer risk. *Phys Ther Rev* 2013;18(2):79-89.

Supplementary Content 8. Meta-analyses results of depression with subgroup analyses for exercise mode, intervention supervision, timing and duration and sensitivity analyses (positive SMD values favour exercise).

Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	4	83 vs. 128	81%	0.53 (0.24, 0.82)	<0.01	⊢
Resistance exercise	2	90 vs. 134	0%	0.04 (-0.23, 0.31)	0.79	⊢
Combined exercise	2	44 vs. 41	9%	0.62 (0.18, 1.06)	<0.01	⊢
Other exercise	6	203 vs. 205	93%	1.16 (0.94, 1.38)	<0.01	⊢∎
Supervision [1]						
Supervised interventions	9	308 vs. 398	93%	0.50 (0.34, 0.66)	<0.01	H
Unsupervised interventions	5	112 vs. 110	0%	1.18 (0.89, 1.47)	<0.01	⊢
Intervention timing						
During treatment	10	320 vs. 410	93%	0.67 (0.51, 0.83)	<0.01	⊢∎→
Post-treatment	4	100 vs. 98	0%	0.62 (0.33, 0.90)	<0.01	⊢_∎(
Mixed	0	-	-	-	-	
Duration						
<12 week interventions	8	250 vs. 268	93%	0.84 (0.65, 1.03)	<0.01	⊢∎⊣
>12 week interventions	6	170 vs. 240	77%	0.44 (0.23, 0.65)	<0.01	⊢∎
Sensitivity analyses [2]						
High quality studies only	42	42	92%	0.46 (0.30, 0.63)	<0.01	⊢∎⊣
Stage II+ only	10	10	21%	1.04 (0.74, 1.34)	<0.01	⊢
Overall	14	420 vs. 508	90%	0.66 (0.52, 0.80)	<0.01	•

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

[2] Sensitivity analyses were performed only on i) high quality studies, and ii) studies with 100% of samples with stage II+ disease.

Supplementary Content 9. Meta-analyses results of upper-body strength with subgroup analyses for exercise mode, intervention supervision, timing and duration and sensitivity analyses (positive SMD values favour exercise).

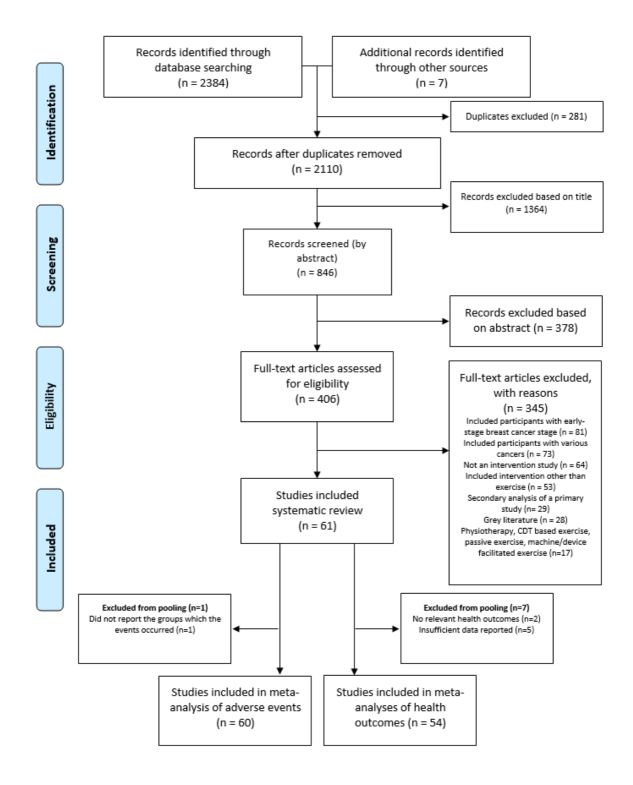
Subgroup	Studies (n=)	Participants (n=)	l² (%)	SMD (95% CI)	P-value	
Exercise mode		Intervention vs. Control				
Aerobic exercise	4	131 vs. 137	63%	0.29 (0.05, 0.54)	0.02	⊢_
Resistance exercise	8	280 vs. 282	50%	0.68 (0.50, 0.85)	<0.01	⊢∎-
Combined exercise	10	395 vs. 375	0%	0.30 (0.16, 0.44)	<0.01	H
Other exercise	0	-	-	-	-	
Supervision [1]						
Supervised interventions	11	406 vs. 410	66%	0.49 (0.35, 0.63)	<0.01	H
Unsupervised interventions	11	400 vs. 384	3%	0.36 (0.22, 0.50)	<0.01	H
Intervention timing						
During treatment	8	490 vs. 484	65%	0.37 (0.25, 0.50)	<0.01	H
Post-treatment	13	235 vs. 231	34%	0.57 (0.38, 0.76)	<0.01	⊢ ∎1
Mixed	1	81 vs. 79	-	0.33 (0.02, 0.64)	0.04	⊢
Duration						
<12 week interventions	8	198 vs. 197	29%	0.49 (0.29, 0.69)	<0.01	⊢ ∎1
>12 week interventions	14	608 vs. 597	58%	0.40 (0.29, 0.52)	<0.01	H
Sensitivity analyses [2]						
High quality studies only	17	727 vs. 722	60%	0.43 (0.32, 0.53)	<0.01	H
Stage II+ only	2	25 vs. 18	0%	0.27 (-0.34, 0.88)	0.39	
Overall	22	806 vs. 794	49%	0.43 (0.33, 0.53)	<0.01	•

