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ORIGINAL ARTICLE

CONSORT Harms 2022 statement, explanation, and elaboration: updated guideline for the reporting of harms in randomized trials

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Abstract

Randomized controlled trials remain the reference standard for healthcare research on effects of interventions, and the need to report both benefits and harms is essential. The Consolidated Standards of Reporting Trials (the main CONSORT) statement includes one item

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Patient and public involvement: Patients were invited to complete the Delphi surveys and given the opportunity to provide inputs and critical review.

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on reporting harms (i.e., all important harms or unintended effects in each group). In 2004, the CONSORT group developed the CONSORT Harms extension; however, it has not been consistently applied and needs to be updated. Here, we describe CONSORT Harms 2022, which replaces the CONSORT Harms 2004 checklist, and shows how CONSORT Harms 2022 items could be incorporated into the main CONSORT checklist. Thirteen items from the main CONSORT were modified to improve harms reporting. Three new items were added. In this article, we describe CONSORT Harms 2022 and how it was integrated into the main CONSORT checklist and elaborate on each item relevant to complete reporting of harms in randomized controlled trials. Until future work from the CONSORT group produces an updated checklist, authors, journal reviewers, and editors of randomized controlled trials should use the integrated checklist presented in this paper. © 2023 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

Keywords: Randomized clinical trials; Harms; Adverse Effects; Adverse Events; Adverse Drug Reaction; Side Effects; Reporting guideline; Transparency; Reproducibility; Checklist

Randomized controlled trials are the reference standard among study designs to investigate the benefits of interventions. These trials are the foundation for regulatory approval of drugs and are also important when evaluating surgical procedures, medical devices, psychological and behavioural interventions, social interventions, and complementary therapies. Ideally, randomized controlled trials should evaluate not only potential benefits of interventions but also potential harms. However, these trials are often limited in their ability to evaluate harms because of the short duration of intervention and follow-up, restricted study populations (e.g., excluding participants with comorbidities or receiving co-interventions), and lack of statistical power to assess rare events [1–4]. Nevertheless, prospectively collected data about harms in randomized controlled trials are important to inform knowledge synthesis and patient and provider decisions. The randomized design offers a clear advantage and a unique opportunity to study harms in a controlled setting. Despite these strengths, trials often fail to report harms [5,6], even serious ones. For example, more than half of the deaths and suicides occurring in trials of psychiatric drugs were not reported in published trial reports [7].

The insufficient reporting of harms outcomes might be the result of inadequate planning and design compounded with major challenges in fully reporting diverse adverse events within a limited amount of space in journal articles. However, harms can seriously affect a patient's quality of life [8] and treatment adherence [9,10] and increase financial costs to patients and health systems [11]. To allow patients and healthcare providers to make truly informed decisions, randomized controlled trial reports should describe evidence on benefits and harms.

To promote better reporting of harms in randomized controlled trials, the Consolidated Standards of Reporting Trials (hereafter referred to the main CONSORT) statement includes one potentially subjective item on reporting harms (i.e., all important harms or unintended effects in each group; for specific guidance see CONSORT for harms) [12]. However, this item did not do full justice to the importance of harms-related issues [13], and in 2004, the CONSORT group developed an extension providing specific guidance for the complete reporting of harms in

randomized controlled trials (the CONSORT Harms statement) [13]. Although the main CONSORT statement was revised in 2001 and 2010, CONSORT Harms has not been updated since 2004. Moreover, although CONSORT is endorsed by journals of diverse areas of healthcare research, CONSORT Harms is not mentioned explicitly in the submission instructions of key scientific journals [5].

Considering the ongoing inadequacies in reporting harms in randomized controlled trials, a CONSORT Harms initiative was formed to update CONSORT Harms to provide current and improved guidance on a minimum set of items about harms to be reported in trials. Furthermore, to enhance harms reporting in clinical trials, these items should be integrated into the main CONSORT statement instead of remaining a standalone extension. Here, we present the CONSORT Harms 2022 checklist, illustrating how these items can be incorporated into the CONSORT statement. Until the CONSORT statement is updated to incorporate more complete guidance for reporting harms,

Summary points

- Prospectively collected data about harms in randomized controlled trials are important to inform knowledge synthesis and patient and provider decisions.
- Randomized controlled trials should measure and report benefits and harms of health interventions.
- This article describes CONSORT Harms 2022, a guideline to support better reporting of harms in randomized controlled trial publications, and elaborates on reporting guidance for each item relevant to the complete reporting of harms assessed in trials.
- The CONSORT Harms 2022 reporting checklist is described in a way to show how it can be integrated into the main CONSORT checklist. Until future work from the CONSORT group produces an updated checklist, trial authors, journal reviewers, and editors should use the integrated checklist presented in this paper.

CONSORT Harms 2022 should replace CONSORT Harms 2004.

1. Development of CONSORT Harms 2022

We followed the strategy recommended by the EQUATOR (Enhancing the quality and transparency of health research) Network for the development of reporting guidelines [14]. The study protocol was approved by the University of Alberta Health Research Ethics Board on March 2, 2018 (Pro00078962). The approval process included a metaepidemiological overview of the literature on reporting of harms and a modified Delphi survey [15,16], including two online rounds followed by an in-person consensus meeting. Consumers and patient representatives participated in the online Delphi rounds, the in-person meeting, and manuscript preparation; their input was received and incorporated into the final document.

We first conducted a comprehensive metaepidemiological overview to identify studies of harms reporting in randomized controlled trials [17]. We assessed 13 reviews of harms reporting in trials. These reviews assessed 522 trials for their reporting of the items in CONSORT Harms 2004, which varied from 9% to 69% [17]. Reporting of harms improved only slightly when comparing clinical trials published before and after the publication of CONSORT Harms 2004 [17]. Data from this overview showed that checklist items comprising multiple components presented challenges for reporting and identified additional relevant items for reporting harms in trials [17]. Examples of items of CONSORT Harms 2004 including multiple components are item 6 (clarify how harms-related information was collected, including (1) mode of data collection, (2) timing, (3) attribution methods, (4) intensity of ascertainment, and (5) harms-related monitoring and stopping rules) and item 8 (describe plans for analyzing harms, including (1) presentation of absolute risk per arm, (2)

per adverse event type, (3) per grade, (4) per seriousness, (5) appropriate metrics for recurrent events, (6) continuous variables, and (7) scale variables). These findings were used to generate new items and potential modifications to the original CONSORT Harms checklist, resulting in a list of 26 items.

