AMSTAR is a reliable and valid measurement tool to assess the methodological quality of systematic reviews

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Abstract

Objective: Our purpose was to measure the agreement, reliability, construct validity, and feasibility of a measurement tool to assess systematic reviews (AMSTAR).

Study Design and Setting: We randomly selected 30 systematic reviews from a database. Each was assessed by two reviewers using: (1) the enhanced quality assessment questionnaire (Overview of Quality Assessment Questionnaire [OQAQ]); (2) Sacks’ instrument; and (3) our newly developed measurement tool (AMSTAR). We report on reliability (interobserver kappas of the 11 AMSTAR items), intraclass correlation coefficients (ICCs) of the sum scores, construct validity (ICCs of the sum scores of AMSTAR compared with those of other instruments), and completion times.

Results: The interrater agreement of the individual items of AMSTAR was substantial with a mean kappa of 0.70 (95% confidence interval [CI]: 0.57, 0.83) (range: 0.38–1.0). Kappas recorded for the other instruments were 0.63 (95% CI: 0.38, 0.78) for enhanced OQAQ and 0.40 (95% CI: 0.29, 0.50) for the Sacks’ instrument. The ICC of the total score for AMSTAR was 0.84 (95% CI: 0.65, 0.92) compared with 0.91 (95% CI: 0.82, 0.96) for OQAQ and 0.86 (95% CI: 0.71, 0.94) for the Sacks’ instrument. AMSTAR proved easy to apply, each review taking about 15 minutes to complete.

Conclusions: AMSTAR has good agreement, reliability, construct validity, and feasibility. These findings need confirmation by a broader range of assessors and a more diverse range of reviews. © 2009 Elsevier Inc. All rights reserved.

Keywords: Systematic reviews; Meta-analysis; Methodological quality; Validity; Reliability; Feasibility

1. Background

Systematic reviews have become the standard approach in assessing and summarizing applied health research, but the quality of systematic reviews has received relatively little attention. Quality can be defined as the likelihood that the design of a systematic review will generate unbiased results [1].

Systematic reviews have appeared in medical journals since the late 1970s. Thousands of systematic reviews are available on all areas of health care, and a substantial portion of them has been produced by the Cochrane Collaboration. High methodological quality is a pre-requisite for valid interpretation and application of review findings. However, systematic reviews are complex exercises, and assessing quality can be a daunting task. Clinicians and policy makers require guidance, which is not provided adequately by the available literature on the quality of systematic reviews. In a previous study, we summarized this literature, tested quality assessment tools, and reached the conclusion that current instruments for conducting methodological quality assessments of systematic reviews were suboptimal and needed revision and updating [2]. No single instrument has achieved dominance in terms of general use.

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One popular instrument (QUality Of Reporting Of Meta-analyses [QUORUM]) is a reporting checklist, not a methodological quality assessment instrument.

Our review revealed a variety of weaknesses in the available instruments [2]. Our intention was not to come up with a truly novel approach, but to bring clarity to the field by: reviewing the available instruments, further developing and updating existing instruments, and providing a model that was validated and useable (in terms of comprehensibility and acceptable time for completion). Based on this evaluation, we created a measurement tool to assess systematic reviews (AMSTAR). This refines and enhances work presented in previously published instruments (by Oxman and Guyatt, 1991 [3] and Sacks et al., 1987 [4]) [5]. The present study concerns the internal validation of AMSTAR using the set of reviews used in its development. Here we focus on parameters of agreement, reliability; construct validity, and feasibility through comparisons with other instruments. An external validation of AMSTAR has been reported separately [6].

2. Methods

We used a computer-generated random sample of 30 (20%) of 151 systematic reviews that were used in the development of the instrument [5]. This sample contained 11 Cochrane and 19 non-Cochrane reviews, including meta-analyses and qualitative reviews. The topics of the reviews ranged across the spectrum of medicine [7–36]. Two reviewers (one without formal training) applied the new AMSTAR instrument and the two quality assessment tools, the enhanced Overview of Quality Assessment Questionnaire (OQAQ, originally developed by Oxman and Guyatt), and the instrument developed by Sacks et al. to all 30 reviews (C.H., B.J.S.) [3,4]. For each reviewer, the data set extracted contained three quality ratings for each review, yielding a total of six ratings per review.

