Eversion technique versus conventional endarterectomy with patch angioplasty in carotid surgery: protocol for a systematic review with meta-analyses and trial sequential analysis of randomised clinical trials

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ABSTRACT

Introduction Traditional carotid endarterectomy is considered to be the standard technique for prevention of a new stroke in patients with a symptomatic carotid stenosis. Use of patch angioplasty to restore the arterial wall after longitudinal endarterectomy is, to date, not unequivocally proven to be superior to eversion technique. A systematic review is needed for evaluation of benefits and harms of the eversion technique versus the traditional endarterectomy with patch angioplasty in patients with symptomatic carotid stenosis.

Methods and outcomes The review will be conducted according to this protocol following the recommendations of the ‘Cochrane Handbook for Systematic Reviews’ and reported according to Preferred Reporting Items for Systematic Reviews and Meta-Analyses. Randomised clinical trials comparing eversion technique versus endarterectomy with patch angioplasty in patients with a symptomatic stenosis of the internal carotid artery will be included. Primary outcomes are all-cause mortality rate, health-related quality of life and serious adverse events. Secondary outcomes are 30-day stroke and mortality rate, symptomatic arterial restenosis or occlusion and non-serious adverse events. The databases Cochrane Central Register of Controlled Trials, PubMed/MEDLINE and EMBASE will be searched (November 2019). We will primarily base our conclusions on meta-analyses of trials with overall low-risk of bias. We will use trial sequential analysis to assist the evaluation of imprecision in Grading of Recommendations, Assessment, Development and Evaluation. However, if pooled point estimates of all trials are similar to pooled point estimates of trials with overall low risk of bias and there is lack of a statistical significant interaction between estimates from trials with overall high risk of bias and trials with overall low risk of bias we will consider the trial sequential analysis adjusted precision of the estimate achieved in all trials as the result of our meta-analyses.

Ethics and dissemination The proposed systematic review will collect and analyse data from published studies, therefore, ethical approval is not required. The results of the review will be disseminated by publication in a peer-review journal and submitted for presentation at conferences.

PROSPERO registration number CRD42019119361.

INTRODUCTION

Carotid artery stenosis occurs due to atherosclerosis and was described to be a pathological substrate for ischaemic diseases of the ipsilateral brain and eye by Fisher. Preventive management of asymptomatic carotid artery stenosis includes antiplatelet, statins,
antihypertensive medication, diabetic control, as well as lifestyle modifications.2–4 Traditional carotid endarterectomy (tCEA) is the preferred guideline treatment for patients with symptomatic stenosis of the carotid artery,5,6 primarily based on the European Carotid Surgery Trial and the North American Symptomatic Carotid Endarterectomy Trial.7–9

Two operation techniques are used globally: the eversion technique (ET) and the tCEA using a longitudinal arteriotomy and patch angioplasty. Both techniques have the same approach to the carotid artery. ET was first reported by De Bakey et al10 and later described by Etheredge.11 This technique has a potential advantage compared with the tCEA, because patch closure is not always necessary, but the downside is the possibility of transection of carotid sinus nerve branches resulting in loss of the baroreceptor reflex. Whereas tCEA using a longitudinal arteriotomy, the incision is made parallel to the nerve branches, with smaller chance of transection of these nerve fibres. Loss of the baroreceptor reflex is associated with postoperative hypertension, a risk factor for cerebral hyper perfusion syndrome. The sympathetic trunk is another structure at risk, damage may result in signs of Horner’s syndrome.

Closure in both techniques can be achieved by either direct suturing of the arterial wall or patch angioplasty in CEA.11 TCEA with patch (for closing the longitudinal arteriotomy of the arterial wall) is suggested to reduce both the risks of restenosis and recurrent ipsilateral stroke.12 Restenosis after tCEA occurs in 6%–36% of patients during long-term follow-up of at least 12 months.13–17 Restenosis after ET occurs in 1.7%–2.5% of patients during long-term follow-up of at least 12 months.18

European guidelines of both the European Society of Vascular Surgery and the Dutch society for vascular surgery consider CEA with patch angioplasty as the reference technique.9,19,20 A meta-analysis of 6 randomised clinical trials (RCTs) including 2790 operations in 2666 patients compared ET with tCEA and concluded that ET may reduce the risks of perioperative stroke and long-term restenosis.21 However, the observed differences in intervention effects may be explained by several confounding factors and/or differential use of cointerventions, such as the use of perioperative transcranial Doppler (TCD) monitoring, perioperative carotid pressure measurement, electroencephalographic (EEG) monitoring, selected use of shunting, regional anaesthesia and variations in materials used for patching.22–29

