Updated Method Guidelines for Cochrane Musculoskeletal Group Systematic Reviews and Metaanalyses


ABSTRACT. The Cochrane Musculoskeletal Group (CMSG), one of 53 groups of the not-for-profit, international Cochrane Collaboration, prepares, maintains, and disseminates systematic reviews of treatments for musculoskeletal diseases. It is important that authors conducting CMSG reviews and the readers of our reviews be aware of and use updated, state-of-the-art systematic review methodology. One hundred sixty reviews have been published. Previous method guidelines for systematic reviews of interventions in the musculoskeletal field published in 2006 have been substantially updated to incorporate methodological advances that are mandatory or highly desirable in Cochrane reviews and knowledge translation advances. The methodological advances include new guidance on searching, new risk-of-bias assessment, grading the quality of the evidence, the new Summary of Findings table, and comparative effectiveness using network metaanalysis. Method guidelines specific to musculoskeletal disorders are provided by CMSG editors for various aspects of undertaking a systematic review. These method guidelines will help improve the quality of reporting and ensure high standards of conduct as well as consistency across CMSG reviews. (First Release Dec 1 2013; J Rheumatol 2014;41:194–205; doi:10.3899/jrheum.121306)

Key Indexing Terms: EVIDENCE-BASED MEDICINE SYSTEMATIC REVIEW MUSCULOSKELETAL ARTHRITIS METAANALYSIS

The aim of the Cochrane Collaboration is to help healthcare providers, patients, patient advocates and carers, and policy makers arrive at well-informed decisions on healthcare treatments by preparing, maintaining, and disseminating methodologically strong systematic reviews1.

The systematic review is an essential tool for managing the vast amount of information generated on the etiology, prognosis, incidence/prevalence, diagnosis, prognosis, and treatment of disease. These method guidelines focus on the assessment of treatments (including both benefits and harms) that are aimed at improving health. Cochrane systematic reviews are increasingly used as the basis for clinical decision support resources such as UpToDate and clinical guidelines [e.g., American College of Rheumatology (ACR) osteoarthritis (OA)2 and rheumatoid arthritis (RA)3 guidelines]. Compared to a narrative literature review, the systematic review uses “scientific strategies that are systematic, and designed to limit bias in the assembly, critical appraisal and synthesis of all relevant studies on a specific topic”4. Most Cochrane reviews are

From the Cochrane Musculoskeletal Group (CMSG), Institute of Population Health, University of Ottawa, Ottawa, Ontario, Canada.

Supported through a grant from the Canadian Institutes of Health Research.

E.A.T. Ghogomu, MD, MSc, Assistant Managing Editor; L.J. Maxwell, MSc, Managing Editor, CMSG, University of Ottawa; R. Buchbinder, MBBS (Hons), MSc, PhD, FRACP, Director, Monash Department of Clinical Epidemiology, Cabrini Hospital, Malvern VIC, Australia, and Professor, Department of Epidemiology and Preventive Medicine, School of Public Health and Preventive Medicine, Monash University, and Joint Coordinating Editor, CMSG; T. Rader, MLIS, Knowledge Translation Specialist, CMSG, University of Ottawa; J. Pardo Pardo, BA, Research Coordinator, Centre for Global Health, University of Ottawa; R.V. Johnston, PhD, Managing Editor, CMSG, Monash Department of Clinical Epidemiology, Cabrini Hospital, Monash University; R.D.K. Christensen, BSc, MSc, PhD, Senior Biostatistician, Head of Musculoskeletal Statistics Unit, The Parker Institute, Copenhagen University Hospital, Bispebjerg and Frederiksberg, Copenhagen, Denmark; A.W.S. Rutjes, PhD, senior research fellow, Institute of Social and Preventive Medicine, University of Bern, Bern, Switzerland; T.M. Winzenberg, MBBS, PhD, FRACGP, MMEdSc(ClinEpi), Senior Research Fellow–General Practice at Menzies Research Institute Tasmania, University of Tasmania, Tasmania, Australia; J.A. Singh, MBBS, MPhil, Birmingham VA Medical Center, Associate Professor, University of Alabama at Birmingham, Alabama, USA; G. Zanoli, MD, PhD, Casa di Cura SM Maddalena, University of Ferrara Occhialbo, Ferrara, Italy; G.A. Wells, PhD, Professor of Epidemiology and Community Medicine, University of Ottawa; P. Tugwell, MD, MSc, FRCP, FCAMS, Canada Research Chair, Professor of Medicine, and Epidemiology and Community Medicine, Faculty of Medicine; Principal Scientist, Institute of Population Health, University of Ottawa, Department of Medicine, Ottawa Hospital, Senior Scientist, Ottawa Hospital Research Institute, Joint Coordinating Editor, CMSG.

