BMJ Open Is exercise-based cardiac rehabilitation effective? A systematic review and meta-analysis to re-examine the evidence

Richard Powell,^{1,2} Gordon McGregor,^{1,3} Stuart Ennis,^{1,4} Peter K Kimani,⁵ Martin Underwood²

ABSTRACT

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¹Department of Cardiac Rehabilitation. Centre for Exercise & Health, University Hospitals, Coventry, UK ²Warwick Clinical Trials Unit, Warwick Medical School. University of Warwick, Coventry, UK ³Faculty of Health and Life Sciences, Coventry University, Coventry, UK ⁴Cardiff Centre for Exercise & Health, Cardiff Metropolitan University, Cardiff, UK ⁵Statistics and Epidemiology Unit, Warwick Medical School, University of Warwick, Coventry, UK

Correspondence to Richard Powell; richard.powell@uhcw.nhs.uk **Objectives** To determine the contemporary effectiveness of exercise-based cardiac rehabilitation (CR) in terms of all-cause mortality, cardiovascular mortality and hospital admissions.

Data sources Studies included in or meeting the entry criteria for the 2016 Cochrane review of exercise-based CR in patients with coronary artery disease.

Study eligibility criteria Randomised controlled trials (RCTs) of exercise-based CR versus a no-exercise control whose participants were recruited after the year 2000. Study appraisal and synthesis methods Two separate reviewers independently screened the characteristics of studies. One reviewer quality appraised any new studies and assessed their risk of bias using the Cochrane Collaboration's recommended risk of bias tool. Data were

reported as the risk difference (95% Cl). **Results** We included 22 studies with 4834 participants

(mean age 59.5 years, 78.4% male). We found no differences in outcomes between exercise-based CR and a no-exercise control at their longest follow-up period for: all-cause mortality (19 studies; n=4194; risk difference 0.00, 95% Cl -0.02 to 0.01, P=0.38) or cardiovascular mortality (9 studies; n=1182; risk difference -0.01, 95% Cl -0.02 to 0.01, P=0.25). We found a small reduction in hospital admissions of borderline statistical significance (11 studies; n=1768; risk difference -0.05, 95% Cl -0.10to -0.00, P=0.05).

Conclusions and implications of key findings Our analysis indicates conclusively that the current approach to exercise-based CR has no effect on all-cause mortality or cardiovascular mortality, when compared with a no-exercise control. There may be a small reduction in hospital admissions following exercise-based CR that is unlikely to be clinically important.

PROSPERO registration number CRD42017073616.

BACKGROUND

Cardiovascular disease is the world's biggest killer, accounting for 15 million deaths in $2015.^{1}$

Secondary prevention of coronary artery disease through exercise-based CR in those who have a diagnosis of coronary artery

Strengths and limitations of this study

- To our knowledge, this is the first systematic review of exercise-based cardiac rehabilitation (CR) that has pooled data relevant to the current medical management of patients diagnosed with coronary artery disease.
- For analysis, we present the data as the risk difference (95% CI), which ensures all studies reporting data on the outcomes of interest were included.
- This systematic review pools data from studies that deliver an intervention recognised as best practice in exercise-based CR, where multiple approaches, including educational/psychosocial components, as well as the exercise component were used.
- We have not done a de novo quality assessment of 21/22 studies included in this review and instead rely on a previous Cochrane assessment.
- We did not include health-related quality of life as an outcome measure as this is unsuitable for metaanalysis.

disease has the potential to reduce mortality, reduce hospital admissions and increase quality of life. Guidelines internationally endorse the use of exercise-based cardiac rehabilitation (CR) programmes.^{2–5}

Typically, exercise-based CR aims to achieve 20–60 min of moderate intensity continuous exercise, 3–5 times a week, with muscular strength and endurance exercises prescribed in conjunction.⁶ Additionally, most programmes include supplementary education (coronary risk factors and cardiac misconceptions), advice on diet and access to psychological support.²⁴⁷⁸ Typically, exercise-based CR is delivered in a supervised centre-based setting, although home-based programmes are used.⁹

A 2016 Cochrane review (63 studies, n=14486 participants) found benefits of exercise-based CR for patients with

Table 1 Risk of bias assessment for additional study Senteularia at a ⁽²⁾							
Santaularia et al ²⁹							
Bias	Authors' judgement	Support for judgement					
Random sequence generation (selection bias)	Low risk	A randomisation list in blocks of 10 was created by a computer random number generator. The randomisation list and the allocation of patients to each group were independently controlled by the Clinical Trials Unit.					
Allocation concealment (selection bias)	Low risk	A randomisation list in blocks of 10 was created by a computer random number generator. The randomisation list and the allocation of patients to each group were independently controlled by the Clinical Trials Unit.					
Blinding of outcome assessment (detection bias): all outcomes	Low risk	An independent committee that was blind to the patients' treatment group assessed the main outcomes. This committee comprised a cardiologist, a rehabilitation cardiologist and a health information manager, all from different centres.					
Incomplete outcome data (attrition bias): all outcomes	Low risk	There was no loss to follow-up.					
Selective reporting (reporting bias)	Low risk	All outcomes described in the methods were reported in the results. Results regarding quality of life are presented in supplementary data but were not required for the current review.					
Groups balanced at baseline	Low risk	No significant differences between groups were observed, with the exception of gender: 23% of the control group were women compared with 7% in the intervention group (P =0.049).					
Intention-to-treat analysis conducted	High risk	No analysis was conducted.					
Groups received same treatment (apart from the intervention)	e Low risk	Patients assigned to the control group received standard care given at the hospital. In addition to standard care, patients randomised to the intervention group.					