To determine which technique, ET or tCEA is more effective for symptomatic carotid stenosis from the patients’ perspective, it is important that all available evidence is evaluated according to the risks of errors in a systematic review in line with the Cochrane Handbook for Systematic Reviews of Interventions.30

Previous reviews such as the one by Paraskevasa et al32 could be considered as ‘similar’ to this review. However, the differences can be found in the methodological approach. Our review will be:

► Conducted after a prepublished protocol.
► We would like to compare ONE technique (eversion CEA) with ONE other technique (CEA with patch closure) to lower the chance of bias (avoid design error).
► We will report numbers of patients with complications instead of the incidence of complications based on number of CEAs.
► Besides performing conventional meta-analysis, we have also planned to perform TSA.
► Conclusions made using TSA (according to the instructions for use35) may very well be more reliable than those using traditional meta-analysis techniques only.

Objective
The objective is to conduct a systematic review with meta-analysis and TSA of randomised clinical trials, evaluating the benefits and harms of the ET versus the tCEA in symptomatic patients according to a prepublished protocol following the Cochrane Handbook for Systematic Reviews of Interventions.30

METHODS
This review will be conducted according to this protocol, registered at PROSPERO34 following the recommendations of the ‘Cochrane Handbook for Systematic Reviews of interventions’50 and will be reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement (www.prisma-statement.org).35

Studies
Only randomised clinical trials comparing ET versus tCEA with patch angioplasty (regardless of the types of patch materials used) will be included. Trials will be considered irrespective of language, blinding, outcomes or publication status. Two authors will screen all the hits from all the searching machines manually for RCTs without the use of a computerised filter. We will also consider quasi-randomised studies, controlled clinical studies and other observational studies for data on harm if retrieved with our searches for randomised clinical trials. So it may occur that trials that are not RCT can be included for data on harm if retrieved. This is because adverse events are rarely reported in randomised clinical trials. Moreover, observational studies may provide information on rare or late occurring adverse events.36 We are aware that the decision not to search for all observational studies may bias our review towards assessment of benefits and may overlook certain harms, such as late or rare harms. However, we will not include observational studies together with RCTs in the meta-analyses.
Patients
According to the current guideline\textsuperscript{7–9} patients with a symptomatic stenosis (>50\%) of the carotid artery will be considered. Only trials which evaluate the ET versus the tCEA in adult patients (≥18 years) will be included.\textsuperscript{39} We are aware of the guideline statement that endarterectomy may be considered in symptomatic internal carotid lesions of <70\%. Studies in children and animals will be excluded.

Experimental intervention
The experimental intervention is the ET. The current ET for resolving the symptomatic stenosis in the (internal) carotid artery is based on the description by Etheredge\textsuperscript{10} and involves an oblique circumferential transection of the internal carotid artery (ICA) at the level of the carotid sinus. After distal eversion the diseased intimal plaque is totally excised from the tunica media and tunica adventitia and subsequently after endarterectomy of the carotid sinus, anatomical reimplantation of ICA follows.\textsuperscript{37}

We wish to compare one experimental intervention to one control intervention to reduce chance for design error (clinical heterogeneity) in the experimental intervention used. Therefore, RCTs which compare primary closure of the arterial wall after longitudinal arteriotomy will be excluded. Other techniques for carotid surgery in symptomatic patients are investigated in separate reviews.\textsuperscript{38,39}

Control intervention
The control intervention is the tCEA with patch closure of the longitudinal incision made in the carotid artery regardless of the type of patch material used.\textsuperscript{11}

Cointerventions and cerebral monitoring
Procedures may either be performed under plexus or general anaesthesia. Patients usually receive 5000 international units of heparin intravenously before cross-clamping the carotid artery. Sometimes protamine sulfate is given after surgery. TCD is used for microemboli detection and look for any increase of blood flow to the brain during the surgery. Patients will or can be monitored intraoperatively with EEG. Patients will or can receive a shunt when EEG changes are observed. Intraoperative monitoring may vary in the trials such as the use of perioperative TCD monitoring, perioperative carotid pressure measurement, EEG monitoring. Other intraoperative cointerventions may also vary in the trials, for example, the selected use of shunting and the variations in the types of materials used.

