SIGN 50

A guideline developer’s handbook

October 2014
Scottish Intercollegiate Guidelines Network

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## Complying with international standards

SIGN seeks to ensure that its methodology complies with international standards as far as possible within its resources. Although there are now a number of published standards for guideline development methodology, SIGN regards AGREE II (Appraisal of Guidelines for Research and Evaluation; [www.agreetrust.org](http://www.agreetrust.org)) as the most evidence based of these. The sections in SIGN 50 that address each criterion are identified below.

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1 Introduction

1.1 AIM AND STRUCTURE OF THIS MANUAL

The principal aim of this manual is to provide a reference tool that may be used by individual members of guideline development groups as they work through the development process. SIGN 50 outlines the key elements of the development process common to all SIGN guidelines. Only where aspects of the topic under consideration require a variation from the standard process will these be reported in the guidelines themselves.

A secondary aim of this manual is to be transparent about the methods used to develop SIGN guidelines, and to instil confidence that the potential biases of guideline development have been addressed adequately, and that the recommendations are both internally and externally valid, and feasible for practice.

1.1.1 REVIEW AND UPDATING OF THIS MANUAL

It is intended that SIGN 50 should be a ‘living’ publication, continually revised to reflect future developments in SIGN methodology. For this reason the definitive version of this handbook is that published on the SIGN website. Comments on either content or presentation of this document are welcome and should be sent to the SIGN Executive, Gyle Square, 1 South Gyle Crescent, Edinburgh, EH12 9EB, email: sign@sign.ac.uk

1.2 CLINICAL GUIDELINES AND SIGN

The Scottish Intercollegiate Guidelines Network (SIGN) was established in 1993 by the Academy of Royal Colleges and their Faculties in Scotland, to develop evidence based clinical guidelines for the National Health Service in Scotland.1,2 Since January 2005 SIGN has been part of NHSScotland, although SIGN retains editorial independence in relation to the guidelines it produces.

The role of guidelines has been described variously over the past two decades. A recent definition of clinical practice guidelines (CPGs) states “CPGs are able to enhance clinician and patient decision making by clearly describing and appraising the scientific evidence and reasoning (the likely benefits and harms) behind clinical recommendations, making them relevant to the individual patient encounter”.3

The accepted criteria for validity of guidelines were first set out as the “essential elements of good guidelines” by the US Institute of Medicine in 1990.4 These recommended “attributes of good guidelines” including validity, reliability, clinical applicability, clinical flexibility, clarity, multidisciplinary process, scheduled review, and documentation. The recommendations were underpinned by the twin themes of credibility and accountability: “The link between a set of guidelines and the scientific evidence must be explicit, and scientific and clinical evidence should take precedence over expert judgement.”4 These attributes have formed the basis of SIGN methodology.

The AGREE (Appraisal of Guidelines for Research and Evaluation) guideline appraisal instrument identifies criteria by which the quality of guideline development may be judged. The AGREE II criteria are reproduced in the introductory material to this manual, with links to those manual sections that explain how SIGN addresses each criterion. The full appraisal instrument can be downloaded from the AGREE website: www.agreetrust.org

1.3 GUIDELINES IN CONTEXT

Guideline development, implementation and review should be seen not as a linear process, but as a cycle of interdependent activities. These in turn are part of a range of complementary activities to translate evidence into practice, set and monitor standards, and promote clinical excellence in NHSScotland, as illustrated in Figure 1-1.
1.4 MEDICO-LEGAL IMPLICATIONS OF SIGN GUIDELINES

Although there has been ongoing discussion about the legal status of clinical guidelines, SIGN guidelines are intended as an aid to clinical judgement not to replace it. Guidelines do not provide the answers to every clinical question, nor guarantee a successful outcome in every case. The ultimate decision about a particular clinical procedure or treatment will always depend on each individual patient's condition, circumstances and wishes, and the clinical judgement of the healthcare team.

To clarify the legal position, all SIGN guidelines carry the following statement of intent:

*This guideline is not intended to be construed or to serve as a standard of care. Standards of care are determined on the basis of all clinical data available for an individual case and are subject to change as scientific knowledge and technology advance and patterns of care evolve. Adherence to guideline recommendations will not ensure a successful outcome in every case, nor should they be construed as including all proper methods of care or excluding other acceptable methods of care aimed at the same results. The ultimate judgement must be made by the appropriate healthcare professional(s) responsible for clinical decisions regarding a particular clinical procedure or treatment plan. This judgement should only be arrived at following discussion of the options with the patient, covering the diagnostic and treatment choices available. It is advised, however, that significant departures from the national guideline or any local guidelines derived from it should be fully documented in the patient's case notes at the time the relevant decision is taken.*
1.5 ORGANISATION OF THE SCOTTISH INTERCOLLEGIATE GUIDELINES NETWORK

Since its establishment in 1993, SIGN has been a collaborative initiative; a network of clinicians and other healthcare professionals, and currently includes all the medical specialties, nursing, pharmacy, dentistry, allied health professions, public partners and Healthcare Improvement Scotland (HIS).

The overall structure of the organisation is shown in Figure 1-2.

Figure 1-2: The structure of SIGN

1.5.1 SIGN COUNCIL

SIGN Council is the policymaking body for SIGN with overall responsibility for methodology and editorial policy. Although SIGN forms part of the Evidence and Technologies Unit at HIS, SIGN is editorially independent from HIS and the Scottish Government which ultimately funds HIS.

Members of SIGN Council are nominated by a particular Royal College or other professional organisation or committee, but also represent their specialty or discipline in a wider sense and consult widely with other specialist societies in their field. Public partners are identified from an open call for interested individuals.

Members of SIGN Council determine the overall direction of SIGN’s development and play a key role in shaping the SIGN guideline programme. Some are also actively involved in aspects of the guideline development process and all provide input into the selection of topics for guideline development and the composition of guideline development groups. The current membership of SIGN Council is noted on the SIGN website.

1.5.2 SIGN EXECUTIVE

The SIGN Executive is made up of a Programme Team working closely with the HIS Knowledge and Information Unit. Together they are responsible for the implementation of decisions taken by SIGN Council and its subgroups, and for delivering the guideline programme to time and on budget. All staff are employees of Healthcare Improvement Scotland.
1.6 FUNDING

Core funding from Healthcare Improvement Scotland supports the SIGN Executive, expenses associated with individual guideline development projects (meeting and literature costs) and the costs of printing and distributing published SIGN guidelines.

Members of SIGN guideline development groups do not receive any payment for their participation, although independent practitioners are entitled to claim locum payments and travel expenses. Patient representatives can also claim travel, subsistence, child care/carer expenses and any other reasonable out of pocket expenses to enable them to attend guideline development group meetings. The expenses of other members of SIGN guideline development groups are met by their employing NHS boards, under an agreement with the Scottish Government Health Directorate.

1.7 INFLUENCE OF FINANCIAL AND OTHER INTERESTS

All individuals involved in the SIGN guideline development process must declare any competing interests (financial and non-financial) on at least an annual basis. This includes all of the following:

- SIGN Council and subcommittee members and deputies
- SIGN Executive staff
- speakers at SIGN events
- guideline development group members
- peer reviewers
- all who submit proposals to SIGN.

Signed copies are retained by the SIGN Executive and are available on the SIGN website.

Full details of the declarations of interest policy are available on the SIGN website.

1.8 REFERENCES


2 The guideline development group

2.1 COMPOSITION OF THE GUIDELINE DEVELOPMENT GROUP

There is international agreement that guideline development groups should be multidisciplinary in their composition, with representation from all relevant professional groups, and participation of patients, carers and appropriate voluntary organisations.¹ ² This facilitates ownership of both the guideline development process and the resulting recommendations.

At the outset of a new guideline development project the SIGN Executive, in discussion with all relevant bodies, aims to bring together a group which will fulfil the following parameters:

- multidisciplinary, with all relevant clinical specialties represented alongside lay input
- relevant to current care practice, with a balance between members actively involved in day-to-day delivery of health care with topic experts and academics where appropriate. Ideally membership should represent the range of care or treatment settings related to the clinical condition (eg primary, secondary and tertiary care centres)
- encompasses the range of skills and expertise required for the specific project. Specialists other than clinicians may be recruited when necessary, for example health economists (see section 6.2.6), social workers
- geographically representative, including participants from across Scotland both from urban centres and rural locations.

Meeting this aim requires an iterative process of seeking nominations, issuing invitations and refining membership depending on the interests and availability of individuals whose participation is sought.

In putting together a guideline development group, SIGN is aware of the many psychosocial factors, including the problems of overcoming professional hierarchies that can affect small group processes. Grimshaw states: “To ensure that guidelines achieve their full potential… requires a programme of research and development that accords at least as much thought to the psychology of group dynamics as the science of systematic reviews”.³ Research into the progress and functioning of SIGN’s own guideline development groups has shown the impact of professional or status differences on members’ contributions to group discussions.⁴ ⁵ A clear relationship between the perceived status of a group member and their level of contribution to group discussions was identified.

SIGN guideline development groups vary in size depending on the scope of the topic under consideration, but generally comprise between 15 and 25 members. There is necessarily a trade-off between the number of organisations or specialties that should be represented on the guideline development group, and achieving a manageable group size for effective decision making. An example of the mix of skills present in a typical guideline group is shown in Figure 2-1. Although their areas of expertise will vary, members of the guideline development group have equal status on the group.
2.2 RESPONSIBILITIES OF GUIDELINE DEVELOPMENT GROUP MEMBERS

2.2.1 GUIDELINE DEVELOPMENT GROUP CHAIR

The role of the guideline development group Chair is crucial to ensure that the group functions effectively and achieves its aims. Chairs of guideline development groups must be sensitive to pre-existing interprofessional tensions and hierarchies and ensure that all members of the group feel able to contribute fully to the guideline development process.

A guideline development group Chair needs to be aware of, and constantly attentive to, small group processes (e.g., how the group interacts and communicates, decision-making processes and chairing strategies). The Chair must be prepared to overcome potentially serious difficulties by careful negotiation.

2.2.2 SIGN TEAM

The SIGN Programme Manager assigned to each guideline helps the Chair to identify potential barriers to successful group work, to plan and progress the guideline development project, and acts as facilitator at group meetings. Some SIGN guideline development groups are co-chaired by the SIGN Programme Manager and the group Chair in order to help reduce potential conflicts.

The SIGN team supporting each guideline development must ensure that clinical knowledge and expertise is appropriately applied to the interpretation of the evidence base and that all group members have the opportunity to actively contribute when the drafting of guideline recommendations is being undertaken.

2.2.3 GUIDELINE DEVELOPMENT GROUP MEMBERS

Guideline development group members in turn must make a full commitment to the group and the tasks involved in guideline development, and are responsible for indicating areas of concern to the Chair.
development group members should also bear in mind that they represent both a geographical region and a specialty or professional group, and must be prepared to consult with colleagues to ensure that the widest possible range of views are considered, whilst maintaining confidentiality around the content of discussions undertaken within the group.

The approximate life span of each guideline development group varies depending on whether it is a new project (around 29 months), an update (around 15 months) or a minor revision (3–6 months). For a full guideline project, groups meet on average once every two to three months, although groups may form subgroups which meet more frequently.

2.3 REFERENCES
3 Selection of guideline topics

3.1 PROPOSING A TOPIC

Producing evidence based clinical practice guidelines is a time and resource intensive process. To make best use of these resources, guidelines should address a specific healthcare need and there should be an expectation that change is possible and desirable and that, if the guidelines are followed, there is potential to improve the quality of care and/or patient outcomes.\(^1,2\) There must also be robust evidence of effective practice on which to base guideline recommendations.

Any group or individual may propose a guideline topic to SIGN. Application is a two-stage procedure and the procedure for application and selection of new topics is illustrated in Figure 3-1.

The application form to request consideration by SIGN of a specific guideline topic and the full guideline proposal form are available from the SIGN Executive or can be downloaded from the \textit{SIGN website}.

3.1.1 INITIAL APPLICATION

The initial application is made using a short, single page application form. When a group or individual proposes a guideline topic to SIGN, their suggestion is discussed initially by the SIGN Senior Management Team (SMT). SMT uses the following criteria to assess the topic:

\begin{itemize}
  \item Is this an appropriate clinical topic for a SIGN guideline? (considering whether the topic is clinical, its breadth and the need for the guideline as identified in the proposal)
  \item Is there a suitable alternative product which would address this topic? (considering whether other Healthcare Improvement Scotland products could better address the topic)
  \item Has this topic been considered before and rejected? (reasons for rejection would be reviewed and assessed for current applicability).
\end{itemize}

If the proposed topic has the potential to meet the selection criteria the proposer is asked to complete a second, more detailed, application form.

3.1.2 FULL PROPOSAL

SIGN's standard guideline application form requests the following information:

\begin{itemize}
  \item Details of the group(s) or institution(s) supporting the proposal.
  \item A brief background to the clinical topic which will be addressed by the proposed guideline.
  \item Evidence of variation in practice in the management of the condition.
  \item An indication of the benefits likely to arise from the development and successful implementation of the guideline.
  \item Key areas of concern for patients, carers and/or the organisations that represent them.
  \item A definition of the patient group to which the guideline will apply. This should include consideration of whether any specific social groups or minorities are likely to be particularly affected, either favourably or adversely, by changes in healthcare provision in the topic area under consideration.
  \item A definition of the aspects of management of the clinical condition which the proposed guideline will address and an indication as to whether the guideline will apply to primary or secondary care, or both.
  \item An indication of the healthcare professionals potentially involved in developing the guideline.
  \item An indication of the size and strength of the evidence base which is available to support recommendations on effective practice, citing key supporting papers.
  \item Details of any existing guidelines or systematic reviews in the field.
\end{itemize}
As part of the preparatory work done before the full guideline proposal is considered by the Guideline Programme Advisory Group (see section 3.2), a scoping search is carried out. This is a very broad search of the literature relevant to the condition that is to be the topic of the guideline. No attempt is made to focus on specific questions at this stage. The intention is only to establish the general extent of the literature in the clinical area to see if there is likely to be sufficient good quality evidence to make an evidence based guideline feasible.
Firstly, a check is made to see if any other good quality guidelines have been produced on the subject by searching the following websites:

Guidelines International Network (www.g-i-n.net)
National Guideline Clearinghouse (www.guideline.gov)
National Institute for Health and Care Excellence guidance (www.nice.org.uk)
National Institute for Health and Care Excellence Evidence Search (www.evidence.nhs.uk)

In addition, a search for existing systematic reviews is carried out in the Cochrane Library. This covers the reviews produced by the Cochrane Collaboration and the Database of Abstracts of Reviews of Effectiveness (DARE).

