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# Reporting guidelines for clinical trial reports for interventions involving artificial intelligence

SPIRIT-AI and CONSORT-AI Working Group

DOI:

10.1016/S2589-7500(20)30218-1

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Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

SPIRIT-AI and CONSORT-AI Working Group 2020, 'Reporting guidelines for clinical trial reports for interventions involving artificial intelligence: the CONSORT-AI extension', *The Lancet Digital Health*, vol. 2, no. 10, pp. E537-E548. https://doi.org/10.1016/S2589-7500(20)30218-1

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# Reporting guidelines for clinical trial reports for interventions involving artificial intelligence: the CONSORT-AI extension



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The CONSORT 2010 statement provides minimum guidelines for reporting randomised trials. Its widespread use has been instrumental in ensuring transparency in the evaluation of new interventions. More recently, there has been a growing recognition that interventions involving artificial intelligence (AI) need to undergo rigorous, prospective evaluation to demonstrate impact on health outcomes. The CONSORT-AI (Consolidated Standards of Reporting Trials-Artificial Intelligence) extension is a new reporting guideline for clinical trials evaluating interventions with an AI component, It was developed in parallel with its companion statement for clinical trial protocols; SPIRIT-AI (Standard Protocol Items: Recommendations for Interventional Trials-Artificial Intelligence). Both guidelines were developed through a staged consensus process involving literature review and expert consultation to generate 29 candidate items, which were assessed by an international multi-stakeholder group in a two-stage Delphi survey (103 stakeholders), agreed upon in a two-day consensus meeting (31 stakeholders), and refined through a checklist pilot (34 participants). The CONSORT-AI extension includes 14 new items that were considered sufficiently important for AI interventions that they should be routinely reported in addition to the core CONSORT 2010 items. CONSORT-AI recommends that investigators provide clear descriptions of the AI intervention, including instructions and skills required for use, the setting in which the AI intervention is integrated, the handling of inputs and outputs of the AI intervention, the human-AI interaction and provision of an analysis of error cases. CONSORT-AI will help promote transparency and completeness in reporting clinical trials for AI interventions. It will assist editors and peer reviewers, as well as the general readership, to understand, interpret, and critically appraise the quality of clinical trial design and risk of bias in the reported outcomes.

#### Introduction

Randomised controlled trials (RCTs) are considered the gold-standard experimental design for providing evidence of the safety and efficacy of an intervention. <sup>1,2</sup> Trial results, if adequately reported, have the potential to inform regulatory decisions, clinical guidelines, and health policy. It is therefore crucial that RCTs are reported with transparency and completeness so that readers can critically appraise the trial methods and findings and assess the presence of bias in the results. <sup>3-5</sup>

The CONSORT statement provides evidence-based recommendations to improve the completeness of the reporting of RCTs. The statement was first introduced in 1996 and has since been widely endorsed by medical journals internationally. Over the past two decades, it has undergone two updates and has demonstrated a substantial positive impact on the quality of RCT reports. The most recent CONSORT 2010 statement provides a 25-item checklist of the minimum reporting content applicable to all RCTs, but it recognises that certain interventions may require extension or elaboration of these items. Several such extensions exist. 8-13

Artificial intelligence (AI) is an area of enormous interest with strong drivers to accelerate new interventions through to publication, implementation, and market.<sup>14</sup> While AI systems have been researched for some time, recent advances in deep learning and neural networks have gained considerable interest for their potential in health applications. Examples of such applications are wide ranging and include AI systems for screening and triage,<sup>15,16</sup> diagnosis,<sup>17–20</sup> prognostication,<sup>21,22</sup> decision support,<sup>23</sup> and treatment recommendation.<sup>24</sup> However, in

the most recent cases, published evidence has consisted of in-silico, early-phase validation. It has been recognised that most recent AI studies are inadequately reported, and existing reporting guidelines do not fully cover potential sources of bias specific to AI systems.<sup>25</sup> The welcome emergence of RCTs seeking to evaluate newer interventions based on, or including, an AI component (called "AI interventions" here)<sup>23,26-31</sup> has similarly been met with concerns about the design and reporting.<sup>25,32-34</sup> This has highlighted the need to provide reporting guidance that is fit for purpose in this domain.

CONSORT-AI (as part of the SPIRIT-AI and CONSORT-AI initiative) is an international initiative supported by CONSORT and the EQUATOR (Enhancing the Quality and Transparency of Health Research) Network to evaluate the existing CONSORT 2010 statement and to extend or elaborate this guidance where necessary, to support the reporting of clinical trials for AI interventions. <sup>15,36</sup> It is complementary to the SPIRIT-AI statement, which aims to promote high-quality protocol reporting for AI trials. This Consensus Statement describes the methods used to identify and evaluate candidate items and gain consensus. In addition, it also provides the CONSORT-AI checklist, which includes the new extension items and their accompanying explanations.

