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Validity of data extraction in evidence synthesis practice of adverse events: reproducibility study

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Additional material is published online only. To view please visit the journal online.

Cite this as: *BMJ* 2022;377:e069155

<http://dx.doi.org/10.1136/bmj-2021-069155>

bmj-2021-069155

Accepted: 10 April 2022

ABSTRACT

OBJECTIVES

To investigate the validity of data extraction in systematic reviews of adverse events, the effect of data extraction errors on the results, and to develop a classification framework for data extraction errors to support further methodological research.

DESIGN

Reproducibility study.

DATA SOURCES

PubMed was searched for eligible systematic reviews published between 1 January 2015 and 1 January 2020. Metadata from the randomised controlled trials were extracted from the systematic reviews by four authors. The original data sources (eg, full text and ClinicalTrials.gov) were then referred to by the same authors to reproduce the data used in these meta-analyses.

ELIGIBILITY CRITERIA FOR SELECTING STUDIES

Systematic reviews were included when based on randomised controlled trials for healthcare interventions that reported safety as the exclusive outcome, with at least one pair meta-analysis that included five or more randomised controlled trials and with a 2×2 table of data for event counts and sample sizes in intervention and control arms available for each trial in the meta-analysis.

MAIN OUTCOME MEASURES

The primary outcome was data extraction errors summarised at three levels: study level, meta-analysis

level, and systematic review level. The potential effect of such errors on the results was further investigated.

RESULTS

201 systematic reviews and 829 pairwise meta-analyses involving 10 386 randomised controlled trials were included. Data extraction could not be reproduced in 1762 (17.0%) of 10 386 trials. In 554 (66.8%) of 829 meta-analyses, at least one randomised controlled trial had data extraction errors; 171 (85.1%) of 201 systematic reviews had at least one meta-analysis with data extraction errors. The most common types of data extraction errors were numerical errors (49.2%, 867/1762) and ambiguous errors (29.9%, 526/1762), mainly caused by ambiguous definitions of the outcomes. These categories were followed by three others: zero assumption errors, misidentification, and mismatching errors. The impact of these errors were analysed on 288 meta-analyses. Data extraction errors led to 10 (3.5%) of 288 meta-analyses changing the direction of the effect and 19 (6.6%) of 288 meta-analyses changing the significance of the P value. Meta-analyses that had two or more different types of errors were more susceptible to these changes than those with only one type of error (for moderate changes, 11 (28.2%) of 39 v 26 (10.4%) 249, P=0.002; for large changes, 5 (12.8%) of 39 v 8 (3.2%) of 249, P=0.01).

CONCLUSION

Systematic reviews of adverse events potentially have serious issues in terms of the reproducibility of the data extraction, and these errors can mislead the conclusions. Implementation guidelines are urgently required to help authors of future systematic reviews improve the validity of data extraction.

Introduction

In an online survey of 1576 researchers by *Nature*, the collected opinions emphasised the need for better reproducibility in research: “More than 70% of researchers have tried and failed to reproduce another scientist’s experiments, and more than half have failed to reproduce their own experiments.”¹

Systematic reviews and meta-analyses have become the most important tools for assessing healthcare interventions. This research involves explicit and standardised procedures to identify, appraise, and synthesise all available evidence within a specific topic.² During the process of systematic reviews, each step matters, and any errors could affect the reliability of the final results. Among these steps, data extraction is arguably one of the most important and is prone to errors because raw data are transferred from original

WHAT IS ALREADY KNOWN ON THIS TOPIC

In evidence synthesis practice, data extraction is an important step and prone to errors, because raw data are transferred from the original studies into the meta-analysis

Data extraction errors in systematic reviews occur frequently in the literature, although these errors generally have a minor effect on the results

However, this conclusion is based on systematic reviews of continuous outcomes, and might not apply to binary outcomes of adverse events

WHAT THIS STUDY ADDS

In a large-scale reproducibility investigation of 201 systematic reviews of adverse events with 829 pairwise meta-analyses, data extraction errors frequently occurred for binary outcomes of adverse events

These errors could be grouped into five categories based on the mechanism: numerical error, ambiguous error, zero assumption error, mismatching error, and misidentification error

The errors can lead to changes in the conclusions of the findings, and meta-analyses that had two or more types of errors were more susceptible to these changes

studies into the systematic review that serves as the basis for evidence synthesis.

To ensure the quality of data extraction, authoritative guidelines, such as the Cochrane Handbook, highlight the importance of independent extraction by two review authors.² Despite this quality assurance mechanism, data extraction error in systematic reviews occurs frequently in the literature.³ Jones et al⁴ reproduced 34 Cochrane reviews published in 2003 (issue 4) and found that 20 (59%) had data extraction errors. Gøtzsche et al⁵ examined 27 meta-analyses of continuous outcomes and reported that 17 (63%) of these meta-analyses had an error for at least one of the two randomly selected trials. In their subsequent study, based on 10 systematic reviews of continuous outcomes, seven (70%) were identified as erroneous data.⁶

Empirical evidence suggests that the effect of data extraction error seems to be minor.^{3,5} However, this conclusion is based on systematic reviews of continuous outcomes, which do not apply to binary outcomes of adverse events. Harms, especially serious harms, tend to be rare, and such data in nature are more susceptible to random or systematic errors than are common outcomes.^{7,8} For example, consider a 1:1 designed trial with a sample size of 100, and the event counts of death are two intervention group and one in the control group. If the review authors incorrectly extracted the number of events in the intervention group as one, the relative risk would drop from two to one, leading to a completely different conclusion. Owing to this feature, in systematic reviews of adverse events, the validity of data extraction can considerably affect the results and even predominate the final conclusion. The erroneous conclusion would further influence the clinical practice guidelines and mislead healthcare practice.

We used a large-scale reproducibility investigation on the reproducibility of data extraction for systematic reviews of adverse events. We propose an empirical classification of the data extraction errors to help methodologists and systematic review authors better understand the sources of data extraction errors. The impact of such errors on the results is also examined based on the reproducibility dataset.

Methods

Protocol and data source

This article is an extension of our previous work describing methods to deal with double-zero-event studies.⁹ A protocol was drafted on 11 April 2021 by a group of core authors (CX, TY, LL, LFK), which was then revised after expert feedback (SV, LZ, RQ, and JZ; see supplementary file). We also record the detailed implementation of this study (supplementary table 1).

