Toward Alignment in the Reporting of Economic Evaluations of Diagnostic Tests and Biomarkers: The AGREEDT Checklist

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Objectives. General frameworks for conducting and reporting health economic evaluations are available but not specific enough to cover the intricacies of the evaluation of diagnostic tests and biomarkers. Such evaluations are typically complex and model-based because tests primarily affect health outcomes indirectly and real-world data on health outcomes are often lacking. Moreover, not all aspects relevant to the evaluation of a diagnostic test may be known and explicitly considered for inclusion in the evaluation, leading to a loss of transparency and replicability. To address this challenge, this study aims to develop a comprehensive reporting checklist.

Methods. This study consisted of 3 main steps: 1) the development of an initial checklist based on a scoping review, 2) review and critical appraisal of the initial checklist by 4 independent experts, and 3) development of a final checklist. Each item from the checklist is illustrated using an example from previous research.

Results. The scoping review followed by critical review by the 4 experts resulted in a checklist containing 44 items, which ideally should be considered for inclusion in a model-based health economic evaluation. The extent to which these items were included or discussed in the studies identified in the scoping review varied substantially, with 14 items not being mentioned in ≥47 (75%) of the included studies.

Conclusions. The reporting checklist developed in this study may contribute to improved transparency and completeness of model-based health economic evaluations of diagnostic tests and biomarkers. Use of this checklist is therefore encouraged to enhance the interpretation, comparability, and indirectly—the validity of the results of such evaluations.

Keywords
approval, biomarkers, checklist, diagnostic test, health economic evaluation, reporting

Detailed evaluation of the clinical utility and also health economic impact of new diagnostic tests prior to their implementation in clinical practice is important to limit overuse of tests, ensure benefits to patients, and support efficient use of health care resources. Different frameworks have been developed for the phased evaluation of diagnostic tests. All these frameworks recognize that after evaluating the safety, efficacy, and accuracy of a diagnostic test, the impact of this test on health outcomes and costs should be determined. Evaluating tests in randomized controlled trials (RCTs), however, is often not feasible for ethical, financial, or other reasons, particularly in early test development stages. Indeed, RCTs evaluating the impact of diagnostic tests on patient outcomes are rare. As an alternative, methods to develop decision-analytic models for the health economic evaluation of diagnostic tests, synthesizing all available evidence from different sources, have long been
available.\textsuperscript{6,12–16} It is widely recognized that such models are a useful and valid alternative to evaluate the impact of new health technologies in general\textsuperscript{17,18} and diagnostic methods in particular.\textsuperscript{12,14}

However, the comprehensive evaluation of the impact of new tests is typically much more complex than, for example, evaluation of the impact of new drugs. Among others, this is due to the indirect impact of tests on health outcomes by improved patient management (also referred to as “clinical utility”\textsuperscript{19}), the use of combinations and sequences of tests in clinical practice (depending on previous test results), and the often complex interpretation of test outcomes. In practice, model-based impact evaluations of tests therefore actually involve the evaluation of diagnostic testing strategies (i.e., test-treatment combinations).

Owing to the complexity of these diagnostic testing strategies, many model-based impact evaluations of tests make use of simplified models that do not incorporate all aspects of clinical practice. Simplified models are used because 1) evidence regarding all aspects involved in health economic test evaluations might be lacking, 2) inclusion of all aspects likely increases model complexity, or 3) researchers may not be aware of all aspects of test evaluation. For example, it is often not reported how the incremental effect of a new test, when used in combination with other tests, is determined and how the correlation between the outcomes of these different tests (applied solo or in sequence) is handled.\textsuperscript{20–23} Similarly, the selection of patients in whom the test is performed, the consequences of incidental findings (also referred to as chance findings), and the occurrence of test failures or indeterminate test results are often not reported.\textsuperscript{24–26}

Although simplifications of the decision-analytic models used for such evaluations may sometimes be necessary and can be adequately justified, implicit simplification due to unawareness of all relevant evaluation aspects or without proper justification may lead to nontransparent and incorrect evaluation results.