#### Methods

The SPIRIT-AI and CONSORT-AI extensions were simultaneously developed for clinical trial protocols and trial reports. An announcement for the SPIRIT-AI and CONSORT-AI initiative was published in October 2019,<sup>35</sup> and the two guidelines were registered as reporting

Lancet Digital Health 2020; 2: e537–548

Published Online September 9, 2020 https://doi.org/10.1016/ \$2589-7500(20)30218-1

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#### Panel: Glossary

#### **Artificial Intelligence**

The science of developing computer systems which can perform tasks normally requiring human intelligence.

#### Al intervention

A health intervention that relies upon an AI/ML component to serve its purpose.

#### CONSORT

Consolidated Standards of Reporting Trials.

#### CONSORT-AI extension item

An additional checklist item to address AI-specific content that is not adequately covered by CONSORT 2010.

#### Class-activation map

Class-activation maps are particularly relevant to image classification AI interventions. Class-activation maps are visualisations of the pixels that had the greatest influence on predicted class, by displaying the gradient of the predicted outcome from the model with respect to the input. They are also referred to as "saliency maps" or "heat maps".

#### Health outcome

Measured variables in the trial that are used to assess the effects of an intervention.

#### **Human-Al interaction**

The process of how users (humans) interact with the AI intervention, for the AI intervention to function as intended.

#### Clinical outcome

Measured variables in the trial that are used to assess the effects of an intervention.

#### Delphi study

A research method that derives the collective opinions of a group through a staged consultation of surveys, questionnaires, or interviews, with an aim to reach consensus at the end.

#### **Development environment**

The clinical and operational settings from which the data used for training the model are generated. This includes all aspects of the physical setting (such as geographical location, physical environment), operational setting (such as integration with an electronic record system, installation on a physical device), and clinical setting (such as primary, secondary and/or tertiary care, patient disease spectrum).

#### Fine-tuning

Modifications or additional training performed on the AI intervention model, done with the intention of improving its performance.

#### Input data

The data that need to be presented to the AI intervention to allow it to serve its purpose.

#### Machine learning

A field of computer science concerned with the development of models/algorithms that can solve specific tasks by learning patterns from data, rather than by following explicit rules. It is seen as an approach within the field of AI.

#### Operational environment

The environment in which the AI intervention will be deployed, including the infrastructure required to enable the AI intervention to function.

#### Output data

The predicted outcome given by the AI intervention based on modeling of the input data. The output data can be presented in different forms, including a classification (including diagnosis, disease severity or stage, or recommendation such as referability), a probability, a class activation map, etc. The output data typically provide additional clinical information and/or trigger a clinical decision.

#### Performance error

Instances in which the AI intervention fails to perform as expected. This term can describe different types of failures, and it is up to the investigator to specify what should be considered a performance error, preferably based on prior evidence. This can range from small decreases in accuracy (compared to expected accuracy) to erroneous predictions or the inability to produce an output, in certain cases.

#### SPIRIT

Standard Protocol Items: Recommendations for Interventional Trials.

#### SPIRIT-AI

An additional checklist item to address AI-specific content that is not adequately covered by SPIRIT 2013.

#### SPIRIT-AI elaboration item

Additional considerations to an existing SPIRIT 2013 item when applied to AI interventions.

Al=artificial intelligence. ML=machine learning.

guidelines under development on the EQUATOR library of reporting guidelines in May, 2019. Both guidelines were developed in accordance with the EQUATOR Network's methodological framework.<sup>37</sup> The SPIRIT-AI

and CONSORT-AI Steering Group, consisting of 15 international experts, was formed to oversee the conduct and methodology of the study. Definitions of key terms are provided in the glossary (panel).

#### Ethical approval

This study was approved by the ethical review committee at the University of Birmingham, UK (ERN\_19-1100). Participant information was provided to Delphi participants electronically before survey completion and before the consensus meeting. Delphi participants provided electronic informed consent, and written consent was obtained from consensus meeting participants.

#### Literature review and candidate-item generation

An initial list of candidate items for the SPIRIT-AI and CONSORT-AI checklists was generated through review of the published literature and consultation with the Steering Group and known international experts. A search was performed on May 13, 2019, using the terms "artificial intelligence", "machine learning", and "deep learning" to identify existing clinical trials for AI interventions listed within the US National Library of Medicine's clinical trial registry (ClinicalTrials.gov). There were 316 registered trials, of which 62 were completed and seven had published results.30,38-43 Two studies were reported with reference to the CONSORT statement, 30,42 and one study provided an unpublished trial protocol.42 The Operations Team (XL, SCR, MJC, and AKD) identified AI-specific considerations from these studies and reframed them as candidate reporting items. The candidate items were also informed by findings from a previous systematic review that evaluated the diagnostic accuracy of deep-learning systems for medical imaging.<sup>25</sup> After consultation with the Steering Group and additional international experts (n=19), 29 candidate items were generated, 26 of which were relevant for both SPIRIT-AI and CONSORT-AI and three of which were relevant only for CONSORT-AI. The Operations Team mapped these items to the corresponding SPIRIT and CONSORT items, revising the wording and providing explanatory text as required to contextualise the items. These items were included in subsequent Delphi surveys.

