



Systematic reviews of prognosis studies I: introduction, design and protocol for systematic reviews of prognostic studies

Carl Moons, Lotty Hooft, Anneke Damen

Katrina Williams, Nicole Skoetz, Marialena Trivella, Angela Aldin, Thomas Debray, Richard Riley, Jill Hayden



We have no actual or potential conflict of interest in relation to this presentation



Outline of workshop

Presentations:

- Introduction to types of prognosis research
- Introduction to types of SR of prognosis studies
- Defining the review question
- Data extraction and Critical appraisal

Lecture + practicals



Systematic reviews (SRs)

- Applicable to all fields of medical research
 - Therapeutic studies (RCTs): Cochrane Intervention Reviews
 - Diagnostic accuracy studies: Cochrane Diagnostic Test Accuracy Reviews
 - Both including meta-analytical approaches
- Next: prognosis studies



Why?

- Increasing interest in and demand for the evaluation of prognostic factors, biomarkers & models
- Growing number of primary studies
- Reviews more challenging: more variation in questions, designs, effect measures, analyses
- Several recent methodological developments & remaining challenges



Group exercise – 5 minutes

1. What is prognosis?
2. Why do we prognosticate?
3. Types of prognosis studies?



Answer

(BMJ series 2009 (Altman, Moons, Royston, Vergouwe) + Progress series
BMJ/Plos Med 2013)

Forecast of the **course** and **outcome** for an **individual** in a **certain health state** (given a **specific treatment** management)

- Not necessarily sick people
- More technical: probable course/prediction of specific future outcomes in subjects with certain health condition
- Disease does not have a prognosis → an individual does



Answer

- Why prognosticate:
 - To provide information to patients
 - Identify groups for treatment or other management – including abstine
 - To target specific prognostic factors that modify treatment effects
 - Select high/low risk patients for inclusion in RCTs
 - Adjust for case-mix differences in comparison health care of institutes
 - Service developers make decisions about what services are needed
 - policy makers what to support/advocate



Answer

Types of prognosis studies?

PROGRESS series 2013: BMJ and Plos Med

1. Average/overall prognosis: 'What is most likely course (outcome) of individuals with certain health condition?'
2. Prognostic factor studies: 'Which factors are associated with specific outcome in individuals with certain health condition?'
3. Prognostic modeling studies: 'What combination of prognostic factors predict, and how well, a certain outcome in individuals with a certain health condition?'
4. Treatment selection factors: 'Which factors lead to/predict different treatment effect in individuals to be treated?'

Focus on 2 + 3



Why SRs Prognosis studies?

1. Number of studied prognostic factors increases per day due to precision/personalized medicine focus

– Biomarkers (all types)



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Product: NycoCard CRP test

NycoCard CRP test

The NycoCard CRP test is a 2-minute Point of Care test to indicate bacterial or viral cause of infection. NycoCard CRP measures C-reactive protein (CRP), an acute phase protein that increases rapidly after onset of infection.

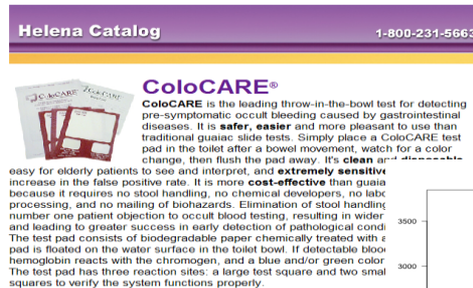
Test specific information

- Sample volume: 5 µL
- Assay time: 2 minutes
- Sample material: Whole blood, serum or plasma
- Measuring range: 0 - 250 mg/L for whole blood samples and 0 - 150 mg/L for serum and plasma samples
- Stability at room temperature: 4 weeks
- Kit size: 24 and 48 test
- NycoCard CRP Control: Positive control provided with the kit

Clinical use of NycoCard CRP

- Reduces unnecessary use of antibiotics
- More rapid induction of treatment
- Fewer hospital admissions
- Healthcare cost savings

See the [CRP Test Procedure](#) for information on how to run a test.



Helena Catalog 1-800-231-5663

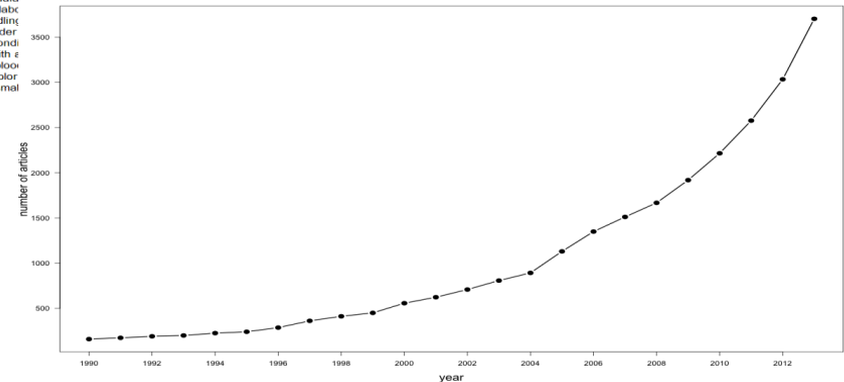
ColoCARE®

ColoCARE is the leading throw-in-the-bowl test for detecting pre-symptomatic occult bleeding caused by gastrointestinal diseases. It is **safer**, **easier** and more pleasant to use than traditional guaiac slide tests. Simply place a ColoCARE test pad in the toilet after a bowel movement, watch for a color change, then flush the pad away. It's **clean** and **disposable**.

easy for elderly patients to see and interpret, and **extremely sensitive** increase in the false positive rate. It is more **cost-effective** than guaiac because it requires no stool handling, no chemical developers, no lab processing, and no mailing of biohazards. Elimination of stool handling number one patient objection to occult blood testing, resulting in wider and leading to greater success in early detection of pathological condi

The test pad consists of biodegradable paper chemically treated with a pad is floated on the water surface in the toilet bowl. If detectable blood hemoglobin reacts with the chromogen, and a blue and/or green color

The test pad has three reaction sites: a large test square and two small squares to verify the system functions properly.



