The RoB 2.0 tool (individually randomized, cross-over trials)

Study design

- □ Randomized parallel group trial
- □ Cluster-randomized trial
- ☑ Randomized cross-over or other matched design

Specify which outcome is being assessed for risk of bias

Specify the numerical result being assessed. In case of multiple alternative analyses being presented, specify the numeric result (e.g. RR = 1.52 (95% CI 0.83 to 2.77) and/or a reference (e.g. to a table, figure or paragraph) that uniquely defines the result being assessed.

Is your aim for this study ...?

- \Box to assess the effect of *assignment to intervention*
- \Box to assess the effect of *starting and adhering to intervention*

Which of the following sources have you <u>obtained</u> to help inform your risk of bias judgements (tick as many as apply)?

- □ Journal article(s) with results of the trial
- □ Trial protocol
- □ Statistical analysis plan (SAP)
- Non-commercial trial registry record (e.g. ClinicalTrials.gov record)
- Company-owned trial registry record (e.g. GSK Clinical Study Register record)
- Grey literature" (e.g. unpublished thesis)
- □ Conference abstract(s) about the trial
- Regulatory document (e.g. Clinical Study Report, Drug Approval Package)
- □ Research ethics application
- Grant database summary (e.g. NIH RePORTER, Research Councils UK Gateway to Research)
- Personal communication with trialist
- Personal communication with the sponsor



Risk of bias assessment

Responses <u>underlined in green</u> are potential markers for low risk of bias, and responses in red are potential markers for a risk of bias. Where questions relate only to sign posts to other questions, no formatting is used.

Bias domain	Signalling questions	Elaboration	Response options
Bias arising from the randomization process	1.1 Was the allocation sequence random?	"Yes" if a random component was used in the sequence generation process such as using a computer generated random numbers, referring to a random number table, minimization, coin tossing; shuffling cards or envelopes; throwing dice; or drawing of lots. Minimization may be implemented without a random element, and this is considered to be equivalent to being random.	<u>Y / PY</u> / PN / N / NI
		"No" if the sequence is non-random, such that it is either likely to introduce confounding, or is predictable or difficult to conceal, e.g. alternation, methods based on dates (of birth or admission) or patient record numbers, allocation decision made by clinicians or participants, based on the availability of the intervention, or any other systematic or haphazard method.	
		If the only information about randomization methods is to state that the study is randomized, then this signalling question should generally be answered as "No information". There may be situations in which a judgement is made to answer "Probably No" or "Probably yes". For example, if the study was large, conducted by an independent trials unit or carried out for regulatory purposes, then it may be reasonable to assume that the sequence was random. Alternatively, if other (contemporary) trials by the same investigator team have clearly used non-random sequences, it might be reasonable to assume that the current study was done using similar methods. Similarly, if participants and personnel are all unaware of intervention assignments throughout/during the trial (blinding or masking), this may be an indicator that the allocation process was also concealed, but this will not necessarily always be the case.	
		If the allocation sequence was clearly concealed but there is no information about how the sequence was generated, it will often be reasonable to assume that the sequence was random (although this will not necessarily always be the case).	
	1.2 Was the allocation sequence concealed until participants were recruited and assigned to interventions?	"Yes" if any form of remote or centrally administered randomization, where the process of allocation is controlled by an outsourced unit or organization, independent of the enrolment personnel (e.g. independent central pharmacy, telephone or internet-based randomization service providers).	<u>Y / PY</u> / PN / N / NI