Next, we deployed a modified Delphi process consisting of two online survey rounds followed by an in-person consensus meeting. The online rounds were completed using a questionnaire developed and managed using the REDCap [18] electronic data capture tool hosted and supported by the Women and Children's Health Research Institute at the University of Alberta. Participants were asked to assess the importance of each item on a five-point Likert scale (ranging from 1 = strongly disagree to 5 = strongly agree). An agreement on the inclusion or exclusion of items of the CONSORT checklist was prespecified and considered if 68% of the respondents agreed or strongly agreed with the inclusion or exclusion of the item. Free text comments were collected to guide the revision of the items. Participants invited to complete the online Delphi rounds were identified by their areas of expertise or stakeholder relevance to the initiative (e.g., methodologists, statisticians, epidemiologists, clinicians, journal editors, consumers or patient representatives, and members of the industry and health regulatory agencies). The list of invitees also included researchers involved in the development of CONSORT, CONSORT Harms 2004 and other CONSORT extensions, and Preferred Reporting Items for Systematic reviews and Meta-Analyses and Preferred Reporting Items for Systematic reviews and Meta-Analyses Harms.

1.1. Delphi rounds findings

We collected feedback from 211 to 92 respondents in the first and second Delphi rounds, respectively. In the first Delphi round, an additional question sought participants'

Table 1. Ambiguity of harms related terminology as currently used in publications of randomized controlled trials that should be avoided

| Terms | Potential concerns |
|--------------------------------------|---|
| Anticipated vs. unanticipated events | These terms are often linked to whether the harm event could be anticipated by the mechanism of action of the intervention or previous reports. However, the exact meaning can be unclear, and the terms do not provide information on whether, when, and how the events were collected. |
| Solicited vs. unsolicited events | These terms can be understood as describing events collected by questioning patients about their experiences with regards to their harm events, for example, in an open-ended fashion (unsolicited): "Have you experienced anything abnormal since the last visit?"; or by answering detailed questionnaires about specific events, such as: "Have you experienced feelings of [specific harms] since the last visit?" Different ways to question about harms might lead to different results. Additional details about who inquired about the harms and when might also be relevant. |
| Attributed events | Attribution is the process of determining a causal relationship between an intervention and a specific event. Causation might be difficult to determine [25], and attribution methods might be of limited value if applied inconsistently [26,27]. In addition, the process might be biased if outcome assessors are not blinded to the intervention either by design or because of unblinding during the trial [27,28]. ^a |
| Unintended events | All harms are "unintended" by definition, so this term lacks specific meaning despite being frequently used in randomized controlled trial reports. |

^a If an attribution assessment was conducted and authors choose to report harms accordingly (attributed vs. nonattributed harms), the attribution methods should be described along with information on who completed such an assessment and how. Please refer to item 6c.

Table 2. CONSORT Harms 2022 items vs. main CONSORT 2010 checklist

| Section and topic | Item No. | CONSORT 2010 | CONSORT Harms 2022 |
|------------------------------------|----------------|---|---|
| Title and abstract | | | |
| Title and abstract | 1a | Identification as a randomized trial in the title | — |
| | 1b (modified) | Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts) | Structured summary of trial design, methods, results of outcomes of benefits and harms, and conclusions (for specific guidance see CONSORT for abstracts) |
| Introduction | | | |
| Background and objectives | 2a | Scientific background and explanation of rationale | — |
| | 2b (modified) | Specific objectives or hypotheses | Specific objectives or hypotheses for outcomes of benefits and harms |
| Methods | | | |
| Trial design | 3a | Description of trial design (such as parallel, factorial) including allocation ratio | — |
| | 3b | Important changes to methods after trial commencement (such as eligibility criteria), with reasons | — |
| Participants | 4a | Eligibility criteria for participants | — |
| | 4b | Settings and locations where the data were collected | — |
| Interventions | 5 | The interventions for each group with sufficient details to allow replication, including how and when they were actually administered | — |
| Outcomes | 6a (modified) | Completely defined prespecified primary and secondary outcome measures, including how and when they were assessed | Completely defined prespecified primary and secondary outcomes, for both benefits and harms, including how and when they were assessed |
| | 6b | Any changes to trial outcomes after the trial commenced, with reasons | — |
| | 6c (new) | — | Describe if and how nonprespecified outcomes of benefits and harms were identified, including any selection criteria, if applicable |
| Sample size | 7a | How sample size was determined | — |
| | 7b | When applicable, explanation of any interim analyses and stopping guidelines | — |
| Randomization: sequence generation | 8a | Method used to generate the random allocation sequence | — |
| | 8b | Type of randomization; details of any restriction (such as blocking and block size) | — |
| Allocation concealment mechanism | 9 | Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned | — |
| Implementation | 10 | Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions | — |
| Blinding | 11a (modified) | If done, who was blinded after assignment to interventions (e.g., participants, care providers, those assessing outcomes) and how | If done, who was blinded after assignment to interventions (e.g., participants, care providers, those assessing outcomes of benefits and harms) and how |
| | 11b | If relevant, description of the similarity of interventions | — |

(Continued)

Table 2. Continued

| Section and topic | Item No. | CONSORT 2010 | CONSORT Harms 2022 |
|--|----------------|---|---|
| Statistical methods | 12a (modified) | Statistical methods used to compare groups for primary and secondary outcomes | Statistical methods used to compare groups for primary and secondary outcomes of both benefits and harms |
| | 12b | Methods for additional analyses, such as subgroup analyses and adjusted analyses | — |
| Results | | | |
| Participant flow (a diagram is strongly recommended) | 13a (modified) | For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analyzed for the primary outcome | For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analyzed for outcomes of benefits and harms |
| | 13b | For each group, losses and exclusions after randomization, together with reasons | — |
| Recruitment | 14a (modified) | Dates defining the periods of recruitment and follow-up | Dates defining the periods of recruitment and follow-up for outcomes of benefits and harms |
| | 14b | Why the trial ended or was stopped | — |
| Baseline data | 15 | A table showing baseline demographic and clinical characteristics for each group | — |
| Numbers analyzed | 16 (modified) | For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups | For each group, number of participants (denominator) included in each analysis of outcomes of benefits and harms and whether the analysis was by original assigned groups and if any exclusions were made |
| Outcomes and estimation | 17a (modified) | For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval) | For each primary and secondary outcome of benefits and harms, results for each group, and the estimated effect size and its precision (such as 95% confidence interval) |
| | 17a2 (new) | — | For outcomes omitted from the trial report (benefits and harms), provide rationale for not reporting and indicate where the data on omitted outcomes can be accessed |
| | 17b (modified) | For binary outcomes, presentation of both absolute and relative effect sizes is recommended | Presentation of both absolute and relative effect sizes is recommended, for outcomes of benefits and harms |
| | 17c (new) | — | Report zero events if no harms were observed |
| Ancillary analyses | 18 (modified) | Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing prespecified from exploratory | Results of any other analyses performed for outcomes of benefits and harms, including subgroup analyses and adjusted analyses, distinguishing prespecified from exploratory |
| Harms | 19 | All important harms or unintended effects in each group (for specific guidance see CONSORT for harms) | — |
| Discussion | | | |
| Limitations | 20 (modified) | Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses | Trial limitations, addressing sources of potential bias related to the approach to collecting or reporting data on harms, imprecision, and, if relevant, multiplicity or selection of analyses |
| Generalizability | 21 | Generalizability (external validity, applicability) of the trial findings | — |
| Interpretation | 22 | Interpretation consistent with results, balancing | — |

