

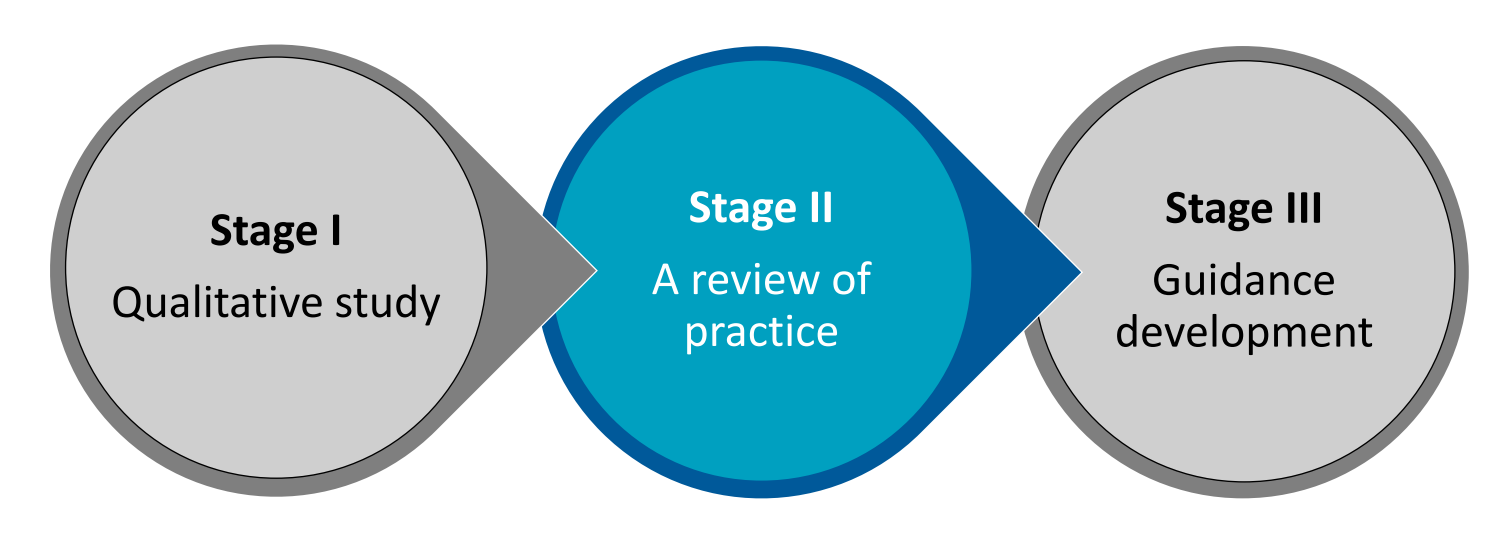
Blinding of Trial Statisticians: A review of current practice

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Background

The Blinding of Trial Statisticians (BOTS) study aimed to develop guidance to advise CTUs on a risk-proportionate approach to blinding trial statisticians (TSs) within clinical trials.



The aim of this part of the study was to compare the outcomes of recently published randomised trials according to blinding status of the statistician.

Methods

Data collection

HTA & EME trials published in NIHR journals library between 2016 & 2020

Trial characteristics

Blinding status of the statistician

Trial outcome and potential sources of bias

Analysis sets

Primary analysis			
Included only those studies where			
	the blinding status of the TS could		
	be confirmed.		

Sensitivity analysis Included all studies, by assuming

that the TS was not blinded for those studies where the blinding status was unclear.