General frameworks and guidelines regarding which aspects to include in decision-analytic modeling and how to report modeling outcomes are available\textsuperscript{27–29} but not specific enough to cover the complexities of diagnostic test evaluation. Furthermore, previous research into (aspects of) diagnostic test evaluation mostly focused on specific diseases or on specific types or combinations of diagnostic tests.\textsuperscript{23,30–35} A generic and comprehensive overview of all potentially relevant aspects in health economic evaluation of diagnostic tests and biomarkers that may be used to guide such evaluations is currently lacking.

The purpose of this article is, therefore, to provide such an overview as a generic checklist, intended to be applicable to all types of diagnostic tests and not specific to a single disease or condition or subgroup of individuals. Thereby, this checklist aims to allow researchers to explicitly consider all aspects potentially relevant to the health economic evaluation of a specific test, from a societal perspective. Therefore, this checklist is referred to as the “AGREEDT” checklist, which is an acronym of “AliGnment in the Reporting of Economic Evaluations of Diagnostic Tests and biomarkers.” Use of the checklist does not need to complicate such evaluations, as some aspects described may not be relevant to particular evaluations, but rather suggests that choices to exclude certain aspects are adequately justified.

Methods

This study consisted of 3 main steps: 1) the development of an initial checklist based on a scoping review, 2) review and critical appraisal of the initial checklist by 4 experts (CEP, MCW, MH, and TM) not involved in the scoping review, and 3) development of a final checklist based on the review by experts. Finally, each item from the checklist is illustrated using an example from previous research.

Scoping Review

In the past decades, hundreds of model-based health economic evaluations of diagnostic tests have been published, across a wide range of medical contexts. A still narrow literature search in PubMed in January 2017 resulted in a total of 1844 articles using the following
combinations of search terms in title and abstract: (health economic OR cost-effectiveness) AND diag* AND (model OR Markov OR tree OR modeling OR modelling). Besides the large number of studies that have been published in this field, systematic identification of health economic evaluations is found to be challenging.\textsuperscript{36} This is partly caused by the multitude of MeSH terms in PubMed related to diagnostic strategies (over 48 MeSH terms exist that include the word diagnostic or diagnosis). Because of these challenges and the fact that different evaluations are very likely to include and exclude the same aspects, a scoping review was performed instead of a systematic literature review, followed by critical appraisal by 4 independent experts. A key strength of a scoping review is that it can provide a rigorous and transparent method for mapping areas of research,\textsuperscript{37} particularly when an area is complex or has not been reviewed comprehensively before.\textsuperscript{38}

This scoping review was performed in PubMed in January 2017, searching for the following combination of search terms in the title of the article: (health economic OR cost-effectiveness) AND diag* AND (model OR Markov OR tree OR modeling OR modelling) NOT diagnosed. The term NOT diagnosed was added to prevent retrieving many articles including patients who are already diagnosed with a certain condition, instead of focusing on the diagnostic process itself. The search was limited to articles published in English or Dutch. Studies were excluded, based on title and abstract, if they did not concern original research or did not evaluate the cost-effectiveness of the use of 1 or more tests (regardless of the effectiveness measure, for example, additional cost per additional correct diagnosis or per additional quality-adjusted life year). In addition, as guidelines for performing health economic evaluations continue to be updated,\textsuperscript{39–41} it was expected that the more recent studies would provide the most comprehensive overview of all potentially relevant items that need to be included in the checklist. To check this assumption, the PubMed search was repeated without limiting the search to studies published ≤5 years ago, resulting in 128 additional articles. Following this, 2 articles that were published >5 years ago were randomly selected.\textsuperscript{42,43} A thorough review of both articles did not result in any additional relevant items for inclusion in the checklist. Therefore, the search was limited to articles published in the past 5 years. One author screened studies for exclusion (MMAK) and consulted with a second author (HK) if necessary.

Design of the reporting checklist

All articles resulting from the scoping review were searched for items related to model-based health economic test evaluation of diagnostic tests that were either included explicitly in the evaluation, or that were only mentioned but not included (mostly in the introduction or discussion sections). Generic items, not specific to diagnostic test evaluation were not included in the new checklist as these are already covered in existing checklists. Examples of such generic items include choosing the time horizon and perspective of the evaluation.\textsuperscript{27–29} However, some overlap remains as the checklist does include items which are considered applicable to diagnostic test evaluation that are only covered partially or at a high level in existing guidelines.

