CHARTER:

DEVELOPMENT OF EVIDENCE-BASED PRACTICE GUIDELINES

AND

DEVELOPMENT OF A SYSTEMATIC REVIEW

By the Centre of Evidence-Based Practice (CEBaP) of the Belgian Red Cross-Flanders
INHOUD

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A. INTRODUCTION

BACKGROUND
Belgian Red Cross-Flanders (BRCF) comes up for the rights of vulnerable people (social assistance, relief, training, international aid, blood transfusion,...). The mission of the Centre of Evidence-Based Practice (CEBaP) consists of formulating evidence-based recommendations to support the activities and interventions of BRCF. The recommendations are collected in evidence-based practice guidelines (developed according to AGREE II) and in systematic reviews (developed according to the Cochrane Handbook), thus guaranteeing knowledge dissemination by publication in peer-reviewed journals. The methodology that is being used by CEBaP is described in detail in this charter, and has been published in International Journal of Evidence Based Healthcare [1].

DEFINITIONS

AGREE II
The AGREE (Appraisal of Guidelines for Research & Evaluation) instrument was developed to promote the quality of practice guidelines [2]. It is a tool that assesses the methodological rigour and transparency with which a guideline is developed. The original AGREE instrument has been refined, resulting in the new AGREE II. The purpose of AGREE II is to provide a framework to (1) assess the quality of guidelines, (2) provide a methodological strategy for the development of guidelines and (3) describe what information ought to be reported in guidelines and in which format.

Bias
In the case of bias, the results or the interpretation of a study differ from reality by a systematic error. Bias can be the result of an error in any of the steps of a study, such as preparing the study, collecting data, analysing and interpreting the results and reporting.

Body of evidence
The quality of all available studies (and not of individual studies) is being assessed and summarised.

Cochrane
An international organisation that gives support in making informed decisions about health care by publishing systematic reviews and meta-analyses about the effect of health care interventions.

Evidence
Scientific data to support the answer to a specific question.

Evidence summary
Transparent and structured summaries of scientific literature related to specific practice questions. In an ‘evidence summary’ the following categories are described: topic; subtopic; intervention; question (PICO); search strategy; search date; inclusion and exclusion criteria; characteristics of included
studies; synthesis of findings; quality assessment; conclusion; references. The Evidence summary template can be found in Appendix 1.

Evidence-based methodology
The process by which decisions are being taken based on the best available scientific evidence, practical experience, preferences of the target group and the available resources.

Expert panel
1 chair and additional panel members that as a minimum have expertise related to the content of the project to give evidence-based support for the practice guideline or recommendation.

Forest plot
A graphical representation of the results of different studies included in a meta-analysis. The point estimates and confidence intervals of every study are shown in a horizontal line below each other and beneath them the pooled result is shown as a diamond, the estimate of the global effect.

Funnel plot
A graphical way to assess publication bias while performing a meta-analysis. Therefore, for every study the effect is plotted against the sample size. The distribution of the points in this graph should take the form of a funnel, in which the dispersion increases as the sample size decreases. In case of asymmetry we can assume that studies are missing (for example, these are not published or not included based on the search strategy).

Good practice point
Where no good-quality evidence is available but consensus among experts with practical experience exists, consensus-based recommendations are given. Such recommendations are called 'Good Practice Points' (GPP).

GRADE
A methodology that can be used to assess the level of evidence of studies and the grade of recommendation based on the corresponding evidence.

Grey literature
Publications that are developed by the government, academics, business and industry, which are not published in easily accessible journals and may not appear in databases or via web searches. Examples are: government reports, conference proceedings, abstracts of the research presented at conferences, technical reports, dissertations or other types of documentation.

Meta-analysis
A statistical method in a systematic review in which the results of a number of comparable studies are pooled and recalculated. By doing this, it is possible to better estimate the true "effect size" and to be more confident about the effect of an intervention or treatment.

NICE
The ‘National Institute for Clinical Excellence’, an independent British organization that develops evidence-based clinical guidelines to improve people’s health and prevent and treat ill health.
PICO
A PICO (population [P], intervention [I], comparison [C] and outcome [O]) question is a specific question which precisely defines the population, intervention, comparison and outcome under consideration.

PRISMA
PRISMA or ‘Preferred Reporting Items for Systematic reviews and Meta-Analyses’ is a checklist, composed of 27 items, that aims to increase transparency and clarity when publishing a systematic review (Appendix 2).

Practice guideline
A document with recommendations, advice and instructions to support daily practice, based on the results of scientific research, discussion and decision making, aimed at good practice.

Publication
Each document that is being published in a (scientific) journal, magazine, newspaper or handbook, each document that is submitted to a scientific congress/symposium, and each document that is present on an information carrier (e.g. CD or DVD).

Recommendation
Advice or statement in which a certain technique, intervention, process or activity is being recommended.

SIGN
The ‘Scottish Intercollegiate Guidelines Network’ which develops evidence-based clinical practice guidelines for the National Health Service in Scotland.

Steering committee
The CEBaP Steering committee is composed of the several directors and managers of the Belgian Red Cross-Flanders: the Director of Humanitarian Services Flanders, the manager of the International Humanitarian Services, the Medical Director of the Blood Service and the CEO.

