BMJ Open is committed to open peer review. As part of this commitment we make the peer review history of every article we publish publicly available.

When an article is published we post the peer reviewers’ comments and the authors’ responses online. We also post the versions of the paper that were used during peer review. These are the versions that the peer review comments apply to.

The versions of the paper that follow are the versions that were submitted during the peer review process. They are not the versions of record or the final published versions. They should not be cited or distributed as the published version of this manuscript.

BMJ Open is an open access journal and the full, final, typeset and author-corrected version of record of the manuscript is available on our site with no access controls, subscription charges or pay-per-view fees (http://bmjopen.bmj.com).

If you have any questions on BMJ Open’s open peer review process please email info.bmjopen@bmj.com
A protocol for development of a reporting guideline (TRIPOD-AI) and risk of bias tool (PROBAST-AI) for diagnostic and prognostic prediction studies based on artificial intelligence

<table>
<thead>
<tr>
<th>Journal:</th>
<th>BMJ Open</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID</td>
<td>bmjopen-2020-048008</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Protocol</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>15-Dec-2020</td>
</tr>
<tr>
<td>Complete List of Authors:</td>
<td>Collins, Gary; University of Oxford, Centre for Statistics in Medicine Dhiman, Paula; University of Nottingham, School of Medicine Andaur Navarro, Constanza L.; Julius Center for Health Sciences and Primary Care, Epidemiology Ma, Ji; University of Oxford Hooft, Lotty; University Medical Center Utrecht, University of Utrecht, Cochrane Netherlands Reitsma, Johannes; University Medical Center Utrecht, Julius Center for Health Sciences and Primary Care Logullo, Patricia; University of Oxford Beam, Andrew; Harvard Medical School, Peng, Lily; Google Inc Van Calster, Ben; KU Leuven, Department of Development and Regeneration van Smeden, Maarten; Universiteit Leiden, Riley, Richard; Keele University Moons, Karel; Julius Center for Health Sciences and Primary Care, Epidemiology</td>
</tr>
<tr>
<td>Keywords:</td>
<td>STATISTICS &amp; RESEARCH METHODS, EPIDEMIOLOGY, GENERAL MEDICINE (see Internal Medicine)</td>
</tr>
</tbody>
</table>
I, the Submitting Author has the right to grant and does grant on behalf of all authors of the Work (as defined in the below author licence), an exclusive licence and/or a non-exclusive licence for contributions from authors who are: i) UK Crown employees; ii) where BMJ has agreed a CC-BY licence shall apply, and/or iii) in accordance with the terms applicable for US Federal Government officers or employees acting as part of their official duties; on a worldwide, perpetual, irrevocable, royalty-free basis to BMJ Publishing Group Ltd (“BMJ”) its licensees and where the relevant Journal is co-owned by BMJ to the co-owners of the Journal, to publish the Work in this journal and any other BMJ products and to exploit all rights, as set out in our licence.

The Submitting Author accepts and understands that any supply made under these terms is made by BMJ to the Submitting Author unless you are acting as an employee on behalf of your employer or a postgraduate student of an affiliated institution which is paying any applicable article publishing charge (“APC”) for Open Access articles. Where the Submitting Author wishes to make the Work available on an Open Access basis (and intends to pay the relevant APC), the terms of reuse of such Open Access shall be governed by a Creative Commons licence – details of these licences and which Creative Commons licence will apply to this Work are set out in our licence referred to above.

Other than as permitted in any relevant BMJ Author’s Self Archiving Policies, I confirm this Work has not been accepted for publication elsewhere, is not being considered for publication elsewhere and does not duplicate material already published. I confirm all authors consent to publication of this Work and authorise the granting of this licence.
A protocol for development of a reporting guideline (TRIPOD-AI) and risk of bias tool (PROBAST-AI) for diagnostic and prognostic prediction studies based on artificial intelligence

Gary S. Collins1,2 *, Paula Dhiman1,2, Constanza L. Andaur Navarro3,4, Jie Ma1, Lotty Hooft3,4, Johannes B. Reitsma3, Patricia Logullo1, Andrew L. Beam5,6, Lily Peng7, Ben Van Calster8,9,10, Maarten van Smeden3, Richard D. Riley11, Karel G.M. Moons3,4 *

1 Centre for Statistics in Medicine, Nuffield Department of Orthopaedics, Rheumatology & Musculoskeletal Sciences, University of Oxford, Oxford, OX3 7LD, United Kingdom
2 NIHR Oxford Biomedical Research Centre, John Radcliffe Hospital, Oxford, United Kingdom NIHR Oxford Biomedical Research Centre, John Radcliffe Hospital, Oxford, United Kingdom
3 Julius Center for Health Sciences & Primary Care, and Cochrane Netherlands, University Medical Center Utrecht, Utrecht University, Utrecht, The Netherlands
4 Cochrane Netherlands, University Medical Center Utrecht, Utrecht University, The Netherlands
5 Department of Epidemiology, Harvard T.H. Chan School of Public Health, Boston, Massachusetts, United States
6 Department of Biomedical Informatics, Harvard Medical School, Boston, Massachusetts, United States
7 Google Health, 3400 Hillview Ave, Palo Alto, CA 94304, United States
8 KU Leuven, Department of Development and Regeneration, Leuven, Belgium
9 Department of Biomedical Data Sciences, Leiden University Medical Centre (LUMC), Leiden, the Netherlands
10 EPI-centre, KU Leuven, Leuven, Belgium
11 Centre for Prognosis Research, School of Medicine, Keele University, Staffordshire, ST5 5BG, United Kingdom.

* Both authors contributed equally

Address for correspondence:

Professor Gary S Collins
Centre for Statistics in Medicine
Nuffield Department of Orthopaedics, Rheumatology & Musculoskeletal Sciences
University of Oxford
Oxford, OX3 7LD
United Kingdom.
Email: gary.collins@csm.ox.ac.uk

Twitter: @GSCollins, @CarlMoons
ORCID: 0000-0002-2772-2316
Abstract

Introduction: The Transparent Reporting of a multivariable prediction model of Individual Prognosis or Diagnosis (TRIPOD) statement and the Prediction model Risk Of Bias ASsessment Tool (PROBAST) were both published to improve the reporting and critical appraisal of prediction model studies for diagnosis and prognosis. This paper describes the processes and methods that will be used to develop an extension to the TRIPOD statement (TRIPOD-AI) and the PROBAST (PROBAST-AI) tool for prediction model studies that applied machine learning techniques.

Methods and Analysis: TRIPOD-AI and PROBAST-AI will be developed following published guidance from the EQUATOR Network, and will comprise five stages. Stage one will comprise two systematic reviews (across all medical fields and specifically in oncology) to examine the quality of reporting in published machine-learning based prediction model studies. In stage two we will consult a diverse group of key stakeholders using a Delphi process to identify items to be considered for inclusion in TRIPOD-AI and PROBAST-AI. Stage three will be a virtual consensus meeting to consolidate and prioritise key items to be included in TRIPOD-AI and PROBAST-AI. Stage four will involve developing the TRIPOD-AI checklist and the PROBAST-AI tool, and writing the accompanying explanation and elaboration papers. In the final stage, stage 5, we will disseminate TRIPOD-AI and PROBAST-AI via journals, conferences, blogs, websites (including TRIPOD, PROBAST and EQUATOR Network), and social media. TRIPOD-AI will provide researchers working on prediction model studies based on machine-learning with a reporting guideline that can help them report key details that readers need to evaluate the study quality and interpret its findings, potentially reducing research waste. We anticipate PROBAST-AI will help researchers, clinicians, systematic reviewers and policymakers critically appraise the design, conduct and analysis of machine learning based prediction model studies, with a robust standardised tool for bias evaluation.

