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## CONSORT 2010 checklist of information to include when reporting a randomised trial $^{\ast}$

Section/Topic	Item No	Checklist item	Reported on page No
Title and abstract			
	1a	Identification as a randomised trial in the title	1
	1b	Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts)	
Introduction			
Background and	2a	Scientific background and explanation of rationale	4
objectives	2b	Specific objectives or hypotheses	4,5
Methods			
Trial design	3a	Description of trial design (such as parallel, factorial) including allocation ratio	6
	3b	Important changes to methods after trial commencement (such as eligibility criteria), with reasons	NA
Participants	4a	Eligibility criteria for participants	6
	4b	Settings and locations where the data were collected	_6
nterventions	5	The interventions for each group with sufficient details to allow replication, including how and when they were actually administered	6,7
Outcomes	6a	Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed	8,9
	6b	Any changes to trial outcomes after the trial commenced, with reasons	NA
Sample size	7a	How sample size was determined	12, 13
	7b	When applicable, explanation of any interim analyses and stopping guidelines	NA
Randomisation:			
Sequence	8a	Method used to generate the random allocation sequence	13
generation	8b	Type of randomisation; details of any restriction (such as blocking and block size)	13
Allocation	9	Mechanism used to implement the random allocation sequence (such as sequentially numbered containers),	13
concealment mechanism		describing any steps taken to conceal the sequence until interventions were assigned	
Implementation	10	Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions	13
Blinding	11a	If done, who was blinded after assignment to interventions (for example, participants, care providers, those	13

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	assessing outcomes) and how	
11b	If relevant, description of the similarity of interventions	NA
12a	Statistical methods used to compare groups for primary and secondary outcomes	13, 14
12b	Methods for additional analyses, such as subgroup analyses and adjusted analyses	13, 14
13a	For each group, the numbers of participants who were randomly assigned, received intended treatment, and	6
	were analysed for the primary outcome	
13b	For each group, losses and exclusions after randomisation, together with reasons	14
14a	Dates defining the periods of recruitment and follow-up	7, 7
14b	Why the trial ended or was stopped	NA
15	A table showing baseline demographic and clinical characteristics for each group	14
16	For each group, number of participants (denominator) included in each analysis and whether the analysis was	13, 14
	by original assigned groups	
17a	For each primary and secondary outcome, results for each group, and the estimated effect size and its	15, 16, 17
	precision (such as 95% confidence interval)	
17b	For binary outcomes, presentation of both absolute and relative effect sizes is recommended	NA
18	Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory	18,19
19	All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)	NA
20	Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses	21
		21, 22, 23
22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence	21, 22, 23
23	Registration number and name of trial registry	3
		6
25	Sources of funding and other support (such as supply of drugs), role of funders	23
	12a 12b 13a 13b 14a 14b 15 16 17a 17b 18 19	<ul> <li>If relevant, description of the similarity of interventions</li> <li>Statistical methods used to compare groups for primary and secondary outcomes</li> <li>Methods for additional analyses, such as subgroup analyses and adjusted analyses</li> <li>For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome</li> <li>For each group, losses and exclusions after randomisation, together with reasons</li> <li>Dates defining the periods of recruitment and follow-up</li> <li>Why the trial ended or was stopped</li> <li>A table showing baseline demographic and clinical characteristics for each group</li> <li>For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups</li> <li>For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)</li> <li>For binary outcomes, presentation of both absolute and relative effect sizes is recommended</li> <li>Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory</li> <li>All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)</li> <li>Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses</li> <li>Generalisability (external validity, applicability) of the trial findings</li> <li>Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence</li> <li>Registration number and name of trial registry</li> </ul>

<sup>\*</sup>We strongly recommend reading this statement in conjunction with the CONSORT 2010 Explanation and Elaboration for important clarifications on all the items. If relevant, we also recommend reading CONSORT extensions for cluster randomised trials, non-inferiority and equivalence trials, non-pharmacological treatments, herbal interventions, and pragmatic trials. Additional extensions are forthcoming: for those and for up to date references relevant to this checklist, see <a href="https://www.consort-statement.org">www.consort-statement.org</a>.

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