A protocol for the development of the STROCSS guideline: Strengthening the Reporting of Cohort Studies in Surgery

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ABSTRACT

Introduction: Strengthening the reporting of observational studies in epidemiology (STROBE) coined in 2007, highlighted the importance of improving the quality of observational research by providing an item checklist in order to avoid inadequate reporting of research. However, currently there are no reporting guidelines specific to surgical cohort studies, which have an extremely important role within the surgical literature. The recent development of surgery specific guidelines has underscored how surgical and procedural interventions require additional detail for readers to have a complete, clear, transparent and reproducible understanding. The objective of this research is to conduct a Delphi consensus exercise to develop the STROCSS guideline (Strengthening the Reporting of Cohort Studies in Surgery).

Methods and analysis: Current guidelines for case series (PROCESS), Cohort Studies (STROBE) and randomised controlled trials (CONSORT) will be analysed to compile items to form baseline material for developing cohort guidelines in the Delphi consensus exercise. The Delphi questionnaire will be administered via Google Forms and conducted using standard Delphi Methodology. Surgeons and individuals with significant experience of reviewing cohort studies as well as those with experience in developing reporting guidelines will be invited to participate. In the first round, existing items from PROCESS and STROBE will be put forward and participants will be invited to augment them or contribute further items for consideration. The provisional guidelines will then be updated in successive rounds using the nine-point Likert scale as proposed by the Grading Recommendations, Assessment, Development, and Evaluations (GRADE) working group. This process will be used to agree Standard definitions for the outcomes.

Dissemination: The work will be published in a peer-reviewed journal and presented at national and international meetings. Findings will be disseminated to interested parties, and journals will be encouraged to endorse the reporting guidelines.

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information on adverse events and risks [3]. Likewise, cohort studies can provide ethical grounds for performing RCTs by providing evidence towards a given association. Cohort studies are able to assess causality by reviewing a large databases of patients, and can provide an opportunity for long follow-up [2]. Cohort studies in the surgical literature are popular. Level 2 or 3 evidence accounted for 20–55% of published research in the top three journals by impact factor in 2013 across six surgical specialties, a significant rise compared with the 2003 figures of 17–50% [4]. However, work by our group has shown them to be poorly reported in Plastic Surgery [5].

Currently there is little guidance to authors with regards to the minimum necessary criteria to report in cohort studies. Journals and authors may adopt the guidelines for other study types, including guidelines for case series (PROCESS) or case reports (Surgical case report ‘SCARE’ guidelines) developed by our group. However, these are not tailored for use in cohort studies where multiple groups exposed to different interventions exist. Not all the points in these criteria are applicable to cohort studies. For example, mentioning the drug history and past medical history of each participant in cohort studies is not feasible, but is essential when reporting on a single case. However, comorbidities can be summarised in order to assess and compare the fitness of different groups. Additionally, cohort studies are often comparative and additional items required for high-quality cohort studies are not included in PROCESS or SCARE. The STROBE guideline provides guidance on reporting cohort studies, but these are not tailored to Surgery [6]. The development of the SCARE guidelines, when compared with the CARE guidelines, illustrates just how important a surgery specific guideline is.

Cohort studies must report on the comparison being made, as well as the clinical relevance of a given comparison [2]. The main form of bias in cohort studies is selection bias. Patients are not randomly allocated to treatment or control groups. Thus, such studies are vulnerable to the possibility that pre-existing baseline differences in prognosis and hidden confounders between two groups may bias the assessment of the effect of an intervention. To minimise selection bias, comparison groups should be carefully selected, for example, by excluding patients with a defined diagnosis or specific characteristics that is known to confound the outcome of interest [7]. All cohort studies should recognise that unknown confounders could affect the results. Effort should be made to identify and measure potential confounders, followed by a control for any pre-existing confounding factors in statistical analysis, often multivariate analyses [2,8]. Authors must then also provide information on the distribution of potential confounders in the intervention and comparison [8].

There is therefore a need to develop guidelines specifically for use in cohort studies. The objective of this research is to conduct a Delphi consensus exercise amongst experienced cohort study reviewers and editors to develop STROCSS (Strengthening the Reporting of Cohort Studies in Surgery) guidelines.

Methods

A similar methodological approach will be taken to that used for developing the SCARE and PROCESS guidelines including Moher et al’s guidance on developing reporting guidelines [1,9]. Relevant points from the PROCESS [1], STROBE [10] and CONSORT guidelines [11], will be used to create a template table to use as baseline material in the Delphi rounds. Guidance from articles appraising cohort studies will also be considered. [2,8,12].

The Delphi process

The Delphi questionnaire will be administered by Google Forms. The questionnaire will be conducted using standard Delphi Methodology [13]. The same questionnaire will be completed by all stakeholders throughout the process. Participants will be invited to recommend adaptations to the current STROBE items and suggest new ones, with a relevance to surgical cohort studies.

In each success round participants will rate the importance of reporting each item according to a nine-point Likert scale, where 1 indicates not important, and 9 indicates critical. This methodology follows the recommendations outlined by the Grading Recommendations, Assessment, Development, and Evaluations (GRADE) working group [13]. Items scoring: 1 to 3 – indicates the item is of little important; 4 to 6 – indicate an item is important but not critical; 7 to 9 – indicates the items is critically important.

Items will proceed to the reporting guideline if they are scored between 7 and 9 by 70% of respondents and between 1 to 3 by fewer than 30% of respondents. Likewise, items will not proceed to reporting guidelines if they score between 1 and 3 by 70% of respondents, and between 7 and 9 by 30% or more of respondents.

The questionnaire will be administered through sequential rounds until a final set of items is agreed. The entire process will be conducted electronically. There is no predetermined number of Delphi rounds, but judging on previous work by our group [1,14] it is expected that two or three rounds will be required.

Participant selection

The panel who helped develop the SCARE and PROCESS guidelines will be invited back. Further participants will be drawn from the Editorial Board of the International Journal of Surgery.

Dissemination

The research will be published in a peer-reviewed journal and will be disseminated electronically and presented at national and international conferences. Journals publishing surgical cohort studies will be encouraged to endorse the statement.

Ethical approval

No ethical approval is required as patient data is not being directly collected.

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Author contribution

RAA – idea conception, revision and final manuscript approval.
MRB – drafting, revision, final manuscript approval.
MV – drafting, revision, final manuscript approval.
RT – drafting, revision, final manuscript approval.

Financial interests

None declared. None of the authors have a financial interest in any of the products, devices or drugs mentioned in the manuscripts.

Conflicts of interest

No conflicts of interest are declared. The authors have no financial, consultative, institutional and other relationships that might lead to bias or conflict of interest.
Guarantor

Riaz A. Agha.

Research registration UIN

This is not a human study.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.isjp.2017.08.001.

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