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	 "Yes" if envelopes or drug containers were used appropriately. Envelopes should be sequentially numbered, sealed with a tamper proof seal and opaque. Drug containers should be sequentially numbered and of identical appearance. This level of detail is rarely provided in reports, and a judgement may be required (e.g. "Probably yes" or "Probably no"). "No" if there is reason to suspect the enrolling investigator or the participant had knowledge of the forthcoming allocation. 	
1.3 Were there baseline imbalances that suggest a problem with the randomization process?	NB Imbalances that are small and compatible with chance should not be highlighted using the RoB 2.0 tool; chance imbalances are not bias.Answer "No" if no imbalances are apparent or if any observed imbalances are compatible with chance	<mark>Y / PY / <u>PN / N</u> /</mark> NI
	 Answer "Yes" if there are imbalances that indicate problems with the randomization process, including: (1) unusually large differences between intervention group sizes; or (2) a substantial excess in statistically significant differences in baseline characteristics than would be expected by chance alone; or (3) imbalance in key prognostic factors (or baseline measures of outcome variables) that are unlikely to be due to chance. 	
	An answer of "Yes/Probably yes" may exceptionally be given if the groups are surprisingly balanced in a way that appears incompatible with chance and the randomization methods, thus raising suspicion about the methods used. In some circumstances, it may be reasonable to answer "Yes/Probably yes" (rather than "No information") when there is a surprising lack of information on baseline characteristics when such information could reasonably be expected to be available/reported.	
	Answer "No information" when there is no <i>useful</i> baseline information available (e.g. abstracts, or studies that reported only baseline characteristics of participants in the final analysis). The answer to this question should not be used to influence answers to questions 1.1 or 1.2. For example, if the trial has large baseline imbalances, but authors report adequate randomization methods, questions 1.1 and 1.2 should still be answered on the basis of the reported adequate methods, and any concerns about the imbalance	

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	should be raised in the answer to the question 1.3 and reflected in the domain-level risk of bias judgement).	
1.4 Is a roughly equal proportion of participants allocated to each of the two groups?	If the allocation ratio is 1:1, then any general trends in outcomes over time (that is, period effects) will cancel. Thus if the answer to this question is yes or probably yes, then the risk of bias is low.	<u>Y / PY</u> / PN / N / NI
	If the answer to this question is no or probably no, a general trend in outcomes over time may lead to bias. For example, if there is a general deterioration in outcomes, imbalance in numbers will lead to bias against the intervention that is "over- represented" in the second period.	
1.5 If N/PN/NI to 1.4: Are period effects included in the analysis?	If period effects are included in the analysis, then any general trend over time should not cause a problem and the risk of bias would be low. If period effects are present but not included in the analysis, then there is a risk of bias.	NA / <u>Y / PY</u> / PN / <mark>N</mark> / NI
Risk of bias judgement	See Figure 1.	Low / High / Some concerns
Optional: What is the predicted direction of bias arising from the randomization process?	If the likely direction of bias can be predicted, it is helpful to state this. The direction might be characterized either as being towards (or away from) the null, or as being in favour of one of the interventions.	Favours experimental / Favours comparator / Towards null /Away from null / Unpredictable

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Bias domain	Signalling questions	Elaboration	Response options
Bias due to	If your aim for this study is to assess the	effect of assignment to intervention, answer the following questions	
deviations from intended interventions	2.1. Were participants aware of their assigned intervention during each period of the trial?	If participants are aware of their intervention assignment, it is more likely that additional health-related behaviours will differ between the assigned interventions, so risk of bias will be higher. Masking participants, which is most commonly achieved through use of a placebo or sham intervention, may prevent such differences.	<mark>Y / PY / <u>PN / N</u> / NI</mark>
	2.2. Were carers and trial personnel aware of participants' assigned intervention during each period of the trial?	If those involved in caring for participants or making decisions about their health care are aware of the assigned intervention, then implementation of the intended intervention, or administration of additional co-interventions, may differ between the assigned interventions. Masking carers and trial personnel, which is most commonly achieved through use of a placebo, may prevent such differences.	<mark>Y / PY / <u>PN / N</u> / NI</mark>
	2.3. <u>If Y/PY/NI to 2.1 or 2.2</u> : Were there deviations from the intended interventions beyond what would be expected in usual practice?	 When interest focusses on the effect of assignment to intervention, it is important to distinguish between: (a) deviations that happen in usual practice following the intervention and so are part of the intended intervention (for example, cessation of a drug intervention because of acute toxicity); and (b) deviations from intended intervention that arise due to expectations of a difference between intervention and comparator (for example because participants have a preference for one intervention over the other). We use the term "usual practice" to refer to the usual course of events in a non-trial context. Because deviations that arise due to expectations of a difference between intervention and comparator are not part of usual practice, they may lead to biased effect estimates that do not reflect what would happen to participants assigned to the interventions in practice. Trialists do not always report (and do not necessarily know) whether deviations that are not part of usual practice actually occurred. Therefore the answer "No information" may be appropriate. However, if such deviations <i>probably</i> occurred you should answer "Probably yes". 	NA / Y / PY / <u>PN /</u> <u>N</u> / NI
	2.4. <u>If Y/PY to 2.3</u> : Were these deviations from intended interventions unbalanced between the two interventions <i>and</i> likely to have affected the outcome?	Deviations from intended interventions that do not reflect usual practice will be important if they affect the outcome, but not otherwise. Furthermore, bias will arise only if there is imbalance in the deviations across the two interventions.	NA / <mark>Y / PY</mark> / <u>PN /</u> <u>N</u> / NI