From this scoping search a report is prepared summarising the available evidence, emphasising the outcomes from systematic reviews and whether these have been positive or have identified significant work that remains to be done.

### 3.2 TOPIC SELECTION PROCESS

SIGN has limited resources for guideline development. As a result it is important to identify and prioritise topics which make best use of those resources. Likewise, when a published guideline is due for review (see section 3.3) it must be judged against potential new topics for inclusion in the SIGN programme.

For information on the current SIGN programme, see the SIGN website.

The Guideline Programme Advisory Group (GPAG) is a subgroup of SIGN Council and oversees development of proposals for new guidelines or for reviewing existing guidelines. GPAG also considers the work programmes of other parts of Healthcare Improvement Scotland, for example, development of standards and health technology assessments (HTAs) as well as other guideline developers, in particular NICE (the National Institute for Health and Care Excellence) in England and Wales, to avoid potential duplication of effort.

GPAG considers all new proposals and prioritises them using a suitability screening and scoring tool. The suitability screening tool identifies the extent to which the proposal fulfils the criteria listed in section 3.2.1, makes an assessment of the extent of evidence on which to base the guideline and considers whether the benefits that were likely to accrue from successful implementation of the guideline recommendations would outweigh the efforts required to develop it.

Using this information and taking into account SIGN's work capacity, GPAG makes recommendations to SIGN Council about which proposals should be accepted onto the work programme and which should be rejected. Topics ranked highest are included in SIGN's proposed programme, depending on capacity. Proposals which are not ranked sufficiently highly to be accepted on to the programme are reconsidered at the next topic prioritisation meeting alongside new and review topics. If the proposal still receives a low ranking on its second reading it is returned to the proposer for reconsideration or revision.

SIGN Council dedicates one meeting each year to approving guideline topic proposals that have been recommended by GPAG as suitable candidates for the SIGN guideline development programme. Council is presented with the guideline proposals, a summary of the suitability screening results and the subsequent discussions of the Guideline Programme Advisory Group before making a decision on which topics should be included in the SIGN programme.
3.2.1 CRITERIA FOR SELECTION OF TOPICS

There is a lack of evidence to guide choice of criteria and methods for prioritising topics, although the criteria used by guideline development organisations are broadly similar.2 Guideline topics selected for inclusion in the SIGN programme are chosen on the basis of the burden of disease, the existence of variation in practice, and the potential to improve outcome. The following criteria are considered by SIGN in selecting and prioritising topics for guideline development:

- Areas of clinical uncertainty as evidenced by wide variation in practice or outcomes.
- Conditions where effective treatment is proven and where mortality or morbidity can be reduced.
- Iatrogenic diseases or interventions carrying significant risks.
- Clinical priority areas for NHSScotland.
- The perceived need for the guideline, as indicated by a network of relevant stakeholders.

3.3 UPDATING PUBLISHED GUIDELINES

As medical practice continues to develop and new options for treatment become available, guidelines inevitably fall behind current best practice. They must therefore be kept under constant review and updated when necessary.2-5

3.3.1 SCHEDULED UPDATES

SIGN considers whether or not published guidelines need to be reviewed after a period of three years and all SIGN guidelines carry a statement indicating that they will be considered for review three years after publication. A full review of a guideline after a fixed time period is not always appropriate as new evidence is published at different rates in different fields. It also imposes a workload for future years that may not be achievable in practice. A further factor that will influence the decision on whether and how to review a guideline is the emergence of any evidence of inequality in access to services between different social groups that can be addressed through guideline recommendations.

3.3.2 SELECTING AND PRIORITISING GUIDELINE FOR UPDATING

As a first step, an update search is carried out looking for evidence based guidelines, HTAs, and systematic reviews produced since publication of the last version of a guideline. These searches are based on the key questions and search strategies used in the original guideline but also include an element of horizon scanning to see if there are new treatments or technologies that should be considered as part of the update. Results are presented in the form of summaries of the findings of the studies that have been identified.

The search results are incorporated into a report that summarises the new evidence and looks at how it will impact on the recommendations made in the existing guideline. This report will also note any new areas or key questions that have emerged since the previous publication.

The review report is then widely circulated for comment within NHSScotland, to Royal Colleges and other professional bodies (through their representatives on SIGN Council), to relevant patient organisations, and to other organisations providing guidance or advice to the NHS in any part of the UK. Responses to this consultation are gathered and presented to the Guideline Programme Advisory Group. GPAG considers the reports using a suitability screening and scoring tool based on the criteria described in section 3.2.1. On the basis of these reports combined with input from their professional networks GPAG then makes recommendations to SIGN Council on which guidelines should be updated.

When a guideline is considered for updating, there are four possible outcomes:

- the guideline, as it stands, will be revalidated
- the guideline will undergo a complete review
- the guideline will undergo a partial or selective review
- the guideline will be withdrawn.
A fifth option, which is likely to be applicable in only a small number of cases, is to make the guideline into a ‘living guideline’. This option involves keeping the evidence under constant review and updating the guideline on a regular basis.

At their Autumn/Winter meeting, SIGN Council will agree which guidelines are to be updated and prioritise the updates along with new guideline proposals for addition to the SIGN guideline programme. This process for selecting guidelines for updating is summarised in Figure 3-2.

Information on the status of guidelines due for updating, or currently being updated, is provided on the SIGN website.

*Figure 3-2: Selection of guidelines for updating*

![Figure 3-2: Selection of guidelines for updating](image_url)
3.3.3 SELECTIVE UPDATE PROCEDURE

It is not always desirable or necessary to completely update a guideline. Updates can apply either to sections of guidelines, or in some circumstances to individual recommendations. Processes have to be in place to address all these possible options.\(^6\)

When a guideline has been accepted for a selective update, the process for carrying out the update will be largely the same as that described elsewhere in this manual. The principal difference is that the update will focus on those sections of the original guideline that have been identified as being in need of updating. The same methodological principles apply, though the nature of the sections being reviewed may necessitate a slightly different composition from the original guideline group. If a section on surgical interventions is a major part of an update, for example, the guideline group is likely to include more surgeons and theatre staff than (say) pharmacists or homecare workers.

Another possible exception is the need for a national meeting. Here the guideline group may decide whether or not the proposed changes are sufficiently far reaching as to justify such wide consultation. If a national meeting is not held, the first draft of the guideline is published on the SIGN website for a fixed period, during which time potentially interested parties will be alerted to its presence and invited to submit comments.

3.3.4 LIVING GUIDELINES

As with a selective update, the process for updating a living guideline is largely the same as that described elsewhere in this manual. The main difference is that a living guideline is developed on a rolling programme of regular updates. The frequency of updating will depend on the rate at which new evidence is emerging, but will normally be annual or biennial. Guideline development group membership will be relatively constant, but only subgroups of the guideline development group with an interest in the topics under review will be actively involved in the development process at any time. A steering group consisting of the Chairs of the subgroups and other relevant individuals oversees guideline development and ensures consistency of approach across the subgroups year on year.

Each update focuses on those areas of the original guideline where new evidence has been identified. The same methodological principles apply and literature searches are based on a series of existing key questions. They seek to update and build on the evidence base used in the original guideline and subsequent updates. The only new questions that may be addressed are any arising from the patient issues search, or that arose from new developments identified during the process of scoping the update.

Once searches are completed, considered judgements are reviewed against the updated evidence base. The text and recommendations of the guideline are revised to take account of any new evidence and flagged as being revised. The other processes used will be the same as those used for a new guideline. A possible exception is, as with a selective update, the need for a national meeting.

SIGN currently develops only one living guideline, the British guideline on the management of asthma in collaboration with the British Thoracic Society (BTS). Updated drafts of this guideline are presented at one of the BTS biannual meetings, as well as being published on the SIGN and BTS websites for a fixed period, during which time comments are invited.

3.3.5 MONITORING AND INTERIM UPDATES

All comments received on published SIGN guidelines, or information on important new evidence in the field, or evidence of impacts on equality groups is considered, either for immediate response or for more detailed consideration on review of the guideline.

Individuals commenting on published guidelines are invited to complete a small change proposal form, which can be downloaded from the SIGN website. Once received small change proposals are processed in the same way as full proposals (see section 3.2).
GPAG considers small change requests on a rolling basis and guidelines will be ‘refreshed’ if a proposal meets the following criteria:

- new evidence substantially changes a small number of recommendations in the guideline (corresponding to no more than two related key questions) OR
- a specific issue such as a new drug therapy or national issue such as a new government policy will give rise to a new key question AND
- the nature of the update may not warrant assembling a multidisciplinary group.

To allow SIGN to be reactive to the needs of healthcare professionals in NHSScotland, small changes to published guidelines are agreed by GPAG rather than SIGN Council and slotted into the programme according to current capacity and workload.

The process for refreshing a guideline is essentially the same as a selective update although the scope of the update is much narrower and the timescales shorter. The level of involvement of a guideline development group and extent of consultation will depend on the nature of the changes to the guideline.

Any updates to the guideline which might be required in the interim period prior to review are noted on the SIGN website.

3.4 WITHDRAWING GUIDELINES

From time to time it is necessary to consider withdrawing guidelines which are outdated or no longer relevant. Proposals to withdraw guidelines are submitted initially to the Guideline Programme Advisory Group and if it agrees with the proposal it is submitted to SIGN Council for final approval.

Once it has been agreed to withdraw a guideline, all versions of the text and any associated material will be removed from the SIGN website. The list of published guidelines will be amended to show the guideline as withdrawn, with a note of the reason for withdrawal and reference to any alternative sources of advice.

Guidelines may be withdrawn for any of the following reasons.

- superseded by a more recent or more comprehensive guideline
- evidence that the guideline is fully complied with by NHSScotland, and has become accepted practice
- emergence of new treatments or preventive measures that render the guideline irrelevant.

3.5 REFERENCES


4 Systematic literature review

Guidelines based on a consensus of expert opinion or on unsystematic literature surveys have been criticised as not reflecting current medical knowledge and being liable to bias.\textsuperscript{1,2} SIGN guidelines are therefore produced using a considered judgement process informed by systematic reviews of evidence. Systematic review is defined as “an efficient scientific technique to identify and summarise evidence on the effectiveness of interventions and to allow the generalisability and consistency of research findings to be assessed and data inconsistencies to be explored”\textsuperscript{3}.

The SIGN approach is to produce a systematic review of the evidence for each key question (KQ) to be addressed in the guideline. Evidence tables are produced as supporting documents and the essential elements of systematic review are met in that the literature is:

- identified according to an explicit search strategy
- selected according to defined inclusion and exclusion criteria
- evaluated against consistent methodological standards.

All the stages of the review process are thoroughly documented (see below).

The benefits of the SIGN approach derive from the close involvement of the guideline development group with the synthesis of the evidence base, allowing them to apply their ‘considered judgment’ when deriving recommendations (see sections 5 and 6), and from encouraging a sense of ownership of the guideline amongst all those involved in the process.

4.1 ADDRESSING PATIENT ISSUES IN THE LITERATURE SEARCH

Incorporating the patient’s perspective from the beginning of the development process is essential if it is to influence the coverage of the final guideline. One of the measures used to achieve this is to conduct a specific search on patient issues in advance of the first meeting of the guideline development group.

This search is designed to cover both quantitative and qualitative evidence, and is not limited to specific study designs. It is carried out over the same range of databases and sources as the main literature review, but will normally include both nursing and psychological literature even where these are not seen as particularly relevant to the later searches of the medical literature. Whereas other literature searches carried out for the guideline attempt to answer focused key questions by filtering out the volume of irrelevant evidence, the patient search is deliberately as broad and inclusive as possible. It focuses entirely on the health condition that is being considered, and makes no attempt to concentrate on any social group or class. As the reviewer develops themes from the literature, (s)he will pay particular attention to anything that suggests there are population groups that are disadvantaged and ensure their interests are specifically considered by the guideline development group.

The use of this literature search is discussed in more detail in SIGN 100: A handbook for patient and carer representatives.\textsuperscript{4}

4.2 USING EXISTING GUIDELINES

As more good quality guidelines are being produced by other agencies, SIGN is making use of the evidence base underlying guidelines produced elsewhere for use in NHSScotland.

The guidelines identified in the scoping search carried out for the original guideline proposal will be presented to an early meeting of the guideline development group to allow it to consider what has been done already.

All guidelines must be evaluated using the AGREE II instrument and be shown to have followed an acceptable methodology before they can be considered for use by SIGN guideline developers.
There is a range of possible ways in which existing guidelines can be used in relation to SIGN guidelines.

- There is an existing guideline that addresses some of the questions in a new guideline, or there is a well produced guideline that is now out of date. If the guideline development group can access the original evidence tables used to develop that guideline, these can be updated and reviewed before being used to form the basis for new recommendations. For example, NICE evidence tables were used by the group developing the SIGN obesity guideline.

- There is an existing guideline that addresses some of the questions in a new guideline, but it is not possible to obtain the evidence tables. In this case the guideline can be included in the body of evidence, as low quality evidence, supporting new recommendations.