Summary of findings table
Contains the major conclusions in a transparent and simple way. In the table the most important outcomes, the effect sizes, and the number of participants are reported.

Systematic review
Literature summaries that aim to answer a specific question on the effectiveness of interventions by performing a systematic search in available literature. The term ‘systematic’ indicates that specific attention is given to formulating the methods of data collection and handling, in order to provide a transparent methodology to the reader who can then make a judgement about the quality of the literature search. This profound method minimises the risk of bias and results in the “best available scientific evidence”.
SEMANTICS

1. Systematic review

A systematic review gives an overview of the best available scientific evidence collected by a literature search on a very specific topic or question and can be used to inform policy makers.

When using the term ‘systematic review’ the semantics are very important: a systematic review literally means ‘performing a literature review in a systematic way’. However, many variations and gradations exist in performing a systematic literature review. Peer-reviewed publications that call themselves ‘systematic review’, thus may differ significantly in methodological quality.

The Cochrane Collaboration uses the strictest methodological criteria for the development of systematic reviews. Any systematic review developed by BRCF will be performed using the methodological principles of Cochrane. These are described in detail in the charter below.

2. Guideline

In a guideline recommendations for practice are being made, to assist practitioners/volunteers on the field. These recommendations are the result of balancing the quality of the evidence, benefits and harms, costs, and the preferences of the target group.

There is also a range of methodological variants among guidelines. A guideline is not an ‘evidence-based guideline’ by definition. Currently the AGREE II checklist is being used as ‘the golden standard’ by guideline developers [2]. A guideline that is developed according to the criteria in this checklist can be called an ‘evidence-based guideline’.

For the development of practice guidelines BRCF uses the AGREE II checklist. This checklist recommends performing the literature search in a systematic way. However, we do not call this a ‘systematic review’ according to the strict Cochrane definition, because we make a compromise between the number of topics on the one hand and a reasonable time span for the development of the practice guideline on the other (cf. SIGN methodology). This results in a methodology that is systematic but less rigorous than is the case for a Cochrane systematic review: a specific search strategy instead of a sensitive search strategy, 1 reviewer instead of 2 reviewers and not handsearching instead of handsearching for evidence. However, for guideline development, additional expert opinion is added and practical recommendations are being formulated. The methodological principles for guideline development used by BRCF are described in detail in the charter below.

An overview of the different methodological aspects used for either guideline development or the development of a systematic review is given in a table in Appendix 3.

PROJECT INITIATION

Following a specific question from a certain Red Cross Service, the CEBaP performs a scoping review before a practice guideline or systematic review project is initiated.

The aim of this scoping review is to get a first idea about the content, quantity and quality of the available evidence. A quick search methodology is being used when performing a scoping review,
using a specific search strategy and limited to one or two databases (The Cochrane Library, MEDLINE). In order to define the research question as good as possible, the input of the requesting Red Cross Service is included.

After finalising the scoping review, the steering group of the CEBaP decides if:

1) A practice guideline project will be initiated.
2) A systematic review will be developed.
3) No new project will be started up.

In order to make this choice, the following criteria are being taken into account:

- urgency
- potential impact
  - impact on practice, society
  - opportunity for a publication
  - intellectual property
  - quality of the body of evidence
- economic and financial impact on BRCF
- relevance for BRCF (does it fit into our core business, in our strategy?)

A systematic review will only be developed if:

- it can be used for policy change
- the answer to the question is not urgent
- there is a major chance that it will result in a peer-reviewed publication
- the quality of the body of evidence is moderate to high (preferably)

A diagram of the project initiation and project flow can be found in Appendix 4.
B. DEVELOPMENT OF AN EVIDENCE-BASED RECOMMENDATION AND PRACTICE GUIDELINE

CHOICE OF THE SUBJECT
In prioritizing a subject for a recommendation or practice guideline we take into account the following criteria:

- Urgency: high
- Potential impact of the practice guideline
  - Impact on practice, society
  - Opportunity for a publication
  - Intellectual property
- Economic and financial impact for BRCF
- Relevance for BRCF (does it fit into our core business, in our strategy?)

DEVELOPMENT OF AN EVIDENCE-BASED PRACTICE GUIDELINE ACCORDING TO AGREE II
The development of an evidence-based practice guideline by BRCF is based on AGREE II [2]. This instrument offers a framework for the development of qualitative guidelines in which the potential biases of guideline development have been adequately addressed. On the other hand, the recommendations are both internally and externally valid, and are feasible for practice. The assessment includes judgments about the methods used for developing the guidelines, the components of the final recommendations, and the factors that are linked to their uptake.

Based on AGREE II 6 domains are described in detail in all guidelines developed by BRCF:

1) SCOPE AND PURPOSE
A. The overall objective of the guideline. This deals with the potential health impact (e.g. prevention) for the target population and the expected outcome.