Hypothesis
We want to relate to the null hypothesis that there is not any difference between the two treatments (H\textsubscript{0}: relative risk reduction (RRR)=0.00\% or risk ratio (RR)=1.00) as well as both the alternative hypotheses (H\textsubscript{1a} and H\textsubscript{1b}) that there is a difference (H\textsubscript{1a} of a 10\% RRR or H\textsubscript{1b} of a 15\% RRR) between ET and tCEA in patients with a symptomatic carotid lesion.

Outcomes
The outcome measures will be graded from the patients’ perspective (GRADE working group 2008, figure 1).\textsuperscript{40} Examples of serious adverse events (SAEs): stroke, bleeding, persisting neurological deficits, for example, patients developing signs of Horner’s syndrome,\textsuperscript{41} hypertension in need for (intravenous) medication.

Stroke and mortality within 30 days were considered as secondary outcomes. These outcomes are important but possibly unreliable especially in surgical interventions as short time outcomes may prevail for one intervention while (with crossing of survival curves) the other intervention prevail on the long term.\textsuperscript{42}

Primary outcomes
\begin{itemize}
\item All-cause mortality.
\item Proportion of participants with one or more SAEs; which is defined as: any untoward medical occurrence that results in death is life threatening, requires hospitalisation or prolongation of existing hospitalisation, results in persistent or significant disability or incapacity (or is a congenital anomaly or birth defect).\textsuperscript{43}
\item Health status (or: quality of life): any scale used by trialists to assess the participants’ reporting of their quality of life (or health status).
\end{itemize}

Secondary outcomes
\begin{itemize}
\item <30 days mortality rate.
\item <30 days stroke rate.
\item Symptomatic (50\%–99\%) arterial restenosis or occlusion.
\item Proportion of participants with one or more non-SAEs: any untoward medical occurrence in a participant that
does not meet the above criteria for a SAE is defined as a non-SAE.\textsuperscript{43}

- Lower importance for patients: asymptomatic (50\%–99\%) arterial restenosis or occlusion.

**Exploratory outcomes**

- Separately reported SAEs.
- Separately reported non-SAEs.

A number of patients with one or more complications were evaluated rather than the numbers of events, depending on the availability of data.

**Search strategy**

The Cochrane Central Register of Controlled Trials (CENTRAL) in the Cochrane Library, PubMed/MEDLINE and EMBASE will be searched. References of the identified trials will be searched to identify any further relevant randomised clinical trials. The search strategies are provided in online supplementary appendix 1. Searches will include MeSH descriptors such as ‘Clinical Trials’, ‘carotid endarterectomy’, ‘eversion’, ‘carotid artery disease’. We will also search online trial registries such as ClinicalTrials.gov (https://clinicaltrials.gov/), European Medicines Agency (EMA) (www.ema.europa.eu/ema/), WHO International Clinical Trial Registry Platform (www.who.int/ictrp) and the Food and Drug Administration (FDA) (www.fda.gov) for ongoing or unpublished trials. In addition, we plan to search Google Scholar (https://scholar.google.nl/) using the terms: eversion and/or Carotid and/or Endarterectomy in the title of the abstract/paper.

**Data collection**

Two authors will perform screening and select the trials for inclusion, independently. Excluded trials and studies will be listed with their reasons for exclusion. When disagreements should occur, a third author will be approached to reconcile. The authors will extract the following data: trial characteristics (year and language of publication, country in which the trial was conducted, year of conduction of the trial, single or multicentre trial, number of patients), patient characteristics (inclusion and exclusion criteria, mean age, mean body mass index and gender, smoking, diabetes mellitus, use of statin and platelet inhibitors), intervention characteristics (primary closure, closure by patch, use of shunting), co-interventions (local or general anaesthesia, perioperative TCD monitoring, perioperative carotid pressure measurement, EEG monitoring) and the outcome measures evaluated. If there are any unclear or missing data, the corresponding authors of the individual trials will be contacted, at least twice, for clarification.

**Risk of bias assessment**

Two authors will assess the risks of bias, without masking for trial names, according to the Cochrane Handbook for Systematic Reviews of Interventions,\textsuperscript{30} including the domains of generation of the allocation sequence, allocation concealment, blinding of participants, personnel and outcome assessors, incomplete outcome data, selective outcome reporting and bias risks such as vested interests (financial interest, academical interest or other parties such as the medical industry). Risk of bias components were scored as low, unclear or high risk of bias. Trials were classified as trials with low overall risk of bias if all risk of bias domains were scored as having low risk of bias. If one or more of the bias domains were scored as unclear or high risk of bias, the trial was considered to have high overall risk of bias. Trials classified as low risk of bias in all domains of sequence generation, allocation concealment, blinding, incomplete data, selective outcome reporting, source of funding and other potential risks of bias will be considered trials at overall low risk of bias. Trials with one or more of these domains scored as unclear or high risk of bias will be considered trials at overall high risk of bias.\textsuperscript{31 44 45}

