National Institute for Health and Care Excellence

Final

Abdominal aortic aneurysm: diagnosis and management

Evidence review J: Pre- and postoperative interventions to optimise outcomes after abdominal aortic aneurysm repair

NICE guideline NG156 Methods, evidence and recommendations March 2020

Final

This evidence review was developed by the NICE Guideline Updates Team



FINAL

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Pre- and postoperative interventions to optimise outcomes after abdominal aortic aneurysm repair

Review questions

What preoperative interventions are effective in optimising surgical outcome in people undergoing surgical repair of an unruptured abdominal aortic aneurysm?

What post-operative interventions are effective in reducing the risk of complications after surgical repair of an abdominal aortic aneurysm, as well as optimising postoperative outcomes and survival?

Introduction

These review questions aim to determine which interventions can be used preoperatively to 'optimise' surgical outcome; and to identify which post-operative interventions are effective in reducing the risk of further aneurysm growth or rupture, cardiovascular events, wound-related complications and graft-related complications (including endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion).

PICO tables:

ultor	
Parameter	Inclusion criteria
Population	People with a confirmed unruptured abdominal aortic aneurysm (AAA) in whom surgery is planned
Interventions	Statins Beta-blockers Tranexamic acid Antiplatelet therapy Iron supplementation Coronary artery revascularisation Supervised exercise program Ischaemic preconditioning Respiratory training, including incentive spirometry and smoking cessation therapy
Comparators	Placebo, no intervention or each other
Outcomes	Mortality Peri- and post-operative complications Adverse effects of intervention Quality of life Resource use, including length of hospital or intensive care stay, and costs

Table 1: Inclusion criteria for preoperative interventions to optimise outcomes after AAA repair

Table 2: Inclusion criteria for postoperative interventions to optimise outcomes after AAA repair

Parameter	Inclusion criteria
Population	People who have undergone surgical repair of an AAA
Interventions	Surgical intervention Antifibrinolytic therapy with tranexamic acid Antiplatelet therapy (aspirin, clopidogrel, ticlopidine, cilostazol, prasugrel, ticagrelor, or any other antiplatelet drugs) Antihypertensive drugs (calcium channel blockers, angiotensin-converting enzyme (ACE) inhibitors, beta-blockers (β-blockers; e.g. metoprolol, propranolol), angiotensin-II receptor antagonists, thiazide/ thiazide-like diuretics, or any other antihypertensive drugs) Lipid-lowering therapy (statins (simvastatin, pravastatin, atorvastatin)) Antibiotics (doxycycline, roxithromycin, azithromycin) Diabetic control, including metformin COPD control Smoking cessation Physical therapy/exercise Diet Weight control Control of alcohol consumption
Comparators	Placebo, no intervention or each other
Outcomes	Incidence of complications (AAA rupture, AAA growth/expansion, cardiovascular events, wound-related complications, endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion) Need for further surgical intervention Mortality (all-cause; AAA-related; cardiovascular; survival) Cardiovascular events Quality of life Adverse effects Resource use and cost

Methods and process

This evidence review was developed using the methods and process described in <u>Developing NICE guidelines: the manual</u>. Methods specific to this review question are described in the review protocol in Appendix A.

Declarations of interest were recorded according to NICE's 2014 conflicts of interest policy.

A 'bulk' search strategy was used to cover review questions relating to pre- and postoperative interventions. Two searches were performed to identify studies that assessed the efficacy of interventions that could potentially be used to improve outcomes of surgical repair of AAAs. The first literature search used a randomised controlled trial (RCT) and systematic review (SR) filter while the second search used an observational study filter to identify potentially relevant studies.

The reviewer sifted the RCT database first to identify systematic reviews, RCTs or quasi-randomised controlled trials exploring the efficacy of preoperative interventions for improving outcomes in people who were due to undergo elective surgical repair of

unruptured AAAs, or postoperative interventions for improving outcomes in people who underwent elective AAA surgery. Studies were included if they met the set of criteria outlined in Tables 1 and 2 (the full protocol is available in Appendix A). If limited evidence was available from systematic reviews, RCTs or quasi-randomised controlled trials, the observational study database was sifted to identify potentially relevant non-randomised controlled trials.

Studies were excluded if they were not:

- were not in English
- were not full reports of the study (for example, published only as an abstract)
- were not peer-reviewed

Clinical evidence

Included studies

Preoperative interventions

From an initial RCT database of 916 abstracts, 24 were identified as being potentially relevant. Following full-text review of these articles, 8 studies were included. Additionally 1 study was identified via examination of the bibliography of an excluded systematic review. From an initial observational study database of 1,216 abstracts, 3 were identified as being potentially relevant. Following full-text review of these articles, no studies were included.

Update searches were conducted in December 2017, to identify any relevant studies published during guideline development. The update RCT and observational study databases contained 70 and 80 abstracts, respectively. Two abstracts from the RCT database and no abstracts from the observational study were considered potentially relevant. Following full text review of the 2 potentially relevant articles, 1 study was included

Overall, 10 studies were included in the evidence review for this review question.

Postoperative interventions

From the RCT database of 916 abstracts, 15 were identified as being potentially relevant. Following full-text review of these articles, 2 studies were included. From the observational study database of 1,216 abstracts, 2 were identified as being potentially relevant. Following full-text review of these articles, no studies were included.

Update searches were conducted in December 2017, to identify any relevant studies published during guideline development. The update RCT and observational study databases contained 70 and 80 abstracts, respectively. No abstracts from either database were considered potentially relevant, and no additional studies were included.

Overall, 2 studies were included in the evidence review for this review question.

Excluded studies

The list of papers excluded at full-text review, with reasons, is given in Appendix H.

Summary of clinical studies included in the evidence review

A summary of the included studies is provided in the tables below.

Preoperative interventions

Table 2: Beta-blockers

Study	Details
Yang H, Raymer K, Butler R, Parlow J, Roberts R. (2006) The	Study design: randomised, placebo-controlled, double-blind trial
effects of perioperative beta-	Location(s): Canada
blockade: results of the Metoprolol after Vascular Surgery (MaVS)	Population: people undergoing elective abdominal aortic surgery (no additional details were provided)
study, a randomized controlled trial. Am Heart J. 152(5):983-90	Sample size: 496; 76% (377/496) male
that. An field 3 . $132(3).903-90$	Follow-up: 30 months
	Intervention: 25 to 100 mg of oral or intravenous metoprolol, administered 2 hours before and after surgery, then continued for 5 days or until hospital discharge.
	Comparators: matched placebo
	Outcomes: the primary outcome was the composite rate of cardiac death, myocardial infarction, congestive heart failure, unstable angina, dysrhythmia requiring treatment, and non-cardiac death at 6 month follow-up. Individual rates were also reported at 30- day follow-up. Secondary outcomes included the need for reoperation, cerebrovascular accidents, new or worsened renal insufficiency, rehospitalisation, and intraoperative adverse events.

Table 3: Exercise

Study	Details
Barakat H M, Shahin Y, Khan J A et al. (2016) Preoperative supervised exercise improves outcomes after elective abdominal aortic aneurysm repair. Annals of Surgery 264, 47-53	Study design: randomised, non-blinded trial Location(s): UK Population: people with AAAs undergoing elective EVAR or open surgical repair Sample size: 124; 89.5% (111/124) male Follow-up: 3 months Intervention: hospital-based exercise classes Comparators: no exercise Outcomes: the primary outcome was the composite rate of cardiac, pulmonary, and renal complications. Secondary outcomes included length of stay, APACHE II scores, occurrence of systematic inflammatory response syndrome, mortality, and bleeding requiring reoperation or transfusion.
Dronkers J, Veldman A, Hoberg E et al. (2008) Prevention of pulmonary complications after upper abdominal surgery by preoperative intensive inspiratory muscle training: a randomized controlled pilot study. Clinical rehabilitation 22, 134-42	Study design: randomised, single-blind trial Location(s): Netherlands Population: people with AAAs undergoing elective surgical repair (not specified) who were considered to have a high risk of pulmonary complications Sample size: 20; 20% (5/15) male Follow-up: 7 days Intervention: inspiratory muscle training

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Study	Details
	Comparators: no exercise Outcomes: incidence of atelectasis, patient satisfaction, and respiratory function
Tew GA, Batterham AM, Colling K, et al. (2017) Randomized feasibility trial of high-intensity interval training before elective abdominal aortic aneurysm repair. The British journal of surgery 104(13), 1791-1801	Study design: randomised, single-blind trial Location(s): UK Population: people with unruptured AAAs between 5.5 and 7.0 cm in diameter who were due to undergo elective EVAR or open surgical repair Sample size: 53; 94.3% (50/53) male Follow-up: 12 weeks Intervention: High intensity interval training Comparators: no exercise Outcomes: adverse events, length of stay, and quality of life

Table 4: Remote ischaemic preconditioning (RIPC)

Study	Details
Ali ZA, Callaghan CJ, Lim E et al. (2007) Remote ischemic preconditioning reduces myocardial and renal injury after elective abdominal aortic aneurysm repair: a randomized controlled trial. Circulation 116, 198-105	Study design: randomised, double-blind trial Location(s): UK Population: people with AAAs undergoing elective open surgical repair Sample size: 82; 93% (76/82) male Follow-up: 7 days Intervention: lower limb RIPC Comparators: conventional open surgical repair without RIPC Outcomes: length of stay, mortality, myocardial injury, myocardial infarction, renal impairment, and adverse events
Li C, Li YS, Xu M et al. (2013) Limb remote ischemic preconditioning for intestinal and pulmonary protection during elective open infrarenal abdominal aortic aneurysm repair: a randomized controlled trial. Anesthesiology 118, 842-52	Study design: randomised, double-blind trial Location(s): China Population: people with AAAs undergoing elective open surgical repair Sample size: 62; 90.1% (55/61) male Follow-up: 24 hours Intervention: upper limb RIPC Comparators: sham RIPC Outcomes: the primary outcomes were haemodynamic data and variables reflecting lung function. Secondary outcomes included mortality, ventilator support time, ICU- and hospital-free days; new arrhythmia, perioperative myocardial infarction, diagnosis of congestive heart failure, symptoms and signs of pulmonary congestion, neurologic events, upper limb ischemia requiring intervention, intestinal injury markers, markers of oxidative stress and systemic inflammatory response, and scores of the severity of intestinal and pulmonary injury.
Mouton R, Pollock J, Soar J et al. (2015) Remote ischaemic preconditioning versus sham procedure for abdominal aortic aneurysm repair: an external	Study design: randomised, double-blind trial Location(s): UK Population: people with AAAs undergoing elective EVAR or open surgical repair.

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Study	Details
feasibility randomized controlled	Sample size: 69; sex-specific proportions were not
trial. Trials 16, 377	reported. Follow-up: 48 hours
	Intervention: upper limb RIPC
	Comparators: sham RIPC
	Outcomes: acute kidney injury scores as classified by the acute injury network (AKIN), mortality, myocardial
	infarction, new postoperative ECG changes, new arrhythmia, troponin T levels above 14 ng/L, and adverse events
Murphy N, Vijayan A, Frohlich S et al. (2014) Remote ischemic	Study design: randomised, double-blind trial Location(s): UK
preconditioning does not affect the incidence of acute kidney injury	Population: people with AAAs undergoing elective open surgical repair
after elective abdominal aortic aneurysm repair. Journal of cardiothoracic and vascular	Sample size: 62; 85.5% (53/62) male Follow-up: 3 days
anaesthesia 28, 1285-92	Intervention: upper limb RIPC Comparators: sham RIPC
	Outcomes: mortality, kidney injury (measured by creatinine levels and AKIN scores), myocardial infarction, and length of hospital stay
Walsh SR, Boyle JR, Tang TY et	Study design: randomised, non-blinded trial
al. (2009) Remote ischemic	Location(s): UK
preconditioning for renal and	Population: people with AAAs undergoing elective
cardiac protection during	open surgical repair
endovascular aneurysm repair: a randomized controlled trial.	Sample size: 40 men
Journal of endovascular therapy:	Follow-up: 48 hours
an official journal of the	Intervention: lower limb RIPC
International Society of Endovascular Specialists 16, 680-	Comparators: conventional open surgical repair without RIPC
9	Outcomes: the primary outcome measure was renal function (measured by urine output, urine retinal binding protein, and creatinine levels). Secondary outcomes included 30-day mortality, myocardial infarction, arrhythmia, congestive heart failure, pneumonia, renal failure, lower limb ischaemia requiring intervention, and postoperative length of stay.
Walsh SR, Sadat U, Boyle JR et al. (2010) Remote ischemic	Study design: randomised, non-blinded trial Location(s): UK
preconditioning for renal protection during elective open	Population: people with AAAs undergoing elective EVAR
infrarenal abdominal aortic aneurysm repair: randomized	Sample size: 40; 85% (34/40) male
ontrolled trial. Vascular and ndovascular surgery 44, 334-40	Follow-up: 48 hours
	Intervention: lower limb RIPC
	Comparators: conventional open surgical repair without RIPC.
	Outcomes: the primary outcome measure was renal function (measured by urine output, urine retinal binding protein, and serum creatinine levels). Secondary outcomes included serum troponin levels and the incidence of major adverse cardiac events
	and the molection of major deverse valuate events

Study	Details
	(cardiac arrest, cardiac death, cardiac failure, unstable angina, or myocardial infarction).

Postoperative interventions

Table 5: Doxycycline

Study	Details
Hackmann AE, Rubin BG, Sanchez LA et al. (2008) A randomized, placebo- controlled trial of doxycycline after endoluminal aneurysm repair. Journal of vascular surgery 48, 519-526	Study design: randomised, placebo-controlled, double-blind trial Location(s): USA Population: people with AAAs undergoing elective EVAR Sample size: 59; sex-specific proportions not reported Follow-up: 6 months Intervention: doxycycline 100 mg b.i.d Comparators: matched placebo Outcomes: aneurysm diameter, graft migration, incidence of endoleak, adverse events

Table 6: Physiotherapy plus walking exercises

Study	Details
Wnuk BR, Durmala J, Ziaja K et al. (2016) A Controlled Trial of the Efficacy of a Training Walking Program in Patients Recovering from Abdominal Aortic Aneurysm Surgery. Advances in clinical and experimental medicine : official organ Wroclaw Medical University 25, 1241-1371	Study design: randomised, single-blind trial Location(s): Poland Population: people with AAAs undergoing surgical repair (type not specified) Sample size: 65 males Follow-up: 2 years Intervention: basic physiotherapy plus backward or forward walking exercises Comparators: basic physiotherapy-alone Outcomes: 6-minute walking test distance, walking speed, spirometry measurements (FVC, FEV1, FEV1/FVC and PEF), length of hospital stay

See Appendix D for full evidence tables.

Quality assessment of clinical studies included in the evidence review

See Appendix F for full GRADE tables, highlighting the quality of evidence from the included studies.

Economic evidence

Included studies

A literature search was conducted jointly for all review questions by applying standard health economic filters to a clinical search for AAA. This search returned a total of 5,173 citations. Following review of all titles and abstracts, no studies were identified as being potentially relevant to these review questions.

An update search was conducted in December 2017, to identify any relevant health economic analyses published during guideline development. The search found 814

abstracts; all of which were not considered relevant to this review. As a result no additional studies were included.

Excluded studies

No studies were retrieved for full-text review.

Evidence statements

Preoperative interventions

Beta-blockers

- Moderate- to high-quality evidence from 1 RCT, including 496 people who underwent elective AAA repair (type not specified), found higher rates of intraoperative hypotension and bradycardia requiring treatment in people who received preoperative beta-blockers compared with those who received placebo.
- Low-quality evidence from 1 RCT, including 496 people who underwent elective AAA repair (type not specified), could not differentiate between rates of cardiac-related mortality, non-cardiac-related mortality, unstable angina, myocardial infarction, congestive heart failure, postoperative cardiovascular accident, dysrhythmia, new or worsened renal insufficiency, and the need for reoperation in people who received preoperative beta-blockers compared with those who received placebo.

