### RESEARCH



**Open Access** 

# CONSORT 2010 Statement: updated guidelines for reporting parallel group randomised trials

Kenneth F Schulz<sup>1\*</sup>, Douglas G Altman<sup>2</sup>, David Moher<sup>3</sup>, the CONSORT Group

### Abstract

The CONSORT statement is used worldwide to improve the reporting of randomised controlled trials. Kenneth Schulz and colleagues describe the latest version, CONSORT 2010, which updates the reporting guideline based on new methodological evidence and accumulating experience.

To encourage dissemination of the CONSORT 2010 Statement, this article is freely accessible on bmj.com and will also be published in the Lancet, Obstetrics and Gynecology, PLoS Medicine, Annals of Internal Medicine, Open Medicine, Journal of Clinical Epidemiology, BMC Medicine, and Trials.

### Introduction

Randomised controlled trials, when appropriately designed, conducted, and reported, represent the gold standard in evaluating healthcare interventions. However, randomised trials can yield biased results if they lack methodological rigour [1]. To assess a trial accurately, readers of a published report need complete, clear, and transparent information on its methodology and findings. Unfortunately, attempted assessments frequently fail because authors of many trial reports neglect to provide lucid and complete descriptions of that critical information [2-4].

That lack of adequate reporting fuelled the development of the original CONSORT (Consolidated Standards of Reporting Trials) statement in 1996 [5] and its revision five years later [6-8]. While those statements improved the reporting quality for some randomised controlled trials, [9,10] many trial reports still remain inadequate [2]. Furthermore, new methodological evidence and additional experience has accumulated since the last revision in 2001. Consequently, we organised a CONSORT Group meeting to update the 2001 statement [6-8]. We introduce here the result of that process, CONSORT 2010.

### Intent of CONSORT 2010

The CONSORT 2010 Statement is this paper including the 25 item checklist in the table (Table 1) and the flow

\* Correspondence: kschulz@fhi.org

diagram (Figure 1). It provides guidance for reporting all randomised controlled trials, but focuses on the most common design type-individually randomised, two group, parallel trials. Other trial designs, such as cluster randomised trials and non-inferiority trials, require varying amounts of additional information. CONSORT extensions for these designs, [11,12] and other CON-SORT products, can be found through the CONSORT website http://www.consort-statement.org. Along with the CONSORT statement, we have updated the explanation and elaboration article, [13] which explains the inclusion of each checklist item, provides methodological background, and gives published examples of transparent reporting.

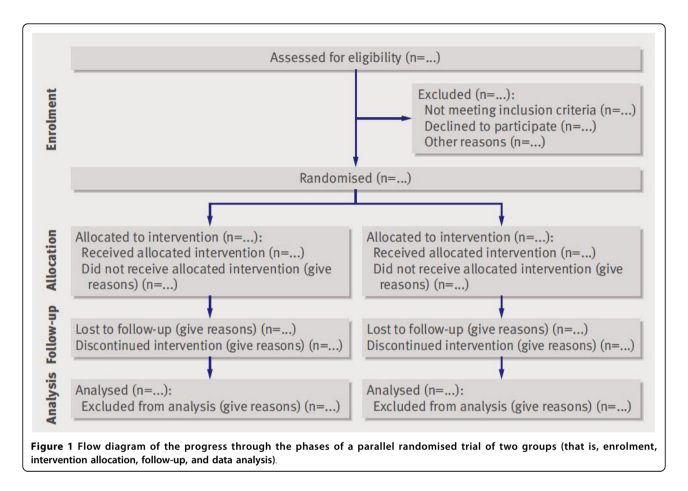
Diligent adherence by authors to the checklist items facilitates clarity, completeness, and transparency of reporting. Explicit descriptions, not ambiguity or omission, best serve the interests of all readers. Note that the CONSORT 2010 Statement does not include recommendations for designing, conducting, and analysing trials. It solely addresses the reporting of what was done and what was found.

Nevertheless, CONSORT does indirectly affect design and conduct. Transparent reporting reveals deficiencies in research if they exist. Thus, investigators who conduct inadequate trials, but who must transparently report, should not be able to pass through the publication process without revelation of their trial's inadequacies. That emerging reality should provide impetus to improved trial design and conduct in the future, a



© 2010 Schulz et al; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

<sup>&</sup>lt;sup>1</sup>Family Health International, Research Triangle Park, NC 27709, USA



secondary indirect goal of our work. Moreover, CON-SORT can help researchers in designing their trial.

### **Background to CONSORT**

Efforts to improve the reporting of randomised controlled trials accelerated in the mid-1990s, spurred partly by methodological research. Researchers had shown for many years that authors reported such trials poorly, and empirical evidence began to accumulate that some poorly conducted or poorly reported aspects of trials were associated with bias [14]. Two initiatives aimed at developing reporting guidelines culminated in one of us (DM) and Drummond Rennie organising the first CONSORT statement in 1996 [5]. Further methodological research on similar topics reinforced earlier findings [15] and fed into the revision of 2001 [6-8]. Subsequently, the expanding body of methodological research informed the refinement of CONSORT 2010. More than 700 studies comprise the CONSORT database (located on the CONSORT website), which provides the empirical evidence to underpin the CONSORT initiative.

Indeed, CONSORT Group members continually monitor the literature. Information gleaned from these

efforts provides an evidence base on which to update the CONSORT statement. We add, drop, or modify items based on that evidence and the recommendations of the CONSORT Group, an international and eclectic group of clinical trialists, statisticians, epidemiologists, and biomedical editors. The CONSORT Executive (KFS, DGA, DM) strives for a balance of established and emerging researchers. The membership of the group is dynamic. As our work expands in response to emerging projects and needed expertise, we invite new members to contribute. As such, CONSORT continually assimilates new ideas and perspectives. That process informs the continually evolving CONSORT statement.

Over time, CONSORT has garnered much support. More than 400 journals, published around the world and in many languages, have explicitly supported the CONSORT statement. Many other healthcare journals support it without our knowledge. Moreover, thousands more have implicitly supported it with the endorsement of the CONSORT statement by the International Committee of Medical Journal Editors http://www.icmje.org. Other prominent editorial groups, the Council of Science Editors and the World Association of Medical Editors, officially support CONSORT. That support seems warranted: when used by authors and journals, CONSORT seems to improve reporting [9].

