National Institute for Health and Care Excellence

Final

Abdominal aortic aneurysm: diagnosis and management

Evidence review L: Anaesthesia and analgesia for people having surgical repair of an abdominal aortic aneurysm

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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Anaesthesia and analgesia for people having surgical repair of unruptured and ruptured abdominal aortic aneurysm

Review questions

What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) endovascular repair (EVAR) and ii) open repair of an unruptured abdominal aortic aneurysm?

What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) EVAR and ii) open repair of a ruptured abdominal aortic aneurysm?

Introduction

Repair of abdominal aortic aneurysms (AAAs) is associated with a variety of risks, including bleeding, infection, nerve or spinal damage, as well as cardiovascular, respiratory, gastrointestinal, and renal complications. People undergoing AAA repair often have cardiovascular and respiratory comorbidities, which can increase the incidence and severity of the aforementioned risks. Optimising how anaesthesia and analgesia are used is an important part of minimising the incidence of complications. This review aims to assess the use of local, regional or general anaesthesia and different analgesic regimens in 'optimising' surgical outcome amongst people undergoing surgery for unruptured and ruptured AAA.

PICO table

Parameter	Inclusion criteria
Population	People undergoing surgery for a confirmed ruptured or unruptured AAA
Interventions	Regional or local anaesthesia and/or analgesia in the surgical repair of a ruptured or unruptured AAA
Comparators	General anaesthesia and/or analgesia in the surgical repair of a ruptured or unruptured AAA
Outcomes	Mortality Adverse events Complications of surgery, including pain, blood loss, wound complications, gut motility, and respiratory complications Need for additional intervention Successful exclusion of the aneurysm, aneurysm rupture, or further aneurysm growth Quality of life Resource use and costs

Table 1: Inclusion criteria

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 focused search strategy was used to pull in all studies that assessed the effectiveness of perioperatively administered local or regional anaesthesia and/or analgesia compared to general anaesthesia and/or analgesia in 'optimising' surgical outcome amongst people undergoing surgery for an AAA. Randomised, quasi-randomised and non-randomised controlled trials were considered for inclusion. Studies were excluded if they:

- were not in English;
- were not full reports of the study (for example, published only as an abstract);
- were not peer-reviewed;
- focused on postoperative anaesthesia and/or analgesia.

Prospective cohort studies were to be considered for inclusion if insufficient trial evidence was identified and if they had sample sizes larger than 500 and were conducted across multiple centres. Full details of the inclusion criteria are available in the review protocol in Appendix A.

Clinical evidence

Included studies

From an initial database of 2,201 abstracts, 116 full-text articles were ordered. Of these 7 studies conducted in people with unruptured AAA met inclusion criteria for this review, whereas no studies were identified relating to ruptured AAA.

An update search was conducted in December 2017, to identify any relevant studies published during guideline development. The search found 222 abstracts; all of which were not considered relevant to this review question. As a result no additional studies were included.

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

Table 2: Summary of included studies

Study	Details
Baron J-F, Bertrand M, Barré E, et	Study design: quasi-randomised controlled trial
al. (1991) Combined epidural and	Location: France
general anesthesia versus general	Population: high-risk surgical patients scheduled for
anesthesia for abdominal aortic	elective abdominal aortic reconstruction

Study	Details
surgery. Anesthesiology, 75: 611-	Sample size: 167
8	Follow-up: not reported
	Intervention: epidural anaesthesia plus general anaesthesia
	Comparator: balanced general anaesthesia
	Outcomes: mortality, cardiovascular morbidity, respiratory morbidity, renal failure, gastrointestinal bleeding, sepsis, major surgical complication & postoperative hospital stay
Broekema AAA, Kuizenga K, Hennis PJ (1996). Does epidural	Study design: double-blind randomised controlled trial Location: Netherlands
sufentanil provide effective analgesia per- and postoperatively for abdominal partia surgery? Acta	Population: people undergoing open surgical repair of unruptured AAA
Anaesthesiol Scandinavica 40	Sample size: 40
20-5	Follow-up: not reported
	Intervention: opioid epidural plus general anaesthesia
	Comparator: general anaesthesia plus placebo
	& need for additional analgesia
Davies MJ, Silbert BS, Mooney PJ	Study design: randomised controlled trial
and general anaesthesia versus	Location: Australia
general anaesthesia for abdominal aortic surgery: A	Population: people undergoing open surgical repair of unruptured AAA
prospective randomised trial.	Sample size: 50
Anaesthesia and Intensive Care	Follow-up: not reported
21: 790-4	anaesthesia
	Comparator: general anaesthesia-alone:
	complications, hepatic complications, renal complications, Length of hospital stay, length of ICU stay, intraoperative blood loss & infections
Davis. (1987) Intrathecal morphine in aortic aneurysm	Study design: randomised controlled trial
surgery. Anaesthesia 42: 491-7	Population: men undergoing open surgical repair of
	Sample size: 30
	Follow-up: not reported
	Intervention: intrathecal opioid plus general anaesthesia:
	Comparator: general anaesthesia-alone
	Outcomes: pain & clinical respiratory depression
Dodds TM, Burns K, DeRoo DB et al. (1997) Effects of anesthetic	Study design: double blind, randomised controlled trial Location: Lebanon
technique on myocardial wall motion abnormalities during	Population: people undergoing open surgical repair of unruptured AAA
abdominal aortic surgery. Journal	Sample size: 73
Anesthesia 11: 129-36	Follow-up: not reported

Study	Details
	Intervention: epidural plus general anaesthesia: Comparator: general anaesthesia-alone Outcomes: in-hospital mortality, cardiac morbidity, respiratory morbidity, renal insufficiency & blood loss
Fleron M-H, Weiskopf RB, Bertrand M et al. (2003) A comparison of intrathecal opioid and intravenous analgesia for the incidence of cardiovascular, respiratory, and renal complications after abdominal aortic surgery. Anesth Analg 97: 2-12	Study design: randomised controlled trial Location: France Population: people undergoing open surgical repair of unruptured AAA or aortoiliac occlusive disease Sample size: 217 Follow-up: not reported Intervention: Intrathecal opioid plus general anaesthesia Comparator: general anaesthesia-alone: Outcomes: major complications, cardiovascular complications, respiratory complications, renal complications & length of hospital stay
Norris EJ, Beattie C, Perler BA et al. (2001) Double-masked randomized trial comparing alternate combinations of intraoperative anesthesia and postoperative analgesia in abdominal aortic surgery. Anesthesiology 95: 1054-67	Study design: double-blind, randomised controlled trial Location: USA Population: patients undergoing open surgery to repair unruptured AAA or surgery for aortoiliac occlusive disease, and visceral and renal arterial reconstruction Sample size: 168 Follow-up: not reported Intervention: epidural anaesthesia combined with a light general anaesthesia Comparator: general anaesthesia plus placebo Outcomes: mortality, cardiac complications, respiratory complications, renal complications, intraoperative blood loss, reoperation & readmission to ICU

See Appendix D for full evidence tables.

Quality assessment of clinical studies included in the evidence review

See Appendix F for full GRADE tables.

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 review question 13 or review question 24.

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 question. As a result no additional studies were included.

Excluded studies

No studies were retrieved for full-text review.

Evidence statements

Use of anaesthesia and analgesia during repair of unruptured AAA general anaesthesia combined with an epidural compared with general anaesthesia alone during elective open repair

Mortality

 Very low-quality evidence from up to 3 RCTs, including up to 400 people undergoing elective open repair of an AAA, could not differentiate levels of inhospital mortality or 12-month mortality between people who received general anaesthesia combined with an epidural and those who received general anaesthesia alone.

Adverse events

- Very low-quality evidence from up to 4 RCTs, including up to 450 people undergoing elective open repair of an AAA, could not differentiate the postoperative incidence of myocardial infarction, congestive heart failure or general cardiovascular morbidity between people who received general anaesthesia combined with an epidural and those who received general anaesthesia-alone.
- Very low-quality evidence from up to 4 RCTs, including up to 450 people undergoing elective open repair of an AAA, could not differentiate the postoperative incidence of acute respiratory failure or pneumonia between people who received general anaesthesia combined with an epidural and those who received general anaesthesia alone.
- Very low-quality evidence from up to 4 RCTs, including up to 327 people undergoing elective open repair of an AAA, could not differentiate the postoperative incidence of renal failure or renal insufficiency between people who received general anaesthesia combined with an epidural and those who received general anaesthesia alone.

Surgical complications

 Very low-quality evidence from up to 5 RCTs, including up to 327 people undergoing elective open repair of an AAA, could not differentiate the levels of surgical complications between people who received general anaesthesia combined with an epidural and those who received general anaesthesia alone. Moderate quality evidence from 1 RCT, including 40 people undergoing elective open repair of an AAA, reported less need for additional analgesia in people who received general anaesthesia combined with an epidural compared with those who received general anaesthesia alone.

Need for reoperation

• Very low-quality evidence from 1 RCT, including 160 people undergoing elective open repair of an AAA, could not differentiate reoperation rates between people who received general anaesthesia combined with an epidural and those who received general anaesthesia alone.

Resource use

 Very low-quality evidence from up to 2 RCTs, including up to 217 people undergoing elective open repair of an AAA, could not differentiate the duration of postoperative hospital stay or postoperative stay in the intensive care unit between people who received general anaesthesia combined with an epidural and those who received general anaesthesia-alone.

General anaesthesia combined with intrathecal opioid compared with general anaesthesia alone during elective open repair

Mortality

• Very low-quality evidence from 1 RCT of 217 people undergoing elective open repair of an AAA could not differentiate levels of in-hospital mortality between people who received general anaesthesia combined with intrathecal opioid injection and those who received general anaesthesia alone.

Adverse events

- Very low quality evidence from 1 RCT of 217 people undergoing elective open repair of an AAA could not differentiate the postoperative incidence of myocardial infarction or congestive heart failure between people who received general anaesthesia combined with intrathecal opioid injection and those who received general anaesthesia alone.
- Very low quality evidence from up to 2 RCTs, including up to 242 people undergoing elective open repair of an AAA, could not differentiate the postoperative incidence of respiratory depression, acute respiratory failure or pneumonia between people who received general anaesthesia combined with intrathecal opioid injection and those who received general anaesthesia alone.

Anaesthesia and analgesia during elective EVAR

No evidence was identified relating to anaesthesia and/or analgesia during elective EVAR.

Use of anaesthesia and analgesia during repair of ruptured AAA

No evidence was identified relating to ruptured AAA.

The committee's discussion of the evidence

Interpreting the evidence

The outcomes that matter most

The guideline committee discussed the relative importance of a variety of outcomes and agreed that the following would be useful to their decision-making:

- Mortality
- The adverse events of anaesthesia or analgesia
- Additional surgical interventions or changes to the approach to anaesthesia and/or analgesia

The quality of the evidence

Use of anaesthesia and analgesia during repair of unruptured AAA

The committee noted that the evidence was limited to the comparison of general anaesthesia alone with general anaesthesia with an epidural or an intrathecal opioid. No evidence was found for other combinations of anaesthesia and analgesia. It was considered that some of the evidence may not have been generalisable to the UK context, primarily because of the formulations and doses of the interventions used. This affected the applicability of the evidence. This applicability was further affected by the populations in a number of the studies, which included people other than those undergoing open repair of an AAA, such as those undergoing surgery for aortoiliac occlusive disease, and those undergoing visceral or renal arterial reconstruction requiring abdominal aortic cross-clamping.

