SHORT ORAL PRESENTATIONS

001 DECIDE: USER INVOLVEMENT IN GUIDELINE DEVELOPMENT

M Callaghan, R Harbour, Healthcare Improvement Scotland, Glasgow, UK
10:1136/bmjqs-2013-002293.32

The European Union funded project, DECIDE aims to improve the dissemination of evidence based recommendations by building on the work of GRADE. Work Package 3 aims to identify approaches to effective dissemination and use of research evidence by the general public, in order to improve shared health-care decision making and person-centred care. This part of the study explores public understanding of clinical guidelines and preferences over future presentation. Focus groups were carried out, to discuss specific guidelines (depression, diabetes, flu vaccination and cervical cancer), and with professionals working in science communication and young people. Questions were informed by a systematic literature review. Data were analysed using the Framework method. Those with experience of a specific illness were better informed and more likely to have actively sought out information than those targeted for screening. Themes emerged including: participants not understanding the concept of ‘strong’ and ‘weak’ evidence; participants feeling information should be presented more clearly with emphasis on self-management and side effects; concern expressed that treatment decisions were based on cost rather than evidence; and participants wanted more general information on the condition. Aspects of the content and presentation of guidelines are not relevant or understood by the public. Several different versions of these guidelines have been created based on these results and user testing is ongoing. Current clinical guidelines will need to be adapted substantially before they are appropriate for the public, and the public should be involved in guideline development on an ongoing basis.

002 PUBLIC ATTITUDES TO AND KNOWLEDGE OF HEALTHCARE GUIDELINES, AND METHODS TO COMMUNICATE GUIDELINE RECOMMENDATIONS TO PATIENTS AND THE PUBLIC – A SYSTEMATIC REVIEW

K Loudon, N Santosco, M Callahan, R Harbour, J Thornton, E McFarlane, S Treweek.
1University of Dundee; 2McMaster University, Hamilton, Canada; 3Healthcare Improvement Scotland; 4National Institute for Health and Clinical Excellence, Manchester, UK; 4University of Aberdeen
10:1136/bmjqs-2013-002293.33

Background Improving patient versions of guidelines is one way to support an increasing role for patients in health decision making.

Objectives To evaluate evidence on the public’s attitudes and knowledge of clinical practice guidelines AND what strategies have been used to communicate guideline recommendations to this group.

Methods We conducted a systematic review of public attitudes and knowledge of clinical practice guidelines to inform the development of dissemination strategies for this population. We searched health databases from 2000 to 01/2013, grey literature, and we contacted guideline producers. Two reviewers independently abstracted, coded, synthesised themes from the studies.

Results We included 25 published studies and three reports (CCPG, NICE, SIGN). There was a huge variation in the public’s awareness of guidelines (12 studies) from 25–75%. The key themes to communicate guidelines (15 studies) to the public included, predictive factors (amount of education), personalisation, access to care, trustworthiness (evidence behind recommendations), and self-management.

Discussion Although there were few studies for thematic analysis there were recurrent themes. When developing patient versions, the danger could be to focus on detailed formatting instead of fundamental issues around whether patients dismiss guidelines as not applicable to their unique situation and restricts care. The results will inform work focused on the public and patients being done in EU FP7 DECIDE project.

Implications Guideline producers will need to increase the public’s awareness of clinical guidelines and developing communication strategies that are clearly personally applicable, trustworthy and useful for patients and carers managing their care.

003 TOWARDS OPTIMAL PATIENT INVOLVEMENT IN GUIDELINE DEVELOPMENT GROUPS

1Athina Institute, Free University, Amsterdam, The Netherlands; 2Integraal Kankercentrum Nederland, Utrecht, The Netherlands; 3Nederlandse Federatie van Kankerpatienorganisaties, Utrecht, The Netherlands
10:1136/bmjqs-2013-002293.34

Background The involvement of patient representatives in guideline development groups (GDGs) could increase legitimacy and quality of clinical practice guidelines (CPGs), since the experiential knowledge of patients could complement scientific evidence. By their involvement, patients have the opportunity to share (consultation) and incorporate their views and experiences into CPGs (decision-making). Although the importance of this approach is emphasised, little methodological support and systematic reflection exist on effective strategies.

Objectives To gain insight in how patient involvement in GDGs can be optimised in order to develop strategies which can be implemented in practice.

Methods The evaluation consisted of a desk study and 23 semi-structured interviews with stakeholders in CPGs, including patient representatives. The acquired insights were used to develop an evaluation framework, which guided monitoring and evaluation of four ongoing oncological guideline development processes. Validation took place through a triangulated approach (e.g. observations, document-analyses, interviews). Two patient representatives were included in the research team.

Results The evaluation revealed that successful patient involvement in GDGs depends on a broad scale of factors (e.g. members of the GDGs, support of patient representatives) which could facilitate or constrain patient involvement. The factors were used to develop practical strategies for patient involvement, ranging from preparation meetings to regular reflections with the patient representatives and dialogue sessions with patients.

Discussion The strategies could lead to more successful patient involvement in GDGs and provide valuable insights on how to involve patients in guideline development processes on other disease areas.