< Favours Control Favours Exercise >

[1] Supervised intervention were classed as interventions where \geq 50% of prescribed exercise involved face-to-face supervision and unsupervised interventions involved <50% of prescribed exercise involving face-to-face supervision.

[2] Sensitivity analyses were performed only on i) high quality studies, and ii) studies with 100% of samples with stage II+ disease.

Supplementary Content 1: Systematic review flow diagram.



Supplementary Content 2: Ratings of all studies included in systematic review using the PEDro scale (n=61).

	PEDro Scale item number											
	1	2	3	4	5	6	7	8	9	10	11	Total
												score
												(Quality)
Ahmed 2006	1	1	0	1	0	0	1	1	0	1	1	6 (High)
Anderson 2012	1	1	1	1	0	0	1	0	0	1	1	6 (High)
Banerjee 2007	1	1	1	1	0	0	0	1	0	1	1	6 (High)
Campbell 2017	1	1	0	1	0	0	1	1	1	1	1	7 (High)
Cantarero-Villanueva 2012a	1	0	0	1	0	0	1	0	0	1	1	4 (Low)
Cantarero-Villanueva 2012b	1	1	1	1	0	0	1	1	1		1	8 (High)
Cantarero-Villanueva 2012c	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Cantarero-Villanueva 2013	1	1	1	1	0	0	1	1	0	1	1	7 (High)
Chandwani 2014	1	1	0	1	0	0	0	0	0	1	1	4 (Low)
Cormie 2013	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Cornette 2016	1	1	0	1	0	0	0	0	1	1	1	5 (Low)
Courneya 2003	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Courneya 2007	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Danhauer 2009	1	1	0	1	0	0	0	0	1	1	1	5 (Low)
De Luca 2016	1	1	0	0	0	0	0	1	0	1	1	4 (Low)
Dethlefsen 2016	1	1	0	1	0	0	0	0	0	1	1	4 (Low)
Dolan 2016	1	1	0	1	0	0	0	1	0	1	1	5 (Low)
Drouin 2005	1	1	0	1	0	0	0	0	0	0	1	3 (Low)
Eakin 2012	1	1	0	1	0	0	0	1	1	1	1	6 (High)
Fernández-Lao 2013	1	0	0	1	0	0	1	0	1	1	1	5 (Low)
Galiano-Castillo 2017	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Gokal 2015	1	1	0	1	0	0	0	1	1	1	1	6 (High)
Guinan 2013	1	1	0	1	0	0	1	0	1	1	1	6 (High)
Hatchett 2013	1	1	0	1	0	0	0	1	0	1	1	5 (Low)
Hayes 2012	1	1	0	1	0	0	1	1	1	1	1	7 (High)
Headley 2004	1	1	0	1	0	0	0	0	0	1	1	4 (Low)
Herrero 2005	1	1	1	1	0	0	0	0	0	1	1	5 (Low)