The committee noted that only 1 outcome in 1 comparison reached significance (the need for additional analgesia in the comparison of epidural plus general anaesthesia and general anaesthesia-alone), though this was likely a result of low event rates and small sample sizes. For this reason, the committee noted that there is an absence of evidence, not evidence of absence with regard to differences in the effects of the interventions and comparators studied.

No evidence was identified for anaesthesia and/or analgesia in people undergoing EVAR for unruptured AAA. The committee agreed that it was not necessary to draft consensus recommendations as they had recommended that EVAR should not be used to treat unruptured infrarenal aneurysms elsewhere in the guideline.

Use of anaesthesia and analgesia during repair of ruptured AAA

Since no evidence was identified for anaesthesia and/or analgesia in people undergoing any type of repair of ruptured AAA, the committee agreed that it was appropriate to draft consensus recommendations based on their collective skills, knowledge and experiences (discussed in the benefits and harms section below).

Benefits and harms Use of anaesthesia and analgesia during repair of unruptured AAA

On the whole, the identified evidence relating to elective open repair did not allow the committee to draw many distinctions between the use of general anaesthesia alone, general anaesthesia with an epidural, and general anaesthesia with an intrathecal injection of opioid. However, the committee noted that the addition of an epidural to general anaesthesia was associated with a lower need for additional analgesia compared with the use of general anaesthesia alone. This preference for the addition of an epidural to general anaesthesia was also supported by the committee's own clinical experience. The superior analgesic effect of adding an epidural has also been demonstrated and accepted in more general terms, such as in abdominal surgery more broadly, and the committee noted that there was no biological reason to expect that this result would be different in this population.

The committee subsequently discussed the possible populations in which the addition of an epidural to general anaesthesia would be contraindicated, but concluded that there were no such populations that could be specified. In the absence of explicit contraindications, possible reasons not to undertake an epidural might include possible side effects (including cardiac, respiratory, or gastrointestinal complications), the failure rate of epidurals, and the need for relatively intensive postoperative management to maximise benefits of an epidural. However, the committee did not feel that these concerns, when properly accounted for in the management of the patient, outweighed the possible benefits of using an epidural in conjunction with general anaesthesia in people undergoing open repair of an unruptured AAA.

The potential complications of epidural mean that some are now trying alternative methods, including the use of wound catheters to apply local anaesthesia, a technique that is being used more and more in abdominal surgery and which some are starting to use in the open repair of unruptured AAAs. However, the group did not feel that they had sufficient evidence or cause to explicitly recommend the use of wound catheters at this point.

No evidence was identified for optimal use of anaesthesia or analgesia in people undergoing elective EVAR. The committee noted that they recommended the procedure should not be performed in elective cases but acknowledged that in some circumstances, such as a hostile abdomen, EVAR may be warranted. In such situations no approach to anaesthesia and/or analgesia is considered superior to another. The committee agreed that it was important to tailor the approach to the individual patient, particularly in the case of people undergoing complex EVAR. They agreed that some important factors that should be considered include the 'ease' of the planned surgery, based upon the size, morphology and position of the aneurysm as well as the estimated duration of surgery, the patient's preference and concerns (for example, general anaesthesia may be preferable to patients who are anxious about being in the operating theatre). The committee agreed any recommendations on the use of anaesthesia or analgesia in people undergoing elective EVAR would be misleading as they had recommended that the procedure should not be performed in elective cases, elsewhere in the guideline.

Use of anaesthesia and analgesia during repair of ruptured AAA

No evidence was identified for the optimal management of anaesthesia and/or analgesia in people having open repair or EVAR of a ruptured AAA.

The committee agreed that the use of general anaesthesia alone is widely accepted as best practice when performing open surgical repair of a ruptured AAA. Furthermore, the committee agreed that the use of epidurals in addition to general anaesthesia is not considered safe or appropriate in the context of ruptured aneurysm. This is for a number of reasons; including a lack of sufficient time to administer an epidural when a patient is losing blood quickly, as well as the fact that people with ruptured AAA are generally not in a condition to tolerate administration of an epidural. The committee agreed that no recommendation was necessary as it is common practice to use anaesthesia alone during open surgery for ruptured AAA.

In the context of EVAR, the committee concluded that the approach of using anaesthesia and/or analgesia should be based primarily on the stability of the patient's condition. The committee felt that there was a lack of awareness among anaesthetists of the potential for effectively using local infiltrative anaesthesia alone in people undergoing EVAR for an AAA, at least at the start of the procedure. General anaesthesia can lead to loss of sympathetic control and muscle tone, which in patients with a ruptured aneurysm can lead to profound hypotension; for this reason, the use of local anaesthesia alone (at least initially) may be preferable.

Support for the use of local infiltrative anaesthesia alone in people undergoing EVAR for a ruptured AAA also came from a supplementary piece of evidence identified by the committee. A subgroup analysis of an included RCT for the question on EVAR versus open repair in ruptured AAA (IMPROVE) found that people who underwent EVAR for a ruptured AAA under local anaesthesia had a lower mortality (13%) than those who underwent the procedure under general anaesthesia (34%). This translated to a meaningful benefit for local anaesthesia (OR 0.25 (95% CI 0.10 to 0.70)), which the committee agreed may indicate a survival advantage associated with the use of local anaesthesia. However, the committee also acknowledged that this was a non-randomised comparison and there is no further evidence to support this. They also acknowledged that local infiltration alone may be distressing for the patient, or that it may not be feasible in all circumstances.

Cost effectiveness and resource use

No cost-effectiveness evidence was identified for this review area, and it was not prioritised for economic modelling.

Use of anaesthesia and analgesia during repair of unruptured AAA

The committee considered that the use of an epidural in addition to general anaesthesia during open surgical repair is already widespread practice, so recommending their use would have a limited impact on resource use.

Use of anaesthesia and analgesia during repair of ruptured AAA

The committee considered that the use of general anaesthesia alone in people undergoing open repair of a ruptured AAA is already widespread practice, so recommending its use will have a limited impact on resource use. The committee also agreed that the recommendation to consider local infiltrative anaesthesia alone for people having EVAR for ruptured AAA is unlikely to lead to any substantial change in resource use.

Other factors the committee took into account

No other factors were discussed by the committee.

Appendices

Appendix A – Review protocols

Review protocol for review question 13: Anaesthesia and analgesia for people having surgical repair of an unruptured abdominal aortic aneurysm

Review question 13	What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) EVAR and ii) open repair of an unruptured abdominal aortic aneurysm?
Objectives	To assess the use of local, regional or general analgesia and anaesthesia in 'optimising' surgical outcome amongst people undergoing surgery for an unruptured abdominal aortic aneurysm
Type of review	Intervention
Language	English only
Study design	Systematic reviews of study designs listed below: Randomised controlled trials Quasi-randomised controlled trials Non-randomised controlled trials If insufficient evidence identified, prospective cohort studies presenting comparative evidence will be considered (n >500; multicentre)
Status	Published papers only (full text) No date restrictions
Population	People undergoing surgery for a confirmed unruptured abdominal aortic aneurysm Subgroups: age, sex, comorbidities (including cardiovascular disease, renal disease, COPD, obesity); fitness/risk for surgery
Intervention	Regional or local anaesthesia and/or analgesia in the elective surgical repair of an unruptured abdominal aortic aneurysm
Comparator	General anaesthesia and/or analgesia in the elective surgical repair of an unruptured abdominal aortic aneurysm
Outcomes	Mortality Adverse events of anaesthesia or analgesia, including renal, pulmonary and cardiac Complications of surgery, including pain, blood loss, wound complications, gut motility, and respiratory complications Need for additional intervention: surgical, conversion from local/regional to general Successful exclusion of the aneurysm, aneurysm rupture, or further aneurysm growth Quality of life Resource use, including length of hospital or intensive care stay and readmissions, and costs

Review question 13	What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) EVAR and ii) open repair of an unruptured abdominal aortic aneurysm?
Other criteria for inclusion / exclusion of studies	Exclusion: Non-English language Abstract/non-published Pharmacological interventions not available in the UK Postoperative anaesthesia and/or analgesia
Baseline characteristics to be extracted in evidence tables	Age, Sex Size of aneurysm Position 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.
	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.

Review protocol for review question 24: Anaesthesia and analgesia for people having surgical repair of a ruptured abdominal aortic aneurysm

Review question 24	What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) EVAR and ii) open repair of a ruptured abdominal aortic aneurysm?
Objectives	To assess the use of local, regional or general analgesia and anaesthesia in 'optimising' surgical outcome amongst people undergoing surgery for a ruptured abdominal aortic aneurysm
Type of review	Intervention
Language	English only
Study design	Systematic reviews of study designs listed below: Randomised controlled trials Quasi-randomised controlled trials Non-randomised controlled trials If insufficient evidence identified, prospective cohort studies presenting comparative evidence will be considered (n >500; multicentre)
Status	Published papers only (full text) No date restrictions
Population	People undergoing surgery for a ruptured abdominal aortic aneurysm Subgroups: age, sex, comorbidities (including cardiovascular disease, renal disease, COPD, obesity)

Review question 24	What is the most effective approach to anaesthesia and/or analgesia in improving surgical outcome in people undergoing i) EVAR and ii) open repair of a ruptured abdominal aortic aneurysm?
Intervention	Regional or local anaesthesia and/or analgesia in the surgical repair of a ruptured abdominal aortic aneurysm
Comparator	General anaesthesia and analgesia in the surgical repair of a ruptured abdominal aortic aneurysm
Outcomes	Mortality Adverse events of anaesthesia or analgesia, including renal, pulmonary and cardiac Complications of surgery, including pain, blood loss, wound complications, gut motility, and respiratory complications Need for additional intervention: surgical, conversion from local/regional to general Successful exclusion of the aneurysm, aneurysm rupture, or further aneurysm growth Quality of life Resource use, including length of hospital or intensive care stay and readmissions, and costs
Other criteria for inclusion / exclusion of studies	Exclusion: Non-English language Abstract/non-published Pharmacological interventions not available in the UK Postoperative anaesthesia and/or analgesia
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. 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.

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 13 and 24

Medline Strategy, searched 11th February 2016
Database: Ovid MEDLINE(R) <1946 to January week 1 2016
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 supra renal* 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 exp Anesthesia/
- 7 (anaesthe* or anesthe*).tw.
- 8 exp Anesthetics/
- 9 Anesthesiology/
- 10 Nurse Anesthetists/
- 11 exp Analgesia/
- 12 Analgesi*.tw.

Medline Strategy, searched 11th February 2016 Database: Ovid MEDLINE(R) <1946 to January week 1 2016 Search Strategy:

- 13 Pain Management/
- 14 (Pain* adj4 (manag* or relie*)).tw.
- 15 or/6-14
- 16 5 and 15
- 17 Animals/ not humans/
- 18 16 not 17
- 19 limit 18 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.