#### Delphi consensus process

In September, 2019, 169 key international experts were invited to participate in the online Delphi survey to vote upon the candidate items and suggest additional items. Experts were identified and contacted via the Steering Group and were allowed one round of "snowball" recruitment in which contacted experts could suggest additional experts. In addition, individuals who made contact following publication of the announcement were included.<sup>35</sup> The Steering Group agreed that individuals with expertise in clinical trials and AI and machine learning (ML), as well as key users of the technology, should be well represented in the consultation. Stakeholders included health-care professionals, methodologists, statisticians, computer scientists, industry representatives, journal editors, policy makers, health "informaticists", experts in law and ethics, regulators, patients, and funders. Participant characteristics are described in the appendix (p 1). Two online Delphi surveys were conducted. DelphiManager software (version 4.0), developed and maintained by the COMET (Core Outcome Measures in Effectiveness Trials) initiative, was used to undertake the e-Delphi survey. Participants were given written information about the study and were asked to provide their level of expertise within the fields of (i) AI/ML and (ii) clinical trials. Each item was presented for consideration (26 for SPIRIT-AI and 29 for CONSORT-AI). Participants were asked to vote on each item using a 9-point scale, as follows: 1–3, not important; 4–6, important but not critical; and 7-9, important and critical. Respondents provided separate ratings for SPIRIT-AI and CONSORT-AI. There was an option to opt out of voting for each item, and each item included space for free text comments. At the end of the Delphi survey, participants had the opportunity to suggest new items. 103 responses were received for the first Delphi round, and 91 responses (88% of participants from round one) were received for the second round. The results of the Delphi survey informed the subsequent international consensus meeting. 12 new items were proposed by the Delphi study participants and were added for discussion at the consensus meeting. Data collected during the Delphi survey were anonymised, and item-level results were presented at the consensus meeting for discussion and voting.

The two-day consensus meeting took place in January, 2020, and was hosted by the University of Birmingham, UK, to seek consensus on the content of SPIRIT-AI and CONSORT-AI. 31 international stakeholders from among the Delphi survey participants were invited to discuss the items and vote on their inclusion. Participants were selected to achieve adequate representation from all the stakeholder groups. 41 items were discussed in turn, comprising the 29 items generated in the initial literature review and item-generation phase (26 items relevant to both SPIRIT-AI and CONSORT-AI; three items relevant only to CONSORT-AI) and the 12 new items proposed by participants during the Delphi surveys. Each item was presented to the Consensus Group, alongside its score from the Delphi exercise (median and interquartile ranges) and any comments made by Delphi participants related to that item. Consensus meeting participants were invited to comment on the importance of each item and whether the item should be included in the AI extension. In addition, participants were invited to comment on the wording of the explanatory text accompanying each item and the position of each item relative to the SPIRIT 2013 and CONSORT 2010 checklists. After open discussion of each item and the option to adjust wording, an electronic vote took place, with the option to include or exclude the item. An 80% threshold for inclusion was pre-specified and deemed reasonable by the Steering Group to demonstrate majority consensus. Each stakeholder voted anonymously using Turning Point voting pads (Turning Technologies, version 8.7.2.14).

See Online for appendix

Section	Item	CONSORT 2010 item*		CONSORT-Al item	Addresse on page number
			Title and abstrac	<u> </u>	
Title and Abstract	1a	Identification as a randomised trial in the title	CONSORT-AI	(i) Indicate that the intervention involves artifical intelligence/machine learning in the title and/or abstract and specify the type of model.	
	1b	Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts)	1a,b Elaboration	(ii) State the intended use of the AI intervention within the trial in the title and/or abstract.	
			Introduction		
Background and objectives	2a	Scientific background and explanation of rationale	CONSORT-AI 2a (i) Extension	Explain the intended use of the AI intervention in the context of the clinical pathway, including its purpose and its intended users (for example, healthcare professionals, patients, public).	
	2b	Specific objectives or hypotheses			
			Methods		
	3a	Description of trial design (such as parallel, factorial) including allocation ratio			
Trial design	3b	Important changes to methods after trial commencement (such as eligibility criteria), with reasons			
Participants	4a	Eligibility criteria for participants	CONSORT-AI 4a (i) Elaboration	State the inclusion and exclusion criteria at the level of participants.	
			CONSORT-AI 4a (ii) Extension	State the inclusion and exclusion criteria at the level of the input data.	
	4b	Settings and locations where the data were collected	CONSORT-AI 4b Extension	Describe how the Al intervention was integrated into the trial setting, including any onsite or offsite requirements.	
		The interventions for each group with sufficient details to allow replication, including how and when they were actually administered	CONSORT-AI	State which version of the AI algorithm was used.	
nterventions	5		5 (i) Extension CONSORT-AI	Describe how the input data were acquired and selected for the Al	
			5 (ii) Extension	intervention.	
			CONSORT-AI 5 (iii) Extension	Describe how poor quality or unavailable input data were assessed and handled.	
			CONSORT-AI 5 (iv) Extension.	Specify whether there was human–Al interaction in the handling of the input data, and what level of expertise was required of users.	
			CONSORT-AI 5 (v) Extension	Specify the output of the AI intervention.	
			CONSORT-AI 5 (vi) Extension	Explain how the Al intervention's outputs contributed to decision-making or other elements of clinical practice.	
Outcomes	6a	Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed	3(1)		
	6b	Any changes to trial outcomes after the trial commenced, with reasons			
	7a	How sample size was determined			
Sample size	7b	When applicable, explanation of any interim analyses			
		and stopping guidelines	Randomisation		
	8a	Method used to generate the random allocation			
Sequence generation	8b	sequence Type of randomisation; details of any restriction (such as blocking and block size)			
Allocation oncealment 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			
plementation	10	Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions			
Blinding	11a	If done, who was blinded after assignment to interventions (for example, participants, care providers, those assessing outcomes) and how			
	11b	If relevant, description of the similarity of interventions			
Statistical methods	12a	Statistical methods used to compare groups for primary and secondary outcomes			
	12b	Methods for additional analyses, such as subgroup analyses and adjusted analyses			