coronary artery disease. Both cardiovascular mortality (27 studies, risk ratio (RR) 0.74, 95% CI 0.64 to 0.86) and hospital readmissions were reduced (15 studies, RR 0.82, 95% CI 0.70 to 0.96), when compared with a no-exercise control. However, in contrast to previous systematic reviews and meta-analyses, there was no significant reduction in risk of reinfarction (36 studies, RR 0.90, 95% CI 0.79 to 1.04) or all-cause mortality (47 studies, RR 0.96, 95% CI 0.88 to 1.04).¹⁰

Over recent decades, the medical management of coronary artery disease has been transformed. The introduction of primary percutaneous coronary intervention has reduced short-term major adverse cardiac events and increased long-term survival.¹¹⁻¹⁴ Simultaneously, there have also been widespread advances in secondary preventative medical therapy. This includes the introduction of aspirin and betablockers in the 1980s,¹⁵¹⁶ lipid-lowering statins and ACE inhibitors in the 1990s^{17 18} and, more recently, the introduction of clopidogrel, a secondary antiplatelet, in 2007.^{19 20} Age-adjusted mortality has decreased substantially in this population.²¹ Systematic reviews and meta-analyses that include data from older studies may not correctly assess the potential effect of exercise-based CR. We hypothesise that previous reviews have overestimated the benefit of exercise-based CR.

OBJECTIVES

To determine the contemporary effectiveness of exercise-based CR on all-cause mortality, cardiovascular mortality and hospital readmissions in patients with coronary artery disease.

METHODS

We conducted and reported this meta-analysis in accordance with Preferred Reporting Items for Systematic Reviews and Meta-Analyses.²²

Search strategy

To identify relevant studies, we started with the latest Cochrane review of exercise-based CR in patients with coronary artery disease.¹⁰ Studies identified as 'awaiting assessment' or 'on-going' in this review were revisited to establish whether publication had been reached. To identify any new studies published since the completion of the Cochrane review, an updated search was run on 28 February 2017. This search used the same search strategies as the latest Cochrane review.¹⁰ We searched Cochrane Central Register of Controlled Trials (online supplementary appendix 1), MEDLINE (Ovid), Embase (Ovid) and CINAHL (EBSCO) databases. This approach allowed us to efficiently identify all relevant studies. Where appropriate, we contacted original authors for clarification of any new included studies.

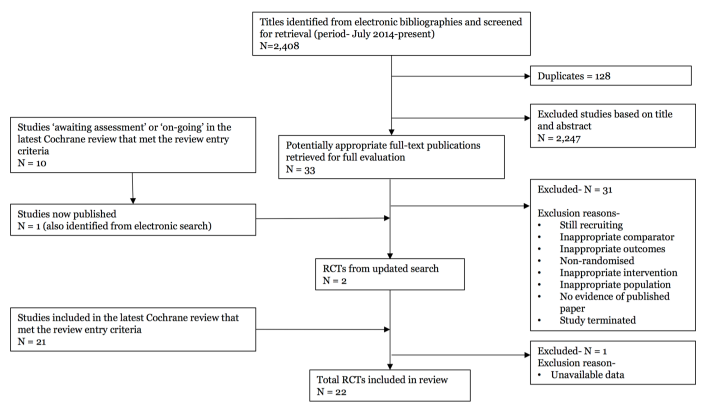


Figure 1 Summary of study selection process. RCTs, randomised controlled trials.

Two separate reviewers (RP and GM) independently screened the characteristics of studies in the latest Cochrane review, studies identified as 'awaiting assessment' or 'on-going' and studies identified in the updated search. Full-text publications were retrieved to allow for further examination and to verify study inclusion. Any discrepancies were resolved by a third reviewer (MU).

Criteria for considering studies

In 1996, The Task Force on the Management of Acute Myocardial Infarction of the European Society of Cardiology first recommended early (within 2 hours) primary percutaneous interventions in preference to thrombolytic therapy for acute myocardial infarction.²³ Two years later, guidelines set by the Joint British recommendations on prevention of Coronary Heart Disease in Clinical Practice were published outlining the recommendations for best practice for secondary prevention medical therapies.²⁴ Although there have been some changes, notably the introduction of a second antiplatelet agent in the early 2000s,^{19 20} the approach to secondary prevention medical therapies has not changed since then. Allowing time for implementation of these guidelines and recommendations, we identified and included studies whose participants were recruited after the year 2000 to represent a contemporary population engaging in exercise-based CR.

Where there was no indication of recruitment period, the diagnosis and the secondary preventative medical therapy received by participants included in the trial determined the inclusion or exclusion of the study in the analysis.

Types of studies

We included randomised controlled trials of exercise-based CR compared with a no-exercise control with a minimum follow-up period of 6 months. Data reported at the longest follow-up period were included in the analysis.

Types of participants

We used the same entry criterion as previous Cochrane reviews:

- people who have had a myocardial infarction or who had undergone revascularisation (coronary artery bypass grafting or percutaneous coronary intervention) or who have angina pectoris or coronary artery disease defined by angiography.
- ► on optimal secondary preventative medical therapy defined by the Joint British recommendations on prevention of Coronary Heart Disease in Clinical Practice.²⁴
- recruited to hospital-based, community-based or home-based CR programmes.

Types of intervention(s)

Randomised controlled trials consisted of supervised or non-supervised exercise-based CR. The intervention was exercise alone or exercise as part of a comprehensive CR programme (consisting of educational/psychosocial components). 'No exercise control' consisted of standard medical care, including optimal secondary preventative medical therapy, education and advice about diet and exercise, psychosocial support but with no formal exercise intervention.

Open Access

Types of outcome measures

We extracted data on: all-cause mortality, cardiovascular mortality and hospital readmissions. We did not include health-related quality of life as the authors of the 2016 Cochrane review found this unsuitable for meta-analysis.

