Section/Topic	ltem	Standard Checklist item	Extension for cluster	Page
	No		designs	No *
Title and abstract				
	1a	Identification as a	Identification as a cluster	1
		randomised trial in the title	randomised trial in the title	
	1b	Structured summary of trial	See table 2	2-3
		design, methods, results, and		
		conclusions (for specific guidance see CONSORT for		
		abstracts) ^{1,2}		
Introduction				
Background and	2a	Scientific background and	Rationale for using a cluster	3-5
objectives	20	explanation of rationale	design	5-5
objectives		explanation of fationale		
	2b	Specific objectives or	Whether objectives pertain to the	5-6
		hypotheses	the cluster level, the individual	
			participant level or both	
Methods				
Trial design	3a	Description of trial design	Definition of cluster and	6
		(such as parallel, factorial)	description of how the design	
		including allocation ratio	features apply to the clusters	
	3b	Important changes to		n/a
		methods after trial		
		commencement (such as		
		eligibility criteria), with		
		reasons		
Participants	4a	Eligibility criteria for	Eligibility criteria for clusters	6
		participants		
	4b	Settings and locations where		6
		the data were collected		
Interventions	5	The interventions for each	Whether interventions pertain to	7-8
	5	group with sufficient details	the cluster level, the individual	
		to allow replication,	participant level or both	
		including how and when they	• • • • • • • • • • • • • • • • • • • •	
		were actually administered		
Outcomes	6a	Completely defined pre-	Whether outcome measures	8-10
		specified primary and	pertain to the cluster level, the	
		secondary outcome	individual participant level or both	
		measures, including how and		

Table 1: CONSORT 2010 checklist of information to include when reporting a cluster randomised trial

		when they were assessed		
	6b	Any changes to trial outcomes after the trial commenced, with reasons		n/a
Sample size	7a	How sample size was determined	Method of calculation, number of clusters(s) (and whether equal or unequal cluster sizes are assumed), cluster size, a coefficient of intracluster correlation (ICC or <i>k</i>), and an indication of its uncertainty	7
	7b	When applicable, explanation of any interim analyses and stopping guidelines		n/a
Randomisation:				
Sequence generation	8a	Method used to generate the random allocation sequence		6
	8b	Type of randomisation; details of any restriction (such as blocking and block size)	Details of stratification or matching if used	6
Allocation concealment mechanism	9	Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned	Specification that allocation was based on clusters rather than individuals and whether allocation concealment (if any) was at the cluster level, the individual participant level or both	6
Implementation	10	Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions	Replace by 10a, 10b and 10c	
	10a		Who generated the random allocation sequence, who enrolled clusters, and who assigned clusters to interventions	6
	10b		Mechanism by which individual participants were included in clusters for the purposes of the trial (such as complete	6

enumeration, random sampling) 10c From whom consent was sought 6-7 (representatives of the cluster, or individual cluster members, or both), and whether consent was sought before or after randomisation	
(representatives of the cluster, or individual cluster members, or both), and whether consent was sought before or after	
Blinding 11a If done, who was blinded n/a after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how	
11bIf relevant, description of then/asimilarity of interventions	
Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes How clustering was taken into 11	
12bMethods for additional11analyses, such as subgroupanalyses and adjustedanalysesanalyses	
Results	
Participant flow (a diagram is strongly recommended)13aFor each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcomeFor each group, the numbers of clusters that were randomly assigned, received intended treatment, and were analysed for the primary outcome6, 12	
13bFor each group, losses and exclusions afterFor each group, losses and exclusions for both clusters and individual cluster members reasons12	
Recruitment 14a Dates defining the periods of recruitment and follow-up 6	
14bWhy the trial ended or was7stopped	
Baseline data 15 A table showing baseline Baseline characteristics for the 12 (term demographic and clinical individual and cluster levels as	xt)

		characteristics for each group	applicable for each group	
Numbers analysed	16	For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups	For each group, number of clusters included in each analysis	12
Outcomes and estimation	17a	For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)	Results at the individual or cluster level as applicable and a coefficient of intracluster correlation (ICC or k) for each primary outcome	13-14; Table 2-4
	17b	For binary outcomes, presentation of both absolute and relative effect sizes is recommended		n/a
Ancillary analyses	18	Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory		13-14; Table 3-4
Harms	19	All important harms or unintended effects in each group (for specific guidance see CONSORT for harms ³)		n/a
Discussion				
Limitations	20	Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses		29
Generalisability	21	Generalisability (external validity, applicability) of the trial findings	Generalisability to clusters and/or individual participants (as relevant)	29
Interpretation	22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence		23-28
Other information				
Registration	23	Registration number and		3

		name of trial registry	
Protocol	24	Where the full trial protocol can be accessed, if available	n/a
Funding	25	Sources of funding and other support (such as supply of drugs), role of funders	30

* Note: page numbers optional depending on journal requirements

Table 2: Extension of CONSORT for abstracts1² to reports of cluster randomised trials

Item	Standard Checklist item	Extension for cluster trials
Title	Identification of study as randomised	Identification of study as cluster randomised
Trial design	Description of the trial design (e.g. parallel, cluster, non-inferiority)	
Methods		
Participants	Eligibility criteria for participants and the settings where the data were collected	Eligibility criteria for clusters
Interventions	Interventions intended for each group	
Objective	Specific objective or hypothesis	Whether objective or hypothesis pertains to the cluster level, the individual participant level or both
Outcome	Clearly defined primary outcome for this report	Whether the primary outcome pertains to the cluster level, the individual participant level or both
Randomization	How participants were allocated to interventions	How clusters were allocated to interventions
Blinding (masking)	Whether or not participants, care givers, and those assessing the outcomes were blinded to group assignment	
Results		
Numbers randomized	Number of participants randomized to each group	Number of clusters randomized to each group
Recruitment	Trial status ¹	
Numbers analysed	Number of participants analysed in each group	Number of clusters analysed in each group
Outcome	For the primary outcome, a result for each group and the estimated effect size and its precision	Results at the cluster or individual participant level as applicable for each primary outcome
Harms	Important adverse events or side effects	
Conclusions	General interpretation of the results	
Trial registration	Registration number and name of trial register	
Funding	Source of funding	

¹ Relevant to Conference Abstracts

REFERENCES

- ¹ Hopewell S, Clarke M, Moher D, Wager E, Middleton P, Altman DG, et al. CONSORT for reporting randomised trials in journal and conference abstracts. *Lancet* 2008, 371:281-283
- ² Hopewell S, Clarke M, Moher D, Wager E, Middleton P, Altman DG at al (2008) CONSORT for reporting randomized controlled trials in journal and conference abstracts: explanation and elaboration. *PLoS Med* 5(1): e20
- ³ Ioannidis JP, Evans SJ, Gotzsche PC, O'Neill RT, Altman DG, Schulz K, Moher D. Better reporting of harms in randomized trials: an extension of the CONSORT statement. *Ann Intern Med* 2004; 141(10):781-788.