Exercise

- Low-quality evidence from 1 RCT, including 20 people who underwent elective AAA repair (type not specified), could not differentiate between atelectasis rates of people who had been performing inspiratory muscle training during the 2 weeks preceding surgery and those who had not been doing any training.
- Moderate-quality evidence from 1 RCT, including 124 people who underwent elective EVAR, indicated that cardiac complications and renal complications were less likely to occur in people who participated in preoperative hospital-based exercise classes compared with those who had not. Low-quality evidence from the same trial could not differentiate between rates of all-cause mortality, pulmonary complications, postoperative bleeding or the need for a blood transfusions of more than 4 units, and the need for reoperation between people who participated in preoperative hospital-based exercise classes and those who did not.
- Low- to moderate-quality evidence from 1 RCT, including 53 people who underwent elective EVAR or open surgical repair, could not differentiate preoperative dizziness, preoperative angina, and postoperative quality of life between people who participated in preoperative hospital-based exercise classes and those who did not.

Remote ischaemic preconditioning

- Low-quality evidence from 5 RCTs, including 273 people who underwent elective EVAR or open AAA repair, found higher rates of arrhythmia in people who received remote ischaemic preconditioning before surgery compared with those who received no preconditioning.
- Very low- to low-quality evidence from up to 6 RCTs, including 355 people who underwent elective EVAR or open AAA repair, could not differentiate rates of 30day mortality, myocardial infarction, congestive heart failure, renal impairment or failure, acute kidney injury, and rates of any type of complication between people who did and did not receive remote ischaemic preconditioning.

Postoperative interventions

Doxycycline versus placebo

- Very low-quality evidence from 1 RCT, including 27 people with AAAs who underwent elective EVAR, could not differentiate mean percentage changes in aneurysm diameters between people who received doxycycline after surgery and those who did not.
- Very low-quality evidence from 1 RCT, including 48 people who underwent elective EVAR of AAAs, could not differentiate endoleak and graft migration rates between people who received doxycycline after surgery and those who did not.

Physiotherapy plus walking exercises versus physiotherapy-alone

• Very low-quality evidence from 1 RCT, including 47 people who underwent elective AAA repair (type not specified) of AAAs, could not differentiate the average length of hospital stay between people who received postoperative physiotherapy plus forward or backward walking exercises and people who received physiotherapy-alone.

Research recommendations

Preoperative interventions

RR4. What is the clinical effectiveness and cost effectiveness of preoperative exercise programmes for improving outcomes of people who are having AAA repair?

Postoperative interventions

RR5. What are the benefits of postoperative use of Direct Oral Anticoagulants (DOACS) for improving outcomes after repair of AAA?

The committee's discussion of the evidence

Interpreting the evidence

The outcomes that matter most

In relation to preoperative interventions, the outcomes that matter most are perioperative morbidity and mortality. With regards to postoperative interventions, the outcomes which matter most are postoperative morbidity and mortality, and the need for re-intervention.

The quality of the evidence

The committee considered that the identified evidence on preoperative exercise interventions was not robust enough to support a recommendation. Although the identified evidence for most outcomes were graded as being low-to-moderate in quality, the committee felt that the small sample sizes of included studies and relatively short follow-up periods precluded confidence in the reported outcomes. In relation to beta-blockers and RIPC, the committee considered that the evidence was of sufficient quality to draft recommendations.

This review excluded evidence on pre- and postoperative interventions for heterogeneous groups of people with vascular diseases who were treated by different types of surgical and non-surgical interventions, including AAA repair. It was noted that excluded studies did not report what proportions of people received AAA surgery or did not stratify analyses according to type of intervention received. As a result, the committee felt that this type of evidence could not be considered because of uncertain applicability to people with AAA.

The committee noted the existence of specialist society (Vascular Society of Great Britain and Ireland) guidelines related to AAA repair. It was noted that some of the recommendations in these guidelines relating to preoperative interventions were not based on evidence specific to people with AAA. The committee took these recommendations into account but refrained from cross-referring to specific recommendations.

With respect to postoperative interventions, the committee considered that the identified evidence was limited in both quantity and quality. The two identified studies were graded as very low to low in quality and indicated that postoperative use of doxycycline and physiotherapy plus walking exercises had no impact on the incidence of postoperative complications. As a result, the committee decided not to make any recommendations regarding these interventions.

Benefits and harms

The committee recognised that most people with AAA are likely to be older people with some form of cardiovascular disease. With this in mind the committee believed that optimisation of pre-existing medical conditions and minimisation of cardiovascular risks would increase the general health of people with AAA, leading to reduced postoperative morbidity and mortality. As a result, the committee felt that general principles of secondary prevention of cardiovascular disease, as outlined in other NICE guidelines, were applicable. The committee also agreed that it is important to reduce the risks of surgical site infections and venous thromboembolism in all people undergoing AAA repair. As result, recommendations were drafted cross-referring to other NICE guidance.

The committee agreed that the evidence on beta-blockers was clear that de novo beta-blockade in the immediate preoperative period was not effective and was potentially harmful. It was also considered that the evidence on beta-blockers in relation to AAA repair was consistent with broader evidence on the use of betablockade in other surgical cohorts, including people undergoing other types of vascular surgery. The committee felt that use of the word "routinely" allowed scope for clinician discretion given that there will be certain indications, for example atrial fibrillation, where beta blockade remains appropriate. It was clear that discontinuation of beta-blockers in such circumstances would be bad practice.

The committee felt that body of evidence on RIPC strongly indicated no benefit to postoperative outcomes, and the potential for harm (arrhythmia). Unlike betablockers, the committee felt that there was no particular circumstance where routine use of RIPC should be considered. Thus, a "do not use" recommendation was made.

The committee recognised the risk of thromboembolic events (such as deep venous thrombosis and pulmonary embolism) after AAA repair, and noted that no evidence was found relating to the use of postoperative anticoagulation in people who have undergone AAA repair. They noted that Direct-acting Oral Anticoagulants (DOACs) have become popular in clinical practice because they are easy to use, have good pharmacokinetic properties associated with fixed dosing, have few interactions with other medications, and require less frequent monitoring. With that in mind, the committee drafted a research recommendation to encourage research on how best to use DOACs in the postoperative period to balance the risk thromboembolic events with that of bleeding.

The committee noted that there is a widespread problem of people with AAA not having their medical therapy optimised, and that it would be good practice for clinicians to optimise medical therapy in all people identified as having an AAA, whether or not they were due to undergo AAA repair. The committee also agreed that it would be good practice for clinicians to perform preoperative medication assessments in order to optimise patient care.

Cost effectiveness and resource use

The committee considered that recommendations relating to secondary prevention of cardiovascular disease, prevention and treatment of surgical site infections, and reduction of the risk of venous thromboembolism were unlikely to have an impact on costs and resource use, because they simply cross-refer to existing guidance and reaffirm best clinical practice.

The committee considered the potential costs of treating intraoperative complications of preoperative beta-blockade (hypotension and bradycardia requiring treatment) and believed that a do not use recommendation would prevent such unnecessary expenses from occurring.

Other factors the committee took into account

The committee felt that NHS providers have already started devoting resources to exercise programmes based on a relatively small body of evidence. Thus, there is a role for further research to inform funding decisions. The committee agreed to make their research recommendation purposely broad, to maximise researcher uptake. It was agreed that the research recommendation should not explicitly state the need to monitor "cardiopulmonary" outcomes because there was some concern that researchers would focus on cardiac outcomes, at the expense of respiratory outcomes.

Appendices

Appendix A – Review protocols

Review protocol for review question 11: preoperative interventions to optimise outcomes after AAA repair

•	
Review question 11	What presurgical interventions are effective in optimising surgical outcome in people undergoing surgical repair of an unruptured abdominal aortic aneurysm?
Objectives	To determine which interventions can be used preoperatively to 'optimise' surgical outcome.
Type of review	Intervention
Language	English
Study design	Systematic reviews of study designs listed below Randomised controlled trials Quasi-randomised controlled trials If insufficient evidence identified, non-randomised controlled trials
Status	Published papers only (full text) No date restrictions
Population	People with a confirmed unruptured abdominal aortic aneurysm in whom surgery is planned
Intervention	Statins Beta-blockers Tranexamic acid Antiplatelet therapy Iron supplementation Coronary artery revascularisation Supervised exercise program Ischaemic preconditioning Respiratory training, including incentive spirometry and smoking cessation therapy
Comparator	Placebo, no intervention or each other
Outcomes	Mortality Peri- and post-operative complications Adverse effects of intervention Quality of life Resource use, including length of hospital or intensive care stay, and costs
Other criteria for inclusion / exclusion of studies	Exclusion: Non-English language Abstract/non-published Pharmacological interventions not available in the UK
Baseline characteristics to be extracted in evidence tables	Age Sex Size of aneurysm Comorbidities
Search strategies	See Appendix B
Review strategies	Appropriate NICE Methodology Checklists, depending on study designs, will be used as a guide to appraise the quality of individual studies.

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Review question 11	What presurgical interventions are effective in optimising surgical outcome in people undergoing surgical repair of an unruptured abdominal aortic aneurysm?
	Data on all included studies will be extracted into evidence tables. Where statistically possible, a meta-analytic approach will be used to give an overall summary effect.
	All key findings from evidence will be presented in GRADE profiles and further summarised in evidence statements.
Key papers	Dronkers, A. Veldman, E. Hoberg, C. van der Waal, N. van Meeteren. Prevention of pulmonary complications after upper abdominal surgery by pre-operative intensive inspiratory muscle training: a randomized controlled pilot study. Clin Rehabil, 22 (2008), pp. 134–142
	Kothmann, A.M. Batterham, S.J. Owen, A.J. Turley, M. Cheesman, A. Parry, et al. Effect of short-term exercise training on aerobic fitness in patients with abdominal aortic aneurysms: a pilot study. Br J Anaesth, 103 (2009), pp. 505–510
	Myers, 2010. Effects of exercise training in patients with AAA: preliminary resits from a randomised trial. J Cardiopulm Rehab Prev, 30 (2010), pp. 374–383
	Tew, 2011. Endurance exercise training in patients with small abdominal aortic aneurysm: a randomised controlled pilot study. Arch Phys Med Rehabil, 93 (2012), pp. 2148–2153
	Ali ZA, Callaghan CJ, Lim E, Ali AA, Nouraei SA, Akthar AM, Boyle JR, Varty K, Kharbanda RK, Dutka DP, Gaunt ME. Remote ischemic preconditioning reduces myocardial and renal injury after elective abdominal aortic aneurysm repair: a randomized controlled trial. Circulation. 2007 Sep 11;116(11 Suppl):I98-105 Mouton R, Pollock J, Soar J, Mitchell DC, Rogers CA. Remote ischaemic preconditioning versus sham procedure for abdominal aortic aneurysm repair: an external feasibility randomized controlled trial. Trials. 2015 Aug 25;16:377 Walsh SR, Sadat U, Boyle JR, Tang TY, Lapsley M, Norden AG, Gaunt ME. Remote ischemic preconditioning for renal protection during elective open infrarenal abdominal aortic aneurysm repair: randomized controlled trial. Vasc Endovascular Surg. 2010 Jul;44(5):334-40
	Walsh SR, Boyle JR, Tang TY, Sadat U, Cooper DG, Lapsley M, Norden AG, Varty K, Hayes PD, Gaunt ME. Remote ischemic preconditioning for renal and cardiac protection during endovascular aneurysm repair: a randomized controlled trial. J Endovasc Ther. 2009 Dec;16(6):680-9

Review protocol for review question 30: postoperative interventions to optimise outcomes after AAA repair

Review question 30	What postoperative interventions are effective in reducing the risk of complications after surgical repair of an abdominal aortic aneurysm, as well as optimising postoperative outcomes and survival?	
Objectives	To identify which postoperative interventions are effective in reducing the risk of further aneurysm growth or rupture, CV events, wound-related complications and graft-related complications (including endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion).	
Type of review	Intervention	
Language	English	
Study design	Systematic reviews of study designs listed below Randomised controlled trials Quasi-randomised controlled trials If insufficient evidence identified, non-randomised controlled trials	
Status	Published papers only (full text) No date restrictions	

Abdominal aortic aneurysm: evidence review for pre- and postoperative interventions to optimise outcomes after abdominal aortic aneurysm repair (March 2020)

What postoperative interventions are effective in reducing the risk of complications after surgical repair of an abdominal aortic aneurysm, as well
as optimising postoperative outcomes and survival?
People who have undergone surgical repair of an abdominal aortic aneurysm
Surgical intervention
Antifibrinolytic therapy with tranexamic acid Antiplatelet therapy (aspirin, clopidogrel, ticlopidine, cilostazol, prasugrel, ticagrelor,
or any other antiplatelet drugs)
Antihypertensive drugs (calcium channel blockers, angiotensin-converting enzyme (ACE) inhibitors, beta-blockers (β -blockers; e.g. metoprolol, propranolol), angiotensin-II receptor antagonists, thiazide/ thiazide-like diuretics, or any other antihypertensive drugs)
Lipid-lowering therapy (statins (simvastatin, pravastatin, atorvastatin))
Antibiotics (doxycycline, roxithromycin, azithromycin)
Diabetic control, including metformin
COPD control
Smoking cessation Physical therapy/exercise
Diet
Weight control
Control of alcohol consumption
Placebo, no intervention or each other
Incidence of complications (AAA rupture, AAA growth/expansion, cardiovascular events, wound-related complications, endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion) Need for further surgical intervention
Mortality (all-cause; AAA-related; cardiovascular; survival) Cardiovascular events Quality of life Adverse effects
Resource use and cost
Exclusion:
Non-English language
Abstract/non-published
Pharmacological interventions not available in the UK
Age
Sex
Size of aneurysm
Comorbidities
See Appendix B
Appropriate NICE Methodology Checklists, depending on study designs, will be used as a guide to appraise the quality of individual studies. Data on all included studies will be extracted into evidence tables. Where statistically possible, a meta-analytic approach will be used to give an overall
summary effect.
All key findings from evidence will be presented in GRADE profiles and further summarised in evidence statements.
Yang H, Raymer K, Butler R, Parlow J, Roberts R. The effects of perioperative beta-blockade: results of the Metoprolol after Vascular Surgery (MaVS) study, a randomized controlled trial. Am Heart J. 2006 Nov;152(5):983-90

Appendix B – Literature search strategies

Clinical search literature search strategy

Main searches

Bibliographic databases searched for the guideline

- Cumulative Index to Nursing and Allied Health Literature CINAHL (EBSCO)
- Cochrane Database of Systematic Reviews CDSR (Wiley)
- Cochrane Central Register of Controlled Trials CENTRAL (Wiley)
- Database of Abstracts of Reviews of Effects DARE (Wiley)
- Health Technology Assessment Database HTA (Wiley)
- EMBASE (Ovid)
- MEDLINE (Ovid)
- MEDLINE Epub Ahead of Print (Ovid)
- MEDLINE In-Process (Ovid)

Identification of evidence for review questions

The searches were conducted between November 2015 and October 2017 for 31 review questions (RQ). In collaboration with Cochrane, the evidence for several review questions was identified by an update of an existing Cochrane review. Review questions in this category are indicated below. Where review questions had a broader scope, supplement searches were undertaken by NICE.

Searches were re-run in December 2017.