Address correspondence to Dr. E.A.T. Ghogomu, 1 Stewart St., Ottawa, Ontario K1N 6N5, Canada. E-mail: cmsg@uottawa.ca

Accepted for publication August 7, 2013.
Defining the question. First, the research question needs to be clearly formulated using the “PICOS” framework, i.e., a clinically relevant or policy-relevant question that takes into account the patient/population, intervention, comparison, outcomes, and study design, and includes both the benefit and harm of the intervention being studied.

Priority topics for new and updated reviews have been identified by CMSG editors and consumers based on criteria including the burden of disease, equity, identification of new interventions, number of new studies, and frequency of access for existing reviews.

Literature search and study selection. The complete search strategy for each database searched is defined a priori and is documented in the review appendices with the date so that the search can be duplicated. The search strategy frequently needs tailoring to the topic, so it is reviewed before implementation by the CMSG Trial Search Coordinator and peer reviewed by information science specialists using the PRESS checklist (Peer Review of Electronic Search Strategies).

It is recommended that, at a minimum, the following databases and trial registers be searched: MEDLINE, EMBASE, the Cochrane Central Register of Controlled Trials (CENTRAL), ClinicalTrials.gov, and the World Health Organization International Clinical Trial Registry Platform portal. In addition, we recommend checking the references in identified relevant systematic reviews and individual studies that meet the review’s inclusion criteria. For systematic reviews of drugs, authors should search for adverse effects in Websites of regulatory authorities such as the US Food and Drug Administration (FDA) and the European Medicines Evaluation Agency (EMEA). For example, the FDA Website contains important trial and observational study data on tuberculosis and fungal infections from the use of biologics, which were included in the Cochrane and BMJ reviews on biologics overview and network metaanalysis. The Trial Search Coordinator may suggest additional sources, such as conference abstracts from ACR and the European League Against Rheumatism (EULAR), depending on the topic.

We do not recommend excluding trials in languages other than English. Some topics such as studies of the effects of medicinal plants may have a significant number of trials published in another language and the CMSG can assist with translation when necessary.

Two people should independently screen the titles and abstracts from the results of the searches for the selection of trials meeting the predefined inclusion criteria. The full text of those articles that appear to meet the inclusion criteria should then be obtained and assessed for eligibility. Those full-text studies that do not meet the inclusion criteria should be added to the Table of Excluded Studies and a reason provided for their exclusion.
INCLUSION CRITERIA

The minimum criteria for trial inclusion in the systematic review should be defined in advance and address several items using the PICO framework.

Population. Participants of trials should be defined by acceptable diagnostic criteria where possible, such as the ACR criteria for OA, RA, gout, systemic lupus erythematosus, and fibromyalgia (FM). Specific exclusions, such as age, sex, and condition, must be detailed.

Trials may report “mixed populations” in which patients with different conditions are enrolled. For example, for reviews on knee OA, randomized controlled trials (RCT) included patients with both hip and knee OA. If such situations are anticipated, review authors should define in advance how to handle these reports. Rather than excluding, a rule may be chosen to include those trials, requiring that at least a given percentage, such as 75%, of patients meet the inclusion criteria. It is also desirable to contact trial authors, to obtain data for the population of interest.

Intervention. Glasziou, et al have pointed out the importance of having sufficient information in papers to be able to apply the intervention to patients. The intervention must be explicitly described. If applicable, the route of administration, dose, timing, duration of treatment, and concomitant treatments should be outlined.

An example of a definition of type of intervention could read like this: Trials were included that investigated treatment with adalimumab 40 mg subcutaneously every week to every other week, alone or in combination with disease-modifying antirheumatic drugs (DMARD) for a minimum of 12 months.

