

TRIPOD+AI statement: updated guidance for reporting clinical prediction models that use regression or machine learning methods

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The TRIPOD (Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis) statement was published in 2015 to provide the minimum reporting recommendations for studies developing or evaluating the performance of a prediction model. Methodological advances in the field of prediction have since included the widespread use of artificial intelligence (AI) powered by machine learning methods to develop prediction models. An update to the TRIPOD statement is thus needed. TRIPOD+AI provides harmonised guidance for reporting prediction model studies, irrespective

of whether regression modelling or machine learning methods have been used. The new checklist supersedes the TRIPOD 2015 checklist, which should no longer be used. This article describes the development of TRIPOD+AI and presents the expanded 27 item checklist with more detailed explanation of each reporting recommendation, and the TRIPOD+AI for Abstracts checklist. TRIPOD+AI aims to promote the complete, accurate, and transparent reporting of studies that develop a prediction model or evaluate its performance. Complete reporting will facilitate study appraisal, model evaluation, and model implementation.

Prediction models are used across different healthcare settings. They are used to estimate an outcome value or risk. Most models estimate the probability of the presence of a particular health condition (diagnostic) or whether a particular outcome will occur in the future (prognostic).¹ Their primary use is to support clinical decision making, such as whether to refer patients for further testing, monitor disease deterioration or treatment effects, or initiate treatment or lifestyle changes. Examples of well known prediction models include EuroSCORE II (cardiac surgery),² the Gail model (breast cancer),³ the Framingham risk score (cardiovascular disease),⁴ IMPACT (traumatic brain injury),⁵ and FRAX (osteoporotic and hip fractures).⁶

Prediction models are abundant in the biomedical literature, with thousands of models published annually (and increasing), and have been developed for many outcomes and health conditions.^{7,8} At least 731 diagnostic and prognostic prediction model studies on covid-19 were published during the first 12 months of the pandemic.⁹ Despite this interest in developing prediction models, there have been longstanding

SUMMARY POINTS

There has been considerable interest and financial investment in developing prediction models by applying artificial intelligence (AI) methods, typically powered by advances in machine learning

To ensure that a prediction model study is valuable to users, authors should prepare a transparent, complete, and accurate account of why the research was done, what they did, and what they found

An update of the TRIPOD (Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis) statement aims to harmonise the landscape of prediction model studies using AI methods and to provide guidance regardless of whether regression models or machine learning methods have been used

The TRIPOD+AI statement consists of a 27 item checklist, an expanded checklist that details reporting recommendations for each item, and a TRIPOD+AI for Abstracts checklist containing 13 items

TRIPOD+AI aims to assist authors in the complete reporting of their study and help peer reviewers, editors, policymakers, end users, and patients understand the data, methods, findings and conclusions of AI driven research

Adherence to the TRIPOD+AI reporting recommendations could encourage the improved use of research time, effort, and money

concerns about transparency and completeness of reporting in the field,^{10 11} and the resulting usability. For readers (including peer reviewers, editors, health professionals, regulators, patients, and the general public), incomplete or inaccurate reporting impairs the ability to critically appraise the study design and methods, have confidence in the findings, and further evaluate or implement a prediction model. Poor reporting of a model might also mask flaws in the design, data collection, or conduct of a study that, if the model was implemented in the clinical pathway, could cause harm. Harm can be perceived to occur when insufficient measures are in place to mitigate bias. Better reporting can create more trust and influence patient and public acceptability of the use of prediction models in healthcare. Authors have an ethical and scientific obligation to honestly report their research in a complete and transparent manner. As noted by the late Doug Altman and colleagues, “Good reporting is not an optional extra; it is an essential component of research”¹²—anything less is little more than avoidable research waste.¹³

In response to concerns about incomplete reporting,^{10 11 14 15} the TRIPOD (Transparent Reporting of a multivariable model for Individual Prognosis Or Diagnosis) statement was published in 2015 (TRIPOD 2015) to provide minimum reporting recommendations.^{16 17} TRIPOD 2015 comprises a checklist of 37 items, which includes 25 items to report in both development and validation studies, and an additional six items for model development studies and six items for validation studies. Accompanying the checklist is an explanation and elaboration document that provides the rationale behind each reporting item; published examples of good reporting; and a discussion of issues relating to the design, conduct, and analysis of prediction model studies.¹⁷ TRIPOD 2015 mainly focused on models developed using regression modelling, which was the prevailing approach at the time. Additional guidance has since been created for reporting abstracts of prediction model studies (TRIPOD for Abstracts¹⁸), studies developing or validating prediction models using clustered data (TRIPOD-Cluster^{19 20}), systematic reviews and meta-analyses of prediction model studies (TRIPOD-SRMA²¹), and guidance in preparation for study protocols (TRIPOD-P²²). All available guidance, as well as template checklists for filling out separately, can also be found on the TRIPOD website (<https://www.tripod-statement.org/>).

Since the publication of TRIPOD 2015, there have been numerous methodological advances in prediction modelling, including sample size guidance for developing models^{23–27} and evaluating their performance,^{28–32} and greater recognition of operationalising fairness,³³ reproducibility,³⁴ and adopting open science principles.³⁵ However, interest and financial investment in applying methods ascribed to artificial intelligence (AI), typically powered by advances in machine learning methods (eg, random forests, deep learning), is where we have seen the

most progress and change. With increasing access to data and availability of off-the-shelf software to apply machine learning methods, developing a prediction model has become faster and easier. Vast numbers of prediction models are now entering the scientific literature for many clinical settings, and for a wide range of outcomes and health conditions, with multiple models often available for the same outcome, health condition, and target population.^{7 8 36} The ability to critically evaluate the quality of prediction models and understand their ability to serve well in a particular setting or for a particular use case is therefore of even greater critical importance. This ability is predicated on complete and transparent reporting.