Where appropriate, study design filters (either designed in-house or by McMaster) were used to limit the retrieval to, for example, randomised controlled trials. Details of the study design filters used can be found in section 4.

Search strategy review questions 11 and 30

Medline Strategy, searched 16th May 2017 Database: Ovid MEDLINE(R) 1946 to May Week 1 2017 Search Strategy:

- 1 Aortic Aneurysm, Abdominal/
- 2 Aortic Rupture/

3 (aneurysm* adj4 (abdom* or thoracoabdom* or thoraco-abdom* or aort* or spontan* or juxtarenal* or juxta-renal* or juxta renal* or paraerenal* or para-renal* or para renal* or suprarenal* or supra-renal* or short neck* or short-neck* or shortneck* or visceral aortic segment*)).tw.

- 4 (AAA* or RAAA*).tw.
- 5 or/1-4
- 6 Preoperative Care/ or Perioperative Care/ or Perioperative Nursing/ or Postoperative Care/
- 7 home care services/ or home care services, hospital-based/

8 (presurg* or pre-surg* or pre surg* or preop* or pre-op* or pre op or periop* or peri-op* or peri op*).tw.

9 ((perianaesthe* or perianesthe* or surgical) adj4 nursing).tw.

Medline Strategy, searched 16th May 2017 Database: Ovid MEDLINE(R) 1946 to May Week 1 2017 Search Strategy:

10 ((before or plan* or electiv* or ahead* or prepar* or prior) adj4 (surg* or operat* or procedure* or repair* or care* or outcome*)).tw.

11 (postsurg* or post-surg* or post surg* or postop* or post-op* or post op*).tw.

12 ((after or follow* or electiv* or post*) adj4 (surg* or operat* or procedure* or repair* or care* or outcome*)).tw.

- 13 (medical* adj4 (therap* or treat* or interven* or manag*)).tw.
- 14 Elective Surgical Procedures/
- 15 Endovascular Procedures/ or Vascular Surgical Procedures/
- 16 (endovascular* adj4 aneurysm* adj4 repair*).tw.
- 17 (endovascular* adj4 aort* adj4 repair*).tw.
- 18 (upper adj4 abdominal adj4 (repair* or surger* or surgic* or operat* or procedur*)).tw.
- 19 (EVAR or EVRAR or FEVAR or F-EAVAR or BEVAR or B-EVAR).tw.
- 20 (Anaconda or Zenith Dynalink or Hemobahn or Luminex* or Memoth-erm or Wallstent).tw.
- 21 (Viabahn or Nitinol or Hemobahn or Intracoil or Tantalum).tw.
- 22 or/6-21
- 23 exp Antifibrinolytic Agents/
- 24 ((antifibrinolytic or anti-fibrinolytic) adj4 (hemostatic or haemostatic or agent*)).tw.
- 25 ((tranexam* or tranex-am* or tranex am* or tranexan or tranex-and or tranex an) adj4 acid*).tw.
- 26 TXA.tw.
- 27 (aminocaproic* adj4 acid*).tw.
- 28 vitamin k*.tw.
- 29 (anti plasmin* or anti-plasmin* or (plasmin* adj4 inhibitor*)).tw.
- 30 Iron/
- 31 iron.tw.
- 32 exp Coronary Artery Bypass/
- 33 ((coronary adj4 arter* adj4 bypass*) or (aortocoronary adj4 bypass*)).tw.
- 34 ((off-pump or off pump) adj4 bypass*).tw.
- 35 CABG.tw.
- 36 ((blood-flow or blood flow or perfus*) adj4 restor*).tw.
- 37 (coronary adj4 (revasculari* or recanali* or reperfus*)).tw.
- 38 Antibiotic Prophylaxis/
- 39 (antibiotic* adj4 (premed* or prophyla*)).tw.

40 (Doxycyclin* or Atridox or Cyclodox or Demix or Doxylar or Efracea or Nordox or Periostat or Ramysis or Vibramycin or Vibramycin).tw.

- 41 Roxithromycin*.tw.
- 42 (Azithromycin* or Azyter or Clamelle or Zedbac or Zithromax).tw.
- 43 Smoking Cessation/
- 44 "Tobacco Use Cessation"/
- 45 ((cigarette* or smok* or tobacco or nicotine*) adj4 (cessation or withdrawal or ceas*)).tw.

46 ((quit* or stop* or giv* or abstin* or abstain*) adj4 (tobacco or cigarette or smoking or nicotine*)).tw.

47 (smoking adj4 (therap* or rehab*)).tw.

48 (cessation adj4 (treat* or therap* or assist* or advice or advis* or program* or interven* or service*)).tw.

49 Motor Activity/

Medline Strategy, searched 16th May 2017 Database: Ovid MEDLINE(R) 1946 to May Week 1 2017 Search Strategy:

50 ((motor or physical* or locomotor or supervis*) adj4 activit*).tw.

- 51 exp Exercise/ or Exercise Therapy/
- 52 (exercise* or exercisi* or kinesiotherap*).tw.
- 53 exp Physical Fitness/
- 54 Physical endurance/
- 55 fitness*.tw.
- 56 (walk* or swim* or jog* or cycl* or bicycl* or bike* or gym*).tw.

57 ((physical* or keep* or cardio* or aerobic or fitness or endurance) adj4 (fit* or activit* or active or train* or therap*)).tw.

- 58 (aerobic adj4 condition*).tw.
- 59 Muscle strength/
- 60 (muscle adj4 strength*).tw.
- 61 Ischemic Preconditioning/
- 62 ((ischemic* or ischaemic* or remote) adj4 (precondition* or pre-condition* or pre condition*)).tw.
- 63 (IPC or RIC or RIPC).tw.
- 64 Respiratory therapy/
- 65 exp Breathing Exercises/

66 ((breath* or respirat* or inhal*) adj4 (exercis* or therap* or train* or alter* or chang* or deepen* or physio* or rehab*)).tw.

- 67 exp Spirometry/
- 68 (spirometr* or bronchospirometr*).tw.
- 69 exp Diet/
- 70 (diet or diets or dieting).tw.
- 71 (health* adj4 eat*).tw.
- 72 exp Food/
- 73 food*.tw.
- 74 (weight adj4 (manag* or control* or maintain* or achiev* or goal* or health*)).tw.
- 75 exp Alcohol-Related Disorders/

76 (alcohol* adj4 (use* or abus* or drink* or reduc* or intake or consum* or control* or abstain* or abstinen* or depend* or addict* or chonic*)).tw.

- 77 ((problem* adj4 drink*) or (alcoholic* or alcoholism)).tw.
- 78 exp Pulmonary Disease, Chronic Obstructive/
- 79 Lung diseases, obstructive/
- 80 (COPD* or COAD* or COBD* or AECB*).tw.
- 81 (chronic adj4 obstruct* adj4 (disease* or airway*)).tw.
- 82 (chronic* adj4 (airflow* or airway* or bronch* or lung* or respirat* or pulmonary) adj4 obstruct*).tw.
- 83 exp Diabetes Mellitus/
- 84 diabet*.tw.
- 85 or/23-84
- 86 5 and 22 and 85
- 87 Aortic Aneurysm, Abdominal/su [Surgery]
- 88 85 and 87
- 89 86 or 88
- 90 animals/ not humans/

Medline Strategy, searched 16th May 2017 Database: Ovid MEDLINE(R) 1946 to May Week 1 2017 Search Strategy:

```
91 89 not 90
```

92 limit 91 to english language

Health Economics literature search strategy

Sources searched to identify economic evaluations

- NHS Economic Evaluation Database NHS EED (Wiley) last updated Dec 2014
- Health Technology Assessment Database HTA (Wiley) last updated Oct 2016
- Embase (Ovid)
- MEDLINE (Ovid)
- MEDLINE In-Process (Ovid)

Search filters to retrieve economic evaluations and quality of life papers were appended to the population and intervention terms to identify relevant evidence. Searches were not undertaken for qualitative RQs. For social care topic questions additional terms were added. Searches were re-run in September 2017 where the filters were added to the population terms.

Health economics search strategy

Medline Strategy Economic evaluations

- 1 Economics/
- 2 exp "Costs and Cost Analysis"/
- 3 Economics, Dental/
- 4 exp Economics, Hospital/
- 5 exp Economics, Medical/
- 6 Economics, Nursing/
- 7 Economics, Pharmaceutical/
- 8 Budgets/
- 9 exp Models, Economic/
- 10 Markov Chains/
- 11 Monte Carlo Method/
- 12 Decision Trees/
- 13 econom*.tw.
- 14 cba.tw.
- 15 cea.tw.
- 16 cua.tw.
- 17 markov*.tw.
- 18 (monte adj carlo).tw.
- 19 (decision adj3 (tree* or analys*)).tw.
- 20 (cost or costs or costing* or costly or costed).tw.
- 21 (price* or pricing*).tw.
- 22 budget*.tw.
- 23 expenditure*.tw.
- 24 (value adj3 (money or monetary)).tw.

Medline Strategy

25 (pharmacoeconomic* or (pharmaco adj economic*)).tw.

26 or/1-25

Quality of life

- 1 "Quality of Life"/
- 2 quality of life.tw.
- 3 "Value of Life"/
- 4 Quality-Adjusted Life Years/
- 5 quality adjusted life.tw.
- 6 (qaly* or qald* or qale* or qtime*).tw.
- 7 disability adjusted life.tw.
- 8 daly*.tw.
- 9 Health Status Indicators/

10 (sf36 or sf 36 or short form 36 or shortform 36 or sf thirtysix or sf thirty six or shortform thirtysix or short form thirtysix or short form thirtysix.

11 (sf6 or sf 6 or short form 6 or shortform 6 or sf six or sfsix or shortform six or short form six).tw.

12 (sf12 or sf 12 or short form 12 or shortform 12 or sf twelve or sftwelve or shortform twelve or short form twelve).tw.

13 (sf16 or sf 16 or short form 16 or shortform 16 or sf sixteen or sfsixteen or shortform sixteen or short form sixteen).tw.

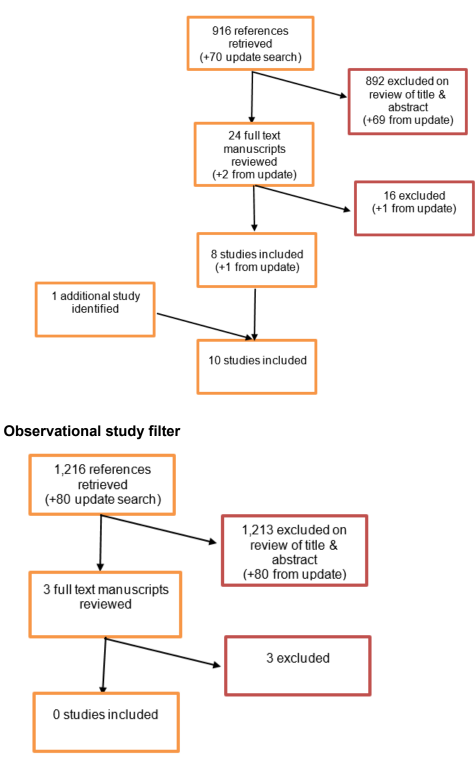
14 (sf20 or sf 20 or short form 20 or shortform 20 or sf twenty or sftwenty or shortform twenty or short form twenty).tw.

- 15 (euroqol or euro qol or eq5d or eq 5d).tw.
- 16 (qol or hql or hqol or hrqol).tw.
- 17 (hye or hyes).tw.
- 18 health* year* equivalent*.tw.
- 19 utilit*.tw.
- 20 (hui or hui1 or hui2 or hui3).tw.
- 21 disutili*.tw.
- 22 rosser.tw.
- 23 quality of wellbeing.tw.
- 24 quality of well-being.tw.
- 25 qwb.tw.
- 26 willingness to pay.tw.
- 27 standard gamble*.tw.
- 28 time trade off.tw.
- 29 time tradeoff.tw.
- 30 tto.tw.
- 31 or/1-30

Appendix C – Clinical evidence study selection

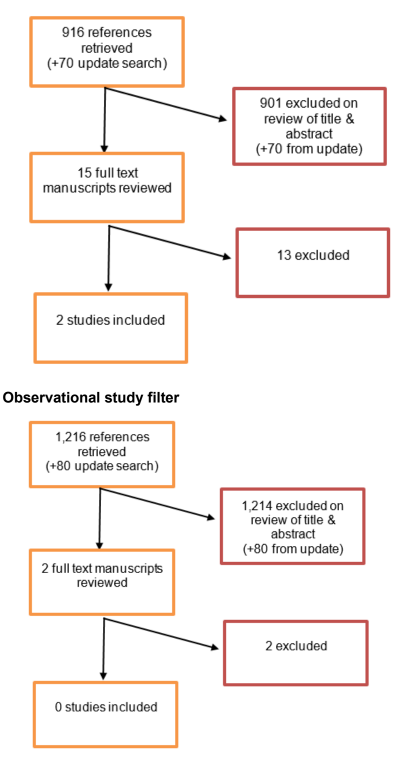
Review question 11 (preoperative interventions) study selection

RCT filter



Review question 30 (postoperative interventions) study selection

RCT filter



Appendix D – Clinical evidence tables

Evidence tables for review question 11 (preoperative interventions)

Beta-blockers

Full citation	Yang H, Raymer K, Butler R, Parlow J, Roberts R. (2006) The effects of perioperative beta-blockade: results of the Metoprolol after Vascular Surgery (MaVS) study, a randomized controlled trial. Am Heart J. 152(5):983-90.
Study details	Study type: randomised, placebo-controlled, double-blind trial Location(s): Canada Aim(s): to assess the efficacy of perioperative metoprolol on postoperative outcomes of patients undergoing abdominal aortic surgery Study dates: 1999 to 2002 Follow-up: 30 months Sources of funding: Heart and Stroke Foundation of Canada
Participants	 Population: patients undergoing elective abdominal aortic surgery (no additional details were provided). Sample size: 496; 76% (377/496) male Inclusion criteria: patients with American Society of Anaesthesiology class of 3 or less undergoing abdominal aortic surgery and infrainguinal or axillofemoral revascularization were included Exclusion criteria: current or recent use of beta-blockers or amiodarone, an airflow obstruction requiring treatment, history of congestive heart failure, a history of atrioventricular block, or previous adverse drug reactions to beta-blockers Baseline characteristics: Mean age: Beta-blocker group, 66.4 years; control group, 65.9 years Sex: Beta-blocker group, 78.5% male; control group, 73.6% male Mean aneurysm size: not reported Prior myocardial infarction: Beta-blocker group, 15.0%; control group, 12.0% Angina: Beta-blocker group, 7.3%; control group, 14.8% Permanent pace maker: Beta-blocker group, 0.4%; control group, 0% Renal insufficiency: Beta-blocker group, 1.2%; control group, 2.8%

Full citation	Yang H, Raymer K, Butler R, Parlow J, Roberts R. (2006) The effects of perioperative beta-blockade: results of the Metoprolol after Vascular Surgery (MaVS) study, a randomized controlled trial. Am Heart J. 152(5):983-90.
Intervention	25 to 100 mg of metoprolol was administered orally or intravenously, 2 hours before and after surgery. Treatment was continued intravenously every 6 hours or orally twice a day for 5 days or until hospital discharge (whichever occurred sooner).
Comparison	Matched placebo
Outcomes measures	The primary outcome was the composite rate of cardiac death, myocardial infarction, congestive heart failure, unstable angina, dysrhythmia requiring treatment, and non-cardiac death at 6 month follow-up. Individual rates were also reported at 30-day follow-up. Secondary outcomes included the need for reoperation, cerebrovascular accidents, new or worsened renal insufficiency, rehospitalisation, and intraoperative adverse events.
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Unclear risk – Authors state that randomisation was constructed in block of 4 by the study statistician; however it is not clear how allocation sequences were generated. Allocation concealment (selection bias): Unclear risk – Insufficient information was provided in the manuscript to ascertain whether appropriate steps were taken to conceal group allocations Blinding of participants and personnel (performance bias): Low risk – Authors state that patients, investigators, and all caretakers were blinded to the study randomisation Blinding of outcome assessment (detection bias): Low risk – Authors state that patients, investigators, and all caretakers were blinded to the study randomisation Incomplete outcome data (attrition bias): Low risk – "Completion of study protocol was 77.6% and 75.2% in the placebo and treatment groups, respectively." All losses to follow-up were accounted for and equally balanced across the 2 groups. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported Other bias: – Unclear risk – Intraoperative use of esmolol was allowed if deemed absolutely necessary. However, it was not clear what proportions of patients in each group received esmolol. Overall risk of bias: Low

