

Measuring the success of blinding in placebo-controlled trials: should we be so quick to dismiss it?

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1 Measuring the success of blinding in placebo-controlled trials: should we be so quick to 2 dismiss it? 3 Running head: Measuring blinding success in placebo-controlled trials 4 Author list and affiliations Rebecca K Webster*, PhD, University of Oxford, Oxford, United Kingdom and University of 5 6 Sheffield, United Kingdom; Department of Psychology, Cathedral Court, 1 Vicar 7 Lane, Sheffield, S1 2LT; r.k.webster@sheffield.ac.uk 8 Felicity Bishop, PhD, University of Southampton, Southampton, United Kingdom Gary S Collins, PhD, University of Oxford, and NIHR Oxford Biomedical Research Centre, 9 Oxford, United Kingdom 10 11 Andrea WM Evers, PhD, Leiden University, Leiden, Netherlands 12 Tammy Hoffmann, PhD, Institute of Evidence-Based Healthcare, Bond University, Queensland, Australia 13 14 J. André Knottnerus, MD PhD, Maastricht University, Maastricht, Netherlands Sarah E Lamb, DPhil, University of Oxford, Oxford, United Kingdom and University of 15 16 Exeter, Exeter, United Kingdom 17 Helen Macdonald, MD, The BMJ, London, United Kingdom 18 Claire Madigan, PhD, University of Oxford, Oxford, United Kingdom 19 Vitaly Napadow, PhD, Harvard Medical School, Boston, United States 20 Amy Price, PhD, Stanford University, Stanford, United States; University of Oxford, Oxford, 21 United Kingdom and The BMJ, London, United Kingdom, 22 Jonathan L Rees, MD, University of Oxford, and NIHR Oxford Biomedical Research Centre 23 Oxford, United Kingdom 24 Jeremy Howick, PhD, University of Oxford, Oxford, United Kingdom *Corresponding author 25 Word count: 2072 26

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1 Background

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- 29 From being almost universally regarded as a methodological virtue of clinical trials and being
- included in the original 2001 Consolidated Standards of Reporting Trials (CONSORT)
- 31 statement (1), measuring the success of blinding has fallen out of fashion. Subsequent
- 32 versions of CONSORT removed this recommendation based on the correct view that it can
- lead to misleading inferences about causes of the failure to blind. (2, 3) In addition, Anand, et
- al. (4) recently questioned the need to blind patients and clinicians or measure and report
- 35 whether blinding was done successfully. While critics are correct to point out problems with
- 36 the view that blinding is a universal methodological virtue, and to point out that measuring
- 37 the success of blinding is not straightforward, they are too quick to dismiss the value of
- testing and reporting on the success of blinding. This is reflected in our findings extending
- 39 the Template for Intervention Description and Replication (TIDieR) statement for
- 40 placebo/sham control components, in which almost all Delphi respondents recommended that
- 41 trials should measure and report whether blinding was successful. (5)
- We are not aware of any publications that set out the case for and against measuring blinding
- success, or that provide mitigating positions. Our experience suggests that confusion about
- blinding inhibits reasonable debates in this area. Here, we attempt to clarify some of the
- 45 confusions surrounding blinding and measuring its success, before providing the case for and
- against, reporting measures of the success of blinding, and suggesting a 'middle road' which
- 47 takes both sides of the debate into account.

2 Measuring blinding success: the case for

- 49 Blinding involves concealing knowledge of treatment assignment to one or more groups
- 50 involved in clinical trials (participants, intervention providers, data collectors, outcome
- assessors, statisticians, and manuscript authors). (6) Trials can be described in a number of
- 52 ways including open (unblinded), single-blind, double-blind or triple-blind. The terminology
- can be confusing however, as a random sample of 200 trials has shown that the term double
- blind can be used to describe blinding up to 18 different combinations of trial personnel. (7)
- As noted in CONSORT, it is important to specify who was blinded in a trial, (2) as blinding
- 56 different people may affect outcomes, especially those which are subjective. For example, if
- 57 participants and data collectors were not blinded this may have more of an impact than an
- unblinded statistician who may have less influence on the outcomes.

Measuring whether blinding was successful involves asking patients and clinicians about their treatment assignment beliefs before the trial is officially unblinded. Successful blinding occurs when there is a balance of expectations and beliefs related to the assigned intervention, demonstrating that those who are blinded are not aware of the (active or inactive) intervention that has been assigned. However, blinding can fail when participants, caregivers, or other groups involved in a trial deduce the intervention allocation at the beginning of the trial (e.g. due to inadequate matching between the placebo and active intervention), or during the trial (e.g. due to adverse events). (8-10) Since the function of blinding is to reduce the impact of expectations, unsuccessful blinding is problematic, as beliefs and expectations of those who correctly guess the intervention allocation *could* then influence the outcome of the trial. (11-14) As such a trial that was designed blinded but in which attempts to blind were unsuccessful may approach the quality of a trial where (complete, double) blinding is ethically and feasibly possible, but is not blinded (see Fig 1).