B. The PICO question(s) covered by the guideline. This contains the setting or context, the population, the intervention, the comparison and the outcome(s). For an extensive practice guideline, it is possible that in the PICO question describes not one but all effective interventions with relevance to a certain target group. No systematic literature search is started when the PICO concerns:

- A ‘good practice point’ or common sense
- The responsibility of professionals (such as a medical doctor or pharmacist)
- Interventions with only a long-term effect
- The practical organisation of activities
- Medico-legal aspects
- Anatomy or physiology
C. **The target population (patients, public, etc.) to whom the guideline is meant to apply.** This contains the age range, sex, clinical condition (if relevant), the severity/stage of the disease (if relevant), comorbidity (if relevant) and excluded populations (if relevant).

### 2) STAKEHOLDER INVOLVEMENT

A. **The composition of the guideline development group including individuals from all relevant professional groups.** The members of the guideline development group are all those involved at some stage of the development process. This item consists of the name, discipline/content expertise, institution or organisation, geographical location and a description of the member’s role in the guideline development group. The guideline development group consists of: the members of the Steering Committee of the CEBaP, the staff members of the CEBaP that are responsible for collecting the evidence, the Red Cross service for whom the guideline is being developed and who is responsible for formulating the draft recommendations, and the expert panel that makes a trade-off between the quality of the evidence, benefits and harm and validates the final recommendations. The expert panel consists of a chairman, with expertise in evidence-based methodology and on the content of the project, and additional panel members, who at a minimum have expertise in the content of the project.

B. **The views and preferences of the target population.** The guideline development group receives information about the views and preferences of the target population from (1) the Red Cross service involved, which has expertise in the content or collects the necessary information (e.g. by composing a reading group or by interviewing the target population) (2) a literature search concerning the values, preferences and experiences of the target population and/or (3) a feedback round or pilot test. In addition the target population is represented in the guideline development group.

C. **The target users of the guideline.** This topic consists of a description of the target group and of how the practice guideline may be used by its target audience.

### 3) RIGOUR OF DEVELOPMENT

The ‘rigour of development’ domain consists of the literature search, the formulation of draft recommendations, the review of the guideline and the updating of the guideline.

#### 3.1 SYSTEMATIC LITERATURE SEARCH

A. **Systematic methods to search for evidence.** We use a stepwise search strategy for collecting literature, which is described in detail. The search strategy consists of the sources of literature, the search terms, the use of methodological filters (if relevant) and the period from which articles are retrieved.

In AGREE II no detailed description is available of the methodology for the literature search. Therefore we based our methodology on that used by SIGN (Scottish Intercollegiate Guidelines Network) and NICE (National Institute for Health and Clinical Excellence) (‘SIGN 50: A guideline

Sources:

- Guidelines: NGC (National Guideline Clearinghouse), GIN (Guidelines International Network) and MEDLINE (via PubMed interface).
- Systematic reviews: the Cochrane Database of Systematic Reviews (via Wiley interface), Database of Abstracts and Reviews of Effects (DARE - via Wiley interface), MEDLINE (via PubMed interface) and BestBETs (containing pragmatic systematic reviews).
- Experimental studies: Cochrane Central Register of Controlled Trials (via Wiley interface), MEDLINE (via PubMed interface), Embase (via Embase.com interface). Optionally: the choice of databases can be expanded if this is relevant for the search question (e.g. SPORTDiscus or PEDro for a practice guideline on the prevention of sports injuries).
- Observational studies: MEDLINE (via PubMed interface) and Embase (via Embase.com interface).

Search terms: The search terms (can) differ for every source (e.g. database of guidelines vs. database of individual studies), but will be described in detail for every source, in order to make the search reproducible (e.g. for the updating of the guideline). For the choice of the search terms, we pay attention to possible synonyms and we consult the MeSH thesaurus to identify possible related terms.

Methodological filters:

If possible we try to avoid the use of methodological filters.

- Guidelines: A non-validated filter for guidelines:
  - Embase: ‘practice guideline’/exp OR ‘practice guideline’ OR ‘practice guidelines’

- Systematic reviews: based on a SIGN (Scottish Intercollegiate Guidelines Network - http://www.sign.ac.uk/) filter for systematic reviews

- Experimental studies: based on an EPOC (Effective Practice and Organisation of Care - http://epoc.cochrane.org/) filter for experimental studies

- Observational studies: a SIGN (Scottish Intercollegiate Guidelines Network - http://www.sign.ac.uk/) filter for observational studies; a validated filter described by Deville et al. [3] for diagnostic studies; a validated filter described by Wilczynski et al. for prognostic studies [4; 5].
**Search period:** We search for guidelines, systematic reviews, experimental studies and observational studies from the date of inception of the database until the date of the current search.

The **search strategy:** this takes into account that a practice guideline of BRCF consists of many different topics (> 40 topics) and therefore makes a compromise between the number of topics on the one hand and a reasonable time span for the development of the practice guideline on the other (cf SIGN methodology).