A thorough screening of all articles was performed by MMAK resulting in an initial list of aspects considered to be potentially relevant. As the checklist was intended to provide a comprehensive overview of all potentially relevant aspects, all of these aspects were added to the checklist, unless it was considered to be already included in currently available guidelines (based upon agreement between MMAK and HK). The definition of each aspect was based on agreement between MMAK and HK.

Critical Appraisal and Validation of the Reporting Checklist

As diagnostic tests and imaging are used for a large variety of (suspected) medical conditions, an expert panel with a broad field of experience was required for critical appraisal of the checklist. Therefore, the expert panel was composed in such a way that at least 1 expert was experienced in each of the different areas of interest (i.e., biomarkers or imaging) and in each of the different purposes of diagnostic testing (i.e., diagnosis, screening, monitoring, and prognosis). In addition, to maximize the likelihood that the final checklist is generalizable to different countries and settings, the experts chosen lived on 3 different continents. Four experts were invited (CEP, MCW, MH, and TM) to participate via email, and none of them declined.

The initial checklist was critically appraised and validated independently by all 4 experts, who received the checklist via email. They were asked to provide individual, qualitative judgments on whether all items in this list were clear and unambiguous, to indicate any missing or redundant items in this list, and to provide suggestions for further improvement.

Finalization of the Reporting Checklist

Based on the experts’ suggestions, several changes were made to the reporting checklist. Those changes involved
the rewording of items, removal of redundant items, and
the addition of missing items to the checklist. As this
checklist is intended to provide an exhaustive list of all
aspects relevant to the health economic evaluation of
diagnostic tests and biomarkers, all suggestions for the
addition of missing items were adopted. All changes
made to the checklist were decided upon agreement
between MMAK and HK (for a full description, see
online Appendix 1). The revised checklist was again criti-
cally appraised by all authors and agreed upon. Finally,
the articles included in the scoping review were reread by
MMAK to assess whether the final checklist items were
included or mentioned.

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This study was not funded.

Results

Results of the Scoping Review
The literature search resulted in 77 articles that were
screened for inclusion in the scoping review, of which 14
articles were excluded. Of these, 4 articles did not specifi-
cally evaluate the cost-effectiveness of a (combination of)
diagnostic test(s), 2 concerned a letter to the editor, and 7
articles focused on methodological aspects of the evalua-
tion of diagnostic strategies (e.g., in the context of single
disease, or on specific types or combinations of diagnos-
tic tests, as mentioned earlier). In addition, 1 article was
excluded because the full text could not be obtained or
purchased by the university library, from online data-
bases, from the website of the publisher, or by contacting
the authors. This resulted in a total of 63 studies that
were included in the scoping review. An overview of this
selection process is provided in Figure 1.

A critical evaluation of the 63 articles resulted in an
initial list of 29 items. These items were divided into 6
main topics: 1) time to presentation of the individual to
the health professional (i.e., the clinical starting point),
2) use of diagnostic tests, 3) test performance and char-
acteristics, 4) patient management decisions, 5) impact
on health outcomes and costs, and 6) wider societal
impact, which may accrue to patients, their families,
and/or health care professionals. This societal impact,
for example, may concern the impact on caregivers (in
terms of time spent on hospital visits and caregiving and
the accompanying impact on productivity), on the health
system or health professional (e.g., in terms of reduced
patient visits), or on society (e.g., measures that aim to
prevent widespread antibiotic resistance). Quantifying
these aspects may provide a broader view on the poten-
tial impact of diagnostic testing.