Exercise

Full citation	Barakat H M, Shahin Y, Khan J A et al. (2016) Preoperative supervised exercise improves outcomes after elective abdominal aortic aneurysm repair. Annals of Surgery 264, 47-53
Study details	Study type: randomised, non-blinded trial Location(s): UK Aim(s): to assess the impact of a preoperative medically supervised exercise programme on postoperative outcomes of elective AAA repair Study dates: September 2009 to January 2014 Follow-up: 3 months Sources of funding: University of Hull (self-funded)
Participants	 Population: patients with AAAs undergoing EVAR or open surgical repair. Sample size: 124; 89.5% (111/124) male Inclusion criteria: patients older than 18 years with AAAs greater than 5.5 cm in diameter were included Exclusion criteria: thoracic aortic aneurysms, presence of factors that would limit exercise participation, patients requiring expedited or urgent aneurysm repair Baseline characteristics: Mean age: Exercise group, 73.8 years; control group, 72.9 years Sex: Exercise group, 90.3% male; control group, 88.7% male Mean aneurysm size: Exercise group 6.0 cm; control group, 6.3 cm Hypertension: Exercise group, 72.6%; control group, 69.4% Coronary artery disease: Exercise group, 38.7%; control group, 37.1% Hyperlipidaemia: Exercise group, 43.5%; control group, 40.3% Peripheral artery disease: Exercise group, 14.5%; control group, 12.9% Diabetes: Exercise group, 6.5%; control group, 37.1% COPD: Exercise group, 29.0%; control group, 37.1%
Intervention	Hospital based exercise classes: Patients attended 1 hour-long classes, 3 times a week. Exercises comprised a 5-minute warm up, using a cycle ergometer, heel- raise repetitions, knee extensions, dumbbells' biceps/arm curls, step-up lunges, knee bends (bodyweight), and 5 minutes for cool down and stretching.
Comparison	No exercise (controls)

Full citation	Barakat H M, Shahin Y, Khan J A et al. (2016) Preoperative supervised exercise improves outcomes after elective abdominal aortic aneurysm repair. Annals of Surgery 264, 47-53
Outcomes measures	The primary outcome was the composite rate of cardiac, pulmonary, and renal complications. Secondary outcomes included length of stay, APACHE II scores, occurrence of systematic inflammatory response syndrome, mortality, and bleeding requiring reoperation or transfusion.
Risk of bias assessment (using Cochrane risk of bias tool)	1. Random sequence generation (selection bias): Low risk – Randomisation was performed using a computer-generated sequence prepared by an independent professional
	 Allocation concealment (selection bias): Low risk – Randomisation was performed using opaque, sealed, identical envelopes containing the treatment allocation
	 Blinding of participants and personnel (performance bias): Low risk – It was not possible to blind participants but this was unlikely to bias results as objective outcomes were measured
	5. Blinding of outcome assessment (detection bias): Low risk – Clinicians including consultant surgeons, anaesthetists, medical staff and interventional radiologists were blinded to group allocations
	6. Incomplete outcome data (attrition bias): Low risk – There were no losses to follow-up.
	7. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported.
	8. Other bias: Low risk – none identified
	Overall risk of bias: Low
	Directness: directly applicable

Full citation	Dronkers J, Veldman A, Hoberg E et al. (2008) Prevention of pulmonary complications after upper abdominal surgery by preoperative intensive inspiratory muscle training: a randomized controlled pilot study. Clinical rehabilitation 22, 134-42
Study details	Study type: randomised, single-blind trial Location(s): Netherlands Aim(s): to investigate the effects of preoperative inspiratory muscle training on the incidence of atelectasis in patients at high risk of pulmonary complications scheduled for elective AAA surgery Study dates: not reported Follow-up: 7 days Sources of funding: not reported
Participants	 Population: patients with AAAs undergoing elective surgical repair (not specified) who were considered to have a high risk of pulmonary complications. Sample size: 20; 20% (5/15) male Inclusion criteria: patients who were due to undergo AAA surgical repair, with a scheduled delay of at least 2 weeks, and at least 1 of the following risk factors were included: age over 65 years, smoking within 2 months before surgery, presence of COPD, and a BMI greater than 27 were included Exclusion criteria: cerebrovascular disorders, neuromuscular diseases, a history of lung surgery, cardiovascular instability, receiving immunosuppressive treatment within 30 days of surgery, or treatment by a physical therapist within 8 weeks of surgery Baseline characteristics: Mean age: Exercise group, 70 years; control group, 59 years Sex: Exercise group, 80% male; control group, 70% male Mean aneurysm size: not reported COPD: Exercise group, 10%; control group, 10%
Intervention	Inspiratory muscle training: Patients took part in a training programme involving one 15-minute exercise session, 6 days a week, for at least 2 weeks prior to surgery. One session per week was supervised by the same physical therapist and the other 5 sessions were unsupervised.
Comparison	No exercise
Outcomes measures	Outcomes included incidence of atelectasis, patient satisfaction, and respiratory function.
Risk of bias assessment (using	 Random sequence generation (selection bias): Unclear risk – Authors state that an independent research assistant randomly assigned patients to treatment groups. No further information was provided. Allocation concealment (selection bias): Low risk – Group allocations were concealed using sealed and numbered envelopes.

Full citation	Dronkers J, Veldman A, Hoberg E et al. (2008) Prevention of pulmonary complications after upper abdominal surgery by preoperative intensive inspiratory muscle training: a randomized controlled pilot study. Clinical rehabilitation 22, 134-42
Cochrane risk of bias tool)	3. Blinding of participants and personnel (performance bias): Low risk – It was not possible to blind participants but this was unlikely to bias results as objective outcomes were measured.
	4. Blinding of outcome assessment (detection bias): Low risk – Assessment of the primary outcome (atelectasis) was performed by radiologists who were blinded to treatment outcomes.
	5. Incomplete outcome data (attrition bias): Low risk – Authors presented results based using an intention-to treat approach and presented final follow up results. All participants were accounted for.
	6. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported.
	7. Other bias: Low risk – none identified.
	Overall risk of bias: Low
	Directness: directly applicable

Full citation	Tew GA, Batterham AM, Colling K, et al. (2017) Randomized feasibility trial of high-intensity interval training before elective abdominal aortic aneurysm repair. The British journal of surgery 104(13), 1791-1801
Study details	Study type: randomised, single-blind trial Location(s): UK
	Aim(s): to assess the feasibility of a preoperative high-intensity interval training (HIT) programme in patients awaiting elective abdominal aortic aneurysm repair
	Study dates: not reported Follow-up: 12 weeks
	Sources of funding: This study was funded by the National Institute for Health Research under its Research for Patient Benefit Programme
Participants	Population: patients with unruptured AAAs undergoing elective EVAR or open surgical repair Sample size: 53; 94.3% (50/53) male
	Inclusion criteria: patients > 18 years, with infrarenal AAAs 5.5 to 7.0 cm in diameter who were due to undergo AAA surgical repair open repair or EVAR were included
	Exclusion criteria: AAA managed non-operatively, not an infrarenal aneurysm (juxtarenal, suprarenal or thoracic), infrarenal AAA diameter exceeding 7.0 cm, emergency AAA repair, contraindication to exercise testing or training
	Baseline characteristics:
	Mean age: Exercise group, 74.6 years; control group, 74.9 years
	 Sex: Exercise group, 92.6% male; control group, 96.2% male Mean aneurysm size: Exercise group 6.0 cm; control group, 5.8 cm
	 Coronary artery disease: Exercise group, 40.7%; control group, 53.8%
	Cerebrovascular disease: Exercise group, 25.9%; control group, 26.9%
	Peripheral arterial disease: Exercise group, 0%; control group, 7.7%
	 Diabetes: Exercise group, 14.8%; control group, 7.7% COPD: Exercise group, 22.2%; control group, 26.9%
Intervention	HIT:
	Patients in the exercise group were invited to complete three hospital-based exercise sessions per week, for the 4 consecutive weeks immediately preceding their intended operation date
Comparison	No exercise
Outcomes measures	Adverse events, quality of life, and length of stay

Full citation	Tew GA, Batterham AM, Colling K, et al. (2017) Randomized feasibility trial of high-intensity interval training before elective abdominal aortic aneurysm repair. The British journal of surgery 104(13), 1791-1801
Risk of bias assessment (using Cochrane risk of bias tool)	1. Random sequence generation (selection bias): Low risk – Authors stated that participants were randomised to groups using minimastion. Minimisation was performed with a 1:1 allocation ratio and equal weighting for the three minimisation factors (sex, type of procedure and study centre).
	Allocation concealment (selection bias): Low risk – Allocation was concealed from those assessing eligibility and recruiting patients, with eligible patients allocated remotely via e-mail by the trial statistician.
	Blinding of participants and personnel (performance bias): Low risk – It was not possible to blind participants but this was unlikely to bias results as objective outcomes were measured.
	 Blinding of outcome assessment (detection bias): Low risk – Authors stated that tests were performed by 2 experienced investigators blinded to group allocations,
	5. Incomplete outcome data (attrition bias): Low risk – All losses to follow-up were reported and accounted for in a consort diagram.
	6. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported
	7. Other bias: Low risk – none identified
	Overall risk of bias: Low
	Directness: directly applicable

Remote ischaemic preconditioning

Full citation	Ali ZA, Callaghan CJ, Lim E et al. (2007) Remote ischemic preconditioning reduces myocardial and renal injury after elective abdominal aortic aneurysm repair: a randomized controlled trial. Circulation 116, I98-105
Study details	Study type: randomised, double-blind trial Location(s): UK Aim(s): to investigate the potential of RIPC on myocardial and renal protection after elective open AAA repair Study dates: February 2003 and December 2005 Follow-up: 7 days Sources of funding: Cambridge University Hospitals NHS Foundation Trust
Participants	 Population: patients with AAAs undergoing elective open surgical repair Sample size: 82; 93% (76/82) male Inclusion criteria: patients referred for primary elective open AAA repair were included. No additional information was provided Exclusion criteria: over 90 years of age, needed concomitant procedures other than AAA repair, history of an acute coronary syndrome or myocardial infraction within 3 months, or taking sulfonylurea oral hypoglycaemic agents or nicorandil drug therapy Baseline characteristics: Mean age: RIPC group, 74 years; control group, 75 years Sex: RIPC group, 93% male; control group, 93% male Mean aneurysm size: not reported History of angina: RIPC group, 24%; control group, 27% History of diabetes: RIPC group, 5%; control group, 5% History of hypercholesterolaemia: RIPC group, 5%; control group, 5%
Intervention	Lower limb RIPC: This involved sequential cross-clamping of the common iliac arteries with 10 minutes ischaemia, followed by 10 minutes of reperfusion (RIPC stimulus). In order to reduce the risk of trash foot, sequential cross-clamping was performed to minimise repeat clamping of a single iliac artery. To prevent prolonged operating times, surgeons used a standardised approach whereby the iliac vessels were dissected before the neck of the aneurysm. The right iliac vessel was cross-clamped for 10 minutes followed by reperfusion during which time the left iliac was prepared. The cross-clamp was then placed to the left iliac vessel for 10 minutes and subsequently released, providing a total of 20 minutes of lower limb ischemia. During this time, the remainder of the operative dissection was carried out until the surgeon was prepared to cross-clamp the aorta before opening the aneurysm sac.
Comparison	Conventional open surgical repair without RIPC

Full citation	Ali ZA, Callaghan CJ, Lim E et al. (2007) Remote ischemic preconditioning reduces myocardial and renal injury after elective abdominal aortic aneurysm repair: a randomized controlled trial. Circulation 116, I98-105
Outcomes measures	Outcomes included length of stay, mortality, myocardial injury, myocardial infarction, renal impairment, and adverse events.
Risk of bias assessment	1. Random sequence generation (selection bias): Low risk – Patients were randomized by a computer-generated list in randomly sequenced blocks of 5, 6, 8, or 12.
(using Cochrane risk	2. Allocation concealment (selection bias): Low risk – Treatment allocations were concealed using numbered, sealed, opaque envelopes.
of bias tool)	3. Blinding of participants and personnel (performance bias): Low risk – Patients and data collectors not present in the operating room were blinded of treatment allocations.
	4. Blinding of outcome assessment (detection bias): Low risk – Results were compared and analysed by 2 blinded groups of assessors, labelled A and B.
	5. Incomplete outcome data (attrition bias): Low risk – No losses to follow-up were reported and all participants were included in the analyses.
	6. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported appropriately.
	7. Other bias: Low risk – none identified
	Overall risk of bias: Low
	Directness: directly applicable

Full citation	Li C, Li YS, Xu M et al. (2013) Limb remote ischemic preconditioning for intestinal and pulmonary protection during elective open infrarenal abdominal aortic aneurysm repair: a randomized controlled trial. Anesthesiology 118, 842-52
Study details	Study type: randomised, double-blind trial
	Location(s): China Aim(s): to assess whether limb RIPC would reduce intestinal and pulmonary injuries in patients undergoing open surgical repair of infrarenal AAAs
	Study dates: January 2008 to June 2011
	Follow-up: 24 hours
	Sources of funding: Sun Yat-Sen University hospital (self-funded)
Participants	Population: patients with AAAs undergoing elective open surgical repair.
	Sample size: 62; 90.1% (55/61) male
	Inclusion criteria: patients less than 80 years who were due to receive open surgical repair. No additional information was provided
	Exclusion criteria: infarction within 3 months, angina pain within 48 hours of surgery, ejection fraction less than 40%, poor pulmonary function (PaO2 < 60 mmHg), COPD, history of inflammatory bowel disease, history of diarrhoea within 1 week of surgery, or intestinal chronic inflammatory disease
	Baseline characteristics:
	Mean age: RIPC group, 62 years; control group, 67 years
	Sex: RIPC group, 93% male; control group, 84% male
	 Mean aneurysm size: RIPC group, 72 mm; control group, 69 mm Hypertension: RIPC group, 77%; control group, 58%
	 Diabetes: RIPC group, 45%; control group, 29%
	Previous myocardial infarction: RIPC group, 16%; control group, 26%
Intervention	Upper limb RIPC:
	A blood pressure cuff was placed on the left upper arm and 3 inflating–deflating cycles were performed. Each cycle consisted of 5 minutes of inflation to 200 mmHg followed by 5 minutes of reperfusion by deflating the cuff. All procedures were consistently performed by the same surgeon.
Comparison	Sham RIPC: an uninflated cuff was placed on the left upper arm for 30 min.
Outcomes measures	The primary outcomes were haemodynamic data and variables reflecting lung function. Secondary outcomes included mortality, ventilator support time, ICU- and hospital-free days; new arrhythmia, perioperative myocardial infarction, diagnosis of congestive heart failure, symptoms and signs of pulmonary congestion, neurologic events, upper limb ischemia requiring intervention,