A subset of the data from the previous study was used in this study. Briefly, we searched PubMed for systematic reviews of adverse events indexed from 1 January 2015 to 1 January 2020. The limit on the search date was arbitrary but allowed us to capture the practice of the most recent systematic reviews.

We did not search in other databases because we did not aim to include all systematic reviews; instead, a representative sample was sufficient for the aim of the current study. The search strategy was developed by an information specialist (supplementary box 1), and the literature search was conducted on 28 July 2020, and has been recorded elsewhere.⁹

Inclusion criteria and screening

We included systematic reviews of randomised controlled trials for healthcare interventions, with adverse events as the exclusive outcome. The term adverse event was defined as “any untoward medical occurrence in a patient or subject in clinical practice,”¹⁰ which could be a side effect, adverse effect, adverse reaction, harm, or complication associated with any healthcare intervention.¹¹ We did not consider systematic reviews based on other types of studies because randomised controlled trials are more likely to be registered with available related summarised data for safety outcomes; this source provided another valid way to assess the reproducibility of data extraction. Additionally, we limited systematic reviews to those with at least one pairwise meta-analysis with five or more studies; the requirement of the number of studies was designed for an ongoing series of studies on synthesis methods to ensure sufficient statistical power.¹² To facilitate the reproducing of the data used in meta-analyses, we considered only systematic reviews that provided a 2×2 table of data of event counts and sample sizes in intervention and control arms of each included study in forest plots or tables. Meta-analyses of proportions and network meta-analyses were not considered. Safety outcomes with continuous type were also not considered because continuous outcomes have been investigated by others.⁴⁻⁶ Systematic reviews in languages other than English and Chinese were excluded.

Two review authors screened the literature independently (XQ and CX). Titles and abstracts were screened first, and then the full texts of the relevant publications were read. For screening of titles and abstracts, only records excluded by both reviewer authors were excluded. Any disagreements were solved by discussion between the two authors.

Data collection

Metadata from the randomised controlled trials were collected from eligible systematic reviews. The following items were extracted: name of the first author, outcome of interest, number of participants and number of events in each group, and detailed information of intervention (eg, type of intervention, dosage, and duration) and control groups. Four experienced authors (CX, TQ, XQ, and HM) extracted the data by dividing the eligible systematic reviews into four equal portions by the initial of the first author, and each extractor led one portion. We had a pilot training for the above items to be extracted through the first systematic review before the formal data extraction. Finally, data were initially checked by

the same extractors for their own portion and double checked by the other two authors separately (CX and TQ) to confirm that no errors were present from the data extraction (supplementary table 1).

Additionally, based on the reporting of each systematic review, we collected the following information according to the good practice guideline of data extraction¹³: how the data were extracted (eg, two extractors independently), whether a protocol was available, whether a clear data extraction plan was made in the protocol, whether any solution for anticipant problems in data extraction was outlined in the protocol, whether a standard data extraction form was used, whether the data extraction form was piloted, whether the data extractors were trained, the expertise of the data extractors, and whether they documented any details of data extraction. CX also collected the methods (eg, inverse variance, fixed effect model), effect estimators used for meta-analysis, and the effect with a confidence interval of the meta-analysis. TQ checked the extraction and any disagreements were solved by discussion between these two authors (with detailed records).

Reproducibility

After we extracted the data from the included systematic reviews, the four authors who extracted the data were required to reproduce the data used in meta-analyses from the original sources, which included the original publications of the randomised controlled trials and their supplementary files, ClinicalTrials.gov, and websites of the pharmaceutical companies. When the trial data used in a meta-analysis were not the same as had been reported from one of its original sources, we classified it as a “data extraction error.” If the authors of the systematic review reported that they had contacted the authors of the original paper and successfully obtained related data, we did not consider the discrepancy a data extraction error, even if the data were not the same as any of the original sources.¹⁴ We recorded the details of the location (that is, event count (r) or total sample size (n), intervention (1) or control group (2), which are marked as r1/n1/r2/n2) and the reasons why the data could not be reproduced. Any enquires or issues that would affect our assessment were resolved by group discussion of the four extractors. Again, reproducibility was initially checked by the data extractors for their own portions of the workload. After data extraction and reproduction, the lead author (CX) and TQ separately conducted two further rounds of double checking (supplementary table 1).¹⁵

Outcomes

Our primary outcome of this study was the proportions of the data extraction errors at the study level, the meta-analysis level, and the systematic review level. The secondary outcomes were the proportion of studies with data extraction error within each meta-analysis and the proportion of meta-analyses with data extraction error within each systematic review.

Statistical analysis

We summarised the frequency of data extraction errors at the study level, the meta-analysis level, and the systematic review level to estimate the aforementioned proportions. For the study level, the frequency was the total number of randomised controlled trials with data extraction errors. For the meta-analysis level, the frequency was the number of meta-analyses with at least one study with data extraction errors. For the systematic review level, the frequency was the number of systematic reviews with at least one meta-analysis with data extraction errors.

Considering that clustering effects might be present (owing to the diverse expertise and experience of the four people who extracted data), a generalised linear mixed model was further used to estimate the extractor adjusted proportion.¹⁶ The potential associations among duplicated data extraction, development of a protocol in advance, and data extraction errors based on systematic review level were examined using multivariable logistic regression. Other recommendations listed in good practice guidelines were not examined because most systematic reviews did not report the information.

Because data extraction errors could have different mechanisms (eg, calculation errors and unclear definition of the outcome), we empirically classified these errors into different types on the basis of consensus after summarising the error information (supplementary fig 1). Then, the percentages of the different types of errors among the total number of errors were summarised based on the study level. We conducted a post-hoc comparison of the difference of the proportions of the total and the subtype errors by two types of interventions: drug interventions; and non-drug interventions (eg, surgery and device). We did this because the safety outcomes are greatly different for these two types of interventions based on our word cloud analysis (supplementary fig 2).

To investigate the potential effect of data extraction errors on the results, we used the same methods and effect estimators that the authors reported based on the corrected dataset. We repeated these meta-analyses and compared the new results to the original results. Some meta-analyses contained errors related to unclear definitions of the outcomes (that is, the ambiguous error defined in table 1). The true number of events is therefore impossible for readers to determine, as is the ability to investigate the effect on the results based on the full empirical dataset. Therefore, we used a subset with meta-analyses free of this type of ambiguous errors. We prespecified a 20% change of the magnitude or more of the effects as moderate impact and a 50% change or more as large impact. We also summarised the proportion of change on the direction of the effects and on the significance of the P value.