Data collection, statistical analysis and quality assessment

We pooled data using Review Manager V.5.3.²⁵ Previous Cochrane reviews have presented the data as individual and pooled risk ratio (95% CI). Using risk ratios automatically removed studies with no events in either study arm from the analysis. Nine studies (n=936 participants) reporting on all-cause mortality, cardiovascular mortality or hospital readmissions were excluded from one or more meta-analyses in the 2016 Cochrane review for this reason. We therefore present the data as the risk difference (95% CI), which ensures all studies reporting data on the outcomes of interest were included.

We applied a random-effects model to all analyses given the clinical heterogeneity of individual studies. Heterogeneity of included studies were tested statistically using the χ^2 test of heterogeneity and I² statistic.²⁶

We did not repeat quality assurance checks already completed by the authors of the Cochrane review. For separate study risk of bias breakdown for these studies, we refer the reader to the existing *characteristics of studies*.¹⁰ For studies identified as 'awaiting assessment' or 'on-going' in the latest Cochrane review or in the updated search, we quality appraised these studies and assessed their risk of bias using the Cochrane Collaboration's recommended risk of bias tool.²⁷

Assessment of risk of bias in additional included study

One reviewer (RP) assessed the risk of bias in any additional included studies (table 1). Assessment of three further quality domains as outlined in the latest Cochrane review was also conducted (groups balanced at baseline, intention-to-treat analysis and groups received comparable treatment (except exercise)). A breakdown of the criteria used for assessing these three domains can be found in the latest Cochrane review. Risk of bias assessments were checked by a second reviewer (GM) and any discrepancies were resolved by a third reviewer (MU).

Patient involvement

No patients were involved in setting the objectives or outcome measures of this review, nor were they involved in the design or implementation. No patients were involved in the analysis or interpretation of the results, nor the writing of any drafts. There are no plans to disseminate the results of the review to participants included in the studies of the review or any relevant patient networks.

RESULTS

Studies retrieved

Of the 63 studies included in the Cochrane review, 21 studies met our entry criteria. We identified two additional relevant papers not included in the 2016 Cochrane review.²⁸ ²⁹ One was excluded because data for our specific research question were not available in a useable format.²⁸ In total, 22 studies (n=4834 participants) contributed to the analysis (figure 1). For the study identified from the updated search,²⁹ there was a low risk of bias in all eight domains, apart from the intention-to-treat analysis, where there was no evidence of this analysis being conducted (table 1).

Three studies (3/22; 14%) reported on all three outcomes of interest, 11 studies (11/22; 50%) reported on two outcomes of interest and 8 studies (8/22; 36%) reported on one outcome of interest.

Two studies for all-cause mortality^{30 31} and one study for cardiovascular mortality³⁰ reported data at varying follow-up periods (6–12 months; >12–36 months; >3 years). Data from these studies were taken at their longest follow-up period. Mean maximum follow-up period was 24.7 months. Maximum follow-up period ranged from 24 weeks to 10 years (table 2).

Sample size, gender, age and study origin

Of our 22 studies, 10 studies were in Europe^{29–38} and 12 from outside of Europe.^{39–50} We included a total of 4834 participants (3788 (78.4%) males). Four studies included males only,^{30 34 45 47} and one study included women only.⁵¹ Participants mean age was 59.5 years. The mean age for individual studies ranged from 47.5 to 76.9 years (table 2).

Incomplete outcome data

The majority of trials (18/22; 82%) reported complete follow-up data, regardless of participants who were lost to follow-up or who dropped out. In four studies, outcome data were incomplete for 75 (75/4,834; 1.6%) participants with no description of withdrawal or dropout.^{41 47 48 50}

Participant diagnosis of coronary artery disease and treatment received

The diagnosis of participants recruited to the studies was described in the majority of studies (21/22; 95%). Thirteen studies enrolled participants with mixed diagnoses, including angina pectoralis or coronary artery disease defined by angiography, myocardial infarction, percutaneous coronary interventions or coronary artery bypass grafts.³² ³⁶⁻⁴² ⁴⁴ ⁴⁶⁻⁴⁸ ⁵⁰ Six studies enrolled participants following acute myocardial infarction only,²⁹³¹ ³³³⁴ ⁴³⁴⁹ and two studies enrolled participants diagnosed with angina pectoralis (unstable and stable angina) only.³⁰ ³⁵ It was unclear from one study whether participants following myocardial infarction were included and instead the population was defined as 'patients after coronary artery bypass graft surgery'⁴⁵ (table 2).

Six studies recruited participants following percutaneous coronary intervention only^{30 32 33 35 41 46} and one study recruited participants following coronary artery bypass grafting only.⁴⁵ Twelve studies included participants who had received thrombolysis, percutaneous coronary intervention, coronary artery bypass grafting and/