– Also prognostic models

Why SRs Prognosis studies?

2. Most studies conflicting results

- much more than in therapeutic trials and in diagnostic test accuracy studies
- **Non-randomised (often not predesigned studies)**
- **Often retrospective using existing data sets**

3. Relatively small studies (compared to therapeutic studies)

- Kyzas Eur J Canc 2007; > 1500 studies cancer prognostic markers in 2005 → largest just over 1000 pts.



Hence ...

... prognosis studies are hot

... SR's and notably MA of prognosis studies as well

- highly desired and well received by journals/policy makers →
- to systematically summarise the existing evidence in the field



Types of prognosis studies

PROGRESS series 2013: BMJ and Plos Med

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4. Treatment selection factors: 'Which factors lead to/predict different treatment effect in individuals to be treated?'



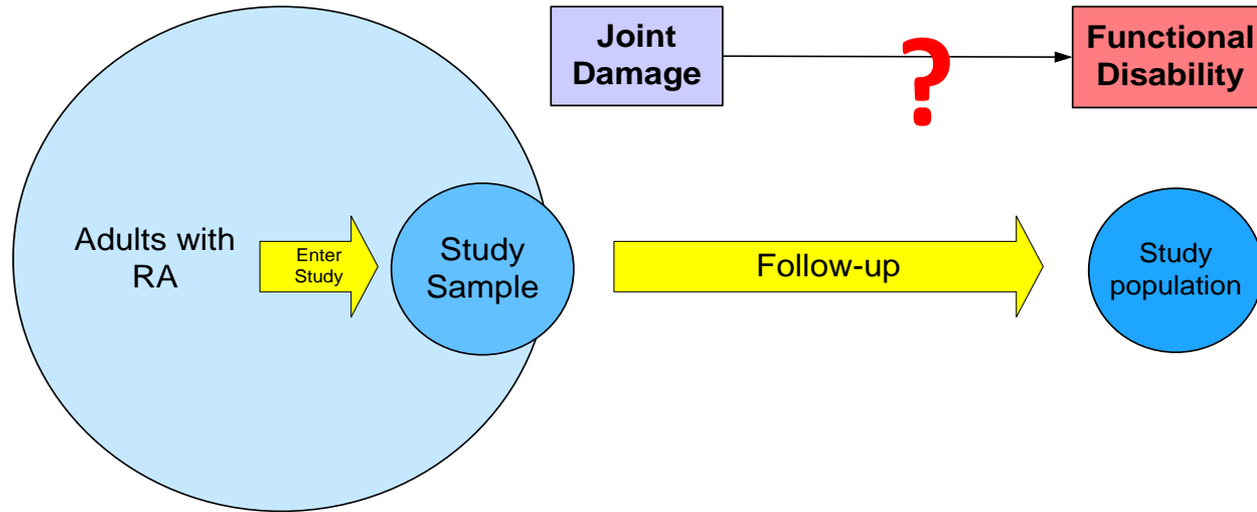
Prognostic Factor Studies

Aim:

- To identify factors associated with subsequent outcomes in subjects with certain health condition
- Not necessarily sick (patients)
- Independent predictors



Prognostic Factor Study Example



Adapted from: *Fletcher & Fletcher, Clinical Epidemiology – The Essentials. Chapter 6. Williams & Wilkins, Baltimore. 1996*



Types of prognosis studies

PROGRESS series 2013: BMJ and Plos Med

1. Average/overall prognosis: 'What is most likely course (outcome) of individuals in certain health condition (often certain disease)?'

2. Prognostic factor studies: 'Which factors are associated with specific outcome in individuals with certain health condition?'

3. Prognostic modeling studies: 'What combination of prognostic factors predict, and how well, a certain outcome in individuals with a certain health condition?'

4. Treatment selection factors: 'Which factors lead to/predict different treatment effect/response in individuals to be treated?'



Prognostic Prediction Model Studies

1. What is a prognostic prediction model study, and what is difference with multivariable analysis of prognostic factors?
2. There are three phases of prediction modelling – which three?
3. What is the biggest difference between phase 1+2 versus 3?



Answers

Risk prediction = foreseeing / foretelling
... (probability) of something that is yet unknown

Combination of 2 or more predictors that convert predictor values into an absolute probability of ...

- ...(presence of disease/result of reference test – diagnostic prediction model)
- ...future occurrence of certain outcome – prognostic prediction models

A prediction model is developed for use in new individuals to estimate their (diagnostic or prognostic) probability. Focus is on accuracy of entire model (discrimination + calibration). Factors of the model not at interest.

Multivariable analysis of prognostic factors not focus on model, but rather on which are independent predictors – Focus is on the HRs of the factors (adjusted HRs)



3 Phases of Prediction Modelling studies

BMJ series 2009/Bouwmeester 2012/PROGRESS series 2013 (BMJ/Plos Med)

1. *Model development studies* – to develop prediction model from data: identify important predictors; estimate predictor weights; construct model for individualised predictions; quantify predictive performance; internal validation
2. *Model validation studies* – test (validate) predictive performance of previously developed model in participant data other than development set
3. *Model impact studies* – quantify effect/impact of actually using model on participant/physician management and health outcomes – relative to not using the model

What is the difference between 3 versus 1 and 2?