Missing data would occur when the original data sources were not available for a few randomised controlled trials in which we were unable to verify data accuracy. For our sensitivity analysis, which investigated the robustness of the results, we removed

Table 1 | Descriptions of the different types of errors during the data extraction

Type of errors	Description	Real life example
Numerical error	Extracted numerical values were incorrect, potentially due to typo, calculation error, or extraction of data of another outcome.	The meta-analysis ¹⁷ investigated an outcome of congestive heart failure. The RCT by Piccart-Gebhart et al ¹⁸ was included under the outcome; the number of participants with heart failure in the intervention group used in the meta-analysis was 482, while the correct number reported in the trial was 68. The number of 482 belongs to another outcome of treatment withdrawals for toxicity.
Ambiguous error	Extracted data could not be reproduced from all available sources (unknown whether it is correct or not) owing to ambiguous definitions of the outcomes, while the review authors did not specify how the data were obtained or calculated. In some situations, the outcomes could not be found in the original study and related materials (eg, supplementary file, ClinicalTrials.gov).	Again, from the same meta-analysis, ¹⁷ with the same outcome above. The authors also included the RCT by Blackwell et al, ¹⁹ and their extracted data were of nine and 17 people with heart failure in the two arms, separately. However, Blackwell's study did not have the outcome of congestive heart failure; the authors only documented the total number of cardiac events of the two arms as 14. The origin of numbers nine and 17 is unclear.
Zero assumption error	This error was a special case of ambiguous error, and generally occurs in safety outcomes. The outcome was not reported in the original study and related materials (eg, supplementary file, ClinicalTrials.gov), while the review authors assumed that no event occurred.	The meta-analysis ²⁰ had an outcome of severe infection. The RCT by Mubarak et al ²¹ was included, and the extracted data of number of participants with severe infection in the meta-analysis in two arms are zero. However, according to their definition, no outcome can be regarded as severe infection in the RCT.
Mismatching error	The extracted data were incorrectly matched to the intervention and exposure groups, but the numerical values were correct. This error could occur in any cells of the summarised table.	A meta-analysis ²² included an outcome of all-grade decrease in left ventricular ejection fraction. The RCT by Flaherty et al ²³ was included. In the metadata, the total participants in the combination intervention group versus monotherapy group were recorded as 53 and 55, respectively. However, in the trial, the numbers were actually 55 and 53 for these two groups, respectively.
Misidentification*	Review authors did not correctly identify the eligibility of the included studies for a certain outcome, categorised into three situations: 1) study reported the outcome and related data but was not included in the meta-analysis of the outcome (this does not apply for double-zero studies as classical methods will exclude such studies by default, although other sophisticated methods can include them for meta-analysis); 2) study with the PICOS did not meet the defined criteria, but was included in the meta-analysis (theoretically, driven from situation 1); and 3) duplicated studies were included as different studies within the same outcome.	The meta-analysis ²⁴ contained an outcome of diarrhoea. Among the included studies, the study by Motzer et al ²⁵ reported the outcome of diarrhoea, but it was not included in the meta-analysis. Also, in the same meta-analysis, the meta-analysis investigated an outcome of high-grade rash. The publications by Hodi et al ²⁶ and Postow et al ²⁷ were both included, but they were referred to the same RCT (ClinicalTrials.gov identifier NCT01927419).

PICOS=patient or population, intervention, comparison, and outcomes. RCT=randomised controlled trial.

*This error is an identification error, and to be strict, not a data extraction error, but it results in errors for the final meta-analytical data and could affect the final pooled effect.

these studies. We used Stata 15/SE for the data analysis. The estimation of the proportions was based on the `meglm` command under the Poisson function with the `log link`²⁸; we set $\alpha=0.05$ as the significance level. We performed the re-evaluation of the meta-analyses by the `admetan` command in Stata and verified by `metafor` command in R 3.5.1 software, and Excel 2013 was used for visualisation.

Patient and public involvement

As this was a technical paper to assess related methodology for data extraction errors of evidence synthesis practice and the impacts of these errors on the analysis, no patients or public members were involved, nor was funding available for the same reason.

Results

Overall, we screened 18 636 records, and initially identified 456 systematic reviews of adverse events.⁹ After a further screening of the full texts, 102 were excluded for having non-randomised studies of intervention and 153 were excluded for not having a pairwise meta-analysis, having fewer than five studies in all meta-analyses, or not reporting 2×2 table data used in meta-analyses (supplementary table 3). As such, 201 systematic reviews were included in the current study (fig 1).

Among the 201 systematic reviews, 156 referred to drug interventions and the other 45 were non-

drug interventions (60% were surgical or device interventions). From the 201 systematic reviews, we identified 829 pairwise meta-analyses with at least five studies involving 10 386 randomised controlled trials. The data extraction error by the four data extractors ranged from 0.5% to 5.4% based on the double-checking process, which suggested that this study had high quality data extraction (supplementary table 1).

Among the 201 systematic reviews, based on the reporting information, 167 (83.1%) stated that they had two data extractors, 31 (15.4%) did not report such information, two (1%) cannot be judged owing to insufficient information, and only one (0.5%) reported that the data were extracted by one person. Fifty four (26.9%) systematic reviews reported a protocol that was developed in advance, whereas most (147, 73.1%) did not report whether they had a protocol. For those with protocols, 32 (59.3%) of 54 had a clear plan for data extraction and 22 (40.7%) outlined a potential solution for anticipant problems for data extraction. Sixty six (32.8%) systematic reviews used a standard data extraction form, while most (135, 67.2%) did not report this information. For the systematic reviews that used a standard extraction form, six (8.8%) piloted this process. No systematic reviews reported the information of whether the data extractor was trained or the expertise of the data extractor. Only seven (3.5%) of 201 systematic reviews documented the details of the data extraction process.

