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Perspective

STROCSS 2021: Strengthening the reporting of cohort, cross-sectional and casecontrol studies in surgery

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ABSTRACT

Introduction: Strengthening The Reporting Of Cohort Studies in Surgery (STROCSS) guidelines were developed in 2017 in order to improve the reporting quality of observational studies in surgery and updated in 2019. In order to maintain relevance and continue upholding good reporting quality among observational studies in surgery, we aimed to update STROCSS 2019 guidelines.

Methods: A STROCSS 2021 steering group was formed to come up with proposals to update STROCSS 2019 guidelines. An expert panel of researchers assessed these proposals and judged whether they should become part of STROCSS 2021 guidelines or not, through a Delphi consensus exercise.

Results: 42 people (89%) completed the DELPHI survey and hence participated in the development of STROCSS 2021 guidelines. All items received a score between 7 and 9 by greater than 70% of the participants, indicating a high level of agreement among the DELPHI group members with the proposed changes to all the items.

Conclusion: We present updated STROCSS 2021 guidelines to ensure ongoing good reporting quality among observational studies in surgery.

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1. Introduction

Observational studies often feature in the surgical literature [1]. However, poor reporting quality among observational studies in surgery has been highlighted [2]. In the absence of good reporting quality, readers are unable to meaningfully assess the research, rendering it less useful [3]. The existence of reporting guidelines and the mandatory implementation of these guidelines by journals have shown to improve the reporting quality among various types of studies [4-6].

Hence, Strengthening The Reporting Of Cohort Studies in Surgery (STROCSS) guidelines were developed in 2017 in order to improve the reporting quality of cohort studies in surgery. Despite the title, STROCSS guidelines aimed to improve the reporting quality of all observational studies in surgery, including casecontrol studies and cross-sectional studies, as well as cohort studies [7]. STROCSS 2017 guidelines were updated in 2019; since

its inception, STROCSS guidelines have been cited over 1000 times illustrating their acceptance within the surgical research community [8]. We aimed to update STROCSS 2019 guidelines in order to maintain relevance and continue upholding good reporting quality among observational studies in surgery.

2. Methods

The DELPHI methodology used in the development of STROCSS 2017 and 2019 guidelines was used in the development of STROCSS 2021 guidelines [9].

2.1. Coming up with proposals to update STROCSS 2019 guidelines

A STROCSS 2021 steering group was formed; members collaborated over email, Google Docs and WhatsApp Messenger to come up with proposals to update STROCSS 2019 guidelines.

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2.2. Delphi process

The proposals to update STROCSS 2019 guidelines were put to an expert panel of researchers; they were asked to assess the proposals and judge whether they should become part of STROCSS 2021 guidelines or not, through a Delphi consensus exercise.

The Delphi questionnaire was sent to all participants using Google Forms. The participants were required to indicate whether they disagreed or agreed with the proposed changes to the 17 items of the STROCSS 2019 guidelines, using a nine-point Likert scale, where 1 indicated "strongly disagree" and 9 indicated "strongly agree". If greater than 70% of participants gave a score between 7 and 9 for a proposed change, this was deemed as consensus and the item was updated. If less than 70% of participants gave a score between 7 and 9 for a proposed change, the item was left unaltered.

2.3. Participants

Researchers who were involved in the development of STROCSS 2017 and 2019 guidelines were invited to participate again. In addition, members of the International Journal of Surgery (IJS) editorial board were invited; IJS has mandated authors submitting surgical research papers using observational methodology to comply with STROCSS guidelines and hence IJS is an ardent supporter of STROCSS guidelines. Participants were accomplished researchers, authors, journal reviewers, editorial board members and

Table 1STROCSS 2021 Delphi participants' scores ranging between 1 (strongly disagree) and 9 (strongly agree). Items listed correspond to individual sections of STROCSS.