3 Phases of Prediction Modelling studies

BMJ series 2009/Bouwmeester 2012/PROGRESS series 2013 (BMJ/Plos Med)

- Big difference = 3 are comparative studies → ideally randomised
- 1 and 2 are by definition single cohort studies- no inherent comparison
- 3 are thus ideally RCTs – for SRs of prediction model impact studies use the Cochrane tools available for RCTs of intervention studies
- This course provides tool for prediction model development and validation studies (type 1 and 2)



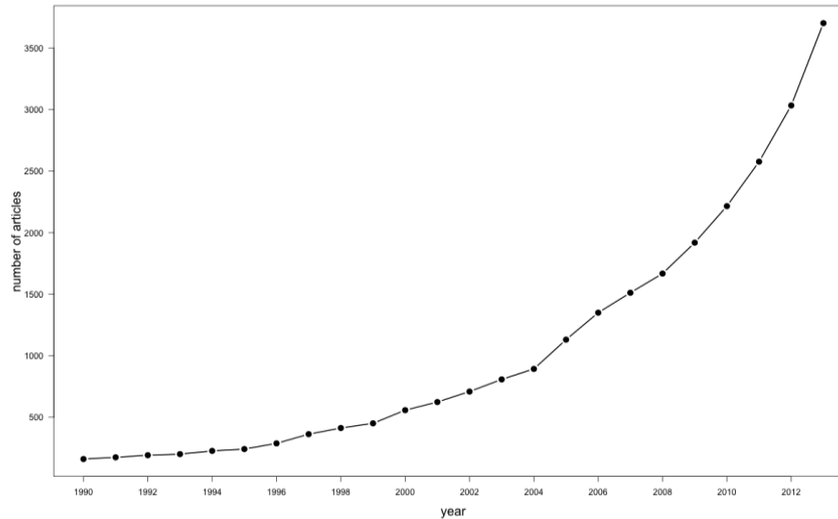
Everything we say from here on also applies to SRs of diagnostic prediction modelling studies

You need no separate course for that!
We use generic term: prediction model

Interesting and booming field – stay in it!



Prediction models are hot



("prognostic model") OR ("prediction model") OR ("risk score") OR
("clinical prediction rule") OR ("decision rule") OR ("prognostic index") OR
("prognostic indices") OR ("prediction index") OR ("risk algorithm") OR
("risk stratification") OR ("multivariable prediction"))



Conducting a systematic review of prognosis studies

1. Formulate review question (PICOTS)
2. Searching for studies
3. Screening and Selection of articles
4. Extraction of data
5. Risk of Bias assessments
6. Synthesis of data (meta-analysis)
7. Interpretation and conclusions

Step 1. Well-formulated review question

Actually: define the PICO → stands for?

Guidance frame review question: CHARMS checklist

Critical Appraisal and Data Extraction for Systematic
Reviews of Prediction Modelling Studies: The CHARMS
Checklist

Plos Med 2014

Karel G. M. Moons^{1*}, Joris A. H. de Groot^{1†}, Walter Bouwmeester¹, Yvonne Vergouwe¹, Susan Mallett²,
Douglas G. Altman³, Johannes B. Reitsma¹, Gary S. Collins³

RESEARCH METHODS AND REPORTING

A guide to systematic review and meta-analysis of prediction
model performance BMJ 2017

Thomas P A Debray,^{1,2} Johanna A A G Damen,^{1,2} Kym I E Snell,³ Joie Ensor,³ Lotty Hooft,^{1,2}
Johannes B Reitsma,^{1,2} Richard D Riley,³ Karel G M Moons^{1,2}

PICOTS SR Prognostic factor(s)

Item	Comments
1. <u>P</u>opulation	Define target population in which prognostic factor(s) under review will be used.
2. <u>I</u>ndex factor(s)	Define the prognostic factor(s) under review.
3. <u>C</u>omparator(s)	If applicable, one can review more than one factor for the target population and outcome under review.
4. <u>O</u>utcome(s)	Define the outcome(s) of interest for the factor(s) under review.
5. <u>T</u>iming	Define at what time-points the prognostic factor(s) are to be used and over what time period the outcome(s) are predicted
6. <u>S</u>etting	Define the intended role or setting of the prognostic factor(s) under review.



PICOTS SR Prognostic (prediction) model(s)

Item	Comments
1. <u>P</u>opulation	Define target population in which prediction model(s) under review will be used.
2. <u>I</u>ndex model(s)	Define the prediction model(s) under review.
3. <u>C</u>omparator(s)	If applicable, one can review more than one model for the target population and outcome under review.
4. <u>O</u>utcome(s)	Define the outcome(s) of interest for the model(s) under review.
5. <u>T</u>iming	Define when prediction model(s) under review is intended to be used and over what time period (notably for prognostic prediction models) the outcome(s) is predicted.
6. <u>S</u>etting	Define the intended role or setting of the prediction model(s) under review.



Practical

Exercise:

- **Define a review question + PICOTS**

Research

BMJ

Value of sentinel node status as a prognostic factor in melanoma: prospective observational study

Stephen Kettlewell, Colin Moyes, Caroline Bray, David Soutar, Alan MacKay, Dominique Byrne, Taimur Shoaib, Barun Majumder, Rona MacKie

Abstract

Objective To establish the prognostic value of knowledge of sentinel node status in melanoma.