(Figure 1 continues on next page)

#### Checklist pilot

Following the consensus meeting, attendees were given the opportunity to make final comments on the wording and agree that the updated SPIRIT-AI and CONSORT-AI items reflected discussions from the meeting. The Operations Team assigned each item as an extension or elaboration item on the basis of a decision tree and produced a penultimate draft of the SPIRIT-AI and CONSORT-AI checklists (appendix p 6). A pilot of the penultimate checklists was conducted with 34 participants

Section	Item	CONSORT 2010 item* CONSORT-Al item			Addressed on page number†				
Results									
Participant flow (a diagram is strongly recommended)	13a	For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome							
	13b	For each group, losses and exclusions after randomisation, together with reasons							
Recruitment	14a	Dates defining the periods of recruitment and follow-up							
	14b	Why the trial ended or was stopped							
Baseline data	15	A table showing baseline demographic and clinical characteristics for each group							
Numbers analysed	16	For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups							
Outcomes and estimation	17a	For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)							
	17b	For binary outcomes, presentation of both absolute and relative effect sizes is recommended							
Ancillary analyses	18	Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory							
Harms	19	All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)	CONSORT-AI 19 Extension	Describe results of any analysis of performance errors and how errors were identified, where applicable. If no such analysis was planned or done, explain why not.					
			Discussion						
Limitations	20	Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses							
Generalisability	21	Generalisability (external validity, applicability) of the trial findings							
Interpretation	22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence							
			Other Informatio	on					
Registration	23	Registration number and name of trial registry							
Protocol	24	Where the full trial protocol can be accessed, if available							
Funding	25	Sources of funding and other support (such as supply of drugs), role of funders	CONSORT-AI 25 Extension	State whether and how the AI intervention and/or its code can be accessed, including any restrictions to access or re-use.					

Figure 1: CONSORT-AI checklist

A = artificial intelligence. ML=machine learning. \*We strongly recommend reading this statement in conjunction with the CONSORT 2010 Explanation and Elaboration for important clarifications on all the items. †Indicates page numbers to be completed by authors during protocol development.

to ensure clarity of wording. Experts participating in the pilot included the following: (a) Delphi participants who did not attend the consensus meeting, and (b) external experts who had not taken part in the development process but who had reached out to the Steering Group after the Delphi study commenced. Final changes were made on wording only to improve clarity for readers, by the Operations Team (appendix p 7).

#### Recommendations

#### CONSORT-AI checklist items and explanation

The CONSORT-AI extension recommends that 14 new checklists items be added to the existing CONSORT 2010 statement (11 extensions and three elaborations). These items were considered sufficiently important for clinical-trial reports for AI interventions that they should be routinely reported in addition to the core CONSORT 2010 checklist items. Figure 1 lists the CONSORT-AI items.

The 14 items below passed the threshold of 80% for inclusion at the consensus meeting. CONSORT-AI 2a, CONSORT-AI 5 (ii), and CONSORT-AI 19 each resulted from the merging of two items after discussion with the Consensus Group. CONSORT-AI 4a was split into two

items (i) and (ii) for clarity and was voted upon separately. CONSORT 5-AI (iii) did not fulfil the criteria for inclusion on the basis of its initial wording (77% vote to include); however, after extensive discussion and rewording, the Consensus Group unanimously supported a re-vote, at which point it passed the inclusion threshold (97% to include). The Delphi and voting results for each included and excluded item are described in the appendix (pp 2–5).