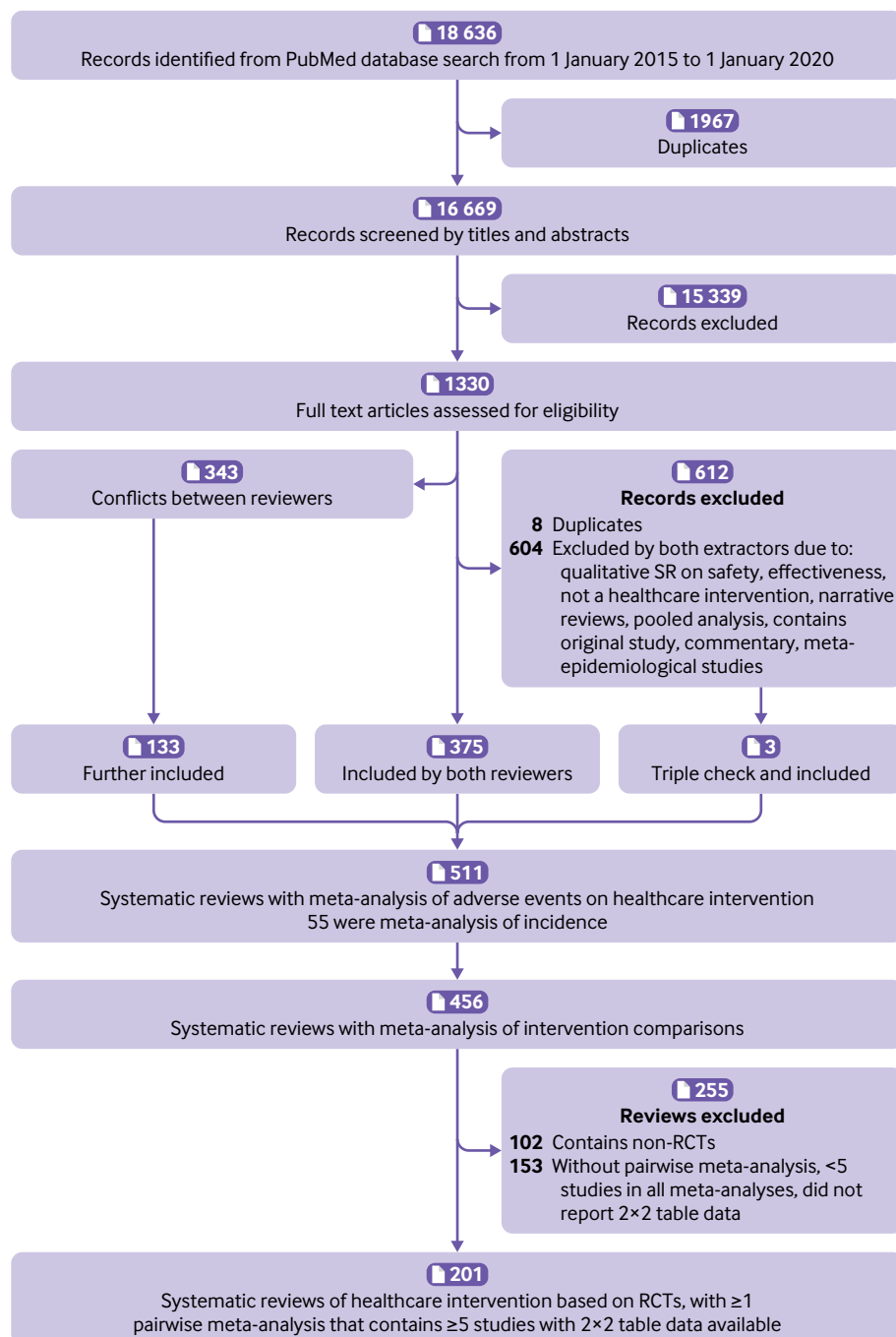


Fig 1 | Flowchart for selection of articles. RCT=randomised controlled trial

Reproducibility of the data extraction

For the reproducibility of the data used in these meta-analyses, at the study level, we could not reproduce 1762 (17.0%) of 10386 studies with an extractor addressed proportion of 15.8%. At the meta-analysis level, 554 (66.8%) of 829 meta-analyses had at least one randomised controlled trial with data extraction errors, with an extractor addressed proportion of 65.5% (fig 2). For meta-analyses with data extraction errors in at least one study, the proportion of studies with data extraction errors within a meta-analysis ranged from 1.9% to 100%, with a median value of 20.6% (interquartile range 12.5-40.0; fig 2).

At the systematic review level, 171 (85.1%) of 201 systematic reviews had at least one meta-analysis with data extraction errors, with an extractor addressed proportion of 85.1% (fig 3). For systematic reviews with data extraction errors in at least one meta-analysis, the proportion of meta-analyses with data extraction errors within a systematic review ranged from 16.7% to 100.0%, with a median value of 100.0% (interquartile range 66.7-100; fig 3).

Based on the multivariable logistic regression, those systematic reviews that reported duplicated data extraction or were checked by another author (odds ratio 0.9, 95% confidence interval 0.3 to 2.5, $P=0.83$)

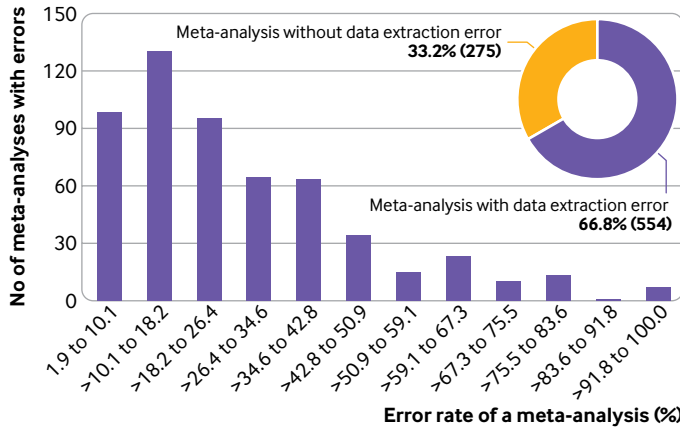


Fig 2 | Data extraction errors at the meta-analysis level. Bar plot is based on studies with data extraction errors (n=554). Error rate within a meta-analysis is calculated by the number of studies with data extraction errors against the total number of studies within a meta-analysis

and developed a protocol in advance (0.7, 0.3 to 1.6, P=0.38) did not show a difference in the odds of errors, but there might be a weak association of errors.

