STARD 2015

Section & Topic	Item No	Item	Reported on Page Number/ Line Number	Reported on Section/ Paragraph
TITLE OR AB	OR ABSTRACT			
	_	Identification as a study of diagnostic accuracy using at least one measure of accuracy (such as sensitivity, specificity, predictive values, or AUC)	Page1-2/1ine1-3, 42-4 4	Title,Abstract/Para3
ABSTRACT				
	2	Structured summary of study design, methods, results, and conclusions (for specific guidance, see STARD for Abstracts)	Page1-3/1ine21-50	Abstract/Para1-4
INTRODUCTION	ON			
	ω	Scientific and clinical background, including the intended use and clinical role of the index test	Page4/line67-79	Introduction/Para1-2
	4	Study objectives and hypotheses	Page4/line82-85	Introduction/Para3
METHODS				
Study design	5	Whether data collection was planned before the index test and reference standard were performed (prospective study) or after (retrospective study)	Page5/line96-98	Methods/Para2
Participants	6	Eligibility criteria	Page5/line96-98	Methods/Para2
	7	On what basis potentially eligible participants were identified (such as symptoms, results from previous tests, inclusion in registry)	Page5/line96-103	Methods/Para2
	8	Where and when potentially eligible participants were identified (setting, location and dates)	Page5/line96-98	Methods/Para2
	9	Whether participants formed a consecutive, random or convenience series	Page5/line96-98	Methods/Para2
Test	10a	Index test, in sufficient detail to allow replication	Page6/1ine112-128	Methods/Para3
methods	10b	Reference standard, in sufficient detail to allow replication	Page9/line188-193	Methods/Para10
	1	Rationale for choosing the reference standard (if alternatives exist)	N/A	N/A
	12a	Definition of and rationale for test positivity cut-offs or result categories of the index test, distinguishing pre-specified from exploratory	Page7-8, 10/line138-1 75, 209-213	Methods/Para6-8,11
	12b	Definition of and rationale for test positivity cut-offs or result categories of the reference standard, distinguishing pre-specified from exploratory	Page9/1ine190-193	Methods/Para10
	13a	Whether clinical information and reference standard results were available to the performers/readers of the index test	Page8-9/line176-180	Methods/Para9
	13b	Whether clinical information and index test results were available to the assessors of the reference standard	Page9/line186-188	Methods/Para10

Acknowledgments	Page17-18/11ne372-37	Sources of funding and other support; role of funders	30	
Online appendix	Page2-3/line23-59	Where the full study protocol can be accessed	29	
Methods/Para1	Page5/line91-93	Registration number and name of registry	28	
		ION	ORMATI	OTHER INFORMATION
Discussion/Para5	Page16/1ine336-348	Implications for practice, including the intended use and clinical role of the index test	27	
Discussion/Para6	Page16-17/11ne349-36	Study limitations, including sources of potential bias, statistical uncertainty, and generalisability	26	
			Z	DISCUSSION
Results/Para1	Page11/1ine235-236	Any adverse events from performing the index test or the reference standard	25	
Results/Para5	Page13/1ine268-273	Estimates of diagnostic accuracy and their precision (such as 95% confidence intervals)	24	
Results/Para3	Page12/1ine261-264	Cross tabulation of the index test results (or their distribution) by the results of the reference standard	23	Test results
Results/Para3	Page12/1ine247-252	Time interval and any clinical interventions between index test and reference standard	22	
Results/Para4	Page12–13/11ne256–26 5	Distribution of alternative diagnoses in those without the target condition	21b	
Results/Para3		Distribution of severity of disease in those with the target condition	21a	
Results/Para1	Page11/1ine233-235	Baseline demographic and clinical characteristics of participants	20	
Results/Para1	Page11/1ine228-234	Flow of participants, using a diagram	19	Participants
				RESULTS
Methods/Para2	Page5/line96-98	Intended sample size and how it was determined	18	
N/A	N/A	Any analyses of variability in diagnostic accuracy, distinguishing pre-specified from exploratory	17	
Methods/Para2	Page5/line103-108	How missing data on the index test and reference standard were handled	16	
Methods/Para9	Page9/1ine178-181	How indeterminate index test or reference standard results were handled	15	
Methods/Paral1	Page10/1ine213-214	Methods for estimating or comparing measures of diagnostic accuracy	14	Analysis

ΑM

Authors can use the list to write informative study reports. Editors and peer-reviewers can use it to evaluate whether the information has been included in manuscripts submitted for publication STARD stands for "Standards for Reporting Diagnostic accuracy studies". This list of items was developed to contribute to the completeness and transparency of reporting of diagnostic accuracy studies.

Explanation

other method for collecting information about the current health status of a patient benefit from therapy, or an event or condition in the future. A medical test can be an imaging procedure, a laboratory test, elements from history and physical examination, a combination of these, or any A diagnostic accuracy study evaluates the ability of one or more medical tests to correctly classify study participants as having a target condition. This can be a disease, a disease stage, response or

An accuracy study can rely on one or more reference standards comparing the distribution of the index test results with those of the reference standard. The reference standard is the best available method for establishing the presence or absence of the target condition. The test whose accuracy is evaluated is called index test. A study can evaluate the accuracy of one or more index tests. Evaluating the ability of a medical test to correctly classify patients is typically done by

estimates of accuracy can then be calculated to quantify the statistical precision of the measurements proportion of participants with the target condition who have a positive index test), and its specificity (the proportion without the target condition who have a negative index test). From this cross tabulation If test results are categorized as either positive or negative, the cross tabulation of the index test results against those of the reference standard can be used to estimate the sensitivity of the index test (the (sometimes referred to as the contingency or "2x2" table), several other accuracy statistics can be estimated, such as the positive and negative predictive values of the test. Confidence intervals around

a single numerical value about the overall diagnostic accuracy of the index test a receiver operating characteristic (ROC) curve which graphically represents the combination of sensitivity and specificity for each possible test positivity cut-off. The area under the ROC curve informs in If the index test results can take more than two values, categorization of test results as positive or negative requires a test positivity cut-off. When multiple such cut-offs can be defined, authors can report

pathway. A replacement test, for example, replaces an existing test. A triage test is used before an existing test; an add-on test is used after an existing test The intended use of a medical test can be diagnosis, screening, staging, monitoring, surveillance, prediction or prognosis. The clinical role of a test explains its position relative to existing tests in the clinical

such as staging or prognosis. The STARD list was not explicitly developed for these other outcomes, statistics, and study types, although most STARD items would still apply. Besides diagnostic accuracy, several other outcomes and statistics may be relevant in the evaluation of medical tests. Medical tests can also be used to classify patients for purposes other than diagnosis,

DEVELOPMENT

list represents an update of the first version, which was published in 2003. More information can be found on http://www.equator-network.org/reporting-guidelines/stard select items that, when reported, would help readers to judge the potential for bias in the study, to appraise the applicability of the study findings and the validity of conclusions and recommendations. The This STARD list was released in 2015. The 30 items were identified by an international expert group of methodologists, researchers, and editors. The guiding principle in the development of STARD was to

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*As the checklist was provided upon initial submission, the page number/line number reported may be changed due to copyediting and may not be referable in the published version. In this case, the section/paragraph may be used as an alternative reference