2.1. Agreement and reliability

We calculated overall agreement and Cohen’s kappa for each item (“yes” scores vs. any other scores) [37]. Bland and Altman’s limits of agreement method explained the agreement graphically [38–40]. We awarded each item scoring “yes” one point and summed these to calculate a total score. Intraclass correlation coefficients (ICCs) assessed the reliability of this total score [41]. We further scrutinized items and reviews with kappa values below 0.50. Finally, we repeated the exercise for the OQAQ and Sacks’ instruments. Kappa values of less than 0 were rated as less than chance agreement; 0.01–0.20, slight agreement; 0.21–0.40, fair agreement; 0.41–0.60, moderate agreement; 0.61–0.80, substantial agreement; and 0.81–0.99, almost perfect agreement. [42]

2.2. Construct validity

The new instrument already has high face and content validity by virtue of its construction process [5]. In the current study, we assessed construct validity by converting the mean total score (mean of two raters C.H. and B.J.S.) of each of the 30 reviews to a percentage of the maximum score for each of the three instruments. ICCs then assessed convergence of the total scores between each pair of instruments (AMSTAR–OQAQ, AMSTAR–Sacks, and OQAQ–Sacks).

2.3. Feasibility

Based on a guideline for assessing feasibility of instrument use developed by the Outcome Measures in Rheumatology (OMERACT) group [43], we compared the feasibility of the new instrument with that of the existing instruments by recording the time it took to complete scoring and the instances where scoring was difficult or impossible. The wording of individual items is critical for the performance of AMSTAR and fine-tuning is expected to be an ongoing task.

SPSS (version 13; SPSS Inc., Chicago, IL, USA) and MedCalc Software (Mariakerke, Belgium) were used to analyze the data, and the results were expressed as means and 95% confidence intervals (CIs) unless otherwise noted.

3. Results

The sample of 30 reviews adequately covered a wide range of quality, albeit with some underrepresentation of poor-quality reviews. Overall quality scores on AMSTAR ranged from 3 to 10 (out of a maximum of 11) with a flat distribution between 3.5 and 10 and a mean percentage score of 49.4%. The overall quality scores on Sacks’ instrument ranged from 5 to 16 (out of a maximum score of 24), with a mean percentage score of 41.6%, and for OQAQ, scores ranged from 3 to 10 (out of a maximum score of 10), with a mean percentage score of 63.3%.
3.1. Agreement and reliability

The interobserver agreement of the individual items in the AMSTAR was high: mean = 0.88 (range: 0.73–1.0) with a mean kappa of 0.70 (95% CI: 0.57, 0.83) (range: 0.38–1.0). However, items 4 (publication status), 7 (report of assessment of scientific quality), and 9 (appropriate method to combine studies) scored fair to moderate at 0.38, 0.42, and 0.45, respectively. On the first two of these items, overall agreement was substantial at 0.80 and the relatively low kappa may be explained by a skewed distribution, that is, a high number of reviews in which the reviewers agreed on the score “no” (item 4) and “yes” (item 7), respectively. On item 8, overall agreement was also satisfactory at 0.74. Compared with the other instruments, agreement on individual items was similar to OQAQ: mean kappa of 0.63 (95% CI: 0.39, 0.78) (range 0.39–0.84), and superior to the Sacks’ instrument: mean kappa of 0.40 (95% CI: 0.29, 0.50) (range: −0.47 to 0.93). In these instruments, fair to moderate agreement was also seen in the items covering assessment of scientific validity, statistical combinability, and comprehensive literature searching (Table 1).

For the AMSTAR total score, the mean difference between the two observers’ scores was 0.2 (0.36–0.91). Agreement was similar in reviews with high- and low-quality scores (Fig. 1).