Reference, country	z	Mean age (years)	Male participants (%)	e, Maximum Berruitment follow-up N (years) (%) period (years) period Pa	Maximum follow-up period	Patient diagnosis	Medication
Aronov <i>et al</i> , ³⁹ Russia	392	61.4	73.5	None specified	1 year	AMI, stable angina, unstable angina or myocardial revascularisation.	Standard medical therapy: β-blocker, acetylsalicyclic acid or other antithrombotic drug, nitrate, ACE inhibitor. Some patients on lipid-lowering drugs.
Belardinelli <i>et al</i> , ³² Italy	118	61	100	None specified	33 months	CAD including AMI. Successful PCI in one or two native epicardial coronary arteries only.	According to international accepted protocols: aspirin, ticlopidine, calcium antagonists and nitrates.
Briffa <i>et al,</i> ⁴⁰ Australia	113	47.5	89.5	None specified	1 year	Uncomplicated AMI or recovery from unstable angina. PCI, CABG, thrombolytic therapy.	Aspirin, antiarrhytmic agent, β -blocker, ACE inhibitor, calcium antagonist, long-acting nitrate and diuretic.
Giallauria <i>et al</i> , ³³ Italy	61	58.5	78.5	None specified	6 months	AMI and undergone primary or rescue PCI only.	Aspirin, $\beta\text{-blocker}$, ACE inhibitor, ARB and statin.
Hambrecht <i>et al</i> , ³⁰ Germany	101	56	87.3	1997–2001	1 year	Stable CAD defined by angina pectoralis and amenable to PCI. AMI patients excluded.	β-receptor antagonists, β-HMG-CoA reductase inhibitors, ACE inhibitor and acetylsalicyclic acid.
Higgins <i>et al,</i> ⁴¹ Australia	105	60.8	81.3	19951997	51 weeks	Post-PCI patients only. No AMI 1-month preprocedure.	Reference to medical therapy, only breakdown for lipid-lowering medication.
Houle <i>et al</i> , ⁴² Canada	65	51.5	100	2007–2008	12 months	Patients hospitalised for an ACS (unstable angina, non-ST-elevation or ST elevation MI). PCI, CABG or no revascularisation procedure.	Reference to medication in usual care group but no breakdown.
Kovoor e <i>t al,⁴³</i> Australia	142	51.5	100	None specified	6 months	AMI only. Thrombolytic therapy, one patient in the exercise treatment group had primary angioplasty.	Aspirin, β-blocker, ACE inhibitor, calcium channel blockers, nitrates and cholesterol-lowering agents.
Maddison e <i>t al</i> , ⁴⁴ New Zealand	171	59	20	2010-2012	24 weeks	Diagnosis of IHD (angina, MI, revascularisation, including angioplasty, stent or CABG).	No description.
Maroto <i>et al</i> , ³⁴ Spain	180	76.9	57.5	(None specified) 2-year enrolment period	10 years	AMI only.	Medication regimens employed in secondary prevention at discharge were clearly insufficient by standard criteria but currently meet Spanish and European guidelines.
Munk <i>et al</i> , ³⁵ Norway	40	56.4	84.8	None specified	6 months	Stable angina and unstable angina, post-PCI only. AMI patients excluded.	Aspirin, β -blocker, ACE inhibitor, ARB, statin and acetylsalicyclic acid.
Mutwalli <i>et al,</i> ⁴⁵ Saudi Arabia	49	69.7	100	2008–2010	6 months	Undergone CABG surgery. Unknown whether AMI patients included.	Undergone CABG surgery. Unknown Participants received advice that focused whether AMI patients included.

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Table 2 Continued							
Reference, country	z	Mean age (years)	Male participants (%)	Recruitment period (years)	Maximum follow-up period	Patient diagnosis	Medication
Oerkild e <i>t al</i> , ³⁶ Denmark	40	63.5	o	2007-2008	12 months (mortality data after 5.5 years)	Recent coronary event defined as AMI, PCI, CABG or without invasive procedure.	β-blocker, antithrombotics, calcium antagonists, lipid-lowering agents and diuretics.
Reid <i>et al</i> , ⁴⁶ Canada 223		54.5	87.3	2004-2007	12 months	ACS including AMI, underwent successful PCI only.	Reference to a 'descriptive summary in supplemental table', no access.
Santaularia <i>et al,</i> ²⁹ Spain	85	59.6	84.7	None specified	12 months	AMI only, no evidence of revascularisation procedure.	Reference to cardiac medication but no breakdown
Seki <i>et al</i> , ⁴⁷ Japan	39	57.8	83.8	None specified	NR	AMI, PCI or CABG.	Reference to 'lipid-lowering drugs and other medications', no breakdown.
Toobert <i>et al</i> , ⁴⁸ USA 25		64.5	0	None specified	24 months	CAD defined as atherosclerosis, AMI, PCI or CABG.	Antihypertensive and hypolipidaemic medications.
Vestfold Heartcare Study Group, ³⁷ Norway	197 (64	75.8	None specified	2 years	AMI, unstable angina pectoris or after PCI or CABG.	Aspirin, β -blocker, statin, ACE inhibitor, calcium antagonist and warfarin.
Wang <i>et al</i> , ⁴⁹ China 160		67	63.5	2005-2007	6 months	AMI only.	Antiplatelet, nitrate, β -blocker, ACE inhibitor, calcium antagonist and statin.
West <i>et al</i> , ³¹ UK	1813	51.9	93.9	1997–2000	7–9 years	AMI only.	Aspirin, β-blocker, ACE inhibitor, diuretic, long-acting nitrate/calcium channel blocker, statin and GTN.
Yu <i>et al</i> , ⁵⁰ China	269	56	83.9	None specified	2 years	Recent AMI, after elective PCI or thrombolytic therapy.	Antiplatelet, β-blocker, calcium channel blocker, nitrate, statin, ACE inhibitor and diuretic.
Zwisler <i>et al</i> , ³⁸ Denmark	446	55.5	72.1	2000-2003	1 year	AMI, angina pectoris or after PCI or CABG.	Antithrombotics, lipid-lowering drugs, β-blocker, calcium antagonists, ACE inhibitor, diuretic and long-acting nitrates.
ACS, acute coronary sy	ndrome;	AMI, acut(e myocardial infar	ction; ARB, angioten	sin receptor blocker	ACS, acute coronary syndrome; AMI, acute myocardial infarction; ARB, angiotensin receptor blockers; CABG, coronary artery bypass graft; CAD, coronary artery disease; GTN, glyceryl	, coronary artery disease; GTN, glyceryl

trinitrate; IHD, ischaemic heart disease; MI, myocardial infarction; PCI, percutaneous coronary intervention.

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or no revascularisation procedure.^{31 36–40 42–44 47 48 50} Three studies did not provide any breakdown of coronary intervention or surgical procedure received by participants prior to enrolment^{29 34 49} (table 2).