Medline Strategy

- 21 (price* or pricing*).tw.
- 22 budget*.tw.
- 23 expenditure*.tw.
- 24 (value adj3 (money or monetary)).tw.
- 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 13 and 24 study selection



Appendix D – Clinical evidence tables

Full citation	Baron J-F, Bertrand M, Barré E, et al. (1991) Combined epidural and general anesthesia versus general anesthesia for abdominal aortic surgery. Anesthesiology 75: 611-8
Study details	Study type: Quasi-randomised controlled trial Location(s): France Aim(s): To determine whether intraoperative thoracic epidural anaesthesia in combination with light general anaesthesia alters postoperative morbidity compared to a standard technique of balanced general anaesthesia Study dates: not reported Follow-up: not reported Sources of funding: not reported
Participants	 Population: High-risk surgical patients scheduled for elective abdominal aortic reconstruction Sample size: 167 Inclusion criteria: patients undergoing elective open repair of AAA or aortoiliac occlusive disease were included. All participants had no contraindications for epidural anaesthesia (preoperative coagulopathy, localised infection or septicaemia and graft sepsis), a left ventricular ejection fraction greater than 35%; and an aortic surgical procedure performed via a midline xiphopubic skin incision. Exclusion criteria: not reported Baseline characteristics: Mean age: epidural plus general anaesthesia group, 62 years; general anaesthesia-alone group, 61 years Sex: epidural plus general anaesthesia group, 94.2% male; general anaesthesia-alone group, 86.4% male Mean aneurysm size: not reported Previous myocardial infarction: epidural plus general anaesthesia group, 15.1%; general anaesthesia-alone group, 17.3% ST-T abnormalities: epidural plus general anaesthesia group, 15.1%; general anaesthesia-alone group, 18.5% Rhythm other than sinus: epidural plus general anaesthesia group, 2.3%; general anaesthesia-alone group, 43.4%
Intervention	 Epidural anaesthesia plus general anaesthesia Intraoperative thoracic epidural anaesthesia in combination with light general anaesthesia: An epidural catheter was inserted via the T8-T9 interspace, and thoracic epidural anaesthesia was induced using an initial 10ml dose of a mixture of plain bupivacaine 0.5% and lidocaine 2%; if necessary, additional incremental doses to a total of up to 16ml were given until a thoracoabdomina sensitive blockade was induced General anaesthesia was induced using fentanyl (6 micrograms/kg), flunitrazepam (0.02 mg/kg) and pancuronium bromide (0.1 mg/kg);

Full citation	Baron J-F, Bertrand M, Barré E, et al. (1991) Combined epidural and general anesthesia versus general anesthesia for abdominal aortic surgery. Anesthesiology 75: 611-8
	 Anaesthesia was maintained under controlled ventilation (50% nitrous oxide in oxygen) by continuous epidural infusion (6-8ml/h) of the bupivacaine-lidocaine mixture described above;
	 When required, a low concentration of isoflurane was administered to maintain anaesthesia; this was increased to control arterial blood pressure during aortic cross clamping.
Comparison	Balanced general anaesthesia:
	 Induced using fentanyl (6 micrograms/kg), flunitrazepam (0.02 mg/kg) and pancuronium bromide (0.1 mg/kg);
	 Maintained under controlled ventilation (50% nitrous oxide in oxygen) by increments of fentanyl (approximately 1.5 micrograms/kg every 20 minutes) and pancuronium bromide;
	 When required, a low concentration of isoflurane was administered to maintain anaesthesia; this was increased to control arterial blood pressure during aortic cross clamping.
Outcomes measures	Mortality, cardiovascular morbidity, respiratory morbidity, renal failure, gastrointestinal bleeding, sepsis, major surgical complication, postoperative hospital stay
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Low risk – Randomisation performed using table of random numbers. Allocation concealment (selection bias): Unclear risk – it was not clear whether appropriate allocation concealment was performed Blinding of participants and personnel (performance bias): Low risk – Authors did not state whether blinding was performed; however this is unlikely to have affected study results as objective outcomes were measured. Blinding of outcome assessment (detection bias): Unclear risk – Authors did not state whether blinding was performed. Incomplete outcome data (attrition bias): Low risk – Although 6 patients from the epidural plus general anaesthesia group were excluded due to non-functioning epidural catheter and subsequent use of general anaesthesia, this was unlikely to have bias study results. Selective reporting (reporting bias): Low risk - All outcomes clearly defined. Other bias: High risk – Postoperative analgesia was not the same in each group (possible performance bias). Furthermore, the study population included some patients who were undergoing surgery for aortoiliac occlusive disease (44%), rather than
	AAA. Overall risk of bias: High
	Directness: partially applicable

Full citation	Broekema AAA, Kuizenga K, Hennis PJ (1996). Does epidural sufentanil provide effective analgesia per- and postoperatively for abdominal aortic surgery? Acta Anaesthesiol Scandinavica, 40: 20-5
Study details	Study type: Double-blind randomised controlled trial Location(s): Netherlands Aim(s): To assess the efficacy of epidural sufentanil in providing peri- and postoperative analgesia Study dates: Not reported Follow-up: Not reported Sources of funding: Not reported
Participants	 Population: people undergoing open surgical repair of unruptured AAA Sample size: 40 Inclusion criteria: people aged 20 to 80 years undergoing open surgical repair of unruptured AAA were included. All participants were categorised as ASA class I, II, or III Exclusion criteria: not reported Baseline characteristics: Mean age: opioid epidural plus general anaesthesia group, 63 years; general anaesthesia-alone group, 67 years Sex: opioid epidural plus general anaesthesia group, 90% male; general anaesthesia-alone group, 70% male Mean aneurysm size: not reported Comorbidities: not reported
Intervention	 Opioid epidural plus general anaesthesia: Intraoperative thoracic epidural of 50 micrograms sufentanil in 10 ml normal saline solution in combination with general anaesthesia Epidural injection of 50 micrograms sufentanil in 10 ml NaCl 0.9% General anaesthesia induced using intravenous midazolam 0.1-0.2 mg * kg-1, sufentanil 0.5 micrograms * kg-1 and vecuronium 0.1 mg * kg-1;
Comparison	 General anaesthesia-alone (Epidural placebo plus general anaesthesia): Intraoperative thoracic epidural of 10 ml normal saline solution in combination with general anaesthesia Epidural injection of 10 ml NaCl 0.9% General anaesthesia induced using intravenous midazolam, sufentanil and vecuronium; Maintained under controlled ventilation (60% nitrous oxide in oxygen and halothane at a 1% inspiratory concentration)
Outcomes measures	Complications, adverse events, blood loss, & need for additional analgesia
Risk of bias assessment (using	 Random sequence generation (selection bias): Unclear risk – Authors stated that randomisation was performed but the method was not reported. Allocation concealment (selection bias): Unclear risk – The approach to allocation concealment not described.

Full citation	Broekema AAA, Kuizenga K, Hennis PJ (1996). Does epidural sufentanil provide effective analgesia per- and postoperatively for abdominal aortic surgery? Acta Anaesthesiol Scandinavica, 40: 20-5
Cochrane risk of bias tool)	 Blinding of participants and personnel (performance bias): Low risk – Authors stated that the trial was double blind, No further details were provided. Blinding of outcome assessment (detection bias): Low risk – Authors stated that the trial was double blind, No further details
	 Were provided. Incomplete outcome data (attrition bias): Low risk – Few losses to follow-up were reported across treatment arms; reasons for follow-up were adequately reported. Selective reporting (reporting bias): Unclear risk Other bias: Low risk – none identified Overall risk of bias: Moderate
	Directness: directly applicable

Full citation	Davies MJ, Silbert BS, Mooney PJ, et al. (1993) Combined epidural and general anaesthesia versus general anaesthesia for abdominal aortic surgery: A prospective randomised trial. Anaesthesia and Intensive Care, 1993, 21: 790-4
Study details	Study type: Randomised controlled trial
	Location(s): Australia
	Aim(s): To examine the potential for combined epidural and general anaesthesia to reduce the incidence of respiratory and cardiovascular complications, decrease the duration of postoperative intensive care stay, and reduce the incidence of postoperative infections and complications.
	Study dates: not reported
	Follow-up: not reported
	Sources of funding: not reported
Participants	Population: people undergoing open surgical repair of unruptured AAA
	Sample size: 50
	Inclusion criteria: Patients undergoing open repair of unruptured AAA
	Exclusion criteria: contraindications to epidural anaesthesia (septicaemia, abnormal coagulation status, infection at the proposed puncture site, neurological disease)
	Baseline characteristics:
	 Mean age: epidural plus general anaesthesia group, 65 years; general anaesthesia-alone group, 67 years Sex: epidural plus general anaesthesia group, 84% male; general anaesthesia-alone group, 92% male Mean aneurysm size: not reported
	Angina: epidural plus general anaesthesia group, 12%; general anaesthesia-alone group, 20%
	Left ventricular failure: epidural plus general anaesthesia group, 4%; general anaesthesia-alone group, 4%
	Hypertension: epidural plus general anaesthesia group, 44%; general anaesthesia-alone group, 52%

Davies MJ, Silbert BS, Mooney PJ, et al. (1993) Combined epidural and general anaesthesia versus general anaesthesia for abdominal aortic surgery: A prospective randomised trial. Anaesthesia and Intensive Care, 1993, 21: 790-4
 Myocardial infarction: epidural plus general anaesthesia group, 20%; general anaesthesia-alone group, 24% COPD: epidural plus general anaesthesia group, 20%; general anaesthesia-alone group, 64%
Thoracic epidural combined with general anaesthesia:
 On arrival in the operating theatre, a 16-guage Tuohy needle was inserted into the epidural space of the lower thoracic spine (usually T9-10); an 18-guage epidural catheter was then inserted
 Following a 2ml test dose of lidocaine 1.5% with 1 in 200,000 adrenaline, a further 5ml was injected preoperatively into the epidural catheter; after this, 5ml was injected each hour intraoperatively
 General anaesthesia was induced by administering fentanyl 1-3 micrograms/kg and thiopental sodium 2-4mg/kg, and the trachea was intubated following pancuronium bromide 0.1mg/kg; the patients lungs were ventilated with 66% N20 in oxygen and eflurane
General anaesthesia-alone:
 General anaesthesia was induced by administering fentanyl 1-3 micrograms/kg and thiopental sodium 2-4mg/kg, and the trachea was intubated following pancuronium bromide 0.1mg/kg; the patients lungs were ventilated with 66% N20 in oxygen and eflurane
Mortality, cardiovascular complications, respiratory complications, hepatic complications, renal complications, Length of hospital stay, length of ICU stay, intraoperative blood loss, infections,
 Random sequence generation (selection bias): Unclear risk – Authors stated that randomisation was performed; however the method of randomisation was not reported. Allocation concealment (selection bias): Unclear risk – The approach to and use of allocation concealment was unclear Blinding of participants and personnel (performance bias): Low risk – Authors did not state whether blinding was performed; however this is unlikely to have affected study results as objective outcomes were measured. Blinding of outcome assessment (detection bias): Unclear risk – Authors did not state whether blinding was performed Incomplete outcome data (attrition bias): Low risk – Losses to follow-up were small and relatively balanced across treatment arms. Selective reporting (reporting bias): Low risk – Low risk - All relevant outcomes were reported. Other bias: High risk – Postoperative analgesia was not the same in each group (possible performance bias). Overall risk of bias: High Directness: directly applicable