- The search strategy is developed by 1 reviewer and evaluated by a second reviewer.
- In a first step we search for guidelines as a source of systematic reviews and individual studies. In this step we also search for systematic reviews as a source of individual studies. In a next step we search for controlled experimental studies. In a third step we search for controlled observational studies. We only go to the next step of the search strategy if no evidence is found (cf. SIGN methodology) or if the evidence cannot be included based on the inclusion and exclusion criteria (see B).
- During the search for guidelines, systematic reviews, experimental studies and observational studies, additional references can be selected by checking the 20 related citations in PubMed and/or handsearching (e.g. in the reference list of an included reference). The additional references (guideline, systematic review, experimental study, observational study) are assessed with the inclusion and exclusion criteria.
- 1 reviewer for each topic selects and evaluates the evidence and describes the literature search in an ‘evidence summary’. The evidence summary is made in a standard template (see Appendix 1). As an internal control the selection of evidence of a random selection of questions is performed periodically (cf. NICE methodology).

**B. Inclusion and exclusion criteria for selecting evidence.** These include criteria for the language, criteria for the content (population, intervention, outcomes and the context), and methodological criteria. No articles are selected if the intervention concerns:

- A ‘good practice point’ or common sense
- The responsibility of professionals (such as a medical doctor or pharmacist)
- Interventions with only a long-term effect
- The practical organisation of activities
- Medico-legal aspects
- Anatomy or physiology

If we search for risk factors, no articles are selected if the risk factor:

- does not precede the outcome
- is common sense
- is a fixed marker (can be project-dependent)
- is no proximal risk factor (can be project-dependent)
• is not valid for healthy people (can be project-dependent)

English, Dutch, French and German literature is selected.

No animal studies, ex vivo or in vitro studies are selected, but depending on the subject we can make an exception to this. Arguments for making this choice are described in the practice guideline.

Methodological criteria depend on the type of study design:

- A guideline: inclusion if the guideline gets an acceptable score in the ‘rigour of development’ domain of AGREE II.
- A systematic review: inclusion if the search strategy and selection criteria are clearly described and if at least the Cochrane Library, MEDLINE and Embase are searched.
- An experimental study: inclusion in case of one of the following study types: (quasi or non-) randomised controlled trial, controlled before and after study or controlled interrupted time series, and the data are available. These study types can be identified using the flowchart in Appendix 5.
- An observational study: inclusion in case of one of the following study types: cohort and case-control study, controlled before and after study or controlled interrupted time series, and the data are available. These study types can be identified using the flowchart in Appendix 5. Depending on the project and context it can be decided to include cross sectional, diagnostic and/or prognostic studies. For all study types the control group should be clearly described in the methods section.

C. The strengths and limitations of the body of evidence. The strengths and limitations of the body of evidence are being assessed using the GRADE approach [6].

The search strategy, inclusion and exclusion criteria, study characteristics, study findings and levels of evidence are described in an ‘evidence summary’. Standard wording for the evidence conclusions in this evidence summary are described in Appendix 6.

3.2 FORMULATING RECOMMENDATIONS

A. The methods for formulating recommendations are clearly described. This topic consists of a clear description of the method that is used for formulating the recommendations, the outcomes of the recommendation development process and a description of how the process influenced the recommendations. A multidisciplinary expert panel is involved in formulating the final recommendations by informal consensus. If a consensus cannot be reached, the decision depends on the opinion of the majority by voting. The expert panel is responsible for reading through the whole guideline and for the assignment of the grades of recommendation.
B. The health benefits, side effects and risks. This topic consists of a description of the trade-off between benefits and harm, side effects or risks in formulating the final recommendations. The multidisciplinary expert panel makes this judgement during the assignment of the grades of recommendation. Therefore, the method of GRADE is being used.

C. Explicit link between the recommendations and the supporting evidence. This consists of a description of how the guideline development group links and uses the evidence to make informed recommendations. During the formulation of the final recommendations and the assignment of the grades of recommendation, the expert panel makes use of a table in which the corresponding evidence is presented for every draft recommendation.

3.3 REVIEW OF THE PRACTICE GUIDELINE

External review by experts. The external experts or peer reviewers are not involved in the guideline development group and can be target population representatives. Reviewers include experts in the clinical area as well as some methodological experts. For every practice guideline the group of reviewers consists of at least 1 expert on the content and 1 methodological expert who is preferably also an expert on the content.

The purpose and intent, methods for the external review, outcomes/information gathered from the external review and a description of how the information was used to inform the guideline, are described in the practice guideline. Furthermore, the name, discipline/content expertise and institution or organization are given.

3.4 UPDATE OF THE PRACTICE GUIDELINE

A procedure for updating the guideline. The practice guideline will be updated every 5 years, unless stated otherwise. To achieve this the literature search will be performed again from the end of the previous literature search until the start of the update.

4) CLARITY OF PRESENTATION

A. The recommendations are specific and unambiguous. This implies a precise description of which option is appropriate in which situation and in what population group. If evidence is inconsistent, this is described in the evidence summaries. In that case, the expert panel decides which option is the most appropriate.

B. The different options for management. The different possible interventions are presented in the guideline.

C. Key recommendations are easily identifiable.

5) APPLICABILITY

Depending on the type of project, context and target group, we can decide to complement the practice guideline with an implementation guide. This implementation guide can contain the following information: the facilitators and barriers to the application of the guideline, advice on how
to put the recommendations into practice, the potential resource implications of applying the recommendations and monitoring and/or auditing criteria.