Medication

A full description and breakdown of the medication received by the participants, comparable with optimal secondary prevention medical therapy defined by the Joint British recommendations on prevention of Coronary Heart Disease in Clinical Practice set in 1998,²⁴ was provided by 13/22 studies (59%).^{30–33 35–40 43 49 50} References to coexisting medical therapies were made in 7/22 (32%), but no breakdowns were provided.^{29 34 41 42 45–47} One study referred to the prescription of antihypertensive and hypolipidaemic medications.⁴⁸ One study failed to provide any description or breakdown of coexisting medical therapies⁴⁴ (table 2).

Clearly defined recruitment period

Seven studies (7/22; 32%) were explicit that they recruited participants after the year 2000.³⁶ ³⁸ ⁴² ⁴⁴⁻⁴⁶ ⁴⁹ In three studies, participants were recruited either just before or during the year 2000.^{30 31 41} Due to participant diagnosis, treatment received and coexisting medical therapies, it was agreed by all reviewers to include these studies.

The remaining 12 studies failed to provide a recruitment period. Following further examination of the full papers, due to adequate description of patient diagnosis, treatment received and coexisting medical therapies, it was agreed by all reviewers to include these studies (table 2).

Content of the interventions

The content of the interventions tested was heterogeneous with multiple approaches being adopted. Sixteen studies (16/22; 73%) compared exercise in combination with other therapies (education and psychosocial management), while six studies compared exercise as a stand-alone intervention, against a no-exercise control. The exercise component alone varied considerably with respect to setting, training modality, duration, session length, frequency and intensity (table 3).

Overall effects of interventions

All-cause mortality

Nineteen studies (n=4194 participants) reported all-cause mortality (figure 2). There was no difference between groups at their longest follow-up (risk difference=0.00, 95% CI -0.02 to 0.01, P=0.38). There was no evidence of statistical heterogeneity across trials (P value=0.91, $I^2=0\%$).

Cardiovascular mortality

Nine studies (n=1182 participants) reported cardiovascular mortality (figure 3). There was no difference between groups at their longest follow-up (risk difference=-0.01, 95% CI -0.02 to 0.01, P=0.25). There was no evidence of statistical heterogeneity across trials (P value=0.44, $I^2=0\%$).

Hospital admissions

Eleven studies (n=1768 participants) reported on proportion with one or more hospital admissions (figure 4). There was a reduction of borderline statistical significance (risk difference=-0.05, 95% CI -0.10 to -0.00, P=0.05). There was evidence of statistical heterogeneity across trials (P value=0.002, I²=64%).

DISCUSSION

The effectiveness of exercise-based CR in patients with coronary artery disease has been determined by Cochrane systematic reviews and meta-analyses, providing clinicians and academics with the highest level of evidence over the last 17 years.^{10 52 53} The latest Cochrane review, conducted in 2016, found benefits of exercise-based CR in terms of reduced cardiovascular mortality and hospital admissions, but unlike previous Cochrane reviews, found no effect on all-cause mortality.¹⁰ We identified that data from studies included in this review dated back as far as 1975.⁵⁴ By including such historical data, this Cochrane review may not be correctly assessing the potential effect of contemporary exercise-based CR.

The current review aimed to assess the effect of exercise-based CR in the era of improved reperfusion strategies and simultaneous advances in pharmacological management, by only including studies whose participants were recruited after the year 2000. The majority of interventions tested in the 22 included trials (table 3) delivered an intervention recognised as best practice in exercise-based CR, where multiple approaches, including educational/psychosocial components, as well as the exercise component were used.^{2 3 8} The interventions were tested against a no exercise control consisting of educational and psychosocial components alone (table 3).

The current analyses demonstrated no improvement in all-cause mortality from participation in exercise-based CR: the risk difference was 0.00 (95% CI -0.02 to 0.01). The largest trial included in our analysis, the UK-based Rehabilitation after myocardial infarction trial (RAMIT) trial, sought to show a 20% reduction in relative risk based on an 11% mortality, that is, a 2.2% risk difference.²⁴ The limits of the 95% CI for the effect size in our analysis do not include the RAMIT trial's prespecified clinically important difference. We therefore conclude that it is extremely unlikely that there is a worthwhile benefit from exercise-based CR on all-cause mortality. Furthermore, it is unlikely that future trials of similar interventions and populations will change this conclusion. This is supported by a recent meta-analysis that included participants with other forms of atherosclerotic cardiovascular disease, that is, peripheral artery disease, ischaemic cerebrovascular accidents, diabetes and hypertension. They too found a zero effect on all-cause mortality (relative risk 1.00, 95% CI 0.88 to 1.14).⁵⁵ With the mean follow-up