The interobserver ICC for the total score was excellent for all instruments: AMSTAR, 0.84 (95% CI: 0.65, 0.92); OQAQ, 0.91 (95% CI: 0.82, 0.96); and Sacks’ instrument, 0.86 (95% CI: 0.71, 0.94). In one non-Cochrane review [12], observers differed by 3 points (6 vs. 9). In this review, the differences were noted on AMSTAR questions addressing duplication study selection and data extraction (item 2), publication status (item 4), and methods used to combine studies (item 9). In one Cochrane review [15], observers differed by 4 points (1 vs. 5). In this review, differences were noted on AMSTAR questions assessing the a priori design (item 1), publication status (item 4), scientific quality (item 7), and methods used to combine studies (item 9). The overall quality of Cochrane reviews included in this data set was somewhat higher than non-Cochrane reviews.

The qualitative analysis of the data on agreement led us to make minor modifications to the wording of some items. In particular, under the original item regarding publication bias, the wording was changed to clarify the purpose of the question, that is, to ask whether the status of publication was used as an inclusion criterion (see item 4 and footnote in Appendix). Additional available electronic databases were also added to the question on literature searching (item 3) and I² was added to the item on methods used to combine findings (item 9).

3.2. Construct validity

Expressed as a percentage of the maximum score, the results of AMSTAR showed convergence with the results of the other instruments. ICC for AMSTAR was 0.66 (95% CI: 0.28, 0.84) against OQAQ and 0.83 (95% CI: 0.64, 0.92) against Sacks’ instrument. The ICC obtained when comparing OQAQ with Sacks’ instrument was 0.86 (95% CI: 0.70, 0.93).

3.3. Feasibility

AMSTAR proved easy to apply, each review taking 14.9 (95% CI: 17.0, 12.8) minutes to complete. OQAQ took, on average, 20.3 (95% CI: 22.5, 18.0) minutes to complete, and Sacks’ instrument 34.4 (95% CI: 37.3, 31.6) minutes (P < 0.0001 for comparison between the three instruments). Two of the reviewers expressed difficulty with

Table 1
Assessment of the interrater agreement for AMSTAR

<table>
<thead>
<tr>
<th>Items</th>
<th>Kappa (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Was an “a priori” design provided?</td>
<td>0.80 (0.63, 0.90)</td>
</tr>
<tr>
<td>2. Was there duplicate study selection and data extraction?</td>
<td>0.80 (0.17, 0.81)</td>
</tr>
<tr>
<td>3. Was a comprehensive literature search performed?</td>
<td>0.72 (0.40, 0.87)</td>
</tr>
<tr>
<td>4. Was the status of publication (i.e., grey literature) used as an inclusion criterion?</td>
<td>0.38 (0.28, 0.70)</td>
</tr>
<tr>
<td>5. Was a list of studies (included and excluded) provided?</td>
<td>0.56 (0.07, 0.79)</td>
</tr>
<tr>
<td>6. Were the characteristics of the included studies provided?</td>
<td>0.74 (0.45, 0.86)</td>
</tr>
<tr>
<td>7. Was the scientific quality of the included studies assessed and documented?</td>
<td>0.42 (0.23, 0.72)</td>
</tr>
<tr>
<td>8. Was the scientific quality of the included studies used appropriately in formulating conclusions?</td>
<td>0.74 (0.45, 0.87)</td>
</tr>
<tr>
<td>9. Were the methods used to combine the findings of studies appropriate?</td>
<td>0.45 (0.12, 0.70)</td>
</tr>
<tr>
<td>10. Was the likelihood of publication bias assessed?</td>
<td>0.88 (0.75, 0.94)</td>
</tr>
<tr>
<td>11. Were potential conflicts of interest included?</td>
<td>0.92 (0.83, 0.96)</td>
</tr>
</tbody>
</table>

Abbreviations: AMSTAR, a measurement tool to assess systematic reviews; CI, confidence interval.
scoring item 4 on publication status: “was the status of publication (i.e., grey literature) used as an inclusion criterion?”