(Continued)

Table 2. Continued

| Section and topic | Item No. | CONSORT 2010 | CONSORT Harms 2022 |
|-------------------|---------------|---|--|
| | | benefits and harms, and considering other relevant evidence | |
| Other information | | | |
| Registration | 23 | Registration number and name of trial registry | — |
| Protocol | 24 (modified) | Where the full trial protocol can be accessed, if available | Where the full trial protocol and other relevant documents can be accessed, including additional data on harms |
| Funding | 25 | Sources of funding and other support (such as supply of drugs), role of funders | — |

Abbreviations: CONSORT, Consolidated Standards of Reporting Trials.

If relevant, we also recommend reading CONSORT extensions including for adaptive designs, cluster randomized trials, noninferiority and equivalence trials, nonpharmacological treatments, pilot and feasibility studies, randomized crossover trials, and nonpharmacological treatment interventions.

opinions on whether CONSORT Harms should be integrated into the main CONSORT statement or remains a standalone extension; 67% of the respondents recommended integration. All items received many comments and suggestions, including recommendations for inclusion of new items or amending the existing ones.

In the first Delphi round, one item did not reach agreement for inclusion (to identify the assessment of harms in the title). In the second Delphi round, respondents disagreed on the inclusion of one item (to describe methods of attribution to determine causality of adverse events). Several respondents voiced concern that the length of CONSORT would be excessive if numerous harms items were incorporated in addition to CONSORT items in the main checklist. To determine whether adding several harms items to the CONSORT statement would make it excessively long, we developed a draft in which all harms items surveyed in the first round were incorporated into the main CONSORT checklist so that respondents could see the potential text in situ. With few exceptions, the CONSORT items only required minor revision to refer to harms. This draft was provided to respondents in the second Delphi round to facilitate the visualization of the integrated checklists.

The next and final iteration was the in-person consensus meeting held in Edmonton, Canada in September 2019. A subset of 18 researchers who had participated in the online rounds or were part of the steering committee attended a two-day meeting. The composition of participants attending the consensus meeting was comparable to the group that completed the online Delphi rounds (participants included methodologists, statisticians, epidemiologists, clinicians, journal editors, members of industry and health regulatory agencies, and consumers or patient representatives). The group discussed the list of items and comments collected in the online Delphi rounds and followed a nominal group technique [19] to reach consensus. The meeting was documented by two note takers and the content was collated and refined to generate the final checklist and explanation document.

1.2. Terminology

Consistent with the guidance provided in CONSORT Harms 2004, we define harms as the totality of possible adverse consequences of an intervention or therapy; they are the direct opposite of benefits, against which they must be compared [13]. We recommend reporting on harms including information on methods of ascertainment (i.e., whether harms were assessed systematically or nonsystematically). Adverse events and adverse drug reactions are other terms used to describe harms and often used in regulatory research. As defined in the main CONSORT guideline, outcomes in a randomized controlled trial are also distinguished as prespecified and nonprespecified, where prespecified refers to outcomes explicitly defined in the protocol [12]. Therefore, in the context of harms reporting, we recommend considering the two related aspects. First, was the case definition for the harmful outcome prespecified? If yes, how was the harmful outcome monitored (systematic or nonsystematic assessment)?

Harms might be assessed systematically by measuring variables for all participants using standardized clinical examinations, questionnaires, and medical instruments [20]. For systematically assessed harms, the trial report should provide the case definition of the outcome assessed (e.g., insomnia), the specific tool or instrument used (e.g., Insomnia Severity Index), the metric (e.g., change from baseline, final value, time to event), the method of aggregation (e.g., median, proportion), and time points for data collection [21]. If cut-off points associated with increased harm (for harms, assessed as continuous variables such as hormone levels or bone mass index) or categories were analyzed, these should also be defined. The nonsystematic assessment of harms relies on the passive or unstructured reporting of adverse events, such as unprompted self-reporting by participants [22], and can meaningfully complement the systematic assessment of harms, particularly when generating signals for unexpected events.

Other terms to describe harms are not recommended. For instance, the use of “safety” can be a misleading term used

to diminish the importance of harms or to imply the absence of harms. Many trials casually note that the intervention assessed was effective and safe. The term “safe” might give the impression that harms are not caused by an intervention or could imply that the trialists or the sponsoring drug company judged that the potential benefits of the intervention assessed outweighed the potential harms (at least under the trial conditions). However, this does not mean that the intervention carries no risk of harms in the trial conditions, let alone during general clinical use in the real world. Therefore, the potentially misleading safe/safety terminology should be avoided. Similarly, the term “side effect” denotes an effect without identifying it as a harmful one and implies that it is related to the mechanism of action of the intervention [23,24]. Importantly, patient representatives participating in all Delphi rounds confirmed they felt the term “side effect” downplays harms and should therefore not be used. Finally, the term “risk” is used colloquially to denote uncertainty, especially of an undesirable event. As a statistical term, risk is a proportion and should not be conflated with the nature of an event; that is, interventions are associated with the “risk” (or probability) of benefits and harms.

Table 1 summarizes some of the terminology problems concerning the reporting of harms in randomized controlled trials. For instance, events described as “anticipated” or “unintended” lack specific meaning. The use of ambiguous terminology is problematic and should be avoided.

1.3. How to use the CONSORT Harms update

Table 2 shows the CONSORT Harms checklist alongside the main CONSORT statement. Subsequently, we provide examples and explanations for the CONSORT Harms items and discuss special considerations for the reporting of harms in randomized controlled trials, including examples of good reporting when available. Several historical examples predate current recommendations and do not adhere entirely to the terminology recommended in this update.