- There are existing guidelines addressing a very specialist area where expertise or resources are limited, and the specialist topic has general applicability. For example, reference in SIGN 88 on management of bacterial urinary tract infections to prescribing guidance produced by the Health Protection Agency or the Infectious Diseases Society of America.

- There is an existing guideline that addresses some aspect of the guideline topic and which can be referred to as an alternative. For example, SIGN 111 on management of hip fracture of the elderly refers to British Orthopaedic Society guidelines for secondary prevention of fragility fractures rather than tackling the topic anew.

In all cases the guideline development group must decide on the best way forward that will address clinical need while avoiding duplication and waste of resources.

### 4.3 DEFINING KEY QUESTIONS

SIGN guideline development groups are encouraged to break down the guideline remit into a series of structured key questions using the PICO format as shown below.5, 6

- **Patients or population** to which the question applies
- **Intervention** (or diagnostic test, exposure, risk factor, etc) being considered in relation to these patients
- **Comparison(s)** to be made between those receiving the intervention and another group who do not receive the intervention
- **Outcome(s)** to be used to establish the size of any effect caused by the intervention

The **patients or population** to be covered by the literature searches is largely defined by the presence of the particular condition that the guideline will cover. It should, however, be made clear which age groups are to be included.

Consideration should also be given to issues of equity, ensuring that any particular subgroup of the patient population that has particular needs in relation to the topic under review has those needs specifically addressed. This should take account not just of the needs of that population, but any evidence of differences in effectiveness of interventions between equality groups.

It is worth emphasising here that, where clinically important, questions should be addressed even if it is not thought there will be any good evidence. If there is in fact no good evidence, then highlighting it as an area for research is a useful outcome in itself. Dealing with uncertainties of this kind will be addressed in the section of this manual covering the later stages of the considered judgment process.

The **interventions** (which in this context includes diagnostic tests, risk factors, risk exposure) must be specified clearly and precisely. The only exception is in drug therapy where drug classes should be used in preference to specific agents unless there is a clear reason for focusing on a named agent.

The decision on **comparisons** is mostly between placebo/no treatment, or comparison with other therapies. It should be borne in mind that, where there is an existing treatment, comparison with placebo or no treatment is not ethically acceptable.
Outcomes must be clearly specified, ideally at the stage of setting the key questions but certainly before making judgments about the quality of evidence. For some questions there will be a wide range of outcomes used in the literature, and, if useful comparisons are to be made across studies, it must be made clear which of these outcomes are expected to influence decisions about healthcare.

Outcomes should be discussed by the guideline development group and rated in terms of their importance. Where existing reviews have been identified in the scoping search, the outcomes used in those reviews should be presented to the guideline development group as an aid to completing this part of the process.

Critical outcomes are those on which the overall quality of evidence for a KQ is based. These are the key outcomes on which a healthcare professional would be expected to base a treatment decision. In osteoporosis, for example, prevention of fracture is likely to be seen as a critical outcome. The number of critical outcomes should be kept low, preferably less than seven per KQ.

Important outcomes are those that a healthcare professional is likely to take into account when making treatment decisions, but which are not the ultimate aim of the intervention under consideration. Often, these will be surrogates for the critical outcome. In the osteoporosis example, improved bone mineral density may be seen as an important outcome as it is a widely accepted surrogate for reduced fracture risk.

In some areas there are a large number of reported outcomes. Some of these are likely to be peripheral to treatment decisions and can be largely ignored in the process of developing guideline recommendations. As far as possible outcomes should be objective and directly related to patient outcomes (eg length of time to next cardiovascular incident or survival time, rather than just reductions in blood pressure). Patient important outcomes should be explicitly considered along with more narrowly defined clinically important outcomes. It is particularly important to include any potential harm associated with the intervention under review so that a balanced view can be taken at the considered judgment stage.

As part of the question setting process, a set of inclusion and exclusion criteria should be drawn up and saved as part of the record of the review. This will provide guidance at a later stage when studies are being selected for review.

Inclusion criteria will include definition of the topic and may include such factors as duration of therapy, drug dosage, and frequency of treatment. Other factors include any geographic or language limits, the types of trials that will be accepted, and date range to be covered. Any equality groups that are expected to have specific needs in relation to the question being addressed should be specified.

Exclusion criteria are likely to be more variable. They are, however, essential in that they help sift out irrelevant studies from the (often very large) initial search result. Equality groups should never be specifically excluded from searches unless a clear justification is provided (eg issues for specific equality groups addressed in a separate question).

Once the questions have been agreed they form the basis of the literature searches to be undertaken by an Evidence and Information Scientist. These searches will focus on the patients, interventions, and (sometimes) comparison parts of the question. Additional references provided by group members or other interested parties may be included for consideration in the evidence base but must be evaluated on the same basis as all other studies (see section 4.5).

Definition of a set of clear and focused clinical questions is fundamental to the successful completion of a guideline development project. It is also important to be realistic about the number of questions that can be addressed in a single guideline if the final product is not to be too large to be useable. A large number of key questions will incur a very high workload for the developers. Care must be taken to ensure this is kept within limits that will allow the guideline to be completed within the agreed timescale. Keeping the number of questions to a minimum is particularly important in areas that are particularly rich in literature, as each question will require review of a large number of studies.

Deciding the key questions is entirely the responsibility of the guideline development group which must apply its knowledge and experience to ensuring the questions address the key issues in the area to be covered by the guideline. The Evidence and Information Scientist working with the group will provide guidance on
question formatting, and ensure they are likely to produce useable results. They will also work with the Patient Involvement Officer to ensure that the key questions address appropriately the issues identified through the patient consultation exercise.  

4.4 IDENTIFYING AND SELECTING THE EVIDENCE

The literature search must focus on the best available evidence to address each key question. SIGN uses a set of standard search filters that identify:

- systematic reviews
- randomised controlled trials
- observational studies
- diagnostic studies
- economic studies.

In order to minimise bias and to ensure adequate coverage of the relevant literature, the literature search must cover a range of sources. As a minimum, SIGN requires searches to cover the following sources:

- Cochrane Library
  - for reviews: Cochrane reviews, Database of Abstracts of Reviews of Effects (DARE)
  - for RCTs: Cochrane Controlled Trials register (CCTR)
- Medline
- NHS Economic Evaluations Database (NEED)
- Internet sites relevant to the topic (including patient organisations)
- WHO International Clinical Trials Registry Platform.

For questions covering drug treatments, searches will also cover:

- Scottish Medicines Consortium (SMC) advice
- NICE Multiple Technology Assessments
- Toxbase
- US Food and Drug Administration (FDA) Register.

Specialised databases such as CINAHL or Psychinfo will only be searched for questions specific to their area of coverage.

Following a review in 2012 SIGN decided not to include Embase in the standard set of databases covered. Evidence suggests that the benefit of searching both Embase and Medline is variable between topics, but can be very low. Inclusion of CCTR in the main search picks up those unique Embase records added by Cochrane and reduces duplication of effort.

SIGN does not undertake hand searching of key journals as part of the literature review. It is accepted that this means some relevant trials may be missed, and introduces the possibility of a degree of bias in the process. However, given time and resource constraints, it is not feasible for this to form part of the process.

The period that the search should cover will depend on the nature of the clinical topic under consideration, and will be discussed with the guideline group. For a rapidly developing field a five year limit to the search may be appropriate, whereas in other areas a much longer time frame might be necessary.

All the main search strategies are available to members of guideline development groups if they want to review them.

A listing of the Medline search strategies used for the guideline, plus notes of any significant variation on other databases, is published on the SIGN website at the time of publication of the guideline.

Before any studies are acquired for evaluation, the search output is sifted to eliminate irrelevant material. Results are sifted in two stages. A preliminary sift of each search result is carried out by the Evidence and Information Scientist, normally by the individual that carried out the search. Studies that are clearly not
relevant to the key questions or not the type of study being considered (e.g., observational studies when the focus is on controlled trials) are eliminated. Abstracts of remaining studies are then examined and any that clearly do not meet the agreed inclusion and exclusion criteria will also be eliminated at this stage. In cases of doubt, the Evidence and Information Scientist will leave abstracts in the output file at this stage.

A final sift is carried out by at least one member of the guideline development group, who will apply clinical judgment to reject any other studies that do not meet the pre-agreed criteria. These will include clinical criteria, but may also consider issues such as size of the study, relevance to practice in the UK, etc. Only when all stages of search result sifting have been completed will the remaining studies be acquired for evaluation.

4.5 EVALUATING THE LITERATURE

Once studies have been selected as potential sources of evidence, the methodology used in each study is assessed to ensure its validity.

The methodological assessment is based on a number of criteria that focus on those aspects of the study design that research has shown to have a significant effect on the risk of bias in the results reported and conclusions drawn. These criteria differ between study types, and a range of checklists is used to bring a degree of consistency to the assessment process. The SIGN checklist for systematic reviews has been updated based on the AMSTAR tool,13,14 while that for RCTs is based on an internal project carried out in 1997.15 Checklists for observational studies are based on the MERGE (Method for Evaluating Research and Guideline Evidence) checklists developed by the New South Wales Department of Health,16 which have been subjected to wide consultation and evaluation. The checklist for diagnostic accuracy studies is based on the QUADAS (Quality Assessment of Diagnostic Accuracy Studies) programme.17

These checklists were subjected to detailed evaluation and adaptation to meet SIGN’s requirements for a balance between methodological rigour and practicality of use. Copies of these checklists and accompanying notes on their use are available on the SIGN website.

The assessment process inevitably involves a degree of subjectivity. The extent to which a study meets a particular criterion, for example an acceptable level of loss to follow-up and, more importantly, the likely impact of this on the reported results from the study will depend on the clinical context and inevitably the judgment of the individual reviewers.

The methodology of studies selected for full consideration will be appraised by at least two people with experience in carrying out such appraisals. The subjective nature of critical appraisal makes double checking essential to minimise the chance of bias and to ensure consistency. Where reviewers cannot agree on the overall quality of a study the Programme Manager will arbitrate before a study goes forward for inclusion in the evidence base. This only applies to studies being actively considered as evidence. There is no need to seek agreement for studies that are not to be included. Any study that has not been included in this process cannot be used as evidence to support a recommendation in the guideline.

4.6 REFERENCES


5 Assessing the quality of evidence

5.1 INTRODUCTION

The previous section of this handbook sets out how individual studies are identified and assessed for methodological rigour. The next step in the guideline development process is to examine the body of evidence associated with each specific key question.

One of the factors likely to influence a practitioner’s decision to implement a recommendation is the degree of confidence that they have in it; that is how certain they are that following the recommendation will produce the expected improvement in outcome for their patients. Not only does this certainty relate to the degree of confidence in the size of effect of an intervention in relation to specific important outcomes, but it also encompasses other issues such as patient preferences and the availability of resources to support introduction of a new intervention. For this reason the guideline development group has to consider both the overall quality of the supporting evidence and the other factors that might influence the strength of the recommendation.

If a reviewer evaluates a guideline using the AGREE instrument they will expect both of these aspects to be clearly addressed. These aspects also need to be addressed properly to comply with the standards set out by the Institute of Medicine. As from June 2014, this is also a requirement for all guidelines included in the National Guidelines Clearinghouse database.

5.2 PRESENTING THE EVIDENCE

This section of the manual considers the practical issues involved in presenting the evidence to the guideline development group. It relates to the SIGN solution, although there are a number of ways of doing this.

5.2.1 EXISTING SYSTEMATIC REVIEW

It is a fundamental principle that each recommendation should be based on systematic review of the literature. For many questions systematic reviews will already exist, and in these cases the guideline development groups are provided with a complete systematic review plus an evidence table summarising more recent studies. Where there are multiple existing reviews, an evidence table summarising the findings of all existing reviews, is provided.

In these circumstances the quality of the studies included in the systematic review has already been established by the systematic reviewers, and, the guideline development group can move on to consider its conclusions (see section 6).

Consideration of the evidence in relation to different outcomes is considerably simplified if a summary of findings (SoF) table is available. Any SoF produced as part of a systematic review should be included in the material submitted to the guideline group. If they are not included in a systematic review, the authors may be contacted to see if SoF tables are available.

5.2.2 INTERNALLY CONDUCTED REVIEWS

A completed evidence table based on an internally conducted systematic review of the literature will be provided for all questions. These will either update existing reviews or provide a review of all relevant literature. Each evidence table will include methodological evaluation of and data from each individual study relevant to a specific key question. Study results will be reported on a per outcome basis wherever possible.

An example of a completed evidence table records appears in Figure 5-1.
5.3 CONSIDERING THE QUALITY OF EVIDENCE

SIGN is committed to following the principles of the GRADE methodology which complies with the standards covered in section 5.1. The process for assessing the overall quality of evidence using GRADE, is described in the Journal of Clinical Epidemiology (JCE) series on GRADE. This chapter will be updated periodically in the light of experience of applying these principles in SIGN guidelines.

From this point in the process the guideline development group is looking at a body of evidence for each question; the collection of studies that help answer the question. This raises a number of issues beyond the methodological quality of the individual studies.

The evaluation of a body of evidence should be completed before deciding what to recommend in the guideline. The focus here is on the quality of the available evidence, not what conclusions may be drawn from it.

The evidence identified in a systematic review of the published literature is first summarised in an evidence table (see Figure 5-1). The features described in the following sections are then reviewed and commented on in part A of the considered judgment form (see Figure 5-2).

In summary, at the end of this stage of the process, the guideline development group will have agreed on the overall quality of the evidence for all critical outcomes for the key questions being addressed.

5.3.1 HOW RELIABLE ARE THE STUDIES IN THE BODY OF EVIDENCE?

The first issue to be considered is the risk of bias in the studies that make up the body of evidence related to a particular question. The methods used for the assessment of risk of bias in individual studies are outlined in section 4 of this manual.