Ethics and Dissemination
Ethical approval has been granted by the Central University Research Ethics Committee (CUREC), University of Oxford on 10-December-2020 (R73034/RE001). Findings from this study will be disseminated through peer review publications.

Keywords: prediction, artificial intelligence, machine learning, reporting guideline, risk of bias, TRIPOD, PROBAST
Article Summary

Strengths and limitations of this study

- The reporting of clinical prediction models using artificial intelligence is poor.
- There are no guidelines for the reporting or risk of bias assessment of clinical prediction models using artificial intelligence.
- The strengths of this study is that it follows published guidance from the EQUATOR Network for developing reporting guidelines. Expert opinion and consensus will be obtained from multiple stakeholders (statisticians, clinician scientists, epidemiologists, computer scientists, funders, healthcare policy makers, patients, and industry leaders).
- Limitations of this study potentially include representativeness of the Delphi panel as those may differ from those who decline to participate.
Background

Models that predict clinical outcomes are abundant in the medical literature and are broadly
categorised as those that estimate the probability of the presence of a particular outcome (diagnostic)
or whether a particular outcome (e.g. event) will occur in the future (prognostic) [1]. Traditionally
these models (herein referred to as prediction model) have been developed using regression-based
methods, typically logistic regression for short-term outcomes and Cox regression for longer-term
outcomes [2]. Numerous reviews have observed that studies describing the development and
validation (including updating) of a prediction model often fail to report key information to help
readers judge the methods and have a complete, transparent and clear picture of the model’s
predictive accuracy and other relevant details such as the target population and the content of the
model itself [3–6]. The absence of full and comprehensive reporting limits the usability of the findings
of these studies, e.g., in subsequent validation studies, evidence synthesis studies, or in daily practice,
and therefore contribute to research waste [7]. In response to this, in 2015, the Transparent Reporting
of a multivariable prediction model for Individual Prognosis Or Diagnosis (TRIPOD) Statement was
published [1, 8]. The TRIPOD Statement is a checklist of 22-items that authors should report with
sufficient detail and clarity to inform how the study was carried out.

Since the publication of the TRIPOD Statement, artificial intelligence (AI) and in particular machine
learning, approaches to clinical prediction have evolved and grown in popularity with the number of
AI and machine learning publications rapidly rising [9–14]. This is evident within a recent review of
COVID-19 related prediction models, where 57 (out of 107 included studies) used machine learning
methods to develop their model [15].

Machine learning, a branch of artificial intelligence, can be broadly described as data analytical
methods that learn from data without being explicitly programmed, with patterns identified based on
the data itself. They are often described as having flexibility to capture complex associations
particularly in large and unstructured data and complexity in modelling. Whilst the vast majority of
the items in the TRIPOD Statement are relevant to machine learning based prediction model studies,
there are some unique challenges with machine learning that are not captured. Due to their
complexity, these prediction models are typically considered to be ‘black box’, unlike say regression-
based models where the full model can be transparently presented (e.g., as an equation containing
all the regression coefficients). Also, whilst many machine learning methods have origins in the
statistical literature, two (overlapping) prediction model cultures have emerged as those from a
statistical/epidemiological background and those from the computer science/data sciences. Although
there is clear overlap, different approaches to model development, validation and updating have appeared, and different and sometimes conflicting terminology have arisen.

Due to the relative novelty of applying machine learning methods to clinical prediction modelling, there is little information on the quality of reporting of these studies. However, the few reviews that have examined the completeness of reporting of have concluded that reporting is poor [16, 17]. In response to these concerns, guidance is required to help authors fully describe their prediction model study when machine learning methods were used. Therefore the TRIPOD group initiated a large international project to develop a consensus based extension of TRIPOD with specific focus on reporting of studies that undertake the development, validation or updating of a diagnostic or prognostic prediction model, using machine learning techniques – herein referred to as TRIPOD-AI [18]. The TRIPOD-AI extension, comprising a checklist and an accompanying elaboration and explanation document will provide researchers, authors, reviewers, editors, users, and other stakeholders of machine learning based prediction model studies, with guidance on the minimal set of items to report, with detailed examples of good reporting for each item.

Complete reporting allows studies to be understood, replicated and used. However, critical appraisal and of the quality of study method is a crucial component of evidence-based medicine as well. Critical appraisal and assessing the quality of studies is a crucial component of evidence-based medicine. In 2019, the Prediction model Risk Of Bias ASsessment Tool (PROBAST) was published [19, 20] to help a variety of stakeholders including, for example, systematic reviewers, researchers, journal editors, manuscript reviewers and policy makers involved in clinical guideline development, critically appraise the study design, conduct and analysis of prediction model studies. PROBAST comprises four domains (participants, predictors, outcome and analysis) and contains 20 signalling questions to facilitate risk of bias assessment. Clearly risk of bias assessment and reporting are intrinsically linked, in that judging risk of bias is predicated on what has been reported in the primary study. Whilst in principle PROBAST is relevant for prediction model studies using machine learning, different approaches to model development and validation, and terminology have appeared, and the ability to critically appraise these studies is crucial before they are implemented [21, 22]. Therefore, in parallel with the development of TRIPOD-AI, we will also develop PROBAST-AI, a tool to assess risk of bias in machine learning based multivariable prediction model studies.
Focus of TRIPOD-AI and PROBAST-AI

The focus of both TRIPOD-AI and PROBAST-AI is on reports of research or endeavours in which a multivariable prediction model is being developed (or updated), or validated (tested) using any machine learning technique. Conforming to the original TRIPOD and PROBAST publications, a multivariable prediction model is defined as any combination or equation of two or more predictors that is to be used for individualized predictions to estimate an individual’s probability of having (diagnosis) or developing (prognosis) a particular health outcome or state. Predictors may have any form and emerge from patient history, physical examination, diagnostic, prognostic or monitoring tests and from undergone treatments. Outcomes may also have any form (dichotomous, categorical, continuous) and of any kind, such as, a particular condition or disorder being present or absent (diagnostic outcome or classification), short-term prognosis outcomes (e.g. hospital mortality, or postoperative complications), and long-term prognostic outcomes such as one year occurrence of treatment complications, five-year occurrence of metastases, or life-long survival).

As per the original publications, TRIPOD-AI and PROBAST-AI will also address prediction model studies from all medical care settings (public health, primary, secondary, tertiary and nursing home care) and all corresponding target populations (healthy individuals, suspected and diseased individuals).

TRIPOD-AI and PROBAST-AI are not meant to address:

- comparative studies that quantify the impact of using a prediction model as compared to not using the model [23];
- so-called predictor finding studies (also known as risk or prognostic factor studies) where multivariable machine learning techniques are used to identify (usually from a wider set of potential predictors) those predictors that are associated with an outcome, but not to develop a model that can be used for individualized predictions in new individuals;
- single medical test studies that use machine learning or AI techniques aimed to read, for example, CT or MRI, images to find which image-parameters are best associated with an outcome (such studies fall under the remit of STARD-AI [24]). If these image-parameters are included as predictors in a multivariable model combined with other predictors, TRIPOD-AI and PROBAST-AI may be useful.