CONSORT Harms 2022 had the goal of updating and incorporating items of the CONSORT Harms 2004 extension into the items of the main CONSORT statement. Overall, 13 of 25 items of the main CONSORT were minimally modified by adding the phrase “benefits and harms” to assure harms are reported alongside potential benefits. Considerable modifications were made to the remaining 12 items of the title and abstract, introduction, methods (topics outcomes, blinding and statistical methods), results (topics participant flow, recruitment, numbers analyzed, outcomes estimation, ancillary analyses), discussion (topic limitations), and other information (topic protocol). Three new items for CONSORT Harms 2022 were developed and incorporated into the reporting checklist (a side-by-side comparison of CONSORT Harms 2004 and 2022 can be found in [Appendix 1](#) in the [Supplementary Material](#)). An elaboration and explanation section specifically concerning harms was also developed. CONSORT Harms

urges authors that data on harms should be completely reported for all assessed and detected harms, and when appropriate to overcome space constraints in journal publications, supplementary information on harms should be made publicly available through online repositories.

2. CONSORT Harms 2022: checklist items, explanation, and elaboration

2.1. Title and abstract

2.1.1. Item 1b

CONSORT—Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts).

CONSORT Harms 2022: Item 1b—Structured summary of trial design, methods, results of outcomes of benefits and harms, and conclusions (for specific guidance see CONSORT for abstracts).

CONSORT Harms explanation—Abstracts are often all readers can openly access from a trial report. Because of barriers to accessing the scientific literature (e.g., paywalls, language), some readers might be able to access only information available in abstracts rather than the full text [29]. Lack of accuracy and completeness of reporting outcomes results in abstracts can result in misleading impression of results to readers [30]. For example, biases favouring the reporting of only benefits and the use of vague general statements (e.g., “safe” or “well tolerated”) are common. This is true for both abstracts of published papers and abstracts presented at conferences [31,32].

Recognizing that there might be challenges in accommodating journal guidance and character limits for abstracts, authors should strive to provide consistent and complete information on important benefits and harms. Finally, if no harms were assessed or detected, this should be stated.

Example of item 1b of CONSORT Harms 2022

“Sixteen (2%) of 958 women in the intravenous iron sucrose group and 13 (1%) of 976 women in the standard therapy group had serious maternal adverse events. Serious fetal and neonatal adverse events were reported by 39 (4%) of 961 women in the intravenous iron sucrose group and 45 (5%) of 982 women in the standard therapy group. At 6 weeks postrandomization, minor side effects were reported by 117 (16%) of 737 women in the intravenous iron sucrose group vs. 155 (21%) of 721 women in the standard therapy group” [33] [In this example, “side effects” is used as a reference to harm outcomes; preferable terms are “harms” or “adverse events”].

2.2. Introduction

2.2.1. Item 2b

CONSORT—Specific objectives or hypotheses.

CONSORT Harms 2022: Item 2b—Specific objectives or hypotheses for outcomes of benefits and harms.

CONSORT Harms explanation—Randomized controlled trials are often designed with a focus on potential benefits, including for sample size estimates. However, it is also relevant to the Introduction to describe what consideration has been given to harms outcomes when planning the trial.

Examples of item 2b of CONSORT Harms 2022

“The LIFE Study is the largest trial to evaluate the benefits of physical activity in older people. Serious fall injury was included in the LIFE Study as one of the prespecified secondary outcomes. By improving gait, balance, and lower extremity strength, physical activity may reduce the likelihood of falling and sustaining a serious injury, but it may also increase opportunities to fall and incur a serious injury. Here, we report results associated with our hypothesis that a long-term physical activity program compared with a health education program reduces the risk of serious fall injuries among sedentary older people with functional limitations.” [34].

“The aim of the present study is to compare the postoperative analgesic efficacy and incidence of nausea between the variable-rate feedback infusion mode and conventional fixed-rate infusion mode of ropivacaine/fentanyl-based PCEA in patients who underwent open gastrectomy.” [35] PCEA = patient-controlled epidural analgesia.

2.3. Methods

2.3.1. Item 6a

CONSORT—Completely defined prespecified primary and secondary outcome measures, including how and when they were assessed.

CONSORT Harms 2022: Item 6a—Completely defined prespecified primary and secondary outcomes, for both benefits and harms, including how and when they were assessed.

2.3.2. Item 6b

CONSORT—Any changes to trial outcomes after the trial commenced, with reasons.

No modifications in Consort Harms 2022.

2.3.3. Item 6c (new item)

CONSORT Harms 2022: Item 6c—Describe if and how nonprespecified outcomes of benefits and harms were identified, including any selection criteria, if applicable.

CONSORT Harms explanation (items 6a and 6c)—Previous recommendations have suggested that randomized controlled trial reports should focus on the most clinically important harms [28]. However, selecting harms for analysis and reporting based on clinical importance can be problematic because it relies on the author’s judgment and allows for the application of selection criteria that might be decided post hoc. There might not be consensus on the most clinically important harms [36], and documentation for any independent evaluation of harms that has informed reporting—for example, the charters for independent data monitoring committee and outcome adjudication committee—should be provided.

For prespecified and systematically assessed harms, authors should identify and define the outcomes as per their domains (case definitions), measurements, metrics, method of aggregation, and time frames [21,25]. For nonsystematically assessed harms (prespecified or nonprespecified), the report should elaborate on the specific methods used to collect nonsystematically assessed harms (e.g., by stating verbatim if indirect questioning was used or including a copy of the blank case report form) and the time points at which participants were asked about harms or had a possibility of reporting harms if no questions were asked. The information provided should be sufficient to allow others to use the same outcomes [37]; and [Supplementary Material](#) could be used to describe the information with the necessary details. For nonsystematically assessed harms (prespecified or nonprespecified), the report should elaborate on the specific methods used to collect nonsystematically assessed harms (e.g., by stating verbatim if indirect questioning was used or including a copy of the blank case report form) and the time points at which participants were asked about harms or had a possibility of reporting harms if no questions were asked. The information provided should be sufficient to allow others to use the same outcomes [37]; and [Supplementary Material](#) could be used to describe the information with the necessary details.

Outcome measurements refer to whether trialists used standardized instruments to assess harms, if and how new scales were developed, and the use of nonvalidated scales. Initiatives like the Consensus-based Standards for the selection of health Measurement Instruments provide specific guidance on how to select outcome measurement instruments; additional guidance on outcomes reporting can be found in CONSORT outcomes [34,38]. Validated scales or consensus guidelines might [39,40] enhance quality of measurement and assist in comparison with similar studies [41]. Metrics refer to the measure used to characterize the results; for example, proportion of patients experiencing an event. For harms reporting, it is important to clarify whether the metric refers to the proportion of patients presenting with at least one harm event or the proportion of patients presenting with specific harm events. This reporting is particularly relevant when considering events that can recur (e.g., headaches). When harms outcomes are

aggregated, the methods must be fully and completely described. These methods could include timing, coding or body system subgrouping, frequency, seriousness, severity, and chronicity, as appropriate. If harms are defined as attributed or not to the intervention, the attribution methods should be described along with information on who completed such assessments (investigators, data safety monitoring boards, patients, sponsors, or any combination thereof). The lack of reporting of such details has been shown to seriously hamper comparability of harms outcomes [42].