**Sequence generation**

- Low risk of bias: The method used (eg, central allocation) is unlikely to induce bias on the final observed effect, such as:
  - Referring to a random number table.
  - Using a computer random number generator.
  - Coin tossing.
  - Shuffling cards or envelopes.
  - Throwing a dice.
  - Drawing of lots.
- Unclear risk of bias: Insufficient information to assess whether the method used is likely to introduce confounders.
- High risk of bias: The method is improper and likely to introduce confounding, for example, based on date of admission, or record number, or by odd or even date of birth.

**Allocation concealment**

Some aspects of the conduct of randomised trials, particularly blinding, are associated with a modest exaggeration of treatment effects on average, but there is little evidence that the average bias differs according to whether the outcome was subjectively or objectively assessed. However, lack of blinding in trials with subjective outcomes leads to increased heterogeneity and hence unpredictable bias in effect estimates. As far as possible, clinical and policy decisions should be cautious when they are based on trials in which blinding was not reported or not feasible and outcome measures were subjectively assessed.\textsuperscript{46}

- Low risk of bias: participants and investigators enrolling participants could not foresee assignment because one of the following, or an equivalent method, was used to conceal allocation:
  - Central allocation (including telephone).
  - Web-based and pharmacy-controlled randomisation.
  - Sequentially numbered drug containers of identical appearance.
  - Sequentially numbered, opaque, sealed envelopes.
- Unclear risk of bias: insufficient information to permit judgement of ‘low risk’ or ‘high risk’. This is usually the case if the method of concealment is not described or not described in sufficient detail to allow a definite judgement.
- High risk of bias: participants or investigators enrolling participants could possibly foresee assignments and thus introduce selection bias, such as allocation based on:
  - An open random allocation schedule.
  - Assignment envelopes were used without appropriate safeguards.
  - Alternation or rotation.
  - Date of birth.
  - Case record number.
  - Any other explicitly unconcealed procedure.

Blinding of participants and personnel
In surgical procedures, it is impossible to blind the surgeon who performs the procedure of CEA, while it is possible to blind the caregivers responsible for postoperative care as well as the patients.60 For this domain, we will consider the caregivers and patients and not the surgeon who performs the procedure, although a certain risk of bias will inevitably be present when evaluating surgical procedures.
- Low risk of bias: no blinding or incomplete blinding, but the review authors judge that the outcome is not likely to be influenced by lack of blinding or blinding of participants and key study personnel ensured, and it is unlikely that the blinding could have been broken.
- Unclear risk of bias: insufficient information to permit judgement of ‘low risk’ or ‘high risk’, or the study did not address this outcome.
- High risk of bias: no blinding or incomplete blinding, and the outcome is likely to be influenced by lack of blinding or blinding of key study participants and personnel attempted, but likely that the blinding could have been broken, and the outcome is likely to be influenced by lack of blinding.

Blinding of outcome assessment
- Low risk of bias: no blinding of outcome assessment, but the review authors judge that the outcome measurement is not likely to be influenced by lack of blinding or blinding of outcome assessment is ensured, and it is unlikely that the blinding could have been broken.
- Unclear risk of bias: insufficient information to permit judgement of ‘low risk’ or ‘high risk’ or the study did not address this outcome.
- High risk of bias: no blinding of outcome assessment, and the outcome measurement is likely to be influenced by lack of blinding, or blinding of outcome assessment, but likely that the blinding could have been broken, and the outcome measurement is likely to be influenced by lack of blinding.

Incomplete outcome data
- Low risk of bias:
  - No missing outcome data.
  - Reasons for missing outcome data unlikely to be related to true outcome (for survival data, censoring unlikely to be introducing bias).
  - Missing outcome data balanced in numbers across intervention groups, with similar reasons for missing data across groups.
  - For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk is not enough to have a clinically relevant impact on the intervention effect estimate.
  - For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes is not enough to have a clinically relevant impact on observed effect size.
  - Missing data have been imputed using appropriate methods.
- Unclear risk of bias: insufficient reporting of attrition/exclusions to permit judgement of ‘low risk’ or ‘high risk’ (eg, number randomised not stated, no reasons for missing data provided) or the study did not address this outcome.
- High risk of bias:
  - Reason for missing outcome data likely to be related to true outcome, with either imbalance in numbers or reasons for missing data across intervention groups.
  - For dichotomous outcome data, the proportion of missing outcomes compared with observed event risk early enough to induce clinically relevant bias in intervention effect estimate.
  - For continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes is not enough to have a clinically relevant impact on the intervention effect estimate.
  - ‘As-treated’ analysis done with substantial departure of the intervention received from that assigned at randomisation.
  - Potentially inappropriate application of simple imputation.