However, systematic reviews evaluating studies of prediction models have shown that they are often poorly conducted (including deficiencies in study design or data collection^{37 38}); use poor methodology^{37 38}; are incompletely reported with key details missing^{39–54}; are consequently at high risk of bias^{41 49 55–57}; rarely adhere to open science practices,⁵⁸ and are susceptible to overinterpretation or so-called spin.^{59 60} These deficiencies cast considerable doubt on models’ usefulness and safety, and raises concerns about their potential to create or widen healthcare disparities.⁶¹ While TRIPOD 2015 is largely agnostic to the type of modelling approach, and much of its reporting recommendations apply equally to non-regression approaches, additional reporting considerations are needed for the growing class of machine learning methods. For example, unlike regression based models, the flexibility and complexity underpinning other machine learning approaches typically means that the resulting prediction models do not result in a simple equation and sometimes even the predictors used remain unclear. Additional reporting considerations are therefore needed that are not currently covered in TRIPOD 2015. Alongside methodological advancements, considerations of fairness,⁶² wider acceptance of open science practices,⁶³ and public and patient involvement in research and implementation of research,^{64 65} an update to the TRIPOD 2015 statement is needed to capture these developments and the consequences for reporting.

The aim of this paper is to describe the development of the updated TRIPOD guidance, present the new TRIPOD+AI checklist, and discuss how to use it. TRIPOD+AI aims to harmonise the landscape of prediction model studies and provide guidance regardless of whether regression models or machine learning methods have been used.⁶⁶ The “+” in TRIPOD+AI indicates that it provides consolidated reporting recommendations for studies of prediction models developed using regression modelling or machine learning (ie, deep learning, random forests) approaches. We also use the additional term “AI” to be consistent with existing reporting guidelines for studies broadly labelled as involving AI. However, for readability, this article will refer to the methods underpinning them as machine learning (table 1). A glossary of terms (box 1) clarifies key concepts used within the TRIPOD+AI reporting guideline.

Table 1 | Reporting guidelines for healthcare studies using machine learning

Reporting guideline	Scope
STARD-AI	Studies evaluating the diagnostic accuracy of an artificial intelligence based test (in preparation) ⁶⁷
TRIPOD+AI	Studies developing or evaluating the performance of a prediction model, using artificial intelligence, including machine learning methods
CLAIM	Medical imaging studies using artificial intelligence ⁶⁸
DECIDE-AI	Early stage clinical evaluation (including safety, human factors evaluation) of decision support systems driven by artificial intelligence ⁶⁹
CHEERS-AI	Studies describing health economic evaluations to estimate the value for money (cost effectiveness) of artificial intelligence interventions ⁷⁰
SPIRIT-AI	Protocols for clinical trials evaluating an intervention with an artificial intelligence component ⁷¹
CONSORT-AI	Clinical trial reports evaluating an intervention with an artificial intelligence component ⁷²
PRISMA-AI	Systematic reviews and meta-analyses of artificial intelligence interventions (in preparation) ⁷³

STARD=Standards for Reporting of Diagnostic Accuracy; TRIPOD=Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis; AI=artificial intelligence; CLAIM=Checklist for Artificial Intelligence in Medical Imaging; DECIDE=Decisions in health Care to Introduce or Diffuse innovations using Evidence; CHEERS=Consolidated Health Economic Evaluation Reporting Standards; SPIRIT=Standard Protocol Items: Recommendations for Interventional Trials; CONSORT=Consolidated Standards of Reporting Trials; PRISMA=Preferred Reporting Items for Systematic Reviews and Meta-Analyses.

Development of TRIPOD+AI

We describe the development of the TRIPOD+AI statement, a guideline to aid the reporting of studies developing prediction models for diagnosis or prognosis using machine learning or regression methods or evaluating (validating) their performance. There is no such thing as a validated prediction model.⁷⁶ To avoid ambiguity and harmonise terminology, we refer to validation as evaluation⁷⁴ in this article (box 1). Existing reporting guidelines and those in development for reporting other types of biomedical studies involving a machine learning component are detailed in table 1. Literature reviews and consensus exercises were used to develop the TRIPOD+AI checklist as recommended by the EQUATOR Network.⁷⁷ A steering group was convened by GSC and KGMM to oversee the guideline development process, with members selected to cover a broad range of expertise and experience (comprising GSC, KGMM, RDR, ALB, JBR, BVC, XL, and PD).

In April 2019, a commentary was published announcing the TRIPOD+AI initiative.⁷⁸ The guideline was registered as a reporting guideline under development with the EQUATOR Network on 7 May 2019 (<https://www.equator-network.org/>). A study protocol was made available on 25 March 2021 on the Open Science Framework (<https://osf.io/zyacb/>), describing the process and methods used to develop the TRIPOD+AI reporting guideline. The protocol, which also describes the development of a quality assessment and risk-of-bias tool for prediction models developed using machine learning methods (PROBAST+AI), was published in 2021.⁷⁹ The reporting of the consensus based methods used in the development of TRIPOD+AI followed the ACCORD (Accurate Consensus Reporting Document) recommendations.⁸⁰

Ethics

This study was approved by the Central University research ethics committee, University of Oxford on 10 December 2020 (R73034/RE001). Participant information was provided to the Delphi survey participants electronically before starting the survey and to the consensus participants before the consensus meeting. Delphi survey participants provided electronic informed consent before completing the survey.

Candidate item list generation

An initial list of items was drafted by GSC and KGMM using TRIPOD 2015.^{16 17} Additional items were identified from TRIPOD-Cluster,^{19 20} TRIPOD for Abstracts,¹⁸ CAIR,⁸¹ MI-CLAIM,⁸² CLAIM,⁶⁸ MINIMAR,⁸³ SPIRIT-AI,⁷¹ and CONSORT-AI,⁷² along with additional literature identified by the steering group.^{34 84-89} The list of items was also informed by the findings of systematic reviews evaluating the reporting, methods, and overinterpretation of prediction model studies using machine learning.^{37-39 48 51 54 59 60} The steering group harmonised the initial list of items to form a final list of 65 unique candidate items covering the title (one item); abstract (one item); introduction (three items); methods (37 items); results (15 items); discussion (five items), and other (three items). This list was used in a modified Delphi exercise as described below.