Finally, the specific time frame for the assessment of harms should be specified. The time point for assessing harms events can be crucial, as illustrated by a systematic review evaluating the risk of heparin-induced thrombocytopenia in postoperative patients [43]. Heparin-induced thrombocytopenia is a prothrombotic disorder related to antibody-mediated platelet activation induced by heparin that occurs within 5–14 days of the start of heparin treatment. In one trial considered for inclusion in the review, heparin-induced thrombocytopenia was assessed on postoperative days 1–5 [44]. The trial was not included in the systematic review because the time point was inappropriate to allow for the reaction to develop.

Criteria to include harms in a trial report are often post hoc decisions based on arbitrary thresholds, such as “harms occurring in at least 5% of the randomized controlled trial participants” or “grade 3 or above events.” We strongly recommend that thresholds are not applied [4]. Instead, authors should provide a summary of the data on harms in the main report and report all harms in detail in [Supplementary Materials](#). If any criteria or rule-based approaches are used to select which harms were identified in a report, they should be prespecified and stated explicitly.

Example of item 6a CONSORT Harms 2022

“As secondary outcomes, we analyzed intraoperative complications, drainage time, length of hospital stay, postoperative pain, postoperative quality of life, and readmissions within 90 days. Drainage time was defined as the interval between surgery and the removal of the chest tube and was measured in days. Length of hospital stay was measured in days after surgery. Postoperative pain was evaluated by a visual analog pain scale on the first, second, and third postoperative days and at the 30-day outpatient visit. We also assessed the need for opioid use at the 30-day outpatient visit. Any hospitalization within the 90-day postoperative period was considered as readmission” [45] [In this example, “complications” is used as a reference to harm outcomes; preferable terms are “harms” or “adverse events”].

When authors choose a subset of harms to report, this can be highly misleading and dramatically impact the public’s ability to identify and synthesize the harms of intervention. For interventions that target healthy individuals (e.g., screening), any harm, however minor, could be important to capture and report because the balance between harms and benefits could easily lean towards harms in a low-risk population. For interventions that improve major outcomes (e.g., survival in patients with cancer), minor harms might not be as important to patients compared with potential benefits. However, harms deemed clinically mild can seriously affect a patient’s quality of life [8] and treatment adherence [9,10] and potentially increase financial costs to patients and health systems [11]. Finally, results should be reported for all prespecified outcomes, not just for the most frequent events or analyses that were statistically significant or interesting.

2.3.4. Item 11a

CONSORT—If done, who was blinded after assignment to interventions (e.g., participants, care providers, those assessing outcomes) and how.

CONSORT Harms 2022: Item 11a—If done, who was blinded after assignment to interventions (e.g., participants, care providers, those assessing outcomes of benefits and harms) and how.

CONSORT Harms explanation—Blinding, or masking, is the process of withholding information about the intervention assignment from people involved in the conduct of a trial after participants are assigned to the intervention. Personnel and participants aware of intervention assignment might be influenced in their delivery of interventions as well as beliefs of intervention effects; therefore, blinding provides protection against bias due to deviations from intended intervention and in outcomes ascertainment [46,47]. People involved in a trial who could be blinded include study participants and personnel such as people administering the intervention, laboratory technicians, data collectors, people assessing benefits and harms, and statisticians. Of note, despite the widespread use of terminologies such as single, double, or triple blind, there is no standard, widespread accepted meaning for these terms; therefore, authors should avoid them [48]. Additionally, “masking” rather than “blinding” is the preferred terminology in ophthalmology trials.

For some trials, it is not possible to keep either the participant or the person administering the intervention unaware of the assignment. In these cases, it might still be possible for people assessing benefits and harms (other than those assessed by the participants themselves) to remain blinded. Authors should report whether or not blinding of harms assessment occurred. In some cases, this assessment involves an external committee, such as an independent safety monitoring board; members of that committee make decisions about attribution of harms and their possible relationship with the intervention under study.

2.3.5. Item 12a

CONSORT—Statistical methods used to compare groups for primary and secondary outcomes.

CONSORT Harms 2022: Item 12a—Statistical methods used to compare groups for primary and secondary outcomes of both benefits and harms.

CONSORT Harms explanation—Statistical methods should be described for each outcome investigated in a randomized controlled trial, including both primary and secondary benefits and harms. Additionally, it is important that a report states whether the methods were prespecified. If a statistical analysis plan is available, it should be dated and have revisions documented. If there are deviations from the statistical analysis plan, they should be documented. [Supplementary Material](#) can be used to provide this information.

For nonsystematically assessed harms, it is a common practice in trials to code and group events into broader categories before conducting statistical analyses. The report should describe how individual events were grouped, including details for any coding dictionary (e.g., MedDRA). Grouping harms could help identify signals for biologically related harms or syndromes, but it might also allow trial investigators to obscure important harms by combining them with less important ones (e.g., migraine might be more severe than headache, and bleeding might be more severe than bruising). Authors should clarify the level of grouping (e.g., high level terms) and whether such groupings were prespecified in a protocol or statistical analysis plan or defined post hoc [49]. Obscuring harms by coding can be problematic [50,51].

When relevant, authors should plan and explain how they accounted for time and duration of events, duration of follow-up, and how recurrent events were managed in the trial analysis. Reporting such information allows a more accurate and comprehensive knowledge of harms and provides a valuable insight into participants' quality of life. Of note, harms occurring repeatedly and over an extended duration might have a

Example of item 12a of CONSORT Harms 2022

“Safety analyses: The primary safety variables were time-to-first treatment emergent adverse event (AE), serious AE, and AE leading to discontinuation from study treatment. The primary variable was analyzed using a Cox regression model stratified by randomization stratum, with treatment group, severity of asthma, and region as fixed factors” [52] [In this example, “safety” is used as a reference to harm outcomes. We recommend against the use of the terminology “safety” variables. Our guidance is for complete reporting of “harms” as the possible adverse consequences of an intervention and the direct opposite of benefits].

detrimental impact on a patient's quality of life, even if the harms are not considered serious or severe.

2.4. Results

2.4.1. Item 13a

CONSORT—For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analyzed for the primary outcome.

Example of item 13a of CONSORT Harms 2022

In this randomized controlled trial of the effect of an intervention (Vosoritide) to treat achondroplasia as compared with placebo, the population analyzed for harms included all patients who received at least one dose of the intervention (Vosoritide) or placebo (the so-called safety analyses) [53]. The population analyzed for potential benefits included all randomized participants, constituting the so-called full analysis set as per intention-to-treat principles.

