The S	The STROCSS 2019 Guideline			
ltem no.	Item description	Page		
TITLE				
1	 Title: The word cohort or cross-sectional or case-controlled is included The area of focus is described (e.g. disease, exposure/intervention, outcome) Key elements of study design are stated (e.g. retrospective or prospective) 	1		
ABST	RACT			
2a	Introduction: the following points are briefly described Background Scientific Rationale for this study 	1		
2b	 Methods: the following areas are briefly described Study design (cohort, retro-/prospective, single/multi-centred) Patient populations and/or groups, including control group, if applicable Interventions (type, operators, recipients, timeframes) Outcome measures 	1		
2c	Results: the following areas are briefly described - Summary data (with statistical relevance) with qualitative descriptions, where appropriate	1		
2d	Conclusion: the following areas are briefly described - Key conclusions - Implications to practice - Direction of and need for future research	2		
INTRO	DUCTION			
3	Introduction: the following areas are described in full Relevant background and scientific rationale Aims and objectives Research question and hypotheses, where appropriate 	2,3		
METH	ODS			
4a	 Registration and ethics Research Registry number is stated, in accordance with the declaration of Helsinki* All studies (including retrospective) should be registered before submission 	3		
	*"Every research study involving human subjects must be registered in a publicly accessible database before recruitment of the first subject" (this can be obtained from: ResearchRegistry.com or ClinicalTrials.gov or ISRCTN)			
4b	 Ethical Approval: the following areas are described in full Necessity for ethical approval Ethical approval, with relevant judgement reference from ethics committees Where ethics was unnecessary, reasons are provided 	3		
4c	 Protocol: the following areas are described comprehensively Protocol (<i>a priori</i> or otherwise) details, with access directions If published, journal mentioned with the reference provided 	3		

4 -1	Definition the second in December	1.
4d	Patient Involvement in Research	3
	- Describe how, if at all, patients were involved in study design e.g. were	
	they involved on the study steering committee, did they provide input	
<u> </u>	on outcome selection, etc.	
5a	Study Design: the following areas are described comprehensively	3
	- 'Cohort' study is mentioned	
<u></u>	- Design (e.g. retro-/prospective, single/multi-centred)	
5b	Setting: the following areas are described comprehensively	3
	- Geographical location	
	- Nature of institution (e.g. academic/community, public/private)	
	- Dates (recruitment, exposure, follow-up, data collection)	
5c	Cohort Groups: the following areas are described in full	3
	- Number of groups	
	- Division of intervention between groups	
5d	Subgroup Analysis: the following areas are described comprehensively	4
	- Planned subgroup analyses	
	- Methods used to examine subgroups and their interactions	
6a	Participants: the following areas are described comprehensively	3
	- Eligibility criteria	
	- Recruitment sources	
	- Length and methods of follow-up	
6b	Recruitment: the following areas are described comprehensively	4
	 Methods of recruitment to each patient group 	
	- Period of recruitment	
6c	Sample Size: the following areas are described comprehensively	3
	- Margin of error calculation	
	 Analysis to determine study population 	
	- Power calculations, where appropriate	
	RVENTION AND CONSIDERATIONS	
7a	Pre-intervention Considerations: the following areas are described	3
	comprehensively	
	- Patient optimisation (pre-surgical measures)	
	- Pre-intervention treatment (hypothermia/-volaemia/-tension; ICU care;	
	bleeding problems; medications)	
7b	Intervention: the following areas are described comprehensively	3
	- Type of intervention and reasoning (e.g. pharmacological, surgical,	
	physiotherapy, psychological)	
	- Aim of intervention (preventative/therapeutic)	
	- Concurrent treatments (antibiotics, analgaesia, anti-emetics, NBM,	
	VTE prophylaxis)	
	Manufacturer and model details where applicable	
7c	Intra-Intervention Considerations: the following areas are described	4
	comprehensively	
	- Administration of intervention (location, surgical details, anaesthetic,	
	positioning, equipment needed, preparation, devices, sutures,	