Recruitment of Delphi panellists

Delphi participants were identified by the steering committee, from authors of relevant publications via a call to participate on social media (eg, Twitter), and through personal recommendations. Including experts recommended by other Delphi participants. The steering group identified participants to achieve geographical and disciplinary diversity and include key stakeholder groups, for example, researchers (statisticians/data scientists, epidemiologists, machine learning researchers/scientists, clinicians, radiologists, and ethicists), healthcare professionals, journal editors, funders, policymakers, healthcare regulators, patients, and the general public as end users of prediction models from a range of settings (eg, universities, hospitals, primary care, biomedical journals, non-profit organisations, and for-profit organisations).

No minimum sample size was placed on the number of Delphi participants. A steering group member checked the expertise or experience of each identified person. Individuals were then invited to participate via email and were sent an information pack with the study description, aims, and contact details. Once participants accepted, they were added to the Delphi panel and received the link to the survey. Delphi panellists did not receive any financial incentive or gift to participate.

Box 1: Glossary of terms used in TRIPOD+AI

The definitions and descriptions given below relate to the specific context of the TRIPOD+AI* guideline; they do not necessarily apply to other areas of research.

Artificial intelligence

Field of computer science that focuses on developing models and algorithms capable of performing tasks that typically require human intelligence.

Calibration

Agreement between observed outcomes and estimated values from the model. Calibration is best assessed graphically with a plot of the estimated values on the x axis and observed values on the y axis, with a smoothed flexible calibration curve in the individual data.

Care pathway

Structured and coordinated plan of care for managing a specific health condition or dealing with a patient's healthcare needs throughout their healthcare journey.

Class imbalance

When the frequency of individuals with and without the outcome event is unequal.

Discrimination

How well the predictions from the model differentiate between individuals with and without the outcome. Discrimination is typically quantified by the c statistic (sometimes referred to as the area under the curve (AUC) or area under the receiver operating characteristics (AUROC)) for binary outcomes, and the c index for time-to-event outcomes.

Evaluation or test data

Data used to estimate the performance of a prediction model, sometimes referred to as test data or validation data.[†] Evaluation data should be distinct from the data used to train the model, tune hyperparameters, or do model selection, such that there is no overlap in participants between the training and evaluation data. Evaluation data should be representative of the population in whom the model is to be used.

Fairness

Property of prediction models that do not discriminate against individuals or groups of individuals based on attributes such as age, race/ethnicity, sex/gender, or socioeconomic status.

Hyperparameters

Values that control the model development or learning process.

Hyperparameter tuning

Finding the best (hyper)parameter settings for a particular model building strategy.

Internal validation

Evaluating the performance of a prediction model on the same population on which the model was developed (eg, train test split, cross validation, or bootstrapping).

Machine learning

A subfield of artificial intelligence that focuses on developing models capable of learning and making predictions or decisions from data, without being explicitly programmed.

Model evaluation

Evaluating predictive accuracy of a model by estimating model discrimination (eg, c statistic), model calibration (eg, calibration plot, calibration slope), and clinical utility (eg, decision curve analysis). This process is referred to as evaluating a prediction model.⁷⁴⁷⁵

Outcome

Diagnostic or prognostic event that is being predicted. In machine learning, this event is often referred to as the target value, response variable, or label.

Predictor

Characteristic that can be measured or attributed at an individual level (eg, age, systolic blood pressure, sex, disease stage, radiomics features) or group level (eg, country). It is also often referred to as an input, feature, independent variable, or covariate.

Training or development data

Data used to train or develop a prediction model. The training data are ideally representative of the population in whom the model is to be used.

*TRIPOD=Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis; AI=artificial intelligence.

[†]Validation data often has different meanings. For example, in machine learning studies, validation data can refer to data used for parameter tuning or data used to evaluate model performance (often referred to as external validation). To avoid any ambiguity, we refer to data used to evaluate model performance as evaluation data.

Delphi process

The Delphi surveys were designed and delivered electronically using the Welphi online platform (www.welphi.com) to be responded to individually, online,

and in English. The platform ensures responses are anonymous by sending a different link to each participant and applying codes to respondents. The panellists received a package of information clarifying the study's

objectives and scope and explaining how to participate, use the platform, and contact the development team with any questions. Participants were asked to rate each item as “can be omitted,” “possibly include,” “desirable for inclusion,” or “essential for inclusion.” Participants were also invited to comment on any item, and to suggest new items. Free text responses were collated and analysed by PL. The themes generated were used by GSC and KGMM to inform item rephrasing, merging, or suggesting new items. All members of the steering group were invited to participate in the Delphi surveys.

Round 1 participants

The invitation and participation link was sent to 292 people. The first round was conducted between 19 April 2021 and 13 May 2021. A reminder message was sent on 5 May 2021. Of 292 people invited, 170 completed the survey, including eight who provided partial responses. Survey participants came from 22 countries, predominantly the UK (n=52), US (n=31), Netherlands (n=23), and Canada (n=20), representing five continents (Europe: 100, South America: 2, North America: 51, Australasia: 4, Asia: 13). Seven participants did not declare their country.

Participants reported their primary fields of research/work and could select more than one field. They indicated statistics and data science (n=70), AI or machine learning (n=69), clinical (n=50) or epidemiology (n=40), prediction (n=18), radiology (n=18), health policy/regulatory (n=10), biomedical research (n=7), journal editor (n=6), meta-research/reporting (n=6), pathology (n=2), funder (n=2), ethics (n=2), technology development/implementation (n=2), genetics/genomics (n=2), biomedical engineering (n=2), and health economics (n=2).