6) EDITORIAL INDEPENDENCE

A. Name of the funding body (BRCF) and statement that the content of the guideline is not influenced by the one that finances the project.

B. Statement of competing interests of guideline development group members.

PUBLICATION OF A PRACTICE GUIDELINE

In the publication of the practice guideline the topics mentioned above are preferably described in detail. In every case the methodology is described or we refer to a document containing the detailed methodology.
C. DEVELOPMENT OF A SYSTEMATIC REVIEW

CHOICE OF THE SUBJECT

During the development of evidence-based recommendations or practice guidelines it becomes clear where an up-to-date overview on the effectiveness of interventions is lacking. In these cases the Steering Committee of the CEBaP can decide to develop a systematic review.

In choosing a subject for a systematic review we take into account the following criteria:

- Urgency: low
- Potential impact of the practice guideline
  - Impact on practice, society
  - Opportunity for a publication in a peer-reviewed journal with a high impact factor (ISI Web of Knowledge – Journal Citation Report)
  - Intellectual property
  - Quality of the body of evidence
- Economic and Financial impact on BRCF
- Relevance for BRCF (does it fit into our core business, in our strategy?)

If more topics are suitable subjects of an evidence-based recommendation or practice guideline, we use the same criteria for prioritization of the question.

DEVELOPMENT OF A SYSTEMATIC REVIEW ACCORDING TO THE COCHRANE HANDBOOK

The systematic review provides an overview of the best available evidence collected by a literature search on a very specific topic or question. A systematic review can be used to inform policy makers. Making a trade-off between the estimated benefits, harm and the estimated costs, and thus making specific recommendations for an action, goes beyond the scope of a systematic review and is typically the task of guideline development group.

The systematic review will be included in the practice guidelines of BRCF when developing or updating a guideline.

For the development of a systematic review we follow the methodology described in the Cochrane handbook (version 5.1.0) (http://www.cochrane-handbook.org/). Below, only the differences with the search strategy as described for practice guideline development will be highlighted.

1) DEFINING THE REVIEW QUESTION AND DEVELOPING CRITERIA FOR INCLUDING STUDIES

A focused research question is described in detail as a PICO question, in which the population, intervention, comparison and outcome are clearly indicated.
In addition, the types of study to be included as the source of evidence are clearly specified. In making this choice we consider a priori what study designs are likely to provide reliable data with which to address the objectives of the review.

2) SEARCHING FOR STUDIES
In the search for studies we closely cooperate with the Trials Search Co-ordinator of the corresponding Cochrane Review Group.

At least the following databases are searched: the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE and Embase. Efforts are made to identify unpublished studies, conference abstracts, grey literature and ongoing trials.

We use a very sensitive search strategy. In the case of using methodological filters, the sensitive filters of Cochrane are used.

3) SELECTING STUDIES AND COLLECTING DATA
The study selection and data extraction are performed by at least 2 independent reviewers. A clear procedure for action is described in case of disagreement between the 2 reviewers.

Preferably the authors of studies are contacted when information in the study is missing.

4) ASSESSING RISK OF BIAS IN INCLUDED STUDIES
The following topics, which may be the source of risk of bias are assessed for every study: sequence generation (selection bias), allocation sequence concealment (selection bias), blinding of participants/personnel (performance bias), blinding of outcome assessment (detection bias), incomplete outcome data (attrition bias), selective outcome reporting (reporting bias) and other potential sources of bias.

To assess the quality and risk of bias in studies we use the ‘Cochrane Collaboration’s tool for assessing risk of bias’.

5) ANALYSING DATA AND UNDERTAKING META-ANALYSIS
If possible a meta-analysis will be conducted to statistically combine the data of several included studies. Heterogeneity between studies is determined statistically. In case of heterogeneity we search for an adequate way to address it (e.g. subgroup analysis).

If a meta-analysis is undertaken, we will also perform a sensitivity analysis to find out if the findings of the systematic review are sufficiently robust.

6) ADDRESSING REPORTING BIASES
If sufficient studies are included, a funnel plot can be used to search for reporting bias.
7) PRESENTING RESULTS AND SUMMARY OF FINDINGS TABLES

The characteristics of the studies are given in a table (‘Characteristics of included studies’). The results, i.e. the effect measures and confidence intervals, of the individual studies are shown in a ‘summary of findings’ table. If possible, meta-analyses are generated and presented in a forest plot. The quality of the individual studies is summarized in a ‘quality of evidence’ table.

For every outcome of each intervention, a level of evidence is assigned to the body of evidence according to the GRADE approach and the evidence conclusion is formulated using a standard wording (Appendix 6).

8) INTERPRETING RESULTS AND DRAWING FINAL CONCLUSIONS FOR PRACTICE

Drawing final conclusions about the practical usefulness of an intervention entails making trade-offs between benefits, harm and costs. Finding this balance, and thus making specific recommendations, goes beyond a systematic review and is typically part of the development of a practice guideline.

In a Cochrane review a suggestion can be made on potential implications for practice by highlighting different possible actions. In addition, the implications for research can be discussed.