#### Title and abstract

CONSORT-AI 1a,b (i) Elaboration: Indicate that the intervention involves artificial intelligence/machine learning in the title and/or abstract and specify the type of model Explanation. Indicating in the title and/or abstract of the trial report that the intervention involves a form of AI is encouraged, as it immediately identifies the intervention as an AI/ML intervention and also serves to facilitate indexing and searching of the trial report. The title should be understandable by a wide audience; therefore, a broader umbrella term such as "artificial intelligence" or "machine learning" is encouraged. More precise terms should be used in the abstract, rather than the title, unless they are broadly recognised as being a form of

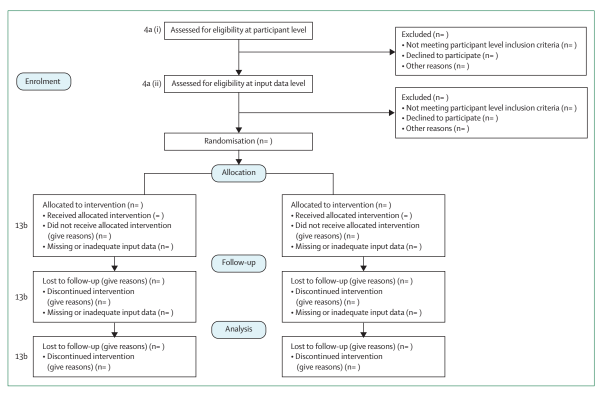


Figure 2: CONSORT 2010 flow diagram, adapted for AI clinical trials

AI=artificial intelligence. CONSORT-AI 4a (ii): State the inclusion and exclusion criteria at the level of participants. CONSORT-AI 4a (ii): State the inclusion and exclusion criteria at the level of the input data. CONSORT 13b (core CONSORT item): For each group, losses and exclusions after randomisation, together with reasons.

AI/ML. Specific terminology relating to the model type and architecture should be detailed in the abstract.

CONSORT-AI 1a,b (ii) Elaboration: State the intended use of the AI intervention within the trial in the title and/or abstract Explanation. Describe the intended use of the AI intervention in the trial report title and/or abstract. This should describe the purpose of the AI intervention and the disease context. Some AI interventions may have multiple intended uses, or the intended use may evolve over time. Therefore, documenting this allows readers to understand the intended use of the algorithm at the time of the trial.

#### Introduction

CONSORT-AI 2a (i) Extension: Explain the intended use for the AI intervention in the context of the clinical pathway, including its purpose and its intended users (for example, health-care professionals, patients, public)

Explanation. In order to clarify how the AI intervention is intended to fit into a clinical pathway, a detailed description of its role should be included in the background of the trial report. AI interventions may be designed to interact with different users, including health-care professionals, patients and the public, and their roles can be wide-ranging (for example, the same AI intervention could theoretically be replacing, augmenting, or adjudicating components of

clinical decision-making). Clarifying the intended use of the AI intervention and its intended user helps readers understand the purpose for which the AI intervention was evaluated in the trial.

#### Methods

CONSORT-AI 4a (i) Elaboration: State the inclusion and exclusion criteria at the level of participants

Explanation. The inclusion and exclusion criteria should be defined at the participant level as per usual practice in non-AI interventional trial reports (figure 2). This is distinct from the inclusion and exclusion criteria made at the input-data level, which is addressed in item 4a (ii).

CONSORT-AI 4a (ii) Extension: State the inclusion and exclusion criteria at the level of the input data

Explanation. "Input data" refers to the data required by the AI intervention to serve its purpose (for example, for a breast cancer diagnostic system, the input data could be the unprocessed or vendor-specific post-processing mammography scan upon which a diagnosis is being made; for an early-warning system, the input data could be physiological measurements or laboratory results from the electronic health record). The trial report should prespecify if there were minimum requirements for the input data (such as image resolution, quality metrics, or data format) that determined pre-randomisation eligibility. It

should specify when, how, and by whom this was assessed. For example, if a participant met the eligibility criteria for lying flat for a CT scan as per item 4a (i), but the scan quality was compromised (for any given reason) to such a level that it was deemed unfit for use by the AI system, this should be reported as an exclusion criterion at the input-data level. Note that where input data are acquired after randomisation, any exclusion is considered to be from the analysis, not from enrolment (CONSORT item 13b and figure 2).

## CONSORT-AI 4b Extension: Describe how the AI intervention was integrated into the trial setting, including any onsite or offsite requirements

Explanation. There are limitations to the generalisability of AI algorithms, one of which is when they are used outside of their development environment. AI systems are dependent on their operational environment, and the report should provide details of the hardware and software requirements to allow technical integration of the AI intervention at each study site. For example, it should be stated if the AI intervention required vendor-specific devices, if there was specialised computing hardware at each site, or if the site had to support cloud integration, particularly if this was vendor specific. If any changes to the algorithm were required at each study site as part of the implementation procedure (such as fine-tuning the algorithm on local data), then this process should also be clearly described.

### CONSORT-AI 5 (i) Extension: State which version of the AI algorithm was used

Explanation. Similar to other forms of software as a medical device, AI systems are likely to undergo multiple iterations and updates during their lifespan. It is therefore important to specify which version of the AI system was used in the clinical trial, whether this is the same as the version evaluated in previous studies that have been used to justify the study rationale, and whether the version changed during the conduct of the trial. If applicable, the report should describe what has changed between the relevant versions and the rationales for the changes. Where available, the report should include a regulatory marking reference, such as a unique device identifier, that requires a new identifier for updated versions of the device.<sup>47</sup>

### CONSORT-AI 5 (ii) Extension: Describe how the input data were acquired and selected for the AI intervention

Explanation. The measured performance of any AI system may be critically dependent on the nature and quality of the input data. A description of the input-data handling, including acquisition, selection, and pre-processing before analysis by the AI system, should be provided. Completeness and transparency of this description are integral to the replicability of the intervention beyond the clinical trial in real-world settings. It also helps readers

identify whether input-data-handling procedures were standardised across trial sites.