5.3.2 ARE THE STUDIES CONSISTENT IN THEIR FINDINGS?

Also known as heterogeneity, this aspect looks at all the studies relating to a particular outcome to see if they all point in the same direction (ie all support or reject the course of action being considered). Sometimes it is very clear that evidence is consistent, but at other times this is far from the case. Sometimes there are clinical reasons to explain this and these will be discussed by the guideline development group. The issue that is of concern here, however, is statistical heterogeneity.

Statistical heterogeneity can only be established through meta-analysis. Often these analyses will provide statistics to indicate the degree of heterogeneity. The most commonly quoted is the \( \chi^2 \) (chi squared) statistic. If this is below 40, inconsistency is not likely to be a problem. Over 90 and it is a very serious issue for consideration. In between these values, you need to consider possible explanations for the variation in results and how they might influence the size of effect in the Scottish population.

In the context of SIGN guideline development, such calculations will normally only be available through published meta-analyses.

5.3.3 ARE THE STUDIES RELEVANT TO OUR TARGET POPULATION?

This is often referred to as directness of evidence, but can also be referred to as applicability or external validity. In this context it relates to how directly applicable the evidence is to NHSScotland. Guidelines should indicate where the studies used as evidence were conducted, if not by listing all the countries involved, at least indicating which parts of the world the evidence came from. For example:

“The main work on this topic has been carried out in Europe and the UK.”

“Most of the evidence in this area comes from the US-based Framingham study.”
Studies carried out in the UK are likely to be directly applicable to the target population for a SIGN guideline. For studies carried out elsewhere some thought has to be given to what factors, if any, might influence relevance of the results in our target population. Studies of diagnosis of heart disease, for example, may be seen as directly relevant no matter where they are conducted as the underlying biology of the condition should be the same. If the studies relate to the risk of heart disease however, applicability may be more problematic.

Examples of factors that can influence the applicability of evidence include:

- variations in baseline risk
- differences in genetic makeup of the population
- differences in culture or lifestyle between populations
- differences in how care is delivered, or availability of technologies or resources
- different outcomes measured in studies to those that the guideline development group see as being of critical importance
- differences in how the intervention(s) studied is/are administered to patients in Scotland
- use of indirect (surrogate) outcomes
- indirect rather than direct comparison of outcomes.

It is worth highlighting that the last two points above relate to different forms of indirectness. Surrogate outcomes reflect a situation where it is difficult or impossible to accurately measure the effect of an intervention on the final patient important outcome. In that case an alternative outcome that can be shown to be related to the important outcome may be measured instead. An example of this is in osteoporosis where studies often report the impact of interventions on bone mineral density, when in fact the outcome of interest is the degree of fracture risk. Increased bone density is associated with a reduced risk of fracture, hence its use as a measure of treatment effect.

The second issue arises where there are no head-to-head comparisons of different options for treatment. It may be, for example, that there is no comparison of A versus B, but there are trials of A versus C and B versus C. In this situation evidence may be addressed through a process referred to variously as indirect comparison reviews, mixed treatment comparisons, or network meta-analyses. These analyses are always indirect, and given their nature are unlikely to give a precise result.

Situations where there is a mix of direct and indirect evidence are the most difficult to judge. The guideline development group has to look at the body of evidence and see where most of the data comes from. If it is clearly from direct evidence there should not be a problem.

In some cases it may be that the group has to consider different sub-populations within the overall body of evidence. People of Asian origin living in the UK, for example, are known to have an increased risk of Type 2 diabetes at an earlier age than the white population. Any evidence in this area would need to take account of the risk difference when considering whether or not it was direct. It may be preferable to ask the question independently for different groups where situations like this are suspected to see if there is direct evidence for any specific group of patients.

### 5.3.4 HOW SURE ARE WE THAT ESTIMATES OF THE SIZE OF EFFECT ARE RELIABLE?

This is often referred to as precision of the estimate of effect. It relates to how confident the user can be in any estimate of the size of the effect to be expected from an intervention or exposure. Precision around an effect estimate is usually presented as 95% confidence intervals.

Trial results are commonly reported in terms of relative effect or relative risk. Wherever possible, estimates of absolute risk or benefit should also be used along with the appropriate confidence intervals.
5.3.5 ARE WE SURE WE HAVE ALL THE RELEVANT EVIDENCE?

This question relates to publication bias, where only some study results (usually the positive ones) have been reported. Unfortunately it is not usually possible to establish the presence or absence of publication bias, and reviewers can only indicate if it is likely or unlikely.

Published reviews should include an assessment by the authors of the likelihood of publication bias.

For internal reviews carried out by SIGN staff, it may be assumed that the literature searches coupled with the knowledge of the guideline development group members have covered the majority of the available literature. Some papers may have been missed, but there will not be any systematic bias in the search results. SIGN searches do not cover unpublished material, and it is a matter of judgment for the guideline development group to decide if there is likely to be a substantial body of unpublished literature that might influence the results.

**Note:** Data and comments in the following figures have been adapted from documents used in an actual guideline. They have, however, been amended for illustrative purposes and should not be taken as accurate and complete summaries of the evidence in this area.
**Guideline topic:** Breast cancer

**Question:** KQ03: In patients with BC (men, pre-menopausal women, postmenopausal women) with an inoperable tumour/unsuitable for breast-conserving surgery, what is the evidence that neoadjuvant chemotherapy is effective and what is the optimal regimen?

**Cuppone, F, Bria, E, Carlini, P, Milella, M, Felici, A, Sperduti, I, et al.**
Taxanes as primary chemotherapy for early breast cancer: Meta-analysis of randomized trials (*Cancer: 2008 113 238-46*)

<table>
<thead>
<tr>
<th>Study type/ evidence level</th>
<th>Study details/limitations</th>
<th>Patient characteristics</th>
<th>Interventions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meta-analysis/ RCT</td>
<td><strong>Countries:</strong></td>
<td>Total no. patients: 2,455</td>
<td>Combination taxane and anthracycline</td>
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<tr>
<td>Evidence level: +</td>
<td><strong>Centres:</strong></td>
<td><strong>Patient characteristics:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Setting:</strong></td>
<td>‘Operable’ breast cancer stage II and III - otherwise not well defined</td>
<td></td>
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<tr>
<td></td>
<td><strong>Funding sources:</strong></td>
<td><strong>Inclusion criteria:</strong></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Not stated</td>
<td>Searched till October 2007</td>
<td></td>
</tr>
<tr>
<td></td>
<td><strong>Dropout rates:</strong></td>
<td><strong>Exclusion criteria:</strong></td>
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<td></td>
<td><strong>Study limitations:</strong></td>
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**Notes:** The primary endpoint is pCR. The included trials use different definitions of pCR (unlisted). A similar criticism exists for other endpoints such as response rate although breast conservation rate should be robust.

Thorough search of the literature. Poor reporting of populations characteristics of the included studies. No quality appraisal carried out. Sensitivity analysis carried out for sequential taxane schedule showing higher probability of achieving a pCR. Author’s comment on the heterogeneity of the studies, including populations, sample size and treatment duration.

**Author’s conclusions:** The combination of taxanes and anthracyclines as neoadjuvant chemotherapy for early breast cancer improved the chance of achieving higher breast-conserving surgery rates and pathologic complete response rates. This main conclusion followed from the evidence presented, but did not clearly acknowledge limitations of the evidence discussed by the review authors, so it may not be entirely reliable.
| Outcome Measures/results | Data for primary endpoints were available for 7 RCTs. The rate of BCS was significantly higher for patients receiving taxanes, with an AD of 3.4% (P=5.012), which translates into 29 patients NNT, without significant heterogeneity. The rate of pCR was higher for patients receiving taxanes (not statistically significant). In the sensitivity analysis, patients receiving taxanes as a sequential schedule had a significantly higher probability to achieve pCR, with an AD of 2.4% (P=5.013), which translates into 41 patients NNT, without significant heterogeneity. Patients receiving taxanes as a concomitant schedule had a significantly higher probability to achieve BCS, with an AD of 5.3% (P=5.027), which translates into 19 patients NNT, without significant heterogeneity. The complete response rate was significantly higher in the taxane arms, regardless of the adopted strategy, with an AD ranging from 6.7% to 15.5%.


<table>
<thead>
<tr>
<th>Study type/ evidence level</th>
<th>Study details/limitations</th>
<th>Patient characteristics</th>
<th>Interventions</th>
</tr>
</thead>
</table>
| Randomised controlled trial | **Countries**: Spain, Italy, Russia, Poland, Estonia, Hungary, Austria, Czech Rep  
**Centres**:  
**Funding sources**: Healthcare Industry, Bristol Meyers Squibb  
**Dropout rates**:  
**Study limitations**: | **Total no. patients**: 1,355  
**Patient characteristics**: 20% had tumours > 4cm, more than half were aged over 50.  
**Inclusion criteria**: Women with primary operable breast tumour >2cm (T2-T3, N0-N1, M0)  
**Exclusion criteria**: | Primary systemic therapy v CMF/ doxorubicin followed by CMF or CMF/ doxorubicin/ paclitaxel followed by CMF |

| Notes: | In multivariate analysis only estrogen receptor (ER) status was significantly associated with pathologic complete response (odds ratio for ER negative, 5.77; 95% confidence interval, 3.49-9.52; P<0.0001). Neoadjuvant chemotherapy induced a significant axillary downstaging (P <0.001), and breast-sparing surgery was feasible in 65% versus 34% (P<0.001). Not a blinded trial. No ITT.  
**Author’s conclusions**: Doxorubicin/paclitaxel followed by CMF is feasible, safe, and well tolerated. Given as PST, it is markedly active, allowing for breast-sparing surgery in a large fraction of patients. |
| Outcome measures/results | Toxicity, adverse events | Grade 3 or 4 National Cancer Institute toxicities were low (<5%) in all arms. Neuropathy was more frequent in the paclitaxel-containing arms (grade 2, 20.5% versus 5.0%; grade 3, 1.3% versus 0.2%). At 31 months of follow-up, asymptomatic drop of left ventricular ejection fraction was similar in all arms, whereas symptomatic cardiotoxicity was recorded in three patients (0.5%) in A and in three patients (0.3%) in B plus C. PST induced clinical complete plus partial remission in 78%, with an in-breast pathologic complete response rate of 23% and an in-breast plus axilla pathologic complete response rate of 20%. In the multivariate analysis, only estrogen receptor (ER) status was significantly associated with pathologic complete response (odds ratio for ER negative, 5.77; 95% confidence interval, 3.49-9.52; P < 0.0001). PTS induced a significant axillary downstaging (P<0.001), and breast sparing surgery was feasible in 65% versus 34% (P< 0.001). |

**Notes:**
- Exploring association of clinico-pathological variables with response - all patients received the same chemotherapy for the first part of the trial. Independent factors for mid-course response and pCR were: young age, non-T4 tumors, high grade, and hormone receptor status, the strongest single predictive factor. Grading and age can identify subgroups within the luminal and triple negative patients who have an increased benefit from NACT.
- These are useful data to answer question of benefit in different populations (ER+ v ER –; HER2+ v HER2–; PR+ v PR–)
- Evaluated as a cohort; evaluation of pre-randomised cohort from Gepartrio trial. No control group. Large number lost to follow-up.
- **Author’s Conclusion:** Grading and age can identify subgroups within the luminal and triple negative patients who have an increased benefit from NACT.

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Effect of neoadjuvant anthracycline-taxane-based chemotherapy in different biological breast cancer phenotypes: Overall results from the gepartrio study (Breast Cancer Research & Treatment: 2010 124 133-40)

<table>
<thead>
<tr>
<th>Study type/evidence level</th>
<th>Study details/limitations</th>
<th>Patient characteristics</th>
<th>Interventions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cohort study</td>
<td><strong>Countries:</strong></td>
<td><strong>Total no. patients:</strong></td>
<td>TAC</td>
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<td></td>
<td><strong>Centres:</strong></td>
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<td></td>
<td><strong>Setting:</strong></td>
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<td></td>
<td><strong>Funding sources:</strong></td>
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<td><strong>Dropout rates:</strong></td>
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- **Author’s Conclusion:** Grading and age can identify subgroups within the luminal and triple negative patients who have an increased benefit from NACT.
| Outcome measures/results | pathological clinical response | The overall pCR rate, defined as no invasive residuals in breast and axilla, was 20.5%. The highest pCR rate of 57% was observed in patients below 40 years of age with triple negative or grade 3 tumors. Independent factors for mid-course response and pCR were: young age, non-T4 tumors, high grade, and hormone receptor status, the strongest single predictive factor. Within the biological subtypes, grading was an independent factor to predict pCR for luminal tumors, clinical tumor stage for the HER2 like tumors and age for the triple negative ones. Grading gave independent information for mid-course response within the triple negative group. No factor predicted mid-course response within the other groups. |
### CONSIDERED JUDGEMENT

**Key question: KQ03**

In patients with BC (men, pre-menopausal women, post-menopausal women) with an inoperable tumour/unsuitable for breast-conserving surgery, what is the evidence that neoadjuvant chemotherapy is effective and what is the optimal regimen?

**A: QUALITY OF EVIDENCE**

1. **How reliable are the studies in the body of evidence?** (see SIGN 50, section 5.3.1, 5.3.4)
   
   If there is insufficient evidence to answer the key question go to section 8

   Comment here on any issues concerning the quantity of evidence available on this topic and its methodological quality. Please include citations and evidence levels.

<table>
<thead>
<tr>
<th>Evidence levels</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1**</td>
</tr>
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</tr>
<tr>
<td>1*</td>
</tr>
<tr>
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2. **Are the studies consistent in their conclusions?** (see SIGN 50, section 5.3.2)

   Comment here on the degree of consistency demonstrated by the evidence. Where there are conflicting results, indicate how the group formed a judgement as to the overall direction of the evidence.

   Overall there does not appear to be any significant inconsistency between the trials.