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2.5 Was there sufficient time carry-over effects to have dia before outcome assessment second period?	or any ppearedCarry-over is a key concern in cross-over trials. It reflects a deviation from the intended intervention, because it acts like a co-intervention during the second period. An understanding of the likelihood of carry-over requires content knowledge, and information to inform this judgement may not be available from the report of the cross- over trial.Y / PY / PN / N / NI
	Carry-over effects can sometimes be detected by comparing imbalance in participant variables at the start of the second period with imbalance in variables at the start of the first period. If there is an exaggerated imbalance at the start of the second period, it may be due to carry over of effects.
	It is important that carry-over effects do not affect outcomes measured in the second period. A long period of wash-out between periods can be used to ensure participants start the second period in a state that is unaffected by what they received in the first period. However, a wash-out period is not essential. The important consideration is whether sufficient time passes before outcome measurement in the second period, such that any carry-over effects have disappeared. (This might sometimes be viewed as the participants having reached "steady state".) If a wash-out period is absent or is too short for carry-over effects to have disappeared, then measurements taken early in the second period may be affected by carry-over
If your aim for this study is t	assess the effect of starting and adhering to intervention, answer the following questions
2.1. Were participants aware allocated intervention during period of the trial?	of their If participants are aware of their intervention assignment, it is more likely that Y / PY / PN / N / each additional health-related behaviours will differ between the intervention groups, so risk NI of bias will be higher. Masking participants, which is most commonly achieved through NI
2.2. Were carers and trial pe aware of participants' alloca intervention during each per trial?	onnelIf those involved in caring for participants and those otherwise involved in the trial are aware of group assignment, then it is more likely that implementation of the intended intervention, or the administration of additional co-interventions, will differ between the interventions. Masking carers and trial personnel, which is most commonly achieved through use of a placebo, may prevent such differences.Y / PY / PN / N / NI
2.3. <u>If Y/PY/NI to 2.1 or 2.2</u> : important co-interventions b across the two interventions	ereRisk of bias will be higher if unplanned co-interventions were implemented in a way that would bias the estimated effect of intervention. Co-interventions will be important if they affect the outcome, but not otherwise. Bias will arise only if there is imbalance in such co-interventions between the interventions. Consider the co-interventions, including any pre-specified co-interventions, that are likely to affect the outcome and to have been administered in this study. Consider whether these co-interventions are balanced between the two interventions.NA / Y / PY / PN / N / NI

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2.4. Was the intervention implemented successfully?	Risk of bias will be higher if the intervention was not implemented as intended by, for example, the health care professionals delivering care during the trial. Consider whether implementation of the intervention was successful for most participants.	<u>Y / PY</u> / PN / N / NI
2.5. Did study participants adhere to the assigned intervention regimen?	 Risk of bias will be higher if participants did not adhere to the intervention as intended. Lack of adherence includes imperfect compliance, cessation of intervention, unintended crossovers to the comparator intervention and switches to another active intervention. Consider available information on the proportion of study participants who continued with their assigned intervention throughout follow up, and answer "No" or "Probably No" if this proportion is high enough to raise concerns. Answer "Yes" for studies of interventions that are administered once, so that imperfect adherence is not possible. One possibility is that the level of adherence will differ by period. For example, participants may adhere less well during the second period. 	<u>Y / PY</u> / PN / N / NI
2.6. <u>If N/PN/NI to 2.3, 2.4 or 2.5</u> : Was an appropriate analysis used to estimate the effect of starting and adhering to the intervention?	It is possible to conduct an analysis that corrects for some types of deviation from the intended intervention. Examples of appropriate analysis strategies include inverse probability weighting or instrumental variable estimation. It is possible that a paper reports such an analysis without reporting information on the deviations from intended intervention, but it would be hard to judge such an analysis to be appropriate in the absence of such information. Note that analyses of the full data from a cross-over trial cannot generally correct for carry-over effects when they are present. If everyone received a co-intervention alongside one of the interventions, adjustments	NA / <u>Y / PY</u> / PN / N / NI
	cannot be made to overcome this. Some examples of analysis strategies that would not be appropriate to estimate the effect of intended intervention are (i) "ITT analysis", (ii) "per protocol analysis", and (iii) "analysis by treatment received".	
2.7 Was there sufficient time for any carry-over effects to have disappeared before outcome assessment in the second period?	Carry-over is a key concern in cross-over trials. It reflects a deviation from the intended intervention, because it acts like a co-intervention during the second period. An understanding of the likelihood of carry-over requires content knowledge, and information to inform this judgement may not be available from the report of the cross-over trial.	<u>Y / PY</u> / PN / N / NI
	Carry-over effects can sometimes be detected by comparing imbalance in participant variables at the start of the second period with imbalance in variables at the start of the	