4. Discussion

There has been a continued proliferation of (largely unvalidated) scales and checklists for assessing the quality of systematic reviews [44]. This causes confusion for those who use reviews in making clinical and policy decisions and who need to be able to distinguish good- from poor-quality reviews. There is a need for a reliable and valid quality assessment instrument that is easy to use. AMSTAR was developed to meet this need. Our aim was not to devise a truly original instrument, but to develop and update the best available from the published literature. Our scan indicated that the OQAQ developed by Oxman and Guyatt [3] and the rating scale of Sacks et al. [4] were among the best out of more than two dozen instruments assessed by us. We found that both instruments had been rigorously developed, but were dated in some respects. We decided to improve the descriptors of the items resulting in the “enhanced OQAQ” that we have applied in the subsequent studies [2]. The checklist developed by Sacks et al. showed good quality and was especially comprehensive but unwieldy in general use [4]. We based AMSTAR on a development of both of these instruments. Full details of this development process are published elsewhere [5].

This study suggests that AMSTAR has good agreement, reliability, construct validity, and feasibility to assess the quality of systematic reviews. Its performance in terms of agreement and reliability was similar to OQAQ and better than Sacks’ instrument; it adds important items that are not present in either instrument (e.g., publication status, conflict of interest), and has better feasibility than OQAQ or Sacks’ instrument. We think AMSTAR can be applied to a wide variety of systematic reviews, but recognize that it has only been tested on systematic reviews of randomized control trials evaluating treatment interventions. We accept that the relatively high reliability of total scores for AMSTAR and OQAQ may be partly because of the raters’ familiarity with both instruments, and this reliability needs to be tested more widely in the field.

AMSTAR showed good (convergent) construct validity in comparison with the two existing instruments. A recently published study concluded that the underlying construction of OQAQ is designed for the assessment of meta-analyses. Thus, it is difficult for any other type of review to score highly on the OQAQ, and if the review does not have a meta-analysis component, it may be deemed to have major flaws [45]. AMSTAR can be scored both individually (components) and as a checklist by summing the item scores (overall score). It was psychometrically developed to score each item as if it was not related to the others. Each component came out separate in the factor analysis. Therefore, all reviews have equal chance of scoring well, but meta-analysis will score slightly higher in the overall results.

The feasibility of AMSTAR is documented in terms of the time required to complete an assessment while using it: about 10—15 minutes, which is substantially less than the time needed to complete the other instruments.

In this study, we did not assess the external validity of AMSTAR. This has been carried out separately and the results have recently been published. In that analysis, we looked at differences in overall scores and compared them with the global assessments made by an informed panel. We found very good correlations on the total scores [6].

There has been considerable discussion regarding the merits of using individual component scores or summary scores when assessing systematic reviews and individual studies [45,46]. From a methodological standpoint, it is worth assessing the component scores as they measure different elements, and some may be more important than others in particular situations. Hence, a summary score may obscure important strengths or weaknesses. In the case of AMSTAR, we have tried to develop an instrument that is pragmatic and of value to decision makers. It is valid and easy to use and the total score is meaningful. By this, we mean that we have ensured that the components are non-overlapping and have separately validated the total score against an external standard [5,6]. Hence, we believe that decision makers can have some confidence that the component scores measure different domains of quality and the overall score is meaningful.

We feel that the main advantage of AMSTAR over the OQAQ and Sacks’ instruments lies in its better compromise between comprehensiveness and feasibility. It adds relevant dimensions to those covered in the OQAQ without becoming unwieldy, as with the Sacks’ instrument. For example, the item “sources of support” was included in the original data set and came out as a component in the factor analysis. Some may doubt the usefulness of the item concerning conflict of interest. We put a lot of thought into the name and description of this item, using all available empirical evidence. In addition to the research previously discussed on this topic, more recent studies also suggest that funding source influences outcomes and quality of research [47]. In a study by Biondi-Zoccai [48], the authors concluded that reviewers who reported previous not-for-profit funding were more likely to carry out higher-quality systematic review. We believe that funding sources are associated with bias in systematic reviews and it is important to rate this aspect of their conduct. The inclusion of unpublished studies is rated by AMSTAR. This remains a controversial topic in systematic reviews. Several examples in the drugs’ field have demonstrated the crucial importance of including nonpublished data. In the case of two Cox-2 inhibitors, rofecoxib and celecoxib, the incomplete reporting of vascular deaths and gastrointestinal events skewed the results of trials that were used as the basis of important decisions [49]. More recently, several systematic
reviews/meta-analyses of antidepressant drugs have shown that the inclusion of unpublished data dramatically alters the perception of benefit in a negative way [50,51]. Based on these examples, we felt it important to include an item in AMSTAR dealing with unpublished studies.