### **Development of CONSORT 2010**

Thirty one members of the CONSORT 2010 Group met in Montebello, Canada, in January 2007 to update the 2001 CONSORT statement. In addition to the accumulating evidence relating to existing checklist items, several new issues had come to prominence since 2001. Some participants were given primary responsibility for aggregating and synthesising the relevant evidence on a particular checklist item of interest. Based on that evidence, the group deliberated the value of each item. As in prior CONSORT versions, we kept only those items deemed absolutely fundamental to reporting a randomised controlled trial. Moreover, an item may be fundamental to a trial but not included, such as approval by an institutional ethical review board, because funding bodies strictly enforce ethical review and medical journals usually address reporting ethical review in their instructions for authors. Other items may seem desirable, such as reporting on whether on-site monitoring was done, but a lack of empirical evidence or any consensus on their value cautions against inclusion at this point. The CONSORT 2010 Statement thus addresses the minimum criteria, although that should not deter authors from including other information if they consider it important.

After the meeting, the CONSORT Executive convened teleconferences and meetings to revise the checklist. After seven major iterations, a revised checklist was distributed to the larger group for feedback. With that feedback, the executive met twice in person to consider all the comments and to produce a penultimate version. That served as the basis for writing the first draft of this paper, which was then distributed to the group for feedback. After consideration of their comments, the executive finalised the statement.

The CONSORT Executive then drafted an updated explanation and elaboration manuscript, with assistance from other members of the larger group. The substance of the 2007 CONSORT meeting provided the material for the update. The updated explanation and elaboration manuscript was distributed to the entire group for additions, deletions, and changes. That final iterative process converged to the CONSORT 2010 Explanation and Elaboration [13].

### Changes in CONSORT 2010

The revision process resulted in evolutionary, not revolutionary, changes to the checklist (Table 1), and the flow diagram was not modified except for one word (Figure 1). Moreover, because other reporting guidelines augmenting the checklist refer to item numbers, we kept the existing items under their previous item numbers except for some renumbering of items 2 to 5. We added additional items either as a sub-item under an existing item, an entirely new item number at the end of the checklist, or (with item 3) an interjected item into a renumbered segment. We have summarised the noteworthy general changes in Appendix 1 and specific changes in Appendix 2. The CONSORT website contains a side by side comparison of the 2001 and 2010 versions.

### Implications and limitations

We developed CONSORT 2010 to assist authors in writing reports of randomised controlled trials, editors and peer reviewers in reviewing manuscripts for publication, and readers in critically appraising published articles. The CONSORT 2010 Explanation and Elaboration provides elucidation and context to the checklist items. We strongly recommend using the explanation and elaboration in conjunction with the checklist to foster complete, clear, and transparent reporting and aid appraisal of published trial reports.

CONSORT 2010 focuses predominantly on the two group, parallel randomised controlled trial, which accounts for over half of trials in the literature [2]. Most of the items from the CONSORT 2010 Statement, however, pertain to all types of randomised trials. Nevertheless, some types of trials or trial situations dictate the need for additional information in the trial report. When in doubt, authors, editors, and readers should consult the CONSORT website for any CONSORT extensions, expansions (amplifications), implementations, or other guidance that may be relevant.

The evidence based approach we have used for CON-SORT also served as a model for development of other reporting guidelines, such as for reporting systematic reviews and meta-analyses of studies evaluating interventions [16], diagnostic studies [17], and observational studies [18]. The explicit goal of all these initiatives is to improve reporting. The Enhancing the Quality and Transparency of Health Research (EQUATOR) Network will facilitate development of reporting guidelines and help disseminate the guidelines: http://www.equatornetwork.org provides information on all reporting guidelines in health research.

With CONSORT 2010, we again intentionally declined to produce a rigid structure for the reporting of randomised trials. Indeed, SORT [19] tried a rigid format, and it failed in a pilot run with an editor and authors [20]. Consequently, the format of articles should abide by journal style, editorial directions, the traditions of the research field addressed, and, where possible, author preferences. We do not wish to standardise the structure of reporting. Authors should simply address checklist

Title and abstract   1a   Identification as a randomised trial in the title     Incoduction   Bit Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts [21,31])     Incoduction   2a   Scientific background and explanation of rationale     2b   Specific objectives or hypotheses     Methods   3a   Description of trial design (such as parallel, factorial) including allocation ratio     3b   important changes to methods after trial commencement (such as eligibility criteria), with reasons     Participants   4a   Eligibility criteria for participants     4b   Settings and locations where the data were collected     Interventions   The interventions for each group with sufficient details to allow replication, including how and when they were assessed     6b   Any changes to trial doctions after their trial commenced, with reasons     Sample size   7a   How sample size was determined     7b   When applicable, explanation of any interim analyses and stopping guidelines     Randomisation:   Sequence generation   8a     Sequence generation   8a   Method used to generate the random allocation sequence (such as sequentilly numbered containes), describing any steps taken to conceal the sequence until intereventions were	Section/Topic	ltem No	Checklist item	Reported on page No
Introduction Exactured summay of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts [2],31])   Introduction 2a Scientific background and explanation of rationale   Background and objectives 2a Scientific background and explanation of rationale   Methods Escription of trial design (such as parallel, factorial) including allocation ratio   Trial design 3a Description of trial design (such as parallel, factorial) including allocation, ratio   Participants 4a Eligibility criteria for participants   Participants 4a Eligibility criterial explanation software collected   Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were assessed   Outcomes 6 Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   Sample size 7a How sample size was determined   Sequence generation 8a Method used to generate the random allocation sequence (such as sequentify numbered containers), description of rationers allocation sequence (such as sequentify numbered containers), description of any interim analyses and stopping guidelines   Balandomisation: 29 Methanismus defined after assignment to interventions (for example, participants, care providers, those assessin	Title and abstract			
selectron and objectives   29   Scientific background and explanation of rationale     Beckground and objectives   29   Specific objectives or hypotheses     Methods		1a	Identification as a randomised trial in the title	
Background and objectives 2a Scientific background and explanation of rationale   2b Specific objectives or hypotheses   Methods 3a Description of trial design (such as parallel, factorial) including allocation ratio   1al design 3a Description of trial design (such as parallel, factorial) including allocation ratio   Participants 4a Eligibility criteria for participants   4b Settings and locations where the data were collected   Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were actually administered   Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were actually administered   Sample size 7a How sample size was determined   7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: 8a Method used to generate the random allocation sequence (such as sequentially numbered containers), description of the similary of interventions for example, participants, and who assigned   Implementation 10 Who generated the random allocation sequence (such as blocking and block size)   Mactains 11 I done, who was blinded after assignment to interventions (for example, participants, care providers, those a		1b		
Wethods   Trial design 3a Description of trial design (such as parallel, factorial) including allocation ratio   Trial design 3a Description of trial design (such as parallel, factorial) including allocation ratio   Participants 4a Eligibility criteria for participants   4b Settings and locations where the data were collected   Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were actually administered   Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   Sample size 7a How sample size was determined   Yue constraint 7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: Sa Method used to generate the random allocation sequence.   Sequence generation 8a Method secription of the ismilarity of interventions   Implementation 10 Who generated the random allocation sequence until interventions were assigned   Implementation 11a if done, who was blinded after assignment to interventions	Introduction			
Methods   Trial design   3a   Description of trial design (such as parallel, factorial) including allocation ratio     Trial design   3a   Description of trial design (such as parallel, factorial) including allocation ratio     Barticipants   4a   Eligibility criteria for participants     4b   Settings and locations where the data were collected     Interventions   5   The interventions for each group with sufficient details to allow replication, including how and when they were actually administered     Outcomes   6a   Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed     Sample size   7b   When applicable, explanation of any interim analyses and stopping guidelines     Sample size   7b   When applicable, explanation of any interim analyses and stopping guidelines     Randomisation:   8a   Method used to generate the random allocation sequence (such as sequentially numbered containers), descripting any steps taken to conceal the sequence until interventions were assigned     Implementation   10   Who generate the random allocation sequence (such as sequence until interventions (for example, participants, care providers, those assessing outcomes) and how     Bilinding   11a   f done, who was blinded after assignment to interventions     10 <td< td=""><td>Background and objectives</td><td>2a</td><td>Scientific background and explanation of rationale</td><td></td></td<>	Background and objectives	2a	Scientific background and explanation of rationale	
Trial design 3a Description of trial design (such as parallel, factorial) including allocation ratio   Participants 4a Eligibility criteria for participants   Participants 4a Eligibility criteria for participants   Networts 5a The interventions for each group with sufficient details to allow replication, including how and when they were actually administered   Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were assesd   Sample size 7a How sample size was determined   Sequence generation 8a Method used to generate the random allocation sequence   Sequence generation 8a Nethod used to generate the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned   Implementation 10 Who generated the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions   Statistical methods 11b If relevant, description of the similarity of interventions   Statistical methods 2a Statistical methods used to compare groups for primary and secondary outcomes intended treatment, and were analysed for the primary outcome   Blinding 11b If relevant, description of the		2b	Specific objectives or hypotheses	
3b Important changes to methods after trial commencement (such as eligibility criteria), with reasons   Participants 4 Eligibility criteria for participants   4 Eligibility criteria for participants 5   45 Settings and locations where the data were collected   Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were assessed   Outcomes 6 Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   Sample size 7 How sample size was determined   Saupe size 7 How sample size was determined   Sequence generation 8 Method used to generate the random allocation sequence   Biolacation concealment mechanism 9 Mechanism used to implement the random allocation sequence (such as sequence until interventions were assigned   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Statistical methods 12 Statistical methods used to compare groups for primary and secondary outcomes   Binding 11a If done, who was blinded after assignment to interventions   Statistical methods 12 Statistical methods u				
Participants 4a Eligibility criteria for participants Settings and locations where the data were collected Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were actually administered Outcomes 6 Completely defined per-specified primary and secondary outcome measures, including how and when they were assessed 6 Any changes to trial outcomes after the trial commenced, with reasons Sample size 7 How sample size was determined 7 b When applicable, explanation of any interim analyses and stopping guidelines Randomisation: 8 Method used to generate the random allocation sequence 8 Method used to generate the random allocation sequence 8 Method used to generate the random allocation sequence 8 Method used to generate the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned 1 Method used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned 1 Method used to implement to interventions 1 f done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how 1 Ib If relevant, description of the similarity of interventions 1 Statistical methods 1 Statistical methods 1 2 Statistical methods used to compare groups for primary and secondary outcomes 1 Methods for additional analyses, such as subgroup analyses and adjusted analyses 8 Results 8 Results 8 Recruitment 14 Dates defining the periods of recruitment and follow-up 14 Why the trial ended or was stopped 8 Baseline data 15 A table showing baseline demographic and clinical characteristics for each group, 16 For each group, losses and exclusions after randominiation, together with reasons 8 Cortice and estimation 17 For each proup, muther of participants (denominator) included in each a	Trial design			
4b Settings and locations where the data were collected   Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were actually administered   Outcomes Ga Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   Sample size 7a How sample isze was determined   7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: #a Method used to generate the random allocation sequence   8a Method used to generate the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until numbered containers), describing any steps taken to conceal the sequence until numbered containers), describing any steps taken to conceal the sequence until numbered containers), describing any steps taken to conceal the sequence until numbered containers), describing any steps taken to conceal the sequence until numbered containers), describing any steps taken to conceal the sequence until numbered containers, description of the similarity of interventions   Statistical methods 10 Who generated the random allocation sequence, who enrolled participants, care providers, those assessing outcomes) and how   Statistical methods 11 If cleavant, description of the similarity of interventions   Statistical methods 23 Setatistical methods used to compare groups for		3b	reasons	
Interventions 5 The interventions for each group with sufficient details to allow replication, including how and when they were actually administered   Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   6b Any changes to trial outcomes after the trial commenced, with reasons   Sample size 7a How sample size was determined   7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: Sequence generation 8a   8a Method used to generate the random allocation sequence (such as sequentially numbered containers), 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   Bilinding 11 If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes   Statistical methods 13a For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome   Statistical methods 15a For each group, losses and exclusions after	Participants	4a	Eligibility criteria for participants	
Outcomes 6a Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed   6b Any changes to trial outcomes after the trial commenced, with reasons   Sample size 7a How sample size was determined   7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: Sequence generation 8a   8b Type of randomisation; details of any restriction (such as blocking and block size)   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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Bilnding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   12b Kretstistical methods 12a Statistical methods used for additional analyses, such as subgroup analyses and adjusted analyses   Results Participant flow (a diagram is strongly recommended) 13a For each group, then numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome   12b Wethods		4b	Settings and locations where the data were collected	
how and when they were assessed 6b Any changes to trial outcomes after the trial commenced, with reasons 5ample size 7a How sample size was determined 7b When applicable, explanation of any interim analyses and stopping guidelines Randomisation: Sequence generation 8a Method used to generate the random allocation sequence 8b Type of randomisation; details of any restriction (such as blocking and block size) 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 Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how 11b If relevant, description of the similarity of interventions Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes 12b Methods for additional analyses, such as subgroup analyses and adjusted analyses <b>Results</b> Recruitment 13a For each group, the numbers of participants who were randomily assigned, received intended treatment, and were analysed for the primary outcome 13b For each group, losses and exclusions after randomisation, together with reasons Recruitment 14a Dates defining the periods of recruitment and follow-up 14b Why the trial ended or was stopped Baseline data 15 A table showing baseline demographic and clinical characteristics 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 Outcomes and estimation 17a For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups	Interventions	5		
Sample size 7a How sample size was determined   7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: 5   Sequence generation 8a Method used to generate the random allocation sequence   Allocation concealment mechanism 7b When applicable, explanation; details of any restriction (such as blocking and block size)   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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes   Statistical methods 12a Statistical methods used for the primary outcome   Storogly recommended) 13a For each group, the numbers of participants who were randomly assigned, received interded treatment, and were analysed for the primary outcome   Recruitment 14a Dates defining the periods of recruitment and follow-up   13	Outcomes	ба		
7b When applicable, explanation of any interim analyses and stopping guidelines   Randomisation: Sequence generation 8a Method used to generate the random allocation sequence   8b Type of randomisation; details of any restriction (such as blocking and block size)   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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   12b If elevant, description of the similarity of interventions   Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes intended treatment, and were analysed for the primary outcome   12b Methods treatment, and were analysed for the primary outcome   13b For each group, losses and exclusions after randomisation, together with reasons   Recruitment 14a Dates defining the periods of recruitment and follow-up   14b Why the trial ended or was stopped 14b   Baseline data 15 <		6b	Any changes to trial outcomes after the trial commenced, with reasons	
Randomisation: Sequence generation 8a Method used to generate the random allocation sequence   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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   11b If devent, description of the similarity of interventions   Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes   Results Participant flow (a diagram is strongly recommended) 13a For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome   13b For each group, losses and exclusions after randomisation, together with reasons   Recruitment 14a Dates defining the periods of recruitment and follow-up   14b Why the trial ended or was stopped Baseline data   15 A table showing baseline demographic and clinical characteristics for each group, Numbers analysed For each group, number of participants (denominator) incl	Sample size	7a	How sample size was determined	
Sequence generation 8a Method used to generate the random allocation sequence   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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   11b If relevant, description of the similarity of interventions   Statistical methods 12a Statistical methods used to random allocation sequence for primary and secondary outcomes   Results 13a For each group, the numbers of participants who were randomly assigned, received interded treatment, and were analysed for the primary outcome   13b For each group, losses and exclusions after randomisation, together with reasons   Recruitment 14a Dates defining the periods of recruitment and follow-up   14b Why the trial ended or was stopped   Baseline data 15 A table showing baseline demographic and clinical characteristics for each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups)   Outcomes and estim		7b	When applicable, explanation of any interim analyses and stopping guidelines	
8b Type of randomisation; details of any restriction (such as blocking and block size)   Allocation concealment 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   Implementation 10 Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions   Blinding 11a If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how   11b If relevant, description of the similarity of interventions   Statistical methods 12a   Statistical methods 12a   Statistical methods 13a   For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome   13b For each group, losses and exclusions after randomisation, together with reasons   Recruitment 14a Dates defining the periods of recruitment and follow-up   14b Why the trial ended or was stopped Baseline data 15 A table showing baseline demographic and clinical characteristics for each group   Numbers analysed 16 For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assign	Randomisation:			
Allocation concealment mechanism9Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assignedImplementation10Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventionsBlinding11aIf done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how 11bStatistical methods12aStatistical methods used to compare groups for primary and secondary outcomes 12bResultsParticipant flow (a diagram is strongly recommended)For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcomeRecultment14aDates defining the periods of recruitment and follow-up 14bNumbers analysed15A table showing baseline demographic and clinical characteristics for each group whether the analysis was by original asigned groupsOutcomes and estimation17aFor each group, number of participants (denominator) included in each analysis and whether the analysis may by original asigned groupsOutcomes and estimation17aFor each group, outcome, presentation of both absolute and relative effect size is	Sequence generation	8a	Method used to generate the random allocation sequence	
mechanismnumbered containers), describing any steps taken to conceal the sequence until interventions were assignedImplementation10Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventionsBlinding11aIf done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and howBlinding11bIf relevant, description of the similarity of interventionsStatistical methods12aStatistical methods used to compare groups for primary and secondary outcomes 12bResults2Vethods for additional analyses, such as subgroup analyses and adjusted analysesRecuitment13aFor each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcomeRecruitment14aDates defining the periods of recruitment and follow-up 14bWhy the trial ended or was stoppedSateline data5A table showing baseline demographic and clinical characteristics for each group whether the analysis was by original assigned groupsOutcomes and estimation17aFor each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each group, spresentation of both absolute and relative effect sizes is		8b	Type of randomisation; details of any restriction (such as blocking and block size)	
Blinding11aIf done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how11bIf relevant, description of the similarity of interventionsStatistical methods12aStatistical methods used to compare groups for primary and secondary outcomes 12bResultsParticipant 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 outcomeRecruitment14aDates defining the periods of recruitment and follow-up 14bWhy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each group whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval) 17bFor binary outcomes, presentation of both absolute and relative effect sizes is		9	numbered containers), describing any steps taken to conceal the sequence until	
providers, those assessing outcomes) and how 11b If relevant, description of the similarity of interventions Statistical methods 12a Statistical methods used to compare groups for primary and secondary outcomes 12b Methods for additional analyses, such as subgroup analyses and adjusted analyses <b>Results</b> Participant flow (a diagram is strongly recommended) 13a For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome 13b For each group, losses and exclusions after randomisation, together with reasons Recruitment 14a Dates defining the periods of recruitment and follow-up 14b Why the trial ended or was stopped Baseline data 15 A table showing baseline demographic and clinical characteristics for each group Numbers analysed 0utcomes 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) 17b For binary outcomes, presentation of both absolute and relative effect sizes is	Implementation	10		
Statistical methods12aStatistical methods used to compare groups for primary and secondary outcomes 12bResultsParticipant 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 outcome 13bFor each group, losses and exclusions after randomisation, together with reasonsRecruitment14aDates defining the periods of recruitment and follow-up 14bWhy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each group whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval) 17bTorFor binary outcomes, presentation of both absolute and relative effect sizes is	Blinding	11a		
12bMethods for additional analyses, such as subgroup analyses and adjusted analysesResultsParticipant 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 outcome 13bRecruitment14aDates defining the periods of recruitment and follow-up 14bMy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each group whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval) 17bFor binary outcomes, presentation of both absolute and relative effect sizes is		11b	If relevant, description of the similarity of interventions	
Results 13a For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome   13b For each group, losses and exclusions after randomisation, together with reasons   Recruitment 14a Dates defining the periods of recruitment and follow-up   14b Why the trial ended or was stopped   Baseline data 15 A table showing baseline demographic and clinical characteristics 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   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)   17b For binary outcomes, presentation of both absolute and relative effect sizes is	Statistical methods	12a	Statistical methods used to compare groups for primary and secondary outcomes	
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 outcome13bFor each group, losses and exclusions after randomisation, together with reasonsRecruitment14aDates defining the periods of recruitment and follow-up14bWhy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each groupNumbers analysed16For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is		12b	Methods for additional analyses, such as subgroup analyses and adjusted analyses	
strongly recommended)intended treatment, and were analysed for the primary outcome13bFor each group, losses and exclusions after randomisation, together with reasonsRecruitment14aDates defining the periods of recruitment and follow-up14bWhy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each groupNumbers analysed16For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is	Results			
Recruitment14aDates defining the periods of recruitment and follow-up14bWhy the trial ended or was stoppedBaseline data15Numbers analysed16For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is		13a		
14bWhy the trial ended or was stoppedBaseline data15A table showing baseline demographic and clinical characteristics for each groupNumbers analysed16For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is		13b	For each group, losses and exclusions after randomisation, together with reasons	
Baseline data 15 A table showing baseline demographic and clinical characteristics 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   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)   17b For binary outcomes, presentation of both absolute and relative effect sizes is	Recruitment	14a	Dates defining the periods of recruitment and follow-up	
Numbers analysed16For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groupsOutcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is		14b	Why the trial ended or was stopped	
Outcomes and estimation17aFor each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)17bFor binary outcomes, presentation of both absolute and relative effect sizes is	Baseline data	15	A table showing baseline demographic and clinical characteristics for each group	
effect size and its precision (such as 95% confidence interval) 17b For binary outcomes, presentation of both absolute and relative effect sizes is	Numbers analysed	16		
	Outcomes and estimation	17a		
recommended		17b		
Ancillary analyses 18 Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory	Ancillary analyses	18		
Harms 19 All important harms or unintended effects in each group (for specific guidance see CONSORT for harms [28])	Harms	19	All important harms or unintended effects in each group (for specific guidance see	
Discussion	Discussion			
Limitations 20 Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses		20		
Generalisability21Generalisability (external validity, applicability) of the trial findings	Generalisability	21		

### Table 1 CONSORT 2010 checklist of information to include when reporting a randomised trial\*

Interpretation	22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence
Other information		
Registration	23	Registration number and name of trial registry
Protocol	24	Where the full trial protocol can be accessed, if available
Funding	25	Sources of funding and other support (such as supply of drugs), role of funders

Table 1: CONSORT 2010 checklist of information to include wh	<b>nen reporting a randomised trial</b> * (Continued)
--	---

\*We strongly recommend reading this statement in conjunction with the CONSORT 2010 Explanation and Elaboration [13] for important clarifications on all the items. If relevant, we also recommend reading CONSORT extensions for cluster randomised trials [11], non-inferiority and equivalence trials [12], non-pharmacological treatments [32], herbal interventions [33], and pragmatic trials [34]. Additional extensions are forthcoming: for those and for up to date references relevant to this checklist, see http://www.consort-statement.org/.

items somewhere in the article, with ample detail and lucidity. That stated, we think that manuscripts benefit from frequent subheadings within the major sections, especially the methods and results sections.

CONSORT urges completeness, clarity, and transparency of reporting, which simply reflects the actual trial design and conduct. However, as a potential drawback, a reporting guideline might encourage some authors to report fictitiously the information suggested by the guidance rather than what was actually done. Authors, peer reviewers, and editors should vigilantly guard against that potential drawback and refer, for example, to trial protocols, to information on trial registers, and to regulatory agency websites. Moreover, the CONSORT 2010 Statement does not include recommendations for designing and conducting randomised trials. The items should elicit clear pronouncements of how and what the authors did, but do not contain any judgments on how and what the authors should have done. Thus, CON-SORT 2010 is not intended as an instrument to evaluate the quality of a trial. Nor is it appropriate to use the checklist to construct a "quality score."

Nevertheless, we suggest that researchers begin trials with their end publication in mind. Poor reporting allows authors, intentionally or inadvertently, to escape scrutiny of any weak aspects of their trials. However, with wide adoption of CONSORT by journals and editorial groups, most authors should have to report transparently all important aspects of their trial. The ensuing scrutiny rewards well conducted trials and penalises poorly conducted trials. Thus, investigators should understand the CONSORT 2010 reporting guidelines before starting a trial as a further incentive to design and conduct their trials according to rigorous standards.

CONSORT 2010 supplants the prior version published in 2001. Any support for the earlier version accumulated from journals or editorial groups will automatically extend to this newer version, unless specifically requested otherwise. Journals that do not currently support CONSORT may do so by registering on the CON-SORT website. If a journal supports or endorses CONSORT 2010, it should cite one of the original versions of CONSORT 2010, the CONSORT 2010 Explanation and Elaboration, and the CONSORT website in their "Instructions to authors." We suggest that authors who wish to cite CONSORT should cite this or another of the original journal versions of CONSORT 2010 Statement, and, if appropriate, the CONSORT 2010 Explanation and Elaboration [13]. All CONSORT material can be accessed through the original publishing journals or the CONSORT website. Groups or individuals who desire to translate the CONSORT 2010 Statement into other languages should first consult the CONSORT policy statement on the website.

We emphasise that CONSORT 2010 represents an evolving guideline. It requires perpetual reappraisal and, if necessary, modifications. In the future we will further revise the CONSORT material considering comments, criticisms, experiences, and accumulating new evidence. We invite readers to submit recommendations via the CONSORT website.

### Appendix 1: Noteworthy general changes in CONSORT 2010 Statement

• We simplified and clarified the wording, such as in items 1, 8, 10, 13, 15, 16, 18, 19, and 21

• We improved consistency of style across the items by removing the imperative verbs that were in the 2001 version

• We enhanced specificity of appraisal by breaking some items into sub-items. Many journals expect authors to complete a CONSORT checklist indicating where in the manuscript the items have been addressed. Experience with the checklist noted pragmatic difficulties when an item comprised multiple elements. For example, item 4 addresses eligibility of participants and the settings and locations of data collection. With the 2001 version, an author could provide a page number for that item on the checklist, but might have reported only eligibility in the paper, for example, and not reported the settings and locations. CONSORT 2010 relieves obfuscations and forces authors to provide page numbers in the checklist for both eligibility and settings.

## Appendix 2: Noteworthy specific changes in CONSORT 2010 Statement

*Item 1b (title and abstract)*-We added a sub-item on providing a structured summary of trial design, methods, results, and conclusions and referenced the CON-SORT for abstracts article [21].

*Item 2b (introduction)*-We added a new sub-item (formerly item 5 in CONSORT 2001) on "Specific objectives or hypotheses"

*Item 3a (trial design)*-We added a new item including this sub-item to clarify the basic trial design (such as parallel group, crossover, cluster) and the allocation ratio

*Item 3b (trial design)*-We added a new sub-item that addresses any important changes to methods after trial commencement, with a discussion of reasons

*Item 4 (participants)*-Formerly item 3 in CONSORT 2001

*Item 5 (interventions)*-Formerly item 4 in CONSORT 2001. We encouraged greater specificity by stating that descriptions of interventions should include "sufficient details to allow replication"[3]

*Item 6 (outcomes)*-We added a sub-item on identifying any changes to the primary and secondary outcome (endpoint) measures after the trial started. This followed from empirical evidence that authors frequently provide analyses of outcomes in their published papers that were not the prespecified primary and secondary outcomes in their protocols, while ignoring their prespecified outcomes (that is, selective outcome reporting) [4,22]. We eliminated text on any methods used to enhance the quality of measurements

*Item 9 (allocation concealment mechanism)*-We reworded this to include mechanism in both the report topic and the descriptor to reinforce that authors should report the actual steps taken to ensure allocation concealment rather than simply report imprecise, perhaps banal, assurances of concealment

*Item 11 (blinding)*-We added the specification of how blinding was done and, if relevant, a description of the similarity of interventions and procedures. We also eliminated text on "how the success of blinding (masking) was assessed" because of a lack of empirical evidence supporting the practice as well as theoretical concerns about the validity of any such assessment [23,24]

*Item 12a (statistical methods)*-We added that statistical methods should also be provided for analysis of secondary outcomes

*Sub-item 14b (recruitment)*-Based on empirical research, we added a sub-item on "Why the trial ended or was stopped" [25]

*Item 15 (baseline data)*-We specified "A table" to clarify that baseline and clinical characteristics of each group are most clearly expressed in a table *Item 16 (numbers analysed)*-We replaced mention of "intention to treat" analysis, a widely misused term, by a more explicit request for information about retaining participants in their original assigned groups [26]

Sub-item 17b (outcomes and estimation)-For appropriate clinical interpretability, prevailing experience suggested the addition of "For binary outcomes, presentation of both relative and absolute effect sizes is recommended" [27]

*Item 19 (harms)*-We included a reference to the CON-SORT paper on harms [28]

*Item 20 (limitations)*-We changed the topic from "Interpretation" and supplanted the prior text with a sentence focusing on the reporting of sources of potential bias and imprecision

*Item 22 (interpretation)*-We changed the topic from "Overall evidence." Indeed, we understand that authors should be allowed leeway for interpretation under this nebulous heading. However, the CONSORT Group expressed concerns that conclusions in papers frequently misrepresented the actual analytical results and that harms were ignored or marginalised. Therefore, we changed the checklist item to include the concepts of results matching interpretations and of benefits being balanced with harms

*Item 23 (registration)*-We added a new item on trial registration. Empirical evidence supports the need for trial registration, and recent requirements by journal editors have fostered compliance [29]

*Item 24 (protocol)*-We added a new item on availability of the trial protocol. Empirical evidence suggests that authors often ignore, in the conduct and reporting of their trial, what they stated in the protocol [4,22]. Hence, availability of the protocol can instigate adherence to the protocol before publication and facilitate assessment of adherence after publication

*Item 25 (funding)*-We added a new item on funding. Empirical evidence points toward funding source sometimes being associated with estimatedeatment effects [30]

### Acknowledgements

The CONSORT Group contributors to CONSORT 2010: DG Altman, Centre for Statistics in Medicine, University of Oxford; Virginia Barbour, *PLoS Medicine*; Jesse A Berlin, Johnson & Johnson Pharmaceutical Research and Development, USA; Isabelle Boutron, University Paris 7 Denis Diderot, Assistance Publique des Hôpitaux de Paris, INSERM, France; PJ Devereaux, McMaster University, Canada; Kay Dickersin, Johns Hopkins Bloomberg School of Public Health, USA; Diana Elbourne, London School of Hygiene & Tropical Medicine; Susan Ellenberg, University of Pennsylvania School of Medicine, USA; Val Gebski, University of Sydney, Australia; Steven Goodman, *Journal of the Society for Clinical Trials*, USA; Peter C Gøtzsche, Nordic Cochrane Centre, Denmark; Trish Groves, *BMJ*; Steven Grunberg, American Society of Clinical Oncology, USA; Brian Haynes, McMaster University, of Oxford; Astrid James, *Lancet*; Peter Juhn, Johnson & Johnson, USA; Philippa

Middleton, University of Adelaide, Australia; Don Minckler, University of California Irvine, USA; D Moher, Ottawa Methods Centre, Clinical Epidemiology Program, Ottawa Hospital Research Institute, Canada; Victor M Montori, Knowledge and Encounter Research Unit, Mayo Clinic College of Medicine, USA; Cynthia Mulrow, *Annals of Internal Medicine*, USA; Stuart Pocock, London School of Hygiene & Tropical Medicine, USA; KF Schulz, *JAMA*, USA; David L Schriger, *Annals of Emergency Medicine*, USA; KF Schulz, Family Health International, USA; Iveta Simera, EQUATOR Network; Elizabeth Wager, Sideview.

Contributors to CONSORT 2010 who did not attend the Montebello meeting: Mike Clarke, UK Cochrane Centre; Gordon Guyatt, McMaster University, Canada.

Funding: We received financial support from United Kingdom National Institute for Health Research and the Medical Research Council; Canadian Institutes of Health Research; Presidents Fund, Canadian Institutes of Health Research; Johnson; *BMJ*; and the American Society for Clinical Oncology. DGA is supported by Cancer Research UK, DM by a University of Ottawa Research Chair, and KFS by Family Health International. None of the sponsors had any involvement in the planning, execution, or writing of the CONSORT documents. Additionally, no funder played a role in drafting the manuscript.

In order to encourage dissemination of the CONSORT 2010 Statement, this article is freely accessible on bmj.com and will also be published in the Lancet, Obstetrics and Gynecology, PLoS Medicine, Annals of Internal Medicine, Open Medicine, Journal of Clinical Epidemiology, BMC Medicine, and Trials. The authors jointly hold the copyright of this article. For details on further use, see the CONSORT website http://www.consort-statement.org.

#### Author details

<sup>1</sup>Family Health International, Research Triangle Park, NC 27709, USA. <sup>2</sup>Centre for Statistics in Medicine, University of Oxford, Wolfson College, Oxford, UK. <sup>3</sup>Ottawa Methods Centre, Clinical Epidemiology Program, Ottawa Hospital Research Institute, Department of Epidemiology and Community Medicine, University of Ottawa, Ottawa, Canada.

#### Authors' contributions

KFS, DM, and DGA participated in meetings and regular conference calls, planned the CONSORT 2007 meeting at Montebello, developed the agenda, prepared background research, identified and invited participants, contributed to the CONSORT meeting, drafted the manuscript, and, after critical review by the CONSORT Group, finalised the text of the manuscript. Members of the CONSORT Group attended the meeting, except for those noted below, and provided input on and review of the revised checklist and text of this article. Some members also prepared background material.

### **Competing interests**

Uniform disclosure of potential conflicts of interest: all authors have completed the ICMJE unified competing interest form at http://www.icmje. org/coi\_disclosure.pdf (available from the corresponding author) and declare (1) DM received grants for this work from Johnson & Johnson, BMJ, and American Society for Clinical Oncology; KFS and DGA received support for travel to meetings for this work from Johnson & Johnson, BMJ, and American Society for Clinical Oncology; (2) KFS and DA had travel expenses reimbursed by the EQUATOR Network; KFS has received honoraria for delivering educational presentations for the American Board of Obstetrics and Gynecology Foundation for Excellence in Women's Health Care, Ortho-McNeii Janssen Scientific Affairs, and the American College of Obstetrics and Gynecology; and has done consultancy for Wyeth. All authors also declare (3) no spouses, partners, or children with relationships with commercial entities that might have an interest in the submitted work; (4) no nonfinancial interests that may be relevant to the submitted work.

### Received: 9 March 2010 Accepted: 24 March 2010 Published: 24 March 2010

#### References

- Jüni P, Altman DG, Egger M: Systematic reviews in health care: assessing the quality of controlled clinical trials. *BMJ* 2001, 323:42-6.
- 2. Chan AW, Altman DG: Epidemiology and reporting of randomised trials published in PubMed journals. *Lancet* 2005, **365**:1159-62.