At this point of the review, it may be beneficial to call on the expertise of relevant external consultants to provide an independent appraisal of the quality and relevance of particular aspects of the review.

PUBLICATION OF A SYSTEMATIC REVIEW

For transparent reporting of the development of a systematic review, we use the PRISMA statements 2009 (http://www.prisma-statement.org/statement.htm). This is a 27-item checklist that aims to guarantee the quality of systematic reviews by clear and transparent reporting in a publication. The checklist can be found in Appendix 2.
D. KNOWLEDGE MANAGEMENT

EVIDENCE SUMMARY DATABASE
Each systematic literature search is documented in an evidence summary (Appendix 1) in the evidence summary database on the CEbaP project site. The evidence summary database is developed in Sharepoint and it ensures rapid document retrieval and offers the ability for future database enhancement.

GUIDELINES DATABASE
For each evidence-based guideline, a guideline summary (Appendix 1) is included in the guideline database. This summary contains a synthesis of the expert opinion and practical experience of the experts in the expert panel. A link to the evidence base, the rationale for the recommendation and the final recommendation including a validation date is provided (validation by the expert panel).
E. REFERENCES


# Appendix 1: Evidence Summary and Guideline Summary Template

## Evidence Summary

<table>
<thead>
<tr>
<th>Topic</th>
<th>Subtopic</th>
<th>Intervention</th>
<th>Question (PICO)</th>
<th>Search Strategy</th>
<th>Search date</th>
<th>In/Exclusion criteria</th>
</tr>
</thead>
</table>

### Characteristics of included studies

<table>
<thead>
<tr>
<th>Author, year</th>
<th>Study design</th>
<th>Population</th>
<th>Comparison/Risk factor</th>
<th>Remarks</th>
</tr>
</thead>
</table>

### Synthesis of findings

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Comparison/Risk factor</th>
<th>Effect Size</th>
<th>#studies, # participants</th>
<th>Reference</th>
</tr>
</thead>
</table>

### Quality of evidence

**Version 1: Quality of Experimental studies**

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Lack of allocation concealment</th>
<th>Lack of blinding</th>
<th>Incomplete accounting of outcome events</th>
<th>Selective outcome reporting</th>
<th>Other limitations</th>
</tr>
</thead>
</table>

**Version 2: Quality of Observational studies**

<table>
<thead>
<tr>
<th>Author, Year</th>
<th>Inappropriate eligibility criteria</th>
<th>Inappropriate methods for exposure and outcome variables</th>
<th>Not controlled for confounding</th>
<th>Incomplete or inadequate follow-up</th>
<th>Other limitations</th>
</tr>
</thead>
</table>

### Level of the body of evidence

<table>
<thead>
<tr>
<th>Limitations of study design</th>
<th>[Initial grading]</th>
<th>Downgrading due to</th>
</tr>
</thead>
<tbody>
<tr>
<td>Imprecision</td>
<td>0</td>
<td>See table ‘Quality of evidence’</td>
</tr>
<tr>
<td>Inconsistency</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Indirectness</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>Publication bias</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>QUALITY (GRADE)</th>
<th>[Final grading]</th>
</tr>
</thead>
</table>

### Conclusion(s)

### Reference(s)

---

*Belgian Red Cross-Flanders*

Effectiviteitsdatum: 27-12-2013
<table>
<thead>
<tr>
<th>Evidence used for</th>
<th>Project</th>
<th>Reviewer(s)</th>
<th>URL</th>
<th>Omleidings-URL2</th>
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</thead>
</table>

**GUIDELINE**

<table>
<thead>
<tr>
<th>Topic</th>
<th>Subtopic</th>
<th>Department</th>
<th>Project</th>
</tr>
</thead>
<tbody>
<tr>
<td>GUIDELINE</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Evidence summaries (basis for recommendation) |
| In- en exclusiecriteria |

| Expert panel |

**Rationale for the recommendation**

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Quality of evidence</th>
<th>Balance of benefits/harms and burdens</th>
<th>Values and preferences</th>
<th>Resource implications</th>
</tr>
</thead>
</table>

**Practical recommendations**

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Recommendation</th>
<th>Grade of Recommendation</th>
</tr>
</thead>
</table>

**Date of validation**

Effectiviteitsdatum: 27-12-2013
## APPENDIX 2: PRISMA CHECKLIST

<table>
<thead>
<tr>
<th>Section/topic</th>
<th>#</th>
<th>Checklist item</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Title</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Title</td>
<td>1</td>
<td>Identify the report as a systematic review, meta-analysis, or both.</td>
</tr>
<tr>
<td><strong>Abstract</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Structured summary</td>
<td>2</td>
<td>Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.</td>
</tr>
<tr>
<td><strong>Introduction</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rationale</td>
<td>3</td>
<td>Describe the rationale for the review in the context of what is already known.</td>
</tr>
<tr>
<td>Objectives</td>
<td>4</td>
<td>Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).</td>
</tr>
<tr>
<td><strong>Methods</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Protocol and registration</td>
<td>5</td>
<td>Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.</td>
</tr>
<tr>
<td>Eligibility criteria</td>
<td>6</td>
<td>Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.</td>
</tr>
<tr>
<td>Information sources</td>
<td>7</td>
<td>Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.</td>
</tr>
<tr>
<td>Search</td>
<td>8</td>
<td>Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.</td>
</tr>
<tr>
<td>Study selection</td>
<td>9</td>
<td>State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).</td>
</tr>
<tr>
<td>Data collection process</td>
<td>10</td>
<td>Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.</td>
</tr>
<tr>
<td>Data items</td>
<td>11</td>
<td>List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.</td>
</tr>
<tr>
<td>Risk of bias in individual studies</td>
<td>12</td>
<td>Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.</td>
</tr>
<tr>
<td>Summary measures</td>
<td>13</td>
<td>State the principal summary measures (e.g., risk ratio, difference in means).</td>
</tr>
</tbody>
</table>
| Synthesis of results  | 14| Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., \( I^2 \)) for
<table>
<thead>
<tr>
<th>Section</th>
<th>Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Results</strong></td>
<td></td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Study selection</td>
<td>17</td>
<td>Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.</td>
</tr>
<tr>
<td>Study characteristics</td>
<td>18</td>
<td>For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.</td>
</tr>
<tr>
<td>Risk of bias within studies</td>
<td>19</td>
<td>Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).</td>
</tr>
<tr>
<td>Results of individual studies</td>
<td>20</td>
<td>For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.</td>
</tr>
<tr>
<td>Synthesis of results</td>
<td>21</td>
<td>Present results of each meta-analysis done, including confidence intervals and measures of consistency.</td>
</tr>
<tr>
<td>Risk of bias across studies</td>
<td>22</td>
<td>Present results of any assessment of risk of bias across studies (see Item 15).</td>
</tr>
<tr>
<td>Additional analysis</td>
<td>23</td>
<td>Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).</td>
</tr>
<tr>
<td><strong>Discussion</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Summary of evidence</td>
<td>24</td>
<td>Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).</td>
</tr>
<tr>
<td>Limitations</td>
<td>25</td>
<td>Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias).</td>
</tr>
<tr>
<td>Conclusions</td>
<td>26</td>
<td>Provide a general interpretation of the results in the context of other evidence, and implications for future research.</td>
</tr>
<tr>
<td><strong>Funding</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Funding</td>
<td>27</td>
<td>Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.</td>
</tr>
</tbody>
</table>
### APPENDIX 3: TABLE OF METHODOLOGICAL PRINCIPLES OF GUIDELINE DEVELOPMENT VERSUS SYSTEMATIC REVIEW DEVELOPMENT (BRCF)

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Practice guideline</th>
<th>Systematic review</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Number of reviewers:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>- building search strategy</td>
<td>-1 reviewer</td>
<td>-2 independent reviewers</td>
</tr>
<tr>
<td>- literature search + selection articles</td>
<td>-1 reviewer; for random topics: 2 reviewers</td>
<td>-2 independent reviewers</td>
</tr>
<tr>
<td>- data extraction</td>
<td>-1 reviewer</td>
<td>-2 independent reviewers</td>
</tr>
<tr>
<td>- quality assessment</td>
<td>-1 reviewer</td>
<td>-2 independent reviewers</td>
</tr>
<tr>
<td><strong>Search formula</strong></td>
<td>Specific search formula; use of methodological filters if necessary</td>
<td>Sensitive search formula</td>
</tr>
<tr>
<td><strong>Databases</strong></td>
<td>GIN, NGC, The Cochrane Library, BestBETs, MEDLINE, Embase if necessary topic-specific databases can be added. No unpublished studies, conference abstracts, grey literature and ongoing trials will be identified.</td>
<td>At least: Central, MEDLINE, Embase Efforts are made to identify conference abstracts, grey literature and ongoing trials.</td>
</tr>
<tr>
<td><strong>Selection criteria with respect to study design</strong></td>
<td>Guidelines, systematic reviews, experimental studies and observational studies We only go to lower type of study design if no evidence is found or if the evidence cannot be included based on the inclusion and exclusion criteria.</td>
<td>Experimental studies or observational studies In making this choice it is considered a priori what study designs are likely to provide reliable data with which to address the objectives of the review.</td>
</tr>
<tr>
<td><strong>Selection criteria with respect to content</strong></td>
<td>Only the most direct and important factors are considered in the inclusion criteria</td>
<td>Depending on the research question</td>
</tr>
<tr>
<td><strong>Quality assessment</strong></td>
<td>Of the ‘body of evidence’</td>
<td>Of each outcome separately</td>
</tr>
<tr>
<td><strong>Meta-analysis</strong></td>
<td>no</td>
<td>If possible</td>
</tr>
<tr>
<td><strong>Involvement of experts</strong></td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td><strong>Formulating recommendations</strong></td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td><strong>Assessment grades of recommendation</strong></td>
<td>yes</td>
<td>no</td>
</tr>
<tr>
<td><strong>External peer review</strong></td>
<td>yes</td>
<td>no</td>
</tr>
</tbody>
</table>
APPENDIX 4: PROJECT INITIATION AND PROJECT FLOW

QUESTION

Input: Operational Red Cross Service
PICO - question(s)

