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Title: Engaging stakeholders and promoting uptake of OMERACT core outcome instrument sets


Abstract [250 words]

Objectives: While there has been substantial progress in the development of core outcomes sets, the degree to which these are used by researchers is variable. We convened a special workshop on knowledge translation at OMERACT 2016 with two main goals. The first focused on the development of a formal knowledge translation framework and the second on promoting uptake of recommended core outcome domain and instrument sets.

Methods: We invited all 189 OMERACT 2016 attendees to the workshop; 86 attended representing the following stakeholders: patient research partners (n=15), healthcare providers/clinician researchers (n=52), industry (n=4), regulatory agencies (n=8), and OMERACT fellows (n=11). Participants were given an introduction to knowledge translation and were asked to propose and discuss recommendations for the OMERACT community to: 1. strengthen stakeholder involvement in the core outcome instrument set development process, and 2. promote uptake of core outcome sets with a specific focus on the potential role of post-regulatory decision makers.

Results: We developed the novel “OMERACT integrated knowledge translation” framework which formalizes OMERACT’s knowledge translation strategies. We produced strategies to improve stakeholder engagement throughout the process of core outcome set development and created a list of creative and innovative ways to promote the uptake of OMERACT’s core outcome sets.
Conclusions: The guidance provided in this paper is preliminary and is based on the views of the participants. Future work will engage OMERACT groups, “post-regulatory decision makers”, and a broad range of different stakeholders to identify and evaluate the most useful methods and processes and revise guidance accordingly.

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A short running footnote of no more than 4 words
OMERACT Knowledge Translation Guidance
INTRODUCTION

To address the critical need for standardized outcome domains and measurement instruments in rheumatology clinical trials, the international organization Outcome Measures in Rheumatology (OMERACT) has developed core outcome sets of domains and instruments using an iterative, data-driven, consensus-based approach (1,2). A key aspect of its research methodology is to obtain input from a broad range of stakeholders, including patient research partners, industry, and regulators. OMERACT’s strategies to promote update of recommended core outcome sets have predominantly focused on reaching clinical trialists, methodologists, regulators, and industry partners via journal publications.

For OMERACT 2016, a special workshop on knowledge translation was convened with two main goals. The first focused on the development of a formal knowledge translation framework to incorporate state of the art innovations in knowledge translation and to strengthen stakeholder engagement and ensure effective input representing a broad range of interests. The second goal focused on promoting uptake of recommended core outcome instrument sets.

Knowledge translation (3) was defined to workshop participants as activities that make users aware of knowledge (i.e. core outcome instrument sets) and that facilitate the use of this knowledge to improve health and health care systems with an aim to close the gap between what we know and what we do (i.e. using OMERACT-endorsed core outcome sets in clinical trials, systematic reviews, etc.).

Knowledge translation can be split into two components: Integrated and End-project knowledge translation. Integrated knowledge translation – also known by such terms as collaborative research, participatory research, engaged scholarship, co-production, and co-creation – is a collaborative or participatory approach that engages end-users in the research process, starting with their involvement in defining the research question (4,5). This engagement occurs with the expectation that it will result in
research outputs that are more relevant, useful, and readily useable to the end-users and therefore more likely to be implemented (4). Effective engagement requires additional considerations both for patients and other stakeholders (6,7). Different stakeholder groups may be broadly defined by the following “Ps”: Patients and their families, the Public, Providers, Payers/Purchasers, Policymakers, Principal investigators [researchers and funders], Product makers, and others, such as. the Press (8,9).

End-project knowledge translation is about translating research findings into policy and practice and is essential for optimizing the impact of research. In OMERACT’s case, the adoption of core outcome sets by clinical trialists, systematic reviewers, guideline developers, regulators, and others helps ensure comparability across studies and improves the ability to synthesize and interpret the evidence base of interventions for rheumatic conditions (10-12). This translation of research into policy and practice is best viewed as a process within which there are various considerations. Among these are defining the specific (current and potential) contexts of use for a given outcome or core set, identification of relevant stakeholders, developing an engagement plan, establishing a strategy for promoting uptake, and enabling plans for implementation and measuring uptake and impact. Table 1 lists specific steps to consider across the different stages.