Full citation	Li C, Li YS, Xu M et al. (2013) Limb remote ischemic preconditioning for intestinal and pulmonary protection during elective open infrarenal abdominal aortic aneurysm repair: a randomized controlled trial. Anesthesiology 118, 842-52
	intestinal injury markers, markers of oxidative stress and systemic inflammatory response, and scores of the severity of intestinal and pulmonary injury.
Risk of bias assessment	1. Random sequence generation (selection bias): Low risk – randomisation was performed by an independent person using a computer random number generator with a 1:1 allocation using blocks of varying sizes.
(using Cochrane risk of bias)	 Allocation concealment (selection bias): Low risk – Allocation details were stored in numbered, sealed, and opaque envelopes. Treatment allocation was revealed by anaesthetists opening the envelope on the morning of surgery.
	 Blinding of participants and personnel (performance bias): Low risk – Patients, investigators, surgeons, critical care teams, and individuals participating in data analysis were all blinded to group allocations.
	4. Blinding of outcome assessment (detection bias): Low risk – as stated above.
	5. Incomplete outcome data (attrition bias): Low risk – No losses to follow-up were reported and all participants were included in the analyses.
	6. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported appropriately.
	7. Other bias: Low risk – none identified.
	Overall risk of bias: Low
	Directness: directly applicable

Full citation	Mouton R, Pollock J, Soar J et al. (2015) Remote ischaemic preconditioning versus sham procedure for abdominal aortic aneurysm repair: an external feasibility randomized controlled trial. Trials 16, 377
Study details	Study type: randomised, double-blind trial Location(s): UK Aim(s): to investigate whether RIPC could be successfully introduced in elective AAA repair Study dates: January 2010 to December 2012 Follow-up: 48 hours Sources of funding: the National Institute of Health Research and the North Bristol NHS Trust
Participants	 Population: patients with AAAs undergoing elective EVAR or open surgical repair. Sample size: 69; sex-specific proportions were not reported. Inclusion criteria: patients referred for a primary elective AAA repair (EVAR or open surgery) were included. No additional information was provided Exclusion criteria: patients taking sulphonylurea oral hypoglycaemic drugs or nicorandil were excluded Baseline characteristics: Mean age: RIPC group, 72 years; control group, 72 years Sex: not reported Mean aneurysm size: not reported Hypertension: RIPC group, 77%; control group, 71% Ischaemic heart disease: RIPC group, 18%; control group, 20% Congestive heart failure: RIPC group, 15%; control group, 3%
Intervention	Upper limb RIPC: A blood pressure cuff was placed on the upper arm (side not specified) and three 10-minute cycles of conditioning were performed. Each cycle consisted of 5 minutes of ischaemia (inflation of a blood pressure cuff to 40 mmHg above the patient's systolic blood pressure) followed by 5 minutes of reperfusion.
Comparison	Sham RIPC: a pressure cuff was inflated for the same periods as the RIPC intervention but only to 40 mmHg.
Outcomes measures	Outcomes included acute kidney injury scores as classified by the acute injury network (AKIN), mortality, myocardial infarction, new postoperative ECG changes, new arrhythmia, troponin T levels above 14 ng/L, and adverse events.
Risk of bias assessment (using	 Random sequence generation (selection bias): Low risk – Randomisation was performed with a 1:1 allocation, using computer-generated randomisation sequences of varying block sizes and stratified by type of surgery. Allocation concealment (selection bias): Low risk – Allocations were concealed and accessed via a secure password protected.

Full citation	Mouton R, Pollock J, Soar J et al. (2015) Remote ischaemic preconditioning versus sham procedure for abdominal aortic aneurysm repair: an external feasibility randomized controlled trial. Trials 16, 377
Cochrane risk of bias tool)	 website and were concealed until sufficient information to uniquely identify the individual had been entered. Blinding of participants and personnel (performance bias): Low risk – With the exception of the in-theatre anaesthetic team who administered the intervention, everyone (participants, surgeons, nursing staff and research nurses) was blinded to the intervention received. Blinding of outcome assessment (detection bias): Low risk – as stated above. Incomplete outcome data (attrition bias): Low risk – All loses to follow-up were adequately explained. Furthermore analyses were performed using an intention to treat approach. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported appropriately. Other bias: Low risk – none identified. Overall risk of bias: Low

Full citation	Murphy N, Vijayan A, Frohlich S et al. (2014) Remote ischemic preconditioning does not affect the incidence of acute kidney injury after elective abdominal aortic aneurysm repair. Journal of cardiothoracic and vascular anaesthesia 28, 1285-92
Study details	Study type: randomised, double-blind trial Location(s): UK Aim(s): to assess the effects of RIPC on renal outcome in patients with AAAs having elective open surgical repair. Study dates: September 2009 to December 2012 Follow-up: 3 days Sources of funding: not reported
Participants	Population: patients with AAAs undergoing elective open surgical repair Sample size: 62; 85.5% (53/62) male Inclusion criteria: adults with AAAs referred for primary elective open surgical repair were included. No additional information was provided
	 Exclusion criteria: myocardial infarction within 2 weeks of surgery, history of upper limb vascular insufficiency, kidney disease requiring renal replacement, or AAAs requiring emergency AAA repair Baseline characteristics: Median age: RIPC group, 75 years; control group, 69 years Sex: RIPC group, 94% male; control group, 77% male Mean aneurysm size: not reported Previous myocardial infarction: RIPC group, 22%; control group, 13% Angina: RIPC group, 13%; control group, 16% Hypertension: RIPC group, 64%; control group, 52% Hypercholesterolemia: RIPC group, 23%; control group, 16% Chronic kidney disease: RIPC group, 61%; control group, 55%
Intervention	Upper limb RIPC: A blood pressure cuff was placed on the upper arm (side not specified) and three 10-minute cycles of conditioning were performed. Each cycle consisted of 5 minutes of ischaemia (inflation of a blood pressure cuff to 100 mmHg above the patient's systolic blood pressure) followed by 5 minutes of reperfusion.
Comparison	Sham RIPC: method not specified
Outcomes measures	Outcomes included mortality, kidney injury (measured by creatinine levels and AKIN scores), myocardial infarction, and length of hospital stay.

Full citation	Murphy N, Vijayan A, Frohlich S et al. (2014) Remote ischemic preconditioning does not affect the incidence of acute kidney injury after elective abdominal aortic aneurysm repair. Journal of cardiothoracic and vascular anaesthesia 28, 1285-92
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Low risk – Patients were assigned randomly, using a random number computer generator in a 1:1 ratio for parallel arms Allocation concealment (selection bias): Unclear risk – Allocations were concealed using sealed envelopes. No further details were assigned
	 were provided. Blinding of participants and personnel (performance bias): Low risk – Authors do not explicitly state that participants were blinded to group allocations. However, the trial is described as a double-blind trial and it is unlikely that patients would have been aware what was being done to them while under general anaesthesia.
	 Blinding of outcome assessment (detection bias): Low risk – Study investigators, attending anaesthetists and surgical staff were blinded to treatment assignments.
	5. Incomplete outcome data (attrition bias): Low risk – Authors presented results based using an intention-to treat approach and presented final follow up results. All participants were accounted for.
	6. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported appropriately.
	 Other bias: Low risk – none identified. Overall risk of bias: Low
	Directness: directly applicable

Full citation	Walsh SR, Boyle JR, Tang TY et al. (2009) Remote ischemic preconditioning for renal and cardiac protection during endovascular aneurysm repair: a randomized controlled trial. Journal of endovascular therapy : an official journal of the International Society of Endovascular Specialists 16, 680-9
Study details	Study type: randomised, non-blinded trial Location(s): UK Aim(s): to determine whether RIPC reduces renal damage in patients with AAAs having elective open surgical repair Study dates: February 2006 to October 2007 Follow-up: 48 hours Sources of funding: The Mouton Charitable Foundation
Participants	 Population: patients with AAAs undergoing elective open surgical repair Sample size: 40 men Inclusion criteria: patients with AAAs and no history of acute renal failure, no history of renal replacement therapy, no previous renal transplant, no history of renal disease, serum creatinine values less than 1.5 mg/dL and a serum urea values less than 20 mmol/L were included Exclusion criteria: a history of previous EVAR, a history of a lower limb amputation, or patients scheduled to receive suprarenal aneurysm repairs Baseline characteristics: Mean age: RIPC group, 74 years; control group, 76 years Sex: 100% in both arms Mean aneurysm size: RIPC group, 60.7 mm; control group, 63.9 mm Diabetes: RIPC group, 17%; control group, 9% Previous myocardial infarction: RIPC group, 18% COPD: RIPC group, 17%; control group, 18% Hypertension: RIPC group, 44%; control group, 55%
Intervention	Lower limb RIPC: A cross-clamp was applied to the right common iliac artery for 10 minutes. Subsequently, the right iliac territory was reperfused and the clamp was applied to the left common iliac artery. Once each common iliac artery territory had undergone one 10-minute cycle of ischemia followed by 10 minutes of reperfusion, the aorta was cross-clamped and the aneurysm sac was opened.
Comparison	Conventional open surgical repair without RIPC

Full citation	Walsh SR, Boyle JR, Tang TY et al. (2009) Remote ischemic preconditioning for renal and cardiac protection during endovascular aneurysm repair: a randomized controlled trial. Journal of endovascular therapy : an official journal of the International Society of Endovascular Specialists 16, 680-9
Outcomes measures	The primary outcome measure was renal function (measured by urine output, urine retinal binding protein, and creatinine levels). Secondary outcomes included 30-day mortality, myocardial infarction, arrhythmia, congestive heart failure, pneumonia, renal failure, lower limb ischaemia requiring intervention, and postoperative length of stay.
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Low risk – Participants were randomised in blocks of 4 using computer- generated sequences. Allocation concealment (selection bias): Low risk – Group allocations were concealed with sealed, opaque, envelopes which were opened on the day of surgery Blinding of participants and personnel (performance bias): Unclear risk – It was unclear whether participants were blinded to treatment allocations. This was unlikely to bias results as objective outcomes were measured. Blinding of outcome assessment (detection bias): High risk – Outcome assessors were not blinded of treatment allocations Incomplete outcome data (attrition bias): Low risk – No losses to follow-up were reported in either treatment arm. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported Other bias: Low risk – none identified Overall risk of bias: Moderate Directness: directly applicable

Full citation	Walsh SR, Sadat U, Boyle JR et al. (2010) Remote ischemic preconditioning for renal protection during elective open infrarenal abdominal aortic aneurysm repair: randomized controlled trial. Vascular and endovascular surgery 44, 334-40
Study details	Study type: randomised, non-blinded trial Location(s): UK Aim(s): to determine whether RIPC reduces renal and cardiac damage in patients with AAAs having elective open surgical repair. Study dates: November 2006 to January 2008 Follow-up: 48 hours Sources of funding: The Mouton Charitable Foundation
Participants	 Population: patients with AAAs undergoing elective EVAR Sample size: 40; 85% (34/40) male Inclusion criteria: patients with AAAs and no history of acute renal failure, no history of renal replacement therapy, no previous renal transplant, no history of renal disease, serum creatinine values less than 1.5 mg/dL and a serum urea values less than 20 mmol/L were included Exclusion criteria: a history of previous EVAR, a history of a lower limb amputation, or patients scheduled to receive fenestrated or branched aneurysm repairs Baseline characteristics: Median age: RIPC group, 75 years; control group, 72 years Sex: RIPC group, 72% male; control group, 100% male Mean aneurysm size: RIPC group, 67.8 mm; control group, 77.4 mm Diabetes: RIPC group, 4.5%; control group, 18%; control group, 22% Angina: RIPC group, 4.5%; control group, 16% COPD: RIPC group, 4.5%; control group, 5.5% Hypertension: RIPC group, 54%; control group, 88%
Intervention	Lower limb RIPC: Ischaemia was induced by placing an inflatable tourniquet around the thigh and inflating it until there was no audible doppler signal in either pedal artery. After 10 minutes the cuff was deflated and the procedure was repeated on the other leg.
Comparison	Conventional open surgical repair without RIPC
Outcomes measures	The primary outcome measure was renal function (measured by urine output, urine retinal binding protein, and serum creatinine levels). Secondary outcomes included serum troponin levels and the incidence of major adverse cardiac events (cardiac arrest, cardiac death, cardiac failure, unstable angina, or myocardial infarction).

Full citation	Walsh SR, Sadat U, Boyle JR et al. (2010) Remote ischemic preconditioning for renal protection during elective open infrarenal abdominal aortic aneurysm repair: randomized controlled trial. Vascular and endovascular surgery 44, 334-40
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Unclear risk – Participants were randomised in blocks of 4 using computer- generated sequences. Allocation concealment (selection bias): Low risk – Group allocations were concealed with sealed, opaque, envelopes which were opened on the day of surgery. Blinding of participants and personnel (performance bias): Unclear risk – It was unclear whether participants were blinded to treatment allocations. This was unlikely to bias results as objective outcomes were measured. Blinding of outcome assessment (detection bias): High risk – Outcome assessors were not blinded of treatment allocations Incomplete outcome data (attrition bias): Low risk – Authors presented results based using an intention-to treat approach and presented final follow up results. All participants were accounted for. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported Other bias: Low risk – none identified Overall risk of bias: Moderate Directness: directly applicable

Evidence tables for review question 30 (postoperative interventions)

Doxycycline versus placebo

Full citation	Hackmann AE, Rubin BG, Sanchez LA et al. (2008) A randomized, placebo-controlled trial of doxycycline after endoluminal aneurysm repair. Journal of vascular surgery 48, 519-526
Study details	Study type: randomised, placebo-controlled, double-blind trial Location(s): USA Aim(s): to evaluate the effect of a MMP inhibitor, doxycycline, on EVAR Study dates: not reported Follow-up: 6 months Sources of funding: Barnes-Jewish Hospital Foundation, National Institutes for Health, Department of Veteran's Affairs, Flight Attendants Medical Research Institute, and the American Heart Association
Participants	 Population: patients with AAAs undergoing elective EVAR Sample size: 59; sex-specific proportions not reported Inclusion criteria: patients with AAAs less than 5.0 cm in diameter were included Exclusion criteria: not reported Baseline characteristics: Mean age: Doxycycline group, 68.9 years; control group, 74.0 years Sex: Doxycycline group, 80% male; control group, 79.2% male Mean aneurysm size: Doxycycline group; 57.2 mm; control group, 57.2 mm Hypertension: Doxycycline group, 90%; control group, 79.2% Coronary artery disease: Doxycycline group, 60%; control group, 45.8% Diabetes: Doxycycline group, 10%; control group, 12.5% Peripheral artery disease: Doxycycline group, 40%; control group, 29.2% COPD: Doxycycline group, 30%; control group, 41.7% Renal insufficiency: Doxycycline group, 10%; control group, 25%
Intervention	Doxycycline 100 mg b.i.d, starting from the day after surgery and continued for 6 months
Comparison	Matched placebo
Outcomes measures	Aneurysm diameter, graft migration, incidence of endoleak, adverse events
Risk of bias assessment	1. Random sequence generation (selection bias): Unclear risk – Authors stated that randomisation was performed in the pharmacy utilising a pre-assigned table of codes. No further details were provided.

Full citation	Hackmann AE, Rubin BG, Sanchez LA et al. (2008) A randomized, placebo-controlled trial of doxycycline after endoluminal aneurysm repair. Journal of vascular surgery 48, 519-526
(using	2. Allocation concealment (selection bias): Unclear risk – It is unclear whether treatment allocations were concealed.
Cochrane risk of bias tool)	 Blinding of participants and personnel (performance bias): Low risk – Participants were blinded to treatment allocations as both doxycycline and placebo tablets had similar packaging and coating.
	4. Blinding of outcome assessment (detection bias): Low risk – Data were collected from the CT scans, by individuals blinded as to treatment group
	 Incomplete outcome data (attrition bias): High risk – At final follow-up, 7 participants in the doxycycline group and 4 participants in the placebo group were either lost to follow-up or withdrew from the study.
	Selective reporting (reporting bias): High risk – Authors reported some outcome measures for the whole study population whereas other outcome measures were only reported for the intervention group; omitting results for the placebo group.
	 Other bias: High risk – Patients in the placebo group were significantly older than those in the doxycycline group. A higher proportion of patients in the doxycycline group were smokers.
	8. Overall risk of bias: High
	Directness: directly applicable

Physiotherapy plus walking exercises versus physiotherapy-alone

Full citation	Wnuk BR, Durmala J, Ziaja K et al. (2016) A Controlled Trial of the Efficacy of a Training Walking Program in Patients Recovering from Abdominal Aortic Aneurysm Surgery. Advances in clinical and experimental medicine : official organ Wroclaw Medical University 25, 1241-1371
Study details	Study type: randomised, single-blind trial Location(s): Poland Aim(s): to evaluate the impact of a physical training (backward walking) programme on patients after AAA surgery Study dates: not specified Follow-up: 2 years Sources of funding: not specified
Participants	 Population: patients with AAAs undergoing surgical repair (not specified) Sample size: 65 males Inclusion criteria: patients with unruptured, non-symptomatic AAAs, between 65 and 75 years, who had a stable cardiologic status, no neurological disorders, and no motor system impairment were included Exclusion criteria: patients with neurological disorders, unstable coronary heart disease, aortic dissection, psychiatric diseases, difficulty in locomotion, or medical contraindications were excluded Baseline characteristics: Mean age: Forward walking exercise group, 68 years; Forward walking exercise group, 70 years; control group, 69 years Sex: 100% male in all groups
Intervention	 Participants were divided into 2 intervention groups: Basic physiotherapy plus backward walking exercises Basic physiotherapy plus forward walking exercises Basic physiotherapy involved general conditioning exercises of low intensity. In addition to basic physiotherapy, participants in the intervention groups performed backward or forward walking exercises, conducted on an interval training cycle. The intensity (workload) of exercises were tailored to each patient by calculating "training heart rates".
Comparison	Basic physiotherapy-alone
Outcomes measures	6-minute walking test distance, walking speed, spirometry measurements (FVC, FEV1, FEV1/FVC and PEF), length of hospital stay
Risk of bias assessment (using	 Random sequence generation (selection bias): Low risk – randomisation was performed by drawing identical sealed envelopes which contained the number of the allocated group. Allocation concealment (selection bias): Low risk – It is unclear whether treatment allocations were concealed.

Full citation	Wnuk BR, Durmala J, Ziaja K et al. (2016) A Controlled Trial of the Efficacy of a Training Walking Program in Patients Recovering from Abdominal Aortic Aneurysm Surgery. Advances in clinical and experimental medicine : official organ Wroclaw Medical University 25, 1241-1371
Cochrane risk of bias tool)	3. Blinding of participants and personnel (performance bias): Low risk – It was not possible to blind participants but this was unlikely to bias results as objective outcomes were measured.
	4. Blinding of outcome assessment (detection bias): Low risk – Outcome assessors were blinded to treatment allocations.
	5. Incomplete outcome data (attrition bias): High risk – During the postoperative period, 17 participants were excluded from the study due to cardiac complications or disorders preventing their participation in exercise training.
	6. Selective reporting (reporting bias): Low risk – All pre-specified outcomes were reported.
	7. Other bias: High risk – It is unclear whether groups were similar at the start of the trial as limited demographic data was reported.
	Overall risk of bias: Moderate
	Directness: directly applicable

Appendix E – Forest plots

Forest plots for review question 11 (preoperative interventions)

Beta-blockers

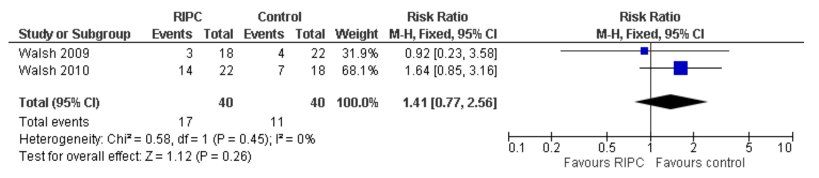
No meta-analysis was performed.

Exercise

No meta-analysis was performed.

RIPC versus sham RIPC or no RIPC (control)

RIPC versus control: any complications



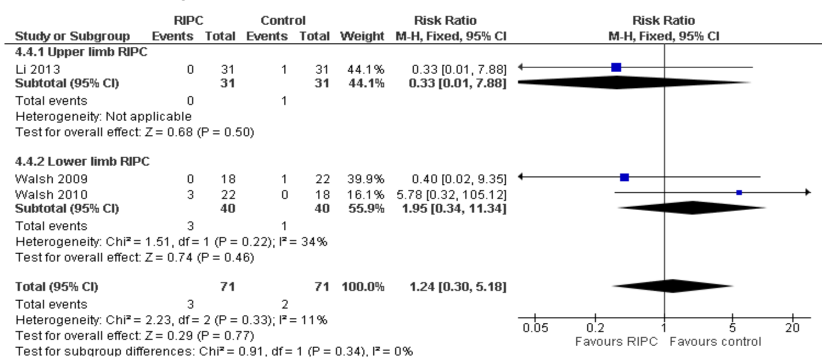
RIPC versus control: myocardial infarction

	RIPO	,	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
4.2.1 Upper limb RIPC							
Li 2013	2	31	1	31	5.6%	2.00 [0.19, 20.93]	
Mouton 2010	5	34	2	35	11.0%	2.57 [0.54, 12.38]	
Murphy 2014	4	31	2	31	11.1%	2.00 [0.39, 10.13]	
Subtotal (95% CI)		96		97	27.7%	2.23 [0.81, 6.15]	
Total events	11		5				
Heterogeneity: Chi ^z = I	0.06, df=	2 (P =	0.97); l² =	= 0%			
Test for overall effect: 2	Z = 1.55 (P = 0.1	2)				
4.2.2 Lower limb RIPC	2						
Ali 2007	2	41	11	41	61.2%	0.18 [0.04, 0.77]	←
Walsh 2009	1	18	1	22	5.0%	1.22 [0.08, 18.20]	
Walsh 2010	1	22	1	18	6.1%	0.82 [0.05, 12.19]	
Subtotal (95% Cl)		81		81	72.3%	0.31 [0.10, 0.90]	
Total events	4		13				
Heterogeneity: Chi ² = 3	2.02, df=	2 (P =	0.36); I ^z =	= 1%			
Test for overall effect: 2	Z = 2.15 (P = 0.0)3)				
Total (95% CI)		177		178	100.0%	0.84 [0.43, 1.62]	-
Total events	15		18				
Heterogeneity: Chi ² = 1	7.97, df=	5 (P =	0.16); I ^z =	= 37%			0.05 0.2 1 5 20
Test for overall effect: 2	Z = 0.52 (P = 0.6	60)				0.05 0.2 1 5 20 Favours RIPC Favours control
Test for subgroup diffe	erences: (Chi ≃ = I	6.87, df=	1 (P =	0.009), I ^z	= 85.5%	

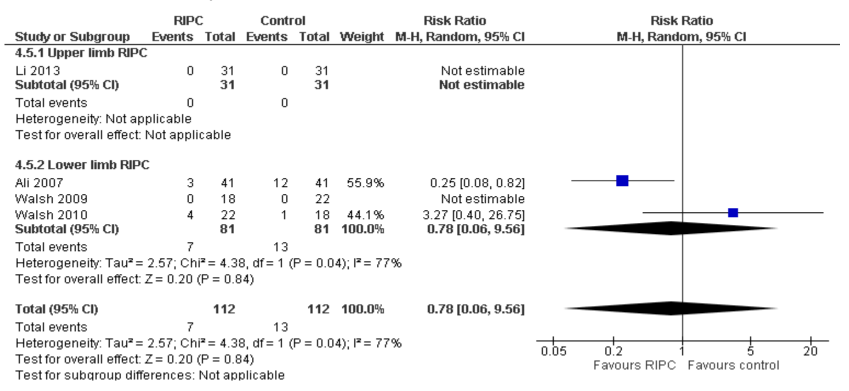
RIPC versus control: arrhythmia

	RIPO	2	Contr	ol		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M-H, Fixed, 95% Cl
4.3.1 Upper limb RIPC								
Li 2013	4	31	3	31	28.6%	1.33 [0.32, 5.47]		
Mouton 2010	7	34	5	35	47.0%	1.44 [0.51, 4.10]		
Murphy 2014 Subtotal (95% Cl)	5	31 96	1	31 97	9.5% 85.2 %	5.00 [0.62, 40.36] 1.80 [0.84, 3.87]		
Total events	16		9					
Heterogeneity: Chi ² =	1.27. df=	2 (P =	0.53); I ² =	= 0%				
Test for overall effect: .								
4.3.2 Lower limb RIPC	2							
Walsh 2009	3	18	0	22	4.3%	8.47 [0.47, 154.04]		
Walsh 2010	2	22	1	18	10.5%	1.64 [0.16, 16.62]		
Subtotal (95% CI)		40		40	14.8%	3.63 [0.66, 19.83]		
Total events	5		1					
Heterogeneity: Chi ² = I	0.78, df=	1 (P =	0.38); I ^z =	= 0%				
Test for overall effect: .	Z=1.49 ((P = 0.1	4)					
Total (95% CI)		136		137	100.0 %	2.07 [1.04, 4.15]		
Total events	21		10					
Heterogeneity: Chi ² = 3	2.47, df=	4 (P =	0.65); l² =	= 0%			0.05	0.2 1 5 20
Test for overall effect: 2	Z = 2.06 ((P = 0.0))4)				0.05	Favours RIPC Favours control
Test for subgroup diffe	erences:	Chi ² = I	0.54, df=	1 (P=	0.46), I ^z =	:0%		

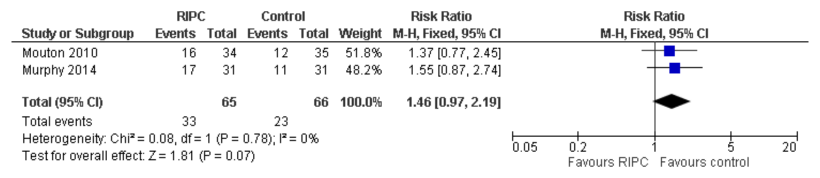
RIPC versus control: congestive heart failure



RIPC versus control: renal impairment or failure



RIPC versus control: acute kidney injury



RIPC versus control: 30-day mortality

	RIPO	С	Contr	ol		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	I M-H, Fixed, 95% Cl
Walsh 2009	1	18	0	22	45.2%	3.63 [0.16, 84.11]	
Walsh 2010	3	22	0	18	54.8%	5.78 [0.32, 105.12]	
Total (95% CI)		40		40	100.0%	4.81 [0.57, 40.68]	
Total events	4		0				
Heterogeneity: Chi² = Test for overall effect:	•			= 0%			0.02 0.1 1 10 50 Favours RIPC Favours control

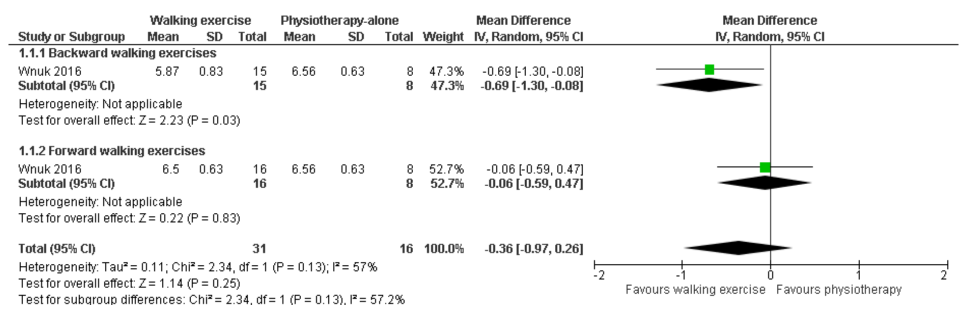
Forest plots for review question 30 (postoperative interventions)

Doxycycline versus placebo

Meta-analysis was not possible.

Physiotherapy plus walking exercises versus physiotherapy-alone

Length of hospital stay



Appendix F – GRADE tables

Grade tables for review question 11 (preoperative interventions)

Beta-blockers versus placebo

Intraoperative complications

		Quality	assessment	No of pa	atients	Effect estimate	Quality					
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results				
Hypotension rec	Hypotension requiring treatment; effect sizes below 1 favour beta blockers											
Yang (2006)	RCT	Not serious	Not serious	N/A	Serious ¹	250	246	RR 1.37 (1.10, 1.71)	Moderate			
Bradycardia req	uiring treat	tment; effect sizes	below 1 favour b	eta blockers								
Yang (2006)	RCT	Not serious	Not serious	N/A	Not serious	250	246	RR 2.81 (1.72, 4.61)	High			
						()))))))))))))))))))						

1. Confidence interval crosses one line of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 1 level.

Postoperative complications

		Quality	assessment	No of pa	tients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results			
Unstable angina; effect sizes below 1 favour beta blockers											
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 3.02 (0.12, 73.88)	Low		
Myocardial infar	Myocardial infarction; effect sizes below 1 favour beta blockers										
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 0.91 (0.50, 1.65)	Low		
Congestive hear	rt failure at	30 days; effect siz	es below 1 favou	ır beta blockers							
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 1.68 (0.41, 6.95)	Low		
Postoperative cerebrovascular accident ; effect sizes below 1 favour beta blockers											
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 1.26 (0.34, 4.64)	Low		
Dysrhythmia; ef	Dysrhythmia; effect sizes below 1 favour beta blockers										

		Quality	assessment	No of pa	itients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results			
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 0.71 (0.27, 1.38)	Low		
New or worsene	ed renal ins	ufficiency; effect s	izes below 1 fav	our beta blockers							
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 1.41 (0.45, 4.39)	Low		
Need for reoper	Need for reoperation; effect sizes below 1 favour beta blockers										
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 1.15 (0.42, 3.13)	Low		
Need for reoperation; effect sizes below 1 favour beta blockers											

1. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.

30-day mortality

		Quality	assessment	No of pa	tients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results			
All-cause mortality; effect sizes below 1 favour beta blockers											
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 0.20 (0.02, 1.69)	Low		
Cardiac death; e	effect sizes	below 1 favour bet	ta blockers								
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 0.33 (0.01, 8.08)	Low		
Non-cardiac death; effect sizes below 1 favour beta blockers											
Yang (2006)	RCT	Not serious	Not serious	N/A	Very serious ¹	250	246	RR 0.14 (0.01, 2.73)	Low		
1. Confidence i	1. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.										

Exercises

Inspiratory muscle training versus no training: postoperative complications

		Quality a	ssessment	No of pa	tients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Intervention	Control	Summary of results						
Atelectasis; effect sizes below 1 favour exercise group											
Dronkers (2008) RCT Serious ¹ Not serious N/A Serious ² 10 10 RR 0.38 (0.14, 1.02) L											
1. Very small sample size, downgrade 1 level.											