Critical Appraisal and Validation of the
Reporting Checklist
Following the critical appraisal of experts, the list was
updated, with 1 item being removed and 15 items being
added; 1 additional item was added based on the sugges-
tion of a reviewer of the manuscript during the submis-
sion process. Finally, this resulted in a reporting checklist
consisting of 44 items, as shown in Table 1. Of these 16
added items, 8 involved a further specification of the
tests’ diagnostic performance, as included below item 3.2
in the checklist. The item that was removed concerned
the generalizability of the results, which was considered
not specific to diagnostic test evaluations. The full report-
ing checklist, including an overview in which of the stud-
ies from the scoping review each of the items was
included or considered, as well as an example for each of
the items, is provided in online Appendix 2. An overview
of this process, including the scoping review and the
Table 1 Reporting Checklist to Indicate Which Items Were Included in the Health Economic Evaluation of Diagnostic Tests and Biomarkers

| Items of the Evaluation of Diagnostic Tests and Biomarkers | Included in Evaluation |
|------------------------------------------------------------|------------------------|
| **Time to presentation**                                  |                        |
| *Onset of complaints or onset of suspicion by physician and start of diagnostic trajectory* |                        |
| 1.1 The study should contain a description of the individuals who enter the diagnostic pathway (i.e., a patient develops a [new] condition or disease, which may or may not result in symptoms or complaints, or undergoes diagnostic testing as part of screening or genetic testing). |                        |
| 1.2 Consider the time to start of the diagnostic trajectory or the time until a monitoring or screening test is (repeatedly) performed (initiated by symptoms/complaints or initiated as part of regular screening or monitoring). |                        |
| *(The time between 1.1 and 1.2 is the time during which individuals are at risk of complications from disease and progression, in the absence of a diagnosis and thus also in the absence of treatment.)* |                        |
| **Use of diagnostic tests**                               |                        |
| *Decision regarding which diagnostic test(s) is/are performed, in which patients, and in what order* |                        |
| 2 Specify for which purpose(s) the test is used (e.g., screening, diagnosing, monitoring, guide dosage, commencement or cessation of therapy, triaging, staging, prognostic) and define the entire diagnostic and clinical pathway. |                        |
| 2.1 Consider whether more than 2 (possible) diagnostic strategies can be compared, each involving a single test or combination of tests. |                        |
| 2.2 Consider whether the evaluated diagnostic strategies include multiple tests, which can be performed in parallel or in sequence. |                        |
| 2.2.1 Consider whether some tests of the diagnostic workup are performed conditional on previous test outcomes, leading to a selection of patients undergoing specific tests. |                        |
| 2.3 Consider whether subgroups can be defined based on explicit criteria or patient characteristics, in which different tests would be performed (not solely dependent on previous test outcomes). |                        |
| 2.4 Consider whether different tests are applied based on implicit (shared) decision making (e.g., perceived condition or risk, or symptom presentation). |                        |
| **Test performance and characteristics**                  |                        |
| *Diagnostic test performance and items related to the sampling and testing* |                        |
| 3.1 Specify the costs of the diagnostic test. |                        |
| 3.2 Specify test performance, in terms of sensitivity, specificity, negative predictive value (and its complement), and/or positive predictive value (and its complement), either or not combined with a decision rule or algorithm. |                        |
| 3.2.1 Describe the evidence base for the estimated test performance. |                        |
| 3.2.2 Describe the positivity criterion (i.e., cutoff value) applied to the test or testing strategy. |                        |
| 3.2.3 Consider whether the estimated test performance may be biased, for example, due to lack of evidence on conditional dependence or independence, lack of a (perfect) gold standard (i.e., classification bias), verification bias, analytic bias, spectrum bias, diagnostic review bias, and incorporation bias. |                        |
| 3.2.3.1 Consider how likely/to what extent bias in the available/applied evidence affects the estimated test performance. |                        |
| 3.2.4 Describe how uncertainty/variation in the test performance (receiver operating characteristic [ROC] curve) was handled or explained, for example, due to interrater and intrarater reliability, or experience of the clinician. |                        |
| 3.2.5 Describe the logic, or analysis, applied to choose the cutoff value (i.e., the point on the ROC curve) for the test, for example, depending on whether the test is used as a single test or part of a sequence of tests. |                        |
| 3.2.6 Describe whether different test performances and cutoff values were considered for different subgroups of patients and/or environmental characteristics. For example: based on specific subgroup(s) of patients, timing of the test in the diagnostic trajectory, or selection of patients based on previous test outcomes (if any). |                        |
| 3.2.7 Consider whether test performance is dependent on disease prevalence (which also includes the impact of spectrum bias on disease prevalence and, as a consequence, on test performance) or affected by other patient characteristics or conditions. |                        |
| 3.2.8 Consider whether test performance is based on a combination of tests (and on a combination of areas under the ROC curves for each test). |                        |