Methods/design

Both TRIPOD-AI and PROBAST-AI will be developed following published guidance from the EQUATOR Network [25]. We will develop the guideline in five stages: (i) systematic reviews to establish the
quality of current reporting, (ii) Delphi exercise, (iii) consensus meeting, (iv) development of the
guidance statement, and (v) guideline dissemination. We have registered our intent to develop the
TRIPOD extension for AI on the EQUATOR Network website (www.equator-network.org), the TRIPOD
website (www.tripod-statement.org) and recently announced it in the Lancet [18], whilst the
PROBAST-AI development has been announced on the PROBAST website (www.probast.org).

TRIPOD-AI/PROBAST-AI working group
The TRIPOD/PROBAST Working Group will include: (1) an Executive Committee (2) an Advisory and
Working Group; and (3) a large international Delphi Panel.

The TRIPOD-AI/PROBAST-AI Executive Committee will be responsible for the leadership and
coordination of all the processes involved in the development and dissemination of the TRIPOD-AI
guideline. The Executive Committee consists of the two lead authors of the TRIPOD reporting guideline
and the PROBAST tool, and also prediction model experts and researchers from the machine learning
community. Key stakeholders for stage 2 (Delphi survey) will be identified and approached to
participate and a subset of these key stakeholders (the Advisory Group) will participate in stage 3
(consensus meeting).

Here the term key stakeholder refers to a cross-sector participant (both industry and public sector)
who falls into at least one of the following categories:

1. researchers who have used machine learning in the context of clinical prediction, have clear
   knowledge and expertise in using machine learning or developed machine learning methods.
   These include applied (bio)medical investigators, statisticians, epidemiologists, and data
   scientists;

2. assessors and approvers of artificial intelligence or machine learning model, such as regulatory
   assessors and ethics committee members;

3. beneficiaries or users of the resultant TRIPOD-AI guidance and PROBAST-AI tool such as
   journal editors and journal reviewers;

4. commissioners of research grants, such as funders;
5. consumers of research results such as healthcare providers and patients and citizens.

Stage 1 – Systematic review of current reporting

Two parallel systematic reviews are ongoing to evaluate the quality of current reporting in published studies developing, validating or updating machine learning based prediction models in the medical domain. Both systematic reviews will assess adherence of the reporting against the original TRIPOD Statement [1, 8], using the TRIPOD adherence checklist [26]. The reviews will also examine the methodological conduct of the primary studies, including a risk of bias assessment using the recently issued risk of bias tool (quality appraisal) for diagnostic and prognostic prediction model studies (PROBAST) [19, 20], and will draw out specific issues, currently not covered by TRIPOD and PROBAST relating to machine learning. The protocols for the two systematic reviews have been registered with the International Prospective Register of Systematic Reviews (PROSPERO IDs CRD42019140361 and CRD42019161764). One review (CRD42019161764) will examine the quality of reporting of machine learning based prediction model studies across all medical fields (between January 2018 to December 2019), whilst the other review (CRD42019140361) will focus on the quality of reporting of machine learning based prediction model studies published in oncology (between January 2019 to September 2019).

Undertaking these reviews serves two purposes: (1) to understand the completeness of current reporting of machine learning based prediction model studies in the medical literature, and (2) to identify unique reporting items for consideration for TRIPOD extension, and unique risk of bias or quality items for PROBAST extension. The data collection for this phase is underway. The reviews will evaluate the current completeness of reporting and the quality of the research and identify additional reporting and quality items to be considered for TRIPOD-AI and PROBAST-AI.

These two reviews will evaluate the current completeness of reporting and the quality of the research. Together with other evidence [3, 4, 16, 17, 27] from existing methodological guidance papers, they will provide important information on the transparency and quality of reporting. Using the original TRIPOD and PROBAST checklists as starting points, the Executive Committee will identify in the literature the preliminary items to consider in Stage 2 (the Delphi study) and therefore inclusion in the eventual TRIPOD-AI checklist and PROBAST-AI tool.
Stage 2 – Delphi exercise

We will perform an extensive Delphi survey among a large international network of relevant stakeholders, with a maximum of three rounds, to help decide on items that could be modified, added to, or removed from the TRIPOD 2015 checklist to form the TRIPOD-AI checklist, and subsequently the PROBAST-AI checklist.

Design

The Delphi process will comprise of a series of rounds where panellists will independently and anonymously evaluate and achieve consensus on the inclusion or exclusion of the proposed reporting and quality items – in addition to suggesting additional items. The process will be repeated for a maximum of three rounds. Following each round, participants will be provided with structured feedback of the previous round to help reconcile individual opinions and achieve group consensus. Items achieving a high level of agreement (≥70%) will be taken forward to the consensus meeting (stage 3).

Selection of potential items

The list of items for TRIPOD-AI (and PROBAST-AI) will be collated by the Executive Committee, including the results of the two systematic reviews, any other available studies on methodology or reporting of machine learning based prediction models, and expert recommendations from the Delphi panellists. Relevant methodological guidance or methodological papers will be retrieved to identify additional candidate reporting and quality items for machine learning based prediction model studies. Pre-selection involves dividing items into those to further consider, those that can be provided as optional guidance (to be outlined in an Explanation and Elaboration accompanying document), or those not to consider for potential inclusion. Delphi participants will have the opportunity to view and provide feedback in each round, and also to suggest new items.

Recruitment process and participants

Delphi participants will be identified through professional networks of the Executive Committee, participation in the Delphi exercise of the original TRIPOD guideline (and TRIPOD for Abstracts and TRIPOD Cluster Delphi surveys), original PROBAST Delphi exercise, via self-response to the Lancet 2019 paper where TRIPOD-AI was announced [18], and responses to social media announcements of TRIPOD-AI (e.g., Twitter).
We will invite international participants with diverse roles (e.g., researchers, healthcare professionals, journal editors, funders, policy makers, healthcare regulators, end users of prediction models) from a range of settings (e.g., universities, hospitals, primary care, biomedical journals, non-profit organisations, and for-profit organisations). Participants will be invited via personalized email that will describe the TRIPOD-AI extension and PROBAST-AI tool development, and explain the objective, process, and timelines of the Delphi exercise. We plan to invite at least 100 participants to the Delphi survey. In all rounds, the survey will remain open for three weeks, with a reminder email sent one week after the initial invitation.

Informed consent from participants will be obtained using an online consent form and participants can withdraw at any time. Individuals who indicate that they wish to opt out of the survey will be removed from subsequent invitations. Participants will not know the identities of other individuals in the Delphi panel, nor will they know the specific answers that any individual provides.

**Participants**

Participants will not know the identities of other individuals in the Delphi panel, nor will they know the specific answers that any individual provides. In all rounds, the survey will remain open for three weeks, with a reminder email sent one week after the initial invitation. Individuals who indicate that they wish to opt out of the survey will be removed from subsequent invitations.

**Procedure for selection of items**

We plan to ask participants to consider the following guiding principles when reviewing existing, new or modified items for inclusion: 1) reporting of the item should facilitate reproducibility of the study (i.e. users should be able to recreate the findings based on the information reported); 2) reporting of the item facilitates assessment of the quality and risk of bias in and applicability of the machine learning study findings, to enhance their uptake and use in subsequent studies, systematic reviews and daily practice; 3) item is likely relevant to nearly all prediction model studies; 4) the set of items represent the minimum that should be reported in all machine learning studies developing, validating or updating a diagnostic or prognostic prediction model.