Round 2 participants

The second round of the Delphi was conducted between 16 December 2021 and 17 January 2022. All participants who completed the first Delphi round were invited to the second Delphi round. Additional participants who did not respond to round 1 were reinvited, as were participants identified or recommended after round 1. Invitations for round 2 were sent to 395 people, of whom 200 completed the survey, including 15 who provided partial responses. Survey participants came from 27 different countries, again predominantly the UK (n=70), US (n=37), Netherlands (n=19), and Canada (n=19), and represented six continents (Europe: 123, South America: 3, North America: 56, Australasia: 7, Asia: 10, Africa: 1). Participants reported their primary fields of research/work and could select more than one field: statistics and data science (n=78), AI or machine learning (n=72), clinical (n=49) or epidemiology (n=51), prediction (n=19), radiology (n=26), health policy (n=12), biomedical research (n=14), journal editor (n=13), meta-research/reporting (n=6), biomedical engineering (n=5), funder (n=2), genetics/genomics (n=4), patient representative/engagement (n=3), health economics (n=2), and ethics (n=1).

Checklist item evolution from round 1 to round 2

In round 1 of the modified Delphi, participants rated 65 initial candidate items generated from literature reviews and other reporting checklists, as described above. Agreement was considered when the individuals agreed an item was desirable or essential for inclusion. As defined in the protocol,⁷⁹ items with agreement of 70% or higher were carried over to round 2. Items that had an agreement rate lower than 70% were excluded, merged, or rephrased to be presented to panellists for revaluation. These modifications were based on or inspired by the hundreds of comments added by panellists.

In round 2, survey participants were given a link to the aggregated ratings from round 1 (<https://osf.io/zyacb/>; supplementary table 3). In round 2, participants rated 59 candidate items, covering the title (one item), abstract (one item), introduction (four items), methods (32 items), results (11 items), discussion (eight items), and other (two items). The item relating to patient and public involvement received 69% agreement for inclusion (supplementary table 4). Despite falling just below the 70% threshold, the steering group agreed to retain this item for discussion during the consensus meeting.

Patient and public involvement and engagement meeting

An online meeting was held on 8 April 2022 with nine members of the Health Data Research UK's group for patient and public involvement and engagement (PPIE) (<https://www.hdruk.ac.uk/about-us/involving-and-engaging-patients-and-the-public/>), chaired by Sophie Staniszewska (University of Warwick, UK). This meeting was not planned in the study protocol and was the only deviation from the published protocol.⁷⁹ Before the meeting, the PPIE group was sent a summary of the TRIPOD+AI project (available at <https://osf.io/zyacb/>), including an executive summary drafted by one member of PPIE group, and the draft checklist. At the meeting, GSC presented details of the TRIPOD+AI initiative, the project status, and draft guidance resulting from round 2 of the Delphi survey. Participants then asked questions and discussed the project aims and scope. Following feedback received at the PPI meeting, and through correspondence written after the meeting, the draft checklist was revised to improve clarity. Three members of the PPI group were invited and two subsequently attended the online consensus meeting with the wider group of stakeholders on 5 July 2022. The manuscript was circulated to the three PPI members for their input and approval.

Consensus meeting

An online consensus meeting was held on 5 July 2022, chaired by GSC and KGMM. Participants were chosen to try to ensure balanced representation of the key stakeholder groups, disciplines, and geographical diversity. Twenty eight participants attended part or all of the meeting, including one non-voting attendee (PL).

Before the meeting, invited participants were emailed a document (available at <https://osf.io/zyacb/>) containing a brief overview of TRIPOD+AI, the consensus

meeting format and instructions, a summary of the aggregated responses from round 2 of the Delphi survey (supplementary table 3), and the draft TRIPOD+AI checklist. The checklist circulated to the consensus meeting participants included 59 items covering: title (one item), abstract (one item), introduction (four items), methods (32 items), results (11 items), discussion (eight items), and other (two items).

Given the high endorsement achieved for many items in round 2, a subset of 17 items were highlighted for plenary discussion and voting during the consensus meeting. After discussion, participants were given one minute to vote to include or exclude the item from the TRIPOD+AI checklist. The voting was registered using

the poll function of the online meeting program. The 17 items included one item that had not achieved consensus in round 2 and 16 items that had undergone rewording after round 2 or were new items that were not included in TRIPOD 2015. After discussion and voting on these 17 items, the final TRIPOD+AI checklist was formed.

TRIPOD+AI statement

TRIPOD+AI comprises a checklist of items that are considered essential for good reporting of studies developing or evaluating (validating) a prediction model using any statistical or machine learning methods (table 2). Box 2 summarises noteworthy additions and changes to TRIPOD 2015. The TRIPOD+AI checklist comprises

Table 2 | TRIPOD+AI checklist for the reporting of prediction model studies

Section/topic	Item	Development/evaluation*	Checklist item
Title			
Title	1	D;E	Identify the study as developing or evaluating the performance of a multivariable prediction model, the target population, and the outcome to be predicted
Abstract			
Abstract	2	D;E	See TRIPOD+AI for Abstracts checklist
Introduction			
Background	3a	D;E	Explain the healthcare context (including whether diagnostic or prognostic) and rationale for developing or evaluating the prediction model, including references to existing models
	3b	D;E	Describe the target population and the intended purpose of the prediction model in the context of the care pathway, including its intended users (eg, healthcare professionals, patients, public)
	3c	D;E	Describe any known health inequalities between sociodemographic groups
Objectives	4	D;E	Specify the study objectives, including whether the study describes the development or validation of a prediction model (or both)
Methods			
Data	5a	D;E	Describe the sources of data separately for the development and evaluation datasets (eg, randomised trial, cohort, routine care or registry data), the rationale for using these data, and representativeness of the data
	5b	D;E	Specify the dates of the collected participant data, including start and end of participant accrual; and, if applicable, end of follow-up
Participants	6a	D;E	Specify key elements of the study setting (eg, primary care, secondary care, general population) including the number and location of centres
	6b	D;E	Describe the eligibility criteria for study participants
	6c	D;E	Give details of any treatments received, and how they were handled during model development or evaluation, if relevant
Data preparation	7	D;E	Describe any data pre-processing and quality checking, including whether this was similar across relevant sociodemographic groups
Outcome	8a	D;E	Clearly define the outcome that is being predicted and the time horizon, including how and when assessed, the rationale for choosing this outcome, and whether the method of outcome assessment is consistent across sociodemographic groups
	8b	D;E	If outcome assessment requires subjective interpretation, describe the qualifications and demographic characteristics of the outcome assessors
	8c	D;E	Report any actions to blind assessment of the outcome to be predicted
Predictors	9a	D	Describe the choice of initial predictors (eg, literature, previous models, all available predictors) and any pre-selection of predictors before model building
	9b	D;E	Clearly define all predictors, including how and when they were measured (and any actions to blind assessment of predictors for the outcome and other predictors)
	9c	D;E	If predictor measurement requires subjective interpretation, describe the qualifications and demographic characteristics of the predictor assessors
Sample size	10	D;E	Explain how the study size was arrived at (separately for development and evaluation), and justify that the study size was sufficient to answer the research question. Include details of any sample size calculation
Missing data	11	D;E	Describe how missing data were handled. Provide reasons for omitting any data
Analytical methods	12a	D	Describe how the data were used (eg, for development and evaluation of model performance) in the analysis, including whether the data were partitioned, considering any sample size requirements
	12b	D	Depending on the type of model, describe how predictors were handled in the analyses (functional form, rescaling, transformation, or any standardisation)
	12c	D	Specify the type of model, rationale, all model building steps, including any hyperparameter tuning, and method for internal validation
	12d	D;E	Describe if and how any heterogeneity in estimates of model parameter values and model performance was handled and quantified across clusters (eg, hospitals, countries). See TRIPOD-Cluster for additional considerations†
	12e	D;E	Specify all measures and plots used (and their rationale) to evaluate model performance (eg, discrimination, calibration, clinical utility) and, if relevant, to compare multiple models
	12f	E	Describe any model updating (eg, recalibration) arising from the model evaluation, either overall or for particular sociodemographic groups or settings
	12g	E	For model evaluation, describe how the model predictions were calculated (eg, formula, code, object, application programming interface)

(Continued)

Table 2 | Continued

Section/topic	Item	Development/evaluation*	Checklist item
Class imbalance	13	D;E	If class imbalance methods were used, state why and how this was done, and any subsequent methods to recalibrate the model or the model predictions
Fairness	14	D;E	Describe any approaches that were used to address model fairness and their rationale
Model output	15	D	Specify the output of the prediction model (eg, probabilities, classification). Provide details and rationale for any classification and how the thresholds were identified
Training versus evaluation	16	D;E	Identify any differences between the development and evaluation data in healthcare setting, eligibility criteria, outcome, and predictors
Ethical approval	17	D;E	Name the institutional research board or ethics committee that approved the study and describe the participant informed consent or the ethics committee waiver of informed consent
Open science			
Funding	18a	D;E	Give the source of funding and the role of the funders for the present study
Conflicts of interest	18b	D;E	Declare any conflicts of interest and financial disclosures for all authors
Protocol	18c	D;E	Indicate where the study protocol can be accessed or state that a protocol was not prepared
Registration	18d	D;E	Provide registration information for the study, including register name and registration number, or state that the study was not registered
Data sharing	18e	D;E	Provide details of the availability of the study data
Code sharing	18f	D;E	Provide details of the availability of the analytical code§
Patient and public involvement			
Patient and public involvement	19	D;E	Provide details of any patient and public involvement during the design, conduct, reporting, interpretation, or dissemination of the study or state no involvement
Result			
Participants	20a	D;E	Describe the flow of participants through the study, including the number of participants with and without the outcome and, if applicable, a summary of the follow-up time. A diagram may be helpful
	20b	D;E	Report the characteristics overall and, where applicable, for each data source or setting, including the key dates, key predictors (including demographics), treatments received, sample size, number of outcome events, follow-up time, and amount of missing data. A table may be helpful. Report any differences across key demographic groups
	20c	E	For model evaluation, show a comparison with the development data of the distribution of important predictors (demographics, predictors, and outcome)
Model development	21	D;E	Specify the number of participants and outcome events in each analysis (eg, for model development, hyperparameter tuning, model evaluation)
Model specification	22	D	Provide details of the full prediction model (eg, formula, code, object, application programming interface) to allow predictions in new individuals and to enable third party evaluation and implementation, including any restrictions to access or reuse (eg, freely available, proprietary)¶
Model performance	23a	D;E	Report model performance estimates with confidence intervals, including for any key subgroups (eg, sociodemographic). Consider plots to aid presentation
	23b	D;E	If examined, report results of any heterogeneity in model performance across clusters. See TRIPOD-Cluster for additional details‡
Model updating	24	E	Report the results from any model updating, including the updated model and subsequent performance
Discussion			
Interpretation	25	D;E	Give an overall interpretation of the main results, including issues of fairness in the context of the objectives and previous studies
Limitations	26	D;E	Discuss any limitations of the study (such as a non-representative sample, sample size, overfitting, missing data) and their effects on any biases, statistical uncertainty, and generalisability
Usability of the model in the context of current care	27a	D	Describe how poor quality or unavailable input data (eg, predictor values) should be assessed and handled when implementing the prediction model
	27b	D	Specify whether users will be required to interact in the handling of the input data or use of the model, and what level of expertise is required of users
	27c	D;E	Discuss any next steps for future research, with a specific view to applicability and generalisability of the model

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*D=items relevant only to the development of a prediction model; E=items relating solely to the evaluation of a prediction model; D;E=items applicable to both the development and evaluation of a prediction model.