3. **Are the studies relevant to our target population?** (see SIGN 50, section 5.3.3)

   For example, do the studies:
   - include similar target populations, interventions, comparators or outcomes to the key question under consideration?
   - report on any comorbidities relevant to the target population?
   - use indirect (surrogate) outcomes
   - use indirect rather than direct comparison of outcomes
The trial results and interventions are applicable to the Scottish population.

4. Are there concerns about publication bias? (see SIGN 50, section 5.3.5)
Comment here on concerns about all studies coming from the same research group, funded by industry etc.

Van de Hage comments that publication bias may be possible. No publication bias reported in the Moja review. Not commented on in Cuppone review.

**B: EVIDENCE TO RECOMMENDATIONS**

5. Balancing benefits and harms (see SIGN 50, section 6.2.2, 6.2.3)
Comment here on the potential clinical impact of the intervention/action – eg magnitude of effect; balance of risk and benefit.

What benefit will the proposed intervention/action have?
Describe the benefits. Highlight specific outcomes if appropriate.

### Neoadjuvant chemotherapy vs surgery/adjuvant chemotherapy

There is sufficient evidence to indicate that neoadjuvant chemotherapy is associated with higher rates of breast conservation than adjuvant chemotherapy, with no difference in overall survival or locoregional recurrence rates, providing surgery is part of the treatment pathway. In total, 5,500 patients treated in 14 studies, with median follow-up of 18-124 months were included. The meta-analysis concluded that OS was equivalent for preoperative chemotherapy versus adjuvant chemotherapy (HR 0.98 (95% CI 0.87-1.09, p=0.67). There were increased breast conservation rates with neoadjuvant chemotherapy but there were associated increased locoregional recurrence rates. However, if surgery was included, even in cases of complete response, locoregional recurrence rates were equivalent (HR 1.12 (95% CI 0.92-1.37, p=0.25). Patients who achieve pathological complete response show improved survival, compared with patients with residual disease (HR 0.48 (95% CI 0.33-0.69, p<0.104).

**Duration of chemotherapy**

A randomised trial of 4 vs 6 cycles epirubicin and docetaxel (Han 2009) of 176 patients showed pCR 11% vs 24% (p=0.047) with a trend to higher clinical response and higher breast conservation. A randomised trial of 3 vs 6 cycles epirubicin and docetaxel (Reitsamer 2005) of 45 patients showed pCR 10% vs 36% (p=0.045) with a trend to increase breast conservation. A randomised trial of 3 vs 6 cycles epirubicin and docetaxel (Steger 2007) of 292 patients showed pCR 7.7% vs 18.6% (p=0.045) with a trend to increased breast conservation.

**Taxanes vs non-taxanes**

There is sufficient evidence to indicate that breast conservation rates and rates of pathological complete response (pCR) are higher in patients treated with a combination of anthracycline and taxane-based neoadjuvant chemotherapy, compared with non-taxane based chemotherapy. Cuppone review – 7 RCTs in the meta-analysis, including concomitant or sequential administration, either docetaxel or paclitaxel used as the taxane (ie the treatment regimes are quite heterogeneous). 2,455 patients in total. BCS significantly higher with a taxane (absolute difference 3.4%, p=0.12). There was a trend to higher pCR for patients treated with taxanes, which reached statistical significance in patients receiving sequential anthracyclines/taxanes (AD 2.4%, p=0.013). The Mazouni pooled analysis of 7 consecutive neoadjuvant chemotherapy trials conducted at MD Anderson (1,079 patients in total) indicated higher pCR in patients receiving a taxane (29% vs 15% in ER negative patients, p<0.001 and 8.8% vs 2.0% in ER positive patients, p<0.001).

**Herceptin in Her2 positive**

There is sufficient evidence to indicate that patients with Her-2 positive (Her2 3+ or Her2 2+ FISH amplified) breast cancer, receiving neoadjuvant chemotherapy, should receive Herceptin, either as adjuvant treatment or with non-anthracycline-based neoadjuvant chemotherapy. In patients with Her2 positive disease, adjuvant or neoadjuvant Herceptin leads to improved DFS and OS. The Cochrane review (Moja 2012) of 8 studies (neoadjuvant/adjuvant studies combined) which included 11,991 patients. This showed improved OS (HR 0.66, 95% CI 0.57-0.77, p=0.00001) and improved DFS (HR 0.060, 95% CI 0.50-0.71, p=0.00001) with no heterogeneity of effect between adjuvant and neoadjuvant administration of Herceptin. A meta-analysis (Valachich) of 5 trials with 515 patients showed higher pCR rates (RR 1.85, 95% CI 1.39-2.46, p<0.001). Data on rate of BCS was available in 4 trials (280 patients). For these patients, no difference was seen in rate of BCS (OR 0.98, 95% CI 0.80-1.19, p=0.82). The NOAH trial (Semiglavzov 2011) has reported a higher rate of BCS in patients with locally advanced breast cancer receiving neoadjuvant Herceptin in addition to chemotherapy (23% vs 13%).
Subtypes
In the ECTO trial (Gianni 2005) ER negativity was significantly associated with pCR (odds ratio for ER negative 5.77, 95% CI 3.49-9.52, p<0.0001). In the GEAPRIO study (Huober 2010), association of clinico-pathological variables with response was explored (not randomised). The strongest single predictive factor for pCR was ER and PR negativity (odds ratio 3.08, 95% CI 2.32-4.09, p<0.001).

What harm might the proposed intervention/action do?
Describe the benefits. Highlight specific outcomes if appropriate.

Neoadjuvant chemotherapy vs surgery/adjuvant chemotherapy
van de Hage found no significant difference between adjuvant and neoadjuvant chemotherapy for post-operative complications, nausea/vomiting or alopecia. Events of leucopenia and infections (RR 0.69, 95% CI 0.56-0.84, p=0.0003) were significantly lower with neoadjuvant chemotherapy.

Duration of chemotherapy
Han showed no similar rates of toxicity (grade 3-4 neutropenia, febrile neutropenia and mucositis) for 3 vs 6 cycles. Reitsamer did not report on toxicity. Steger reported no differences in haematological, GI, neurological, cardiac or other serious toxicities between the 2 groups.

Taxanes vs non-taxanes
The Cuppone, Mazouni and Nowak reviews did not report toxicity. Trudeau reported increased haematological toxicity, including neutropenia and febrile neutropenia, with taxane-containing regimes. There was no evidence of significant difference in other adverse events, other than peripheral neurotoxicity.

Herceptin in Her2 positive
A combined analysis of neoadjuvant and adjuvant trials (Moja) reported a significantly increased risk of congestive heart failure (RR 5.11, 90% CI 3.00-8.72, p<0.00001) and left ventricular ejection fraction decline (RR 1.83, 90% CI 1.36-2.47, p=0.0008) when Herceptin is added to chemotherapy. There was no difference in haematological toxicities.

Impact on patients (see SIGN 50, section 6.2.4, 6.2.5)
Is the intervention/action acceptable to patients and carers compared to comparison? Consider benefits vs harms, quality of life, other patient preferences (refer to patient issues search if appropriate).

Are there any common comorbidities that could have an impact on the efficacy of the intervention?

The use of adjuvant chemotherapy in patients with breast cancer at moderate/high risk of disease recurrence reduces the risk of relapse and improves survival. The use of adjuvant trastuzumab in patients with Her-2 positive breast cancer reduces the risk of relapse and improves survival. Healthcare professionals and patients place high value on these outcomes.

Neoadjuvant chemotherapy +/- trastuzumab is associated with higher rates of breast conservation than adjuvant chemotherapy, with no difference in overall survival or locoregional recurrence rates, providing surgery is part of the treatment pathway. In addition, there is no evidence of any significant difference between toxicity of neoadjuvant or adjuvant chemotherapy.

Patients and healthcare professionals place high value on recurrence/survival outcomes and, in suitable patients, increased breast conservation rates.

Feasibility (see SIGN 50, section 6.2.6)
Is the intervention/action implementable in the Scottish context? Consider existing SMC advice, cost effectiveness, financial, human and other resource implications.

Neoadjuvant chemotherapy +/- trastuzumab is generally well tolerated and the regimes used are similar to adjuvant chemotherapy. Patients with inoperable breast cancer are, by definition, incurable and therefore treatment that renders such a patient operable and potentially curable is cost effective. Breast conservation is less morbid than mastectomy and, arguably, offers improved quality of life and neoadjuvant chemotherapy with this aim is, therefore, cost effective.

In line with SMC/NICE recommendations

Recommendation (see SIGN 50, section 6.3)
What recommendation(s) does the guideline development group agree are appropriate based on this evidence?
**Strong** recommendations should be made where there is confidence that, for the vast majority of people, the intervention/action will do more good than harm (or more harm than good). The recommendation should be clearly directive and include 'should/should not' in the wording.

Conditional recommendations, should be made where the intervention/action will do more good than harm, for most patients, but may include caveats eg on the quality or size of the evidence base, or patient preferences. Conditional recommendations should include 'should be considered' in the wording.

- Neoadjuvant chemotherapy should be offered for all patients with breast cancer whose disease is either inoperable but localized to the breast/locoregional lymph node groups, or high T/N stage, or where the only surgical option is mastectomy and downstaging might offer the patient the opportunity for breast conservation.
- Anthracycline-taxane based chemotherapy combinations increase the rate of breast conservation rates and pCR rates, compared with anthracycline-alone regimes and should be offered for all patients receiving neoadjuvant chemotherapy.
- Trastuzumab should be offered to patients with Her-2 positive breast cancer, either as adjuvant treatment or with non-anthracycline based neoadjuvant chemotherapy.
- Anthracyclines and trastuzumab should be avoided or used with extreme caution in patients with cardiac comorbidity. In less fit patients, the benefits of chemotherapy outweigh the likelihood of doing harm, and treatment should only be recommended after careful consideration.

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**Anthracyclines and trastuzumab should be avoided or used with extreme caution in patients with cardiac comorbidity. In less fit patients, the benefits of chemotherapy outweigh the likelihood of doing harm, and treatment should only be recommended after careful consideration.**

**Briefly justify the strength of the recommendation**

In patients with breast cancer, whose disease is either inoperable, higher stage or there is the possibility of increasing breast conservation rates, the likelihood of doing good outweighs the likelihood of doing harm.

9. **Recommendations for research**

List any aspects of the question that have not been answered and should therefore be highlighted as an area in need of further research.

Studies into the efficacy of therapies for men with breast cancer are required.
CONSIDERED JUDGEMENT

Key question: KQ03

In patients with BC (men, pre-menopausal women, post-menopausal women) with an inoperable tumour/unsuitable for breast-conserving surgery, what is the evidence that neoadjuvant chemotherapy is effective and what is the optimal regimen?

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1. How reliable are the studies in the body of evidence? (see SIGN 50, section 5.3.1, 5.3.4)

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2. Are the studies consistent in their conclusions? (see SIGN 50, section 5.3.2)

Comment here on the degree of consistency demonstrated by the evidence. Where there are conflicting results, indicate how the group formed a judgement as to the overall direction of the evidence.

Overall there does not appear to be any significant inconsistency between the trials.

3. Are the studies relevant to our target population? (see SIGN 50, section 5.3.3)

For example, do the studies:

- include similar target populations, interventions, comparators or outcomes to the key question under consideration?
- report on any comorbidities relevant to the target population?
- use indirect (surrogate) outcomes
5.5 REFERENCES


6 Making recommendations

6.1 INTRODUCTION

The work covered in earlier sections of this manual covers the identification and evaluation of evidence relating to specific key questions. By this point of the process the guideline development group knows how much evidence there is (or is not) available to help answer their questions. Though there is clearly room for debate about some issues, a framework of basic rules for identifying and appraising evidence has been in place. The process now moves on to an area where there is more scope for opinion to guide the final conclusion.

A newcomer to the field of evidence based guidelines may wonder what place opinion has in applying evidence. For an explanation, we need to go back to a basic definition of evidence based medicine (EBM). In the introduction to their landmark book on EBM,1 David Sackett and his co-authors defined it as:

“…the integration of best research evidence with clinical expertise and patient values”.

So far, we have dealt with gathering the research evidence. What now needs to happen is for clinical expertise and patient values, among other things, to be applied to that evidence to arrive at a recommendation that is in line with the evidence, is practical to deliver, and takes account of patient preferences. In other words a recommendation that is likely to be implemented and to be acceptable to patients.

This section is based on the Evidence to Decision (EtD) tool developed as part of the DECIDE project,2 which is in turn based on the work of the GRADE group.3,4

6.1.1 STRONG VERSUS WEAK

Before going further, it is worth focusing on the outcome of a GRADE decision-making process. It is to produce a recommendation that is rated as either strong or weak (which, in the SIGN implementation of GRADE, we will refer to as ‘conditional’ recommendations).

A strong recommendation is made where:

- the evidence is of high quality (see section 5)
- estimates of the effect of an intervention are precise (ie there is a high degree of certainty that effects will be achieved in practice)
- there are few downsides of therapy
- there is a high degree of acceptance among patients.

A conditional recommendation is made where:

- there are weaknesses in the evidence base
- there is a degree of doubt about the size of the effect that can be expected in practice
- there is a need to balance the upsides and downsides of therapy
- there are likely to be varying degrees of acceptance among patients.

6.2 EVIDENCE TO RECOMMENDATION

The following sections are based on DECIDE EtD frameworks. At the time of writing these were at an advanced stage of development, but may be further refined. This chapter will be updated periodically in the light of further developments in the DECIDE work and experience of applying these principles in SIGN guidelines.

It is worth keeping in mind at all times that fundamental to this approach to guideline development is the issue of transparency. Different guideline developers will allocate greater or fewer resources to developing their guidelines, and the detail of the work they do will vary accordingly. The important point in all that follows is to be clear about what was actually done at each stage of the process. Justifications can be provided if thought necessary, but the key point is to produce a structured summary of the complete process that reviewers or guideline users can check when they are considering implementation of the guideline.
6.2.1 IS THIS QUESTION A PRIORITY?