Full citation	Davis. (1987) Intrathecal morphine in aortic aneurysm surgery. Anaesthesia, 42: 491-7
Study details	Study type: Randomised controlled trial Location(s): UK
	Aim(s): The present study compares low-dose intrathecal morphine with balanced anaesthesia in aortic aneurysm surgery. Study dates: not reported
	Follow-up: not reported
	Sources of funding: not reported
Participants	Population: men undergoing open surgical repair of unruptured AAA Sample size: 30
	Inclusion criteria: Male patients who presented for aortic aneurysm surgery (open repair), who were in sinus rhythm, were not taking beta-adrenoceptor blocking drugs or calcium antagonists and had not sustained a recognised myocardial infarction in the preceding 6 months were included
	Exclusion criteria: not reported
	Baseline characteristics:
	 Mean age: epidural plus general anaesthesia group, 65.6 years; general anaesthesia-alone group, 53.8 years Sex: 100% male
	Mean aneurysm size: not reported
	Comorbidities: not reported
Intervention	Intrathecal opioid plus general anaesthesia:
	• Intrathecal injection of 0.8 mg preservative-free morphine in 4ml of 0.9% saline, without barbotage, at the L2-3 level through a 25-G needle immediately before pre-oxygenation. They received no further analgesia in theatre.
Comparison	General anaesthesia-alone:
	 Papaveretum 0.1 mg/kg by slow intravenous injection during preoxygenation and additional doses of the same drug during surgery to a total dose of 0.25-0.5 mg/kg depending upon body weight and pre-operative condition: the mean dose (standard deviation) was 30±10 mg with a range of 10-40 mg.
Outcomes measures	Pain & clinical respiratory depression
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Unclear risk – Authors stated that randomisation was performed; however the method of randomisation was not reported. Allocation concealment (selection bias): Unclear risk – The approach to and use of allocation concealment was not reported Blinding of participants and personnel (performance bias): Low risk – It is unclear whether participants were blinded to treatment allocations; however this is unlikely to have affected study results as objective outcomes were measured. Blinding of outcome assessment (detection bias): Low risk – Assessors blinded to intervention allocation. Incomplete outcome data (attrition bias): Low risk – There were low rates of losses to follow-up across treatment arms Selective reporting (reporting bias): Low-risk All relevant outcomes were reported

Full citation	Davis. (1987) Intrathecal morphine in aortic aneurysm surgery. Anaesthesia, 42: 491-7
	7. Other bias: Low risk – none identified
	Overall risk of bias: Low
	Directness: directly applicable
Full citation	Dodds TM, Burns K, DeRoo DB, ET AL. (1997). Effects of anesthetic technique on myocardial wall motion abnormalities during abdominal aortic surgery. Journal of Cardiothoracic and Vascular Anesthesia, 11: 129-36
Study details	Study type: Double blind, randomised controlled trial Location(s): Lebanon Aim(s): to assess whether supplementation of general anaesthesia with epidural anaesthesia would decrease the incidence of new left ventricular segmental wall motion abnormalities during abdominal aortic surgery
	Study dates: not reported Follow-up: not reported Sources of funding: not reported
Participants	 Population: people undergoing open surgical repair of unruptured AAA Sample size: 73 Inclusion criteria: patients scheduled for open repair of unruptured infrarenal AAA via an anterior, transperitoneal approach were included Exclusion criteria: a primary diagnosis of aortic occlusive disease, previous coronary artery bypass surgery, and contraindications to placement of an epidural catheter (coagulopathy, localized infection at site of insertion) or a pre-existing neurological deficit Baseline characteristics: Mean age: epidural plus general anaesthesia group, 71 years; general anaesthesia-alone group, 71 years Sex: epidural plus general anaesthesia group, 80% male; general anaesthesia-alone group, 80.7% male Mean aneurysm size: not reported Myocardial infarction: epidural plus general anaesthesia group, 67%; general anaesthesia-alone group, 54%
Intervention	 Epidural plus general anaesthesia: All patients were sedated with intravenous midazolam, as needed, while in a holding area outside the operating room, during placement of invasive catheters and the epidural; Before induction of general anaesthesia, an epidural catheter was placed between the tenth thoracic and second lumbar interspace, using a loss-of-resistance technique, in all patients; Induction of anaesthesia was similar in both study groups and was accomplished, after preoxygenation, with fentanyl, 2 to 5 micrograms, followed by thiopental sodium, 2 to 4mg/kg, endotracheal intubation followed administration of vecuronium, 0.1mg/kg, or suxamethonium, 1mg/kg

Full citation	Dodds TM, Burns K, DeRoo DB, ET AL. (1997). Effects of anesthetic technique on myocardial wall motion abnormalities during abdominal aortic surgery. Journal of Cardiothoracic and Vascular Anesthesia, 11: 129-36
	 Anaesthesia was maintained with a nitrous oxide/oxygen ratio of 1:1 and enflurane, 0.5 to 1.0 MAC (end-tidal concentration); vecuronium was used to maintain surgical relaxation
	 After the induction of general anaesthesia, patients were administered (in divided doses) 6 to 9 mL of 1.5% lidocaine with 1:200,000 adrenaline which served as a test dose and to establish initial epidural blockade; subsequently, a further 5 to 8 mL of 0.5% bupivacaine was administered followed by an infusion at 6 to 8 mL/h.
Comparison	General anaesthesia-alone
	 All patients were sedated with intravenous midazolam, as needed, while in a holding area outside the operating room, during placement of invasive catheters and the epidural;
	 Before induction of general anaesthesia, an epidural catheter was placed between the tenth thoracic and second lumbar interspace, using a loss-of-resistance technique, in all patients;
	 Induction of anaesthesia was similar in both study groups and was accomplished, after preoxygenation, with fentanyl, 2 to 5 micrograms, followed by thiopental sodium, 2 to 4mg/kg, endotracheal intubation followed administration of vecuronium, 0.1mg/kg, or suxamethonium, 1mg/kg
	 Anaesthesia was maintained with a nitrous oxide/oxygen ratio of 1:1 and enflurane, 0.5 to 1.0 MAC (end-tidal concentration); supplemental doses of fentanyl, 1 to 2 micrograms/kg/h, were administered as needed, and vecuronium was used to maintain surgical relaxation
Outcomes measures	In-hospital mortality, cardiac morbidity, respiratory morbidity, renal insufficiency & blood loss
Risk of bias assessment	 Random sequence generation (selection bias): Unclear risk – Authors reported that randomisation was performed; however no information was provided as to how it was performed.
(using Cochrane risk of bias tool)	 Allocation concealment (selection bias): Unclear risk – The approach to and use of allocation concealment was not reported. Blinding of participants and personnel (performance bias): Low risk –The anaesthetist caring for the patient was aware of group assignment, but patients were blinded to treatment group. This is unlikely to have affected study results as objective outcomes were measured.
	 Blinding of outcome assessment (detection bias): Low risk – Assessors were blinded to treatment allocations. Incomplete outcome data (attrition bias): Low risk – There were low rates of losses to follow-up across treatment arms. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported. Other bias: Low risk – none identified
	Overall risk of bias: Low
	Directness: directly applicable

Full citation	Fleron M-H, Weiskopf RB, Bertrand M, Mouren S et al. (2003) A comparison of intrathecal opioid and intravenous analgesia for the incidence of cardiovascular, respiratory, and renal complications after abdominal aortic surgery. Anesth Analg, 97: 2-12
Study details	Study type: Randomised controlled trial Location(s): France Aim(s): to evaluate whether the administration of neuraxial opioids, in the intraoperative and immediate postoperative periods, would reduce the combined incidence of major cardiac, respiratory, and renal complications after major abdominal aortic surgery Study dates: not reported Follow-up: not reported Sources of funding: not reported
Participants	 Population: people undergoing open surgical repair of unruptured AAA or aortoiliac occlusive disease Sample size: 217 Inclusion criteria: patients undergoing elective open repair of AAA or aortoiliac occlusive disease were included. Exclusion criteria: contraindications to dural puncture (clinical signs of coagulopathy, localized infection, septicaemia, graft infection, previous lumbar spinal surgery). Baseline characteristics: Mean age: epidural plus general anaesthesia group, 67 years; general anaesthesia-alone group, 66 years Sex: epidural plus general anaesthesia group, 89% male; general anaesthesia-alone group, 88% male Mean aneurysm size: not reported Aortic disease: epidural plus general anaesthesia group, 58%; general anaesthesia-alone group, 62% Coronary artery disease: epidural plus general anaesthesia group, 51%; general anaesthesia-alone group, 58% Congestive heart failure: epidural plus general anaesthesia group, 30%; general anaesthesia-alone group, 88% COPD: epidural plus general anaesthesia group, 34%; general anaesthesia-alone group, 30% Diabetes: epidural plus general anaesthesia group, 11%; general anaesthesia-alone group, 16%
Intervention	 Intrathecal opioid plus general anaesthesia: Balanced general anaesthesia with intravenous sufentanil, isoflurane, and 50% nitrous oxide combined with intrathecal opioid (1 micrograms/kg sufentanil with 8 micrograms/kg preservative-free morphine injected at the L4-5 interspace)
Comparison	 General anaesthesia-alone: Balanced general anaesthesia with intravenous sufentanil, isoflurane and 50% nitrous oxide. Anaesthesia was induced with intravenous 0.5 micrograms/kg IV sufentanil, and 1–2 mg/kg IV propofol
Outcomes measures	Major complications, cardiovascular complications, respiratory complications, renal complications & length of hospital stay
Risk of bias assessment	1. Random sequence generation (selection bias): Low risk – randomisation was performed using computer-generated random sequences
Abdominal aorl	tic aneurysm: evidence review for anaesthesia and analgesia for

people having surgical repair of unruptured and ruptured abdominal aortic aneurysm (March 2020)

Full citation	Fleron M-H, Weiskopf RB, Bertrand M, Mouren S et al. (2003) A comparison of intrathecal opioid and intravenous analgesia for the incidence of cardiovascular, respiratory, and renal complications after abdominal aortic surgery. Anesth Analg, 97: 2-12
(using Cochrane risk of bias tool)	 Allocation concealment (selection bias): Unclear risk – The approach to and use of allocation concealment was not reported. Blinding of participants and personnel (performance bias): Low risk – patients and those taking care of them were not blinded; however this is unlikely to have affected study results because objective outcomes were assessed. Blinding of outcome assessment (detection bias): Unclear risk – Authors did not report whether outcome assessors were blinded to treatment allocations. Incomplete outcome data (attrition bias): Low risk – There were low rates of losses to follow-up across treatment arms. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported. Other bias: High risk – Study population included some patients who were undergoing surgery for aortoiliac occlusive disease (35%), rather than AAA. Overall risk of bias: Moderate Directness: partially applicable

Full citation	Norris EJ, Beattie C, Perler BA et al. (2001) Double-masked randomized trial comparing alternate combinations of intraoperative anesthesia and postoperative analgesia in abdominal aortic surgery. Anesthesiology, 95: 1054-67
Study details	Study type: Double-blind, randomised controlled trial
	Aim(s): To compare alternate combinations of intraoperative anaesthesia and postoperative analgesia with respect to postoperative outcomes in patients undergoing surgery of the abdominal aorta
	Study dates: August 1993 to July 1997 Follow-up: not reported
	Sources of funding: not reported
Participants	Population: patients undergoing open surgery to repair unruptured AAA or surgery for aortoiliac occlusive disease, and visceral and renal arterial reconstruction
	Sample size: 168
	Inclusion criteria: patients undergoing elective abdominal aortic reconstructive surgery were included. Procedures included open abdominal aortic surgery for unruptured AAA or aortoiliac occlusive disease, as well as visceral and renal arterial reconstruction requiring abdominal aortic cross-clamping.
	Exclusion criteria: patients whose procedure required clamping of the thoracic aorta, contraindication to any feature of the proposed clinical management (including epidural anaesthesia, previous surgery or severe deformity of the thoraco-lumbar spine, previous or current neurologic disease affecting the lower hemithorax or below) opioid dependence and major surgery in the previous 14 days
	Baseline characteristics:

Full citation	Norris EJ, Beattie C, Perler BA et al. (2001) Double-masked randomized trial comparing alternate combinations of intraoperative anesthesia and postoperative analgesia in abdominal aortic surgery. Anesthesiology, 95: 1054-67
	 Mean age: epidural plus general anaesthesia group, 68 years; general anaesthesia-alone group, 69 years Sex: unclear as se Mean aneurysm size: not reported Comorbidities: not adequately reported
Intervention	 Thoracic epidural anaesthesia combined with a light general anaesthesia: Thoracic epidural catheter placement was performed via the midline approach using a standard loss-of-resistance technique at the T8–T9 interspace for patients requiring a left flank incision, and at the T10–T11 interspace for patients requiring a midline incision. Epidural bolus: 6ml (left flank incision) or 8ml (midline incision) of 0.5% bupivacaine with 50 micrograms fentanyl General anaesthesia was achieved as follows: each subject received 10–15ml/kg of lactated Ringer's solution before induction, followed by incremental doses of sodium thiopental (up to 500mg) and fentanyl (up to 250 micrograms, including sedation fentanyl) until unconsciousness was achieved; general anaesthesia was maintained using 50% nitrous oxide in oxygen and enflurane (0.2–0.8% end tidal)
Comparison	 General anaesthesia plus placebo: General anaesthesia was achieved as follows: each subject received 10–15ml/kg of lactated Ringer's solution before induction, followed by incremental doses of sodium thiopental (up to 500mg) and fentanyl (up to 250 micrograms, including sedation fentanyl) until unconsciousness was achieved; general anaesthesia was maintained using 50% nitrous oxide in oxygen and enflurane (0.2–0.8% end tidal)
Outcomes measures	Mortality, cardiac complications, respiratory complications, renal complications, intraoperative blood loss, reoperation, readmission to ICU
Risk of bias assessment (using Cochrane risk of bias tool)	 Random sequence generation (selection bias): Unclear risk – Authors reported that randomisation was performed but the methods were not specified Allocation concealment (selection bias): Unclear risk – The approach to and use of allocation concealment was not reported. Blinding of participants and personnel (performance bias): Low risk – it is unclear whether patients were blinded to treatment allocations; however, this is unlikely to have affected study results. Blinding of outcome assessment (detection bias): Unclear risk – Authors did not report whether outcome assessors were blinded to treatment allocations. Incomplete outcome data (attrition bias): Low risk – There were low rates of losses to follow-up across treatment arms. Selective reporting (reporting bias): Low risk – All relevant outcomes were reported. Other bias: patients undergoing elective abdominal aortic reconstructive surgery were included. Procedures included open abdominal aortic surgery for unruptured AAA or aortoiliac occlusive disease, as well as visceral and renal arterial reconstruction. Overall risk of bias: Moderate Directness: partially applicable

Appendix E – Forest plots

Epidural plus general anaesthesia versus general anaesthesia-alone/plus placebo during open repair of unruptured AAA

Mortality

	Epidural plus general anae	esthesia	General anaesthesia alone/plus	placebo		Risk Ratio		Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M–H, Fixe	d, 95% CI		
Baron	3	81	4	86	38.4%	0.80 [0.18, 3.45]					
Dodds	2	36	2	37	19.5%	1.03 [0.15, 6.91]					
Norris	5	85	4	75	42.1%	1.10 [0.31, 3.96]			•		
Total (95% CI)		202		198	100.0%	0.97 [0.41, 2.29]					
Total events	10		10								
Heterogeneity. Chi² = Test for overall effect:	0.11, df = 2 (P = 0.95); l ² = Z = 0.07 (P = 0.95)	0%					0.01	0.1 Favours epidural plus general anaesthesia	1 Favours general anaesthes) ia alone/plus placebo	10

Cardiovascular adverse events

Any cardiovascular adverse event

	Epidural plus general anae	sthesia	General anaesthesia alone/plus	; placebo		Risk Ratio		Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl		M–H, Fix	ed, 95% CI		
Baron	19	81	22	86	68.4%	0.92 [0.54, 1.56]			—		
Dodds	11	36	10	37	31.6%	1.13 [0.55, 2.33]			•		
Total (95% CI)		117		123	100.0%	0.98 [0.64, 1.51]		•			
Total events	30		32								
Heterogeneity. Chi² = Test for overall effect:	0.21, df = 1 (P = 0.65); $I^2 = Z = 0.07$ (P = 0.94)	0%					0.01	0.1 Favours epidural plus general anaesthesia	1 Favours general anaesthes) ia alone/plus placeł	100

Myocardial infarction

	Epidural plus general anae	esthesia	General anaesthesia alone/plus p	olacebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	CI M-H, Fixed, 95% CI
Baron	5	81	5	86	44.4%	1.06 [0.32, 3.53]	3]
Davies	2	25	1	25	9.1%	2.00 [0.19, 20.67]	7]
Dodds	3	36	3	37	27.1%	1.03 [0.22, 4.76]	5]
Norris	3	85	2	75	19.4%	1.32 [0.23, 7.71]	L]
Total (95% CI)		227		223	100.0%	1.19 [0.54, 2.60]	
Total events	13		11				
Heterogeneity. Chi ² =	0.27, df = 3 (P = 0.96); l ² =	0%					
Test for overall effect:	Z = 0.43 (P = 0.66)						Favours epidural plus general anaesthesia Favours general anaesthesia alone/plus placebo

Congestive heart failure

	Epidural plus general anae	sthesia	General anaesthesia alone/plus p	lacebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Baron	5	81	7	86	59.7%	0.76 [0.25, 2.29]	
Davies	3	25	2	25	17.6%	1.50 [0.27, 8.22]	
Dodds	2	36	1	37	8.7%	2.06 [0.19, 21.69]	
Norris	0	85	1	75	14.0%	0.29 [0.01, 7.12]	• • • • • • • • • • • • • • • • • • • •
Total (95% CI)		227		223	100.0%	0.94 [0.42, 2.10]	
Total events	10		11				
Heterogeneity. Chi ² =	1.37, df = 3 (P = 0.71); l ² =	0%					
Test for overall effect:	Z = 0.16 (P = 0.87)						Favours epidural plus general anaesthesia Favours general anaesthesia alone/plus placebo

Ventricular tachyarrhythmia

	Epidural plus general anae	esthesia	General anaesthesia alone/plus	placebo		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M–H, Fixed, 95% Cl	
Baron	1	81	0	86	24.4%	3.18 [0.13, 77.03]		
Davies	0	25	1	25	75.6%	0.33 [0.01, 7.81]		_
Norris	0	85	0	75		Not estimable		
Total (95% CI)		191		186	100.0%	1.03 [0.15, 7.07]		
Total events	1		1					
Heterogeneity: Chi² = Test for overall effect:	0.97, df = 1 (P = 0.32); l ² = Z = 0.03 (P = 0.98)	0%					0.01 0.1 1 Favours epidural plus general anaesthesia Favours general anae	10 100

Respiratory adverse events

Acute respiratory failure (prolonged ventilation)



Pneumonia

	Epidural plus general ana	esthesia	General anaesthesia alone/plus	s placebo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% Cl	M-H, Fixed, 95% Cl
Baron	8	81	16	86	75.6%	0.53 [0.24, 1.17]	
Dodds	1	36	4	37	19.2%	0.26 [0.03, 2.19]	
Norris	1	85	1	75	5.2%	0.88 [0.06, 13.86]]
Total (95% CI)		202		198	100.0%	0.50 [0.24, 1.01]	
Total events	10		21				
Heterogeneity. Chi ² =	0.56, df = 2 (P = 0.76); I^2 =	0%					
Test for overall effect:	Z = 1.93 (P = 0.05)						Favours epidural plus general anaesthesia Favours genreal anaesthesia alone/plus placebo

Renal adverse events

Renal failure



Renal insufficiency

	Epidural plus general anae	esthesia	General anaesthesia alone/plu	s placebo		Risk Ratio		Ris	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		M-H, Fix	ed, 95% CI		
Davies	2	25	1	25	30.4%	2.00 [0.19, 20.67]			•		
Dodds	1	36	2	27	69.6%	0.38 [0.04, 3.92]					
Total (95% CI)		61		52	100.0%	0.87 [0.19, 3.98]					
Total events	3		3								
Heterogeneity: Chi ² = Test for overall effect:	0.98, df = 1 (P = 0.32); $I^2 = Z = 0.18$ (P = 0.86)	0%					0.01	0.1 Favours epidural plus general anaesthesia	1 Favours general anaest	10 hesia alone/plus place	100 2 bo

Gastrointestinal adverse events

Gastrointestinal bleeding



Surgical complications

Blood loss



Sepsis

	Epidural plus general anaes	thesia	General anaesthesia alone/plus placebo			Risk Ratio		Risk	Ratio	
Study or Subgroup	Events	Total	Events Tota	l Weig	jht	M-H, Fixed, 95% CI		M-H, Fixe	d, 95% CI	
Baron	2	81	4 86	5 78.	5%	0.53 [0.10, 2.82]				
Norris	1	85	1 75	5 21.	5%	0.88 [0.06, 13.86]				
Total (95% CI)		166	161	1 100.	0%	0.61 [0.15, 2.51]				
Total events	3		5							
Heterogeneity: Chi² = Test for overall effect:	0.10, df = 1 (P = 0.76); I ² = 0 Z = 0.69 (P = 0.49))%					0.01	0.1 Favours epidural plus general anaesthesia	10 Favours general anaesthesia a	100 lone/plus placebo

Resource use

Length of stay



Intrathecal opioid plus general anaesthesia versus general anaesthesia-alone during open repair of unruptured AAA

Respiratory adverse events: respiratory depression

	Intrathecal plus general anaesthesia		General anaesthesia alone/plus placebo			Risk Ratio	Risk Ratio				
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		M-H, Fixe	ed, 95% CI		
Davis	0	13	0	12		Not estimable					
Fleron	2	105	2	112	100.0%	1.07 [0.15, 7.44]					
Total (95% CI)		118		124	100.0%	1.07 [0.15, 7.44]					
Total events	2		2								
Heterogeneity: Not app	blicable							01		10	100
Test for overall effect:	Z = 0.07 (P = 0.95)						0.01	Favours intrathecal plus general anaesthesia	Favours general anaesthe	esia alone/plus placeb	0 100

Appendix F – GRADE tables

Epidural plus general anaesthesia versus general anaesthesia-alone/plus placebo during open repair of unruptured AAA

Mortality

		Quality ass	essment			No of	patients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results					
In-hospital mor	hospital mortality; effect sizes below 1 favour epidural plus general anaesthesia group												
3 (Baron 1991, Dodds 1997, Norris 2001)	RCTs	Serious ¹	Very Serious ^{2,3}	Not serious	Very serious ⁴	202	198	RR 0.97 (0.41, 2.29)	Very low				
Cardiovascular	mortality; effect si	izes below 1 favo	ur epidural plus	general anaesthe	esia group								
1 Davies 1993	RCT	Serious ¹	Serious ⁵	N/A	Very serious ⁴	25	25	RR 2.00 (0.19, 20.7)	Very low				
12-month morta	lity; effect sizes b	elow 1 favour epi	dural plus gener	al anaesthesia g	roup								
1 Norris	NorrisRCTNot seriousSerious ² N/AVery serious ⁴ 8575RR 0.88 (0.30, 2.62)Very low												
1. Different pos	stoperative analgesi	a were used in ea	ch treatment arm	(Baron 1991 and	Davies 1993), dov	vngrade 1 level							

2. Study samples (in Baron 1991 and Norris 2001) included patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.

3. Intervention (in Baron 1991) includes flunitrazepam, which is not available in the UK, downgrade 1 level.

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

5. Lidocaine-adrenaline formulation & dosing (in Davies 1993 and Dodds 1997) varies significantly from that used in UK practice, downgrade 1 level.