Hornsby 2014	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Husebø 2014	1	1	1	1	0	0	0	0	0	1	1	5 (Low)
Hutnick 2005	1	0	0	1	0	0	0	0	0	0	1	2 (Low)
Kilbreath 2012	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Kim 2006	1	1	0	1	0	0	0	0	0	1	1	4 (Low)
Ligibel 2008	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Ligibel 2016	1	1	0	1	0	0	1	0	0	1	1	5 (Low)
Loudon 2014	1	1	1	1	0	0	1	0	0	1	1	6 (High)
Macvicar 1989	0	1	0	1	0	0	0	0	0	1	0	3 (Low)
Maryam 2010	1	0	0	1	0	0	0	0	0	1	1	3 (Low)
Milne 2008	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Moadel 2007	1	1	0	1	0	0	0	0	1	1	1	5 (Low)
Mohan Rao 2015	1	1	1	1	0	0	0	0	1	1	1	6 (High)
Mulero Portela 2008	1	1	0	1	0	0	1	0	0	1	1	5 (Low)
Murtezani 2014	1	1	1	1	0	0	1	1	0	1	1	7 (High)

Musanti 2012	1	1	1	1	0	0	1	0	1	1	1	7 (High)
Naraphong 2015	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Naumann 2012	1	0	0	1	0	0	0	1	1	1	1	5 (Low)
Pinto 2005	1	1	0	1	0	0	0	1	1	1	1	6 (High)
Pinto 2013	1	1	1	1	0	1	1	0	1	1	1	8 (High)
Pinto 2015	1	1	1	1	0	0	1	1	0	1	1	7 (High)
Raghavendra 2007	1	1	1	1	0	0	0	0	0	1	1	5 (Low)
Rao 2012	1	1	1	1	0	0	1	1	1	1	1	8 (High)
Rogers 2009	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Schmidt 2015	1	1	1	1	0	0	0	1	1	1	1	7 (High)
Schwartz 2007	1	1	0	1	0	0	0	1	1	1	1	6 (High)
Short 2015	1	1	1	0	0	0	0	1	1	1	1	6 (High)
Vallance 2007	1	1	1	1	0	0	0	1	1		1)	7 (High)
Vallance 2015	1	1	1	1	0	0	0	0	1	1	_1	6 (High)
Van Waart 2016	1	1	0	1	0	0	0	1	1	1	1	6 (High)
Wang 2011	1	1	0	1	0	0	0	1	0	1	1	5 (Low)
Winters Stone 2011	1	1	0	1	0	0	1	0	1	1	1	6 (High)
Winters-Stone 2013	1	1	1	1	0	0	1	0	1	1	1	7 (High)
Yang 2010	1	1	0	1	0	0	0	1	0	1	1	5 (Low)
Pedro scale items: 1 Eligibilit	v crit	eria	2 SI	ihiec	ts rai	ndom	lv al	locat	ed· 4	Gro	nine e	similar at

Pedro scale items: 1. Eligibility criteria; 2. Subjects randomly allocated; 4. Groups similar at baseline; 5. Subject blinding; 6. Therapist blinding; 7. Assessor blinding; 8. Outcome obtained from >85% of subjects; 9. Intention to treat; 10. Results of between-group comparisons; 11. Point and variability measures