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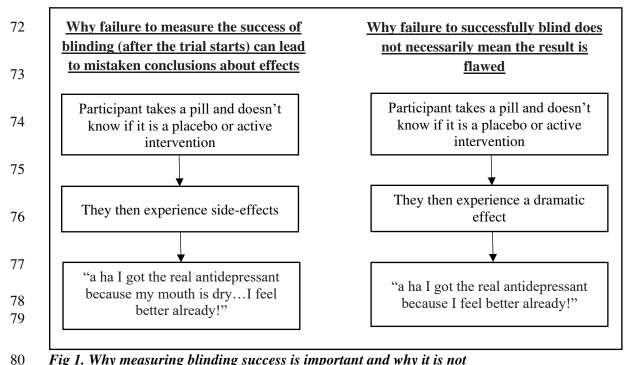


Fig 1. Why measuring blinding success is important and why it is not

A number of meta-epidemiological studies have investigated differences between trials (reported as) blinded and those that are not (reported as) blinded. (15-24) Some (but not all) of those found that lack of reporting of blinding led to larger effect sizes. Recently, Moustgaard, et al. (15) found inconsistent effects of blinding on treatment effect sizes. However, there are methodological concerns regarding the study's sample selection and classifications of reporting of blinding. (25) Like randomisation and allocation concealment, blinding can reasonably be expected to have a small average effect, possibly with an

unpredictable direction. (26, 27) In an era when marginal gains from many of our medical interventions suffice to change policy and practice, (28) ruling out small biases or errors is becoming more important. In addition, small average effects are compatible with larger effects in some instances, for example trials of treatments for disorders that are placebo responsive, such as pain. Additional meta-epidemiological studies with large sample sizes, together with well-defined outcomes, disease areas, and classifications of reporting of blinding are required to address this important issue. Such studies cannot be conducted unless trials report whether blinding was successful (where this is feasible). Aside from the importance of blinding itself, the importance of measuring (see Box 1) and reporting blinding success is apparent in various trials. For example, Karlowski, et al. (29) compared Vitamin C with placebo for treating the common cold, and found Vitamin C to be apparently effective. However, because of the sour taste of Vitamin C and sweet taste of the lactose placebo pills, the trial was not successfully blinded. When the authors carried out a subgroup analysis in which they divided participants into those who remained blinded and to those who were not, they found that there was no benefit of Vitamin C in the blinded group. Although ideally the authors should have ensured both placebo and active intervention were adequately matched, this example still shows the importance of measuring and reporting blinding success. Otherwise, it would have been mistakenly concluded that Vitamin C was superior. More recently, a unsuccessfully blinded trial of zinc for treating common cold symptoms found that zinc significantly reduced the duration of cold symptoms compared to placebo. (30) Whereas, another trial with successful blinding, found that zinc did not reduce symptom duration. (31) This difference may be due to significantly more side-effects being reported to Zinc than placebo in the first trial, (30) which led to unblinding and subsequent bias. As such the success of blinding reported in these studies could be useful for those appraising them and looking for reasons for their discrepant results.

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A common approach to measuring the success of blinding uses chi-square tests of independence, where successful blinding is indicated by a null finding (patient guesses are not related to their intervention allocation). (32) However, this lacks sensitivity and does not provide any directional information about the pattern of participant guesses. (33) James' (34) and Bang's (33) blinding index (BI) have addressed some of these concerns by asking participants to guess their intervention assignment using three responses (active, placebo or do not know). James' provides a single value that combines data from all arms ranging from 0 to 1, 0 being total lack of blinding, 1 being complete blinding and 0.5 being completely random blinding. Bang's BI aims to provide a more sensitive measure of blinding within each experimental arm compared to James' by calculating a score from -1 to 1, 1 being complete lack of blinding, 0 being consistent with perfect blinding and -1 indicating opposite guessing which may be related to unblinding. (33) As such, it can be used to detect where blinding may have failed, while still assessing overall success. An even newer method is the use of video surveillance. This involves video-recording procedures in the trial and asking a professional familiar with the procedure to guess the intervention allocation. (35) However, in practice, blinding success is rarely measured, with only 2-24% of trials reporting the success of blinding. (36, 37). In addition, these methods fall short as they do not consider why unblinding may have occurred.

Box 1. How to measure blinding success?