**Round 1**

Participants will be asked to rate on a 5-point Likert scale, the extent to which they agree with the inclusion of each checklist item in the TRIPOD-AI extension and PROBAST-AI tool (1=strongly disagree, 2=somewhat disagree, 3=I don’t know, 4=somewhat agree, 5=strongly agree). A free-text box will be
provided for general comments on each item (to justify their decision or suggest wording changes), and a free-text box will be provided at the end of the survey to suggest additional checklist items or provide general comments on the checklist. The survey will be pilot-tested for usability and clarity to a small number of individuals familiar with prediction models or machine learning but not involved in the TRIPOD-AI guideline extension or PROBAST-AI tool, and revised accordingly based on their feedback.

**Round 2**
The same participants involved in round 1 will be invited to participate in round 2. Participants will be provided with their first-round responses on each item, an anonymised summary of the group ratings and anonymised comments to justify ratings. Using the same format as round 1, participants will be presented with each item, including any new items suggested during round 1, and again express the extent to which they agree with the inclusion of the item in the TRIPOD-AI checklist or PROBAST-AI tool, considering the structured feedback to inform their responses. Participants who were invited to participate in round 1, but who did not respond will be invited to participate in round 2, and will be presented with an anonymised summary of the group ratings. Items that reached a high-level of agreement (scoring 4 or 5) in round 1 (≥ 70%) will be presented for information purposes only, with no voting on these items, though a free-text box will be provided for any comments. A third Delphi round will be used if deemed necessary by the Executive Committee.

**Results from the Delphi survey**
Item scores will be summarised for the entire panel as a whole, as appropriate (e.g., frequency and proportions across the rating categories) accompanied by a narrative summary of findings, comments, and suggestions. Results from both rounds of the survey will be discussed by the Executive Committee. For items where there was no consensus following the second Delphi round will be discussed by the Executive Committee, and will be considered for discussion at the subsequent consensus meeting.

**Stage 3 – Consensus meeting**
Two virtual consensus meetings (separately for TRIPOD-AI and PROBAST-AI), both spread over 2 days, will be held with the objective of discussing the results from the Delphi exercise and finalising items to be included in the reporting guideline and risk of bias tool. The composition of the consensus group will reflect the diversity of the key stakeholders addressed above. Key experts participating in the Delphi exercise will be considered to participate in the consensus meeting. We will also consider...
inviting experts who did not contribute to the Delphi to participate in the consensus. A total of around 25-30 international participants are expected to contribute to the virtual consensus meeting.

**Procedure**

The agenda and any material (e.g., results from the systematic reviews and Delphi) for the consensus meeting will be prepared by the Executive Committee and will be shared with attendees in advance. Members of the Executive Committee will facilitate a structured discussion on the rationale behind each item identified in the Delphi exercise. Consensus meeting participants will then be given the opportunity to discuss each item (reporting item for TRIPOD-AI and signalling question for PROBAST-AI), and vote on each item. The decision to retain an item in the TRIPOD-AI and PROBAST-AI will be based on achieving at least 70% support from the consensus meeting participants. The group will agree on the draft list of reporting items for the final TRIPOD-AI extension and PROBAST-AI tool. Specific item wording will not be discussed during the meeting, though participants can suggest and the group to agree on general intent and meaning of the item. Plans for dissemination will be discussed at the end of the consensus meeting.

**Pilot testing**

We will invite authors of machine learning prediction model studies in the medical domain, doctoral students undertaking prediction model, machine learning courses or workshops, and peer reviewers and editors of journals who frequently publish such prediction model studies, to pilot the use of a draft version of the TRIPOD-AI checklist and PROBAST-AI tool. We will ask those who pilot the checklist and tool whether the wording of items is ambiguous or difficult to interpret.

**Stage 4 – Development of the draft TRIPOD-AI statement, PROBAST-AI and Explanation & Elaboration documents**

The Executive Committee will lead the development of the TRIPOD-AI reporting guidance and PROBAST-AI signalling questions based on the agreed list of items from the consensus meeting (stage 3). The Executive Committee will invite a subset of members from the consensus meeting (to form a writing group) to help draft the Explanation & Elaboration paper.

The Executive Committee will reserve the right to update (that is remove or add) additional items to the TRIPOD-AI checklist during the development of the TRIPOD-AI statement, if and as necessary (as a result of the pilot testing).
For each of the TRIPOD-AI extension and the PROBAST-AI risk of bias tool, two manuscripts will be
developed; (1) the Statement paper, presenting the checklist/tool and describing the process of how
it was developed and (2) an Explanation & Elaboration paper. The Explanation & Elaboration papers
will outline the rationale of the reporting items (TRIPOD-AI) and signalling questions (PROBAST-AI),
examples of good reporting (TRIPOD-AI) and examples of how to use PROBAST-AI. Drafts of the papers
will be circulated to all participants of the consensus meeting for their comments.

Stage 5 – Guideline dissemination
The dissemination strategy will be informed by discussions at the consensus meeting. We will aim to
seek simultaneous publication in key journals to target different readerships. To increase visibility and
aid uptake, the TRIPOD-AI checklist and PROBAST-AI tool will be published open access, and made
available on the TRIPOD website along with other TRIPOD extensions (www.tripod-statement.org),
and on the PROBAST website (www.probast.org) respectively, as well on the PROGRESS website
(www.prognosisresearch.com). The TRIPOD-AI extension will be indexed on the EQUATOR website
(www.equator-network.org). Social media will be used to help disseminate the extension. The
Executive Committee will (and consensus participants will be encouraged to) publicise the TRIPOD-AI
statement and PROBAST-AI tool at key conferences and courses.

Publication plan
It is envisaged that the following publications will arise from the TRIPOD-AI and PROBAST-AI initiative:

- Publication 1: Study protocol
- Publication 2: Systematic review protocol (with registration on PROSPERO)
- Publication 3 & 4: Systematic reviews
- Publication 5 & 6: TRIPOD-AI statement and the Explanation and Elaboration paper
- Publication 7 & 8: PROBAST-AI tool and the Explanation and Elaboration paper

Conclusion
The number of prediction model studies using machine learning methods is rapidly increasing,
including developed, validated or updated prediction models. Ensuring that key details are reported
is important so that readers can evaluate the study quality, and interpret its findings including the
developed, validated or updated prediction model to enhance their uptake in subsequent research
(e.g. validation studies), evidence synthesis projects (e.g. systematic reviews of prediction models),
and in daily practice by healthcare professionals, patients or citizens. We anticipate that TRIPOD-AI
will help authors transparently report their study and help reviewers, editors, policy makers and end-users understand the methods and findings, and thereby reduce research waste. Similarly, we anticipate PROBAST-AI will help researchers, clinicians, systematic reviewers and policy-makers critically appraise the design, conduct and analysis of machine learning based prediction model studies.

**Patient and Public Involvement**

No patients or members of the public were involved in the development of this protocol. Patient and public involvement is planned for the virtual consensus meeting.

**Ethics**

Ethical approval has been granted by the Central University Research Ethics Committee (CUREC), University of Oxford on 10-December-2020 (R73034/RE001).

**Contributor statement**

Gary S. Collins, Paula Dhiman, Constanza L. Andaur Navarro, Jie Ma, Lotty Hooft, Johannes B. Reitsma, Patricia Logullo, Andrew L. Beam, Lily Peng, Ben Van Calster, Maarten van Smeden, Richard D. Riley, Karel G.M. Moons were involved in the planning and design of the study. Gary Collins drafted the manuscript with all authors contributing to the writing.

**Competing interests**

The authors declare no competing interests.

**Data sharing statement**

There is no data to share.

**Funding**

This research was supported by Health Data Research UK, an initiative funded by UKResearch and Innovation, Department of Health and Social Care (England) and the devolved administrations, and leading medical research charities, Cancer Research UK programme grant (C49297/A27294), the NIHR Biomedical Research Centre, Oxford, and the Netherlands Organisation for Scientific Research. The funders have no role in the development of this protocol and will not have a role in the data collection, analyses, interpretation of the data, and publication of the findings.