Design Single centre prospective observational study, with sentinel nodes identified by lymphoscintigraphy, γ probe, and intraoperative blue dye and examined by both conventional histopathology and immunopathology

multicentre randomised trial (MSLT1) is in progress with the aim of determining if patients with melanoma who have a positive SNB and proceed immediately to full node dissection have a superior disease-free survival or overall survival compared with patients who have node dissection only when nodes draining the site of the primary melanoma are clinically palpable. Definitive results are awaited.¹²



Suggested answer

Population	<ul style="list-style-type: none">• Patients with melanoma
Index factor	<ul style="list-style-type: none">• Sentinel node status
Comparator	<ul style="list-style-type: none">• Not applicable
Outcomes	<ul style="list-style-type: none">• Recurrence• Mortality
Timing	<ul style="list-style-type: none">• Prediction at preoperative visit• 3 months prediction of outcomes
Setting	<ul style="list-style-type: none">• Patients scheduled for surgery to remove the melanoma



BMC Medical Informatics and Decision Making



Research article

Open Access

Systematic review of prognostic models in traumatic brain injury

Pablo Perel*, Phil Edwards, Reinhard Wentz and Ian Roberts

Address: Nutrition and Public Health Intervention Research Unit, Epidemiology and Population Health Department, London School of Hygiene

Different clinical questions possible → different aims of SR of prediction models?

Group exercise:

- **Define a review question + PICOTS**

Suggested answer

Population	<ul style="list-style-type: none">• Patients with TBI (e.g. surviving the first 24 hours)
Index model(s)	<ul style="list-style-type: none">• All developed+validated models for patients with TBI (surviving the first 24 hours)
Comparator	<ul style="list-style-type: none">• All existing developed+validated models
Outcomes	<ul style="list-style-type: none">• Mortality• Or: Daily functioning
Timing	<ul style="list-style-type: none">• Prediction T0 24 hours after accident/injury• 3 months prediction of outcome (or: 12 months)
Setting	<ul style="list-style-type: none">• Patients in hospital surviving a TBI after 24 hours• Or: battlefield TBI• Or: prediction in ambulance or at the site of the accident



Types of SR prognostic/prediction model questions

- Review all models for specific outcome in specific target population
 - Models predicting fatal/non-fatal CHD in general population; models predicting stroke in general population;
 - Models predicting survival after cardiac surgery ; predicting Length of stay after cardiac surgery ; predicting QoL after surgery
- Review all existing models in a particular clinical field
 - e.g. all models for any CVD outcome in general population ; all developed models in obstetrics.



Types of SR prognostic/prediction model questions

- How good is predictive performance of specific model for specific target population (validation studies only)
 - Predictive performance Framingham risk model / GAIL model
- Review on added predictive value of specific predictor/biomarker/test to a specific model
 - Adding CRP to Framingham risk score; D-dimer to Wells Rule
 - Adding imaging results to 'basic risk scores' (cancer models)



Conducting a systematic review of prognosis studies

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Search Strategies

- No optimal, reliable methods for searching the literature for prognostic information
 - As for RCTs and Diagnostic Test Accuracy Studies
- A few published
 - Altman DG (2001): single prognostic factors
 - Wong SS (2003): very generic
 - Ingui BJ (2001): prediction models
 - Geersing (2012): validation Ingui (2001) and updated (new) search strategy

Search Filters for Finding Prognostic and Diagnostic Prediction Studies in Medline to Enhance Systematic Reviews

Geert-Jan Geersing^{1*}, Walter Bouwmeester^{1,9}, Peter Zuithoff¹, Rene Spijker^{2,4}, Mariska Leeflang^{3,4}, Karel Moons¹

1 Julius Center for Health Sciences and Primary Care, University Medical Center Utrecht, Utrecht, The Netherlands, **2** Medical Library Academic Medical Center, University of Amsterdam, Amsterdam, The Netherlands, **3** Department of Clinical Epidemiology and Bio-Informatics, Academic Medical Center, University of Amsterdam, Amsterdam, The Netherlands, **4** Dutch Cochrane Center, Academic Medical Center, University of Amsterdam, Amsterdam, The Netherlands

Table 1. Search strategies for finding prediction research in Medline.

Filter	Search terms included in the filter*	Sensitivity# (95% CI)	Specificity# (95% CI)
Ingui filter	(Validat\$ OR Predict\$.ti. OR Rule\$) OR (Predict\$ AND (Outcome\$ OR Risk\$ OR Model\$)) OR ((History OR Variable\$ OR Criteria OR Scor\$ OR Characteristic\$ OR Finding\$ OR Factor\$) AND (Predict\$ OR Model\$ OR Decision\$ OR Identif\$ OR Prognos\$)) OR (Decision\$ AND (Model\$ OR Clinical\$ OR Logistic Models/)) OR (Prognostic AND (History OR Variable\$ OR Criteria OR Scor\$ OR Characteristic\$ OR Finding\$ OR Factor\$ OR Model\$))	0.98 (0.92–1.0)	0.86 (0.85–0.87)
Haynes broad filter	(Predict*[tiab] OR Predictive value of tests[mh] OR Scor*[tiab] OR Observ*[tiab] OR Observer variation[mh])	0.96	0.79

*Using the Pubmed interface for MEDLINE.

#Sensitivity and specificity as reported by Ingui and Haynes in their original publication; CI= confidence interval, for the Haynes broad filter no confidence intervals were given in the original publication.

doi:10.1371/journal.pone.0032844.t001



Geersing et al 2012

Conclusions

- Updated search strategy for prognosis research good in retrieving "Prediction model studies" (Se 0.78 to 0.89)
- Less value in retrieving "Predictor Finding/prognostic factor" and "Prediction Model Impact Studies"

Table 4. Updated search string for finding prediction research.

"Stratification" OR "ROC Curve"[Mesh] OR "Discrimination" OR "Discriminate" OR "c-statistic" OR "c statistic" OR "Area under the curve" OR "AUC" OR "Calibration" OR "Indices" OR "Algorithm" OR "Multivariable"

doi:10.1371/journal.pone.0032844.t004

- Strategy for "Predictor Finding / prognostic factor" studies still sub-optimal but good starting point!