Given that a question has survived through the processes of topic selection and key question setting, it might be taken as read that it is a priority. The intention here, however, is to indicate why the question is being addressed.

- What risks will be reduced?
- To what extent is there a need to improve on current treatments?
- How many patients are likely to be affected?
- Could improvement in this condition reduce the risk/impact of common comorbid conditions?

These are some of the types of issue to be addressed here.

Members of the guideline development group have a key role to play as they will be aware of the main issues that make a question important, as well as some of the key information that will illustrate that importance. Their knowledge may be supplemented by evidence from official data, published sources, or research studies.

6.2.2 HOW SURE ARE WE THAT ANY GIVEN OPTION WILL WORK?

At this point the guideline development group relies on the summarised evidence produced at the previous stage in the process (see section 5). The factors described in the following sections are then considered in part B of the considered judgment form (see Figure 5-2) to allow recommendations to be formed from the evidence. Ideally this table can be taken from a summary of findings (SoF), but this is unlikely to be available in every case. For those key questions where an SoF is not available, an alternative short format presenting non-pooled results (for example, an evidence table) will suffice. The guideline development group should focus on (for each outcome):

- outcome
- impact
- number of studies
- quality/certainty of the body of evidence.

6.2.3 BALANCING BENEFITS AND HARMS

Fundamental to making any recommendation is the need to ensure that any benefit to the patient outweighs, preferably by a substantial margin, any risks or harms associated with the treatment.

In order to make such judgments, the guideline development group has to have a clear understanding of how substantial the expected benefits of an intervention are likely to be in practice. They also need to consider how substantial the downsides are. These may range from physical side effects to an increased risk of developing additional health problems.

The evidence supporting benefits will often come from stronger study designs than that supporting harms. This makes judgments more difficult, but it is nonetheless essential to explicitly consider the size of effect for both sides of the balance. A detailed presentation of the evidence from a summary of findings or similar table (see section 5.3) is essential when making such decisions.

Once the size of all effects has been established, a judgment must be made as to whether the benefits outweigh the harms. This is not just a clinical judgment but must take into account patient values (see section 6.2.4) if a realistic assessment is to be achieved.

6.2.4 HOW DO PATIENTS VALUE THE DIFFERENT OUTCOMES?

For a recommendation to be implemented effectively, it is important that the outcomes are sufficiently valued by patients for them to be willing to adhere to the treatment. The science of assessing patient values and preferences, however, remains largely undeveloped. When developing guideline recommendations, the focus should be on questions where the application of values is likely to affect outcomes and should rely on practical and achievable methods.
In the case of venous leg ulcers, for example, there is strong evidence that using compression stockings is an effective treatment, and the higher the compression the better the results. Compression stockings have various drawbacks, however, and some patients either cannot or will not tolerate the highest levels of compression. It then becomes a question of balancing these preferences against the risk of larger or longer lasting ulcers. A recommendation based entirely on trial evidence without taking into account patient preferences is unlikely to be widely adhered to, and therefore ineffective.

Assessing patient values and preferences can focus on the extent to which they are likely to follow a recommended course of action, though there is some evidence that wider social values can play a part in such decisions.\(^7\) A first step should be to consult patient representatives on the guideline development group, and through them a wider body of patient opinion.

If time and resources allow, a literature search can be carried out looking specifically for information on patient values in relation to the question being addressed. It is worth noting that there is an increasing literature on patient values related to specific conditions.

If acceptability of a recommendation to patients is seen as critical to its effective implementation, and no clear idea of patient views has been identified by the above methods, it may be necessary to run a series of focus groups to establish patient values and preferences.

### 6.2.5 EQUITY

Under the **Equality Act 2010** all public bodies in Scotland are required to take into account the needs of equality groups. This applies to all guidelines and other publications produced by SIGN. The equality groups identified in the Act are:

- age
- disability
- gender reassignment
- marriage and civil partnership
- race
- religion or belief
- sex
- sexual orientation.

Guideline groups are therefore required by law, as well as good practice, to consider whether any recommendations they make will have a differential impact on any of these groups.

Some aspects of equality issues have been addressed earlier in this manual (**see sections 4.3 and 5.3.3**). At this later stage in the process, it may be necessary to analyse the evidence for specific subgroups of the population to see if and how it differs from the main results. If there are substantial differences it will be necessary to make separate recommendations for these subgroups taking these differences into account.

Apart from issues of social equity, subgroups may need to be considered for clinical reasons such as specific comorbidities, or issues around polypharmacy where separate recommendations may be required for these groups.

### 6.2.6 COSTS AND BENEFITS

There are two aspects to the consideration of costs and benefits in relation to guideline recommendations.\(^8,9\)

The first relates to cost effectiveness of a single proposed intervention, and involves assessing the incremental cost of applying the new intervention compared to current practice and relating it to the net benefit of the intervention. In many cases this will not be a major issue, but where it is an assessment should be made to ensure introduction of the new approach is worthwhile. This may be achieved through a review of the existing
literature, or through economic modelling. For these issues to be addressed adequately a health economist should be available to advise the group on which questions are likely to require an assessment, and if possible to conduct that assessment and report the results to the group prior to their making a recommendation.

The second issue relates to the resources required to implement a recommendation across the NHS in Scotland. Again, this may not be an issue in a lot of cases but where very expensive treatments or interventions requiring substantial investment in equipment or changes to working practices are involved an assessment of the cost impact is important if the guideline is to be implemented.

In this second case the cost assessment may not influence specific recommendations directly, but should be produced along with the guideline to inform decision makers who need to allocate resources within individual health boards. If the potential cost is very high and may not be achievable in the short term, a ‘next best’ option may be recommended in the guideline. The guideline should, however, always identify the most cost-effective option, with the ‘next best’ as an interim option only.

### 6.3 MAKING RECOMMENDATIONS

Balancing all the issues described above is a matter of considerable complexity, and presents a challenge to any guideline group. High quality evidence from well conducted studies should lead to a strong recommendation, but relating the trial populations to the target population of a guideline and taking into account issues of cost and patient acceptability may lead to a recommendation that is much weaker than first thought. Equally, there will be circumstances where the evidence is flawed but there are few or no downsides to treatment and the clinical importance of the topic is such that a strong recommendation is justifiable.

It is not possible for SIGN or any other guideline organisation to advise or direct a guideline group as to the conclusions they should reach. All that can be asked is that the group considers all the issues and uses a transparent process to reach their conclusion.

Particularly where considerations of equity or comorbidity are involved, the guideline development group may have to make more than one recommendation; one for each subgroup discussed.

In all situations, however, the overall judgment of the guideline development group can only lead to one of the five possible conclusions shown in Table 6-1, each related to a particular form of recommendation.

**Table 6-1: Forms of recommendation**

<table>
<thead>
<tr>
<th>Judgment</th>
<th>Recommendation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Undesirable consequences clearly outweigh desirable consequences</td>
<td>Strong recommendation against</td>
</tr>
<tr>
<td>Undesirable consequences probably outweigh desirable consequences</td>
<td>Conditional recommendation against</td>
</tr>
<tr>
<td>Balance between desirable and undesirable consequences is closely balanced or uncertain.</td>
<td>Recommendation for research and possibly conditional recommendation for use restricted to trials</td>
</tr>
<tr>
<td>Desirable consequences probably outweigh undesirable consequences</td>
<td>Conditional recommendation for</td>
</tr>
<tr>
<td>Desirable consequences clearly outweigh undesirable consequences</td>
<td>Strong recommendation for</td>
</tr>
</tbody>
</table>
Whatever the conclusion, the published guideline and supporting documentation should contain a justification for the recommendation highlighting the supporting evidence and the factors that have been taken into account when arriving at a conclusion.

Where decisions are particularly complex, such a justification may be quite lengthy. In these cases the full justification can be included in supporting material with a shortened version included in the published guideline.

Recommendations are differentiated from other text in the published guideline by presenting as a single paragraph in bold text. A capital ‘R’ is used alongside to emphasise that the associated text is a recommendation.

\[ R \] Patients with larger tumours may be considered for oncoplastic surgery instead of mastectomy.

6.4 GOOD PRACTICE POINTS AND CONSENSUS RECOMMENDATIONS

Good Practice Points (GPP) are intended to assist guideline users by providing short pieces of advice which may not have an evidence base, but which are seen as essential to good clinical practice.

Examples of acceptable GPPs:

- Healthcare professionals should refer to the WHO medical eligibility criteria for contraceptive use prior to offering contraceptive advice to women with diabetes.
- Healthcare professionals should signpost patients to self help resources, identified and recommended by local pain services, at any point throughout the patient journey.

If the group feels strongly that they want to make a recommendation even though there is no significant evidence, this should be done as a weak recommendation based on very low quality evidence. Note that there must be some evidence of opinion supporting the recommendation from outside the guideline group. If no such evidence exists, formal methods should be used to develop a consensus based recommendation which will be clearly identified as such within the guideline by a statement accompanying the recommendation.

The methods used to reach consensus may vary between guideline groups. Whatever method is used, it is essential that it is described either in an Annex to the guideline or as a supporting document linked to the guideline on the SIGN website.

6.5 KEY RECOMMENDATIONS

The guideline group will identify a small number of key recommendations to be listed in a separate section of the guideline (see section 8). These key recommendations are identified by the guideline development group as the recommendations that, in order to improve patient outcomes, should be prioritised for implementation. They appear in the main text as well as section 2 of the guideline, and will appear in the quick reference guide and associated app.

A consensus based recommendation may be included as a key recommendation.
6.6 REFERENCES


7 Consultation and peer review

7.1 NATIONAL OPEN MEETING

As an early stimulus to testing the feasibility of implementation, SIGN holds a national open meeting to discuss the draft recommendations of each guideline. This takes place whilst the guideline is still in development and gives the guideline development group the opportunity to present its preliminary conclusions and draft recommendations to a wider audience. The benefits of the national open meeting are twofold:

1. the guideline development group obtains valuable feedback and suggestions for additional evidence which group members might consider, or alternative interpretation of that evidence
2. the participants are able to contribute to and influence the form of the final guideline, generating a sense of ownership over the guideline across geographical and disciplinary boundaries.

SIGN national open meetings are widely publicised and are usually attended by between 100 and 300 healthcare professionals and others interested in the guideline topic, including patient representatives, from across Scotland. Advertising of the meetings is targeted on those professional and patient representative groups most likely to have an interest in the topic. Particular efforts are made to ensure that all equality groups with a potential interest in the topic are represented.

The draft guideline is also available on the SIGN website for a month at this stage to allow those unable to attend the meeting to submit comments on the guideline. Social media is also used as a forum for discussion around the consultation meeting and content of the draft guideline.

The national open meeting is the main consultative phase of SIGN guideline development. Corporate interests, whether commercial, professional, or societal, have an opportunity to make representations at the national meeting or open consultation stage where they can send representatives to the meeting or provide comment on the consultation draft.

For selective updates, national open meetings are only held if the content of the guideline has significantly changed. Otherwise, the guideline is made available for open consultation on the SIGN website for one month.

No consultation meeting is held for guidelines that are only being refreshed. The revised section of the guideline is sent directly to appropriate expert reviewers (see section 7.2).

7.2 PEER REVIEW

All SIGN guidelines are reviewed in draft form by independent referees who are selected for their expertise and to reflect the multidisciplinary nature of the guideline. The draft is also sent to at least two lay reviewers in order to obtain comments from the patient’s perspective. Reviewers are asked to comment primarily on the comprehensiveness and accuracy of interpretation of the evidence base supporting the recommendations in the guideline.

It should be noted that all reviewers are invited to comment as individuals, not as representatives of any particular organisation or group. Comments from peer reviewers will not be considered unless an accompanying declaration of interests form has also been submitted.

The comments received from peer reviewers and others are compiled in a report and discussed by the guideline development group. Each point must be addressed and any changes to the guideline as a result noted or, if no change is made, the reasons for this recorded.

The names and designations of all reviewers are published in the guideline and their comments, along with the guideline development group response, and declaration of interest forms are available from SIGN on request.

As a final quality control check prior to publication, the guideline and the summary of peer reviewers’ comments are reviewed by the SIGN Editorial Group, consisting of the SIGN Chair, Director, Programme
Lead and representatives of SIGN Council. The Editorial Group ensures that each point raised by the peer reviewers has been addressed adequately and that any risk of bias in the guideline development process as a whole has been minimised. Each member of the guideline development group is then asked formally to approve the final guideline for publication.

The full editorial and consultation phase is illustrated in Figure 7-1. This process of extended consultation greatly enhances the validity of the final SIGN guideline and increases the likelihood that the guideline will be implemented successfully into local practice for the benefit of patients.

*Figure 7-1: Consultation and peer review phases of guideline development*

<table>
<thead>
<tr>
<th>CONSULTATION AND PEER REVIEW</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SYSTEMATIC LITERATURE REVIEW AND DRAFT RECOMMENDATIONS</strong></td>
</tr>
<tr>
<td>(see sections 5 and 6)</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>Draft guideline</strong></td>
</tr>
<tr>
<td>• presented and discussed at national open meeting (optional for selective update)</td>
</tr>
<tr>
<td>• available for comment on SIGN website for one month</td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>Feedback discussed by guideline development group and draft guideline revised</strong></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>Revised draft sent to expert peer reviewers</strong></td>
</tr>
<tr>
<td><strong>Draft circulated for information to various health service organisations</strong></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>Feedback discussed by guideline development group and draft guideline revised</strong></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>SIGN Editorial Group reviews guideline and peer review comments</strong></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td><strong>PUBLICATION</strong></td>
</tr>
</tbody>
</table>
8 Presentation and publication

8.1 PRESENTATION OF THE GUIDELINE

There is little information available on the effect that style and presentation have on the adoption and utility of guidelines. Clarity of definitions, language, and format is likely to be important. Guidelines should, therefore, be written in unambiguous language and should define all terms precisely. The most appropriate format for presenting guidelines will vary depending on the target group(s), the subject matter, and the intended use of the guideline. Ideally, end users should be consulted on methods of presentation. This is an additional function of the extensive peer review process to which all SIGN guidelines are subject (see section 7).