**[INSERT TABLE 1]**

This original research article:

- Describes the novel “OMERACT integrated knowledge translation” framework which formalizes OMERACT’s knowledge translation strategies
- Provides specific strategies on whom to involve and how to involve them in order to improve stakeholder engagement throughout the process of core outcome instrument set development
- Describes creative and innovative ways to promote the uptake of OMERACT’s core outcome instrument sets
• Explains the potential influence of “post-regulatory decision makers” in increasing the uptake of OMERACT’s core outcome instrument sets
• Offers examples of stakeholder engagement and uptake strategies used by two current OMERACT Working Groups throughout their projects

METHODS

We held a workshop session during OMERACT 2016 to generate ideas for developing “best practices” in stakeholder engagement for OMERACT working groups and to promote the uptake of core outcome instrument sets. All 189 OMERACT 2016 attendees were invited to the workshop and 86 attended. Workshop participants included patient research partners (n=15), healthcare providers/clinician researchers (n=52), industry (n=4) and regulatory agency (n=4) representatives, and OMERACT fellows (n=11). Two presentations at the start of the session provided participants with a broad introduction to knowledge translation. These were followed by presentations from two current OMERACT Working Groups on their strategies for stakeholder engagement and promoting uptake of their work. The Rheumatoid Arthritis Flare Working Group has been working for several years to establish a means to capture clinically significant worsening of rheumatoid arthritis disease activity, primarily as an outcome measure for use in clinical trials, but also for potential use in other settings including clinical practice (13). The Worker Productivity Working Group has sought to identify instruments that could be used to measure at work productivity loss due to rheumatologic conditions (14). Examples from these two groups are presented in this paper as case studies.

Workshop participants then moved to six breakout groups each led by two OMERACT Executive members. Participants were asked to develop and discuss recommendations on strategies for the OMERACT community to: 1) strengthen stakeholder involvement in core outcome set development, and
2) promote the uptake of core outcome sets. A rapporteur from each breakout group presented key findings back to the entire group.

RESULTS

Goal 1. Strengthening engagement with stakeholders during the development of core outcome sets

Who to involve?
Establishing the context(s) of use serves as an important starting point for how a group would begin to consider who should be engaged in a research project to ensure utilization. For maximum effectiveness, broad engagement should occur throughout the entire core outcome set development process, from conceptualizing the question, to developing the research agenda and protocol, conducting the research itself, seeking interpretation and comments on the results, and creating audience-specific information to promote the uptake and use of the recommendations (Figure 1). However, the simple formulation that every stakeholder should be equally involved from beginning to end is likely not the most effective or efficient approach.

[INSERT FIGURE 1]

Optimal stakeholder engagement requires identification of the right people, their involvement at the appropriate phases of the core outcome set research process and the integration of their perspectives in the best possible way to maximize the impact of their input. The overall research program to develop a Core Outcome Instrument Set represents an ongoing and iterative process, with different types of input required along the way (e.g. qualitative expertise in domain identification and content validation, psychometric expertise in instrument evaluation and development). It is not necessary to involve every conceivable stakeholder at each stage to an equal extent. It is important to create a shared understanding among stakeholders and researchers concerning their expected roles within the overall process, and the commitments that will be asked of them (e.g. time, travel, etc.). It should also be
understood that their involvement may differ depending on the stage of the research project.

OMERACT has an extensive history of engaging with patient research partners (people living with a disease or condition who actively and equally contribute to research projects) to ensure that the patient perspective is captured when identifying important outcome domains (15-20). The Grading of Recommendations Assessment, Development and Evaluation (GRADE) Working Group has a ‘GRADE Stakeholders Group’ project that is currently underway. The group provides guidance on which stakeholders should be involved in the GRADE guideline development process, and how to identify, engage, and involve them. This GRADE guidance may be relevant for OMERACT to consider when developing integrated knowledge translation strategies.