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	first period. If there is an exaggerated imbalance at the start of the second period, it may be due to carry over of effects.	
	It is important that carry-over effects do not affect outcomes measured in the second period. A long period of wash-out between periods can be used to ensure participants start the second period in a state that is unaffected by what they received in the first period. However, a wash-out period is not essential. The important consideration is whether sufficient time passes before outcome measurement in the second period, such that any carry-over effects have disappeared. (This might sometimes be viewed as the participants having reached "steady state".) If a wash-out period is absent or is too short for carry-over effects to have disappeared, then measurements taken early in the second period may be affected by carry-over.	
Risk of bias judgement	See Figure 2 and Figure 3.	Low / High / Some concerns
Optional: What is the pred direction of bias due to dev intended interventions?	cted viations from If the likely direction of bias can be predicted, it is helpful to state this. The direction might be characterized either as being towards (or away from) the null, or as being in favour of one of the interventions.	Favours experimental / Favours comparator / Towards null /Away from null / Unpredictable

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Bias domain	Signalling questions	Elaboration	Response options
Bias due to missing	3.1 Were outcome data available for all, or nearly all, participants randomized?	The appropriate study population for an analysis of the intention to treat effect is all randomized patients.	<u>Y / PY</u> / PN / N / NI
outcome data		Note that imputed data should be regarded as missing data, and not considered as "outcome data" in the context of this question.	
		"Nearly all" (equivalently, a low or modest amount of missing data) should be interpreted as "enough to be confident of the findings", and a suitable proportion depends on the context.	
		For continuous outcomes, availability of data from 95% (or possibly 90%) of the participants would often be sufficient. For dichotomous outcomes, the proportion required is directly linked to the risk of the event. If the observed number of events is much greater than the number of participants with missing outcome data, the bias would necessarily be small.	
	3.2 If N/PN/NI to 3.1: Are the proportions of missing outcome data and reasons for missing outcome data similar across interventions?	"Similar" (with regard to proportion and reasons for missing outcome data) includes some minor degree of discrepancy across intervention groups as expected by chance. Assessment of comparability of reasons for missingness requires the reasons to be reported.	NA / <u>Y / PY</u> / PN / <mark>N</mark> / NI
		Bias would be introduced if, for example, the participants omitted from the analysis were those for whom one treatment is superior, leaving in the analysis only those in whom the treatments have the same effect. This is an instance of participants with missing data differing importantly between groups. It would be difficult to address this in an analysis – it would require strong assumptions about informative missingness.	
	3.3. If N/PN/NI to 3.1: Is there evidence that results were robust to the presence of missing outcome data?	Evidence for robustness may come from how missing data were handled in the analysis and whether sensitivity analyses were performed by the trial investigators, or from additional analyses performed by the systematic reviewers.	NA / <u>Y / PY</u> / PN / N / NI
		Use of last observation carried forward imputation may be particularly problematic if the observations being carried forward were made before carry-over effects had disappeared.	
		A common debate in analysis of a cross-over trial is between having the patient effect as fixed or random. The former will automatically exclude (for an AB/BA design) all patients with missing data in either period. The latter will permit the recovery of inter- patient information and can thus in theory lead to more precise inferences (although in practice the effect is small). Validity of either approach rests on an assumption of data being missing at random.	