(continued)
Table 1 (continued)

| Items of the Evaluation of Diagnostic Tests and Biomarkers | Included in Evaluation |
|------------------------------------------------------------|------------------------|
| 3.3 Consider the feasibility of obtaining (sufficient) sample and/or usability of the sample that is obtained. | Yes\(^a\) No\(^b\) |
| 3.4 Consider the occurrence of test failures or indeterminate/not assessable results. | Yes\(^a\) No\(^b\) |
| 3.5 Consider costs of retesting (after obtaining insufficient/unsuitable sample or after test failure or indeterminate/not assessable result). | Yes\(^a\) No\(^b\) |
| 3.6 Consider complications, risks, or other negative/positive aspects directly related to obtaining the sample and/or performing the diagnostic test (either in the intervention or in the control strategy). | Yes\(^a\) No\(^b\) |
| 3.7 Consider the time taken to perform the test (including waiting time) until the test result is available or until a management decision or treatment is initiated based on this test result (either in the intervention or in the control strategy). | Yes\(^a\) No\(^b\) |
| 3.8 Consider the impact of additional knowledge gained by performing the diagnostic test (i.e., for a genetic test) or the occurrence and impact of incidental findings (i.e., the unintentional discovery of a previously undiagnosed condition during the evaluation of another condition). | Yes\(^a\) No\(^b\) |
| 3.8.1 The impact of incidental findings on performing additional tests is addressed. | Yes\(^a\) No\(^b\) |

Patient management decisions

**Impact of a test on the diagnosis and/or patient management strategy (based on this diagnosis)**

4.1 Clearly specify the impact of the test in selecting the patient management strategy. | Yes\(^a\) No\(^b\) |
4.2 Consider whether other aspects besides test results themselves are part of the decision algorithm (and included in the evaluation). These may involve a shared decision-making process of the physician with patients/relatives or aspects including coverage or physician adherence to treatment guidelines. | Yes\(^a\) No\(^b\) |
4.3 Consider whether the impact of the test result on resulting/selected diagnosis or management strategy varies across subgroups (this difference should not only be caused by differences in diagnostic performance of the test and does not need to include the impact on costs and/or health outcomes within this subgroup). | Yes\(^a\) No\(^b\) |
4.4 Consider the consistency of test results over time (e.g., genetic mutations may be affected by treatment prescribed after the initial diagnosis). | Yes\(^a\) No\(^b\) |
4.5 Consider the impact of performing the test and providing and interpreting the result on the time spend/capacity of the health care professional(s) or the patient. | Yes\(^a\) No\(^b\) |

Impact on health outcomes and costs

**Impact of the patient management strategy on diseased and nondiseased individuals, in terms of health outcomes and costs**

5 Evaluate the direct impact of the chosen patient management strategy on the number of (in)correctly diagnosed individuals, health outcomes, and/or costs. | Yes\(^a\) No\(^b\) |
5.1 Consider the direct impact of the chosen patient management strategy on health outcomes and/or costs. This concerns the entire period in which patient management may affect a patient’s health and/or costs and does not only involve the testing strategy itself. | Yes\(^a\) No\(^b\) |
5.2 Consider whether the direct impact of the chosen patient management strategy on health outcomes and/or costs varies across subgroups. (This does not include only varying the incidence of a certain condition in a sensitivity analysis. The subgroups should be clearly defined and preferably be identifiable based on patient characteristics.\(^c\)) | Yes\(^a\) No\(^b\) |
5.3 Consider patient’s adherence to treatment (which includes aspects that may indicate (partial) nonadherence (e.g., following only some of the treatment recommendations, as well as aspects that affect the degree of administration of treatment)). | Yes\(^a\) No\(^b\) |
5.4 Consider the occurrence (and consequences) of treatment-related adverse events. | Yes\(^a\) No\(^b\) |
5.5 Describe the probability or time it takes to observe that the patient management strategy proves to be effective over time or that the patient cures spontaneously (regardless of whether the patient received a correct or an incorrect diagnosis). | Yes\(^a\) No\(^b\) |
5.6 Describe the probability or time it takes to repeat or extend the diagnostic workup when the patient management strategy proves to be ineffective, either directly or over time (regardless of whether the patient received a correct or an incorrect diagnosis). This also includes the situation in which the patient receives no treatment or unnecessary treatment. | Yes\(^a\) No\(^b\) |
5.7 Describe the impact of ineffective or unnecessary treatment or management on health outcomes and/or costs (including both side effects and costs and regardless of whether the patient received a correct or an incorrect diagnosis). This also includes the situation in which incorrectly no treatment is provided or in which the treatment is delayed. | Yes\(^a\) No\(^b\) |