The flow diagram (Fig. 1) details the numbers of participants who were randomly assigned and received intended treatment as defined for each population analyzed.

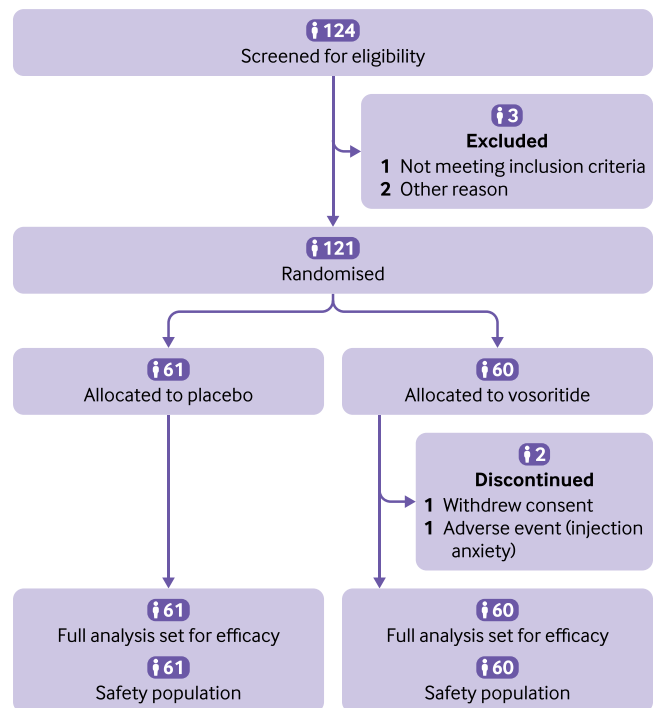


Fig. 1. Flow diagram, relating to example of 13a of CONSORT Harms 2022 [In this example, “safety” is used as a reference to harm outcomes. We recommend against the use of “safety analysis” or “safety population” terms. Our guidance is for complete reporting of the randomized population and number analyzed for outcomes of benefits and harms regardless of whether harms were the primary or secondary outcome in the trial].

CONSORT Harms 2022: Item 13a—For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analyzed for outcomes of benefits and harms.

CONSORT Harms explanation—The flow of participants in the trial and the population analyzed for the outcomes measured might not be identical for benefits and harms. Censoring or attrition might lead to different analyzable populations, which should be described clearly. For harms, additional information on dose reductions of the allocated intervention might also be relevant to report, with emphasis on reasons, including if related to participants experiencing harms. It should be acknowledged that attrition is often due to a combination of harms and (perceived) lack of benefits, and it might be difficult to disentangle the relative contribution of these different reasons. Also, trials with long-term follow-up should differentiate and describe attrition issues related to harms at early and late time points to improve accuracy of the information related to harms. Information on time to discontinuation due to harms might also be useful in some circumstances.

2.4.2. Item 14a

CONSORT—Dates defining the periods of recruitment and follow-up.

CONSORT Harms 2022: Item 14a—Dates defining the periods of recruitment and follow-up for outcomes of benefits and harms.

CONSORT Harms explanation—Information on the date of the randomization and the completion of the study for the primary efficacy outcome does not suffice for the assessment of harms. For instance, the assessment of harms might be planned to take place during the entire study through nonsystematic assessment, might occur during only part of the study duration, might occur at specific time points using systematic or nonsystematic assessment, or might continue after the completion of follow-up for the main efficacy outcome. Reporting the periods of recruitment and follow-up for benefits and

Example of item 14a of CONSORT Harms 2022

“The trial involved five visits: Visit 1 on day 1 (screening, randomization, and initial dosing), Visit 2 on day 2 (assessment of the primary end point), Visit 3 on day 4 (assessment of efficacy and safety parameters), Visit 4 on day 6 (end-of-treatment visit), and Visit 5 on day 8 to day 10 (follow-up by telephone interview). Patients were asked to return all unused trial medication and their diaries at each visit” [54] [In this example, “safety” is used as a reference to harm outcomes. We recommend against the use of “safety”; preferable terms are “harms” or “adverse events”].

harms is crucial to allow comprehensive and accurate interpretation of the trials results.

2.4.3. CONSORT Harms 2022 special consideration: run-in periods

A metaepidemiological study [55] identified randomized trials with run-in periods and evaluated the characteristics of these run-in periods, and the completeness of reporting with regard to exclusions (number of and reasons for exclusions and baseline characteristics of those excluded). Of 470 PubMed-indexed randomized controlled trial publications from 2014, 25 (5%) included some type of run-in period. In 23 of 25 trials (88%), the run-in period was incompletely reported, mostly because of missing baseline characteristics. The authors recommended trial publications should state the number of excluded patients, reasons for exclusion, and baseline characteristics of the excluded patients.

Run-in periods have been used in some randomized controlled trials to exclude patients before randomization. A frequent approach is to give patients the trial drug and to exclude those who experience harms [56]. Trials estimating the effects of initiating unknown interventions (vs. initiating placebo) might obtain different results compared with trials estimating the effect of continuing well-tolerated interventions. Therefore, when run-in periods are used, adequate reporting of exclusions due to harms during or after the run-in period is essential to avoid underestimating the true incidence of harms [57]. Harms should be reported for all the study periods, including the run-in period.

2.4.4. Item 16

CONSORT—For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups.

CONSORT Harms 2022: Item 16—For each group, number of participants (denominator) included in each analysis of outcomes of benefits and harms and whether

Example of item 16 of CONSORT Harms 2022

“Results, patient disposition, baseline demographics, and study drug exposure. The integrated safety database included a total of 4,439 patients who received at least one dose of study drug (placebo, $n = 1,262$; lasmiditan, $n = 3,177$). Of those who received a study dose, 97% assigned either placebo or to a lasmiditan dose group completed the study (i.e., all required follow-ups)” [60] [In this example, we recommend against the use of the terminology “safety” database. Our guidance is for complete reporting of “harms” as the possible adverse consequences of an intervention and the direct opposite of benefits].

the analysis was by original assigned groups and if any exclusions were made.

CONSORT Harms explanation—Information about which participants were included in which analyses is essential. It is a common practice in randomized controlled trial reports to describe the population analyzed using a variety of terms, such as intention to treat, modified intention to treat, and per protocol analysis. In most trials, each of these will include a different set of participants and corresponding denominator; however, it is often not clear from trial reports which study population was included in the analyses of different outcomes. The most common terminology, the intention-to-treat analysis, by principle should analyze all participants in the group randomized regardless of noncompliance, protocol deviations, withdrawal, and any other reason for exclusion after randomization [58]. Nevertheless, it is not always possible to measure outcome data

on all participants [59]. “Safety population” is an additional term often used in reports of randomized controlled trials to define an analysis approach that includes patients who received at least one dose of the drug or intervention. However, heterogeneity in the use of the different terms and missing data might confuse readers trying to identify the population at risk that was analyzed.