Any adverse event

		Quality as	sessment			No of	patients	Effect estimate	Quality				
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results					
Adverse events gastrointestinal	Adverse events (myocardial infarction, congestive heart failure, ventricular tachyarrhythmias, supraventricular tachyarrhythmias, respiratory failure, renal insufficiency, gastrointestinal haemorrhage, hepatic failure, sepsis); effect sizes below 1 favour epidural plus general anaesthesia group												
1 Davies 1993	RCT	Serious ¹	Serious ²	N/A	Very serious ³	25	25	RR 1.27 (0.73, 2.23)	Very low				
 Different pos Lidocaine-ad 	Different postoperative analgesia were used in each treatment arm, downgrade 1 level Lidocaine-adrenaline formulation & dosing used varies significantly from that used in UK practice, downgrade 1 level.												

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

Cardiovascular adverse events

		Quality ass	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Any postoperat	ive cardiovascular	adverse event; e	effect sizes below	v 1 favour epidur	al plus general a	naesthesia grou	qı		
2 (Baron 1991, Dodds 1997)	RCTs	Serious ¹	Very Serious ^{2,3}	Not serious	Very serious ⁴	117	123	RR 0.98 (0.64, 1.51)	Very low
Cardiac death; e	effect sizes below	1 favour epidura	l plus general an	aesthesia group					
1 Norris 2001	RCT	Not serious	Serious ²	N/A	Very serious ⁴	85	25	RR 0.30 (0.01, 7.22)	Very low
Myocardial infa	rction; effect sizes	below 1 favour e	epidural plus gen	eral anaesthesia	group				
4 (Baron 1991, Davies 1993 Dodds 1997, Norris 2001)	RCTs	Serious ¹	Very Serious ^{2,3}	Not serious	Very serious ⁴	227	223	RR 1.19 (0.54, 2.60)	Very low
Congestive hea	rt failure; effect siz	zes below 1 favor	ur epidural plus g	jeneral anaesthe	sia group				
4 (Baron 1991, Davies 1993 Dodds 1997, Norris 2001)	RCTs	Serious ¹	Very Serious ^{2,3}	Not serious	Very serious ⁴	227	223	RR 0.94 (0.42, 2.10)	Very low
Prolonged myo	cardial ischaemia;	effect sizes belo	w 1 favour epidu	ral plus general	anaesthesia grou	р			
1 Baron 1991	RCT	Serious ¹	Very Serious ^{2,3}	N/A	Very serious ⁴	81	86	RR 1.06 (0.57, 1.98)	Very low

		Quality ass	essment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Ventricular tach	yarrhythmia; effec	t sizes below 1 fa	avour epidural pl	us general anaes	sthesia group				
3 (Baron 1991, Dodds 1997, Norris 2001)	RCTs	Serious ¹	Very Serious ^{2,3}	Not serious	Very serious ⁴	191	186	RR 1.03 (0.15, 7.07)	Very low
Supraventricula	r tachyarrhythmia	; effect sizes belo	ow 1 favour epidu	ural plus general	anaesthesia gro	up			
1 Davies 1993	RCT	Serious ¹	Serious ⁵	N/A	Very serious ⁴	25	25	RR 1.00 (0.22, 4.49)	Very low
Unstable angina	a; effect sizes belo	w 1 favour epidu	ral plus general a	anaesthesia grou	p				
1 Norris 2001	RCT	Not serious	Serious ²	N/A	Very serious ⁴	85	75	RR 0.88 (0.02, 44.0)	Very low
 Different pos Study sampled owngrade for the second s	stoperative analgesi es (in Baron 1991 a 1 level.	a were used in eac nd Norris 2001) in	ch treatment arm cluded patients w	(Baron 1991 and I ho are undergoing	Davies 1993), dow g surgery for aorto	vngrade 1 level iliac occlusive di	sease as well as s	some undergoing AAA re	pair,

Intervention in Baron 1991 includes flunitrazepam, which is not available in the UK, 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.
 Lidocaine-adrenaline formulation & dosing used (in Davies 1993 and Dodds 1997) varies significantly from that used in UK practice, downgrade 1 level.

Respiratory adverse events

		Quality ass	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Any respiratory	adverse event; ef	fect sizes below [•]	1 favour epidural	plus general ana	esthesia group				
1 Baron 1991	RCT	Serious ¹	Serious ^{2,3}	N/A	Serious ⁴	81	86	RR 0.92 (0.71, 1.19)	Very low
Acute respirato	ry failure (prolong	ed ventilation); e	ffect sizes below	1 favour epidura	l plus general an	aesthesia grou	р		
4 (Baron 1991, Davies 1993 Dodds 1997, Norris 2001)	RCT	Serious ¹	Serious ^{2,3}	Not serious	Very serious⁵	227	223	RR 0.81 (0.48, 1.39)	Very low
Duration of ven	tilation (hours); ef	fect sizes below	0 favour epidural	plus general ana	esthesia group				
1 Broekema 1996	RCT	Not serious	Not serious	N/A	Serious ⁶	20	20	MD 2.20 (-2.79, 7.19)	Moderate
Minor atelectas	is; effect sizes bel	ow 1 favour epid	ural plus general	anaesthesia gro	up				
1 Baron 1991	RCT	Serious ¹	Serious ^{2,3}	N/A	Very serious ⁵	81	86	RR 1.06 (0.74, 1.52)	Very low
Major atelectas	is; effect sizes belo	ow 1 favour epide	ural plus general	anaesthesia gro	up				
1 Baron 1991	RCT	Serious ¹	Serious ^{2,3}	N/A	Very serious ⁵	81	86	RR 2.65 (0.53, 13.3)	Very low
Pneumonia; eff	ect sizes below 1 f	avour epidural pl	lus general anae	sthesia group					
3 (Baron 1991, Dodds 1997, Norris 2001)	RCTs	Serious ¹	Serious ^{2,3}	Not serious	Serious ⁴	202	198	RR 0.50 (0.24, 1.01)	Very low
 Different positive Study samp downgrade 	stoperative analgesi les (in Baron 1991 a 1 level.	a were used in ea and Norris 2001) ir	ch treatment arm ncluded patients w	(Baron 1991 and I /ho are undergoing	Davies 1993), dow g surgery for aorto	/ngrade 1 level iliac occlusive di	sease as well as	some undergoing AAA re	pair,

Intervention (in Baron 1991) includes flunitrazepam, which is not available in the UK, 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.

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

6. Non-significant result, downgrade 1 level.

Renal adverse events

		Quality ass	sessment			No of	patients	Effect estimate	Quality	
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results		
Renal failure; et	ffect sizes below 1	favour epidural	plus general ana	esthesia group						
2 (Baron 1991, Norris 2001)	RCTs	Serious ¹	Serious ^{2,3}	Not serious	Very serious ⁴	166	161	RR 1.46 (0.41, 10.95)	Very low	
Renal insufficie	ncy; effect sizes b	elow 1 favour ep	idural plus gener	al anaesthesia g	roup					
2 (Dodds 1997, Davies 1993)	RCTs	Serious ¹	Serious ⁵	Not serious	Very serious ⁴	61	52	RR 0.87 (0.19, 3.98)	Very low	
1. Different pos 2. Study samp	Different postoperative analgesia were used in each treatment arm (Baron 1991 and Davies 1993), downgrade 1 level Study samples (in Baron 1991 and Norris 2001) included nationte who are undergoing surgery for aertailing acquisive disease as well as some undergoing AAA repair									

- Study samples (in Baron 1991 and Norris 2001) included patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair downgrade 1 level.
- 3. Intervention in Baron 1991 includes flunitrazepam, which is not available in the UK, downgrade 1 level.
- 4. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.
- 5. Lidocaine-adrenaline formulation & dosing used (in Davies 1993 and Dodds 1997) varies significantly from that used in UK practice, downgrade 1 level.

Gastrointestinal adverse events

		Quality as:	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Gastrointestina	I bleeding; effect s	izes below 1 fav	our epidural plus	general anaesth	esia group				
2 (Baron 1991, Davies 1993)	RCTs	Serious ¹	Very serious ^{2,3}	Serious ⁴	Very serious ⁵	106	111	RR 2.07 (0.39, 10.95)	Very low
1. Differer 2. Study s downgr	nt postoperative ana amples (in Baron 19 ade 1 level.	lgesia were used 991 and Norris 200	in each treatment 01) included patier	arm (Baron 1991 and the second s	and Davies 1993) going surgery for a	, downgrade 1 le aortoiliac occlusi	vel ve disease as wel	II as some undergoing AA	∖A repair,

- 3. Intervention in Baron 1991 includes flunitrazepam, which is not available in the UK, downgrade 1 level.
- 4. I² value between 33.3% and 66.7%, downgrade 1 level.
- 5. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.

Surgical complications

		Quality as	sessment		No of	patients	Effect estimate	Quality	
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Any major surg	ical complication;	effect sizes belo	w 1 favour epidu	ral plus general a	naesthesia grou	р			
1 Baron 1991	RCT	Serious ¹	Very serious ^{2,3}	N/A	Very serious ⁴	81	86	RR 0.69 (0.26, 1.85)	Very low
Blood loss; effe	ect sizes below 0 fa	avour epidural pl	us general anaes	thesia group					
3 (Broekema 1996, Davies 1993, Dodds 1997)	RCTs	Serious ¹	Serious ⁵	Not serious	Serious ⁶	81	82	MD -0.03 (-0.60, 0.54)	Very low
1 Norris 2001	RCT	Not serious	Serious ²	N/A	Very serious ⁷	85	75	Median difference = 0	Very low
Need for addition	onal analgesia; effe	ect sizes below 1	favour epidural	plus general ana	esthesia group				
1 Broekema 1996	RCT	Not serious	Not serious	N/A	Serious ⁸	20	20	RR 0.38 (0.17, 0.88)	Moderate
Sepsis; effect s	izes below 1 favou	ır epidural plus g	eneral anaesthes	sia group					
2 (Baron 1991, Norris 2001)	RCTs	Serious ¹	Very serious ^{2,3}	Not serious	Very serious ⁴	166	161	RR 0.61 (0.15, 2.51)	Very low
Wound infectio	n; effect sizes belo	ow 1 favour epidu	iral plus general	anaesthesia grou	ıp				
1 Davies 1993	RCT	Serious ¹	Serious ⁵	N/A	Very serious ⁴	25	25	RR 1.50 (0.27, 8.22)	Very low
Urinary tract in	fection; effect size	s below 1 favour	epidural plus ge	neral anaesthesia	a group				
1 Davies 1993	RCT	Serious ¹	Serious ⁵	N/A	Very serious ⁴	25	25	RR 3.00 (0.13, 70.3)	Very low
Pulmonary infe	ction; effect sizes	below 1 favour e	pidural plus gene	eral anaesthesia	group				
1 Davies 1993	RCT	Serious ¹	Serious ⁵	N/A	Very serious ⁴	25	25	RR 2.00 (0.19, 20.7)	Very low
 Different positive Study samp 	stoperative analgesi les (in Baron 1991 a	ia were used in ea and Norris 2001) ir	ich treatment arm ncluded patients w	(Baron 1991 and l /ho are undergoing	Davies 1993), dov g surgery for aorto	vngrade 1 level iliac occlusive di	sease as well as	some undergoing AAA re	pair,

downgrade 1 level.3. Intervention in Baron 1991 includes flunitrazepam, which is not available in the UK, downgrade 1 level.

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

5. Lidocaine-adrenaline formulation & dosing used (in Davies 1993 and Dodds 1997) varies significantly from that used in UK practice, downgrade 1 level.