\(^{a}\) \(^{b}\) \(^{c}\)
critical appraisal by the experts, is shown in Figure 1. The final list of items in this reporting checklist, in chronological order from the start of the diagnostic trajectory and onward, is illustrated in Figure 2.

Results indicate that health economic evaluations of diagnostic tests or biomarkers differ considerably in the items that have been explicitly included (or considered for inclusion) in the corresponding decision-analytic model (Table 1). Some of the items from the checklist were only included (or considered) in a few studies from the scoping review. For example, the impact of incidental findings on performing additional tests, the consistency of test results over time, and the impact of test outcomes on relatives themselves were each only addressed in 3 of the 63 included studies. These items may not have been included in other studies because they were considered not relevant to the specific context, because (scientific) evidence was lacking, or because these items were not considered due to unawareness of their relevance by the authors.

Discussion

Strengths

A strength of this study is that it combines evidence from multiple sources, including a review of literature, as well as a validation by experts. As the items included in the checklist are defined in general terms and not limited to specific diseases, tests, care providers, or patient management strategies, this reporting checklist can potentially be useful in performing and appraising health economic evaluations worldwide and across a broad spectrum of (novel) diagnostic technologies. In addition, as this checklist specifically focuses on health economic evaluations of diagnostic tests or biomarkers, an area for which no reporting checklists are yet available, it may be a useful extension to existing reporting checklists, such as the Consolidated Health Economic Evaluation Reporting Standards (CHEERS) checklist.27 Finally, use of this checklist can also support development of health economic models through increased awareness of all potentially relevant evaluation aspects.

In addition, use of this checklist does not necessarily require more resources to be allocated to the evaluation or increase the complexity of the resulting decision-analytic model. In general, deliberation on the relevance of all aspects is key, and aspects may be excluded from the evaluation whenever this can be adequately justified. For example, when evaluating the cost-effectiveness of a new point-of-care troponin test used by the general practitioner compared to an existing, older point-of-care troponin test (in the context where the new test would replace the old test), aspects such as

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Table 1 (continued)

| Items of the Evaluation of Diagnostic Tests and Biomarkers | Included in Evaluation |
|----------------------------------------------------------|-----------------------|
| 5.7.1 Consider the impact of delay in treatment initiation on health outcomes and/or costs. | Yes\(^a\) No\(^b\) |

\(^a\)If an item is included in the quantitative analysis, indicate the corresponding model parameter(s) and evidence source(s).
\(^b\)If an item is excluded from the quantitative analysis, please explain why the exclusion was necessary.

Existing guidelines indicate that subgroup analyses are relevant when different strategies are likely to be (sub)optimal in different subgroups. Subgroup specific analyses can then be performed to address multiple decision problems. Here we consider scenarios where different tests may be used in different subgroups, depending on patient characteristics or previous test outcomes.
Figure 2 Overview of steps in diagnostic trajectory. This figure gives a conceptual outline of the steps involved in the diagnostic trajectory, in chronological order from top to bottom. The numbers shown at the several steps correspond to the item numbers presented in Table 1. The dashed lines represent steps of which the duration may vary substantially, for example, the time between symptom onset and presentation to a clinician (which may vary from minutes in case of severe symptoms to years for mild and gradually developing conditions). The arrows indicate situations in which either the diagnostic test (result) was not usable or indeterminate (items 3.3–3.5) or situations in which the treatment proves to be ineffective (items 5.5–5.7). As this may be caused by an incorrect diagnosis, the patient may undergo a subsequent round of diagnostic testing and (possibly) treatment. Alternatively, the diagnosis may be correct but the treatment incorrect, in which an alternative treatment may be initiated. *Although the (wider) societal impact of diagnostic testing often involves long-term effects, these effects may sometimes also become apparent in the short term. Dx, diagnostic test; Tx, treatment.
“time to start of the diagnostic trajectory” and “purpose of the test” will not differ between both strategies. In addition, “complication risks” associated with taking the blood sample (in both point-of-care tests) are likely extremely small, which could justify excluding these aspects from the analysis.