Confidence interval crosses one line of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 1 level

Physical exercise versus no exercise: preoperative intervention-related adverse events

		Quality as	ssessment	No of pa	tients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results			
Dizziness; effect sizes below 1 favour exercise group											
Tew (2016)	RCT	Not serious	Not serious	N/A	Very serious ¹	27	26	RR 2.89 (0.12, 67.69)	Low		
Angina; effect size	s below 1 fa	vour exercise grou	ıp								
Tew (2016)	RCT	Not serious	Not serious	N/A	Very serious ¹	27	26	RR 2.89 (0.12, 67.69)	Low		
1. Confidence interval crosses 2 lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 level											

Physical exercise versus no exercise: postoperative complications

		Quality	assessment			No of pa	atients	Effect estimate	Quality			
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results				
Cardiac complie	Cardiac complications (including myocardial infarction, prolonged inotropic support, new onset arrhythmia, and unstable angina); effect sizes below 1 favour exercise group											
Barakat (2016)	RCT	Not serious	Not serious	N/A	Serious ¹	62	62	RR 0.36 (0.14, 0.93)	Moderate			
Pulmonary com	plications (including pneumo	nia, pneumonia r	equiring reintuba	tion, exacerbation	of COPD, and	reintubation)	; effect sizes below 1 favo	ur exercise group			
Barakat (2016)	RCT	Not serious	Not serious	N/A	Very serious ²	62	62	RR 0.54 (0.23, 1.26)	Low			
Renal complication	tions (inclu	ding acute renal fa	ilure and renal ir	nsufficiency); effe	ct sizes below 1 fa	vour exercise g	group					
Barakat (2016)	RCT	Not serious	Not serious	N/A	Serious ¹	62	62	RR 0.31 (0.11, 0.89)	Moderate			
Postoperative b	leeding or	need for a blood tr	ansfusion of mo	re than 4 units; eff	ect sizes below 1	favour exercise	e group					
Barakat (2016)	RCT	Not serious	Not serious	N/A	Very serious ²	62	62	RR 0.57 (0.18, 1.85)	Low			
Need for reoperation; effect sizes below 1 favour exercise group												
Barakat (2016)	RCT	Not serious	Not serious	N/A	Very serious ²	62	62	RR 0.67 (0.12, 3.84)	Low			
	 Confidence interval crosses one line of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 1 level. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels. 											

2. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.

Physical exercise versus no exercise: 30-day mortality

	Quality assessment						tients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
All-cause mortality; effect sizes below 1 favour exercise group									
Barakat (2016)	RCT	Not serious	Not serious	N/A	Very serious ²	62	62	RR 1.10 (0.15, 6.88)	Low
()	-	Not serious			,				Low

1. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.

Physical exercise versus no exercise: length of stay

		Quality a	ssessment			No of pa	itients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
Median length of s	tay								
Tew (2016)	RCT	Not serious	Not serious	N/A	Very serious ¹	27	26	Difference in medians: 1 day (Statistical significance not reported)	Low
1. Level of statistical significance not reported, downgrade 2 levels									

Physical exercise versus no exercise: quality of life

		Quality as	ssessment			No of pa	tients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
SF-36 Physical fun	F-36 Physical function subscale scores at 12 weeks; effect sizes below 0 favour exercise group								
Tew (2016)	RCT	Not serious	Not serious	N/A	Serious ¹	27	26	MD -0.3 (-2.7, 2.1)	Moderate
SF-36 mental healt	h subscale :	scores at 12 weeks	; effect sizes bel	ow 0 favour exerc	ise group				
Tew (2016)	RCT	Not serious	Not serious	N/A	Serious ¹	27	26	MD -0.5 (-3.3, 2.3)	Moderate
1. Non-significant	I. Non-significant result, downgrade 1 level.								

RIPC

Postoperative complications

		Quality	assessment			No of pa	tients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
Any complication	ons; effect s	sizes below 1 favo	ur RIPC						
2 studies	RCT	Serious ¹	Not serious	Serious ²	Very serious ³	40	38	RR 1.41 (0.77, 2.56)	Very low
Myocardial infa	rction; effe	ct sizes below 1 fa	our RIPC						
6 studies	RCT	Not serious	Not serious	Very serious ^{2,3}	Very serious ⁴	177	178	RR 0.84 (0.43, 1.62)	Very low
Arrhythmia; effe	ect sizes be	low 1 favour RIPC							
5 studies	RCT	Not serious	Not serious	Serious ²	Serious ⁵	136	137	RR 2.07 (1.04, 4.15)	Low
Congestive hea	rt failure; e	ffect sizes below 1	favour RIPC						
3 studies	RCT	Serious ¹	Not serious	Serious ²	Very serious ⁴	71	71	RR 1.24 (0.30, 5.18)	Very low
Renal impairme	nt or failure	; effect sizes belo	w 1 favour RIPC						
4 studies	RCT	Serious ¹	Not serious	Very serious ^{2,6}	Very serious ⁴	71	71	RR 0.78 (0.06, 9.56)	Very low
Acute kidney in	jury; effect	sizes below 1 favo	our RIPC						
2 studies	RCT	Not serious	Not serious	Serious ²	Serious ⁵	146	147	RR 1.46 (0.97, 2.19)	Low
1. Outcome as	sessors wer	e not blinded of trea	tment allocations.	downgrade 1 level	Ι.				

Outcome assessors were not blinded of treatment allocations, downgrade 1 level.
 Different surgical techniques (EVAR or open surgical repair) were used across included studies, downgrade 1 level.

3. I² between 33% and 66.7%, downgrade 1 level.

Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.
 Confidence interval crosses one line of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 1 levels.

6. I² between >66.7%, downgrade 2 levels.

Mortalitv

	Quality assessment						tients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
30-day mortality	; effect siz	es below 1 favour F	RIPC						
2 studies	RCT	Serious ¹	Not serious	Serious ²	Very serious ³	40	40	RR 4.81 (0.57, 40.68)	Very low
1 Outeene ee	1 Outcome appagars were not blinded of treatment allocations, devingrade 1 lovel								

1. Outcome assessors were not blinded of treatment allocations, downgrade 1 level.

2. Different surgical techniques (EVAR or open surgical repair) were used across included studies, downgrade 1 level.

	Quality assessment						atients	Effect estimate	Quality
No of studies	No of studies Design Risk of bias Indirectness Inconsistency Imprecision					Intervention	Control	Summary of results	
3. Confide	3. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.								

Grade tables for review question 30 (postoperative interventions)

Doxycycline versus placebo

		Quality	assessment			No of pa	itients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
Mean percentag	Mean percentage change in aneurysm diameter								
Hackmann (2008)	RCT	Very serious ¹	Not serious	N/A	Very serious ²	12	15	Non-significant (MD not reported)	Very low
Presence of end	loleak								
Hackmann (2008)	RCT	Very serious ¹	Not serious	N/A	Very serious ²	20	24	Non-significant (RR not reported)	Very low
Occurrence of g	raft migrat	ion							
Hackmann (2008)	RCT	Very serious ¹	Not serious	N/A	Very serious ²	20	24	Non-significant (RR not reported)	Very low

1. Patients in the placebo group were significantly older than those in the doxycycline group. A higher proportion of patients in the doxycycline group were smokers. Finally, authors reported some outcome measures for the whole study population whereas other outcome measures were only reported for the intervention group; omitting results for the placebo group. Downgrade 2 levels.

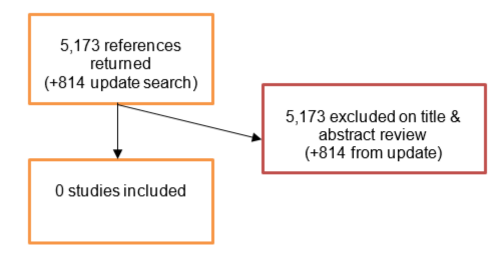
2. Risk ratio and measures of dispersion not reported. Downgrade 2 levels.

Physiotherapy plus walking exercises versus physiotherapy-alone

	Quality assessment							Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Intervention	Control	Summary of results	
Hospital length	Hospital length of stay (days); effect sizes below 0 favour exercise group								
Wnuk (2016)	RCT	Serious ¹	Not serious	Serious ²	Serious ³	31	16	MD -0.36 (-0.97, 0.26)	Very low

	Quality assessment						atients	Effect estimate	Quality
No of studies	No of studies Design Risk of bias Indirectness Inconsistency Imprecision						Control	Summary of results	
excluded fro 2. I ² value betw	om the stud veen 33.39	vention arms were dy during the posta % and 66.7%, dow Downgrade 1 leve	operative perioc ngrade 1 level.	due to cardiac co				ses to follow-up: 17 part	icipants were

Appendix G – Economic evidence study selection



Review questions 11 and 30 study selection

Appendix H – Excluded studies

Clinical studies

Rev	iew quest	ion 11	(preoperative	interventions)	

No.	Study	Reason for exclusion
1	Alreja G, Bugano D, and Lotfi A (2012) Effect of remote ischemic preconditioning on myocardial and renal injury: meta-analysis of randomized controlled trials. The Journal of invasive cardiology 24, 42-8	Meta-analysis assessed the efficacy of remote ischaemic preconditioning by pooling data from 3 studies of patients undergoing AAA repair and 2 studies of patients undergoing coronary artery bypass grafting. No subgroup or sensitivity analysis was performed.
2	Bani-Hani M, Titi M A, Jaradat I, and al- Khaffaf H (2008) Interventions for preventing venous thromboembolism following abdominal aortic surgery (Cochrane review) [with consumer summary]. Cochrane Database of Systematic Reviews 2008, andIssue 1,	The systematic review included studies of patients who underwent aortic surgery not related to AAAs: aortic bifurcation graft surgery and aortic reconstruction surgery.
3	Chello M, Mastroroberto P, Romano R et al. (1996) Protection by coenzyme Q10 of tissue reperfusion injury during abdominal aortic cross-clamping. The Journal of cardiovascular surgery 37, 229-35	Intervention (antioxidant supplement) is not outlined in the review protocol.
4	Desai M, Gurusamy K, Ghanbari H et al. (2011) Remote ischaemic preconditioning does not improve morbidity or mortality following open or endovascular aneurysm repair: A meta- analysis. Interactive cardiovascular and thoracic surgery 12, S139	Conference abstract.
5	Kertai M D, Boersma E, Westerhout C et al. (2004) A combination of statins and beta-blockers is independently associated with a reduction in the incidence of perioperative mortality and nonfatal myocardial infarction in patients undergoing abdominal aortic aneurysm surgery. European journal of vascular and endovascular surgery : the official journal of the European Society for Vascular Surgery 28, 343-52	Not a controlled trial. The study is a retrospective study which assessed the perioperative outcomes of patients with AAAs that had been taking statins and compared with those who had not been taking statins.
6	Kothmann E, Batterham A M, Owen S J et al. (2009) Effect of short-term exercise training on aerobic fitness in patients with abdominal aortic aneurysms: a pilot study. British Journal of Anaesthesia 2009 Oct, and103(4):505-510,	Participants in this study did not go on to receive surgery. As a result, this study does not assess whether exercise training is effective in optimising surgical outcomes in people undergoing surgical repair.
7	Hayashi K, Hirashiki A, Kodama A et al. (2016) Impact of preoperative regular	Not a controlled trial. This is a prospective cohort study which assessed whether

No.	Study	Reason for exclusion
	physical activity on postoperative course after open abdominal aortic aneurysm surgery. Heart and vessels 31, 578-83	patients preoperative physical activity levels affected postoperative outcomes of people undergoing AAA surgery.
8	Holzheimer R G (2003) Oral antibiotic prophylaxis can influence the inflammatory response in aortic aneurysm repair: results of a randomized clinical study. Journal of chemotherapy (Florence, and Italy) 15, 157-64	Outcome measure not of interest. Study assesses how oral antibiotic administration affects circulating inflammatory markers. No definitive outcomes were assessed.
9	Lo Sapio, P, Chechi T, Gensini GF et al. (2014) Impact of two different cardiac work-up strategies in patients undergoing abdominal aortic aneurysm repair. International journal of cardiology 175, e1-e3	Study is not directly relevant to this review question. This is a non-randomised comparative study comparing 2 algorithms for preoperative work-up: no comparisons were made with a control group (standard care).
10	McElrath M, Myers J, Chan K, et al. (2017) Exercise adherence in the elderly: Experience with abdominal aortic aneurysm simple treatment and prevention. Journal of vascular nursing : official publication of the Society for Peripheral Vascular Nursing 35(1), 12- 20	Participants in this study did not go on to receive surgery. As a result, this study does not assess whether exercise training is effective in optimising surgical outcomes in people undergoing surgical repair.
11	Mouton R, Pollock J, Soar J et al. (2014) Remote ischaemic preconditioning for elective abdominal aortic aneurysm (AAA) repair: a randomized controlled trial to assess feasibility. Applied cardiopulmonary pathophysiology 18, 35	Conference abstract.
12	Myers JN, White JJ, Narasimhan B et al. (2010) Effects of exercise training in patients with abdominal aortic aneurysm: preliminary results from a randomized trial. Journal of cardiopulmonary rehabilitation and prevention 30, 374-83	Participants in this study did not go on to receive surgery. As a result, this study does not assess whether exercise training is effective in optimising surgical outcomes in people undergoing surgical repair.
13	Myers J, McElrath M, Jaffe A et al. (2014) A randomized trial of exercise training in abdominal aortic aneurysm disease. Medicine and science in sports and exercise 46, 2-9	Participants in this study did not go on to receive surgery. As a result, this study does not assess whether exercise training is effective in optimising surgical outcomes in people undergoing surgical repair.
14	Pouwels S, Willigendael EM, van Sambeek M R H et al. (2015) Beneficial Effects of Pre-operative Exercise Therapy in Patients with an Abdominal Aortic Aneurysm: A Systematic Review. European journal of vascular and endovascular surgery : the official journal of the European Society for Vascular Surgery 49, 66-76	No quantitative synthesis was performed. Instead authors discussed the results of individual studies. Identified studies were assessed to ascertain their relevance to this NICE review question.

No.	Study	Reason for exclusion
15	Railton CJ, Wolpin J, Lam-McCulloch J et al. (2010) Renin-angiotensin blockade is associated with increased mortality after vascular surgery. Canadian journal of anaesthesia = Journal canadien d'anesthesie 57, 736-44	Not a controlled trial. The study is a cohort study which assessed outcomes of patients with preoperative renin-angiotensin system blockade, achieved either by angiotensin converting enzyme inhibitors or angiotensin receptor blocking agents.
16	Richardson K, Sanders G, Hayden P et al. (2014) The effect of preoperative exercise on postoperative outcome in abdominal aortic aneurysm (AAA) patients: Pilot study. Intensive care medicine 40, S136	Conference abstract.
17	Robertson L, Atallah E, and Stansby G (2017) Pharmacological treatment of vascular risk factors for reducing mortality and cardiovascular events in patients with abdominal aortic aneurysm. Cochrane Database of Systematic Reviews	Systematic review included one RCT which is already considered in this NICE review.
18	Tew G A, Moss J, Crank H et al. (2012) Endurance exercise training in patients with small abdominal aortic aneurysm: a randomised controlled pilot study. Archives of Physical Medicine and Rehabilitation 2012 Dec, and93(12):2148-2153,	Participants in this study did not go on to receive surgery. As a result, this study does not assess whether exercise training is effective in optimising surgical outcomes in people undergoing surgical repair.
19	Wijnen M, Vader HL, Van Den Wall Bake, A et al. (2002) Can renal dysfunction after infra-renal aortic aneurysm repair be modified by multi- antioxidant supplementation?. The Journal of cardiovascular surgery 43, 483-8	Intervention (antioxidant supplements) is not outlined in the review protocol.
20	Wijnen M, Roumen R, Vader HL, et al. (2002) A multiantioxidant supplementation reduces damage from ischaemia reperfusion in patients after lower torso ischaemia. A randomised trial. European journal of vascular and endovascular surgery : the official journal of the European Society for Vascular Surgery 23, 486-90	Intervention (antioxidant supplements) is not outlined in the review protocol.