**References**


# A protocol for development of a reporting guideline (TRIPOD-AI) and risk of bias tool (PROBAST-AI) for diagnostic and prognostic prediction studies based on artificial intelligence

<table>
<thead>
<tr>
<th>Journal:</th>
<th>BMJ Open</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manuscript ID</td>
<td>bmjopen-2020-048008.R1</td>
</tr>
<tr>
<td>Article Type:</td>
<td>Protocol</td>
</tr>
<tr>
<td>Date Submitted by the Author:</td>
<td>14-Apr-2021</td>
</tr>
<tr>
<td>Complete List of Authors:</td>
<td>Collins, Gary; Oxford University, Centre for Statistics in Medicine Dhiman, Paula; Oxford University, Nuffield Department of Orthopaedics, Rheumatology &amp; Musculoskeletal Sciences Andaur Navarro, Constanza L.; Julius Center for Health Sciences and Primary Care, Epidemiology Ma, Ji; University of Oxford Hooft, Lotty; Julius Center for Health Sciences and Primary Care, Cochrane Netherlands Reitsma, Johannes; Julius Center for Health Sciences and Primary Care Logullo, Patricia; University of Oxford Beam, Andrew; Harvard Medical School, Peng, Lily; Google Inc Van Calster, Ben; KU Leuven, Department of Development and Regeneration van Smeden , Maarten; Universiteit Leiden, Riley, Richard; Keele University Moons, Karel; Julius Center for Health Sciences and Primary Care, Epidemiology</td>
</tr>
<tr>
<td>&lt;b&gt;Primary Subject Heading&lt;/b&gt;:</td>
<td>Medical publishing and peer review</td>
</tr>
<tr>
<td>Secondary Subject Heading:</td>
<td>Epidemiology</td>
</tr>
<tr>
<td>Keywords:</td>
<td>STATISTICS &amp; RESEARCH METHODS, EPIDEMIOLOGY, GENERAL MEDICINE (see Internal Medicine)</td>
</tr>
</tbody>
</table>
I, the Submitting Author has the right to grant and does grant on behalf of all authors of the Work (as defined in the below author licence), an exclusive licence and/or a non-exclusive licence for contributions from authors who are: i) UK Crown employees; ii) where BMJ has agreed a CC-BY licence shall apply, and/or iii) in accordance with the terms applicable for US Federal Government officers or employees acting as part of their official duties; on a worldwide, perpetual, irrevocable, royalty-free basis to BMJ Publishing Group Ltd (“BMJ”) its licensees and where the relevant Journal is co-owned by BMJ to the co-owners of the Journal, to publish the Work in this journal and any other BMJ products and to exploit all rights, as set out in our licence.

The Submitting Author accepts and understands that any supply made under these terms is made by BMJ to the Submitting Author unless you are acting as an employee on behalf of your employer or a postgraduate student of an affiliated institution which is paying any applicable article publishing charge (“APC”) for Open Access articles. Where the Submitting Author wishes to make the Work available on an Open Access basis (and intends to pay the relevant APC), the terms of reuse of such Open Access shall be governed by a Creative Commons licence – details of these licences and which Creative Commons licence will apply to this Work are set out in our licence referred to above.

Other than as permitted in any relevant BMJ Author’s Self Archiving Policies, I confirm this Work has not been accepted for publication elsewhere, is not being considered for publication elsewhere and does not duplicate material already published. I confirm all authors consent to publication of this Work and authorise the granting of this licence.
A protocol for development of a reporting guideline (TRIPOD-AI) and risk of bias tool (PROBAST-AI) for diagnostic and prognostic prediction studies based on artificial intelligence

Gary S. Collins¹,² *, Paula Dhiman¹,², Constanza L. Andaur Navarro³,⁴, Jie Ma¹, Lotty Hooft³,⁴, Johannes B. Reitsma³, Patricia Logullo¹, Andrew L. Beam⁵,⁶, Lily Peng⁷, Ben Van Calster⁸,⁹,¹⁰, Maarten van Smeden³, Richard D. Riley¹¹, Karel G.M. Moons³,⁴ *

¹ Centre for Statistics in Medicine, Nuffield Department of Orthopaedics, Rheumatology & Musculoskeletal Sciences, University of Oxford, Oxford, OX3 7LD, United Kingdom
² NIHR Oxford Biomedical Research Centre, John Radcliffe Hospital, Oxford, United Kingdom NIHR Oxford Biomedical Research Centre, John Radcliffe Hospital, Oxford, United Kingdom
³ Julius Center for Health Sciences & Primary Care, and Cochrane Netherlands, University Medical Center Utrecht, Utrecht, The Netherlands
⁴ Cochrane Netherlands, University Medical Center Utrecht, Utrecht University, The Netherlands
⁵ Department of Epidemiology, Harvard T.H. Chan School of Public Health, Boston, Massachusetts, United States
⁶ Department of Biomedical Informatics, Harvard Medical School, Boston, Massachusetts, United States
⁷ Google Health, 3400 Hillview Ave, Palo Alto, CA 94304, United States
⁸ KU Leuven, Department of Development and Regeneration, Leuven, Belgium
⁹ Department of Biomedical Data Sciences, Leiden University Medical Centre (LUMC), Leiden, the Netherlands
¹⁰ EPI-centre, KU Leuven, Leuven, Belgium
¹¹ Centre for Prognosis Research, School of Medicine, Keele University, Staffordshire, ST5 SBG, United Kingdom.

* Both authors contributed equally

Address for correspondence:

Professor Gary S Collins
Centre for Statistics in Medicine
Nuffield Department of Orthopaedics, Rheumatology & Musculoskeletal Sciences
University of Oxford
Oxford, OX3 7LD
United Kingdom.
Email: gary.collins@csm.ox.ac.uk

Twitter: @GSCollins, @CarlMoons
ORCID: 0000-0002-2772-2316
Abstract

Introduction: The Transparent Reporting of a multivariable prediction model of Individual Prognosis Or Diagnosis (TRIPOD) statement and the Prediction model Risk Of Bias ASsessment Tool (PROBAST) were both published to improve the reporting and critical appraisal of prediction model studies for diagnosis and prognosis. This paper describes the processes and methods that will be used to develop an extension to the TRIPOD statement (TRIPOD-AI) and the PROBAST (PROBAST-AI) tool for prediction model studies that applied machine learning techniques.

Methods and Analysis: TRIPOD-AI and PROBAST-AI will be developed following published guidance from the EQUATOR Network, and will comprise five stages. Stage one will comprise two systematic reviews (across all medical fields and specifically in oncology) to examine the quality of reporting in published machine-learning based prediction model studies. In stage two we will consult a diverse group of key stakeholders using a Delphi process to identify items to be considered for inclusion in TRIPOD-AI and PROBAST-AI. Stage three will be a virtual consensus meeting to consolidate and prioritise key items to be included in TRIPOD-AI and PROBAST-AI. Stage four will involve developing the TRIPOD-AI checklist and the PROBAST-AI tool, and writing the accompanying explanation and elaboration papers. In the final stage, stage 5, we will disseminate TRIPOD-AI and PROBAST-AI via journals, conferences, blogs, websites (including TRIPOD, PROBAST and EQUATOR Network), and social media. TRIPOD-AI will provide researchers working on prediction model studies based on machine-learning with a reporting guideline that can help them report key details that readers need to evaluate the study quality and interpret its findings, potentially reducing research waste. We anticipate PROBAST-AI will help researchers, clinicians, systematic reviewers and policymakers critically appraise the design, conduct and analysis of machine learning based prediction model studies, with a robust standardised tool for bias evaluation.