Item	1-3 (%)	4-6 (%)	7-9 (%)
1	2.4	7.2	90.5
2a	0.0	2.4	97.6
2b	0.0	9.6	90.4
2c	2.4	7.2	90.5
2d	0.0	19.1	81.0
3	2.4	7.2	90.5
4a	2.4	7.2	90.5
4b	7.2	14.3	78.5
4c	0.0	11.9	88.2
4d	0.0	7.2	92.8
5a	0.0	7.2	92.8
5b	0.0	14.3	85.7
5c	2.4	4.8	92.8
5d	0.0	19.1	80.9
6a	0.0	4.8	95.2
6b	4.8	14.2	80.9
6c	2.4	9.5	88.1
7a	0.0	9.5	90.4
7b	0.0	14.2	85.7
7c	0.0	11.9	88.1
7d	4.8	9.5	85.7
7e	0.0	14.3	85.7
7f	0.0	11.9	88.1
8	0.0	9.5	90.5
9	2.4	9.6	88.0
10a	0.0	2.4	97.6
10b	0.0	9.5	90.4
10c	0.0	11.9	88.1
11a	0.0	19.0	80.9
11b	0.0	16.7	83.4
11c	0.0	14.3	85.7
12	0.0	9.6	90.4
13	2.4	19.1	78.5
14	0.0	9.5	90.5
15	0.0	14.3	85.7
16	2.4	14.3	83.3
17a	2.4	14.3	83.3
17b	0.0	4.8	95.2
17c	0.0	2.4	97.5

editors representing countries across North America, South America, Europe, Africa, Asia, and Australia.

3. Results

47 people agreed to participate in the development of STROCSS 2021 guidelines; 42 people (89%) completed the DELPHI survey and hence participated in the development of STROCSS 2021 guidelines. Table 1 shows a summary of the scores given by the Delphi participants to indicate agreement or disagreement with the proposed changes to each item of the STROCSS 2019 guidelines. All items received a score between 7 and 9 by greater than 70% of the participants, indicating consensus with the proposed changes to all the items. The revised STROCSS 2021 guidelines are shown in Table 2.

4. Discussion

Since the publication of STROCSS guidelines, it has been cited over 1000 times and thus enjoyed great acceptance within the surgical research community. We present the updated STROCSS 2021 guidelines to continue ensuring good reporting quality among observational studies in surgery; we encourage authors, reviewers, editors, and journals to adopt them.

Authors should cite STROCSS 2021 guidelines in their methods section; additionally, they should submit a completed STROCSS 2021 guidelines checklist alongside their manuscript for reviewers and editors to inspect and ensure compliance. STROCSS website (https://www.strocssguideline.com) has provided the STROCSS 2021 guidelines checklist in various formats to ensure accessibility.

5. Conclusion

We present updated STROCSS 2021 guidelines for authors, reviewers, editors, and journals to implement, with a view to ensuring good reporting quality among observational studies in surgery.

International Journal of Surgery author disclosure form

The following additional information is required for submission. Please note that failure to respond to these questions/statements will mean your submission will be returned. If you have nothing to declare in any of these categories, then this should be stated.

Please state any conflicts of interest

None declared - the authors have no financial, consultative, institutional, and other relationships that might lead to bias or conflict of interest.

Please state any sources of funding for your research

None.

Please state whether ethical approval was given, by whom and the relevant Judgement's reference number

Not applicable.

Research registration Unique Identifying number (UIN)

- 1. Name of the registry: Not applicable
- 2. Unique Identifying number or registration ID: Not applicable
- 3. Hyperlink to your specific registration (must be publicly accessible and will be checked): Not applicable

