The ST	The STROCSS 2021 Guideline		
ltem	Item description	Page	
no.			
TITLE			
1	<ul> <li>Title</li> <li>The word cohort or cross-sectional or case-control is included*</li> <li>Temporal design of study is stated (e.g. retrospective or prospective)</li> <li>The focus of the research study is mentioned (e.g. population, setting, disease, exposure/intervention, outcome etc.)</li> </ul>	Title Page	
	*STROCSS 2021 guidelines apply to cohort studies as well as other observational studies (e.g. cross-sectional, case-control etc.)		
ABSTR	ACT		
2a	Introduction – briefly describe: • Background • Scientific rationale for this study • Aims and objectives	$\checkmark$	
2b	<ul> <li>Methods - briefly describe:</li> <li>Type of study design (e.g. cohort, case-control, cross-sectional etc.)</li> <li>Other key elements of study design (e.g. retro-/prospective, single/multi-centred etc.)</li> <li>Patient populations and/or groups, including control group, if applicable</li> <li>Exposure/interventions (e.g. type, operators, recipients, timeframes etc.)</li> <li>Outcome measures – state primary and secondary outcome(s)</li> </ul>	✓	
2c	<ul> <li>Results - briefly describe:</li> <li>Summary data with qualitative descriptions and statistical relevance, where appropriate</li> </ul>	$\checkmark$	
2d	Conclusion - briefly describe: <ul> <li>Key conclusions</li> <li>Implications for clinical practice</li> <li>Need for and direction of future research</li> </ul>	$\checkmark$	
INTRO	DUCTION	•	
3 METHO	<ul> <li>Introduction – comprehensively describe:         <ul> <li>Relevant background and scientific rationale for study with reference to key literature</li> <li>Research question and hypotheses, where appropriate</li> <li>Aims and objectives</li> </ul> </li> </ul>	3	
		2	
4a	<ul> <li>Registration         <ul> <li>In accordance with the Declaration of Helsinki*, state the research registration number and where it was registered, with a hyperlink to the registry entry (this can be obtained from ResearchRegistry.com, ClinicalTrials.gov, ISRCTN etc.)</li> <li>All retrospective studies should be registered before submission; it should be stated that the research was retrospectively registered</li> </ul> </li> </ul>	3	
	* "Every research study involving human subjects must be registered in a publicly accessible database before recruitment of the first subject"		
4b	<ul> <li>Ethical approval</li> <li>Reason(s) why ethical approval was needed</li> <li>Name of body giving ethical approval and approval number</li> <li>Where ethical approval wasn't necessary, reason(s) are provided</li> </ul>	3	

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 ar	nd 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 outcomes,	
		dissemination of results etc.) and describe the extent to which they were	
		involved.	
5a	Study des	-	3
	•	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 a	nd timeframe of research – comprehensively describe:	3
	•	Geographical location	
	•	Nature of institution (e.g. primary/secondary/tertiary care setting, district	
		general hospital/teaching hospital, public/private, low-resource setting	
		etc.)	
5.	•	Dates (e.g. recruitment, exposure, follow-up, data collection etc.)	4
5c	Study gro		4
	•	Total number of participants	
	•	Number of groups	
	•	Detail exposure/intervention allocated to each group	
5d	• Cubarour	Number of participants in each group	4
Su	Subgroup	o analysis – comprehensively describe:	4
	•	Planned subgroup analyses	
6a	Participa	Methods used to examine subgroups and their interactions <b>nts</b> – comprehensively describe:	4
Ua	Faiticipai	Inclusion and exclusion criteria with clear definitions	4
	•	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	Recruitm	ent – comprehensively describe:	3-4
•••	•	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 s	ize – comprehensively describe:	-
	•	Analysis to determine optimal sample size for study accounting for	
		population/effect size	
	•	Power calculations, where appropriate	
	•	Margin of error calculation	
		RVENTION AND CONSIDERATIONS	I
7a	Pre-interv	vention 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, ICU	
1		care etc.)	1