Study selection

- Selecting studies involves judgement, and is highly influential on the outcomes of the review
- Two (or more...) reviewers, independently
 - minimizing bias
 - pilot selection on a few papers first: substantial variation
 - Determine how will disagreements be managed
- Examine titles and abstracts
- Flow chart of included/excluded studies
- Retrieve and examine full text reports



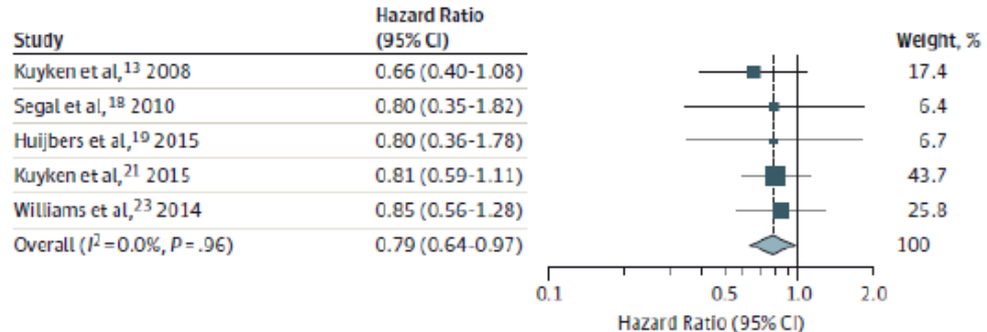
Intermezzo Challenge

Meta-analysis/Pooling of prognostic factor studies

Exercise 10 minutes:

1. Assume this forest plot is of RCTs on intervention X to prevent outcome Y in patients with disease Z.

- Is this pooling ok?
- Why or why not?



2. Assume this forest plot is of studies on prognostic factor X, to predict outcome Y in patients with disease Z.

- Is this pooling ok?
- Why or why not?



Meta-analysis/Pooling in prognostic factor studies

Answers:

- If RCTs
 - Pooling is ok – provided correctly randomised
 - Then the 3 HRs are unbiased (provided no other risks of biases) so can easily pool them
 - Clear effect of intervention X to prevent outcome Y
 - In frequentistic world, at alpha 0.05 – even statistically significant result.
- If prognostic factor studies?
 - Non randomised → even if a study was based on a RCT – the prognostic factor analysis is per arm and thus non randomised
 - Can not assume that the 3 HRs are unbiased
 - Only pool them if studies have adjusted for the same co-variates – or largely for the same co-variates – e.g. the same big 6 or 7 (the eighth co variate probably did not change the HR further)
 - So pooling of prognostic factor studies only if same adjustment -- otherwise do stratified pooling (e.g. over studies with similar adjustment)



Conducting a systematic review of prognosis studies

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CHARMS

- Extraction of characteristics/data of included studies + Critical appraisal
 - **CHARMS** – Table 2

Critical Appraisal and Data Extraction for Systematic Reviews of Prediction Modelling Studies: The CHARMS Checklist

Karel G. M. Moons^{1†*}, Joris A. H. de Groot^{1†}, Walter Bouwmeester¹, Yvonne Vergouwe¹, Susan Mallett², Douglas G. Altman³, Johannes B. Reitsma¹, Gary S. Collins³

- Does not exist for prognostic factor studies – though can use CHARMS for that



Data Extraction Key issues CHARMS checklist

Domain	Key items	Reported on page #
SOURCE OF DATA	Source of data (e.g., cohort, case-control, randomized trial participants, or registry data)	
PARTICIPANTS	Participant eligibility and recruitment method (e.g., consecutive participants, location, number of centers, setting, inclusion and exclusion criteria)	
	Participant description	
	Details of treatments received, if relevant	
	Study dates	
OUTCOME(S) TO BE PREDICTED	Definition and method for measurement of outcome	
	Was the same outcome definition (and method for measurement) used in all patients?	
	Type of outcome (e.g., single or combined endpoints)	
	Was the outcome assessed without knowledge of the candidate predictors (i.e., blinded)?	
	Were candidate predictors part of the outcome (e.g., in panel or consensus diagnosis)?	
	Time of outcome occurrence or summary of duration of follow-up	
CANDIDATE PREDICTORS (OR INDEX TESTS)	Number and type of predictors (e.g., demographics, patient history, physical examination, additional testing, disease characteristics)	
	Definition and method for measurement of candidate predictors	
	Timing of predictor measurement (e.g., at patient presentation, at diagnosis, at treatment initiation)	
	Were predictors assessed blinded for outcome, and for each other (if relevant)?	
	Handling of predictors in the modelling (e.g., continuous, linear, non-linear transformations or categorised)	
SAMPLE SIZE	Number of participants and number of outcomes/events	
	Number of outcomes/events in relation to the number of candidate predictors (Events Per Variable)	
MISSING DATA	Number of participants with any missing value (include predictors and outcomes)	
	Number of participants with missing data for each predictor	
	Handling of missing data (e.g., complete-case analysis, imputation, or other methods)	



Data Extraction Key issues CHARMS checklist

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MODEL DEVELOPMENT	Modelling method (e.g., logistic, survival, neural network, or machine learning techniques)	
	Modelling assumptions satisfied	
	Method for selection of predictors for inclusion in multivariable modelling (e.g., all candidate predictors, pre-selection based on unadjusted association with the outcome)	
	Method for selection of predictors during multivariable modelling (e.g., full model approach, backward or forward selection) and criteria used (e.g., p-value, Akaike Information Criterion)	
	Shrinkage of predictor weights or regression coefficients (e.g., no shrinkage, uniform shrinkage, penalized estimation)	
MODEL PERFORMANCE	Calibration (calibration plot, calibration slope, Hosmer-Lemeshow test) and Discrimination (C-statistic, D-statistic, log-rank) measures with confidence intervals	
	Classification measures (e.g., sensitivity, specificity, predictive values, net reclassification improvement) and whether a-priori cut points were used	
MODEL EVALUATION	Method used for testing model performance: development dataset only (random split of data, resampling methods e.g. bootstrap or cross-validation, none) or separate external validation (e.g. temporal, geographical, different setting, different investigators)	
	In case of poor validation, whether model was adjusted or updated (e.g., intercept recalibrated, predictor effects adjusted, or new predictors added)	
RESULTS	Final and other multivariable models (e.g., basic, extended, simplified) presented, including predictor weights or regression coefficients, intercept, baseline survival, model performance measures (with standard errors or confidence intervals)	
	Any alternative presentation of the final prediction models, e.g., sum score, nomogram, score chart, predictions for specific risk subgroups with performance	
	Comparison of the distribution of predictors (including missing data) for development and validation datasets	
INTERPRETATION AND DISCUSSION	Interpretation of presented models (confirmatory, i.e., model useful for practice versus exploratory, i.e., more research needed)	
	Comparison with other studies, discussion of generalizability, strengths and limitations.	