	operative time)	
	- Pharmacological therapies include formulation, dosages, routes and	
	durations	
	 Figures and other media are used to illustrate 	

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7d	Operator Details: the following areas are described comprehensively	3
	- Training needed	
	- Learning curve for technique	
_	- Specialisation and relevant training	
7e	Quality Control: the following areas are described comprehensively	3
	- Measures taken to reduce variation	
	 Measures taken to ensure quality and consistency in intervention 	
	delivery	
7f	Post-Intervention Considerations: the following areas are described	4
	comprehensively	
	 Post-operative instructions and care 	
	- Follow-up measures	
	- Future surveillance requirements (e.g. imaging, blood tests)	
8	Outcomes: the following areas are described comprehensively	4
	 Primary outcomes, including validation, where applicable 	
	- Definitions of outcomes	
	 Secondary outcomes, where appropriate 	
	 Follow-up period for outcome assessment, divided by group 	
9	Statistics: the following areas are described comprehensively	5
	 Statistical tests, packages/software used, and interpretation of 	
	significance	
	 Confounders and their control, if known 	
	 Analysis approach (e.g. intention to treat/per protocol) 	
	- Sub-group analysis, if any	
RESU	LTS	
10a	Participants: the following areas are described comprehensively	5
	- Flow of participants (recruitment, non-participation, cross-over and	_
	withdrawal, with reasons)	
	- Population demographics (prognostic features, relevant socioeconomic	
	features, and significant numerical differences)	
10b	Participant Comparison: the following areas are described comprehensively	5
	- Table comparing demographics included	Ū
	- Differences, with statistical relevance	
	- Any group matching, with methods	
10c	Intervention: the following areas are described comprehensively	5
	- Changes to interventions, with rationale and diagram, if appropriate	Ŭ
	- Learning required for interventions	
	- Degree of novelty for intervention	
11a	Outcomes: the following areas are described comprehensively	5
	- Clinician-assessed and patient-reported outcomes for each group	5
	- Relevant photographs and imaging are desirable	
	- Confounders to outcomes and which are adjusted	
11b	Tolerance: the following areas are described comprehensively	5
	- Assessment of tolerance	5
	 Loss to follow up, with reasons (percentage and fraction) 	
	 Cross-over with explanation 	
11c	Complications: the following areas are described comprehensively	
110	- Adverse events described	-
	 Adverse events described Classified according to Clavien-Dindo classification* 	
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 Mitigation for adverse events (blood loss, wound care, revision surg should be specified) *Dindo D, Demartines N, Clavien P-A. Classification of Surgical Complications. A New Proposal with Evaluation in a Cohort of 6336 Patien and Results of a Survey. Ann Surg. 2004; 240(2): 205-213 Key Results: the following areas are described comprehensively Key results, including relevant raw data 	
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12 Key Results: the following areas are described comprehensively - Key results, including relevant raw data	5
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	1
 Statistical analyses with significance 	
DISCUSSION	
13 Discussion: the following areas are described comprehensively	6
- Conclusions and rationale	0
- Reference to relevant literature	
- Implications to clinical practice	
- Comparison to current gold standard of care	
- Relevant hypothesis generation	
14 Strengths and Limitations: the following areas are described comprehensiv	/ely 7,8
- Strengths of the study	,,0
- Limitations and potential impact on results	
 Assessment of bias and management 	
15 Implications and Relevance: the following areas are described	9
comprehensively	-
 Relevance of findings and potential implications to clinical practice a 	are
detailed	
 Future research that is needed is described, with study designs 	
detailed	
CONCLUSION	
16 Conclusions:	10
 Key conclusions are summarised 	
 Key directions for future research are summarised 	
DECLARATIONS	
17a Conflicts of interest	
 Conflicts of interest, if any, are described 	10
17b Funding	
 Sources of funding (e.g. grant details), if any, are clearly stated 	10