6. Non-significant result, downgrade 1 level.

7. Median reported with level of statistical significance not reported, downgrade 2 levels.

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

Need for reoperation

		Quality as	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Reoperation; ef	fect sizes below 1	favour epidural p	olus general anae	esthesia group					
1 Norris 2001	RCT	Not serious	Serious ¹	N/A	Very serious ²	85	75	RR 1.06 (0.34, 3.33)	Very low
1. Study samp	les included patient	s who are undergo	oing surgery for ac	ortoiliac occlusive	disease as well as	some undergoir	ng AAA repair, do	wngrade 1 level.	

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

Resource use

		Quality as	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Duration of pos	toperative hospita	l stay (days); effe	ect sizes below 0	favour epidural	plus general ana	esthesia group			
2 (Baron 1991, Davies 1993)	RCTs	Serious ¹	Very serious ^{2,3}	Not serious	Serious ⁴	106	111	MD -0.10 (-1.24,1.05)	Very low
Duration of pos	toperative stay in i	intensive care ur	nit (hours); effect	sizes below 0 fav	vour epidural plu	us general anae	sthesia group		
1 Baron 1991	RCT	Serious ¹	Very serious ^{2,3}	N/A	Serious ⁴	25	25	MD 3.00 (-14.6, 20.6)	Very low
Readmission to	intensive care uni	it; effect sizes be	low 0 favour epi	dural plus genera	al anaesthesia gr	oup			
1 Broekema 1996	RCT	Not serious	Not serious	N/A	Very serious ⁵	85	75	RR 1.76 (0.33, 9.36)	Low
1. Different pos	stoperative analgesi	a were used in ea	ch treatment arm	(Baron 1991 and	Davies 1993), dov	vngrade 1 level			

2. Study samples (in Baron 1991 and Norris 2001) included patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.

3. Intervention in Baron 1991 includes flunitrazepam, which is not available in the UK, downgrade 1 level.

4. Non-significant result, downgrade 1 level.

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

Intrathecal opioid plus general anaesthesia versus general anaesthesia-alone during elective open repair

Mortality

		Quality ass	essment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Summary of results					
30-day mortality	; effect sizes belo	w 1 favour epidu	ral plus general a	anaesthesia grou	р				
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	RR 0.30 (0.06, 1.43)	Very low
1. Study sampl 2. Confidence	. Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level. 2. Confidence interval crosses two lines of a defined minimum clinically important difference (RR MIDs of 0.8 and 1.25), downgrade 2 levels.								

Any adverse event

		Quality ass	sessment			No of	patients	Effect estimate	Quality	
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results		
Adverse events respiratory dep	(myocardial dama ression, acute res	ge or infarction, piratory failure, o	congestive hear r renal insufficie	t failure, new car ncy); effect sizes	diac arrhythmia, below 1 favour e	new segmental epidural plus ge	or lobar atelecta eneral anaesthes	isis, confirmed pneumo ia group	onia, severe	
1 Fleron 2003 RCT Not serious Serious ¹ N/A Serious ² 105 112 RR 0.74 (0.48, 1.14) Low										
1. Study sampl	1. Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.									

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

Cardiovascular adverse events

		Quality ass	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Myocardial infa	rction; effect sizes	below 1 favour e	epidural plus ger	neral anaesthesia	group				
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	RR 0.53 (0.17, 1.72)	Very low
Congestive hea	rt failure; effect siz	zes below 1 favo	ur epidural plus g	general anaesthe	sia group				
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	RR 1.07 (0.22, 5.17)	Very low

		Quality ass	essment			No of patients		Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
New cardiac arr	hythmia; effect siz	es below 1 favou	ır epidural plus g	eneral anaesthes	sia group				
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	RR 0.53 (0.14, 2.08)	Very low
4 Otradia	a transformed and a second and and	the second s	a mana a succeder a succede		a sector strate attacks as		- · · · · · · · · · · · · · · · · · · ·	• • • • • • • • • • • • • • • • • • •	1

1. Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.

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

Respiratory adverse events

		Quality ass	sessment			No of	patients	Effect estimate	Quality
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Respiratory dep	pression; effect siz	es below 1 favou	ır epidural plus g	eneral anaesthes	sia group				
2 (Fleron 2003, Davis 1997)	RCTs	Not serious	Serious ¹	Not serious	Serious ²	118	124	RR 0.92 (0.71, 1.19)	Low
Acute respirato	ry failure (prolong	ed ventilation); e	ffect sizes below	1 favour epidura	al plus general ar	aesthesia grou	р		
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ³	105	112	RR 0.81 (0.48, 1.39)	Very low
Major atelectas	is; effect sizes bel	ow 0 favour epidu	ural plus general	anaesthesia gro	up				
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Serious ⁴	105	112	MD 2.20 (-2.79, 7.19)	Low
Pneumonia; eff	ect sizes below 1 f	avour epidural pl	us general anaes	sthesia group					
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ³	105	112	RR 1.06 (0.74, 1.52)	Very low

1. Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.

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

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

4. Non-significant result, downgrade 1 level.

Renal adverse events

Quality assessment					No of patients		Effect estimate	Quality	
No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
Renal failure; effect sizes below 1 favour epidural plus general anaesthesia group									
1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	RR 0.83 (0.32, 2.15)	Very low
1. Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.									

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

Resource use

Quality assessment					No of patients		Effect estimate	Quality		
	No of studies	Design	Risk of bias	Indirectness	Inconsistency	Imprecision	Epidural plus general anaesthesia	General anaesthesia- alone	Summary of results	
	Renal failure; effect sizes below 1 favour epidural plus general anaesthesia group									
	1 Fleron 2003	RCT	Not serious	Serious ¹	N/A	Very serious ²	105	112	median difference = 0	Very low
					e					

Study sample included a proportion patients who are undergoing surgery for aortoiliac occlusive disease as well as some undergoing AAA repair, downgrade 1 level.
 Median reported with level of statistical significance not reported, downgrade 2 levels

Appendix G – Economic evidence study selection



Appendix H – Excluded studies

Clinical studies

Short Title		Reason for exclusion
Abdallah (2013)	Analgesic benefits of preincisional transversus abdominis plane block for abdominal aortic aneurysm repair	Not a controlled trial or systematic review of controlled trials
Aivatidi (2011)	Oxidative stress during abdominal aortic aneurysm repair - Biomarkers and antioxidant's protective effect: A review	Study does not contain a relevant intervention
Ansley (2006)	Is anesthesia good for you? Timing is everything!	Not a controlled trial or systematic review of controlled trials
Arar (2015)	Combined spinal-epidural anesthesia or local anesthesia + Sedoanalgesia in abdominal aortic Aneurism Repair?	Not a controlled trial or systematic review of controlled trials (retrospective study)
Asakura (2010)	In reply: The anesthetic technique of choice for better outcomes in high-risk elderly patients undergoing endovascular repair of aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Ballantyne (2004)	Does epidural analgesia improve surgical outcome?	Not a controlled trial or systematic review of controlled trials
Barbieri (2011)	Analgesia and endocrine surgical stress: effect of two analgesia protocols on cortisol and prolactin levels during abdominal aortic aneurysm endovascular repair	Study does not contain a relevant intervention
Baril (2007)	Endovascular Abdominal Aortic Aneurysm Repair: Emerging Developments and Anesthetic Considerations	Not a controlled trial or systematic review of controlled trials
Baril (2009)	The management of ruptured infrarenal abdominal aortic aneurysms	Not a controlled trial or systematic review of controlled trials
Barker (2005)	High thoracic epidural with general anesthesia for combined simultaneous on-pump coronary artery bypass grafts and abdominal aortic aneurysm repair	Not a peer-reviewed publication
Bayly (2001)	In-hospital mortality from abdominal aortic surgery in Great Britain and Ireland: Vascular Anaesthesia Society audit	Study does not contain a relevant intervention
Berggren (1989)	Eleven years of aortic aneurysm surgery: changes in techniques and results	Not a controlled trial or systematic review of controlled trials
Blake (1998)	Patient-controlled epidural versus intravenous pethidine to supplement epidural bupivacaine after abdominal aortic surgery	Study does not contain a relevant intervention (postoperative pain management)
Blay (2006)	Efficacy of low-dose intrathecal morphine for postoperative analgesia after abdominal aortic surgery: A double-blind randomized study	Study does not contain a relevant intervention (postoperative pain management)

Short Title		Reason for exclusion
Bookallil (1968)	Anaesthetic management of aortic aneurysms	Not a controlled trial or systematic review of controlled trials
Botney (1998)	Comparison of lumbar and thoracic epidural narcotics for postoperative analgesia in patients undergoing abdominal aortic aneurysm repair	Study does not contain a relevant intervention
Boylan (1998)	Epidural bupivacaine-morphine analgesia versus patient-controlled analgesia following abdominal aortic surgery. Analgesic, respiratory, and myocardial effects	Study does not contain a relevant intervention
Brady (2005)	Perioperative beta-blockade (POBBLE) for patients undergoing infrarenal vascular surgery: results of a randomized double-blind controlled trial	Study does not contain a relevant intervention
Brimacombe (1993)	A review of anaesthesia for ruptured abdominal aortic aneurysm with special emphasis on preclamping fluid resuscitation	Not a controlled trial or systematic review of controlled trials
Bull (1964)	Anaesthetic Problems in Resection of Abdominal Aortic Aneurysms	Not a controlled trial or systematic review of controlled trials
Carli (1997)	Combined epidural/general anaesthesia	Not a controlled trial or systematic review of controlled trials
Chiesa (2013)	Open repair of juxtarenal aortic aneurysms	Not a controlled trial or systematic review of controlled trials
Chlebowski (1999)	Anesthesia for abdominal aortic aneurysm surgery part I: Preoperative evaluation	Not a controlled trial or systematic review of controlled trials
Chlebowski (1999)	Anesthesia for abdominal aortic surgery part II: Intraoperative and postoperative management	Not a controlled trial or systematic review of controlled trials
Chuter (1999)	Abdominal aortic aneurysm in high-risk patients: Short- to intermediate- term results of endovascular repair	Not a controlled trial or systematic review of controlled trials
Crawford (1982)	A comparison of intercostal block with general anesthesia for abdominal aortic aneurysm resection	Not a controlled trial or systematic review of controlled trials
Crosby (1990)	A randomized double-blind comparison of fentanyl- and sufentanil-oxygen anesthesia for abdominal aortic surgery	Study does not contain a relevant intervention
Cunningham (1989)	Anaesthesia for abdominal aortic surgery: A review (Part I)	Not a controlled trial or systematic review of controlled trials
Cunningham (1989)	Anaesthesia for abdominal aortic surgerya review (Part II)	Not a controlled trial or systematic review of controlled trials
Cunningham (1991)	Abdominal aortic surgery: Anesthetic implications	Not a controlled trial or systematic review of controlled trials