Table 3 Overview	Overview of exercise interventions					
	Exercise intervention					
Reference, country Exercise	y Exercise	Modality	Study duration	Session duration/frequency/ intensity	Additional	Control (comparator)
Aronov <i>et al,</i> ³⁹ Russia	Moderate intensity physical training (unknown setting).	Cycle ergometer.	12 months	45–60 min/three sessions per week/50%–60% of the performed capacity by bicycle ergometry.	None specified.	Standard medical therapy.
Belardinelli <i>et al</i> , ³² Italy	Moderate intensity exercise (supervised in hospital gym).	Cycle ergometer.	6 months	53 min/three sessions per week/60% of peak oxygen uptake (VO ₂ peak).	None specified.	Recommended to perform basic daily mild physical activities but to avoid any physical training.
Briffa <i>et al</i> , ⁴⁰ Australia	Aerobic circuit training (supervised in hospital).	Aerobic circuit training.	6 weeks	60–90 min/three sessions per week/ not specified.	Education and psychosocial counselling.	Education, pharmacotherapy and lifestyle counselling.
Giallauria <i>et al,</i> ³³ Italy	Moderate intensity exercise (supervised in centre).	Cycle ergometer.	6 months	40 min/three sessions per week/60%-70% of peak oxygen uptake (VO ₂ peak).	None specified.	Generic instructions on maintaining physical activity and a correct lifestyle.
Hambrecht <i>et al</i> , ³⁰ Germany	Moderate intensity exercise (supervised in hospital and unsupervised at home). Cycle ergometer.	Cycle ergometer.	12 months	10 min, 42 sessions per week (hospital), 20 min, seven sessions per week (home) plus 60 min group training, one session per week/70% of symptom-limited max HR.	None specified.	Standard medical therapy.
Higgins <i>et al,</i> ⁴¹ Australia	Moderate intensity walking programme (unsupervised at home). Walking.	Walking.	Not specified	Not specified/not specified/ notspecified.	Psychological plus education.	Psychological support, education, counselling and guidance.
Houle <i>et al</i> , ⁴² Canada	Pedometer-based walking programme (unsupervised at home). Walking.	Walking.	12 months	Not specified/not specified/ notspecified.	Education plus sociocognitive.	Sociocognitive support and advice regarding physical activity, diet and medication.
Kovoor <i>et al,</i> ⁴³ Australia	Standard cardiac rehabilitation programme (unknown setting).	Not specified.	5 weeks	Not specified/2–4 sessions per week/not specified.	Education and counselling.	Encouraged to exercise at home and return to normal activities.
Maddison <i>et al,</i> ⁴⁴ New Zealand	Automated package of text messages to increase exercise behaviour (unsupervised at home).	Moderate to vigourous aerobic exercise, for example, walking and household chores.	24 weeks	Minimum of 30 min/at least five sessions per week/not specified.	Optional access to other CR service or support.	Behaviour change therapy, encouragement to be physically active and advice to attend a cardiac club.
						Continued

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Table 3 Continued						
	Exercise intervention					
Reference, country Exercise	r Exercise	Modality	Study duration	Session duration/frequency/ intensity	Additional	Control (comparator)
Maroto Montero <i>et</i> al, ³⁴ Spain	Individualised physical training (supervised in hospital gym).	Physiotherapy and aerobic training on mats or a cycle ergometer.	3 months	60 min/three sessions per week/75%-85% max HR.	Psychological support, education plus return to work counselling.	Psychological support, education plus return to work counselling.
Munk <i>et al</i> , ³⁵ Norway	Moderate/high intensity interval training (supervised in centre).	Cycle ergometer or running.	6 months	60 min/three sessions per week/60%-70% and 80%- 90% max HR.	Spine and abdominal resistance training.	Usual care, including drug therapy.
Mutwalli <i>et al,</i> ⁴⁵ Saudi Arabia	Moderate intensity walking programme (unsupervised at home). Walking.	Walking.	6 months	30 min/seven sessions per week/ not specified.	Education.	Education, standard hospital care
Oerkild <i>et al</i> , ³⁶ Denmark	Moderate intensity exercise (unsupervised at home).	Individualised.	12 months	30min/six sessions per week/11– 13 on the Borg Scale.	Risk factor management.	Usual care, no exercise education or dietary counselling.
Reid <i>et al</i> , ⁴⁶ Canada	Internet-based physical activity plan and motivational tool to increase physical activity (unsupervised at home).	Not specified.	20 weeks	Not specified/not specified/ notspecified.	None specified.	Online education, physical activity guidance and an education booklet.
Santaularia <i>et al,</i> ²⁹ Spain	Outpatient exercise training programme (supervised in hospital). Cycle ergometer.	Cycle ergometer.	10 weeks	60min/threesessions per week/75%-90% max HR (RPE 11-15 on Borg Scale)	Resistance training, education and risk factor management.	Standard care, risk factor management, guidance on physical activity and adherence to medication.
Seki <i>et al</i> ,47 Japan	Moderate intensity aerobic exercise (supervised in centre and unsupervised at home).	Walking, cycle ergometer and jogging.	6 months	50–110 min, one session per week (centre), ≥30 min, two sessions per week (home)/12–13 on the standard Borg Scale.	Education.	Education and outpatient follow-up with physician.
Toobert <i>et al</i> , ⁴⁸ USA	Walking or aerobics (supervised in centre and unsupervised at home).	Walking or aerobics.	24 months	60 min, seven sessions per week (centre), 60 min, three sessions per week (home)/individually prescribed.	Education and psychological support.	Cooking classes, stress management and education.
Vestfold Heartcare Study Group, ³⁷ Norway	Dynamic endurance physical activity (supervised, group sessions in centre).	Dynamic endurance training.	ance 15 weeks	55 min/two sessions per week/RPE 11–13 on the Borg Scale, increased to 13–15 after 6 weeks.	Education and psychological support.	Education and psychological support.
						Continued

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Table 3 Continued						
	Exercise intervention					
Reference, country Exercise	Exercise	Modality	Study duration	Session duration/frequency/ intensity	Additional	Control (comparator)
Wang <i>et al</i> , ⁴⁹ China Not specified.	Not specified.	Not specified.	Not specified	Not specified/not specified/ notspecified.	Education.	Education.
West <i>et al</i> , ³¹ UK	Not specified, multicentre (supervised in centre).	Varied by centre (exercise equipment in physiotherapy gyms).	6–8 weeks	Averaged 20hours over 6–8 weeks/1–2 sessions per week/ notspecified.	Education plus psychological support.	Education plus psychological support.
Yu <i>et al</i> , ⁵⁰ China	Ambulatory and aerobic cardiovascular training (supervised in hospital and centre, unsupervised at home).	Walking, treadmill, cycle ergometry, rowing, stepper, arm ergometry and dumbbell	8 1/2 months	2 hours/two sessions per week (centre), not specified (home)/65%– 85% of maximal aerobic capacity (VO ₂ peak).	Resistance training and education.	Conventional medical therapy and education.
Zwisler <i>et al</i> , ³⁸ Denmark	Intensive CR programme (supervised in centre).	Not specified.	6 weeks	Not specified/two sessions per week/not specified.	Education and psychosocial support.	Education and psychosocial support.
CR, cardiac rehabilitatio	CR, cardiac rehabilitation; HR, heart rate; RPE, rating of perceived exertion; VO, Peak, peak oxygen uptake.	ng of perceived exertion;	VO, Peak, pea	ik oxygen uptake.		

period for all studies included in our review being 24.7 months, it may be that any benefits on mortality will accrue over a longer follow-up. However, the absence of any kind of signal in this review means a substantial longer term benefit is unlikely.