LITERATURE REVIEW

Input: CEBaP Steering group considering:
- Urgency
- Potential impact on practice and society
- Economic and financial impact on BRC-F
- Relevance for BRC-F

Guideline:
multiple reviews
Highly relevant with potential high impact
Urgency: high

Systematic review:
one (highly sensitive) review
Highly relevant with potential high impact
(In addition: use for policy change, acceptable evidence quality, chance for peer-review publication)
Urgency: Low

No new project:
no further review
Relevant for Internal
BRC-F decision(s)
Not relevant for BRC-F with (no) potential impact

EXPERT

Multidisciplinary expert panel
Expert(s)

RESULT

Evidence-based guideline (according to AGREE II)
Systematic review (according to Cochrane standards)
Action: internal decision(s)
No action
APPENDIX 5: FLOWCHART STUDY DESIGNS

Start

Information on outcome and exposure collected simultaneously?

N

Y

Cross-sectional study

More than 1 group studied?

N

Y

Case report, case series
Before-after study
Interrupted time series

Interventions/exposures assigned randomly?

N

Y

Case-control study

Groups defined by interventions/exposures?

N

Y

Controlled before-after study
OR
Controlled interrupted time series

Was there a comparison over time?

N

Y

Information on outcome and exposure collected simultaneously?

N

Y

Non-randomised trial

Cohort study

Interventions/exposures (attempted) assigned randomly?

N

Y

Cohort study

Controlled before-after study
OR
Controlled interrupted time series
APPENDIX 6: STANDARD WORDING OF EVIDENCE CONCLUSIONS

A) No Evidence
No studies on the effect of the intervention on the outcome, and fulfilling the selection criteria, were found: “No relevant studies were identified using the above search strategy and criteria.”

B) Conflicting evidence
The results of the different studies are not consistent and no explanation for the differences can be found: “There is conflicting evidence from # experimental studies and/or # observational studies...”

C) (Limited) evidence:

<table>
<thead>
<tr>
<th>P-value</th>
<th>Level of evidence</th>
<th>Imprecision (GRADE methodology)</th>
<th>Evidence conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; α (significant)</td>
<td>A or B</td>
<td>No</td>
<td>There is evidence from # experimental studies and/or # observational studies in favour of [intervention] (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; resulted in a statistically significant increase/decrease of &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of high/moderate quality.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Yes</td>
<td>There is limited evidence from # experimental studies and/or # observational studies in favour of [intervention] (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; resulted in a statistically significant increase/decrease of &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of moderate quality and results cannot be considered precise due to limited sample size, lack of data and/or large variability of results.</td>
</tr>
<tr>
<td>C or D</td>
<td>No</td>
<td></td>
<td>There is limited evidence from # experimental studies and/or # observational studies in favour of [intervention] (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; resulted in a statistically significant increase/decrease of &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of low/very low quality.</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td></td>
<td>There is limited evidence from # experimental studies and/or # observational studies in favour of [intervention] (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; resulted in a statistically significant increase/decrease of &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of low/very low quality and results cannot be considered precise due to limited sample size, lack of data and/or large variability of results.</td>
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<tr>
<td>P-value</td>
<td>Level of evidence</td>
<td>Imprecision (GRADE methodology)</td>
<td>Evidence conclusion</td>
</tr>
<tr>
<td>---------</td>
<td>-------------------</td>
<td>---------------------------------</td>
<td>---------------------</td>
</tr>
<tr>
<td>&gt; α (not significant)</td>
<td>A or B</td>
<td>No</td>
<td>There is evidence from # experimental studies and/or # observational studies, showing no difference between intervention and control (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; did not result in a statistically significant difference in &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of high/moderate quality.</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td></td>
<td>There is limited evidence from # experimental studies and/or # observational studies, neither in favour of the intervention nor the control (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). A statistically significant increase/decrease on &lt;outcome&gt;, using &lt;intervention&gt; compared to &lt;comparison&gt;, could not be demonstrated (&lt;Author&gt; &lt;year&gt;). Evidence is of moderate quality and results of this study/these studies are imprecise due to limited sample size, lack of data and/or large variability of results.</td>
</tr>
<tr>
<td>C or D</td>
<td>No</td>
<td></td>
<td>There is limited evidence from # experimental studies and/or # observational studies, neither in favour of the intervention nor the control (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). It was shown that &lt;intervention&gt; did not result in a statistically significant increase/decrease of &lt;outcome&gt;, compared to &lt;comparison&gt; (&lt;Author&gt; &lt;year&gt;). Evidence is of low/very low quality.</td>
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<td></td>
<td>Yes</td>
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<td>There is limited evidence from # experimental studies and/or # observational studies, neither in favour of the intervention nor the control (&lt;Author&gt; &lt;year&gt;, &lt;Author&gt; &lt;year&gt;, etc.). A statistically significant increase/decrease on &lt;outcome&gt;, using &lt;intervention&gt; compared to &lt;comparison&gt;, could not be demonstrated (&lt;Author&gt; &lt;year&gt;). Evidence is of low/very low quality and results of this study/these studies are imprecise due to limited sample size, lack of data and/or large variability of results.</td>
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</tbody>
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