2.4.5. Item 17a

CONSORT—For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval).

CONSORT Harms 2022: Item 17a—For each primary and secondary outcomes of benefits and harms, results for each group, and the estimated effect size and its precision (such as 95% confidence intervals).

Example of items 17a and 17b of CONSORT Harms 2022

Harms experienced by patients treated with albiglutide were compared with harms assessed in patients receiving placebo in a randomized controlled trial conducted by Hernandez et al. [63]. The trial publication reports the number and proportion of patients who experienced systematically assessed harms among patients who received at least one dose of albiglutide or placebo, along with the relative risk of harms with 95% confidence interval.

Table 3 presents the absolute risks and relative risks reported by Hernandez et al. and the estimates of risk difference, which are considered a preferable statistical measure when dealing with rare events. This is because relative risk estimates for rare events can be misleading in representing effects sizes; Table 3 shows how the relative risk of severe hypoglycemia of 0.56 relates to a risk difference of 0.005. The use of relative risks is also problematic when dealing with zero events in one or two arms.

Table 3. Prespecified systematic harms

| Harms | Albiglutide (n = 4,717) | Placebo (n = 4,715) | RR (95% CI) | RD (95% CI) |
|--|-------------------------|---------------------|---------------------|---------------------------|
| Severe hypoglycaemia | 31 (0.7) | 55 (1.2) | 0.56 (0.36 to 0.87) | −0.005 (−0.009 to −0.001) |
| Pancreatitis | 10 (0.2) | 7 (0.1) | 1.43 (0.54 to 3.75) | 0.001 (−0.001 to 0.002) |
| Injection site reactions | 86 (1.8) | 29 (0.6) | 2.96 (1.95 to 4.51) | 0.012 (0.007 to 0.016) |
| Thyroid cancer | 0 (0) | 0 (0) | - | 0 (−0.001 to 0.001) |
| Hematological neoplasia | 9 (0.2) | 5 (0.1) | 1.80 (0.60 to 5.36) | 0.001 (−0.001 to 0.002) |
| Pancreatic cancer | 6 (0.1) | 5 (0.1) | 1.20 (0.37 to 3.93) | 0 (−0.001 to 0.001) |
| Hypersensitivity syndrome or symptoms | 45 (1) | 48 (1) | 0.94 (0.63 to 1.40) | −0.001 (−0.005 to 0.003) |
| Hepatobiliary disorders | 51 (1.1) | 41 (0.9) | 1.24 (0.83 to 1.87) | 0.002 (−0.002 to 0.006) |
| Alanine aminotransferase of at least three times the ULN | 17 (0.4) | 30 (0.6) | 0.57 (0.31 to 1.03) | −0.003 (−0.006 to 0.000) |
| Alanine aminotransferase of at least five times the ULN | 6 (0.1) | 17 (0.4) | 0.35 (0.14 to 0.89) | −0.002 (−0.005 to 0.000) |
| Bilirubin of at least twice the ULN | 12 (0.3) | 7 (0.1) | 1.71 (0.68 to 4.35) | 0.001 (−0.001 to 0.003) |
| Serious gastrointestinal events | 92 (2) | 87 (1.8) | 1.06 (0.79 to 1.41) | 0.001 (−0.005 to 0.006) |
| Appendicitis | 3 (0.1) | 8 (0.2) | 0.37 (0.10 to 1.41) | −0.001 (−0.003 to 0.000) |
| Atrial fibrillation or flutter | 108 (2.3) | 131 (2.8) | 0.82 (0.64 to 1.06) | −0.005 (−0.011 to 0.001) |
| Pneumonia | 121 (2.6) | 138 (2.9) | 0.95 (0.75 to 1.20) | −0.004 (−0.010 to 0.003) |
| Renal impairment | 279 (5.9) | 319 (6.8) | 0.87 (0.75 to 1.02) | −0.009 (−0.019 to 0.001) |
| Diabetic retinopathy | 78 (1.7) | 89 (1.9) | 0.88 (0.65 to 1.18) | −0.002 (−0.008 to 0.003) |

Abbreviations: CI, confidence interval; RD, risk difference; RR, relative risk; UNL, upper limit of normal. Data are number of patients (%) in those who took at least one dose.

2.4.6. Item 17a2 (new item)

CONSORT Harms 2022: Item 17a2—For outcomes omitted from the trial report (benefits and harms), provide rationale for not reporting and indicate where the data on omitted outcomes can be accessed.

2.4.7. Item 17b

CONSORT—For binary outcomes, presentation of both absolute and relative effect sizes is recommended.

CONSORT Harms 2022: Item 17b—Presentation of both absolute and relative effect sizes is recommended for outcomes of benefits and harms.

2.4.8. Item 17c (new item)

CONSORT Harms 2022: Item 17c—Report zero events if no harms were observed.

CONSORT Harms explanation (items 17a, 17a2, and 17c)—Harms are often dichotomized into binary or count data (see explanation for items 6a, 6c, and 12a), in which case it could be appropriate to follow the main CONSORT guidance to present relative (risk ratio [relative risk] or odds ratio) and absolute effect (risk difference) metrics with confidence intervals [12]. However, for nonsystematically assessed harms, measures of relative or absolute risk difference should be used with caution because these outcomes were not actively determined. Additionally, harms should be reported even when they are not directly comparable among intervention groups, thus precluding computing measures of relative or absolute effect. Therefore, for harms, we recommend that the reporting of items 17a and 17b be considered as appropriate and not solely based on whether the outcome was analyzed as binary data.

When presenting the results of harms, whether using binary or continuous data, it is important to separate the reporting into systematically and nonsystematically assessed harms. Additionally, authors should always report events of serious harms. If attribution methods were applied, it is informative and adds to transparency that all

harms are reported, even if events are deemed not related to the intervention (see explanation for items 6a and 6c). We also recommend that authors consider available visualization methods for the communication of adverse events (illustrating data through figures rather than tables), which can offer a useful alternative to help summarize harm profiles [61,62].

Reporting incidence (e.g., proportion of people experiencing an event) can be misleading for recurring events. Therefore, it should be clear whether authors report number of participants presenting with at least one event (incidence) or number of events per unit of time at risk (rate), and whether recurrent events were included.

Concerning the relevance of the time frame to specify and define harms (see item 6), authors should report exposure time and any differences in follow-up or exposure and how they accounted for these differences in the analysis. Studies with differential follow-up between intervention arms (e.g., because of participant withdrawals) only provide the opportunity to collect information on harms for participants who remain in the study. In this scenario, using person time for exposure can be misleading if participants who continued an intervention are a subgroup who experienced few harms.