Having a well developed and defined template for presentation of the final guideline can greatly facilitate the development process, enabling guideline development groups to plan at the outset what type of information will be required and also to envisage what format the content will take. By following the model for systematic review and formation of guideline recommendations outlined in sections 4, 5 and 6, guideline development groups will find that most of the required information will then be produced in a structured, accessible format, ready to slot into the guideline template.

8.2 CONTENT OF THE GUIDELINE

8.2.1 INTRODUCTION

Each SIGN guideline has an introduction, outlining the need for the guideline, including evidence of variation in practice and the potential for the guideline to improve patient care. The remit of the guideline is carefully defined, detailing definitions, the patient population and target users of the guideline. The key clinical questions covered in the guideline are detailed in an Annex with clear reference to the section to which the question refers and where methodological limitations of the evidence base are discussed. A statement of intent makes clear the purpose of the guideline.

8.2.2 EVIDENCE AND RECOMMENDATIONS

Within the main body of the guideline, the structure should as far as possible reflect the development process that the guideline development group has followed, (ie for each section):

• A clear statement of the issue under consideration
• An explanation of the treatment options available
• A summary of the conclusions drawn from the critical appraisal of the evidence (the evidence statement, annotated with the quality of evidence and key references). This should provide the justification for the recommendation to follow; that is, the evidence for improved patient outcome resulting from the recommended action or for harms or contraindications relating to treatment options (see section 5)
• The recommendations that the group has derived from this evidence (see section 6)
• A brief discussion of any practical points (eg resource/geographical considerations to be taken up in the discussion of local guidelines for implementation), or treatment options for which there is no evidence (the last should be stated clearly)
• Finally, if the group feels it is important to give guidance in any of these latter areas, a ‘good practice point’ may be presented alongside the recommendations.

8.2.3 KEY RECOMMENDATIONS

The guideline development group highlights a small number of recommendations, which may include good practice points, as the key recommendations that should be prioritised for implementation. It is important to note that the key recommendations will not necessarily be those with the strongest supporting evidence, but those considered by the guideline development group as having the greatest potential impact on patient care (see section 6).
8.2.4 INFORMATION FOR PATIENTS

All SIGN guidelines include a ‘Provision of information’ section, which gives examples of the information patients and carers may find helpful at the key stages of the patient journey. The information in this section is provided for use by health professionals when interacting with patients and carers and for guiding the production of locally produced information materials. The issues highlighted in this section are informed by:

- patient views gathered earlier in the development process (see section 4)
- discussion with patient representatives on the development group
- input from other guideline development group members.

In cases where there are strong and diverse views among patients, focus groups may be used to identify the most widely needed information that patients require.

This section also includes details of appropriate help lines, support groups and reading materials.

8.2.5 IMPLEMENTATION RESOURCES

During the development of the guideline, the development group identifies or develops tools and activities that will aid implementation of the guideline (see section 9). Resource implications of implementing the key recommendations and key points for audit are also developed as part of an implementation strategy for the guideline (see section 9).

8.2.6 GUIDELINE DEVELOPMENT

Brief details of the systematic review on which the guideline recommendations are based are also provided, with full details of the main search strategy available on the SIGN website. Stakeholder involvement is demonstrated through listing the guideline development group members, specialist peer reviewers and others commenting at the consultations stage of guideline development, and the SIGN Editorial Group.

8.2.7 RECOMMENDATIONS FOR RESEARCH

SIGN guidelines themselves may act as a stimulus to research. An important subsidiary outcome of the guideline development process is in highlighting gaps in the evidence base and guidelines contain a section listing the guideline development group’s recommendations for research.

As each guideline is published, SIGN uses it as a basis for adding records to the Database of Uncertainties about the Effects of Treatments (UKDUETS). Records are added for research recommendations and, in some cases, for treatments the guideline group has recommended but where better evidence is still required.

The review of a guideline is an opportunity to discover whether any of the gaps in the evidence base have been filled.

8.3 PUBLISHING THE GUIDELINE

All SIGN guidelines are available free of charge on the SIGN website in PDF or HTML format. Updates including any corrections are made to the web version of the guideline, which is the definitive version at all times.

The search strategy and register of interests declared by the guideline development group are published alongside the guideline. Other supporting material may include:

- implementation resources, eg patient pathways, costing tools
- patient resources, eg sample leaflets
- learning resources, eg slide sets, online tutorials.

8.3.1 QUICK REFERENCE GUIDE

Each SIGN guideline is published with an accompanying Quick Reference Guide (QRG). This provides a summary of the key recommendations and other information from the guideline, often following a loosely algorithmic format illustrating the recommended care pathway.
8.3.2 GUIDELINE APP

The SIGN guideline app for iPhone, iPad, iPod Touch and Android phones and tablets contains QRGs of recently published SIGN guidelines. The QRG content is enhanced with material from the main guideline and online resources, linked to the SIGN website. Each new SIGN QRG is available as an update through the Apple App Store or Google Play as it is published.

The app features keyword search, bookmarking, and a function allowing the user to attach a PDF of the guideline they are interested in to an email message.

8.3.3 PATIENT VERSION

SIGN patient versions of guidelines are lay translations of the clinical guidelines. They are intended to:

- help patients and carers understand what the latest evidence supports around diagnosis, treatment and self care
- empower patients to participate fully in decisions around management of their condition in discussion with healthcare professionals
- highlight for patients where there are areas of uncertainty.

The decision to develop a patient version will depend on the clinical topic, the availability of existing good quality patient information, for example from the voluntary sector, and the capacity of the guideline development group members to assist in the interpretation of the evidence based recommendations for a lay audience.

As part of SIGN’s commitment to the equality agenda of NHSScotland, patient versions of guidelines can be produced in languages other than English upon receipt of requests from users. Languages covered include those community languages identified by the Scottish Government, Gaelic, or British Sign Language (BSL). Large print versions can also be made available.

A small selection of patient versions has been published in alternative electronic formats such as Apple and Android apps and ebooks.
9 Implementation

9.1 GETTING GUIDELINES INTO PRACTICE

To achieve the objectives set out in section 1.2 it is important not only to develop valid guidelines by a sound methodology, but also to ensure the implementation of the evidence based recommendations. As one of a range of tools to help healthcare professionals and organisations to improve clinical effectiveness and patient outcomes, guidelines provide an opportunity for practitioners to improve shared clinical decision making, increase team working, expand their evidence based knowledge, and reduce variation in practice. They can also enable professionals to keep up to date and to assess their own clinical performance against the recommendations for best practice.

However, there is often a gap between the development of guidelines, as set out in the previous sections of this handbook, and their implementation into practice. Just as guidelines help provide a bridge between research and practice, this section outlines the strategies that can assist practitioners, and health services to bridge the gap between guideline development and implementation.

9.2 DISSEMINATION

Guidelines must be made as widely available as possible in order to facilitate implementation. SIGN Quick Reference Guides and patient booklets are distributed free of charge throughout NHSScotland. Our focus is on electronic distribution and all SIGN guidelines can also be downloaded free of charge from the SIGN website. The Quick Reference Guides are available on the SIGN guideline Smartphone app.

Dissemination of SIGN guidelines in NHSScotland is organised within each NHS board by local distribution coordinators, who are responsible for disseminating guidelines across their board. The distribution coordinators are notified of all new guidelines and updates to published guidelines and given an opportunity to order Quick Reference Guides to distribute within their board. It is important to maintain a strong relationship with the distribution coordinators and keeping contact details up to date is key to a robust dissemination process. Notification of new guidelines is also sent to the Royal Colleges in Scotland, the chairs of NHS boards, the chief executives of NHS boards, the chief scientist’s office, other guideline development organisations, postgraduate college deans and voluntary organisations listed in the guideline.

Copies of all SIGN publications are deposited with the British Library and the Agency for Legal Deposit.

9.3 IDENTIFYING BARRIERS TO IMPLEMENTATION

There are two types of barriers to the implementation of guidelines: those internal to the guideline itself, and the external barriers relating to the clinical environment and particular local circumstances. SIGN addresses the internal barriers by developing guidelines according to a robust methodology, described in detail in the earlier sections. Potential external barriers to guideline implementation include:

- Structural factors (eg budget constraints, significant service redesign required)
- Organisational factors (eg inappropriate skill mix, lack of facilities or equipment)
- Peer group (eg local standards of care not in line with desired practice)
- Individual factors (eg knowledge attitudes, skills)
- Patient perceptions and treatment preferences
- Professional-patient interaction (eg problems due to language or social origin, mental health issues)
- Disadvantaged patient populations (eg poverty, homelessness).

Disadvantaged populations are known to have poorer health and health care and external barriers to implementation contribute to inequalities in health care. For successful implementation, and to achieve the aim of reducing variation in practice, external barriers also need to be assessed and implementation strategies developed to address them.1
9.4 IMPLEMENTATION SUPPORT STRATEGIES

Implementation of guidelines is a local responsibility and many local initiatives have already been successful in overcoming these barriers to implementation. Most clinical governance support teams in NHS boards now have audit and clinical effectiveness facilitators with some resources to help local implementation. This is an opportunity to encourage team working and cooperation within primary and secondary care and at the interface between them.

Initiatives both nationally and locally have taken into account evidence on the effectiveness of different strategies to implementation: “evidence based medicine requires evidence based implementation”. Implementing guidelines is not simple or straightforward. Difficulties often centre on the need for personal, organisational or cultural change. However, such change is being carried through in many areas of clinical practice and information to support a local evidence based strategy is available from a variety of sources.

The Cochrane Effective Practice and Organisation of Care (EPOC) group has published a summary of 44 systematic reviews of implementation interventions, giving an indication of the most effective approaches as summarised in Figure 9-1. The authors emphasised that there are “no magic bullets”. Each implementation strategy is effective under certain circumstances, and a multifaceted approach is most likely to achieve change. The approach should be tailored to suit local circumstances taking into account any particular potential barriers. Characteristics of the patient population and any potential health inequities also need to be considered. It is important to build in support and incentives and to consider the resources needed for successful implementation.

Figure 9-1: Effectiveness of interventions to promote implementation

<table>
<thead>
<tr>
<th>Variable effectiveness</th>
<th>Largely effective</th>
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</thead>
<tbody>
<tr>
<td>Audit and feedback</td>
<td>Reminders</td>
</tr>
<tr>
<td>Local consensus conferences</td>
<td>Educational outreach (for prescribing)</td>
</tr>
<tr>
<td>Opinion leader</td>
<td>Interactive educational workshops</td>
</tr>
<tr>
<td>Patient-mediated interventions</td>
<td>Multifaceted interventions</td>
</tr>
</tbody>
</table>

A Health Technology Assessment (HTA) review of dissemination and implementation strategies suggests that the evidence for educational outreach is equivocal and that dissemination of educational materials may have greater impact than originally considered, and that multifaceted intervention comparison is problematic. The review makes it clear that there is an imperfect evidence base to support decisions about dissemination and implementation and that any approach should always take account of local circumstances.

Stakeholder consultation showed what type of support healthcare professionals in NHSScotland would like to aid implementation of SIGN guidelines. Results of the survey fell into the following four key domains, which form the basis for our implementation support; improving SIGN processes, awareness raising and education, networking and implementation support resources.

9.4.1 IMPROVING PROCESSES

Robust dissemination

Implementation can only begin once the appropriate people are made aware of the guideline and receive and read a copy. As discussed in section 9.2, SIGN works closely with local distribution coordinators to ensure that all staff who need to know about a new guideline or an update, can access it easily.
9.4.2 AWARENESS RAISING AND EDUCATION

Awareness raising activities

An important element of implementation is making people aware of the guideline and its recommendations. SIGN staff, lay representatives, guideline development group members and SIGN Council members raise awareness by presenting at conferences, workshops and educational events.

SIGN often publishes guideline summaries in medical journals, which allows a much wider audience to be made aware of SIGN and its latest recommendations.

Local clinical champions

Having a powerful clinical champion can be an effective way to raise awareness of a guideline. Clinical champions have a high profile, are widely respected, have a good understanding of policy and have local contacts. SIGN supports clinical champions, usually guideline development group members, to take a leading role in supporting implementation of specific guidelines in their area.

Patients as champions for change

Patients are a powerful agent for change in the health service. Many guidelines are published with an accompanying patient and carer version of the guideline and by being aware of a clinical guideline, patients can ask for their care to be in line with the latest recommendations. Making use of connections with patient groups and voluntary organisations also affords more opportunities to raise awareness of guidelines.

Lay representatives on guideline development groups are supported to raise awareness at conferences and other events.

Education and training modules

SIGN has worked with NHS Education for Scotland and the Royal College of Physicians and Surgeons of Glasgow to develop training modules based on guidelines. By linking these to Continuous Professional Development, healthcare professionals are encouraged to complete the training. The modules are available on the SIGN website.

9.4.3 NETWORKING

Linking with existing networks and projects

In many areas of clinical practice, there are existing professional networks and/or national projects that aim to improve the quality of care and to put evidence into practice. Examples include Managed Clinical Networks (MCNs) for cancer, coronary heart disease (CHD) and epilepsy and the national Mental Health Collaborative. Building relationships with the various professional networks, Scottish Government, NHS Education for Scotland and others as part of a wider cohesive approach to improving patient care should facilitate implementation. For example, the stroke guideline was developed in coordination with revision of the clinical standards, the production of key performance indicators and the production of a resource calculator.