Empirical classification of errors

Based on the mechanism of the data extraction errors, we empirically classified these errors into five types: numerical error, ambiguous error, zero assumption error, mismatching error, and misidentification error. Table 1 provides the definitions of these five types of data extraction errors, with detailed examples.¹⁷⁻²⁷

Numerical error was the most prevalent data extraction error, which accounted for 867 (49.2%) of 1762 errors recorded in the studies (fig 4). The second most prevalent data extraction error was the ambiguous error, accounting for 526 (29.9%) errors. Notably, zero assumption errors accounted for as much as 221 (12.5%) errors. Misidentification accounted for 115 (6.5%) errors and mismatching errors accounted for 33 (1.9%) errors.

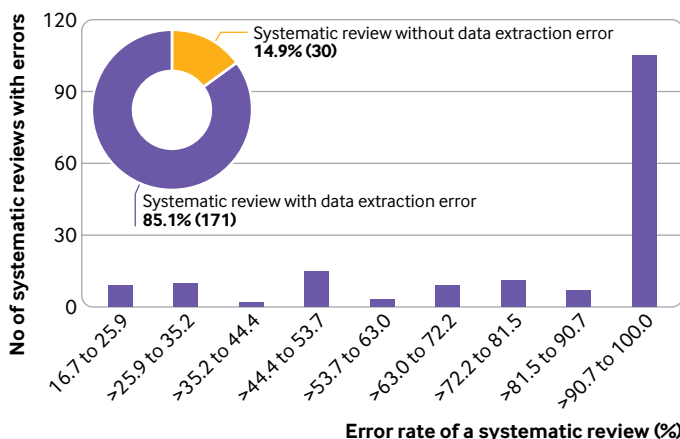


Fig 3 | Data extraction errors at the systematic review level. Bar plot is based on studies with data extraction errors (n=171). Error rate within a systematic review is calculated by the number of meta-analyses with data extraction errors against the total number of meta-analyses within a systematic review

Subgroup analysis by the intervention type suggested that meta-analyses with drug interventions were more likely to have data extraction errors than those involving non-drug interventions: total error (19.9% v 8.9%; P<0.001), ambiguous error (6.1% v 2.4%; P<0.001), numerical error (9.4% v 5.4%; P<0.001), zero assumption error (2.6% v 0.9%; P<0.001), and misidentification errors error (1.5% v 0.1%; P<0.001; supplementary fig 3). Although mismatching errors showed the same pattern, the data were not significantly different (0.4% v 0.2%; P=0.09).

Impact of data extraction errors on the results

After removing meta-analyses with ambiguous errors and without errors, 288 meta-analyses could be used to investigate the impact of data extraction errors on the results (supplementary table 4). Among them, 39 had two or more types of errors (mixed), and 249 had only one type of error (single). For the 249 meta-analyses, 200 had numerical errors, 25 had zero assumption errors, 16 had misidentification errors, and eight had mismatching errors. Because of the limited sample size of each subtype, we only summarised the total impact and the effect grouped by the number of types (that is, single type of error or mixed type of errors).

In total, in terms of the magnitude of the effect, when using corrected data for the 288 meta-analyses, 151 (52.4%) had decreased effects, whereas 137 (47.6%) had increased effects; 37 (12.8%) meta-analyses had moderate changes (with ≥20% changes), and 13 (4.5%) had large changes (with ≥50% changes) in the effect estimates (fig 5). For those 37 studies with moderate changes, the effects in 26 (70.2%) increased, whereas those in 11 (29.7%) decreased when using corrected data. For those 13 studies with large changes, nine (69.2%) showed increased effects, whereas four (30.8%) showed decreased effects. Ten (3.5%) of the 288 meta-analyses had changes in the direction of the effect, and 19 (6.6%) of the 288 meta-analyses changed the significance of the P value. For those studies that had changes in the direction, two (20.0%) of 10 changed from beneficial to harmful effects, and eight (80.0%) of the 10 changed from harmful to beneficial effects. For studies that changed in significance, 10 (52.6%) of 19 changed from non-significance to significance, and nine (47.4%) of 19 changed from significance to non-significance. Some examples are presented in table 2. Studies with two or more types of errors had higher proportions of moderate (28.2% v 10.4%, P=0.002) and large changes (12.8% v 3.2%, P=0.01; fig 5) than did with only a single error.

Sensitivity analysis

For 318 (3.1%) of 10386 studies in the total dataset, we could not obtain full texts or had no access to the original data source to verify data accuracy. After treating them as missing values and removing them from the analyses, no changes were obvious in the proportions of data extraction errors: 16.2% for the study level, 65.7% for the meta-analysis level, and

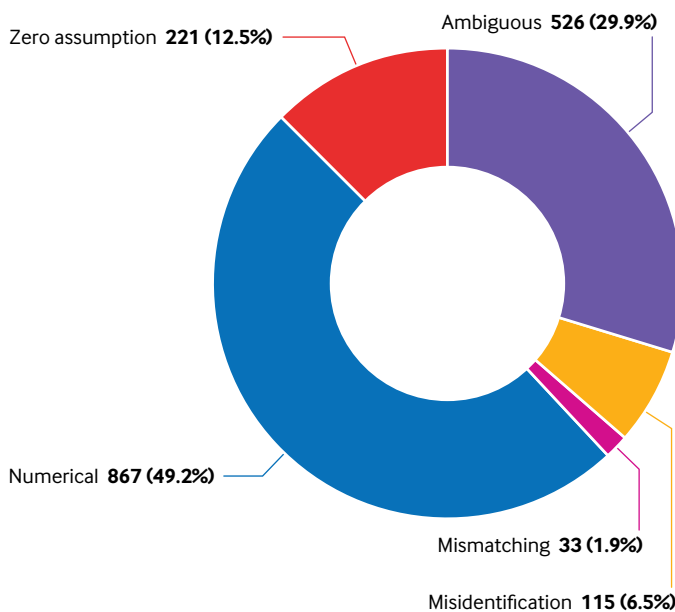


Fig 4 | Proportion of 1762 studies classified by five types of data extraction error

85.1% for the systematic review level (addressed by extractor clustering effects).

Discussion

Principal findings

We investigated the reproducibility of the data extraction of 829 pairwise meta-analyses within 201 systematic reviews of safety outcomes by repeating the data extraction from all the included studies. Our results suggested that as much as 85% of the systematic reviews had data extraction errors in at least one meta-analysis. From the point of meta-analysis level, as many as 67% of the meta-analyses had at least one study with data extraction error. Our findings support the seriousness of the findings from the survey conducted by *Nature* regarding reproducibility of basic science research (70%).¹ At the systematic review level, the problem is even more serious.

Our subgroup analysis showed that data for the safety outcomes of drug interventions had a higher proportion of extraction error (19.9%) than did data for non-drug interventions (8.9%). One important

reason could be that safety outcomes of different types of interventions vary considerably (supplementary fig 1). For non-drug interventions, most interventions were surgical or a device, where safety outcomes might be easier to define. For example, a common safety outcome in surgical intervention is bleeding during surgery, whereas a common outcome of drug interventions is liver toxicity, which might be more complex to define and measure. Additionally, the reporting of adverse events in surgical interventions heavily relies on the surgical staff, whereas for adverse events of a drug, patients might also participate in the reporting process. Selective reporting could exist for adverse events of surgical interventions without patients' participation,²⁹ and mild but complex adverse events (eg, muscular pain) might be neglected and further make reported adverse events appear more straightforward.

We classified data extraction errors into five types based on the mechanism. Based on this classification, we further found that numerical errors, ambiguous errors, and zero assumption errors accounted for 91% of the total errors. The classification and related findings are important because these data provide a theoretical basis for researchers to develop implementation guidelines and help systematic review authors to reduce the frequency of errors during the data extraction process. Another important reason for data extraction errors might be the poor reporting of adverse events in randomised controlled trials, which have varying terminology, poorly defined categories, and diverse data sources.³⁰⁻³² If trials did not clearly define an adverse outcome and report it transparently, then systematic review authors would face difficulties during data extraction and the process would be prone to errors, especially with regard to the ambiguous types. We believe that with proper implementation guidance and more explicit trial reporting guidelines for adverse events, these errors can substantially be reduced.

The classification also provides a theoretical basis for methodologists to investigate the potential impact of different types of data extraction errors on the results. The impact of different types of errors on the results could vary. For example, the zero assumption error is expected to push the final effect towards the null when related studies have balanced sample sizes in two arms.³³ The mismatching error has a similar effect because the error pushes the effect towards the opposite direction. By contrast, the direction of the effect is difficult to predict in the other three types of errors. In our empirical data, because of the small number of meta-analyses in each category, we were unable to investigate the impact of each single type of error on the results. One of the most important reasons is that many meta-analyses have ambiguous errors. Nevertheless, we were able to compare the effect of multiple error types against a single error type for meta-analyses. Our results suggested that meta-analyses with multiple types of data extraction errors were prone to be affected. Because different methods can

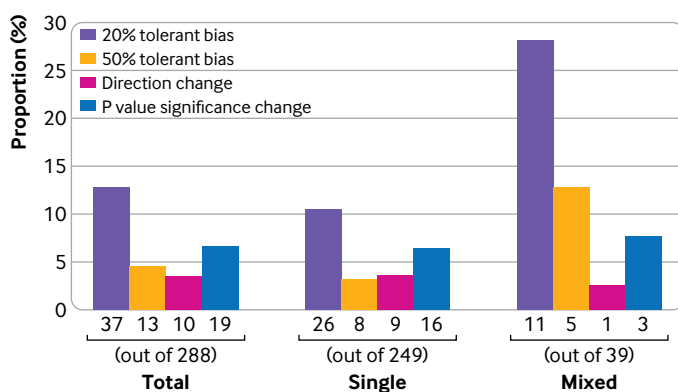


Fig 5 | Impact of data extraction errors on results

Table 2 | Examples of changes in the effects and significance when using corrected data

Example	Original result (with error)		Corrected result (without error)		Difference*	
	Effect (95% CI)	P value	Effect (95% CI)	P value	Relative effect (%)	P value of absolute data
Moderate change						
Increased effect (risk ratio)	2.51 (1.21 to 5.22)	0.01	3.19 (1.34 to 7.59)	0.01	27.17	0
Decreased effect (odds ratio)	1.59 (0.63 to 4.02)	0.33	1.17 (0.42 to 3.25)	0.76	-26.40	0.43
Large change						
Increased effect (odds ratio)	1.21 (1.05 to 1.40)	0.01	3.24 (1.24 to 8.43)	0.02	167.77	0.01
Decreased effect (risk ratio)	0.17 (0.13 to 0.22)	<0.01	0.09† (0.04 to 0.17)	<0.01	-50.00	0
Changed of the effect direction						
Benefit to harm (risk ratio)	0.92 (0.78 to 1.08)	0.32	1.11 (0.93 to 1.32)	0.25	20.70	-0.07
Harm to benefit (odds ratio)	1.14 (0.95 to 1.35)	0.14	0.94 (0.83 to 1.06)	0.29	-17.50	0.15
Change of significance						
Significance to non-significance (odds ratio)	3.82 (1.27 to 11.45)	0.02	0.93 (0.63 to 1.37)	0.71	-75.65	0.73
Non-significance to significance (odds ratio)	1.35 (0.75 to 2.44)	0.32	1.65 (1.04 to 2.63)	0.03	22.20	-0.29

*Difference in effects calculated by: $(\theta_{\text{corrected}} - \theta_{\text{original}}) \div \theta_{\text{original}} \times 100\%$, where θ is the estimated pooled effects. †0.09 has been rounded from 0.085.

vary on this assumption (eg, two-stage methods with odds ratios assume that double-zero studies are non-informative⁹), the use of different synthesis methods and effect estimates might have different impacts.^{34 35} The impact of data extraction errors on the results is expected to be thoroughly investigated by simulation research.

Strengths and limitations

This large empirical study investigates the reproducibility of the data extraction of systematic reviews of adverse events and its impact on the results of related meta-analyses. The findings of our study pose a serious warning to the community that much progress is needed to achieve high quality, evidence-based practice. We are confident that the results of our findings are reliable because the data have been through five rounds of cross-checking within our tightly structured collaborative team. Additionally, this study is the first time that data extraction errors were defined based on their mechanism, which we think will benefit future methodological research in this area.