Confidence intervals are useful for reporting systematically assessed harms and sometimes for nonsystematically assessed harms (*P* values for results of nonsystematically assessed harms are of little relevance because hypothesis testing would be of limited value). Confidence intervals could indicate that the result does not rule out an important clinical difference in one or both directions. Although *P* values might be provided in addition to confidence intervals, results should not be reported solely as *P* values.

Finally, for prespecified and systematically assessed harms, authors should report results even if zero events were observed to promote transparency and to facilitate synthesis across trials.

Example of item 17c of CONSORT Harms 2022

Table 4. Primary and secondary cardiovascular outcomes and safety outcomes in the overall population [64]

| Variable | Dapagliflozin (n = 3,131) | | Placebo (n = 3,132) | |
|--|---------------------------|-------------------------|---------------------|-------------------------|
| | Values | Events/100 patient year | Values | Events/100 patient year |
| Safety outcomes—No./total No. (%) | | | | |
| Any definite or probable diabetic ketoacidosis | 2/3,126 (0.1) | — | 0 | — |
| Fournier's gangrene | 0 | — | 0 | — |

Solomon SD et al. Dapagliflozin in heart failure with mildly reduced or preserved ejection fraction. *New Engl J Med* 2022;387:1089–98 [In this example, we recommend against the use of “safety”; preferable terms are “harms” or “adverse events”].

2.4.9. Item 18

CONSORT—Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing prespecified from exploratory.

CONSORT Harms 2022: Item 18—Results of any other analyses performed for outcomes of benefits and harms, including subgroup analyses and adjusted analyses, distinguishing prespecified from exploratory.

CONSORT Harms Explanation—When subgroup analyses are done, authors should report which subgroups were examined, the rationale for choosing these subgroups, and whether the direction and magnitude of differences between subgroups were prespecified. Selective reporting of subgroup analyses is an important source of bias. Subgroup analysis investigates if the intervention effects vary among the levels of some factor of interest, but very few trials are powered to detect subgroup differences, especially for harms outcomes.

If multiple analyses are done, authors should indicate which analysis (e.g., unadjusted or adjusted) is the primary one, and if it was prespecified in the statistical analysis plan or protocol. For example, whether variables adjusted for were prespecified in the protocol and known a priori to be strongly or moderately associated with the outcome should also be stated.

2.5. Discussion

2.5.1. Item 20

CONSORT—Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses.

CONSORT Harms 2022: Item 20—Trial limitations, addressing sources of potential bias related to the approach to collecting or reporting data on harms, imprecision, and, if relevant, multiplicity or selection of analyses.

CONSORT Harms explanation—When discussing trial limitations, authors should indicate whether the data are similarly valid and reliable for both benefits and harms. If data on harms were not systematically assessed, authors should draw particular attention to limitations in the assessment of harms, including the chance of both false positive and false negative results (lack or poor monitoring of harms does not provide evidence that no harms occurred) [65].

Example of item 20 of CONSORT Harms 2022

“Finally, although we found no increase in the risk of congenital abnormalities among babies of women treated with progesterone, the trial was not powered for such rare outcomes” [71].

In addition to risk of bias and imprecision that apply to all outcomes, trials often have specific limitations related to assessing, reporting, and analyzing harms. Trials with adequate power to detect potential benefits might be underpowered to detect important differences in harms, including prespecified harms. Trials of short duration might not be able to detect harms that develop after prolonged treatment, and trials reporting only the proportion of participants who experienced one or more harms might conceal information about event rates that is important for decision-making [5]. Finally, threats to internal validity might affect harms and benefits differently. For instance, data are commonly missing for participants who discontinue interventions because of harms.

Although harms are more likely to be under-reported than over-reported, estimates of the occurrence of harms might also be influenced by a nocebo effect, where the communication about potential harms inflates the reporting of these events [66]. From this perspective, the informed consent process can contribute to the detection of specific harms if they are communicated with negative framing or in a way that promotes the reporting of placebo-induced harms [67–70]. Harms can also be overlooked or disregarded by investigators who are not convinced of attribution; all harms should be reported rather than only those felt by investigators to be causally attributed to an intervention.

2.5.2. Item 24

CONSORT—Where the full trial protocol can be accessed, if available.

CONSORT Harms 2022: Item 24—Where the full trial protocol and other relevant documents can be accessed, including additional data on harms.

CONSORT Harms explanation—A large portion of data about harms are not published [72–75]. The inadequate reporting of harms is prevalent among pharmaceutical and nonpharmaceutical sponsored trials, a troublesome and widespread practice [76,77]. It might not be practical to report all data on harms in journal publications, but these data should be made freely available elsewhere for

Example of item 24 of CONSORT Harms 2022

“Data sharing: The statistical analysis plan is available at <https://doi.org/10.6084/m9.figshare.5375026.v1>. The raw trial data are provided by the authors on a secure online repository (see [Supplementary Appendix](#) for link). Data include anonymized individual patient variables for results reported here, a read-me file with data dictionary and analyzes scripts used in this study” [79].

systematically and nonsystematically assessed outcomes, ideally including anonymized individual participant data.

Trial authors should report where data can be found. The FAIR data principles (findability, accessibility, interoperability, and reuse) are relevant to harms data and should be considered as open science practices and evolving legislation become more accepted [78].

3. Discussion

Investigators continue to under-report harms [4,17,75,76,80–84]. The CONSORT extension for harms was published in 2004 to highlight the inadequate reporting of harms in trials and to promote guidance on the minimum set of information on harms trials should provide [13]. A guideline is only helpful if its guidance is followed. Although the CONSORT statement has been endorsed by more than 500 medical journals, CONSORT extensions are endorsed by a small fraction of these. We evaluated the adherence to CONSORT Harms items in published clinical trials and found that half of the items failed to reach more than 50% compliance after 2004 [17].

Because interventions have the potential to cause beneficial and harmful effects, we proposed to the CONSORT group to incorporate harms items into the main CONSORT statement. The CONSORT group requested that Delphi participants in this study be offered the opportunity to comment. Most Delphi participants supported the incorporation of CONSORT Harms extensions into the main CONSORT checklist and the CONSORT Executive agreed. The main CONSORT statement will likely be revised, at which stage the removal of item 19 might be recommended considering the integration of CONSORT and CONSORT Harms. Until future work from the CONSORT group produces an updated checklist, trial authors, journal reviewers, and editors should use the integrated checklist presented in this paper (Table 2).

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Supplementary Data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jclinepi.2023.04.005>.

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