MODEL DEVELOPMENT	Modelling method (e.g., logistic, survival, neural network, or machine learning techniques)	
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	Method for selection of predictors for inclusion in multivariable modelling (e.g., all candidate predictors, pre-selection based on unadjusted association with the outcome)	
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MODEL PERFORMANCE	Shrinkage of predictor weights or regression coefficients (e.g., no shrinkage, uniform shrinkage, penalized estimation)	
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MODEL EVALUATION	Classification measures (e.g., sensitivity, specificity, predictive values, net reclassification improvement) and whether a-priori cut points were used	
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Conducting a systematic review of prognosis studies

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Risk of Bias tools (this afternoon)

Prognostic factor/predictor finding studies

- **QUIPS** → J Haydn, Ann Int Med 2006 + 2013

Prediction model studies (development and validation)

- **PROBAST** – Ann Int Med (fall 2018)



RoB tools: QUIPS & PROBAST

RESEARCH AND REPORTING METHODS | **Annals of Internal Medicine**

Assessing Bias in Studies of Prognostic Factors

Jill A. Hayden, DC, PhD; Danielle A. van der Windt, PhD; Jennifer L. Cartwright, MSc; Pierre Côté, DC, PhD; and Claire Bombardier, MD

PROBAST: a tool to assess risk of bias and applicability of prediction model studies – explanation and elaboration

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This afternoon



Cochrane PMG title registration form for SRs of prognostic studies



Prognosis Studies review proposal form

Review Proposal Form

Please complete this form to outline your proposal for a Cochrane systematic review. Email the completed form to [email address], or send to [name], Managing Editor, Cochrane XXX Group, [postal address]. P: +XX XXXXXXXX Fax: +XX XXXXXXXX.

Before completing this form:

- Read "Managing expectations: what does The Cochrane Collaboration expect of authors, and what can authors expect of The Cochrane Collaboration?" (see <http://community.cochrane.org/editorial-and-publishing-policy-resource/cochrane-review-development/managing-expectations>) Note: this information is particularly for systematic reviews of intervention studies. A page for prognosis reviews is under construction.)
- Note that a Cochrane review of prognosis studies clearly differs from that of intervention studies and diagnostic test accuracy studies, in, e.g., searching, data extraction, critical appraisal and meta-analysis. Step-by-step guidance to help you understanding prognosis studies and the processes of conducting a review of prognosis studies is given in the papers in the reference list below.
- Cochrane reviews of prognosis require a multidisciplinary team. Below you find several question addressing the available expertise in the author team, and whether external expertise (e.g. from information specialists or methodologists) is needed to conduct this review. If additional expertise is needed, e.g. an information specialist, or methodological or statistical expertise, please provide this request to the Prognosis Methods Group (PMG) timely.

Proposed title

Choose one of the formats below. See also the generic guidance on defining a review question for prognosis studies in the CHARMS checklist.

Incidence of [outcome] within [time] in [population]
 [Prognostic factors] for predicting incidence of [outcome] in [population]
 Prediction of [outcome] in [population] using [prognostic factors]
 Prognostic models for predicting [outcome] in [population]
 Predictive performance of [prognostic model] for predicting [outcome] in [population]
 Added value of [prognostic factor] on top of [existing prognostic factors/prognostic model] for predicting [outcome] in [population]
 [Predictive factors] predicting the [outcome of treatment] in [population]
 [Factors / Models] predicting differential treatment response in [population]
 [Factors / Models] for predicting treatment response in [population]

Short description of review proposal

Provide brief but enough information to make sure that the clinical context and the actual question that is being asked is clear for non-content experts as well.

For explicit guidance to help filling in this title registration form and for the conduct of the review, from framing the review question, search strategy, study in/exclusion criteria, critical appraisal, risk of bias assessment, meta-analysis and reporting, please see the papers mentioned in the reference list below.

Type of prognosis review
 Indicate what type of review you are going to perform (double click to check a box). See PROGRESS series in the reference list.

Overall prognosis
 Prognostic factors
 Prognostic models
 Predictive/Treatment selection factors

Motivation for the review
 For example, is this going to be part of a PhD thesis; is it part of a larger project; is it particularly topical at the present time?

Background
 i) The clinical problem.
 A short description of the existing clinical pathway of the targeted individuals/patients; their starting condition and moment of prognostication (time point in the clinical pathway); what prognostic outcomes are relevant to the targeted individuals. For predictive factor reviews also refer to the role of treatment.
 ii) Why is this review relevant, including how might the results of the review be used: e.g., the prognostic or predictive factor(s) or model(s) under review may be used to determine treatment allocation or abstention, decide on closer follow-up or monitoring, etc. Reference to an existing systematic review on this topic outside Cochrane is helpful.

Review objective(s)
 What is the review question, according to the PICOTS format? (see Box 1 in the paper of Debray et al, BMJ 2017, see reference list below.)