### CONSORT-AI 5 (iii) Extension: Describe how poor-quality or unavailable input data were assessed and handled

Explanation. As with CONSORT-AI 4a (ii), "input data" refers to the data required by the AI intervention to serve its purpose. As discussed in CONSORT-AI 4a (ii), the performance of AI systems may be compromised as a result of poor-quality or missing input data<sup>49</sup> (for example, excessive-movement artifact on an electrocardiogram). The trial report should report the amount of missing data, as well as how this was identified and handled. The report should also specify if there was a minimum standard required for the input data and, where this standard was not achieved, how this was handled (including the impact on, or any changes to, the participant care pathway).

Poor-quality or unavailable data can also affect non-AI interventions. For example, suboptimal quality of a scan could affect a radiologist's ability to interpret it and make a diagnosis. It is therefore important that this information is reported equally in the control intervention, where relevant. If this minimum-quality standard was different from the inclusion criteria for input data used to assess eligibility pre-randomisation, this should be stated.

### CONSORT-AI 5 (iv) Extension: Specify whether there was human–AI interaction in the handling of the input data, and what level of expertise was required of users

Explanation. A description of the human-AI interface and the requirements for successful interaction when input data are handled should be provided—for example, clinician-led selection of regions of interest from a histology slide that is then interpreted by an AI diagnostic system,50 or an endoscopist's selection of colonoscopy video clips as input data for an algorithm designed to detect polyps.28 A description of any user training provided and instructions for how users should handle the input data provides transparency and replicability of trial procedures. Poor clarity on the human-AI interface may lead to a lack of a standard approach and may carry ethical implications, particularly in the event of harm. 51,52 For example, it may become unclear whether an error case occurred due to human deviation from the instructed procedure, or if it was an error made by the AI system.

### CONSORT-AI 5 (v) Extension: Specify the output of the AI intervention

Explanation. The output of the AI intervention should be clearly specified in the trial report. For example, an AI system may output a diagnostic classification or probability, a recommended action, an alarm alerting to an event, an instigated action in a closed-loop system (such as titration of drug infusions), or another output. The nature of the AI intervention's output has direct implications on its usability and how it may lead to downstream actions and outcomes.

CONSORT-AI 5 (vi) Extension: Explain how the AI intervention's outputs contributed to decision-making or other elements of clinical practice

Explanation. Since health outcomes may also critically depend on how humans interact with the AI intervention, the report should explain how the outputs of the AI system were used to contribute to decision-making or other elements of clinical practice. This explanation should include adequate description of downstream interventions that can affect outcomes. As with CONSORT-AI 5 (iv), any effcts of human-AI interaction on the outputs should be described in detail, including the level of expertise required to understand the outputs and any training and/or instructions provided for this purpose. For example, a skin-cancer-detection system that produced a percentage likelihood as its output should be accompanied by an explanation of how this output was interpreted and acted upon by the user, specifying both the intended pathways (for example, skin-lesion excision if the diagnosis is positive) and the thresholds for entry to these pathways (for example, skinlesion excision if the diagnosis is positive and the probability is >80%). The information produced by comparator interventions should be similarly described, alongside an explanation of how such information was used to arrive at clinical decisions on patient management, where relevant. Any discrepancy in how decisionmaking occurred versus how it was intended to occur (that is as specified in the trial protocol) should be reported.

#### Results

CONSORT-AI 19 Extension: Describe results of any analysis of performance errors and how errors were identified, where applicable. If no such analysis was planned or done, explain why not

Explanation. Reporting performance errors and failure case analysis is especially important for AI interventions. AI systems can make errors that may be hard to foresee but that, if allowed to be deployed at scale, could have catastrophic consequences.<sup>53</sup> Therefore, reporting cases of error and defining risk-mitigation strategies are important for informing when, and for which populations, the intervention can be safely implemented. The results of any performance-error analysis should be reported and the implications of the results should be discussed.

#### Other information

CONSORT-AI 25 Extension: State whether and how the AI intervention and/or its code can be accessed, including any restrictions to access or re-use

Explanation. The trial report should make it clear whether and how the AI intervention and/or its code can

be accessed or re-used. This should include details about the license and any restrictions to access.

#### Discussion

CONSORT-AI is a new reporting-guideline extension developed through international multi-stakeholder consensus. It aims to promote transparent reporting of AI intervention trials and is intended to facilitate critical appraisal and evidence synthesis. The extension items added in CONSORT-AI address a number of issues specific to the implementation and evaluation of AI interventions, which should be considered alongside the core CONSORT 2010 checklist and other CONSORT extensions.<sup>54</sup> It is important to note that these are minimum requirements and there may be value in including additional items not included in the checklists (appendix pp 2–5) in the report or in supplementary materials.