9.4.4 IMPLEMENTATION SUPPORT RESOURCES

Each guideline development group develops tools or signposts useful resources that will support implementation. An implementation resource is any tool or activity that contributes towards putting the recommendations into practice. They are generally targeted towards recommendations that will have the maximum impact on patient care and can include:

Algorithms, care pathways and integrated care pathways

Algorithms and care pathways describe the typical journey of care and provide a visual representation of a group of recommendations. They can be a useful tool for people wishing to implement a change in practice and can be used for educational purposes. Similarly, integrated care pathways, by their design, can ensure that specific recommendations are implemented in practice.
Resource implication tools

Where a key recommendation is likely to result in significant resource changes a resource implications calculator or costing tools can be developed to help NHS boards identify the potential costs and savings of implementation. Resource tools for the CHD and stroke guidelines can be found on the SIGN website.

Datasets

Datasets that support the implementation of key recommendations are often included with guidelines. Wherever possible SIGN works with other agencies to support the incorporation of recommendations in national datasets and audit tools.

Electronic decision support tools

Incorporating recommendations into local electronic decision support systems is an efficient way to assist implementation. For example, some referrals made via the Scottish Care Information (SCI) Gateway are based on SIGN recommendations for referral, which are embedded in the system.

Other tools

Other simple tools such as posters highlighting key recommendations, audit proforma, easily accessible and editable lists of the recommendations, slide sets and case studies may also be developed with each guideline and made available on the SIGN website.

9.5 PRACTICAL STEPS

The first step in this process is to prioritise the topic for the team. This may be decided by the NHS board through their Local Health Plan, or a local service or practice may identify a priority clinical area in which they wish to examine care and identify areas for improvement. It is important to recognise that clinical teams can only tackle one guideline at a time for an active implementation strategy. In fact it may be that only certain key recommendations within the guideline are prioritised for implementation. The clinical team, however, should identify the strengths and weaknesses of present provision and not merely choose those areas that are most easily implementable. It is encouraging to identify what is being done well but also important to identify where services could be improved ensuring that any changes that are planned are achievable.

Figure 9-2 outlines the likely steps that a local implementation group might take, adapted from the Royal College of Nursing Guidelines7 and the SPICEpc (Scottish Programme for Improving Clinical Effectiveness in Primary Care) project (www.ceppc.org/spice/index.shtml).

Figure 9-2: Practical steps towards guideline implementation

| Step 1 | Decide who will lead and coordinate the team and identify stakeholder representatives for the implementation group. It is often helpful to have a key facilitator for this process. The team should be multidisciplinary in composition. |
|-----------------------------------------------|
| Step 2 | Determine the current position. It is essential to be aware of current practice and to identify where changes need to be made. It is helpful to audit current clinical practice. It is also important to review the local environment considering people, systems, structures and internal and external influences. Through this process it is possible to identify potential barriers and facilitators to implementation. |
Step 3

Prepare the people and the environment for guideline implementation. It is important to ensure that the professionals are receptive with a positive attitude to the initiative and have the skills and knowledge to carry out the procedures. This requires time, enthusiasm and commitment with good communication and offers of tangible help. It is important to also involve patient groups in planning the initiative so they are involved from the outset and can influence the way that the guideline is implemented into local services. Patient preferences and views eg Scottish Health Council publications, local surveys should be taken into account. In preparing the environment it may be necessary to acquire new equipment or change forms or access services in a different way. It may be possible to consider the inclusion of reminder notes or computer-assisted reminders.

Step 4

Decide which implementation techniques to use to promote the use of the clinical guidelines in practice. This should take into account the potential barriers already identified and use the research evidence on effective strategies.

Step 5

Pulling it all together. This requires an action plan for the improvement process. It requires everyone to agree the aims with a named person responsible for the action plan and a time scale identified with contingency plans to deal with any problems along the way.

Step 6

Evaluate progress through regular audit and review with feedback to the team. Rewarding achievements is important. Plans may be required to be modified in the light of difficulties or surprises found during the implementation process. It is always important though to celebrate successes and aim for small achievable steps along the way to improve the quality of patient care.

9.6 REFERENCES


10 Involving patients and their representatives

10.1 PATIENT INVOLVEMENT IN GUIDELINE DEVELOPMENT

The term patients is used throughout this section as a generic term to describe patients, carers, lay representatives and those who represent and/or support patients in the voluntary sector.

Patient involvement is “the appropriate, active participation of patients, carers and patient representatives as partners in their own care and in the planning, monitoring and development of health services”1 The potential contribution of patient representatives has been recognised for some time, as well as the difficulties in making that contribution effective.2

Patients may have different perspectives on healthcare processes, priorities, and outcomes from those of health professionals. The involvement of patients in guideline development is therefore important to ensure that guidelines reflect their needs and concerns. The purpose of patient involvement is to ensure that the guideline addresses issues that matter to them and that their perspectives are reflected in the guideline. Patients can identify issues that may be overlooked by health professionals, can highlight areas where the patient’s perspective differs from the views of health professionals, and can ensure that the guideline addresses key issues of concern to patients.

A wide range of other issues can be drawn out by patient representatives to make sure a guideline addresses the needs of all those affected by a condition. The influence of religion/belief on compliance with treatment, for example, complying with a recommended diet or medication, or a different approach to sexually transmitted infection (STI) screening being required for people in prison and those who are homeless.

Patient representatives can also assist the group on the use of clear and sensitive language in the guideline.

10.2 IDENTIFYING PATIENTS’ VIEWS

10.2.1 LITERATURE SEARCH

SIGN has developed a literature search strategy to identify both qualitative and quantitative studies that reflect patients’ experiences and preferences in relation to the clinical topic (see section 4.1). This search is performed at least three months prior to the first group meeting to ensure adequate time to obtain relevant articles and summarise their findings for presentation at the first guideline group meeting.

The types of studies identified generally include patients’ views on:

• positive and negative experiences of the condition, including diagnosis, medication and other treatments, follow-up care and quality of life
• unfulfilled needs
• information needs and preferences
• participation in decision making about treatment
• overall satisfaction with care received.

A copy of the Medline version of the patient search strategy is available on the SIGN website.
10.2.2 PATIENT ORGANISATIONS AND THE SIGN PATIENT NETWORK

SIGN writes to the organisations and charities that aim to represent and/or lobby for patients at least four months before the first meeting of the guideline development group, asking them to inform SIGN of the issues they think the guideline should address. A form is supplied to enable them to structure their feedback in a useful way and, importantly, to indicate the source(s) of their suggestions (e.g. telephone help line data, surveys).

SIGN also writes to members of the Patient Network asking them which issues they think the guideline should address. The Patient Network is a database of patient, carer and other service user representatives. The Network includes contacts for both individuals and organisations, including NHS board Designated Directors for patient and public involvement, equality and diversity group stakeholders (e.g. REACH community health project), previous and current patient representatives on SIGN guideline development groups, representatives from patient advocacy services, representatives from patient support organisations, and representatives from relevant Scotland-wide groups.

10.2.3 OTHER NHS ORGANISATIONS

SIGN writes to various other NHS organisations at least four months before the first meeting of the guideline development group to find out if any local research on patient views has been carried out. This might include, for example, patient focus groups to help in the redesign of services, or questionnaire studies to gauge levels of patient satisfaction with existing services. Reports such as this tend not to be published even though they are in the public domain and can be very useful as a snapshot into current patient issues and concerns regarding particular NHS services and treatments.

10.2.4 DIRECT FEEDBACK FROM USERS OF THE SERVICE

Where published evidence is scarce and inadequate feedback from patient organisations has been received, patient and carer views may be sought through direct contact with users of the service. Techniques used to date have included focus groups with patients in different regions of Scotland, attending patient support group meetings, and SIGN organised meetings for patients and carers. All of these approaches have provided valuable information that has been fed back directly to guideline groups to influence the remit and key questions underpinning the guideline. Often the guideline development group identifies a need for further input from patients and carers at a later stage of the guideline development process. Focus groups can be carried out and the findings used to complement the scientific evidence.

Running focus groups requires expert facilitation. Views are sought from both men and women of different age groups, in both rural and urban communities. Special efforts are made to include those who are socially excluded and may be less likely to join a local or national organisation. SIGN does this by working with healthcare professionals, local community groups and schools who can help identify people to take part.

10.2.5 PRESENTING THE FINDINGS

The Patient Involvement Officer reviews the results of the patient literature search, and seeks to identify common themes that emerge from the literature. These themes are then integrated with the issues that emerge from the other approaches described and presented at the first meeting of the guideline development group by the Patient Involvement Officer.

The group is asked to take account of these issues when it drafts its key questions. Once a first draft of the key questions has been prepared, the Evidence and Information Scientist working with the group along with the Patient Involvement Officer compares the questions with the issues highlighted through the consultative process and highlights any that have not been included in the key questions. At a subsequent group meeting the results of this comparison are presented to the group, and they are asked to consider whether the questions should be revised.

Guideline groups are not obliged to take on board all the issues raised through the patient consultative process, but they are expected to give explicit reasons if they choose to omit particular topics that have arisen from this source.
10.3 RECRUITMENT OF PATIENTS TO GUIDELINE DEVELOPMENT GROUPS

SIGN recruits a minimum of two patient representatives to guideline development groups by inviting nominations from the relevant ‘umbrella’, national and/or local patient focused organisations in Scotland. Where organisations are unable to nominate, patient representatives are sought via other means, for example from consultation with health board public involvement staff. Where patients have been consulted directly (eg if a focus group has been held) this may also provide a source of possible future patient and carer representatives.

Details of the role of the patient representatives, the support they will be given, the commitment required and useful attributes for representatives are provided to allow informed nominations to be made.

10.4 ROLE OF PATIENT REPRESENTATIVES ON GUIDELINE DEVELOPMENT GROUPS

Although their areas of expertise will vary, members of the guideline development group have equal status on the group. A key role for patient and carer representatives is to ensure that patient views and experiences inform the group’s work. This includes:

- ensuring that key questions are informed by issues that matter to patients
- identifying outcome measures they think are important for each key question
- considering the extent to which the evidence presented by group members has measured and taken into account these outcome measures
- identifying areas where patients’ preferences and choices may need to be acknowledged in the guideline
- making sure that the degree to which the evidence addresses patients’ concerns is reflected in the guideline
- helping to write the ‘Provision of information’ section of the guideline, including identifying sources of further information
- raising awareness of patient issues at the National Open Meeting by preparing a presentation and speaking at this meeting
- assisting SIGN with the identification of voluntary organisations and charities to invite to the National Open Meeting
- helping to ensure that the guideline is sensitively worded (eg treating patients as people and not as objects of tests or treatments)
- identifying individuals to take part in the peer review process
- assisting SIGN with the collection of patient views (eg by helping to prepare questions for focus groups)
- helping SIGN with consultation arrangements
- appraising literature (if the individual chooses to do so)
- raising awareness of the SIGN guideline among members of their support group and members of the public.

No formal qualifications are needed but it may be helpful if patient representatives have some of the following:

- experience of the guideline condition (eg as someone who has, or has had the condition, or a carer or relation of someone who has or has had the condition)
- an understanding of the experiences and needs of a wider network of patients (eg as a member of a patient support group)
- time to commit to the work of the group (eg attending meetings, background reading, commenting on drafts)
- some familiarity with medical and research language (although members of the guideline group should help with specific technical terms)
- willingness to feed in the views of patient/carer groups not represented on the guideline group
- ability to be objective
- good communication and team working skills.
10.5 SUPPORT FOR PATIENT REPRESENTATIVES ON GUIDELINE DEVELOPMENT GROUPS

SIGN supports patient representatives by:

- delivering 'Introduction to SIGN' training, based on SIGN 100: A handbook for patient and carer representatives for patient representatives
- offering telephone and email support
- inviting new patient representatives to join the SIGN Patient Network
- providing clear guidance on their roles and responsibilities within the group
- ensuring opportunities to attend training events are open to all guideline development group members, including patient representatives
- inviting patient representatives to informal events.

The Chair of each guideline development group is asked to support patient representatives by:

- ensuring patient representatives are fully engaged with the group
- addressing the group if contributions by patient representatives are not acknowledged appropriately
- welcoming and encouraging contributions from patient representatives.

10.6 WIDER CONSULTATION WITH PATIENTS AND CARERS

Further patient and public participation in guideline development is achieved by involving patients, carers and voluntary organisation representatives at the National Open Meeting which is held to discuss each draft guideline (see section 7.1). The meetings are advertised widely and are free of charge.

Patient representatives are invited to take part in the peer review stage of each guideline and specific guidance for lay reviewers has been produced.

Members of the SIGN patient network are also invited to comment on draft documents such as patient versions of guidelines, patient sections of guidelines and other literature aimed at patients.

10.7 REFERENCES


11 Development of the manual

11.1 INTRODUCTION
SIGN is a collaborative network of clinicians, other healthcare professionals and patient organisations and is part of Healthcare Improvement Scotland.


11.2 ACKNOWLEDGEMENTS
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All members of the SIGN Executive and SIGN Council make yearly declarations of interest. Registers of interests for each are available on the contacts page and on the SIGN Council membership page of the SIGN website www.sign.ac.uk

11.3 SIGN EDITORIAL REVIEW
As a final quality control check, this manual was reviewed by an editorial group.

Dr Roberta James SIGN Programme Lead; Co-Editor
Professor John Kinsella Chair of SIGN; Co-Editor
Dr Sara Twaddle Director of SIGN; Co-Editor

All members of the SIGN Executive make yearly declarations of interest. A register of interests is available on the contacts page of the SIGN website www.sign.ac.uk
The Healthcare Environment Inspectorate, the Scottish Health Council, the Scottish Health Technologies Group, the Scottish Intercollegiate Guidelines Network (SIGN) and the Scottish Medicines Consortium are key components of our organisation.