The current analyses do not quite exclude a worthwhile benefit of exercise-based CR on hospital admissions. While a risk difference of -0.05 (95% CI -0.10 to -0.00) is of borderline statistical significance, it is probably clinically unimportant in the context of no change in all-cause mortality.

From the studies included in this review, we do not know if there is a worthwhile benefit on quality of life, as a meta-analysis was not conducted. However, the authors of the 2016 Cochrane review reported that in 4 of the 22 studies included in this review, there was a significantly higher quality of life in at least half or more of the subscales.^{32 45 46 49}

Based on the present data, we are also unable to comment on whether exercise-based CR might be cost-effective. Five of the studies in this review included a within-trial health economic evaluation.^{30 40 43 44 50} Of these five papers, three studies showed no difference in healthcare costs between groups,^{40 43 50} one found healthcare costs to be lower for exercise-based CR³⁰ and one failed to report a P value for cost difference, ¹⁴ While a decrease, of borderline statistical significance, in hospital admissions may improve quality of life for patients, it is unclear if this confers any economic benefit, in the absence of robust cost-effectiveness analyses.

It may be that exercise-based CR has an effect on other outcomes, not specifically addressed in this review, such as cardiorespiratory fitness, lifestyle risk factor management, adherence to medication, diet, smoking cessation, psychosocial health and return to work.^{78,56,57} If the focus of future research is on measuring and improving these outcomes, attention will be needed to develop the best multicomponent intervention.

Strengths and limitations

To our knowledge, this is the first systematic review of exercise-based CR that has pooled data relevant to contemporary medical management of patients diagnosed with coronary artery disease. Although we have not done a de novo quality assessment of 21/22 studies included in this review and instead are relying on a previous Cochrane assessment, it is unlikely that we would have drawn different conclusions from such an assessment.¹⁰

The current review does not provide information on participant baseline characteristics. In the majority of studies (20/22; 91%), however, baseline characteristics were comparable between the intervention and control groups.^{10 29}

While there was no evidence of statistical heterogeneity across trials for all outcome measures (P value <0.01, $I^2 > 30\%$), except for hospital admissions, there was substantial context and interventional

	Exercise-ba	sed CR	No exercise	control		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
Briffa 2005	0	57	2	56	3.3%	-0.04 [-0.09, 0.02]	-+
Hambrecht 2004	1	51	2	50	2.5%	-0.02 [-0.09, 0.05]	
Higgins 2001	1	54	0	49	4.2%	0.02 [-0.03, 0.07]	+-
Houle 2012	0	32	0	33	3.3%	0.00 [-0.06, 0.06]	+
Kovoor 2006	0	72	0	70	15.1%	0.00 [-0.03, 0.03]	+
Maddison 2014	0	85	0	86	21.7%	0.00 [-0.02, 0.02]	+
Maroto 2005	7	90	16	90	1.2%	-0.10 [-0.20, -0.00]	
Munk 2009	0	20	0	20	1.3%	0.00 [-0.09, 0.09]	
Mutwalli 2012	0	28	1	21	0.8%	-0.05 [-0.16, 0.07]	
Oerkild 2012	4	19	5	21	0.2%	-0.03 [-0.29, 0.23]	
Reid 2012	0	115	2	108	11.9%	-0.02 [-0.05, 0.01]	-
Santaularia 2017	0	41	0	44	5.5%	0.00 [-0.04, 0.04]	+
Seki 2008	0	18	0	16	1.0%	0.00 [-0.11, 0.11]	
Toobert 2000	1	17	0	11	0.4%	0.06 [-0.11, 0.23]	
VHSG 2003	2	98	1	99	9.5%	0.01 [-0.02, 0.04]	+
Wang 2012	1	80	3	80	4.8%	-0.02 [-0.07, 0.02]	-+
West (RAMIT) 2012	245	903	243	910	6.7%	0.00 [-0.04, 0.05]	+
Yu 2004	4	132	4	72	3.0%	-0.03 [-0.09, 0.04]	-+
Zwisler 2008	24	227	20	219	3.6%	0.01 [-0.04, 0.07]	+
Total (95% CI)		2139		2055	100.0%	-0.00 [-0.02, 0.01]	
Total events	290		299				
Heterogeneity: Tau ² =	= 0.00; Chi ² =	10.60, df	= 18 (P = 0.92)	L); $I^2 = 0\%$	6		
Test for overall effect	: Z = 0.87 (P =	0.38)					-1 -0.5 0 0.5 1 Favours [Exercise-CR] Favours [No exercise-CR]

Figure 2 All-cause mortality for studies at their longest follow-up period. Filled squares represent the risk difference for individual studies at the longest reported follow-up. The boxes are proportional to the weight of each study, and the lines represent their 95% CI. The filled diamond represents the pooled risk difference. Weights are from random effects analysis. CR, cardiac rehabilitation.