In both CONSORT-AI and its companion project SPIRIT-AI, a major emphasis was the addition of several new items related to the intervention itself and its application in the clinical context. Items 5 (i)-5 (vi) were added to address AI-specific considerations in descriptions of the intervention. Specific recommendations were made pertinent to AI systems related to algorithm version, input and output data, integration into trial settings, expertise of the users, and protocol for acting upon the AI system's recommendations. It was agreed that these details are critical for independent evaluation or replication of the trial. Journal editors reported that despite the importance of these items, they are currently often missing from trial reports at the time of submission for publication, which provides further weight for their inclusion as specifically listed extension items.

A recurrent focus of the Delphi comments and Consensus Group discussion was the safety of AI systems. This was in recognition that AI systems, unlike other health interventions, can unpredictably yield errors that are not easily detectable or explainable by human judgement. For example, changes to medical imaging that are invisible, or appear random, to the human eye may change the likelihood of the diagnostic output entirely.55,56 The concern is that given the theoretical ease with which AI systems could be deployed at scale, any unintended harmful consequences could be catastrophic. CONSORT-AI item 19, which requires specification of any plans to analyse performance errors, was added to emphasise the importance of anticipating systematic errors made by the algorithm and their consequences. Beyond this, investigators should also be encouraged to explore differences in performance and error rates across population subgroups. It has been shown that AI systems may be systematically biased toward different outputs, which may lead to different or even unfair treatment, on the basis of extant features. 53,57-59

The topic of "continuously evolving" AI systems (also known as "continuously adapting" or "continuously learning" AI systems) was discussed at length during the

consensus meeting, but it was agreed that this be excluded from CONSORT-AI. These are AI systems with the ability to continuously train on new data, which may cause changes in performance over time. The group noted that, while of interest, this field is relatively early in its development without tangible examples in health-care applications, and that it would not be appropriate for it to be included in CONSORT-AI at this stage.60 This topic will be monitored and will be revisited in future iterations of CONSORT-AI. It is worth noting that incremental software changes, whether continuous or iterative, intentional or unintentional, could have serious consequences on safety performance after deployment. It is therefore of vital importance that such changes be documented and identified by software version and that a robust postdeployment surveillance plan is in place.

This study is set in the current context of AI in health; therefore, several limitations should be noted. First, there are relatively few published interventional trials in the field of AI for health care; therefore, the discussions and decisions made during this study were not always supported by existing examples of completed trials. This arises from our stated aim of addressing the issues of poor reporting in this field as early as possible, recognising the strong drivers in the field and the specific challenges of study design and reporting for AI. As the science and study of AI evolves, we welcome collaboration with investigators to co-evolve these reporting standards to ensure their continued relevance. Second, the literature search for AI RCTs used terminology such as "artificial intelligence", "machine learning", and "deep learning", but not terms such as "clinical decision support systems" or "expert systems", which were more commonly used in the 1990s for technologies underpinned by AI systems and share risks similar to those of recent examples.61 It is likely that such systems, if published today, would be indexed under "artificial intelligence" or "machine learning"; however, clinicaldecision support systems were not actively discussed during this consensus process. Third, the initial candidate-items list was generated by a relatively small group of experts consisting of Steering Group members and additional international experts; however, additional items from the wider Delphi group were taken forward for consideration by the Consensus Group, and no new items were suggested during the consensus meeting or post-meeting evaluation.

As with the CONSORT statement, the CONSORT-AI extension is intended as a minimum reporting guidance, and there are additional AI-specific considerations for trial reports that may warrant consideration (appendix pp 2–5). This extension is aimed particularly at investigators and readers reporting or appraising clinical trials; however, it may also serve as useful guidance for developers of AI interventions in earlier validation stages of an AI system. Investigators seeking to report studies developing and validating the diagnostic and predictive

properties of AI models should refer to TRIPOD-ML (Transparent Reporting of a Multivariable Prediction Model for Individual Prognosis or Diagnosis-Machine Learning) and STARD-AI (Standards for Reporting of Diagnostic Accuracy Studies-Artificial Intelligence), both of which are currently under development. 32,62 Other potentially relevant guidelines, which are agnostic to study design, are registered with the EQUATOR Network.63 The CONSORT-AI extension is expected to encourage careful early planning of AI interventions for clinical trials and this, in conjunction with SPIRIT-AI, should help to improve the quality of trials for AI interventions. The development of the CONSORT-AI guidance does not include additional items within the discussion section of trial reports. The guidance provided by CONSORT 2010 on trial limitations, generalisability, and interpretation was deemed to be translatable to trials for AI interventions.