During the development of AMSTAR, careful consideration was given to the wording of the individual items and minor adjustments were made where necessary; despite this process, agreement between observers was disappointingly low on three items. One of these items assessed publication restriction. After discussion between observers, we reworded the descriptor slightly and this has improved agreement. The reworded version is provided in Appendix. The other two items describe “report of assessment of scientific quality” and “appropriate method to combine studies.” Agreement was low on similar items in the other instruments assessed here. Subjective judgment comes into play when one is asked to assess whether quality of included studies was assessed adequately. Conceivably, one could increase reliability of assessment by providing more detailed instructions or by adding more items or criteria. This would, however, decrease feasibility. It should also be noted that overall agreement on these items was good; hence, their relatively low kappa values are likely caused by skewness in the responses, that is, most of the responses in either the “yes” or the “no” category. This is a well-known limitation of the kappa statistic [52].

Our study has other limitations. We did not compare AMSTAR with the current state-of-the-art reporting quality of meta-analysis (QUOROM) [53]. The reason for this is that QUOROM is not specifically designed to assess methodological quality. Rather, it is specifically focused on the quality of reporting (not conduct) of the review. This does not detract from the utility of QUOROM, but its limited focus made it unsuitable for our study. A further limitation of the present study is the fact that the sample of reviews used is derived from the original source used to develop AMSTAR (Table 2), and one of the assessors is the principal investigator. Thus, application to other reviews and by other assessors is necessary to discover the full potential of this tool. Finally, the number of reviews used to validate AMSTAR was rather small.

Our new instrument builds on previous work. Methodologists continue to struggle with methodological quality issues, whereas decision makers struggle with the challenge of basing policy, clinical, or resource planning decisions on the available evidence. The personal feedback received on AMSTAR has been supportive. AMSTAR is now being used by a number of groups, including the Canadian Agency for

<table>
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<tr>
<th>Author</th>
<th>Year</th>
<th>Journal type</th>
<th>Topic area</th>
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<tbody>
<tr>
<td>Anonymous</td>
<td>1989</td>
<td>NEJM</td>
<td>Breast cancer</td>
</tr>
<tr>
<td>Appel</td>
<td>1993</td>
<td>Arch Intern Med</td>
<td>Blood pressure reduction</td>
</tr>
<tr>
<td>Buring</td>
<td>1988</td>
<td>Rev Inf Dis</td>
<td>Aminoglycoside antibiotics</td>
</tr>
<tr>
<td>Chalmers</td>
<td>1977</td>
<td>NEJM</td>
<td>Acute myocardial infarction</td>
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<tr>
<td>Clagett</td>
<td>1988</td>
<td>Ann Surg</td>
<td>Venous thromboembolism</td>
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<tr>
<td>Counsell</td>
<td>1996</td>
<td>Cochrane</td>
<td>Carotoid surgery</td>
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<td>Daya</td>
<td>1996</td>
<td>Cochrane</td>
<td>FSH and hMG in IVF</td>
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<td>Duley</td>
<td>1996</td>
<td>Cochrane</td>
<td>Anticonvulsants for pre-eclampsia</td>
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<tr>
<td>Fanning</td>
<td>1992</td>
<td>Obstet Gynecol</td>
<td>Ovarian Carcinoma</td>
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<tr>
<td>Gent</td>
<td>1986</td>
<td>Chest</td>
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<td>Gotzsche</td>
<td>1995</td>
<td>BMJ</td>
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<td>1996</td>
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<td>Arzneim-Forsch</td>
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<td>Fertil Steril</td>
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<td>Kaufmann</td>
<td>1988</td>
<td>Health Psychol</td>
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<td>1996</td>
<td>Cochrane</td>
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<td>Renfrew</td>
<td>1996</td>
<td>Cochrane</td>
<td>Infant discharge times</td>
</tr>
<tr>
<td>Renfrew</td>
<td>1996</td>
<td>Cochrane</td>
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<td>Soares</td>
<td>1996</td>
<td>Cochrane</td>
<td>Tardive dyskinesia and GABA</td>
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<td>Thacker</td>
<td>1985</td>
<td>BJOG</td>
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<td>Velanovich</td>
<td>1989</td>
<td>Surgery</td>
<td>Resuscitation</td>
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<tr>
<td>Wilson</td>
<td>1992</td>
<td>J Hosp Infect</td>
<td>Surgical prophylaxis</td>
</tr>
</tbody>
</table>