Review question 30 (postoperative interventions)

No.	Study	Reason for exclusion
1	Abdul-Hussien H, Hanemaaijer R, Verheijen JH et al. (2009) Doxycycline therapy for abdominal aneurysm: Improved proteolytic balance through	Outcome measure not of interest. Study assesses how doxycycline affects aortic wall expression of the enzyme, matrix metalloproteinase.

No.	Study	Reason for exclusion
	reduced neutrophil content. Journal of vascular surgery 49, 741-9	
2	Aoki A, Suezawa T, Yamamoto S et al. (2014) Effect of antifibrinolytic therapy with tranexamic acid on abdominal aortic aneurysm shrinkage after endovascular repair. Journal of vascular surgery 59, 1203-8	Not a controlled trial. The study involved a retrospective review of medical records of patients treated before and after 5 institutions started administering tranexamic acid as part of their EVAR treatment protocols.
3	Boker A, Haberman CJ, Girling L et al. (2004) Variable ventilation improves perioperative lung function in patients undergoing abdominal aortic aneurysmectomy. Anesthesiology 100, 608-16	Perioperative intervention: study assessed the efficacy of variable ventilation delivered during surgery.
4	Brinkmann S J. H, Buijs N, Vermeulen M A et al. (2016) Perioperative glutamine supplementation restores disturbed renal arginine synthesis after open aortic surgery: A randomized controlled clinical trial. American Journal of Physiology - Renal Physiology 311, F567-f575	Outcome measure not of interest. Study assesses how perioperative glutamine administration affects arginine biosynthesis.
5	de Bruin JL, Baas AF, Heymans MW et al. (2014) Statin therapy is associated with improved survival after endovascular and open aneurysm repair. Journal of vascular surgery 59, 39-44.e1	Post-hoc analysis of an RCT comparing EVAR and open aneurysm repair. One of the secondary outcomes assessed was whether statin therapy reduced the risk of cardiovascular deaths. Unfortunately, it was not clear whether patients received statins before or after surgery.
6	Duffy MJ, O'Kane CM, Stevenson M et al. (2015) A randomized clinical trial of ascorbic acid in open abdominal aortic aneurysm repair. Intensive Care Medicine Experimental 3,	Perioperative intervention: study assessed the efficacy of parenteral ascorbic acid, administered during surgery.
7	Jones CI, Payne DA, Hayes PD et al.(2008) The antithrombotic effect of dextran-40 in man is due to enhanced fibrinolysis in vivo. Journal of vascular surgery 48, 715-22	Perioperative intervention: study assessed the efficacy of dextran-40, administered over 1 hour during surgery.
8	Kalimeris K, Nikolakopoulos N, Riga M et al. (2014) Mannitol and renal dysfunction after endovascular aortic aneurysm repair procedures: a randomized trial. Journal of cardiothoracic and vascular anesthesia 28, 954-9	Perioperative intervention: study assessed the efficacy of mannitol, administered within 15 minutes of surgery commencement.
9	Kertai MD, Boersma E, Westerhout CM et al. (2004) Association between long- term statin use and mortality after successful abdominal aortic aneurysm surgery. The American journal of medicine 116, 96-103	Not a controlled trial. The study is a retrospective study which assessed the outcomes of patients with AAAs that had been taking statins and compared with those who had not been taking statins.

No.	Study	Reason for exclusion
10	Leijdekkers VJ, Vahl AC, Mackaay A J et al. (2006) Aprotinin does not diminish blood loss in elective operations for infrarenal abdominal aneurysms: A randomized double-blind controlled trial. Annals of Vascular Surgery 20, 322-329	Perioperative intervention: study assessed the efficacy of aprotinin, administered during surgery.
11	Nicholson ML, Baker DM, Hopkinson BR et al. (1996) Randomized controlled trial of the effect of mannitol on renal reperfusion injury during aortic aneurysm surgery. The British journal of surgery 83, 1230-3	Perioperative intervention: study assessed the efficacy of mannitol, administered during surgery.
12	Rittoo D, Gosling P, Burnley S et al. (2004) Randomized study comparing the effects of hydroxyethyl starch solution with Gelofusine on pulmonary function in patients undergoing abdominal aortic aneurysm surgery. British journal of anaesthesia 92, 61-6	Perioperative intervention: study compared the efficacy hydroxyethyl starch solution with gelofusine, administered during surgery.
13	Smaka TJ, Cobas M, Velazquez OC et al. (2011) Perioperative management of endovascular abdominal aortic aneurysm repair: update 2010. Journal of cardiothoracic and vascular anesthesia 25, 166-76	Literature review.
14	Tisi PV, and Shearman CP (1997) Randomized controlled trial of the effect of mannitol on renal reperfusion injury during aortic aneurysm surgery. The British journal of surgery 84, 587	Letter to editor.
15	West MA, Parry M, Asher R et al. (2015) The Effect of beta-blockade on objectively measured physical fitness in patients with abdominal aortic aneurysmsA blinded interventional study. British journal of anaesthesia 114, 878-85	Study did not assess postoperative outcomes.

Economic studies

No full text papers were retrieved. All studies were excluded at review of titles and abstracts.

Appendix I – Research recommendations

Preoperative exercise programmes

Research recommendation	What is the clinical and cost-effectiveness of preoperative exercise programmes for improving outcomes of people who are having repair of an AAA?
Population	People with a confirmed unruptured abdominal aortic aneurysm in whom surgery is planned.
Intervention(s)	Exercise programmes incorporating physical exercise, preoperative physiotherapy or respiratory muscle training.
Comparator(s)	Each other, or no exercise
Outcomes	 Perioperative morbidity and mortality Incidence of postoperative complications (AAA rupture, AAA growth/expansion, cardiovascular events, wound-related complications, endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion) Need for further surgical intervention Mortality (all-cause; AAA-related; cardiovascular; survival) Cardiovascular events Quality of life Adverse effects Resource use and cost
Study design	Randomised controlled trial

Potential criterion	Explanation
Importance to patients, service users or the population	NHS providers have started devoting resources to exercise programmes, based on a relatively small body of evidence. Further research on the effectiveness of these programmes is needed to inform funding decisions.
Relevance to NICE guidance	Medium priority: no recommendations were made in this guideline due to limited evidence, and further research would allow for recommendations to be possible in future guideline updates.
Current evidence base	There is a growing body of evidence on preoperative exercise interventions for people undergoing various types of surgical procedures; however, the evidence relating to people with AAA was limited in quantity. Identified studies evaluating preoperative exercise interventions, in people with AAAs, were not considered robust enough to draft recommendations. The study evaluating the efficacy of inspiratory muscle training, by Dronkers et al. (2008), was considered low in quality as it had a small sample size (20 participants) and a short follow-up period. The study assessing the efficacy on supervised exercise, by Barakat et al (2016) was considered moderate in quality; however, reporting of composite outcomes, made it difficult to establish specific benefits (or harms) associated with the intervention.
Equality	No specific equality concerns are relevant to this research recommendation.
Feasibility	Postoperative growth and rupture is very rare, such that the committee suggested that it would require a very large RCT to detect an effect.

Research recommendation	What are the benefits of postoperative use of Direct Oral Anticoagulants (DOACS) for improving outcomes after repair of AAA?
Population	People who have undergone surgical repair of an abdominal aortic aneurysm.
Intervention(s)	 Apixaban Dabigatran Edoxaban Rivaroxaban Betrixaban
Comparator(s)	Each otherMatched placebo
Outcomes	 Incidence of postoperative complications (AAA rupture, AAA growth/expansion, cardiovascular events, wound-related complications, endoleak, graft migration, graft kinking, incisional hernia, graft occlusion, aortic neck expansion) Need for further surgical intervention Mortality (all-cause; AAA-related; cardiovascular; survival) Cardiovascular events Quality of life Adverse effects Resource use and cost
Study design	Randomised controlled trial

Postoperative use of Direct Oral Anticoagulants (DOACs)

Potential criterion	Explanation
Importance to patients, service users or the population	The committee recognised the risk of thromboembolic events (such as deep venous thrombosis and pulmonary embolism) following AAA surgery, and noted that postoperative anticoagulation, with or without the use of mechanical devices, can safely reduce the risk of such complications. DOACs are becoming increasingly popular because they are easy to use, have good pharmacokinetic properties associated with fixed dosing, have few interactions with other medications, and require less frequent monitoring. With that in mind, it is important to establish how best to use DOACs in the postoperative period to balance the risk thromboembolic events with that of bleeding.
Relevance to NICE guidance	Medium priority: no recommendations were made in this guideline due to the lack of evidence, and studies would allow for recommendations to be possible in future guideline updates.
Current evidence base	No studies were identified that specifically assessed the efficacy of postoperative use of DOACs for improving outcomes after repair of AAA.
Equality	No specific equality concerns are relevant to this research recommendation.
Feasibility	Postoperative growth and rupture is very rare, such that the committee suggested that it would require a very large RCT to detect an effect.

Appendix J – Glossary

Abdominal Aortic Aneurysm (AAA)

A localised bulge in the abdominal aorta (the major blood vessel that supplies blood to the lower half of the body including the abdomen, pelvis and lower limbs) caused by weakening of the aortic wall. It is defined as an aortic diameter greater than 3 cm or a diameter more than 50% larger than the normal width of a healthy aorta. The clinical relevance of AAA is that the condition may lead to a life-threatening rupture of the affected artery. Abdominal aortic aneurysms are generally characterised by their shape, size and cause:

- Infrarenal AAA: an aneurysm located in the lower segment of the abdominal aorta below the kidneys.
- Juxtarenal AAA: a type of infrarenal aneurysm that extends to, and sometimes, includes the lower margin of renal artery origins.
- Suprarenal AAA: an aneurysm involving the aorta below the diaphragm and above the renal arteries involving some or all of the visceral aortic segment and hence the origins of the renal, superior mesenteric, and celiac arteries, it may extend down to the aortic bifurcation.

Abdominal compartment syndrome

Abdominal compartment syndrome occurs when the pressure within the abdominal cavity increases above 20 mm Hg (intra-abdominal hypertension). In the context of a ruptured AAA this is due to the mass effect of a volume of blood within or behind the abdominal cavity. The increased abdominal pressure reduces blood flow to abdominal organs and impairs pulmonary, cardiovascular, renal, and gastro-intestinal function. This can cause multiple organ dysfunction and eventually lead to death.

Cardiopulmonary exercise testing

Cardiopulmonary Exercise Testing (CPET, sometimes also called CPX testing) is a noninvasive approach used to assess how the body performs before and during exercise. During CPET, the patient performs exercise on a stationary bicycle while breathing through a mouthpiece. Each breath is measured to assess the performance of the lungs and cardiovascular system. A heart tracing device (Electrocardiogram) will also record the hearts electrical activity before, during and after exercise.

Device migration

Migration can occur after device implantation when there is any movement or displacement of a stent-graft from its original position relative to the aorta or renal arteries. The risk of migration increases with time and can result in the loss of device fixation. Device migration may not need further treatment but should be monitored as it can lead to complications such as aneurysm rupture or endoleak.

Endoleak

An endoleak is the persistence of blood flow outside an endovascular stent - graft but within the aneurysm sac in which the graft is placed.

- Type I Perigraft (at the proximal or distal seal zones): This form of endoleak is caused by blood flowing into the aneurysm because of an incomplete or ineffective seal at either end of an endograft. The blood flow creates pressure within the sac and significantly increases the risk of sac enlargement and rupture. As a result, Type I endoleaks typically require urgent attention.
- Type II Retrograde or collateral (mesenteric, lumbar, renal accessory): These
 endoleaks are the most common type of endoleak. They occur when blood bleeds
 into the sac from small side branches of the aorta. They are generally considered
 benign because they are usually at low pressure and tend to resolve spontaneously
 over time without any need for intervention. Treatment of the endoleak is indicated if
 the aneurysm sac continues to expand.
- Type III Midgraft (fabric tear, graft dislocation, graft disintegration): These endoleaks occur when blood flows into the aneurysm sac through defects in the endograft (such as graft fractures, misaligned graft joints and holes in the graft fabric). Similarly to Type I endoleak, a Type III endoleak results in systemic blood pressure within the aneurysm sac that increases the risk of rupture. Therefore, Type III endoleaks typically require urgent attention.
- Type IV– Graft porosity: These endoleaks often occur soon after AAA repair and are associated with the porosity of certain graft materials. They are caused by blood flowing through the graft fabric into the aneurysm sac. They do not usually require treatment and tend to resolve within a few days of graft placement.
- Type V Endotension: A Type V endoleak is a phenomenon in which there is continued sac expansion without radiographic evidence of a leak site. It is a poorly understood abnormality. One theory that it is caused by pulsation of the graft wall, with transmission of the pulse wave through the aneurysm sac to the native aneurysm wall. Alternatively it may be due to intermittent leaks which are not apparent at imaging. It can be difficult to identify and treat any cause.

Endovascular aneurysm repair

Endovascular aneurysm repair (EVAR) is a technique that involves placing a stent –graft prosthesis within an aneurysm. The stent-graft is inserted through a small incision in the femoral artery in the groin, then delivered to the site of the aneurysm using catheters and guidewires and placed in position under X-ray guidance.

- Conventional EVAR refers to placement of an endovascular stent graft in an AAA where the anatomy of the aneurysm is such that the 'instructions for use' of that particular device are adhered to. Instructions for use define tolerances for AAA anatomy that the device manufacturer considers appropriate for that device. Common limitations on AAA anatomy are infrarenal neck length (usually >10mm), diameter (usually ≤30mm) and neck angle relative to the main body of the AAA
- Complex EVAR refers to a number of endovascular strategies that have been developed to address the challenges of aortic proximal neck fixation associated with complicated aneurysm anatomies like those seen in juxtarenal and suprarenal AAAs.

These strategies include using conventional infrarenal aortic stent grafts outside their 'instructions for use', using physician-modified endografts, utilisation of customised fenestrated endografts, and employing snorkel or chimney approaches with parallel covered stents.

Goal directed therapy

Goal directed therapy refers to a method of fluid administration that relies on minimally invasive cardiac output monitoring to tailor fluid administration to a maximal cardiac output or other reliable markers of cardiac function such as stroke volume variation or pulse pressure variation.

Post processing technique

For the purpose of this review, a post-processing technique refers to a software package that is used to augment imaging obtained from CT scans, (which are conventionally presented as axial images), to provide additional 2- or 3-dimensional imaging and data relating to an aneurysm's, size, position and anatomy.

Permissive hypotension

Permissive hypotension (also known as hypotensive resuscitation and restrictive volume resuscitation) is a method of fluid administration commonly used in people with haemorrhage after trauma. The basic principle of the technique is to maintain haemostasis (the stopping of blood flow) by keeping a person's blood pressure within a lower than normal range. In theory, a lower blood pressure means that blood loss will be slower, and more easily controlled by the pressure of internal self-tamponade and clot formation.

Remote ischemic preconditioning

Remote ischemic preconditioning is a procedure that aims to reduce damage (ischaemic injury) that may occur from a restriction in the blood supply to tissues during surgery. The technique aims to trigger the body's natural protective functions. It is sometimes performed before surgery and involves repeated, temporary cessation of blood flow to a limb to create ischemia (lack of oxygen and glucose) in the tissue. In theory, this "conditioning" activates physiological pathways that render the heart muscle resistant to subsequent prolonged periods of ischaemia.

Tranexamic acid

Tranexamic acid is an antifibrinolytic agent (medication that promotes blood clotting) that can be used to prevent, stop or reduce unwanted bleeding. It is often used to reduce the need for blood transfusion in adults having surgery, in trauma and in massive obstetric haemorrhage.