Selective outcome reporting
- Low risk of bias: the study protocol is available and all the studies prespecified (primary and secondary) outcomes that are of interest in the review have been reported in the prespecified way, or the study protocol is not available but it is clear that the published reports include all expected outcomes, including those that were prespecified.
- Unclear risk of bias: insufficient information to permit judgement of ‘low risk’ or ‘high risk’. It is likely that the majority of studies will fall into this category.
- High risk of bias:
  - Not all of the studies prespecified primary outcomes have been reported.
- One or more primary outcomes is reported using measurements, analysis methods or subsets of the data (eg, subscales) that were not prespecified.
- One or more reported primary outcomes were not prespecified (unless clear justification for their reporting is provided, such as an unexpected adverse effect).
- One or more outcomes of interest in the review are reported incompletely so that cannot be entered in a meta-analysis.
- The study report fails to include results for a key outcome that would be expected to have been reported for such a study.

Other bias
► Low risk of bias: the study appears to be free of other sources of bias.
► Unclear risk of bias: there may be a risk of bias, but there is either insufficient information to assess whether an important risk of bias exists or insufficient rationale or evidence that an identified problem will introduce bias.
► High risk of bias: there is at least one important risk of bias.

Statistical methods
Meta-analyses will be performed according to the Cochrane Handbook for Systematic Reviews of Interventions.30 The software package Review Manager (RevMan) V.5.3 will be used.47 Significant levels will be adjusted due to multiplicity of several outcomes. The results of each outcome will require an adjusted statistical significance level (threshold). An alpha of (0.05/((1+3)/2)=) 0.025 will be used for the primary outcomes to keep the family wise error rate (FWER) below 0.05. For the secondary outcomes, this will be 0.017,48 49 For exploratory outcomes, we will consider a \( p < 0.05 \) as significant, because we view these outcomes as only hypothesis-generating outcomes. For dichotomous variables, the RR with TSA-adjusted CIs will be calculated. For continuous variables, the mean difference (MD) or the standardised MD with 95% CI will be calculated.

For the outcome of SAE, we plan to estimate the proportion of patients with one or more SAE in each group and to analyse this outcome in a binary meta-analysis. However, as we anticipate the reporting of SAEs in trials to vary considerably we plan to do two analyses:
1. The cumulated SAE analysis: Assuming that only one SAE is reported per patient. We will summarise all reported SAE in each trial and calculate the proportion of summed SAE divided with number of randomised patients in the experimental and control intervention group, the number of patients in each group will be a maximum.
2. To avoid multiple counts of SAE in the same patients (SAE counting is not a statistical independent outcome) we will also analyse the most frequent SAE as if it represents the total number of SAEs in the experimental and control intervention group (best case scenario).

The impact of attrition bias will be explored using best/worst and worst/best-case scenarios: a best/worst-case scenario is one where all patients lost to follow-up in the intervention group are supposed to have survived while all patients lost to follow-up in the control intervention group have died. A worst/best-case scenario is the reverse.

Heterogeneity will be explored by \( X^2 \) test with significance set at \( p \) value of 0.10, and the quantity of heterogeneity will be measured by \( I^2 \). We will conduct both random-effects model and fixed-effect model meta-analyses. In case of discrepancies, the results of both models will be presented and we will primarily stress the result of the model with the result closest to null effect due to the principle of cautiousness.45 The analyses will be performed on an intention-to-treat basis whenever possible.