heterogeneity. The studies came from a wide range of clinical environments and countries, and the interventions delivered ranged greatly in quality. When compared with both the BACPR 'minimum standards and core components'8 and Association of Chartered Physiotherapists in Cardiac Rehabilitation (ACPICR) guidelines,⁶ there was considerable variation in the exercise interventions delivered (table 3). Critics have questioned the exercise component reported in the largest included study-the RAMIT trial (n=1813).³¹ They argued that underdosage of exercise intensity and duration may have led to the inconclusive result.⁴ Several other studies included in this review also fail to report on the intensity, modality and/or duration of the exercise intervention. Exercise and physical activity has a 'dose-response' relationship with cardiovascular disease risk.⁵⁹ Moreover, a higher exercise capacity (VO₉ peak) is associated with an improvement

in mortality risk.⁶⁰ ⁶¹ If patients engaging in exercise-based CR do not achieve the correct dose of exercise, a physiological benefit is unlikely. It is a legitimate concern that participants in many included trials may not have received an adequate dose of exercise. In the era of contemporary medical management, higher intensity exercise protocols might be appropriate and effective.⁶²

One major concern is the reporting of adherence to, and fidelity of, exercise interventions.¹⁰ While the majority of studies included in this review report the intended prescription exercise dose^{29 30 32–37 39 40 47 50} (table 3), it is not possible to determine adherence and fidelity. Without basic reporting of these parameters, the actual exercise dose received cannot be quantified. This may have a significant bearing on intervention efficacy and the results of this meta-analysis. Moving forward, the introduction of checklists and reporting

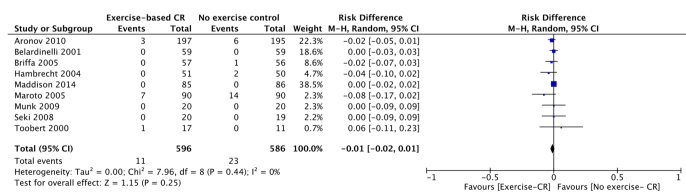


Figure 3 Cardiovascular mortality for studies at their longest follow-up period. Filled squares represent the risk difference for individual studies at the longest reported follow-up. The boxes are proportional to the weight of each study and the lines represent their 95% CI. The filled diamond represents the pooled risk difference. Weights are from random effects analysis. CR, cardiac rehabilitation.

Exercise-based CR

No exercise control

	LACICISE-Das	eu cr	NU EXERCISE C	Untrol		KISK DITTETETICE	Kisk Difference			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl			
Belardinelli 2001	11	59	21	59	6.1%	-0.17 [-0.33, -0.01]				
Briffa 2005	19	57	19	56	5.3%	-0.01 [-0.18, 0.17]				
Giallauria 2008	3	30	7	31	5.0%	-0.13 [-0.31, 0.06]				
Hambrecht 2004	1	51	7	50	9.9%	-0.12 [-0.22, -0.02]				
Maddison 2014	0	85	0	86	17.9%	0.00 [-0.02, 0.02]	+			
Mutwalli 2012	4	28	11	21	3.0%	-0.38 [-0.63, -0.13]				
Reid 2012	4	115	6	108	15.0%	-0.02 [-0.08, 0.03]	-			
Santaularia 2017	4	41	6	44	7.4%	-0.04 [-0.17, 0.10]				
VHSG 2003	11	98	14	99	10.9%	-0.03 [-0.12, 0.06]				
Yu 2004	34	132	16	72	8.4%	0.04 [-0.09, 0.16]				
Zwisler 2008	95	227	94	219	11.0%	-0.01 [-0.10, 0.08]				
Total (95% CI)		923		845	100.0%	-0.05 [-0.10, -0.00]	•			
Total events186201Heterogeneity: Tau ² = 0.00; Chi ² = 27.74, df = 10 (P = 0.002); l ² = 64%Test for overall effect: Z = 1.97 (P = 0.05)Favours [Exercise-CR]Favours [Exercise-CR]Favours [A Hospital admissions for studies at their longest follow up periodFigure 4										
Figure 4 Hospital admissions for studies at their longest follow-up period. Filled squares represent the risk difference for individual studies at the longest reported follow-up. The boxes are proportional to the weight of each study and the lines represent their 95% CI. The filled diamond represents the pooled risk difference. Weights are from random effects analysis. CR, cardiac rehabilitation.										
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Risk Difference

CONCLUSION

Based on the outcomes of all-cause mortality and cardiovascular mortality, our analysis indicates conclusively that the current approach to exercise-based CR has no effect when compared with a no-exercise control. There may be a small reduction in hospital admissions following exercise-based CR that is unlikely to be clinically important.

The continued delivery of exercise-based CR needs to be supported by new research to show its impact on health-related quality of life and whether it is a cost-effective intervention.

Contributors RP and MU were principally responsible for the study concept and design. RP and GM were responsible for study selection, data extraction and risk of bias assessment. With the assistance of University Hospital Coventry & Warwickshire library services, RP updated and ran the searches. RP, MU and PKK were responsible for statistical analysis and interpretation of data. GM and SE provided clinical advice. RP and MU wrote the first draft of the review, and all coauthors contributed to review and editing of drafts of the report. All authors approved the final manuscript. RP is the study guarantor and had full access to all trial level data in the review, takes responsibility for the integrity of the data, and accuracy of the data analysis, and had final responsibility to submit for publication.

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Data sharing statement No additional data available.

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Correction: Is exercise-based cardiac rehabilitation effective? A systematic review and meta-analysis to re-examine the evidence

Powell R, McGregor G, Ennis S, *et al.* Is exercise-based cardiac rehabilitation effective? A systematic review and meta-analysis to re-examine the evidence. *BMJ Open* 2018;8:e019656. doi: 10.1136/bmjopen-2017-019656.

The author was unaware of a protocol paper that changes a couple of things in this paper. It does not change the conclusions.

In Table 1, the 'Intention-to treat analysis conducted' is currently listed as 'High risk' but should read '**Low risk'**. The 'Support for judgement' column for this row should read '**Outcomes will be analysed on an intention-to-treat basis**.'

In Table 2, in the 'Santaularia *et al*,²⁹ Spain' row, the 'Recruitment period (years)' currently reads 'None specified' but this should read '**2010–2012**'.

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