004 PARTNERING TO TRANSFORM CLINICAL RESEARCH INTO EVIDENCE-BASED HEALTH CARE GUIDELINES

N Wasson, C Chang, MEB Smith, A Qaseem, M Starkey, O Buckley, R Chou, S Saha.
1The Pacific Northwest Evidence-based Practice Center, OHSU, Portland, USA;
Background A guideline-making body nominated pressure ulcer risk assessment, prevention, and treatment as evidence review topics to support the development of clinical practice guidelines, partnering with a funding agency and a systematic review team to conduct the research.

Context Collaboration may enhance evidence-based health care given that multiple organisations bring diverse resources and expertise to the process of guideline development. By partnering to develop systematic reviews (SR) with focused research questions, funders, review teams, and guideline committees can effectively evaluate and synthesise the voluminous evidence required to inform guidelines.

Description of Best Practice We describe our processes for linking reviews to guideline development, including: Nomination/refinement of focused review topics; Clearly defined roles for each participant; Development of comprehensive SRs; Well-defined processes to preserve the scientific integrity of the review while allowing for input from stakeholders; Stakeholder and funder participation throughout the review process; Development of the guideline; Publication of the research results and guidelines.

Lessons for Guideline Developers, Adaptors, Implementers, and/or Users Challenges include balancing the interests of the nominator/guideline developer and a broader stakeholder audience; answering the clinically important questions needed to develop a guideline; effectively presenting the findings; and coordinating among groups. Collaboration ensures that SRs are focused and relevant to guideline committees, aiding in the development of research that meaningfully informs clinical guidelines. Synergy between partner organisations can lead to wider dissemination of findings and facilitate timely guideline development for implementing best practices to improve health outcomes.

005 CLINICAL PRACTICE GUIDELINES AND SYSTEMATIC REVIEWS: POINT OF INTERSECTION?

1L Wilson, 1K Robinson. 1Department of Health Policy and Management, Johns Hopkins University, Baltimore, USA; 2Department of Medicine, Johns Hopkins University, Baltimore, USA

Background Guideline committees (GCs) rely on the evidence synthesised in systematic reviews (SRs) to develop evidence-based guidelines. Institute of Medicine standards for clinical practice guidelines include an interaction between the GC and the team conducting the SR.

Context In 2005, the Cystic Fibrosis Foundation moved from consensus-based to evidence-based guideline development. SRs are now commissioned to inform specific guidelines. A methodologist, serving as a member of each multidisciplinary GC and as the lead investigator for the SR teams, provides the link for the scope, approach, and output of both processes.

Description of Best Practice The methodologist, as part of the GC, facilitates the definition of the scope and refines the questions for the SRs. The methodologist oversees the conduct of the SR, ensuring that the review team addresses relevant questions, appropriately conducts searches, and establishes inclusion criteria and provides informative details to the GC. The methodologist provides training, where needed, and ensures consistency across guidelines in the drafting and grading of the recommendations. The methodologist also helps to address peer review comments and draft the guideline documents.

Lessons for Guideline Developers, Adaptors, Implementers, and/or Users Having a methodologist serve on both the GC and the SR team ensures that there is appropriate and timely intersection of the guideline and SR processes. The methodologist can ensure that the SR team meets the needs of the GC and illuminate for the GC the methods and outcomes of the SR.

006 TRANSLATING KNOWLEDGE INTO PRACTICE: A SYSTEMATIC REVIEW OF BARRIERS, FACILITATORS AND INTERVENTIONS IMPACTING ON UPTAKE OF SYSTEMATIC REVIEWS

1J Wallace, 1C Byrne, 1M Clarke. 1Oxford University, Oxford, UK; 2Health Service Executive, Dublin, Ireland

Background The increased uptake of evidence from systematic reviews is advocated because of their potential to improve the quality of decision making for patient care and their use in clinical practice guidelines.

Objectives To identify how uptake of evidence from systematic reviews can be enhanced.

Methods Data sources: We searched 19 databases covering the full range of publication years. Study selection: Studies of barriers and natural facilitators to uptake of evidence from systematic reviews and meta-analyses were eligible. These studies encompassed survey, focus group and interview designs. Intervention, or outcome, studies were also included. Data extraction: Two reviewers independently assessed quality and extracted data that were summarised and then analysed. Using a pre-established taxonomy, the barriers and facilitators were organised into a framework according to their effect on knowledge, attitudes, or behaviour. For the intervention studies, two reviewers also independently assessed quality and extracted data. Data synthesis: Twenty-seven studies dealing with barriers were detected and 15 studies that included investigation of natural facilitators. Ten publications addressing interventions met inclusion criteria. A synthesis of findings was conducted to find out to what extent the interventions overcome the perceived barriers and built on the facilitators detected.

Results Educational visits, summaries of systematic reviews, and targeted messaging had a significant impact on systematic review uptake and also addressed a range of identified barriers and facilitators.

Conclusion On the basis of this study, specific strategies addressing a range of barriers and facilitators are recommended to enhance uptake of systematic reviews and meta-analyses. Promising interventions are also identified.

007 PARTNERING TO TRANSFORM CLINICAL RESEARCH INTO EVIDENCE-BASED HEALTH CARE GUIDELINES

1L Al-Ansary, 1Y Amer, 1R Fattouh, 2Y Adi. 1Department of Family and Community Medicine, College of Medicine, King Saud Univ, Riyadh, Saudi Arabia; 2Bahamdan’s Research Chair for Evidence-Based Health Care & Knowledge Translation, Riyadh, Saudi Arabia; 2Quality Management Department, King Saud University Medical City,