- Glasziou P, Meats E, Heneghan C, Shepperd S: What is missing from descriptions of treatment in trials and reviews? BMJ 2008, 336:1472-4.
- Dwan K, Altman DG, Arnaiz JA, Bloom J, Chan AW, Cronin E, Decullier E, Easterbrook PJ, Von Elm E, Gamble C, Ghersi D, Ioannidis JP, Simes J, Williamson PR: Systematic review of the empirical evidence of study publication bias and outcome reporting bias. *PLoS ONE* 2008, 3:e3081.
- Begg C, Cho M, Eastwood S, Horton R, Moher D, Olkin I, Pitkin R, Rennie D, Schulz KF, Simel D, Stroup DF: Improving the quality of reporting of randomized controlled trials. The CONSORT statement. JAMA 1996, 276:637-9.
- Moher D, Schulz KF, Altman DG: The CONSORT statement: revised recommendations for improving the quality of reports of parallel-group randomised trials. *Lancet* 2001, 357:1191-4.
- Moher D, Schulz KF, Altman DG: The CONSORT statement: revised recommendations for improving the quality of reports of parallel-group randomized trials. *Ann Intern Med* 2001, 134:657-62.
- Moher D, Schulz KF, Altman D: The CONSORT statement: revised recommendations for improving the quality of reports of parallel-group randomized trials. JAMA 2001, 285:1987-91.
- Plint AC, Moher D, Morrison A, Schulz K, Altman DG, Hill C, Gaboury : Does the CONSORT checklist improve the quality of reports of randomised controlled trials? A systematic review. *Med J Aust* 2006, 185:263-7.
- Hopewell S, Dutton S, Yu L-M, Chan A-W, Altman DG: The quality of reports of randomised trials in 2000 and 2006: a comparative study of articles indexed by PubMed. *BNJ* 2010, 340:c723.
- 11. Campbell MK, Elbourne DR, Altman DG: CONSORT statement: extension to cluster randomised trials. *BMJ* 2004, **328**:702-8.
- Piaggio G, Elbourne DR, Altman DG, Pocock SJ, Evans SJ: Reporting of noninferiority and equivalence randomized trials: an extension of the CONSORT statement. JAMA 2006, 295:1152-60.
- Moher D, Hopewell S, Schulz KF, Montori V, Gøtzsche PC, Devereaux PJ, et al: CONSORT 2010 Explanation and Elaboration: updated guidelines for reporting parallel group randomised trials. *BMJ* 2010, 340:c869.
- Schulz KF, Chalmers I, Hayes RJ, Altman DG: Empirical evidence of bias. Dimensions of methodological quality associated with estimates of treatment effects in controlled trials. JAMA 1995, 273:408-12.
- Moher D, Pham B, Jones A, Cook DJ, Jadad AR, Moher M, Tugwell P, Klassen TP: Does quality of reports of randomised trials affect estimates of intervention efficacy reported in meta-analyses? *Lancet* 1998, 352:609-13.
- Moher D, Liberati A, Tetzlaff J, Altman DG, for the PRISMA Group: Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *BMJ* 2009, 339:b2535.
- Bossuyt PM, Reitsma JB, Bruns DE, Gatsonis CA, Glasziou PP, Irwig LM, Lijmer JG, Moher D, Rennie D, de Vet HC, Standards for Reporting of Diagnostic Accuracy: Towards complete and accurate reporting of studies of diagnostic accuracy: the STARD initiative. *BMJ* 2003, 326:41-4.
- von Elm E, Altman DG, Egger M, Pocock SJ, Gøtzsche PC, Vandenbroucke JP, for the STROBE Initiative: Strengthening the reporting of observational studies in epidemiology (STROBE) statement: guidelines for reporting observational studies. *BMJ* 2007, 335:806-8.
- Standards of Reporting Trials Group: A proposal for structured reporting of randomized controlled trials. JAMA 1994, 272:1926-31.
- 20. Rennie D: Reporting randomized controlled trials. An experiment and a call for responses from readers. *JAMA* 1995, **273**:1054-5.
- Hopewell S, Clarke M, Moher D, Wager E, Middleton P, Altman DG, Schulz KF, CONSORT Group: CONSORT for reporting randomised trials in journal and conference abstracts. *Lancet* 2008, 371:281-3.
- Chan AW, Hróbjartsson A, Haahr MT, Gøtzsche PC, Altman DG: Empirical evidence for selective reporting of outcomes in randomized trials: comparison of protocols to published articles. JAMA 2004, 291:2457-65.
- 23. Sackett DL: Commentary: Measuring the success of blinding in RCTs: don't, must, can't or needn't?. Int J Epidemiol 2007, 36:664-5.
- 24. Schulz KF, Grimes DA: Blinding in randomised trials: hiding who got what. *Lancet* 2002, **359**:696-700.
- Montori VM, Devereaux PJ, Adhikari NK, Burns KE, Eggert CH, Briel M, Lacchetti C, Leung TW, Darling E, Bryant DM, Bucher HC, Schünemann HJ, Meade MO, Cook DJ, Erwin PJ, Sood A, Sood R, Lo B, Thompson CA, Zhou Q, Mills E, Guyatt GH: Randomized trials stopped early for benefit: a systematic review. JAMA 2005, 294:2203-9.

- 26. Hollis S, Campbell F: What is meant by intention to treat analysis? Survey of published randomised controlled trials. *BMJ* 1999, **319**:670-4.
- Nuovo J, Melnikow J, Chang D: Reporting number needed to treat and absolute risk reduction in randomized controlled trials. *JAMA* 2002, 287:2813-4.
- Ioannidis JP, Evans SJ, Gøtzsche PC, O'Neill RT, Altman DG, Schulz K, Moher D, CONSORT Group: Better reporting of harms in randomized trials: an extension of the CONSORT statement. Ann Intern Med 2004, 141:781-8.
- De Angelis C, Drazen JM, Frizelle FA, Haug C, Hoey J, Horton R, Kotzin S, Laine C, Marusic A, Overbeke AJ, Schroeder TV, Sox HC, Weyden Van Der MB: International Committee of Medical Journal Editors. Clinical trial registration: a statement from the International Committee of Medical Journal Editors. *Lancet* 2004, 364:911-2.
- Lexchin J, Bero LA, Djulbegovic B, Clark O: Pharmaceutical industry sponsorship and research outcome and quality: systematic review. *BMJ* 2003, 326:1167-70.
- Hopewell S, Clarke M, Moher D, Wager E, Middleton P, Altman DG, Schulz KF, CONSORT Group: CONSORT for reporting randomized controlled trials in journal and conference abstracts: explanation and elaboration. *PLoS Med* 2008, 5:e20.
- Boutron I, Moher D, Altman DG, Schulz KF, Ravaud P: Extending the CONSORT statement to randomized trials of nonpharmacologic treatment: explanation and elaboration. *Ann Intern Med* 2008, 148:295-309.
- Gagnier JJ, Boon H, Rochon P, Moher D, Barnes J, Bombardier C: Reporting randomized, controlled trials of herbal interventions: an elaborated CONSORT statement. Ann Intern Med 2006, 144:364-7.
- Zwarenstein M, Treweek S, Gagnier JJ, Altman DG, Tunis S, Haynes B, Oxman AD, Moher D, CONSORT group: Pragmatic Trials in Healthcare (Practihc) group. Improving the reporting of pragmatic trials: an extension of the CONSORT statement. *BMJ* 2008, 11(337):a2390.

### doi:10.1186/1745-6215-11-32

**Cite this article as:** Schulz *et al.*: **CONSORT 2010 Statement: updated guidelines for reporting parallel group randomised trials.** *Trials* 2010 11:32.

### Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

BioMed Central

Submit your manuscript at www.biomedcentral.com/submit