Comparator. The comparator intervention should be explicitly defined (e.g., placebo, another treatment). An example of a definition of type of comparator: Studies were included comparing leflunomide treatment (as monotherapy or in combination with other DMARD) at a dose of 20 to 25 mg/day (with or without a loading daily dose of 100 mg given in the first 1 to 3 days) to placebo or other DMARD.

Outcomes. Cochrane reviews are only as useful as their outcomes are relevant and accurate. Cochrane reviews now report results by outcome. They should include all outcomes that are likely to be meaningful, and not include trivial outcomes. At the time of the title registration, the authors should list all patient-important outcomes, including both benefits and harms relevant to the intervention, organized from the most important to the least important. The major outcomes (up to a maximum of 7) to be presented in the “summary of findings” (SoF) table are selected at the protocol stage.

Review authors must choose at the protocol stage what they consider the main timepoint of interest for each outcome. This does not imply that they should extract only 1 timepoint; to the contrary, analyzing and depicting results over time is very informative. However, defining the timing in advance forces the review author to think about short-term and long-term effects and to consider whether both are relevant for their intervention in question. It helps to plan statistical analyses and to define the focus of the review.

The SoF table (Figure 1), which may be created using GRADEProfiler software, is now shown on the first page of every Cochrane review, along with the matching abstract and plain language summary. Although review authors may complete an SoF table for each major comparison in their review, to best convey the main “evidence-based actionable message” to users, the Cochrane Library format places the single most important SoF table on the first page of the review. This means that the accuracy and consistency of the numbers and wording across the SoF table, plain language summary, and abstract are pivotal.

The CMSG is in the process of developing default outcome templates for SoF tables for classes of interventions for each condition. Standardizing the outcomes presented in these tables will improve consistency for readers and also permit easier production of overviews of reviews using network metaanalyses. The CMSG editorial team has drafted preliminary default guidelines for which outcomes should be included in SoF tables for pharmacologic and complementary interventions in the following conditions: RA, OA, FM, and ankylosing spondylitis. The preliminary default templates for pharmacologic and complementary interventions for RA and OA are shown in Table 1. These may need tailoring for specific interventions or for specific research questions and may require a different set of major outcomes. For example, in a review on a biologic for RA, radiographic progression would be an important outcome of interest; in a review of arthroplasty, the imaging outcomes would be very different; while in a review on the effect of patient education programs for RA, imaging changes may not be of key interest.

Common core sets established and validated by groups such as OMERACT (Outcome Measures in Rheumatology — an international initiative to improve outcome measurement in rheumatology) and their associated groups are encouraged. This has yet to be done for regional musculoskeletal disorders such as shoulder and elbow disorders, where currently a set of standardized measures does not exist; until then we favor a description of the most relevant ones from the patient’s perspective. The CMSG has a joint working group with OMERACT and their partners to develop standardized outcomes by both condition and intervention for use in CMSG SoF tables.

The CMSG accepts surrogate outcomes if other outcomes are not available and if they meet the following conditions: (1) they have been shown to be on the causal
Abatacept (2 and 10 mg/kg) + DMARD/biologic versus placebo + DMARD/biologic for rheumatoid arthritis

**Patient or Population:** patients with rheumatoid arthritis  
**Settings:** International; clinic/hospital  
**Intervention:** Abatacept (2 and 10 mg/kg) + DMARD/biologic  
**Comparison:** Placebo +DMARD /biologic

<table>
<thead>
<tr>
<th>Outcomes</th>
<th>Placebo +DMARD/biologic</th>
<th>Abatacept (2 &amp;10 mg/kg) +DMARD/biologic</th>
<th>Relative Effect (95% CI)</th>
<th>No of Participants (Studies)</th>
<th>Quality of Evidence (GRADE)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACR 50% improvement Follow-up: 12 months</td>
<td>168 per 1000 (291 to 474)</td>
<td>371 per 1000 (1.73 to 2.82)</td>
<td>993 (3)</td>
<td>Moderate</td>
<td>Absolute difference =21% (16% to 27%). NNT=5 (4 to 7) Relative percent change =121% (73% to 182%)</td>
<td></td>
</tr>
<tr>
<td>Pain measured at end of study on a 100 mm visual analog scale. Scale from 0 (better) to 100 (worse). Follow-up: 12 months.</td>
<td>Mean pain in the control group was 49.24 mm</td>
<td>Mean pain in the intervention group was 10.71 mm lower (12.97 to 8.45)</td>
<td>1425 (1)</td>
<td>Moderate</td>
<td>Absolute difference = -11% (-13% to -8.5%). NNT=5 (4 to 6) Relative percent change = 18% (-22% to -14%)</td>
<td></td>
</tr>
<tr>
<td>Improvement in physical function (HAQ: greater than 0.3 increase from baseline, 0-3 scale) Follow-up: 12 months</td>
<td>393 per 1000 (531 to 766)</td>
<td>637 per 1000 (1.35 to 1.95)</td>
<td>638 (1)</td>
<td>Moderate</td>
<td>Absolute difference = -24% (16% to 32%) NNT=5 (4 to 7) Relative percent change = 62% (35% to 198%)</td>
<td></td>
</tr>
<tr>
<td>Achievement of low disease activity state (DAS 28 less than 3.2, scale 0-10) Follow-up: 12 months</td>
<td>98 per 1000 (278 to 646)</td>
<td>424 per 1000 (2.84 to 6.59)</td>
<td>683 (1)</td>
<td>Moderate</td>
<td>Absolute difference = 33% (26% to 39%) NNT=4 (3 to 5) Relative percent change = 333% (184% to 559%)</td>
<td></td>
</tr>
<tr>
<td>Total serious adverse events Follow-up: 6 to 12 months</td>
<td>121 per 1000 (105 to 155)</td>
<td>127 per 1000 (0.87 to 1.28)</td>
<td>3151 (6)</td>
<td>Moderate</td>
<td>Absolute difference = 1% (-2% to 3%). NNT=n/a Relative percent change = 1% (-14% to 29%)</td>
<td></td>
</tr>
<tr>
<td>Long-term serious adverse events Follow-up: 2 years</td>
<td>See comment</td>
<td>See comment</td>
<td>Not estimable</td>
<td>950 (2)</td>
<td>Low</td>
<td>Number of patients with SAE: Genovesse 2005: 103/357; 23.4 SAE/100 patient-years; 70% completed the LTE Kremers 2006: 149/593; 16.3 SAE/100 patient-years; 90.5% completed the LTE</td>
</tr>
</tbody>
</table>
STUDY DESIGN

Review authors should consider what study designs are likely to provide valid data to answer their questions. The study designs included will depend upon the question, the context, and the resources of the systematic review team. Reviews should define selection criteria for study designs according to their “fitness for purpose” for the research question being posed, rather than just follow an evidence hierarchy. The rationale for the fitness for purpose should be clearly stated and explained.

RCT, where 2 or more groups are formed by randomly allocating participants so that any differences between groups can be attributed to the intervention, should always be included. Controlled clinical trials (CCT) are trials where allocation to treatment and control groups is quasi-random, for example, alternation, date of birth, or case record number. In some treatment settings, such as educational interventions, it is not possible to randomize individuals because of the risk of 1 group receiving some or all of the intervention of the other group (i.e., contamination) as a result of being in the same setting or place; cluster RCT overcome this contamination by randomizing the different individual practices to different groups. Crossover RCT, in which each patient is allocated to a sequence of treatment and control interventions, can also be included, but their analyses need special attention.

Other study designs can also be included, whenever possible, according to their “fitness for purpose”. In 2009, 6% of Cochrane reviews included nonrandomized study designs and it has become a priority to develop the skills and best-practice methods to ensure that this component of systematic reviews is useful. The Cochrane Non-Randomised Trial Methods Group has developed guidelines for nonrandomized studies to standardize searching and assessment of rare and delayed adverse effects that will not be detected in short-term trials. A series of 6 papers has been published that provides an update on the increasing consensus on how these studies should be assessed and synthesized. The last article provides useful checklists to help authors.

An example of study design: To assess benefits and harms we included RCT. To further assess harms, we included the following types of studies as long as they reported at least 1 year of followup for patients taking anti-tumor necrosis factor agents, had a sample size > 100 patients, and reported an adverse effect outcome: CCT, cohort studies (prospective, e.g., longterm extension of RCT, or retrospective), case-control studies, case series, and published registry data.

Assessment of risk of bias. In RCT, because an included study may be performed to the highest possible standards but some individual patient-important outcomes may be underpowered and/or still have an important risk of bias, the 2011 handbook also recommends the assessment of risk of bias by outcomes.

---

Table 1. Rheumatoid arthritis (RA) and osteoarthritis (OA) preliminary default outcome templates for pharmacologic and complementary interventions.

<table>
<thead>
<tr>
<th>RA: Major Outcomes for the SoF Table</th>
<th>OA: Major Outcomes for the SoF Table</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACR 50</td>
<td>Pain</td>
</tr>
<tr>
<td>DAS (low or remission)</td>
<td>Physical function</td>
</tr>
<tr>
<td>HAQ for function</td>
<td>Quality of life</td>
</tr>
<tr>
<td>Radiograph or appropriate imaging changes</td>
<td>Radiograph or appropriate imaging changes</td>
</tr>
<tr>
<td>Patients who withdraw because of adverse events</td>
<td>Patients who withdraw because of adverse events</td>
</tr>
<tr>
<td>Serious adverse events</td>
<td>Serious adverse events</td>
</tr>
<tr>
<td>Longterm adverse events</td>
<td>Longterm adverse events</td>
</tr>
</tbody>
</table>

* Where possible, biomarkers such as radiographs should be transformed into an outcome. ACR: American College of Rheumatology; SoF: summary of findings; DAS: Disease Activity Score; HAQ: Health Assessment Questionnaire.
Risk of bias is assessed in a 2-step process (Figure 2, Figure 3).

Step 1: The risk-of-bias tool addresses 7 different domains: (1) sequence generation; (2) allocation concealment; (3) blinding of participants and personnel; (4) blinding of outcome assessment; (5) incomplete outcome data; (6) selective outcome reporting; and (7) “other sources of bias”. Review authors need to specify in their protocol which issues they will consider for “other sources of bias”. Other potential sources of bias should address issues that may affect the internal validity of the study. The handbook provides further details on these issues such as significant baseline imbalances between groups, or situations where a cointervention is not administered evenly between groups. Each domain includes 1 or more specific entries in a risk-of-bias table. Within each entry, the first part of the tool involves assigning a judgment of low risk, high risk, or unclear risk, and the second part involves providing an explanation of the judgment. A summary table of review authors’ judgments for each risk-of-bias item for each study is shown in Figure 2 for the “Abatacept for rheumatoid arthritis” review. Independent assessment of risk of bias should be undertaken by at least 2 review authors. Where differences in assessment cannot be resolved, arbitration by a third person is warranted.

Step 2: The handbook suggests summarizing risk of bias for each important outcome within and across studies using 3 categories — low, unclear, and high risk of bias. Within a single trial, different outcomes may be at different risk of bias given that different studies may contribute to each
outcome. Figure 3 shows a plot of the distribution of review authors’ judgments across studies for each risk-of-bias item in the “Abatacept for rheumatoid arthritis” review.

In nonrandomized (observational) studies, assessment of risk of bias is more difficult than assessment in an RCT. Risk-of-bias assessment methods for systematic reviews of nonrandomized studies are under development for the Cochrane Collaboration. Meanwhile, 6 existing useful tools have been identified\textsuperscript{3,48}. One tool is the Newcastle-Ottawa Scale\textsuperscript{49} (www.ohri.ca/programs/clinical_epidemiology/oxford.htm), which assesses cohort and case-control studies and takes 5–10 min to complete. The second, by Downs and Black\textsuperscript{30}, is a longer tool taking about 10–20 min to complete. CMSG authors should also consider the methodological checklists developed by the Scottish Intercollegiate Guidelines Network (www.sign.ac.uk/methodology/checklists.html).

DATA COLLECTION AND ANALYSIS

The CMSG recommends that at least 2 review authors independently extract data from included studies. Data collection forms should be used on all CMSG reviews and it is recommended that they be piloted on a sample of studies. It is important that key characteristics and contextual factors of each study be identified for entry into the “Table of Included Studies.” At the editorial office we have developed data collection forms that can be modified for new reviews.

Priority is given to extracting the information on up to 7 outcomes predefined for inclusion in the SoF table that is presented on the first page of the Cochrane review; however, the review authors should be alert to the possibility of important, unexpected findings, particularly serious adverse effects that may need to be recorded.

It is usually desirable to collect summary data separately for each intervention group and to enter these into RevMan (Review Manager, the software used for preparing and maintaining Cochrane Reviews), where effect estimates can be calculated. Examples are frequency summary data upon which effect estimates such as risk ratio (RR), OR, and risk difference (RD) can be calculated, or mean and SD for continuous data upon which effect estimates such as mean difference (MD) and standardized mean difference (SMD) can be calculated. Chapter 7 in The Cochrane Handbook describes how data should be extracted and converted when necessary to obtain an effect estimate.

Cluster randomized trials are often incorrectly analyzed\textsuperscript{41}. When including cluster randomized trials in a review, we recommend that a CMSG statistical editor be consulted.

Metaanalysis. Because we want to provide the best numerical estimate of the probability of each patient-important outcome, metaanalysis should be undertaken when data are sufficiently clinically homogeneous. Straightforward statistical analyses should be performed using RevMan, if data are available and sufficiently similar. The timing of outcome measures should be provided for the most clinically relevant time frame. It may be appropriate to provide summary estimates for short, medium, or long term, depending on the intervention. For example, in the CMSG network metaanalysis of biologics for RA\textsuperscript{18}, the methods section defined the following timing of outcomes: short (≤ 6 mo), intermediate (> 6 to 12 mo), or long duration (> 1 yr). Estimates based on these different timings were presented in the SoF tables.

Effect measures. The “effect sizes” for dichotomous outcomes may be expressed in RR, OR, Peto OR, or RD. Although absolute differences in important outcomes is the preferred way of presenting the magnitude of the benefit or harm of an intervention to patients and their clinicians, an absolute measure such as the RD is very vulnerable to heterogeneous baseline rates; a relative effect size measure is more stable, so these are preferred for the statistical estimation. The CMSG recommends that RR be used to express dichotomous outcomes because they are easier to understand\textsuperscript{51}. When events are rare, the Peto OR is recommended\textsuperscript{52}. There is no consensus on the definition of “rare;” a working rule of an event rate of < 10% can be used, but special care is needed when studies within the metaanalysis do not have any events.

For continuous outcomes, relative differences are again used, such as MD between the postintervention values, or the difference between baseline values and postintervention values, of the intervention and control groups. SMD should be used when results for continuous outcomes measuring the same concept are presented on different scales; for example, visual analog scale (VAS) and Likert pain scales. One important caveat associated with the use of SMD values in metaanalyses is that few clinicians, patients, journalists, or policy makers understand how to interpret them; we recommend transforming them back to a well-known scale for the SoF table, abstract, and plain language summary (e.g., VAS pain). For examples of this conversion, see Bjirdaal and Christensen\textsuperscript{53}.

Although relative difference metrics are used in the RevMan statistic calculations, patients and their clinicians also need to be provided with absolute differences in the patient-important benefits and harms as listed in the SoF table. The frequency of events without treatment (i.e., the baseline prevalence) makes a marked difference. For example, a relative 50% success in achieving a patient-important reduction in severe pain in a group of 100 patients, 90 of whom report severe pain without the treatment of interest, gives an absolute patient-important reduction in severe pain in 45 patients out of 100 patients with pain. This is substantively different from the same relative 50% success only providing a patient-important reduction in severe pain in 5 of a group of 100 patients when only 10 report severe pain without the treatment of interest.
Subgroup analyses. Subgroups are frequently of clinical or policy importance, e.g., to determine the effects of dosage or disease severity on the response to treatment. The problem is that they may show spurious differences, i.e., by chance alone. However, if the review authors have some clear objectives that justify this in advance — to confirm clinically sound hypotheses — the CMSG endorses this prespecified behavior. Therefore, as few subgroups as possible should be prespecified. These should be justified against the criteria proposed by Sun, et al (Table 2).

Table 2. Criteria to assess the credibility of subgroup analyses. The greater the extent to which these criteria are met, the more plausible the putative subgroup effect.