†Separately for all model building approaches.

‡TRIPOD-Cluster is a checklist of reporting recommendations for studies developing or validating models that explicitly account for clustering or explore heterogeneity in model performance (eg, at different hospitals or centres).^{19,20}

§Relates to the analysis code, for example, any data cleaning, feature engineering, model building, and evaluation.

¶Relates to the code to implement the model to get estimates of risk for a new individual.

27 main items about the title (item 1), abstract (item 2), introduction (items 3 and 4), methods (items 5-17), open science practises (item 18), patient and public involvement (item 19), results (items 20-24), and discussion (items 25-27). Some items included multiple subitems, totalling to 52 checklist subitems.

TRIPOD+AI covers studies that describe the development of a prediction model, the evaluation (validation) of prediction model performance, or both. Any items denoted D;E apply to all studies regardless

of whether they are developing a prediction model or evaluating the performance of a prediction model (table 2). Items in the checklist denoted D apply to studies that describe the development of a prediction model, while items denoted E apply to studies that evaluate the performance of a prediction model. For studies both developing and evaluating the performance of a prediction model, all checklist items apply.

A separate checklist for journal or conference abstracts of prediction model studies is included in

Box 2: Noteworthy changes and additions to TRIPOD 2015

- New checklist of reporting recommendations to cover prediction model studies using any regression or machine learning method (eg, random forests, deep learning), and harmonise nomenclature between regression and machine learning communities
- New TRIPOD+AI checklist supersedes the TRIPOD 2015 checklist, which should no longer be used
- Particular emphasis on fairness (box 1) to raise awareness and ensure that reports mention whether specific methods were used to deal with fairness. Aspects of fairness are embedded throughout the checklist
- Inclusion of TRIPOD+AI for Abstracts for guidance on reporting abstracts
- Modification of the model performance item recommending that authors evaluate model performance in key subgroups (eg, sociodemographic)
- Inclusion of a new item on patient and public involvement to raise awareness and prompt authors to provide details on any patient and public involvement during the design, conduct, reporting (and interpretation), and dissemination of the study
- Inclusion of an open science section with subitems on study protocols, registration, data sharing and code sharing

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TRIPOD+AI. This checklist updates the TRIPOD for Abstracts statement,¹⁸ reflecting new content and maintaining consistency with TRIPOD+AI (table 3).

The recommendations in TRIPOD+AI are for transparently reporting how prediction model research was conducted; it does not prescribe how to develop or evaluate a prediction model. The checklist is not a quality appraisal tool. Readers are referred to PROBAST^{90 91} and the forthcoming PROBAST+AI⁷⁹ to assess the quality and risk of bias of prediction models (<https://www.probast.org/>).

How to use TRIPOD+AI

The TRIPOD+AI checklist supersedes the TRIPOD 2015 checklist, which should no longer be used. For prediction model studies that have accounted

for clustering (eg, multiple hospitals, multiple datasets), authors should consult TRIPOD-Cluster for additional reporting recommendations.^{19 20} The 2015 explanation and elaboration document remains an important document to provide background and examples for most of the TRIPOD+AI reporting items¹⁷ (because many items have not changed or have been minimally changed), while we produce a detailed and updated document for TRIPOD+AI. We recommend using TRIPOD+AI early in the writing process to ensure that all key details are addressed and reported. An expanded checklist in a bullet point structure has been developed (supplementary table 1) to facilitate implementation of TRIPOD+AI by providing a brief rationale and guidance for each item in the checklist.

Although many of the items in the TRIPOD+AI checklist have a natural order and sequence in a report, some do not. We do not stipulate a structured format or dictate where each individual reporting recommendation should appear in a prediction model report or publication, because this order might also depend on journal formatting policies.

The recommendations contained within TRIPOD+AI are the minimum reporting recommendations, and authors may provide additional information. If journal word limits and restrictions on number of tables and figures in the main body of the manuscript complicate reporting, authors can report and reference some of the requested or additional information in supplementary material. If the information required is already reported in a publicly accessible study protocol, then referring to that document may suffice. If a particular checklist item cannot be discussed in the report because the information is unknown or irrelevant, then this should be acknowledged and clearly stated. Additional files and study materials not included in the supplementary material should be deposited in

Table 3 | Essential items to include for the reporting of prediction model studies in a journal or conference abstract (TRIPOD+AI for Abstracts*)

Section and item	Checklist item
Title	
1	Identify the study as developing or evaluating the performance of a multivariable prediction model, the target population, and the outcome to be predicted
Background	
2	Provide a brief explanation of the healthcare context and rationale for developing or evaluating the performance of all models
Objectives	
3	Specify the study objectives, including whether the study describes model development, evaluation, or both
Methods	
4	Describe the sources of data
5	Describe the eligibility criteria and setting where the data were collected
6	Specify the outcome to be predicted by the model, including time horizon of predictions in case of prognostic models
7	Specify the type of model, a summary of the model-building steps, and the method for internal validation ^t
8	Specify the measures used to assess model performance (eg, discrimination, calibration, clinical utility)
Results	
9	Report the number of participants and outcome events
10	Summarise the predictors in the final model ^t
11	Report model performance estimates (with confidence intervals)
Discussion	
12	Give an overall interpretation of the main results
Registration	
13	Give the registration number and name of the registry or repository

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*This checklist is based on the TRIPOD for Abstracts statement published in 2020,¹⁷ but has been revised and updated for consistency with the TRIPOD+AI statement.

^tRelevant only to studies describing the development of a prediction model.

general purpose (eg, Open Science Framework, Dryad, figshare) or institutional open access repositories that provide free access in perpetuity. Details of access to any additional files should be referenced and linked, for example, with a doi number in the main study report or publication.

We recommend that authors submit a completed checklist indicating the page or line where each requested item can be found, to help the editorial and peer review process. A template for the TRIPOD+AI checklist for filling out separately can be found in supplementary table 2 and is available to download from www.tripod-statement.org.

