

CONSORT 2010 checklist of information to include when reporting a randomised trial*

Section/Topic	Ite m 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,	2	
		results, and conclusions (for specific guidance		
		see CONSORT for abstracts)		
Introduction				
Background and	2a	Scientific background and explanation of rationale	3	
objectives	2b	Specific objectives or hypotheses	3	
Methods	ı			
Trial design	За	Description of trial design (such as parallel,	4	
		factorial) including allocation ratio		
	3b	Important changes to methods after trial	N/A	
		commencement (such as eligibility criteria), with		
		reasons		
Participants	4a	Eligibility criteria for participants	4	
	4b	Settings and locations where the data were		
		collected		
Interventions	5	The interventions for each group with sufficient	4-5	
		details to allow replication, including how and		
		when they were actually administered		
Outcomes	6a	Completely defined pre-specified primary and	5	
		secondary outcome measures, including how and		
		when they were assessed		
	6b	Any changes to trial outcomes after the trial	N/A	
		commenced, with reasons		
Sample size	7a	How sample size was determined	5	
	7b	When applicable, explanation of any interim	N/A	

		analyses and stopping guidelines		
Randomisation:				
Sequence	8a	Method used to generate the random allocation	4	
generatio		sequence		
n	8b	Type of randomisation; details of any restriction	4	
		(such as blocking and block size)		
Allocation	9	Mechanism used to implement the random	4	
concealm		allocation sequence (such as sequentially		
ent		numbered containers), describing any steps taken		
mechanis		to conceal the sequence until interventions were		
m		assigned		
	10	Who generated the random allocation sequence,	4	
Implementation		who enrolled participants, and who assigned		
		participants to interventions		
Blinding	11a	If done, who was blinded after assignment to	N/A	
		interventions (for example, participants, care		
		providers, those assessing outcomes) and how		
	11b	If relevant, description of the similarity of	N/A	
		interventions		
Statistical	12a	Statistical methods used to compare groups for	5-6	
methods		primary and secondary outcomes		
	12b	Methods for additional analyses, such as	N/A	
		subgroup analyses and adjusted analyses		
Results				
Participant flow	13a	For each group, the numbers of participants who	6	
(a diagram is		were randomly assigned, received intended		
strongly		treatment, and were analysed for the primary		
recommended)		outcome		
	13b	For each group, losses and exclusions after	Figure 1	
		randomisation, together with reasons		
Recruitment	14a	Dates defining the periods of recruitment and	4	
		follow-up		
	14b	Why the trial ended or was stopped		

howing baseline demographic and haracteristics for each group group, number of participants	Table 1	
group, number of participants		
	Tables 2-4	
nator) included in each analysis and		
the analysis was by original assigned		
primary and secondary outcome, results	6-7, Tables	
group, and the estimated effect size and	2-4	
ion (such as 95% confidence interval)		
ry outcomes, presentation of both	N/A	
and relative effect sizes is recommended		
of any other analyses performed,	N/A	
subgroup analyses and adjusted		
s, distinguishing pre-specified from		
ory		
tant harms or unintended effects in each	N/A	
or specific guidance see CONSORT for		
tations, addressing sources of potential	8	
precision, and, if relevant, multiplicity of		
•		
sability (external validity, applicability) of	8	
indings		
ation consistent with results, balancing	7-8	
and harms, and considering other		
evidence		
tion number and name of trial registry	2	
ne full trial protocol can be accessed, if	2, 4	
,		
of funding and other support (such as	9	
f drugs), role of funders		
	a primary and secondary outcome, results group, and the estimated effect size and sion (such as 95% confidence interval) ry outcomes, presentation of both and relative effect sizes is recommended of any other analyses performed, g subgroup analyses and adjusted s, distinguishing pre-specified from cry tant harms or unintended effects in each or specific guidance see CONSORT for specific guidance see CONSORT for stations, addressing sources of potential precision, and, if relevant, multiplicity of sability (external validity, applicability) of sindings ation consistent with results, balancing and harms, and considering other evidence tion number and name of trial registry me full trial protocol can be accessed, if so of funding and other support (such as f drugs), role of funders	primary and secondary outcome, results group, and the estimated effect size and sion (such as 95% confidence interval) ry outcomes, presentation of both and relative effect sizes is recommended of any other analyses performed, subgroup analyses and adjusted so, distinguishing pre-specified from any other analyses are CONSORT for specific guidance see CONSORT for specific guidance see CONSORT for sability (external validity, applicability) of sindings ation consistent with results, balancing and harms, and considering other evidence tion number and name of trial registry 2 ne full trial protocol can be accessed, if and of trianding and other support (such as 9)

*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 www.consort-statement.org.