Ethics and Dissemination

Ethical approval has been granted by the Central University Research Ethics Committee (CUREC), University of Oxford on 10-December-2020 (R73034/RE001). Findings from this study will be disseminated through peer review publications.

Keywords: prediction, artificial intelligence, machine learning, reporting guideline, risk of bias, TRIPOD, PROBAST
Article Summary

Strengths and limitations of this study

- The reporting of clinical prediction models using artificial intelligence is poor.
- There are no guidelines for the reporting or risk of bias assessment of clinical prediction models using artificial intelligence.
- The strengths of this study is that it follows published guidance from the EQUATOR Network for developing reporting guidelines.
- Expert opinion and consensus will be obtained from multiple stakeholders (statisticians, clinician scientists, epidemiologists, computer scientists, funders, healthcare policy makers, patients, and industry leaders).
Background

Models that predict clinical outcomes are abundant in the medical literature and are broadly
categorised as those that estimate the probability of the presence of a particular outcome (diagnostic)
 or whether a particular outcome (e.g. event) will occur in the future (prognostic) [1]. Traditionally
these models (herein referred to as prediction model) have been developed using regression-based
methods, typically logistic regression for short-term outcomes and Cox regression for longer-term
outcomes [2]. Numerous reviews have observed that studies describing the development and
validation (including updating) of a prediction model often fail to report key information to help
readers judge the methods and have a complete, transparent and clear picture of the model’s
predictive accuracy and other relevant details such as the target population and the content of the
model itself [3–6]. The absence of full and comprehensive reporting limits the usability of the findings
of these studies, e.g., in subsequent validation studies, evidence synthesis studies, or in daily practice,
and therefore contribute to research waste [7]. In response to this, in 2015, the Transparent Reporting
of a multivariable prediction model for Individual Prognosis Or Diagnosis (TRIPOD) Statement was
published [1, 8]. The TRIPOD Statement is a checklist of 22-items that authors should report with
sufficient detail and clarity to inform how the study was carried out.

Since the publication of the TRIPOD Statement, artificial intelligence (AI) and in particular machine
learning, approaches to clinical prediction have evolved and grown in popularity with the number of
AI and machine learning publications rapidly rising [9–14]. This is evident within a recent review of
COVID-19 related prediction models, where 57 (out of 107 included studies) used machine learning
methods to develop their model [15].

Machine learning, a branch of artificial intelligence, can be broadly described as data analytical
methods that learn from data without being explicitly programmed, with patterns identified based on
the data itself. They are often described as having flexibility to capture complex associations
particularly in large and unstructured data and complexity in modelling. Whilst the vast majority of
the items in the TRIPOD Statement are relevant to machine learning based prediction model studies,
there are some unique challenges with machine learning that are not captured. Due to their
complexity, these prediction models are typically considered to be ‘black box’, unlike say regression-
based models where the full model can be transparently presented (e.g., as an equation containing
all the regression coefficients). Also, whilst many machine learning methods have origins in the
statistical literature, two (overlapping) prediction model cultures have emerged as those from a
statistical/epidemiological background and those from the computer science/data sciences [16].
Although there is clear overlap, different approaches to model development, validation and updating have appeared, and different and sometimes conflicting terminology have arisen.

Due to the relative novelty of applying machine learning methods to clinical prediction modelling, there is little information on the quality of reporting of these studies. However, the few reviews that have examined the completeness of reporting of have concluded that reporting is poor [17, 18]. In response to these concerns, guidance is required to help authors fully describe their prediction model study when machine learning methods were used. Therefore the TRIPOD group initiated a large international project to develop a consensus based extension of TRIPOD with specific focus on reporting of studies that undertake the development, validation or updating of a diagnostic or prognostic prediction model, using machine learning techniques – herein referred to as TRIPOD-AI [19]. The TRIPOD-AI extension, comprising a checklist and an accompanying elaboration and explanation document will provide researchers, authors, reviewers, editors, users, and other stakeholders of machine learning based prediction model studies, with guidance on the minimal set of items to report, with detailed examples of good reporting for each item.

Complete reporting allows studies to be understood, replicated and used. However, critical appraisal and of the quality of study method is a crucial component of evidence-based medicine as well. Critical appraisal and assessing the quality of studies is a crucial component of evidence-based medicine. In 2019, the Prediction model Risk Of Bias ASsessment Tool (PROBAST) was published [20, 21] to help a variety of stakeholders including, for example, systematic reviewers, researchers, journal editors, manuscript reviewers and policy makers involved in clinical guideline development, critically appraise the study design, conduct and analysis of prediction model studies. PROBAST comprises four domains (participants, predictors, outcome and analysis) and contains 20 signalling questions to facilitate risk of bias assessment. Clearly risk of bias assessment and reporting are intrinsically linked, in that judging risk of bias is predicated on what has been reported in the primary study. Whilst in principle PROBAST is relevant for prediction model studies using machine learning, different approaches to model development and validation, and terminology have appeared, and the ability to critically appraise these studies is crucial before they are implemented [22, 23]. Therefore, in parallel with the development of TRIPOD-AI, we will also develop PROBAST-AI, a tool to assess risk of bias in machine learning based multivariable prediction model studies.
Focus of TRIPOD-AI and PROBAST-AI

The focus of both TRIPOD-AI and PROBAST-AI is on reports of research or endeavours in which a multivariable prediction model is being developed (or updated), or validated (tested) using any (supervised) machine learning technique. Conforming to the original TRIPOD and PROBAST publications, a multivariable prediction model is defined as any combination or equation of two or more predictors that is to be used for individualized predictions to estimate an individual’s probability of having (diagnosis) or developing (prognosis) a particular health outcome or state. Predictors may have any form and emerge from patient history, physical examination, diagnostic, prognostic or monitoring tests and from undergone treatments. Outcomes may also have any form (dichotomous, categorical, continuous) and of any kind, such as, a particular condition or disorder being present or absent (diagnostic outcome or classification), short-term prognosis outcomes (e.g., hospital mortality, or postoperative complications), and long-term prognostic outcomes such as one year occurrence of treatment complications, five-year occurrence of metastases, or life-long survival).

As per the original publications, TRIPOD-AI and PROBAST-AI will also address prediction model studies from all medical care settings (public health, primary, secondary, tertiary and nursing home care) and all corresponding target populations (healthy individuals, suspected and diseased individuals).

TRIPOD-AI and PROBAST-AI are not meant to address:
- comparative studies that quantify the impact of using a prediction model as compared to not using the model [24];
- so-called predictor finding studies (also known as risk or prognostic factor studies) where multivariable machine learning techniques are used to identify (usually from a wider set of potential predictors) those predictors that are associated with an outcome, but not to develop a model that can be used for individualized predictions in new individuals;
- single medical test studies that use machine learning or AI techniques aimed to read, for example, CT or MRI, images to find which image-parameters are best associated with an outcome (such studies fall under the remit of STARD-AI [25]). If these image-parameters are included as predictors in a multivariable model combined with other predictors, TRIPOD-AI and PROBAST-AI may be useful.