News, announcements, and information relating to TRIPOD+AI can be found on the TRIPOD website (www.tripod-statement.org) and on social media accounts such as X (formerly known as Twitter; @TRIPODStatement). The Enhancing the Quality and Transparency Of health Research (EQUATOR) Network (<https://www.equator-network.org/>) will also disseminate and promote the TRIPOD+AI statement. Translation of TRIPOD+AI into different languages is welcomed and encouraged, please contact the corresponding author. Translations should use the structured and predefined process that includes authors of the original publication and receives their approval. The TRIPOD website contains further details on translation (www.tripod-statement.org).

Discussion

TRIPOD+AI has been developed through an international multi-stakeholder consensus process. It provides minimum reporting recommendations for studies describing the development or evaluation (validation) of prediction models using any regression or machine learning methods. At the time of guideline development, foundation and large language models (such as ChatGPT) that are rapidly gaining momentum were not considered—the TRIPOD+AI guidance is primarily aimed at non-generative models. However, many of the principles are applicable for driving transparency in generative AI studies in health. Periodic updating of TRIPOD+AI will be needed to remain relevant and reflect advancements in AI and machine learning methods, for example, by explicitly looking at generative approaches.

TRIPOD+AI was developed by updating TRIPOD 2015, with recommendations informed by systematic reviews of the literature, a Delphi survey, and an online consensus meeting. Reporting TRIPOD+AI items can help users to understand and appraise the quality of the study methods, increasing transparency around the study findings, reducing overinterpretation of study findings, facilitating replication and reproducibility, and aiding implementation of the prediction model. The checklist items are minimum reporting recommendations, and authors will typically provide additional details on the data, study design, methods, analysis, results, and discussion.

TRIPOD+AI emphasises fairness issues throughout the checklist, which was lacking or not explicitly

stated in TRIPOD 2015.³³ Fairness in prediction model research is particularly important in healthcare, which has gained prominence with AI and machine learning methods being used to develop models to assist in decision making. Fairness in this context means that prediction models are designed and used in a way that does not adversely discriminate against any particular group of individuals and does not create or exacerbate (and ideally mitigates or reduces) existing inequalities in healthcare provision or patient outcomes.⁹² One important aspect of fairness is ensuring that the data used to develop or evaluate prediction models are representative and diverse, and that limitations of data bias are acknowledged, dealt with, and mitigated during model development. The STANDING Together initiative is in the process of developing standards for data diversity, inclusivity, and generalisability to tackle bias in AI health datasets.⁶²

Data should ideally include information from individuals of different ages, sexes/genders, and races/ethnicities, with different health conditions or comorbidities and from different geographical locations. These differences should be representative of the population in whom the prediction model is intended to be used. If the data used to develop the models do not adequately represent the full diversity of the intended use population, the resulting model might not perform as expected in those missing from the data, which should be clearly stated. If the data used to evaluate a model are not representative of the target population, then the estimated predictive accuracy in subgroups (eg, defined by relevant personal, social, or clinical attributes) could be biased and misleading.

While adequate representation of minoritised and underserved groups within datasets is one key element to achieving fairness goals, representation alone does not guarantee fairness.^{61 93} Therefore, TRIPOD+AI has embedded items on fairness throughout, including in the background (item 3c), methods (items 5a, 7, 8a, 8b, 9c, 12f, 14), results (items 20b, 23a), and discussion (items 25, 26).

Fairness in healthcare also means involving diverse stakeholders, including patients, the general public, and clinicians, in the development, evaluation, implementation, and deployment of a prediction model into the clinical pathway.⁹⁴ Involving a variety of perspectives will help to ensure that the prediction model is, in principle, designed to meet the needs of all individuals and is used in a way that is fair and equitable, promoting health equity. TRIPOD+AI includes item 19 on public and patient involvement to incentivise the integration of patient involvement in prediction model studies beyond a mere tick box exercise, to encourage and promote the principles of open science and engagement, and to ensure better clinical and public acceptability of the work.

TRIPOD+AI prominently features open science practices.³⁵ Open science practices are crucial for prediction model research in healthcare as they promote transparency, reproducibility, and collaboration between researchers.⁹⁵ By registering

research and making study materials such as protocols, data, code, and the prediction model open available, other researchers can verify the findings and evaluate model performance in new data to ensure that models are accurate, and evaluate models for safety. Open science practices also enable researchers to build on each other's work, leading to more efficient progress in healthcare. These practices can have a considerable impact on patient outcomes by improving the accuracy, integrity, and reliability of prediction models. If data are openly shared, clinicians and researchers can develop or evaluate models on larger and more diverse sets of patient data,⁹⁶ potentially leading to more accurate predictions and better informed decisions for patient healthcare. Therefore, TRIPOD+AI includes a section on open science, covering issues such as

funding declarations (item 18a), conflicts of interest (18b), protocol availability (18c), study registration (18d), data sharing (18e), and code sharing (18f).

We anticipate that the key users and beneficiaries of TRIPOD+AI will be researchers writing papers, journal editors and peer reviewers who evaluate research papers, and other stakeholders (eg, academic institutions, policy makers, funders, regulators, patients, study participants, and the broader public) who will benefit from the increased quality of prediction model research (table 4). The guideline is relevant for any reports related to clinical prediction model development and validation studies, including medical research articles and other areas where evidenced reports are needed, for example, to accompany software and tools.