However, some limitations are still present. Firstly, owing to the large amount of work, data collection was divided into four portions, and each portion was conducted by a separate author. Although all authors undertook pilot training in advance, their judgments might still differ. Nevertheless, our analysis used the generalised linear mixed model, which accounted for the potential clustering effect by different extractors, of which the findings suggested no obvious impact on the results. Secondly, our study covered only systematic reviews published in the five year period from 2015 to 2020; therefore, the validity of the data extraction in earlier studies is unclear. Whether this issue has deteriorated or improved over time could not be assessed. Thirdly, a small proportion of studies could not have reproducibility checked, and these studies were treated as if no data extraction errors existed, which could lead to a slight underestimation of data extraction error overall.³⁶

Furthermore, we only focused on systematic reviews of randomised controlled trials and did not consider observational studies. Because the sample sizes of

randomised controlled trials tend to be small, the impact might be exacerbated. Finally, poor reporting has been commonly investigated in literature^{37 38}; owing to the limited information of the data extraction process reported by review authors, we could not fully investigate the association between good practice recommendations and the likelihood of data extraction. For the same reason, the association among duplicated data extraction, development of a protocol in advance, and data extraction errors should be interpreted with caution. Further studies based on randomised controlled design might be helpful. However, we believe these limitations have little impact on our main results and conclusions.

Conclusions

Systematic reviews of adverse events face serious issues in terms of the reproducibility of their data extraction. Prevalence of data extraction errors is high among these systematic reviews and these errors could lead to the changing of the conclusions and further mislead the healthcare practice. A series of expanded reproducibility studies on other types of meta-analyses might be useful for further evidence-based practice. Additionally, implementation guidelines on data extraction for systematic reviews are urgently required to help future review authors improve the validity of their findings.

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We thank Riaz Qureshi from Johns Hopkins University and Zhang Jiaxin from Guizhou Provincial People's Hospital for their comments and edits on our protocol. We also thank Lu Cuncun from Lanzhou University for developing the search strategy for the whole project.

Contributors: CX and SV conceived and designed the study; CX collected the data, analysed the data, and drafted the manuscript; ZXQ and CX screened the literature; YTQ, CX, DHM, ZXQ extracted and reproduced the data; YTQ and CX contributed to the data checking; CX, SV, LFK, LL, LZ, and YL provided methodological comments, and revised the manuscript. All authors approved the final version to be published. CX and SV are the study guarantors. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

Funding: LFK is funded by an Australian National Health and Medical Research Council Fellowship (APP1158469). LL is funded by the US National Institutes of Health/National Library of Medicine grant R01 LM012982 and the National Institutes of Health/National Institute of Mental Health grant R03 MH128727. The funding body had no role in any process of the study (that is, study design, analysis, interpretation of data, writing of the report, and decision to submit the article for publication).

Competing interests: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare: support from the Australian National Health and Medical Research Council Fellowship, US National Institutes of Health, National Library of Medicine, and National Institute of Mental Health for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work.

Ethical approval: Not required.

Data sharing: A subset of the data can be found at <https://osf.io/czyqa/>. The dataset could be obtained from the first author (xuchang2016@runbox.com) or the corresponding author (svohra@ualberta.ca) on request.

The lead author affirms that the manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as originally planned (and, if relevant, registered) have been explained.

Dissemination to participants and related patient and public communities: We plan to present our findings at national and international scientific meetings and to use social media outlets to disseminate findings.

Provenance and peer review: Not commissioned; externally peer reviewed.