Methods/design

Both TRIPOD-AI and PROBAST-AI will be developed following published guidance from the EQUATOR Network [26]. We will develop the guideline in five stages: (i) systematic reviews to establish the
quality of current reporting, (ii) Delphi exercise, (iii) consensus meeting, (iv) development of the
guidance statement, and (v) guideline dissemination. We have registered our intent to develop the
TRIPOD extension for AI on the EQUATOR Network website (www.equator-network.org), the TRIPOD
website (www.tripod-statement.org) and recently announced it in the Lancet [19], whilst the
PROBAST-AI development has been announced on the PROBAST website (www.probast.org).

TRIPOD-AI/PROBAST-AI working group

The TRIPOD/PROBAST Working Group will include: (1) an Executive Committee (2) an Advisory and
Working Group; and (3) a large international Delphi Panel.

The TRIPOD-AI/PROBAST-AI Executive Committee will be responsible for the leadership and
coordination of all the processes involved in the development and dissemination of the TRIPOD-AI
guideline. The Executive Committee consists of the two lead authors of the TRIPOD reporting guideline
and the PROBAST tool, and also prediction model experts and researchers from the machine learning
community. Key stakeholders for stage 2 (Delphi survey) will be identified and approached to
participate and a subset of these key stakeholders (the Advisory Group) will participate in stage 3
(consensus meeting).

Here the term key stakeholder refers to a cross-sector participant (both industry and public sector)
who falls into at least one of the following categories:

1. researchers who have used machine learning in the context of clinical prediction, have clear
   knowledge and expertise in using machine learning or developed machine learning methods.
   These include applied (bio)medical investigators, statisticians, epidemiologists, and data
   scientists;

2. assessors and approvers of artificial intelligence or machine learning model, such as regulatory
   assessors and ethics committee members;

3. beneficiaries or users of the resultant TRIPOD-AI guidance and PROBAST-AI tool such as
   journal editors and journal reviewers;

4. commissioners of research grants, such as funders;
5. consumers of research results such as healthcare providers and patients and citizens.

**Stage 1 – Systematic review of current reporting**

Two parallel systematic reviews are ongoing to evaluate the quality of current reporting in published studies developing, validating or updating machine learning based prediction models in the medical domain. Both systematic reviews will assess adherence of the reporting against the original TRIPOD Statement [1, 8], using the TRIPOD adherence checklist [27]. The reviews will also examine the methodological conduct of the primary studies, including a risk of bias assessment using the recently issued risk of bias tool (quality appraisal) for diagnostic and prognostic prediction model studies (PROBAST) [20, 21], and will draw out specific issues, currently not covered by TRIPOD and PROBAST relating to machine learning. The protocols for the two systematic reviews have been registered with the International Prospective Register of Systematic Reviews (PROSPERO IDs CRD42019140361 and CRD42019161764). One review (CRD42019161764) will examine the quality of reporting of machine learning based prediction model studies across all medical fields (between January 2018 to December 2019), whilst the other review (CRD42019140361) will focus on the quality of reporting of machine learning based prediction model studies published in oncology (between January 2019 to September 2019).

Undertaking these reviews serves two purposes: (1) to understand the completeness of current reporting of machine learning based prediction model studies in the medical literature, and (2) to identify unique reporting items for consideration for TRIPOD extension, and unique risk of bias or quality items for PROBAST extension. The data collection for this phase is underway. The reviews will evaluate the current completeness of reporting and the quality of the research and identify additional reporting and quality items to be considered for TRIPOD-AI and PROBAST-AI.

These two reviews will evaluate the current completeness of reporting and the quality of the research. Together with other evidence [3, 4, 17, 18, 28] from existing methodological guidance papers, they will provide important information on the transparency and quality of reporting. Using the original TRIPOD and PROBAST checklists as starting points, the Executive Committee will identify in the literature the preliminary items to consider in Stage 2 (the Delphi study) and therefore inclusion in the eventual TRIPOD-AI checklist and PROBAST-AI tool.
Stage 2 – Delphi exercise

We will perform an extensive Delphi survey among a large international network of relevant stakeholders, with a maximum of three rounds, to help decide on items that could be modified, added to, or removed from the TRIPOD 2015 checklist to form the TRIPOD-AI checklist, and subsequently the PROBAST-AI checklist.

Design

The Delphi process will comprise of a series of rounds where panellists will independently and anonymously evaluate and achieve consensus on the inclusion or exclusion of the proposed reporting and quality items – in addition to suggesting additional items. The process will be repeated for a maximum of three rounds. Following each round, participants will be provided with structured feedback of the previous round to help reconcile individual opinions and achieve group consensus. Items achieving a high level of agreement (≥70%) will be taken forward to the consensus meeting (stage 3).

Selection of potential items

The list of items for TRIPOD-AI (and PROBAST-AI) will be collated by the Executive Committee, including the results of the two systematic reviews, any other available studies on methodology or reporting of machine learning based prediction models, and expert recommendations from the Delphi panellists. Relevant methodological guidance or methodological papers will be retrieved to identify additional candidate reporting and quality items for machine learning based prediction model studies. Pre-selection involves dividing items into those to further consider, those that can be provided as optional guidance (to be outlined in an Explanation and Elaboration accompanying document), or those not to consider for potential inclusion. Delphi participants will have the opportunity to view and provide feedback in each round, and also to suggest new items.

Recruitment process and participants

Delphi participants will be identified through professional networks of the Executive Committee, participation in the Delphi exercise of the original TRIPOD guideline (and TRIPOD for Abstracts and TRIPOD Cluster Delphi surveys), original PROBAST Delphi exercise, via self-response to the Lancet 2019 paper where TRIPOD-AI was announced [19], and responses to social media announcements of TRIPOD-AI (e.g., Twitter).
We will invite international participants with diverse roles (e.g., researchers, healthcare professionals, journal editors, funders, policy makers, healthcare regulators, end users of prediction models) from a range of settings (e.g., universities, hospitals, primary care, biomedical journals, non-profit organisations, and for-profit organisations). Participants will be invited via personalized email that will describe the TRIPOD-AI extension and PROBAST-AI tool development, and explain the objective, process, and timelines of the Delphi exercise. We plan to invite at least 200 participants to the Delphi survey. In all rounds, the survey will remain open for three weeks, with a reminder email sent one week after the initial invitation. In round two of the Delphi exercise, additional participants may be sought to ensure fair representation of all key stakeholders [29].

Informed consent from participants will be obtained using an online consent form and participants can withdraw at any time. Individuals who indicate that they wish to opt out of the survey will be removed from subsequent invitations. Participants will not know the identities of other individuals in the Delphi panel, nor will they know the specific answers that any individual provides.

Procedure for selection of items
We plan to ask participants to consider the following guiding principles when reviewing existing, new or modified items for inclusion: 1) reporting of the item should facilitate reproducibility of the study (i.e. users should be able to recreate the findings based on the information reported); 2) reporting of the item facilitates assessment of the quality and risk of bias in and applicability of the machine learning study findings, to enhance their uptake and use in subsequent studies, systematic reviews and daily practice; 3) item is likely relevant to nearly all prediction model studies; 4) the set of items represent the minimum that should be reported in all machine learning studies developing, validating or updating a diagnostic or prognostic prediction model.

Round 1
Participants will be asked to rate on a 5-point Likert scale, the extent to which they agree with the inclusion of each checklist item in the TRIPOD-AI extension and PROBAST-AI tool (1=strongly disagree, 2=somewhat disagree, 3=I don’t know, 4=somewhat agree, 5=strongly agree). A free-text box will be provided for general comments on each item (to justify their decision or suggest wording changes), and a free-text box will be provided at the end of the survey to suggest additional checklist items or provide general comments on the checklist. The survey will be pilot-tested for usability and clarity to a small number of individuals familiar with prediction models or machine learning but not involved in
the TRIPOD-AI guideline extension or PROBAST-AI tool, and revised accordingly based on their feedback.