There is also recognition that AI is a rapidly evolving field, and there will be the need to update CONSORT-AI as the technology, and newer applications for it, develop. Currently, most applications of AI involve disease detection, diagnosis, and triage, and this is likely to have influenced the nature and prioritisation of items within CONSORT-AI. As wider applications that utilise "AI as therapy" emerge, it will be important to continue to evaluate CONSORT-AI in light of such studies. Additionally, advances in computational techniques and the ability to integrate them into clinical workflows will bring new opportunities for innovation that benefits patients. However, they may be accompanied by new challenges around study design and reporting. In order to ensure transparency, minimise potential biases, and promote the trustworthiness of the results and the extent to which they may be generalisable, the SPIRIT-AI and CONSORT-AI Steering Group will continue to monitor the need for updates.

#### Contributors

All authors contributed to the concept and design of the study and the acquisition, analysis, and interpretation of data. XL, SCR, DM, MJC, and AKD contributed to the drafting of the manuscript. AKD, MJC, CY, and CHo obtained funding. The SPIRIT-AI and CONSORT-AI Group consists of two working groups that have been key in the development of the guidelines: the SPIRIT-AI and CONSORT-AI Steering Group, which was responsible for overseeing the consensus process and guidelines development methodology, and the SPIRIT-AI and CONSORT-AI Consensus Group, which was responsible for reaching consensus on the content and wording of the items within the checklists.

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#### Declaration of interests

MJC has received personal fees from Astellas, Takeda, Merck, Daiichi Sankyo, Glaukos, GlaxoSmithKline, and the Patient-Centered Outcomes Research Institute (PCORI), outside the submitted work. ADa is an advisor for Google DeepMind, outside the submitted work. LF reports personal fees from Allergan, Bayer, and Novartis, outside the submitted work. JF reports personal fees from British Medical Journal, during the conduct of the study. HH reports that he is Managing Director at Hardian Health, consultancy for health technology firms. PAK reports personal fees from DeepMind Technologies, Roche, Novartis, Apellis, Bayer, Allergan, Topcon, and Heidelberg Engineering, outside the submitted work. AYL reports personal fees from Genentech, US Food and Drug Administration, and Verana Health, grants from Microsoft, NVIDIA, Carl Zeiss Meditec, and Santen, outside the submitted work. CSL reports grants from National Institute of Health/ National Institute

on Aging, outside the submitted work. CJK is an employee of Google and owns Alphabet stock. AE is an employee of Salesforce CRM. RiS is an employee of Pinpoint Science. JMa was an employee of AstraZeneca PLC at the time of this study. RuS is Editor-in-Chief of The Lancet Digital Health and reports personal fees from The Lancet Group, during the conduct of the study. JMo is Chief Editor of the journal Nature Medicine; he has recused himself from any aspect of decision-making on this manuscript and played no part in the assignment of this manuscript to in-house editors or peer reviewers, and was also separated and blinded from the editorial process from submission inception to decision. SJV reports funding from IQVIA. All other authors declare no competing interests.

#### Data sharing

Data requests should be made to the corresponding author and release will be subject to consideration by the SPIRIT-AI and CONSORT-AI Steering Group.

#### Acknowledgments

We thank the participants who were involved in the Delphi study and Pilot study (Supplementary Note), Eliot Marston for providing strategic support (University of Birmingham, Birmingham, UK), and Charlotte Radovanovic (University Hospitals Birmingham NHS Foundation Trust, UK) and Anita Walker (University of Birmingham, UK) for administrative support. The views expressed in this publication are those of the authors, Delphi participants and stakeholder participants and may not represent the views of the broader stakeholder group or host institution. This work was funded by a Wellcome Trust Institutional Strategic Support Fund: Digital Health Pilot Grant, Research England (part of UK Research and Innovation), Health Data Research UK, and the Alan Turing Institute. The study was sponsored by the University of Birmingham, UK. The study funders and sponsors had no role in the design and conduct of the study; collection, management, analysis and interpretation of the data; preparation, review or approval of the manuscript; or decision to submit the manuscript for publication. MJC is a National Institute for Health Research (NIHR) Senior Investigator and receives funding from the NIHR Birmingham Biomedical Research Centre; the NIHR Surgical Reconstruction and Microbiology Research Centre and NIHR ARC West Midlands at the University of Birmingham and University Hospitals Birmingham NHS Foundation Trust; Health Data Research UK; Innovate UK (part of UK Research and Innovation); the Health Foundation; Macmillan Cancer Support; and UCB Pharma. ADa and JJD are also NIHR Senior Investigators. The views expressed in this article are those of the author(s) and not necessarily those of the NIHR, or the Department of Health and Social Care. DM is supported by a University of Ottawa Research Chair. MKEZ is supported by the US Food and Drug Administration (FDA), and DP is supported in part by the Office of the Director at the National Library of Medicine (NLM), US National Institutes of Health (NIH). AB is supported by an NIH award 7K01HL141771-02. PAK received grants from UKRI Future Leaders Fellowship and from Moorfields Eye Charity Career Development Award. SJV received funding from the Engineering and Physical Sciences Research Council, UK Research and Innovation (UKRI), Accenture, Warwick Impact Fund, Health Data Research UK, and European Regional Development Fund. SR is an employee of the UKRI. This article may not be consistent with NIH and/or FDA's views or policies. It reflects only the views and opinions of the authors.

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