Table 2The full revised STROCSS 2021 checklist

The STROCSS	3 2021 Guideline	
Item no.	Item description	Pag
TITLE	-	
1	 Title The word cohort or cross-sectional or case-control is included* Temporal design of study is stated (e.g. retrospective or prospective) The focus of the research study is mentioned (e.g. population, setting, disease, exposure/intervention, outcome etc.) 	
	*STROCSS 2021 guidelines apply to cohort studies as well as other observational studies (e.g. cross-sectional, case-control etc.)	_
ABSTRACT		
2a	Introduction — briefly describe: • Background • Scientific rationale for this study • Aims and objectives	
2 b	Methods - briefly describe: Type of study design (e.g. cohort, case-control, cross-sectional etc.) Other key elements of study design (e.g. retro-/prospective, single/multi-centred etc.) Patient populations and/or groups, including control group, if applicable Exposure/interventions (e.g. type, operators, recipients, timeframes etc.) Outcome measures — state primary and secondary outcome(s)	
2c	Results - briefly describe: • Summary data with qualitative descriptions and statistical relevance, where appropriate	
2d	Conclusion - briefly describe: Key conclusions Implications for clinical practice Need for and direction of future research	
INTRODUCTI	ON	
3	Introduction — comprehensively describe: • Relevant background and scientific rationale for study with reference to key literature • Research question and hypotheses, where appropriate • Aims and objectives	
METHODS		
4a	 Registration In accordance with the Declaration of Helsinki*, state the research registration number and where it was registered, with a hyperlink to registry entry (this can be obtained from ResearchRegistry.com, ClinicalTrials.gov, ISRCTN etc.) All retrospective studies should be registered before submission; it should be stated that the research was retrospectively registered * "Every research study involving human subjects must be registered in a publicly accessible database before recruitment of the first subject" 	the
4b	Ethical approval Reason(s) why ethical approval was needed Name of body giving ethical approval and approval number Where ethical approval wasn't necessary, reason(s) are provided	
4c	Protocol Give details of protocol (a priori or otherwise) including how to access it (e.g. web address, protocol registration number etc.) If published in a journal, cite and provide full reference	
4d	Patient and public involvement in research • Declare any patient and public involvement in research • State the stages of the research process where patients and the public were involved (e.g. patient recruitment, defining research outcome dissemination of results etc.) and describe the extent to which they were involved.	nes,
5a	Study design State type of study design used (e.g. cohort, cross-sectional, case-control etc.) Describe other key elements of study design (e.g. retro-/prospective, single/multi-centred etc.)	
5b	Setting and timeframe of research – comprehensively describe: • Geographical location • Nature of institution (e.g. primary/secondary/tertiary care setting, district general hospital/teaching hospital, public/private, low-resource set etc.) • Dates (e.g. recruitment, exposure, follow-up, data collection etc.)	ting
5c	Study groups Total number of participants Number of groups Detail exposure/intervention allocated to each group Number of participants in each group	
5d	Subgroup analysis — comprehensively describe: • Planned subgroup analyses • Methods used to examine subgroups and their interactions	_

Table 2 (continued)

Idama e -	2021 Guideline	ъ.
Item no.	Item description	Pag
6a	Participants – comprehensively describe:	
	 Inclusion and exclusion criteria with clear definitions Sources of recruitment (e.g. physician referral, study website, social media, posters etc.) 	
	Length, frequency and methods of follow-up (e.g. mail, telephone etc.)	
6b	Recruitment – comprehensively describe:	
ob	• Methods of recruitment to each patient group (e.g. all at once, in batches, continuously till desired sample size is reached etc.)	
	• Any monetary incentivisation of patients for recruitment and retention should be declared; clarify the nature of any incentives provided	
	Nature of informed consent (e.g. written, verbal etc.)	
	Period of recruitment	_
6c	Sample size – comprehensively describe:	
	 Analysis to determine optimal sample size for study accounting for population/effect size Power calculations, where appropriate 	
	Margin of error calculation	
METHODS -	INTERVENTION AND CONSIDERATIONS	-
7a	Pre-intervention considerations — comprehensively describe: • Preoperative patient optimisation (e.g. weight loss, smoking cessation, glycaemic control etc.)	
	 Pre-intervention treatment (e.g. medication review, bowel preparation, correcting hypothermia/-volemia/-tension, mitigating bleeding risk, ICI 	U
	care etc.)	
7b	Intervention – comprehensively describe:	
	Type of intervention and reasoning (e.g. pharmacological, surgical, physiotherapy, psychological etc.)	
	Aim of intervention (preventative/therapeutic)	
	Concurrent treatments (e.g. antibiotics, analgesia, anti-emetics, VTE prophylaxis etc.) Manufactures and model details unless analicables.	
	Manufacturer and model details, where applicable	
7c	Intra-intervention considerations — comprehensively describe:	
	 Details pertaining to administration of intervention (e.g. anaesthetic, positioning, location, preparation, equipment needed, devices, sutures operative techniques, operative time etc.) 	S,
	 Details of pharmacological therapies used, including formulation, dosages, routes, and durations 	
	• Figures and other media are used to illustrate	
7d	Operator details — comprehensively describe:	_
	Requirement for additional training	
	Learning curve for technique	
	Relevant training, specialisation and operator's experience (e.g. average number of the relevant procedures performed annually)	
7e	Quality control – comprehensively describe:	
	Measures taken to reduce inter-operator variability Measures taken to preve consistency in other assects of intervention delivery.	
	 Measures taken to ensure consistency in other aspects of intervention delivery Measures taken to ensure quality in intervention delivery 	
7f		
/1	Post-intervention considerations — comprehensively describe: • Post-operative instructions (e.g. avoid heavy lifting) and care	
	Follow-up measures	
	Future surveillance requirements (e.g. blood tests, imaging etc.)	
8	Outcomes – comprehensively describe:	
	Primary outcomes, including validation, where applicable	
	Secondary outcomes, where appropriate	
	Definition of outcomes If any unlidated outcome management tools are used, give full reference.	
	 If any validated outcome measurement tools are used, give full reference Follow-up period for outcome assessment, divided by group 	
9	Statistics – comprehensively describe:	
J	Statistics – comprehensively describe. Statistical tests and statistical package(s)/software used	
	Confounders and their control, if known	
	Analysis approach (e.g. intention to treat/per protocol)	
	Any sub-group analyses Level of statistical simplificance	
DEGI W	Level of statistical significance	
RESULTS		
10a	Participants – comprehensively describe:	
	• Flow of participants (recruitment, non-participation, cross-over and withdrawal, with reasons). Use figure to illustrate.	
	 Population demographics (e.g. age, gender, relevant socioeconomic features, prognostic features etc.) Any significant numerical differences should be highlighted 	
106	• • •	
10b	 Participant comparison Include table comparing baseline characteristics of cohort groups 	
	Give differences, with statistical relevance	
	Describe any group matching, with methods	

Table 2 (continued)

The STROCSS	2021 Guideline	
Item no.	Item description	Page
10c	Intervention — comprehensively describe: • Degree of novelty of intervention • Learning required for interventions • Any changes to interventions, with rationale and diagram, if appropriate	
11a	Outcomes — comprehensively describe: • Clinician-assessed and patient-reported outcomes for each group • Relevant photographs and imaging are desirable • Any confounding factors and state which ones are adjusted	
11b	Tolerance — comprehensively describe: • Assessment of tolerability of exposure/intervention • Cross-over with explanation • Loss to follow-up (fraction and percentage), with reasons	
11c	Complications — comprehensively describe: • Adverse events and classify according to Clavien-Dindo classification* • Timing of adverse events • Mitigation for adverse events (e.g. blood transfusion, wound care, revision surgery etc.) *Dindo D, Demartines N, Clavien P-A. Classification of Surgical Complications. A New Proposal with Evaluation in a Cohort of 6336 Patients and Results of a Survey. Ann Surg. 2004; 240(2): 205-213	ı
12	Key results — comprehensively describe: Key results with relevant raw data Statistical analyses with significance Include table showing research findings and statistical analyses with significance	
DISCUSSION		_
13	Discussion — comprehensively describe: Conclusions and rationale Reference to relevant literature Implications for clinical practice Comparison to current gold standard of care Relevant hypothesis generation	
14	Strengths and limitations — comprehensively describe: Strengths of the study Weaknesses and limitations of the study and potential impact on results and their interpretation Assessment and management of bias Deviations from protocol, with reasons	
15	Relevance and implications — comprehensively describe: Relevance of findings and potential implications for clinical practice Need for and direction of future research, with optimal study designs mentioned	
CONCLUSION	l .	-
16	Conclusions • Summarise key conclusions • Outline key directions for future research	
DECLARATIO	INS	
17a	Conflicts of interest Conflicts of interest, if any, are described	
17b	Funding • Sources of funding (e.g. grant details), if any, are clearly stated • Role of funder	
17c	ContributorshipAcknowledge patient and public involvement in research; report the extent of involvement of each contributor	

Author contribution

RA: Concept and design, data interpretation and analysis, drafting, revision and approval of final manuscript. GM: Design, data collection, data interpretation and analysis, drafting, revision and approval of final manuscript.

Guarantor

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