**Round 2**

The same participants involved in round 1 will be invited to participate in round 2. Participants will be provided with their first-round responses on each item, an anonymised summary of the group ratings and anonymised comments to justify ratings. Using the same format as round 1, participants will be presented with each item, including any new items suggested during round 1, and again express the extent to which they agree with the inclusion of the item in the TRIPOD-AI checklist or PROBAST-AI tool, considering the structured feedback to inform their responses. Participants who were invited to participate in round 1, but who did not respond will be invited to participate in round 2, and will be presented with an anonymised summary of the group ratings. Items that reached a high-level of agreement (scoring 4 or 5) in round 1 (≥ 70%) will be presented for information purposes only, with no voting on these items, though a free-text box will be provided for any comments. A third Delphi round will be used if deemed necessary by the Executive Committee.

**Results from the Delphi survey**

Item scores will be summarised for the entire panel as a whole, as appropriate (e.g., frequency and proportions across the rating categories) accompanied by a narrative summary of findings, comments, and suggestions. Results from both rounds of the survey will be discussed by the Executive Committee. For items where there was no consensus following the second Delphi found will be discussed by the Executive Committee, and will be considered for discussion at the subsequent consensus meeting.

**Stage 3 – Consensus meeting**

Two virtual consensus meetings (separately for TRIPOD-AI and PROBAST-AI), both spread over 2 days, will be held with the objective of discussing the results from the Delphi exercise and finalising items to be included in the reporting guideline and risk of bias tool. The composition of the consensus group will reflect the diversity of the key stakeholders addressed above. Key experts participating in the Delphi exercise will be considered to participate in the consensus meeting. We will also consider inviting experts who did not contribute to the Delphi to participate in the consensus. A total of around 25-30 international participants are expected to contribute to the virtual consensus meeting.

**Procedure**
The agenda and any material (e.g., results from the systematic reviews and Delphi) for the consensus meeting will be prepared by the Executive Committee and will be shared with attendees in advance. Members of the Executive Committee will facilitate a structured discussion on the rationale behind each item identified in the Delphi exercise. Consensus meeting participants will then be given the opportunity to discuss each item (reporting item for TRIPOD-AI and signalling question for PROBAST-AI), and vote on each item. The decision to retain an item in the TRIPOD-AI and PROBAST-AI will be based on achieving at least 70% support from the consensus meeting participants. The group will agree on the draft list of reporting items for the final TRIPOD-AI extension and PROBAST-AI tool. Specific item wording will not be discussed during the meeting, though participants can suggest and the group to agree on general intent and meaning of the item. Plans for dissemination will be discussed at the end of the consensus meeting.

Pilot testing
We will invite authors of machine learning prediction model studies in the medical domain, doctoral students undertaking prediction model, machine learning courses or workshops, and peer reviewers and editors of journals who frequently publish such prediction model studies, to pilot the use of a draft version of the TRIPOD-AI checklist and PROBAST-AI tool. We will ask those who pilot the checklist and tool whether the wording of items is ambiguous or difficult to interpret.

Stage 4 – Development of the draft TRIPOD-AI statement, PROBAST-AI and Explanation & Elaboration documents
The Executive Committee will lead the development of the TRIPOD-AI reporting guidance and PROBAST-AI signalling questions based on the agreed list of items from the consensus meeting (stage 3). The Executive Committee will invite a subset of members from the consensus meeting (to form a writing group) to help draft the Explanation & Elaboration paper.

The Executive Committee will reserve the right to update (that is remove or add) additional items to the TRIPOD-AI checklist during the development of the TRIPOD-AI statement, if and as necessary (as a result of the pilot testing).

For each of the TRIPOD-AI extension and the PROBAST-AI risk of bias tool, two manuscripts will be developed; (1) the Statement paper, presenting the checklist/tool and describing the process of how it was developed and (2) an Explanation & Elaboration paper. The Explanation & Elaboration papers
will outline the rationale of the reporting items (TRIPOD-AI) and signalling questions (PROBAST-AI), examples of good reporting (TRIPOD-AI) and examples of how to use PROBAST-AI. Drafts of the papers will be circulated to all participants of the consensus meeting for their comments.

Stage 5 – Guideline dissemination

The dissemination strategy will be informed by discussions at the consensus meeting. We will aim to seek simultaneous publication in key journals to target different readerships. To increase visibility and aid uptake, the TRIPOD-AI checklist and PROBAST-AI tool will be published open access, and made available on the TRIPOD website along with other TRIPOD extensions (www.tripod-statement.org), and on the PROBAST website (www.probast.org) respectively, as well on the PROGRESS website (www.prognosisresearch.com). The TRIPOD-AI extension will be indexed on the EQUATOR website (www.equator-network.org). Social media will be used to help disseminate the extension. The Executive Committee will (and consensus participants will be encouraged to) publicise the TRIPOD-AI statement and PROBAST-AI tool at key conferences and courses.

Publication plan

It is envisaged that the following publications will arise from the TRIPOD-AI and PROBAST-AI initiative:

- Publication 1: Study protocol
- Publication 2: Systematic review protocol (with registration on PROSPERO)
- Publication 3 & 4: Systematic reviews
- Publication 5 & 6: TRIPOD-AI statement and the Explanation and Elaboration paper
- Publication 7 & 8: PROBAST-AI tool and the Explanation and Elaboration paper

Conclusion

The number of prediction model studies using machine learning methods is rapidly increasing, including developed, validated or updated prediction models. Ensuring that key details are reported is important so that readers can evaluate the study quality, and interpret its findings including the developed, validated or updated prediction model to enhance their uptake in subsequent research (e.g. validation studies), evidence synthesis projects (e.g. systematic reviews of prediction models), and in daily practice by healthcare professionals, patients or citizens. We anticipate that TRIPOD-AI will help authors transparently report their study and help reviewers, editors, policy makers and end-users understand the methods and findings, and thereby reduce research waste. Similarly, we anticipate PROBAST-AI will help researchers, clinicians, systematic reviewers and policy-makers
critically appraise the design, conduct and analysis of machine learning based prediction model studies.

**Patient and Public Involvement**

No patients or members of the public were involved in the development of this protocol. Patient and public involvement is planned for the virtual consensus meeting.

**Ethics and Dissemination**

Ethical approval has been granted by the Central University Research Ethics Committee (CUREC), University of Oxford on 10-December-2020 (R73034/RE001). Findings from this study will be disseminated through peer review publications.

**Contributor statement**

Gary S. Collins, Paula Dhiman, Constanza L. Andaur Navarro, Jie Ma, Lotty Hooft, Johannes B. Reitsma, Patricia Logullo, Andrew L. Beam, Lily Peng, Ben Van Calster, Maarten van Smeden, Richard D. Riley, Karel G.M. Moons were involved in the planning and design of the study. Gary Collins drafted the manuscript with all authors contributing to the writing.

**Competing interests**

The authors declare no competing interests.

**Data sharing statement**

There is no data to share.

**Funding**

This research was supported by Health Data Research UK, an initiative funded by UKResearch and Innovation, Department of Health and Social Care (England) and the devolved administrations, and leading medical research charities, Cancer Research UK programme grant (C49297/A27294), the NIHR Biomedical Research Centre, Oxford, and the Netherlands Organisation for Scientific Research. The funders have no role in the development of this protocol and will not have a role in the data collection, analyses, interpretation of the data, and publication of the findings.

**References**


