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4d	Patient involvement in Research	
τu	- 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	3
	on outcome selection, etc.	5
5a	Study Design: the following areas are described comprehensively	
°u	- 'Cohort' study is mentioned	2
	- Design (e.g. retro-/prospective, single/multi-centred)	2
5b	Setting: the following areas are described comprehensively	
00	- Geographical location	
	- Nature of institution (e.g. academic/community, public/private)	2
	- Dates (recruitment, exposure, follow-up, data collection)	
5c	Cohort Groups: the following areas are described in full	
	- Number of groups	2-3
	- Division of intervention between groups	2-3
5d	Subgroup Analysis: the following areas are described comprehensively	
04	- Planned subgroup analyses	3
	- Methods used to examine subgroups and their interactions	
6a	Participants: the following areas are described comprehensively	
ou	- Eligibility criteria	
	- Recruitment sources	2-3
	- Length and methods of follow-up	
6b	Recruitment: the following areas are described comprehensively	
0.0	- Methods of recruitment to each patient group	2-3
	- Period of recruitment	
6c	Sample Size: the following areas are described comprehensively	
	- Margin of error calculation	
	- Analysis to determine study population	4
	- Power calculations, where appropriate	
Interv	vention and Considerations	
7a	Pre-intervention Considerations: the following areas are described	
	comprehensively	
	- Patient optimisation (pre-surgical measures)	3-4
	- Pre-intervention treatment (hypothermia/-volaemia/-tension; ICU care;	
	bleeding problems; medications)	
7b	Intervention: the following areas are described comprehensively	
	- Type of intervention and reasoning (e.g. pharmacological, surgical,	
	physiotherapy, psychological)	
	- Aim of intervention (preventative/therapeutic)	3-4
	- Concurrent treatments (antibiotics, analgaesia, anti-emetics, NBM,	
	VTE prophylaxis)	
	 Manufacturer and model details where applicable 	
7c	Intra-Intervention Considerations: the following areas are described	
	comprehensively	
	- Administration of intervention (location, surgical details, anaesthetic,	
	positioning, equipment needed, preparation, devices, sutures,	3-4
	operative time)	
	- Pharmacological therapies include formulation, dosages, routes and	
	durations	
	- Figures other media are used to illustrate	

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7d	Operator Details: the following areas are described comprehensively	
	- Training needed	3
	- Learning curve for technique	
	- Specialisation and relevant training	
7e	Quality Control: the following areas are described comprehensively	
	- Measures taken to reduce variation	
	 Measures taken to ensure quality and consistency in intervention 	4
	delivery	
7f	Post-Intervention Considerations: the following areas are described	
	comprehensively	
	 Post-operative instructions and care 	4
	- Follow-up measures	
	- Future surveillance requirements (e.g. imaging, blood tests)	
8	Outcomes: the following areas are described comprehensively	
	 Primary outcomes, including validation, where applicable 	
	- Definitions of outcomes	3-4
	- Secondary outcomes, where appropriate	
	- Follow-up period for outcome assessment, divided by group	
9	Statistics: the following areas are described comprehensively	
	- Statistical tests, packages/software used, and interpretation of	
	significance	4
	- Confounders and their control, if known	
	- Analysis approach (e.g. intention to treat/per protocol)	
	- Sub-group analysis, if any	
RESU		•
10a	Participants: the following areas are described comprehensively	
	- Flow of participants (recruitment, non-participation, cross-over and	
	withdrawal, with reasons)	4-5
	- Population demographics (prognostic features, relevant socioeconomic	
	features, and significant numerical differences)	
10b	Participant Comparison: the following areas are described comprehensively	
	- Table comparing demographic included	
	- Differences, with statistical relevance	4-5
	- Any group matching, with methods	
10c	Intervention: the following areas are described comprehensively	
100	- Changes to interventions, with rationale and diagram, if appropriate	
	- Learning required for interventions	4-5
	- Degree of novelty for intervention	
11a	Outcomes: the following areas are described comprehensively	
Пă	- Clinician-assessed and patient-reported outcomes for each group	4-5
	 Relevant photographs and imaging are desirable 	45
	 Confounders to outcomes and which are adjusted 	
11b		
UD	Tolerance: the following areas are described comprehensively Assessment of tolerance 	
		5
	- Loss to follow up, with reasons (percentage and fraction)	
44-	- Cross-over with explanation	
11c	Complications: the following areas are described comprehensively	
	- Adverse events described	5
	 Classified according to Clavien-Dindo classification* 	

	- Mitigation for adverse events (blood loss, wound care, revision surgery	
	should be specified)	
	*Diada D. Damantin et N. Olavian D.A. Olavaitian et Oumited	
	*Dindo D, Demartines N, Clavien P-A. Classification of Surgical	
	Complications. A New Proposal with Evaluation in a Cohort of 6336 Patients	
10	and Results of a Survey. Ann Surg. 2004; 240(2): 205-213	
12	Key Results: the following areas are described comprehensively	
	- Key results, including relevant raw data	4-5
	- Statistical analyses with significance	
	ISSION	
13	Discussion: the following areas are described comprehensively	
	- Conclusions and rationale	
	- Reference to relevant literature	
	 Implications to clinical practice 	5-7
	 Comparison to current gold standard of care 	
	- Relevant hypothesis generation	
14	Strengths and Limitations: the following areas are described comprehensively	
	- Strengths of the study	6-7
	 Limitations and potential impact on results 	
	 Assessment of bias and management 	
15	Implications and Relevance: the following areas are described	
	comprehensively	
	- Relevance of findings and potential implications to clinical practice are	_
	detailed	7
	- Future research that is needed is described, with study designs	
	detailed	
CONC	LUSION	
16	Conclusions:	
	- Key conclusions are summarised	7
	- Key directions for future research are summarised	
DECL	ARATIONS	
17a	Conflicts of interest	7
	- Conflicts of interest, if any, are described	/
17b	Funding	
~	- Sources of funding (e.g. grant details), if any, are clearly stated	7