Short Title		Reason for exclusion
Curley (1996)	Rectus sheath bupivacaine analgesia after aortic surgery	Not a controlled trial or systematic review of controlled trials
Elisha (2014)	Anesthesia case management for endovascular aortic aneurysm repair	Not a controlled trial or systematic review of controlled trials
Ellard (2013)	Anaesthesia for vascular emergencies	Not a controlled trial or systematic review of controlled trials
Ellis (2005)	Pro: vascular stents in the radiology suite-an anesthesiologist is needed	Not a controlled trial or systematic review of controlled trials
Faggioli (2011)	Preferences of patients, their family caregivers and vascular surgeons in the choice of abdominal aortic aneurysms treatment options: The PREFER study	Not a controlled trial or systematic review of controlled trials
Fitzgerald (2003)	Perioperative Anaesthesiological Management and Outcome of the Ruptured Abdominal Aortic Aneurysm	Not a controlled trial or systematic review of controlled trials
Flaherty (2014)	Regional anesthesia for vascular surgery	Not a controlled trial or systematic review of controlled trials
Florence (1978)	Neuroleptanaesthesia for surgery of the abdominal aorta	Not a controlled trial or systematic review of controlled trials
Galt (1991)	The effect of ibuprofen on cardiac performance during abdominal aortic cross-clamping	Study does not contain a relevant intervention
Gamulin (1991)	Renal consequences of infrarenal aortic cross- clamping in humans: Influence of different anesthetic techniques	Not in English
Gold (1994)	The effect of lumbar epidural and general anesthesia on plasma catecholamines and hemodynamics during abdominal aortic aneurysm repair	No relevant outcomes reported
Gold (1997)	Comparison of lumbar and thoracic epidural narcotics for postoperative analgesia in patients undergoing abdominal aortic aneurysm repair	Study does not contain a relevant intervention
Gottlieb (2014)	Anesthesia for major vascular surgery	Not a controlled trial or systematic review of controlled trials
Haljamae (1999)	Anaesthesia in non-cardiac vascular surgery	Not a controlled trial or systematic review of controlled trials
Hartman (1997)	Anesthesia for abdominal aortic reconstruction	Not a controlled trial or systematic review of controlled trials
Her (1990)	Combined epidural and general anesthesia for abdominal aortic surgery	Not a controlled trial or systematic review of controlled trials

Short Title		Reason for exclusion
Herring (2013)	Anaesthesia for abdominal vascular surgery	Not a controlled trial or systematic review of controlled trials
Houweling (1992)	A haemodynamic comparison of epidural versus intrathecal sufentanil to supplement general anaesthesia for abdominal aortic surgery	Not a population of people undergoing surgery for an abdominal aortic aneurysm
Houweling (1993)	A haemodynamic comparison of intrathecal morphine and sufentanil supplemented with general anaesthesia for abdominal aortic surgery	Not a population of people undergoing surgery for an abdominal aortic aneurysm
Javid (2007)	Should all patients with a ruptured AAA be anaesthetised by a vascular specialist?	Study does not contain a relevant intervention
Joseph (1973)	Bloood loss and acid-base balance during elective abdominal aortic aneurysmectomy	Not a controlled trial or systematic review of controlled trials
Joshi (1997)	Ruptured aortic aneurysm and cardiac arrest associated with spinal anesthesia	Not a controlled trial or systematic review of controlled trials
Karkos (2011)	A meta-analysis and metaregression analysis of factors influencing mortality after endovascular repair of ruptured abdominal aortic aneurysms	Not a controlled trial or systematic review of controlled trials
Karlsen (1999)	Anaesthesia for abdominal aortic surgery	Not a controlled trial or systematic review of controlled trials
Karnwal (2009)	Endovascular abdominal aneurysm repair in nonagenarians: never beyond the limits	Not a peer-reviewed publication
Karthikesalingam (2012)	Locoregional anesthesia for endovascular aneurysm repair (Structured abstract)	Not a controlled trial or systematic review of controlled trials
Kilickan (2002)	Abdominal aortic aneurism operation in a high risk patient under combined spinal epidural anesthesia	Not a controlled trial or systematic review of controlled trials
Knight (1963)	Anaesthesia for the leaking abdominal aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Kothandan (2016)	Anesthetic considerations for endovascular abdominal aortic aneurysm repair	Not a controlled trial or systematic review of controlled trials
Koyama (2012)	Efficacy of oral clonidine premedication on postoperative management for open abdominal aortic surgery	Not a peer-reviewed publication
Krajcer (2012)	Single-center experience of percutaneous abdominal aortic aneurysm repair with local anesthesia and conscious sedation: technique and results	Not a controlled trial or systematic review of controlled trials
Kunisawa (2014)	The dexmedetomidine concentration required after remifentanil anesthesia is three-fold higher than that after fentanyl anesthesia or that for general sedation in the ICU	Not a controlled trial or systematic review of controlled trials

Short Title		Reason for exclusion
Lachat (2000)	Regarding "Feasibility of endovascular repair of abdominal aortic aneurysms with local anesthesia with intravenous sedation"	Not a peer-reviewed publication
Lachat (2000)	Regarding 'Feasibility of endovascular repair of abdominal aortic aneurysms with local anesthesia with intravenous sedation'	Not a peer-reviewed publication
Lichtor (2005)	Depth of anesthesia monitors and shock	Not a controlled trial or systematic review of controlled trials
Lindahl (2011)	Should I choose open surgery or EVAR for my aortic aneurysm repair? reflections on the PREFER study on patients' preferences	Not a controlled trial or systematic review of controlled trials
Lippman (2003)	Anesthesia for endovascular repair of abdominal and thoracic aortic aneurysms: A review article	Not a controlled trial or systematic review of controlled trials
Lippmann (2010)	The anesthetic technique of choice for better outcomes in high-risk elderly patients undergoing endovascular repair of aortic aneurysms	Not a peer-reviewed publication
Lippmann (2015)	An alternative anaesthetic technique on nonagenerians undergoing endovascular aortic surgery and long term outcomes	Not a peer-reviewed publication
Lombardo (2009)	Epidural plus general anesthesia vs general anesthesia alone for elective aortic surgery: effects on gastric electrical activity and serum gastrin secretion	No relevant outcomes reported
Lorentz (2008)	Anesthesia for endovascular surgery of the abdominal aorta	Not a controlled trial or systematic review of controlled trials
Lubarsky (1998)	The impact of choice of muscle relaxant on postoperative recovery time (multiple letters)	Not a controlled trial or systematic review of controlled trials
Mathes (2000)	Continuous spinal anesthetic technique for endovascular aortic stent graft surgery	Not a controlled trial or systematic review of controlled trials
Mehta (2010)	Endovascular aneurysm repair of ruptured abdominal aortic aneurysm: the Albany Vascular Group approach	Not a controlled trial or systematic review of controlled trials
Mehta (2010)	Ruptured Abdominal Aortic Aneurysm: Endovascular Program Development and Results	Not a controlled trial or systematic review of controlled trials
Miller (1989)	Continuous alfentanil infusion for abdominal aortic surgery	Not a controlled trial or systematic review of controlled trials
Muehling (2008)	Prospective randomized controlled trial to evaluate "fast-track" elective open infrarenal aneurysm repair	Study does not contain a relevant intervention
Muehling (2009)	A prospective randomized trial comparing traditional and fast-track patient care in elective open infrarenal aneurysm repair	Study does not contain a relevant intervention
Panaretou (2009)	Combined anaesthesia and postoperative epidural analgesia in patients with chronic	Not a peer-reviewed publication

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Short Title		Reason for exclusion
	obstructive pulmonary disease undergoing abdominal aortic aneurysm repair	
Panaretou (2009)	The effect of combined anaesthesia with epidural postoperative analgesia on splanchnic perfusion in patients undergoing abdominal aortic aneurysm repair	Not a peer-reviewed publication
Panaretou (2009)	Ropivacaine 0.2% vs. Levobupivacaine 0.125% combined with fentanyl for epidural analgesia after abdominal aortic aneurysm repair	Not a peer-reviewed publication
Panaretou (2012)	Combined general-epidural anesthesia with continuous postoperative epidural analgesia preserves sigmoid colon perfusion in elective infrarenal aortic aneurysm repair	No relevant outcomes reported
Paries (2002)	A multicenter experience with the Talent endovascular graft for the treatment of abdominal aortic aneurysms	Not a controlled trial or systematic review of controlled trials
Park (2002)	Anesthesia for endovascular repair of abdominal aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Pichel (2008)	Focus on: Vascular anaesthesia	Not a controlled trial or systematic review of controlled trials
Pol (2014)	Frailty should determine type of anesthesia in reducing postoperative delirium after vascular surgery and not vice versa	Not a controlled trial or systematic review of controlled trials
Primieri (1991)	[A comparison of the hemodynamic effects of midazolam and propofol during anesthetic induction in patients at vascular risk]	Not in English
Rasmussen (1946)	Paravertebral injection of procaine for pain produced by aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Riddell (2005)	Endovascular abdominal aortic aneurysm repair	Not a controlled trial or systematic review of controlled trials
Robertson (2011)	Anaesthesia for endovascular surgery	Not a controlled trial or systematic review of controlled trials
Saleh (1980)	Anesthesia and monitoring for aortic aneurysm surgery	Not a controlled trial or systematic review of controlled trials
Salman (2013)	Comparison of effects of epidural bupivacaine and intravenous meperidine analgesia on patient recovery following elective abdominal aortic surgery	Study does not contain a relevant intervention
Saratzis (2013)	Acute kidney injury after endovascular repair of abdominal aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Schurmann (2012)	Tips and tricks: Patient selection, when to carry on and when to stop	Not a controlled trial or systematic review of controlled trials

Short Title		Reason for exclusion
Seeling (1985)	Infrarenal aortic bypass operations - influence of neuroleptanaesthesia and continuous epidural anaesthesia on cardiovascular responses during surgery	Not in English
Shigematsu (1985)	Evaluation of anesthetic management for the surgery of the aortic aneurysm	Not a controlled trial or systematic review of controlled trials
Sitzman (2000)	Combined general and epidural anesthesia for abdominal aortic aneurysm surgery	Not a controlled trial or systematic review of controlled trials
Smaka (2011)	Perioperative management of endovascular abdominal aortic aneurysm repair: update 2010	Not a controlled trial or systematic review of controlled trials
Smeets (1993)	Endocrine-metabolic response to abdominal aortic surgery: A randomized trial of general anesthesia versus general plus epidural anesthesia	No relevant outcomes reported
Stoneham (2014)	IR relevant locoregional techniques	Not a controlled trial or systematic review of controlled trials
Svensson (1992)	Aortic dissection and aortic aneurysm surgery: clinical observations, experimental investigations, and statistical analyses. Part I	Not a controlled trial or systematic review of controlled trials
Telford (2010)	Anaesthesia for abdominal vascular surgery	Not a controlled trial or systematic review of controlled trials
Tham (1997)	Back pain following postoperative epidural analgesia: an indicator of possible spinal infection	Not a controlled trial or systematic review of controlled trials
Tsakiliotis (2011)	Evaluation of hemodynamic parameters in endovascular treatment of ruptured abdominal aortic aneurysms (RAAAs) using different anaesthetic techniques. Preliminary study	Not a peer-reviewed publication
Varty (2011)	Comments regarding 'Local anaesthesia for endovascular repair of infra-renal aortic aneurysms'	Not a controlled trial or systematic review of controlled trials
Wozniak (2005)	Anesthesia for open abdominal aortic surgery	Not a controlled trial or systematic review of controlled trials
Xue (2014)	Comparing cardioprotective effects of anesthesia methods in patients undergoing elective abdominal aortic surgery	Not a controlled trial or systematic review of controlled trials
Young (1988)	Anaesthesia for elective abdominal aortic surgery	controlled trial or systematic review of controlled trials
Zaugg (2014)	Sevoflurane-Compared with propofol-based anesthesia reduces the need for inotropic support in patients undergoing abdominal aortic aneurysm repair: Evidence of cardioprotection by volatile anesthetics in noncardiac surgery	Not a controlled trial or systematic review of controlled trials

Economic studies

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

Appendix I – 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 60 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.