A funnel plot will be used to explore small trial bias and to use asymmetry in funnel plot of trial size against treatment effect to assess this bias. Begg’s and Egger’s tests will be used to test for asymmetry in funnel plots.50

Trial sequential analyses
Meta-analyses may result in type I errors and type II errors due to an increased risk of random error when sparse data are analysed and due to repeated significance testing when a cumulative meta-analysis is updated with new trials.51 52 To assess the risk of type I and type II errors, TSA will be used. The vast majority of meta-analyses (nearly 80%) in Cochrane systematic reviews have less than the required information size to conclude on a 30% RRR and less than 2% have sufficient power to conclude on a 10% RRR.53-55

TSA combines information size estimation for meta-analysis (cumulated sample size of included trials) with an adjusted threshold for statistical significance of meta-analysis.51 52 56 The latter, called trial sequential monitoring boundaries (TSMBs), reduce type I errors. In TSA, the addition of each trial in a cumulative meta-analysis is regarded as an interim analysis and helps to clarify whether additional trials are needed or not. The idea in TSA is that when the cumulative z-curve crosses the TSMB, a sufficient level of evidence has been reached and no further trials may be needed. If the z-curve does not cross the boundary of benefit and the required information size has not been reached, there may be insufficient evidence to reach a conclusion.51 52 57 58 -TSA can also be used for the evaluation of type II errors, that is, to evaluate whether further randomised trials are futile to show or discard the anticipated intervention effect (RRR or MD). This happens when the cumulative z-curve does cross the TSMBs for futility. TSA will be applied since it controls the risks of type I and type II errors in a cumulative meta-analysis and may provide important information on
how many more patients need to be included in further trials. The information size will be calculated as diversity-adjusted required information size (DARIS).59 We will do the primary analysis calculating the DARIS based on an a priori anticipated intervention effect of a 10% RRR which is close to a minimal important difference. We will conduct sensitivity analyses for a 15% RRR as well as the RRR suggested by the meta-analysis of the included trials.60 If the estimated Diversity of the meta-analysis is 0%, a sensitivity analysis with TSA using a diversity of 25% will be conducted. TSA will be performed on all outcomes. The required information size for primary outcomes will be calculated based on an a priori RRR of 10% and appropriately adjusted for diversity according to an overall type I error of 2.5% for the coprimary outcomes and 1.7% for the secondary outcomes to account for a FWER of 5% in all, we will use a power of 90% considering sparse data and repetitive testing.59 For secondary outcomes, the DARIS will be calculated using a power of 90%.59 As a sensitivity analysis, the DARIS will be calculated using the estimated intervention effect from the trials at low risk of bias in a conventional meta-analysis. If the required information size is surpassed for the TSA using the estimated intervention effect in the conventional meta-analysis or a TSMB is crossed a TSA with an anticipated intervention effect equal to the confidence limit closest to the null effect in the effect estimate from the conventional meta-analysis will be performed. The TSAs will be conducted using the control event proportion calculated from the unweighted control event proportion from the control groups of the actual meta-analyses.

Subgroup analyses
The following subgroup analyses will be performed:

Trials at overall low risk of bias (all except blinding of surgeons scored as low risk of bias) compared with trials at high overall risk of bias (two or more of the bias domains (excluding blinding of surgeons) scored as unclear or high risk).

Different patch materials may be used including venous, polytetrafluorethylene, Dacron and biopatches (bovine/porcine).29 Subgroup analyses will be conducted according availability of data on different types of materials.

Grading of Recommendations, Assessment, Development and Evaluation
We will use summary of findings tables to summarise the results of the trials with overall low risk of bias and for all trials, separately. Reasons for downgrading the quality of the available evidence are: risk of bias evaluation of the included bias domains, publication bias, heterogeneity, imprecision and indirectness (eg, length of stay is a surrogate outcome measure).62–65 We will compare the imprecision assessed according to GRADE with that of TSA.54

Patient and public involvement
Patients and/or public were not involved in this study.

Ethics and dissemination
The results of the systematic review will be disseminated by publication in a peer-review journal and submitted for presentation at relevant conferences. This protocol will be online available prior to the start of the review process, and at the PROSPERO website.34

Resemblances in literature
Our previous published protocol30 may show overlap with this current protocol. This overlap is because of the basic methods to conduct a review process in line with the Cochrane Handbook.30 Nevertheless, this review topic differs on specific important details for the well informed vascular community. We would like to emphasise that it is important to compare ONE (experimental) technique with ONE other technique (control intervention) to reduce the chance for design error. In many other reviews, all kinds of techniques are compared with other more or less similar techniques. This is wrong. The important point (one-to-one technique comparisons) was recently highlighted during a global Vascular Congress (VEITH) November 2019, New York, USA. Together with this important point, the rigid methodological approach and the sophisticated and specific recommendations were followed during the design process of this protocol. Together with the Copenhagen Trial Unit, we managed to reach consensus and a detailed description of specific outcomes. To some readers, this may only look as if there is a lot of duplication. However, the differences are in the details and are specific and needed to proceed in evaluating important techniques to reduce complication rates.

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