Primary objective: []
 Secondary objective(s): []

Participants / setting
 Short outline of the targeted population and clinical setting, to be included and excluded for the review.



Cochrane PMG Protocol Template for SRs of prognostic studies



Protocol Cochrane Review Prognosis Studies

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*Prognosis exemplar protocols are published in the Cochrane Library using the “Flexible (Prognosis)” type. The Prognosis Methods Group recommends inclusion of specific sub-headers relevant to the type of prognostic review being undertaken. This document includes the recommended sub-headers for exemplar reviews of prognostic model(s). See at the end of this document relevant references that may be helpful when writing the protocol.

Header*	Description
Title	Choose preferably one of the following formats: Incidence of [outcome] within [time] in [population] [Prognostic factors] for predicting incidence of [outcome] in [population] Prediction of [outcome] in [population] using [prognostic factors] Prognostic models for predicting [outcome] in [population] Performance of [prognostic model] for predicting [outcome] in [population] Added/Incremental value of [prognostic factor] on top of [existing prognostic factors/prognostic model] for predicting [outcome] in [population]



Protocol Cochrane Review Prognosis Studies

	disease recurrence, or even lifelong incidence of certain outcome events.
Why it is important to do this review [Fixed, level 2 heading]	Explain the rationale for the review and why the prognosis questions being asked are important.
Objectives [Fixed, level 1 heading]	
Primary objectives [Optional, level 2 heading]	State the review question, including a table in the PICOTS format. (See Box 1 in the paper of Debray et al, BMJ 2017, and Table 1 of the CHARMS guidance Moons et al, PLOS Med 2014). The PICOTS format consists of the following elements: <ul style="list-style-type: none"> • Population—define the target population in which the overall prognosis or factor(s)/model(s) will be used. • Intervention (model/factor)—define the factor(s)/model(s) under review. • Comparator—if applicable, one can address competing factor(s)/model(s) for the factor(s)/model(s) under review.

Available via <http://methods.cochrane.org/prognosis>

Background [Fixed, level 1 heading]	
Description of the health condition and context [Fixed, level 2 heading]	A description of the targeted health condition and clinical context for which the (overall) prognosis or prognostic/predictive factor or model under review is intended (frequency, severity, and possible treatments). A health condition can for example be people undergoing surgery, having a certain disease or diagnosis, being pregnant, or healthy individuals of the general population within a certain age range. Also clearly define the moment of prognostication or prediction in the targeted population. For example, within two weeks after receiving a certain diagnosis, the day of intensive care admission, being 3 months pregnant, or visiting the emergency department with a trauma. If there are existing Cochrane reviews of interventions or diagnostic tests for the targeted health condition they should be cross-referenced here.
Description of the prognostic / predictive model(s) / factor(s) [Fixed, level 2 heading]	Not applicable for review on overall prognosis. Clearly state in which of the types of prognosis studies you are interested in: prognostic factor, prognostic model, or predictive factor (see PROGRESS series for definitions, see below for references). Describe the factor(s) or model(s) under review in more detail.
Health outcomes	Description of the health outcomes that are being studied in the targeted

Secondary objectives [Optional, level 2 heading]	Elaboration of the factor(s)/model(s). Reviews that investigate multiple prognosis questions may categorise their objectives as ‘Primary Objectives’ and ‘Secondary Objectives’. For example, the primary objectives may be to quantify the added predictive value of several biomarkers to an existing prognostic model; the secondary objective may be to compare the performance of this existing prognostic model to the performance of the biomarkers alone. Secondary objectives related to investigating heterogeneity between study results should not be listed under this subheading but under the next subheading.
Investigation of sources of heterogeneity between studies [Fixed, level 2 heading]	Heterogeneity investigations explore factors which may affect, e.g. the overall prognosis or the prognostic accuracy of factors or models. These explorations are essential because they provide a framework by which the observed heterogeneity may be explained a priori and to provide a more clinically useful review. For example, the predictive performance of a certain prognostic model for predicting 10-year cardiovascular disease outcomes in the adults above 40 in the general population, may vary when different definitions of cardiovascular disease outcomes are applied, when different age ranges, ethnic groups or genders have been studied, or when different study designs were used in the prognostic model studies.