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- Baker M. 1,500 scientists lift the lid on reproducibility. *Nature* 2016;533:452-4. doi:10.1038/533452a
- Chandler J, Cumpston M, Thomas J, et al. Introduction. In: Higgins JPT, Thomas J, Chandler J, eds. *Cochrane Handbook for Systematic Reviews of Interventions version 6.2 (updated February 2021)*. Cochrane, 2021. www.training.cochrane.org/handbook.
- Mathes T, Klaffen P, Pieper D. Frequency of data extraction errors and methods to increase data extraction quality: a methodological review. *BMC Med Res Methodol* 2017;17:152. doi:10.1186/s12874-017-0431-4
- Jones AP, Remington T, Williamson PR, Ashby D, Smyth RL. High prevalence but low impact of data extraction and reporting errors were found in Cochrane systematic reviews. *J Clin Epidemiol* 2005;58:741-2. doi:10.1016/j.jclinepi.2004.11.024
- Götzsche PC, Hróbjartsson A, Maric K, Tendal B. Data extraction errors in meta-analyses that use standardized mean differences. *JAMA* 2007;298:430-7. doi:10.1001/jama.298.4.430
- Tendal B, Higgins JP, Juni P, et al. Disagreements in meta-analyses using outcomes measured on continuous or rating scales: observer agreement study. *BMJ* 2009;339:b3128. doi:10.1136/bmj.b3128
- Austin PC, Steyerberg EW. Events per variable (EPV) and the relative performance of different strategies for estimating the out-of-sample validity of logistic regression models. *Stat Methods Med Res* 2017;26:796-808. doi:10.1177/0962280214558972
- Ju K, Lin L, Chu H, Cheng LL, Xu C. Laplace approximation, penalized quasi-likelihood, and adaptive Gauss-Hermite quadrature for generalized linear mixed models: towards meta-analysis of binary outcome with sparse data. *BMC Med Res Methodol* 2020;20:152. doi:10.1186/s12874-020-01035-6
- Xu C, Zhou X, Zorzela L, et al. Utilization of the evidence from studies with no events in meta-analyses of adverse events: an empirical investigation. *BMC Med* 2021;19:141. doi:10.1186/s12916-021-02008-2
- Zorzela L, Golder S, Liu Y, et al. Quality of reporting in systematic reviews of adverse events: systematic review. *BMJ* 2014;348:f7668. doi:10.1136/bmj.f7668
- Zorzela L, Loke YK, Ioannidis JP, et al, PRISMAHarms Group. PRISMA harms checklist: improving harms reporting in systematic reviews. *BMJ* 2016;352:i157. doi:10.1136/bmj.i157
- Jackson D, Turner R. Power analysis for random-effects meta-analysis. *Res Synth Methods* 2017;8:290-302. doi:10.1002/jrsm.1240
- Taylor KS, Mahtani KR, Aronson JK. Summarising good practice guidelines for data extraction for systematic reviews and meta-analysis. *BMJ Evid Based Med* 2021;26:88-90. doi:10.1136/bmjebm-2020-111651
- Berstock J, Beswick A. Importance of contacting authors for data on adverse events when compiling systematic reviews. *BMJ* 2014;348:g1394. doi:10.1136/bmj.g1394
- Büchter RB, Weise A, Pieper D. Development, testing and use of data extraction forms in systematic reviews: a review of methodological guidance. *BMC Med Res Methodol* 2020;20:259. doi:10.1186/s12874-020-01143-3
- Lin L, Xu C, Chu H. Empirical Comparisons of 12 Meta-analysis Methods for Synthesizing Proportions of Binary Outcomes. *J Gen Intern Med* 2022;37:308-17. doi:10.1007/s11606-021-07098-5
- Hao S, Tian W, Gao B, et al. Does dual HER-2 blockade treatment increase the risk of severe toxicities of special interests in breast cancer patients: A meta-analysis of randomized controlled trials. *Oncotarget* 2017;8:19923-33. doi:10.18632/oncotarget.15252
- Piccari-Gebhart M, Holmes E, Baselga J, et al. Adjuvant Lapatinib and Trastuzumab for Early Human Epidermal Growth Factor Receptor 2-Positive Breast Cancer: Results From the Randomized Phase III Adjuvant Lapatinib and/or Trastuzumab Treatment Optimization Trial. *J Clin Oncol* 2016;34:1034-42. doi:10.1200/JCO.2015.62.1797
- Blackwell KL, Burstein HJ, Storniolo AM, et al. Overall survival benefit with lapatinib in combination with trastuzumab for patients with human epidermal growth factor receptor 2-positive metastatic breast cancer: final results from the EGF104900 Study. *J Clin Oncol* 2012;30:2585-92. doi:10.1200/JCO.2011.35.6725
- Tong S, Fan K, Jiang K, et al. Increased risk of severe infections in non-small-cell lung cancer patients treated with pemetrexed: a meta-analysis of randomized controlled trials. *Curr Med Res Opin* 2017;33:31-7. doi:10.1080/03007995.2016.1232705
- Mubarak N, Gaafar R, Shehata S, et al. A randomized, phase 2 study comparing pemetrexed plus best supportive care versus best supportive care as maintenance therapy after first-line treatment with pemetrexed and cisplatin for advanced, non-squamous, non-small cell lung cancer. *BMC Cancer* 2012;12:423. doi:10.1186/1471-2407-12-423
- Mincu RI, Mahabadi AA, Michel L, et al. Cardiovascular adverse events associated with BRAF and MEK Inhibitors: a systematic review and meta-analysis. *JAMA Netw Open* 2019;2:e198890. doi:10.1001/jamanetworkopen.2019.8890
- Flaherty KT, Infante JR, Daud A, et al. Combined BRAF and MEK inhibition in melanoma with BRAF V600 mutations. *N Engl J Med* 2012;367:1694-703. doi:10.1056/NEJMoa1210093
- Zhang B, Wu Q, Zhou YL, Guo X, Ge J, Fu J. Immune-related adverse events from combination immunotherapy in cancer patients: A comprehensive meta-analysis of randomized controlled trials. *Int Immunopharmacol* 2018;63:292-8. doi:10.1016/j.intimp.2018.08.014
- Motzer RJ, Tannir NM, McDermott DF, et al, CheckMate 214 Investigators. Nivolumab plus ipilimumab versus sunitinib in advanced renal-cell carcinoma. *N Engl J Med* 2018;378:1277-90. doi:10.1056/NEJMoa1712126
- Hodi FS, Chesney J, Pavlick AC, et al. Combined nivolumab and ipilimumab versus ipilimumab alone in patients with advanced melanoma: 2-year overall survival outcomes in a multicentre, randomised, controlled, phase 2 trial. *Lancet Oncol* 2016;17:1558-68. doi:10.1016/S1470-2045(16)30366-7
- Postow MA, Chesney J, Pavlick AC, et al. Nivolumab and ipilimumab versus ipilimumab in untreated melanoma. *N Engl J Med* 2015;372:2006-17. doi:10.1056/NEJMoa1414428

- 28 Xu C, Furuya-Kanamori L, Lin L. Synthesis of evidence from zero-events studies: A comparison of one-stage framework methods. *Res Synth Methods* 2022;13:176-89. doi:10.1002/jrsm.1521
- 29 Saini P, Loke YK, Gamble C, Altman DG, Williamson PR, Kirkham JJ. Selective reporting bias of harm outcomes within studies: findings from a cohort of systematic reviews. *BMJ* 2014;349:g6501. doi:10.1136/bmj.g6501
- 30 Pitrou I, Boutron I, Ahmad N, Ravaud P. Reporting of safety results in published reports of randomized controlled trials. *Arch Intern Med* 2009;169:1756-61. doi:10.1001/archinternmed.2009.306
- 31 Hodkinson A, Kirkham JJ, Tudur-Smith C, Gamble C. Reporting of harms data in RCTs: a systematic review of empirical assessments against the CONSORT harms extension. *BMJ Open* 2013;3:e003436. doi:10.1136/bmjopen-2013-003436
- 32 Favier R, Crépin S. The reporting of harms in publications on randomized controlled trials funded by the "Programme Hospitalier de Recherche Clinique," a French academic funding scheme. *Clin Trials* 2018;15:257-67. doi:10.1177/1740774518760565
- 33 Kuss O. Statistical methods for meta-analyses including information from studies without any events-add nothing to nothing and succeed nevertheless. *Stat Med* 2015;34:1097-116. doi:10.1002/sim.6383
- 34 Xu C, Furuya-Kanamori L, Zorzela L, Lin L, Vohra S. A proposed framework to guide evidence synthesis practice for meta-analysis with zero-events studies. *J Clin Epidemiol* 2021;135:70-8. doi:10.1016/j.jclinepi.2021.02.012
- 35 Bender R, Friede T, Koch A, et al. Methods for evidence synthesis in the case of very few studies. *Res Synth Methods* 2018;9:382-92. doi:10.1002/jrsm.1297
- 36 Kahale LA, Khamis AM, Diab B, et al. Potential impact of missing outcome data on treatment effects in systematic reviews: imputation study. *BMJ* 2020;370:m2898. doi:10.1136/bmj.m2898
- 37 Jung RG, Di Santo P, Clifford C, et al. Methodological quality of COVID-19 clinical research. *Nat Commun* 2021;12:943. doi:10.1038/s41467-021-21220-5
- 38 Abbott R, Bethel A, Rogers M, et al. Characteristics, quality and volume of the first 5 months of the COVID-19 evidence synthesis infodemic: a meta-research study. *BMJ Evid Based Med* 2021;bmjebm-2021-111710. doi:10.1136/bmjebm-2021-111710.

Web appendix: Supplementary materials