<table>
<thead>
<tr>
<th>Design</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Is the subgroup variable a characteristic measured at baseline or after randomization?</td>
</tr>
<tr>
<td>• Is the effect suggested by comparisons within rather than between studies?</td>
</tr>
<tr>
<td>• Was the hypothesis specified a priori?</td>
</tr>
<tr>
<td>• Was the direction of the subgroup effect specified a priori?</td>
</tr>
<tr>
<td>• Was the subgroup effect one of a small number of hypothesized effects tested?</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Does the interaction test suggest a low likelihood that chance explains the apparent subgroup effect?</td>
</tr>
<tr>
<td>• Is the significant subgroup effect independent?</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Context</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Is the size of the subgroup effect large?</td>
</tr>
<tr>
<td>• Is the interaction consistent across studies?</td>
</tr>
<tr>
<td>• Is the interaction consistent across closely related outcomes within the study?</td>
</tr>
<tr>
<td>• Is there indirect evidence that supports the hypothesized interaction (biological rationale)?</td>
</tr>
</tbody>
</table>

Another way to interpret an SMD value is to convert it into a number needed to treat (NNT) through a transformation to an OR^{54,55}. This OR can then be combined with an assumed control group risk to obtain an absolute benefit as in NNT^{56}. To do this for continuous or categorical data, the review authors need to estimate a reasonable control event rate — the percentage of patients who would be expected to respond to placebo/sham therapy.

Diversity/heterogeneity of effect sizes across available studies. Following the terminology of the handbook, the terms “heterogeneity” or “diversity” may be used to describe variability among studies included in a systematic review. Clinical diversity (or heterogeneity) is the most important — that is, the variability in participants, interventions (e.g., dose), context, comparator (including differences in “usual care”), and outcomes (both surrogate and clinical). Variability in study design is termed “methodological diversity (or heterogeneity)”. “Statistical heterogeneity” (or conventionally just heterogeneity) is the term used when the variation in intervention effects between studies is greater than that expected by chance. “Inconsistency” is the term used for quantifying the effect of heterogeneity on the metaanalysis.

This issue is characterized by the expression “one cannot combine apples and oranges”. It is important to take an initial look at the results for both clinical diversity and methodological diversity. Clinical diversity is assessed by checking that the patients, interventions, and comparators are not too different from each other such that combining them is clinically useless. Methodological diversity means checking that the studies are similar in terms of study design and risk of bias. Once satisfied that the studies are minimally diverse and that it makes sense to combine them in a metaanalysis, an assessment of the statistical heterogeneity must be undertaken by examining the forest plot and result of the $I^2$ statistic and the $\tau^2$ statistic, described below.

A forest plot provides a visual sense of heterogeneity because one can easily see whether the different point estimates of the effect size of each trial all show either a benefit or harm. RevMan calculates $I^2$ and $\tau^2$ statistics, used to indicate the presence of statistical heterogeneity. The $I^2$ statistic provides an estimate of the between-study variance. The $I^2$ statistic describes the percentage of total variation across studies due to heterogeneity, and it does not inherently depend on the number of studies in the metaanalysis^{58}, although the size of trials included in the metaanalysis should be taken into consideration for proper interpretation^{59}. In Figure 4, at 12 months the effect size has an $I^2$ of 0%, which is consistent with a $\tau^2$ of 0.

If the effects observed across trials are inconsistent and vary to a large extent (say, $I^2 > 50\%$), it is important that the review authors explore the results again and try to assess whether the differences can be explained by some clinical or methodological heterogeneity^{60}. Inconsistency that cannot be explained (i.e., reduced) by prespecified stratified analyses will lead to an overall estimate with less confidence when interpreting the inference from the metaanalysis. In this case, instead of the fixed-effect approach, a suitable, more conservative approach would be a random-effects metaanalysis, so that the between-study variance is considered and the uncertainty of the effect estimate is reflected in wider CI in the model.

Sensitivity analyses. Sensitivity analyses should be performed to examine the strength of the results to risk of bias and the influence of other variables. Authors should prespecify in their protocol which key domains of the
risk-of-bias criteria will be used to perform a sensitivity analysis, by outcome. For example, for each major outcome, those studies contributing data to that outcome that are judged at low risk of bias for the domains of allocation concealment, blinding of patients and outcome assessors, and incomplete outcome data may be compared with all studies to check the strength of the result of including all studies versus a restricted set of studies with a stronger methodological design. Effectiveness/pragmatic studies may need additional sensitivity analyses of considerations such as different populations, differences in interventions, or patient adherence.