7b	Intervention – comprehensively describe:	4
	<ul> <li>Type of intervention and reasoning (e.g. pharmacological, surgical,</li> </ul>	
	physiotherapy, psychological etc.)	
	<ul> <li>Aim of intervention (preventative/therapeutic)</li> </ul>	
	<ul> <li>Concurrent treatments (e.g. antibiotics, analgesia, anti-emetics, VTE</li> </ul>	
	prophylaxis etc.)	
	<ul> <li>Manufacturer and model details, where applicable</li> </ul>	
7c	Intra-intervention considerations – comprehensively describe:	4
	<ul> <li>Details pertaining to administration of intervention (e.g. anaesthetic,</li> </ul>	
	positioning, location, preparation, equipment needed, devices, sutures,	
	operative techniques, operative time etc.)	
	<ul> <li>Details of pharmacological therapies used, including formulation,</li> </ul>	
	dosages, routes, and durations	
7.1	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	
7.	number of the relevant procedures performed annually)	
7e	Quality control – comprehensively describe:	-
	Measures taken to reduce inter-operator variability	
	<ul> <li>Measures taken to ensure consistency in other aspects of intervention delivery</li> </ul>	
	<ul> <li>delivery</li> <li>Measures taken to ensure quality in intervention delivery</li> </ul>	
7f	Post-intervention considerations – comprehensively describe:	4
/ 1	<ul> <li>Post-operative instructions (e.g. avoid heavy lifting) and care</li> </ul>	-
	<ul> <li>Follow-up measures</li> </ul>	
	<ul> <li>Future surveillance requirements (e.g. blood tests, imaging etc.)</li> </ul>	
8	Outcomes – comprehensively describe:	4
	<ul> <li>Primary outcomes, including validation, where applicable</li> </ul>	
	<ul> <li>Secondary outcomes, where appropriate</li> </ul>	
	<ul> <li>Definition of outcomes</li> </ul>	
	<ul> <li>If any validated outcome measurement tools are used, give full reference</li> </ul>	
	<ul> <li>Follow-up period for outcome assessment, divided by group</li> </ul>	
9	Statistics – comprehensively describe:	5
	Statistical tests and statistical package(s)/software used	
	<ul> <li>Confounders and their control, if known</li> </ul>	
	<ul> <li>Analysis approach (e.g. intention to treat/per protocol)</li> </ul>	
	<ul> <li>Any sub-group analyses</li> </ul>	
	Level of statistical significance	
RESUL		
10a	Participants – comprehensively describe:	5
	Flow of participants (recruitment, non-participation, cross-over and	
	withdrawal, with reasons). Use figure to illustrate.	
	<ul> <li>Population demographics (e.g. age, gender, relevant socioeconomic</li> </ul>	
	features, prognostic features etc.)	
	Any significant numerical differences should be highlighted	
10b	Participant comparison	5
	<ul> <li>Include table comparing baseline characteristics of cohort groups</li> </ul>	
	<ul> <li>Give differences, with statistical relevance</li> </ul>	
	Describe any group matching, with methods	
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:	6
	Clinician-assessed and patient-reported outcomes for each group	
	Relevant photographs and imaging are desirable	
	<ul> <li>Any confounding factors and state which ones are adjusted</li> </ul>	
11b	Tolerance – comprehensively describe:	-
	Assessment of tolerability of exposure/intervention	
	Cross-over with explanation	
	<ul> <li>Loss to follow-up (fraction and percentage), with reasons</li> </ul>	
11c	<b>Complications</b> – comprehensively describe:	5-6
	<ul> <li>Adverse events and classify according to Clavien-Dindo classification*</li> </ul>	
	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	
12	Key results – comprehensively describe:	6
	<ul> <li>Key results with relevant raw data</li> </ul>	
	<ul> <li>Statistical analyses with significance</li> </ul>	
	<ul> <li>Include table showing research findings and statistical analyses with</li> </ul>	
	significance	
DISCU		T
13	Discussion – comprehensively describe:	7
	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:	9
	Strengths of the study	
	<ul> <li>Weaknesses and limitations of the study and potential impact on results</li> </ul>	
	and their interpretation	
	Assessment and management of bias	
. –	Deviations from protocol, with reasons	
15	Relevance and implications – comprehensively describe:	8-9
	Relevance of findings and potential implications for clinical practice	
	Need for and direction of future research, with optimal study designs	
	mentioned	
		0
16	Conclusions	9
	Summarise key conclusions	
	Outline key directions for future research	
	ARATIONS	Title nega
17a	Conflicts of interest	Title page
476	Conflicts of interest, if any, are described	Title is a set
17b	Funding	Title page
	<ul> <li>Sources of funding (e.g. grant details), if any, are clearly stated</li> </ul>	
1	Role of funder	

17c	Contributorship	
	Acknowledge patient and public involvement in research; report the extent of	submission
	involvement of each contributor	system

Table 2: The full revised STROCSS 2021 checklist