FSH, follicle stimulating hormone; hMG, human menopausal gonadotrophin; IVF, in vitro fertilization; GABA, gamma-aminobutyric acid.
Drugs and Technologies in Health, and The Cochrane Effective Practice and Organization of Care Group (EPOC) [54]. With publication of data on reliability and validity in a peer-reviewed journal, we hope that it will help many reviewers with their task of assessing the methodological quality and incorporating the results into their systematic reviews.

In summary, AMSTAR is an empirically developed instrument for documenting the quality of systematic reviews. It was found to have good agreement, reliability, and construct validity in a limited test setting. It combines one instrument a level of comprehensiveness and feasibility not found in existing instruments. We encourage others to test our new instrument on other samples of systematic reviews. Its ongoing application in the assessment of the quality of systematic reviews will provide further confirmation of its utility.

Further validation is needed to replicate the initial promising results, and this should involve a broader range of assessors and a broader range of reviews to assess whether the reliability and validity are confirmed in diverse circumstances.

Appendix: A measurement tool to assess systematic reviews (AMSTAR)

1. Was an “a priori” design provided?
   - Yes
   - No
   - Can’t answer
   - Not applicable

2. Was there duplicate study selection and data extraction?
   - Yes
   - No
   - Can’t answer
   - Not applicable

3. Was a comprehensive literature search performed?
   - Yes
   - No
   - Can’t answer
   - Not applicable

4. Was the status of publication (i.e., grey literature) used as an inclusion criterion?
   - Yes
   - No
   - Can’t answer
   - Not applicable

5. Was a list of studies (included and excluded) provided?
   - Yes
   - No
   - Can’t answer
   - Not applicable

6. Were the characteristics of the included studies provided?
   - Yes
   - No
   - Can’t answer
   - Not applicable

7. Was the scientific quality of the included studies assessed and documented?
   - Yes
   - No
   - Can’t answer
   - Not applicable

8. Was the scientific quality of the included studies used appropriately in formulating conclusions?
   - Yes
   - No
   - Can’t answer
   - Not applicable

9. Were the methods used to combine the findings of studies appropriate?
   - Yes
   - No
   - Can’t answer
   - Not applicable

10. Was the likelihood of publication bias assessed?
    - Yes
    - No
    - Can’t answer
    - Not applicable

11. Was the conflict of interest included?
    - Yes
    - No
    - Can’t answer
    - Not applicable

“Can’t answer” is chosen when the item is relevant but not described by the authors; “not applicable” is used when the item is not relevant, such as when a meta-analysis has not been possible or was not attempted by the authors.
* The original wording for question #4: Was the status of publication (i.e., grey literature) used/not used as an exclusion criterion? The authors should state that they searched for reports regardless of their publication type. The authors should state whether or not they excluded any reports (from the systematic review), based on their publication status, language etc.

References


[2] Shea B, Dubé C, Moher D. Assessing the quality of reports of systematic reviews: the QUOROM statement compared to other tools. In:


