Protocol for a systematic review of reporting standards of anaesthetic interventions in randomised controlled trials

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ABSTRACT

Introduction There is significant variation in how anaesthesia is defined and reported in clinical research. This lack of standardisation complicates the interpretation of published evidence and planning of future clinical trials. This systematic review will assess the reporting of anaesthesia as an intervention in randomised controlled trials (RCT) against the Consolidated Standards of Reporting Trials for Non-Pharmacological Treatments (CONSORT-NPT) framework.

Methods and analysis Online archives of the top six journals ranked by impact factor for anaesthesia and the top three general medicine and general surgery journals will be systematically hand searched over a 42-month time period to identify RCTs describing the use of anaesthetic interventions for any invasive procedure. All modes of anaesthesia and anaesthesia techniques will be included. All study data, including the type of anaesthetic intervention described, will be extracted in keeping with the CONSORT-NPT checklist. Descriptive statistics will be used to summarise general study details including types/modes of anaesthetic interventions, and reporting standards of the trials.

Ethics and dissemination No ethical approval is required. The results will be used to inform a funding application to formally standardise general, local, regional anaesthesia and sedation for use in clinical research. The systematic review will be disseminated via peer-reviewed manuscript and conferences.

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INTRODUCTION

The choice of anaesthetic technique for different types of surgery and invasive procedures, and their suitability for individual patients, relies largely on evidence from high-quality randomised controlled trials (RCT) and clinician expertise. Despite the widespread use of anaesthesia, there is significant variation in how it is defined and reported in clinical research and consensus definitions are lacking. For example, there is significant overlap between definitions of deep sedation and general anaesthesia (GA). The American Society of Anesthesiologists define GA as a ‘drug-induced loss of consciousness during which patients are not arousable, even by painful stimulation’ and sedation is defined by the Royal College of Anesthesiologists as ‘a continuum of depressed conscious state with unpredictable inter-individual dose responses to the drugs used, which may result in unconsciousness’. This may explain why standardised scales to routinely measure, record and standardise depth of sedation are lacking.

Clinician autonomy is acknowledged as a necessity in anaesthesia and is a fundamental reason for variation in practice. There is emerging evidence that the choice of mode of anaesthesia (GA, local anaesthesia, regional or sedation) is multifactorial, formulated around clinicians’ expertise, preference, habit, policies, practicalities and may also be influenced by other healthcare professionals and patients. While the autonomous nature of anaesthesia is partly unavoidable, the lack of standardisation and consistency in how anaesthetic techniques are defined, administered and reported complicates the
interpretation of published evidence and planning of future RCTs.

The Consolidated Standards of Reporting Trials (CONSORT) makes recommendations for improving the quality of reporting in clinical trials, and is endorsed by many journals. Although published in 1996, several reviews have established that reporting standards of RCTs relating to anaesthetic interventions remain poor and concluded that clinicians and researchers would benefit from more uniform reporting. The more recent CONSORT extension for Non-Pharmacological Treatments (CONSORT-NPT) recognises the complexity of non-pharmacological interventions (which includes anaesthesia) that comprise multiple interacting components. Specific additions include the need to fully describe interventions, and standardise and monitor their delivery (ie, intervention fidelity) during RCTs, to facilitate reproducibility and ensure that effective interventions can be successfully implemented in clinical practice.

STUDY AIM
Anaesthesia provides an example of a complex intervention that the CONSORT-NPT extension targets for improved reporting in RCTs. To date, no study has assessed the quality of reporting of anaesthesia in relation to CONSORT-NPT. The aim of this study, therefore, is to systematically review and summarise the reporting of anaesthesia as an intervention in RCTs.

METHODS
This systematic review protocol follows the Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (PRISMA-P) guidelines. This review protocol has been registered with the International Prospective Register of Systematic Reviews (PROSPERO), and the protocol will be updated with amendments if required.

DATA SOURCES AND SEARCH STRATEGY
We will employ a hand-search methodology similar to that previously described by Blencowe et al in a systematic review of intervention design and delivery in surgery. Online archives of selected journals will be systematically hand searched to identify relevant articles. Articles published in the top journals by impact factor as listed in the Scimago Journal & Country Rank for anaesthesia (n=6), general medicine (n=3) and general surgery (n=3) will be included (table 1). Thomson Reuters Journal Citation Reports: Web of Knowledge will be accessed to confirm impact factor by citation. A hand-searching methodology will be used whereby content pages of relevant journals will be screened to identify eligible articles, rather than undertaking a formal search using MeSH terms and text words.

STUDY SELECTION INCLUSION AND EXCLUSION CRITERIA
Full-text RCTs describing the use of anaesthetic interventions within any invasive procedure in humans will be included. Studies where techniques comparing anaesthesia in cadavers, laboratory and animal studies will be excluded.

We define invasive procedures as ‘purposeful/deliberate access to the body is gained via an incision, percutaneous puncture, where instrumentation is used in addition to the puncture needle, or instrumentation via a natural orifice. It begins when entry to the body is gained and ends when the instrument is removed, and/or the skin is closed. Invasive procedures are performed by trained healthcare professionals using instruments, which include, but are not limited to, endoscopes, catheters, scalpels, scissors, devices and tubes. All modes of anaesthesia and anaesthesia techniques will be included. This will include studies comparing different modes of anaesthesia (eg, general, local, regional anaesthesia or sedation), as well as studies comparing different types/techniques of the same mode of anaesthesia (volatile vs intravenous GA).

Case reports, non-randomised studies, retrospective and other non-comparative studies will be excluded. Abstracts and conference proceedings will be excluded due to high probability of incomplete data. The hand-search strategy is shown in online supplementary appendix 1.
STUDY MANAGEMENT
Electronic article information will be downloaded into EndNote software. Abstracts and titles will be screened independently by two researchers to identify articles that are potentially relevant, for which the full-text articles will be obtained. Full-text articles will be screened against the inclusion criteria by one researcher.

DATA EXTRACTION AND ASSESSMENT
Data will be extracted using a prespecified form in keeping with the standard CONSORT checklist for reporting trials. Study data will be collected and managed using REDCap electronic data capture tools hosted at the University of Bristol, UK. Data extraction will be completed by one researcher and verified by a second independent researcher. Disagreements will be resolved through discussion with the senior researcher/other members of the research team.

The country of study, number and type of included centres and the volume of invasive procedures undertaken in each setting will be recorded.

Descriptions of the following study demographics will be recorded: journal of publication, anaesthetic mode and subtype as stated in the paper, availability of study protocol and, if available, the format of the protocol (weblink, published article or obtained through contacting authors directly).

The type of anaesthetic intervention described in each study will be recorded in keeping with the CONSORT-NPT checklist in as much detail as is published in either the included study or extracted study protocol.

The anaesthetic intervention will be recorded according to an initial draft typology of anaesthetic interventions developed by the authors. Reporting of anaesthetic technique will be categorised according to whether it was performed pre-procedurally, intra-procedurally or post-procedurally. This will be to allow as much information as is included in each study to be recorded and subsequently categorised. All studies providing information about any aspect of the anaesthetic intervention will be classified as reporting a description, regardless of the included level of detail. Any citations to anaesthetic interventions within the studies will be recorded separately.

Any reference to standardisation with regard to any anaesthetic technique will be recorded, including how and why this was done, and to which standard. For the purpose of this review, standardisation will be defined as a process ‘to establish a standard consisting of regulations for how something is to be done’.

The invasive procedure for which the anaesthetic was being used will be recorded, but no details regarding how the invasive procedure was performed will be recorded.

FIDELITY
Fidelity will be defined as ‘how far those responsible for delivering an intervention actually adhere to the intervention as it is outlined by its designers’ as previously described by Blencowe et al. For each study, the reporting of fidelity to the anaesthetic intervention will be recorded as per the CONSORT-NPT guideline. Assessment of fidelity will be performed through details of any strategies implemented in the study to improve fidelity and details of how it was measured. This will include any crossover between trial arms of participants.

ASSESSMENT OF RISK OF BIAS
The Cochrane Collaboration’s revised risk of bias tool will be used to assess the internal validity of selected RCTs. The tool includes an assessment of sequence generation and allocation concealment, blinding of participants and outcome assessment, incomplete outcome data and selective outcome reporting. Given that no meta-analysis will be performed for this review, the risk of bias assessment will be reported as an appendix within the main review.

DATA ANALYSIS
A PRISMA flow chart of search and study selection with included and excluded studies will be presented. Reasons for exclusion of studies will be given. Extracted data will be presented in tables.

Descriptive statistics will be used to summarise general study details including types and modes of anaesthetic interventions and reporting standards of the trials. The reporting of anaesthetic interventions against the CONSORT-NPT checklist will be reported qualitatively and in tabulated form. The reporting of anaesthetic interventions for the anaesthesia and non-anaesthesia journals will be reported both in combination and separately, to examine reporting standards between these journal types.

Formal statistical comparisons will not be undertaken in this review as the aim is to summarise reporting standards and not analyse specific trial results. This is in keeping with published systematic reviews that have summarised reporting standards in other research fields.

PATIENT AND PUBLIC INVOLVEMENT
There was no involvement of patients or the public in the design of this systematic review, and the research question was not formulated or developed with patient or public involvement. Additionally, no patients or public members will be required in order to complete the systematic review.

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Contributors LE contributed to conception and design of the study, has written and reviewed the manuscript before submission. KC, NSB, RM and RJH contributed to conception and design of the study and have reviewed the manuscript before submission. MQ and SW have reviewed the manuscript before submission. All authors have given final approval for publication and agree to be accountable for all aspects of the work.
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