**Limitations**

Performing a systematic literature review was considered not possible given the large number of published economic evaluations of diagnostic tests. Therefore, a scoping review was performed instead by 1 reviewer. As the judgment regarding whether an aspect was incorporated in a health economic evaluation was sometimes found to be difficult, it cannot be excluded that these judgments may have differed slightly when performed by a different reviewer. In addition, as the decision to limit the search strategy to the past 5 years was based on reviewing 2 studies published >5 years ago, this small sample (i.e., 1.6% of studies published >5 years ago) cannot rule out the possibility that items have been missed by excluding all older studies. Also, the scoping review may have been subject to publication bias, as it may have omitted potentially relevant aspects from unpublished studies, as well as from method manuals (including those focusing on economic evaluations of other interventions or technologies in health care). Despite the abovementioned limitations, the critical review of the checklist by 4 independent experts from different countries makes it unlikely that important items have been missed.

In addition, the expert appraisal resulted in the addition of 16 items to the checklist. Although this may seem to be a large extension to the items already identified in the scoping review, 8 of these added items actually involved a further specification (i.e., a subitem) of the test’s diagnostic performance. It was found useful to further specify “test performance” (i.e., item 3.2, which initially integrated several performance measures) into 8 subitems to further increase the transparency and comparability of health economic test evaluations.

**Implications for Practice**

This study was intended to design a reporting checklist without formulating a quality judgment of the studies included in the scoping review, based on which items of the checklist they did or did not incorporate. Furthermore, some items may have been included *implicitly* in the health economic evaluations identified in the scoping review, which could thus not be identified by the reviewer. As scientific articles are often restricted in their length, there may often be insufficient space to mention the inclusion (or justified exclusion) of each of the items from this checklist. In these situations, authors are recommended to describe their use of this checklist in an appendix. More specifically, authors are recommended to describe which items from the checklist they included in their evaluation and what evidence was used to inform them. Furthermore, they are recommended to explicitly state the reason(s) for excluding checklist items from their evaluation. Although it may be considered time-consuming to consider all 44 items of this checklist, it should be noted that most of these items are actually subitems, which do not need to be considered if the overarching (higher-level) item is (justifiably) excluded from the evaluation.

In addition, it should be noted that not all items in this checklist can be considered of equal importance. For example, diagnostic performance will typically have a larger impact on health outcomes and costs compared to considering the occurrence of test failures or the consistency of test results over time. However, this checklist is designed to provide an exhaustive overview of all potentially relevant items, regardless of importance. Therefore, use of this checklist will likely increase the chance that all relevant aspects will be included in health economic evaluations of diagnostic tests and biomarkers. Ultimately, it is up to the researchers to make a justifiable decision on which items to incorporate and which to exclude.

Finally, experiences regarding the use of this reporting checklist in practice may be valuable to further enhance its completeness and usability. Furthermore, given the rapid methodological developments in the field of health economic evaluation of diagnostic tests, regular updating of this checklist may be warranted.

**Conclusion**

Given the complexity and dependencies related to the use of diagnostic tests or biomarkers, researchers may not always be fully aware of all the different aspects potentially influencing the result of a model-based health economic evaluation. The use of the reporting checklist developed in this study may remedy this by increasing awareness of all potentially relevant aspects involved in such model-based health economic evaluations of diagnostic tests and biomarkers and thereby also increase the transparency, comparability, and—indirectly—the validity of the results of such evaluations.
Authors’ Note
This research was conducted at the department of Health Technology and Services Research, University of Twente, Enschede, the Netherlands.

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