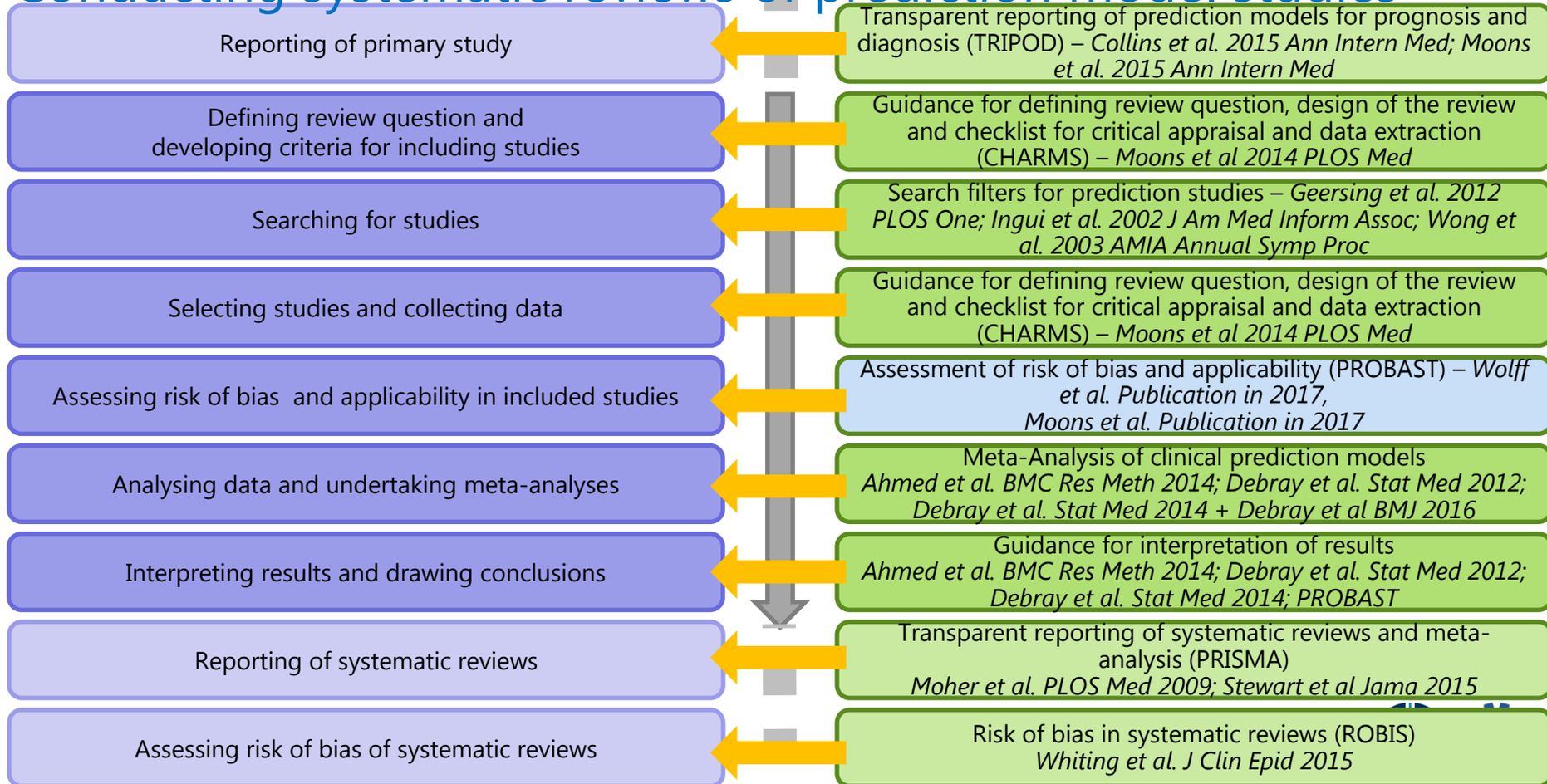


Take home messages

- 4 Main types of prognostic studies
- 3 Main types of prognostic model studies
- Systematic reviews of prognostic factor and model studies largely same as for intervention SRs
- Different and indeed more challenges in SRs of prognostic studies
- Tools available for all familiar steps of SR → prognostic studies



Conducting systematic reviews of prediction model studies



Reporting guideline prognostic studies

REporting recommendations for tumor MARKer prognostic studies (REMARK)

Lisa M. McShane · Douglas G. Altman · Willi Sauerbrei · Sheila E. Taube ·
Massimo Gion · Gary M. Clark for the Statistics Subcommittee of the
NCI-EORTC Working Group on Cancer Diagnostics

Annals of Internal Medicine RESEARCH AND REPORTING METHODS

Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis (TRIPOD): The TRIPOD Statement

Gary S. Collins, PhD; Johannes B. Reitsma, MD, PhD; Douglas G. Altman, DSc; and Karel G.M. Moons, PhD *Ann Intern Med.* 2015;162:55-63. doi:10.7326/M14-0697

Annals of Internal Medicine RESEARCH AND REPORTING METHODS

Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis (TRIPOD): Explanation and Elaboration

Ann Intern Med. 2015;162:W1-W73. doi:10.7326/M14-0698

Karel G.M. Moons, PhD; Douglas G. Altman, DSc; Johannes B. Reitsma, MD, PhD; John P.A. Ioannidis, MD, DSc;
Petra Macaskill, PhD; Ewout W. Steyerberg, PhD; Andrew J. Vickers, PhD; David F. Ransohoff, MD; and Gary S. Collins, PhD

www.tripod-statement.org



Example SRs of prognostic factor studies

Is P53 a prognostic (molecular) marker for bladder cancer (Malats, Lancet Oncol, 2005)

SR on all molecular and biological prognostic markers of tumours in the Ewing's Sarcoma Family (Riley Eur J Canc 2003)

Prognostic markers for death or tumour recurrence in patients with neuroblastoma (Riley, Br J Canc 2003)

Prognostic factors of sequelae and death after bacterial meningitis in childhood (de Jonge, BMC Infectious Diseases 2010)

Added value of carotid imaging markers in the prediction of fatal or non-fatal CVD events in general population (Peters, Heart 2011)



Examples SRs of prognostic modeling studies

Risk prediction models for the development of type 2 DM: SR (Collins 2011, Plos Med)

Prediction models for CVD in patients with type 2 DM (van Dieren, Heart 2011)

Risk prediction models for prolonged ICU stay after cardiac surgery (Ettema, Circulation 2010)

Risk prediction models for outcome after traumatic brain injury (Perel BMC Med Informatic and Decis Making 2006)

Prediction models for cardiovascular disease risk in the general population: systematic review (Damen, BMJ 2016)



Other workshops

- Systematic reviews of prognostic studies I: introduction, design and protocol for systematic reviews of prognostic studies (Sunday, September 16th, 11:00)
- Systematic reviews of prognostic studies II: risk of bias assessment in systematic reviews of prognostic studies (Sunday, September 16th, 16:00)
- Systematic reviews of prognostic studies III: meta-analytical approaches in systematic reviews of prognostic studies (Monday, September 17th, 11:00)
- Systematic reviews of prognostic studies IV: meta-analysis of prognostic studies using individual participant data (Tuesday, September 18th, 11:00).



Upcoming course

SR and MA van prognosis studies

- Medium – advanced level
- 3-day face-to-face course