Table 4 | Adherence to the TRIPOD+AI reporting guideline: potential benefits from stakeholders' actions

User/stakeholder	Proposed action	Potential benefits
Academic institutions	Promote or require adherence of TRIPOD+AI by investigators developing, evaluating, or implementing prediction models	Enhance a culture of transparency in the design, analysis, and reporting of prediction model research
	Provide training for early career researchers on the importance and benefits of transparent and complete reporting, including requiring doctoral students to write their thesis and manuscripts in accordance with the full TRIPOD+AI guideline	Improve the quality, accountability, reproducibility, replicability, and usefulness of produced research
Researchers	Adhere to TRIPOD+AI when writing studies for publication	Improved completeness and quality of reporting Increased awareness of the minimal detail required and expected when writing a prediction model publication Improved quality, accountability, reproducibility, replicability, and usefulness of produced research Improved reporting of details that facilitate independent evaluation of the model
Journal editors	Require and enforce authors to use TRIPOD+AI and submit a completed a checklist when writing the manuscript Recommend peer reviewers use TRIPOD+AI	Improved understanding of journal requirements and expectations for prediction model publications Increased efficiency of peer review resulting from improved author understanding of journal requirements for prediction model publications Improved quality, accountability, reproducibility, replicability, and usefulness of published research
Peer reviewers	Use TRIPOD+AI to evaluate completeness of reporting	Improve the efficiency and quality of peer review Facilitate and direct specific feedback to authors on where important details are missing
Funders	Recommend or mandate use of TRIPOD+AI by investigators when receiving a grant for prediction model research	Increase the usefulness of research outputs Reduce avoidable research waste due to incomplete reporting Ensure that funded research can be used by others
Patients, public, and study participants	Advocate use of TRIPOD+AI by authors, peer reviewers, journals, and funders	Improved trust in research findings Improved understanding of prediction model research Promote health equity considerations in research Align patient reported outcomes and patient experience with clinical research outcomes for precision medicine and personalised disease management
Systematic reviewers and meta-researchers	Use TRIPOD+AI to assess completeness of reporting Use TRIPOD+AI as an aid when assessing quality and risks of bias	Improved evaluation of study quality when used alongside risk of bias tools (eg, PROBAST) Improved availability of data needed for meta-analysis
Policy makers	Use or promote TRIPOD+AI to ensure research is transparently and completely reported	Ensure decisions to evaluate or implement a prediction model are based on complete and transparently reported information Add integrity for evidence based policy recommendations
Regulators	Clinical reviewers use TRIPOD+AI to assess completeness of clinical investigation reporting for "software as a medical device" regulatory submissions where the operating principle of the product is a prediction model	Align reported intended use with regulatory intended purpose Align medical device regulatory review and pivotal investigational reporting with best practice Encourage manufacturers to publish clinical investigation reports by encouraging one common standard
Technology and medical device manufacturers	Verify whether sufficient details about a model are available to enable development and manufacturing of technology and devices	Encourages manufacturers to publish clinical investigation reports by encouraging one common standard
Healthcare professionals	Verify whether sufficient details about a model are available before purchasing or using a model to support clinical use	Improved understanding of the target population of a model and the clinical decision it is intended to support Improved understanding of model predictions and awareness of limitations Improved trust in research findings

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We encourage editors and publishers to support adherence to TRIPOD+AI by referring to it in journals' instructions to authors, enforcing its use during the submission and peer review process, and making adherence to the recommendations an expectation. We also encourage funders to require that funding applications for prediction model studies include a plan to report their prediction model according to the TRIPOD+AI recommendations, thereby minimising research waste and ensuring value for money.

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and declare: support from the funding bodies listed above for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work. GSC is a National Institute for Health and Care Research (NIHR) senior investigator, the director of the UK EQUATOR Centre, editor-in-chief of *BMC Diagnostic and Prognostic Research*, and a statistics editor for *The BMJ*. KGMM is director of Health Innovation Netherlands and editor-in-chief of *BMC Diagnostic and Prognostic Research*. RDR is an NIHR senior investigator, a statistics editor for *The BMJ*, and receives royalties from textbooks *Prognosis Research in Healthcare* and *Individual Participant Data Meta-Analysis*. AKD is an NIHR senior investigator. EWL is the head of research at *The BMJ*. BG is a part time employee of HeartFlow and Kheiron Medical Technologies and holds stock options with both as part of the standard compensation package. SR receives royalties from Springer for the textbooks *Targeted Learning: Causal Inference for Observational and Experimental Data* and *Targeted Learning: Causal Inference for Complex Longitudinal Studies*. JCC receives honorariums as a current lay member on the UK NICE covid-19 expert panel and a citizen partner on the COVID-END Covid-19 Evidence Network to support decision making; was a lay member on the UK NIHR AI AWARD panel in 2020-22 and is a current lay member on the UK NHS England AAC Accelerated Access Collaborative NHS AI Laboratory Evaluation Advisory Group; is a patient fellow of the European Patients' Academy on Therapeutic Innovation and a EURORDIS rare disease alumni; reports grants from the UK National Institute for Health and Care Research, European Commission, UK Cell Gene Catapult, University College London, and University of East Anglia; reports patient speaker fees from MEDABLE, Reuters Pharma events, Patients as Partners Europe, and EIT Health Scandinavia; reports consultancy fees from Roche Global, Smith, the FutureScience Group and Springer Healthcare (scientific publishing), outside of the scope of the present work; and is a strategic board member of the UK Medical Research Council IASB Advanced Pain Discovery Platform initiative, Plymouth Institute of Health, and EU project Digipredict Edge AI-deployed Digital Twins for covid-19 Cardiovascular Disease. ALB is a paid consultant for Generate Biomedicines, Flagship Pioneering, Porter Health, FL97, Tessera, FL85; has an equity stake in Generate Biomedicines; and receives research funding support from Smith, National Heart, Lung, and Blood Institute, and National Institute of Diabetes and Digestive and Kidney Diseases. No other conflicts of interests with this specific work are declared.

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Supplementary table 1: TRIPOD+AI Expanded Checklist (Explanation & Elaboration Light)

Supplementary table 2: Fillable TRIPOD+AI checklist

Supplementary table 3: Aggregated TRIPOD+AI responses from Delphi round 1

Supplementary table 4: Aggregated TRIPOD+AI responses from Delphi round 2