Results



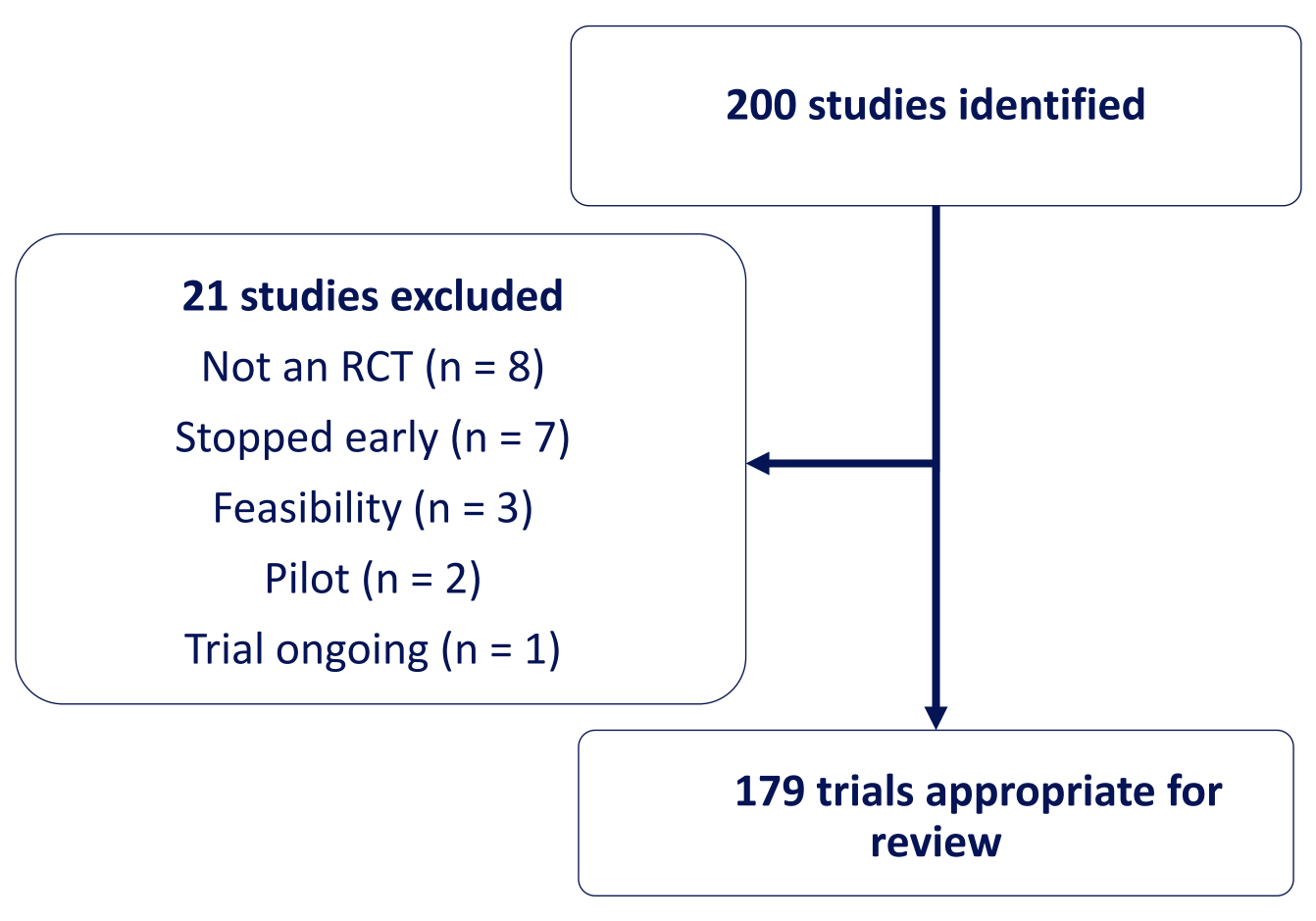


Table 1: Statistically significant findings			
	Statistician blinded prior to the final analysis		
	No	Yes	Unclear
Statistically significant finding reported			
for primary outcome, n/N (%)	23/83 (28%)	19/69 (28%)	8/27 (30%)

Table 2: Odds ratio describing the association between selected study design features and the reporting of statistically significant findings

	Primary analysis (N = 152)	Sensitivity analysis (N = 179)
Blinded statistician	0.98 (0.47, 2.05)	0.96 (0.48, 1.92)
Multiple comparisons ¹	1.34 (0.63, 2.86)	1.38 (0.69, 2.78)
Sample size not achieved	0.87 (0.38, 1.97)	0.72 (0.34, 1.52)
Blinded trial ²	0.33 (0.13, 0.86)	0.36 (0.15, 0.88)
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¹ A composite of multiple treatment groups and multiple primary outcomes.

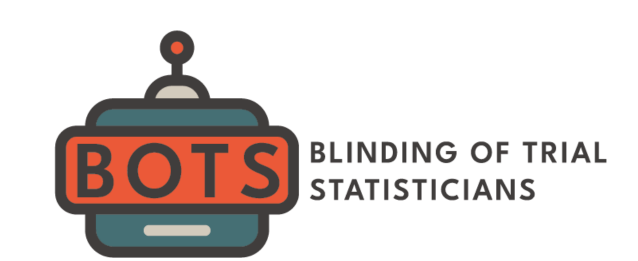
Discussion

- There is no evidence that the blinding status of the statistician is associated with the probability of a significant finding being reported.
- There remains some uncertainty around this finding; however, it appears risk of bias is certainly smaller compared to when other groups are unblinded.
- The reporting of blinding methodology was often absent or of low quality.

A risk-proportionate approach

Alongside the BOTS qualitative findings, this study provides evidence for a *risk-proportionate* approach to blinding statisticians. That is, the decision about whether to blind or not blind the statistician should be based upon the specific risks and merits for a given trial.







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² Participants, clinicians, and outcome assessors all blinded; where blinding status is unclear this was assumed as "no blinding".