**Forest plots.** Using RevMan, the results of individual studies should be presented graphically in forest plots (Figure 4). The overall effect size is shown as a diamond (individual studies as a square), and the horizontal points of the diamond (horizontal line in an individual study) illustrate the 95% CI. The treatment effect is determined by the location of the square in relation to the vertical middle line that indicates the null hypothesis; an effect size is considered to have no statistical significance when the CI crosses the vertical middle line. When appropriate, data from more than 1 trial may be combined in a metaanalysis, and the diamond at the bottom of the graph provides an estimate of effect of this pooled data.

**Grading of the evidence.** In an effort to make it easier for the end user to understand the quality of the evidence or the “degree of confidence” in the reported results included in the review, we recommend that a rating or grade of the evidence for each major outcome be provided in each review. The GRADE approach now replaces the simplified grading system that was derived by the editors of Evidence-based Rheumatology.

The GRADE approach specifies 4 levels of quality: high, moderate, low, and very low to quantify the “degree of confidence” in the reported results per outcome (Table 3). Note that this requires a decision for all the studies included in an SoF table (and hence is distinct from the assessment of the risk of bias or methodological strength of the individual studies). The highest quality rating is for a body of evidence based on data from randomized trials without important limitations and the lowest quality rating is for a body of evidence based on case series/case reports.

A detailed description of the factors that reduce or increase the quality of the evidence is provided in Chapter 12 of the Cochrane Handbook.
These guidelines on developing and performing a systematic review will help improve the quality of reporting and promote high standards of conduct as well as consistency across CMSG reviews.

ACKNOWLEDGMENT

CMSG editors who contributed to the preparation of this manuscript: Isabelle Boutron, Angela Busch, Ernest Choy, Robin Christensen, Rob de Bie, Rhian Goodfellow, Tracey Howe, Anne Lyddiatt, Mário Lenza, Philippe Ravaud, Raphaèle Seror, Beverley Shea, Maria Suarez-Almazor, and Karine Toupin-April.

REFERENCES


Table 3. Levels of quality and definition of a body of evidence in the Grading of Recommendation Assessment Development and Evaluation (GRADE) approach

<table>
<thead>
<tr>
<th>Underlying Methodology</th>
<th>Quality of Rating or Grade</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Randomized trials; or double-upgraded observational studies</td>
<td>High</td>
<td>Further research is very unlikely to change our confidence in the estimate of effect</td>
</tr>
<tr>
<td>Downgraded randomized trials; or single upgraded observational studies</td>
<td>Moderate</td>
<td>Further research may change the estimate (and is likely to have an important influence on our confidence in the estimate of effect)</td>
</tr>
<tr>
<td>Double-downgraded randomized trials; or observational studies</td>
<td>Low</td>
<td>Further research is likely to change the estimate (and is very likely to have an important influence on our confidence in the estimate of effect)</td>
</tr>
<tr>
<td>Triple-downgraded randomized trials; or single downgraded observational studies; or case series/case reports</td>
<td>Very low</td>
<td>Any estimate of effect is very uncertain</td>
</tr>
</tbody>
</table>

Table 4. Factors that may influence the quality level of a body of evidence.

<table>
<thead>
<tr>
<th>Decrease Quality</th>
<th>Increase Quality</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Within study, risk of bias causes the overall analysis to be biased</td>
<td>1. A large magnitude of effect</td>
</tr>
<tr>
<td>2. Indirectness of evidence (i.e., considering each of the following 4 PICO letters — participant, intervention, control, and outcome, especially surrogates). If any of the PICO factors are not directly clinically relevant, the evidence might be judged appropriate for downgrading</td>
<td>2. All plausible confounding would reduce a demonstrated effect or suggest a spurious effect when results show no effect</td>
</tr>
<tr>
<td>3. Unexplained heterogeneity or inconsistency of results (including problems with subgroup analyses, making it difficult to interpret the overall metaanalysis)</td>
<td>3. Evidence of a dose-response gradient</td>
</tr>
<tr>
<td>4. Lack of precision of the overall effect estimates (wide CI), when the overall analysis does not confirm that the effect size is evident</td>
<td></td>
</tr>
<tr>
<td>5. Risk of publication bias</td>
<td></td>
</tr>
</tbody>
</table>

Ghogomu, et al: Cochrane systematic review methods

Personal non-commercial use only. The Journal of Rheumatology Copyright © 2014. All rights reserved.


42. Konnerup M, Kongsted H. Do Cochrane reviews provide a good model for social science? The role of observational studies in systematic reviews. Evid Policy 2012;8:79-96.