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Risk of bias judgement	See Figure 4.	Low / High /
		Some concerns
Optional: What is the predicted	If the likely direction of bias can be predicted, it is helpful to state this. The direction	Favours
direction of bias due to missing	might be characterized either as being towards (or away from) the null, or as being in	experimental /
outcome data?	favour of one of the interventions.	Favours
		comparator /
		Towards null
		/Away from null /
		Unpredictable

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Bias domain	Signalling questions	Elaboration	Response options
Bias in measurement of the	4.1 Were outcome assessors aware of the intervention received by study participants?	"No" if outcome assessors were blinded to intervention status. In studies where participants report their outcomes themselves (i.e., participant-reported outcome), the outcome assessor is the study participant.	<mark>Y / PY / <u>PN / N</u> / NI</mark>
outcome	4.2 If Y/PY/NI to 4.1: Was the assessment of the outcome likely to be influenced by knowledge of intervention received?	Knowledge of the assigned intervention may impact on participant-reported outcomes (such as level of pain), observer-reported outcomes involving some judgement, and intervention provider decision outcomes, while not impacting on other outcomes such as observer reported outcomes not involving judgement such as all-cause mortality. In many circumstances the assessment of <i>observer reported outcomes not involving judgement</i> such as all-cause mortality might be considered to be unbiased, even if outcome assessors were aware of intervention assignments.	NA / <mark>Y / PY</mark> / <u>PN /</u> <u>N</u> / NI
	Risk of bias judgement	See Figure 5.	Low / High / Some concerns
	Optional: What is the predicted direction of bias due to measurement of the outcome?	If the likely direction of bias can be predicted, it is helpful to state this. The direction might be characterized either as being towards (or away from) the null, or as being in favour of one of the interventions.	Favours experimental / Favours comparator / Towards null /Away from null / Unpredictable

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Bias domain	Signalling questions	Elaboration	Response options
Bias in selection of the reported	Are the reported outcome data likely to have been selected, on the basis of the results, from		
result	5.1 multiple outcome measurements (e.g. scales, definitions, time points) within the outcome domain?	A particular outcome domain (i.e. a true state or endpoint of interest) may be measured in multiple ways. For example, the domain pain may be measured using multiple scales (e.g. a visual analogue scale and the McGill Pain Questionnaire), each at multiple time points (e.g. 3, 6 and 12 weeks post-treatment). If multiple measurements were made, but only one or a subset is reported on the basis of the results (e.g. statistical significance), there is a high risk of bias in the fully reported result. A response of "Yes/Probably yes" is reasonable if:	<mark>Y / PY / <u>PN / N</u> / NI</mark>
		There is clear evidence (usually through examination of a trial protocol or statistical analysis plan) that a domain was measured in multiple ways, but data for only one or a subset of measures is fully reported (without justification), and the fully reported result is likely to have been selected on the basis of the results. Selection on the basis of the results arises from a desire for findings to be newsworthy, sufficiently noteworthy to merit publication, or to confirm a prior hypothesis. For example, trialists who have a preconception or vested interest in showing that an experimental intervention is beneficial may be inclined to selectively report outcome measurements that are favourable to the experimental intervention.	
		A response of "No/Probably no" is reasonable if:	
		There is clear evidence (usually through examination of a trial protocol or statistical analysis plan) that all reported results for the outcome domain correspond to all intended outcome measurements.	
		or There is only one possible way in which the outcome domain can be measured (hence there is no opportunity to select from multiple measures). or	
		Outcome measurements are inconsistent across different reports on the same trial, but the trialists have provided the reason for the inconsistency and it is not related to the nature of the results. A response of "No information" is reasonable if:	

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	Analysis intentions are not available, or the analysis intentions are not reported in sufficient detail to enable an assessment, and there is more than one way in which the outcome domain could have been measured.	
5.2 multiple analyses of the data?	A particular outcome domain may be analysed in multiple ways. Examples include: unadjusted and adjusted models; final value vs change from baseline vs analysis of covariance; transformations of variables; conversion of continuously scaled outcome to categorical data with different cut-points; different sets of covariates for adjustment; different strategies for dealing with missing data. Application of multiple methods generates multiple effect estimates for a specific outcome domain. If multiple estimates are generated but only one or a subset is reported on the basis of the results (e.g. statistical significance), there is a high risk of bias in the fully reported result. In a cross-over trial, it is possible that trial authors might decide between presenting a paired analysis and an unpaired analysis on the basis of the results. A decision between presenting a paired analysis and an unpaired analysis of the first period only might be made on the basis of whichever produces the preferred results. If there is truly no effect, then either of these might produce results that is more extreme than the other. Alternatively, a decision might be made between presenting a paired analysis and an (inappropriate) unpaired analysis of the full dataset . The expected analysis in this situation is a paired analysis. An unpaired analysis will be less precise, so a decision to present an unpaired analysis is likely to be made only if the trialists were keen to show a lack of effect or equivalence of interventions. Such behaviour is probably unusual.	Y / PY / <u>PN / N</u> / NI
	A response of "Yes/Probably yes" is reasonable if:	
	There is clear evidence (usually through examination of a trial protocol or statistical analysis plan) that a domain was analysed in multiple ways, but data for only one or a subset of analyses is fully reported (without justification), and the fully reported result is likely to have been selected on the basis of the results. Selection on the basis of the results arises from a desire for findings to be newsworthy, sufficiently noteworthy to merit publication, or to confirm a prior hypothesis. For example, trialists who have a preconception or vested interest in showing that an experimental intervention is beneficial may be inclined to selectively report analyses that are favourable to the experimental intervention. A response of "No/Probably no" is reasonable if:	