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3 Measuring blinding success: the case against

The case against measuring the success of blinding can be traced to Dave Sackett, who cited a 2x2 factorial trial of aspirin and sulfinpyrazone for stroke prevention. In the trial, blinded clinicians largely distinguished aspirin from sulfinpyrazone. (38) But, because of prior 'hunches' that sulfinpyrazone would be more effective, they mistakenly believed that patients with better outcomes had received sulfinpyrazone, when in fact the trial showed aspirin was more effective. In this example, the results of tests for blinding can be ambiguous. Hence, Sackett and others following him argued that tests for the success of blinding should not be conducted. Sackett is correct that in this example (and perhaps others like it), that the test for the success of blinding was confounded by mistaken beliefs about which intervention was effective (or a misattributed response to treatment). However, if these (mistaken) hunches about efficacy were different (unbalanced) in the intervention and control groups, then they could have confounded the study no matter how mistaken they were. Or, their beliefs were the same (balanced) across the groups, in which case there was no confounding (even if the beliefs were mistaken). Either way, the test for the success of blinding will reveal useful information, namely about whether expectations might have confounded the results. There are some cases in which failure to successfully blind does not imply that the study was methodologically lacking. For example, a dramatically effective treatment can cause

unblinding, however it should not lead us to conclude that a trial of the treatment was

methodologically lacking. On the contrary, as Senn (39) argued: 'The whole point of a 152 successful double-blind trial is that there should be unblinding through efficacy." The 153 problem remains however, that if a trial reports that the cause of unblinding was dramatic 154 effectiveness, a report of 'failed' blinding could mislead some into thinking the trial was less 155 156 trustworthy. Secondly, measuring the success of blinding at the wrong time (for example before follow-157 up or trial completion) may raise suspicion among participants and cause the problem it is 158 159 intended to prevent. (40) (41) 160 Thirdly, some trials cannot feasibly or ethically be blinded, for example, non-drug 161 interventions such as exercise, behavioural therapy and nutritional advice. (Aside: trials of these interventions can be rigorous by using other methodological tools to reduce bias (42), 162 163 such as pre-registering trials, following a pre-specified analysis plan, adequate sample size 164 and using randomisation, to reach the best achievable research practice.) Also, in some cases 165 unblinding is an ethical requirement, for example due to hypothesized toxicity, and blinding 166 itself could increase research waste, with some evidence indicating that patients are less likely to enrol in blinded trials. (4) 167 168 4 Discussion Demanding that all trials attempt to use and measure the success of blinding is too strong 169 because blinding is sometimes impossible, unethical, or misleading. Future research is 170 171 required to determine how to best interpret findings from assessing the success of blinding. 172 On the other hand, blinding has the potential to rule out bias, and failure to recommend that 173 the success of blinding be reported when it is measured, seems like wilful withholding of 174 information that potentially useful. In addition, the change in the CONSORT recommendation from asking researchers to report 175 176 on success of blinding (if measured) to not asking, seems to have been based on arguments that may deserve revisiting. Of course, the fact that CONSORT does not explicitly 177 178 recommend reporting on the success of blinding does not prevent reviewers from reporting it. 179 However, the fact that CONSORT sites a paper by Sackett as the reason for removing it, in 180 which he claims that testing the success of blinding is a 'mug's game' could be interpreted as a reason to avoid reporting on the success of blinding. 181 Also, while measuring the success of blinding at many (or the wrong) points may cause some 182 183 problem, this does not imply that measuring success of blinding at a single (roughly) correct

184	point is not useful. Moreover, empirical research suggests that getting the 'correct' point may
185	not be required. Rees, et al. (43) have shown that the difference between a six-point
186	assessment of blinding success during a trial and a two-point model is not significant.
187	Overall, the fact that difficulties, ethical problems, or ambiguity in measuring its success does
188	not imply that it should be given up altogether.
189	5 Conclusion and recommendation? A middle ground
190	While we acknowledge there are a dearth of studies that have investigated this issue, more
191	definitive evidence can only come from studies that measure the success of blinding. We
192	recognise that some trials cannot feasibly or ethically be blinded, but it is important that trials
193	that could have introduced blinding and measured its success, are distinguished from trials
194	that could not have. Our suggestion for a way forward considers the current state of evidence
195	for and against measuring the success of blinding. We hope this stimulates further discussion,
196	and that future iterations of CONSORT reflect on our arguments and revisits this issue.
197	We suggest that:
198	1. Authors should make every attempt to match the placebo and active intervention to
199	avoid unblinding at the start of the trial and subsequent research waste.
200	2. When authors have measured the success of blinding they should report the results.
201	3. Critical appraisers should consider reasons why unblinding may have arisen before
202	condemning a trial as having a high risk of bias, or if blinding success has not been
203	reported, they should assess whether it is possible that blinding has been compromised.
204	4. Future development of measures to assess the success of blinding should ask those
205	intended to be blinded what their intervention allocation beliefs were and why. This
206	can help disentangle the reasons (dramatic effects or side-effects), although the reason
207	may not always be known for sure.
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210	Contributor statement
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