20/10/2016 There is clear evidence (usually through examination of a trial protocol or statistical analysis plan) that all reported results for the outcome domain correspond to all intended analyses. or There is only one possible way in which the outcome domain can be analysed (hence there is no opportunity to select from multiple analyses). or Analyses are inconsistent across different reports on the same trial, but the trialists have provided the reason for the inconsistency and it is not related to the nature of the results. A response of "No information" is reasonable if: Analysis intentions are not available, or the analysis intentions are not reported in sufficient detail to enable an assessment, and there is more than one way in which the outcome domain could have been analysed. 5.3 ... the outcome of a statistical test for Selective reporting of results from the first period only is likely to be due to carry-over Y/PY/PN/N/ carry-over? having been identified. The test for carry-over is affected importantly by baseline NI differences in the randomized groups at the start of the cross-over trial. If a statistically significant result is obtained, it might therefore reflect such baseline differences. Reporting only the first period data in this situation is particularly problematic given the possibility that the two groups differ in their baseline characteristics; the benefits of the cross-over design in making intervention comparisons within individuals is lost. **Risk of bias judgement** See Figure 6. Low / High / Some concerns Optional: What is the predicted If the likely direction of bias can be predicted, it is helpful to state this. The direction Favours might be characterized either as being towards (or away from) the null, or as being in direction of bias due to selection of the experimental / favour of one of the interventions. reported result? Favours comparator / Towards null /Away from null / Unpredictable

Bias domain	Signalling questions	Elaboration	Response options			
Overall bias	Risk of bias judgement	See Table 1.	Low / High / Some concerns			
	Optional: What is the overall predicted direction of bias for this outcome?	If the likely direction of bias can be predicted, it is helpful to state this. The direction might be characterized either as being towards (or away from) the null, or as being in favour of one of the interventions.	Favours experimental / Favours comparator / Towards null /Away from null / Unpredictable			



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Figure 1. Suggested algorithm for reaching risk of bias judgements for bias arising from the randomization process. This is only a suggested decision tree: all default judgements can be overridden by assessors.



Figure 2. Suggested algorithm for reaching risk of bias judgements for bias due to deviations from intended interventions (*effect of assignment to intervention*). This is only a suggested decision tree: all default judgements can be overridden by assessors.



Figure 3. Suggested algorithm for reaching risk of bias judgements for bias due to deviations from intended interventions (*effect starting and adhering to intervention*). This is only a suggested decision tree: all default judgements can be overridden by assessors.



Figure 4. Suggested algorithm for reaching risk of bias judgements for bias due to missing outcome data. This is only a suggested decision tree: all default judgements can be overridden by assessors



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Figure 5. Suggested algorithm for reaching risk of bias judgements for bias in measurement of the outcome. This is only a suggested decision tree: all default judgements can be overridden by assessors.



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Figure 6. Suggested algorithm for reaching risk of bias judgements for bias in selection of the reported result. This is only a suggested decision tree: all default judgements can be overridden by assessors



Overall risk of bias judgement	Criteria
Low risk of bias	The study is judged to be at low risk of bias for all domains for this result.
Some concerns	The study is judged to be at some concerns in at least one domain for this result.
High risk of bias	The study is judged to be at high risk of bias in at least one domain for this result.
	Or
	The study is judged to have some concerns for multiple domains in a way that substantially lowers confidence in the result.

Table 1. Reach	ing an overall risk of	bias judgement	for a spe	cific outcome.
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