Participants at the OMERACT 2016 workshop identified various stakeholders that should be considered (Table 2). In order to ascertain who best to engage, they recommended identifying and networking with key opinion leaders in relevant clinical areas.

[INSERT TABLE 2]

Case Studies: Who to involve?

1. Rheumatoid Arthritis Flare Working Group

- Brought together an international, multidisciplinary group consisting of patients, providers (physicians, nurses, psychologists, and other allied health professionals), clinicians who perform clinical trials and those who design studies from academia and industry, funders, research experts (qualitative researchers, biostatisticians, clinical epidemiologists, and psychometricians), and pharma and regulatory representatives.
• Engaged fellows and trainees to ensure a pipeline of individuals committed to ultimate adoption of the outcomes being studied, and for them to observe engagement of partners in research.

2. Worker Productivity Working Group

• Engaged with an international and interdisciplinary group of stakeholders including patient research partners and representatives from epidemiology, health economics, industry partners, rehabilitation, rheumatology, work disability research field, the International Canadian Arthritis Network for Work Outcomes (I CAN Work) and a wide array of arthritis researchers.

How to engage?

During the OMERACT workshop, participants suggested the following strategies for better stakeholder engagement:

1. Involve the “right” stakeholders from the start of the project and throughout the development of the core outcome set, while acknowledging that consideration should be given to which stakeholders should be involved, and to what extent, at each phase of the core outcome set development project.

2. Provide stakeholders in a timely manner with necessary information such as pre-reading materials, to ensure informed engagement.

3. Hold working group meetings at large national and international conferences where different stakeholders often meet to increase opportunities for face-to-face involvement; consider paying expenses for key stakeholders to attend.

4. Expand the stakeholder community using virtual meetings and voting.
Case studies: How to engage?

1. Rheumatoid Arthritis Flare Working Group

- Held ongoing interactions between face-to-face meetings via tele/web conferences and other means of communication (e.g. email).
- Developed pre-briefing/debriefing calls and specific education sessions for patient research partners; tools such as the OMERACT glossary were found to be particularly helpful.
- Engaged patient research partners throughout the process in the following ways with major roles in participation/leadership in the Working Group: participating as members of the Steering Committee, analyzing qualitative data, developing the questionnaire, debriefing the questionnaire, interpreting results, facilitating and moderating OMERACT plenaries, presenting research findings at other international meetings, authoring publications.
- Presented research results at various points in the instrument development process at biannual OMERACT meetings; used marketing materials to raise awareness and obtain endorsement from OMERACT participants. At meetings, working group members wore t-shirts, brooches and an extra name tag that said “ASK ME ABOUT RA FLARE”. A wide-ranging audience representing multiple constituencies in small breakout groups provided important feedback.

2. Worker Productivity Working Group

- Started knowledge translation engagement efforts early, first to introduce the domain, establish need and identify people eager to be involved; included a highly active and encouraging patient group from the beginning; continued to hold meetings at biannual OMERACT meetings as well as international workshops in between.
• Held an “overt” engagement blitz at OMERACT 2014 using the branding “It Works!” with group members wearing t-shirts to advertise whom to ask questions of throughout the conference, along with post-it notes, pamphlets and tables summarizing the evidence. A fun, high-energy breakout session was held using the format of speed dating with various working group members providing the OMERACT Filter evidence on different instruments as session participants moved around the room.

• Patient research partners contributed equally to design and proof of questionnaires and surveys, interpretation of findings, and assistance with study recruitment through their networks. However, their biggest influence was identifying contextual factors and expanding understanding of the concept of interest so that measuring productivity meant understanding the job situation, e.g. “If they could give me flexible hours I would be at 100%, but now I am at 75%”.

Goal 2. Promote the uptake of core outcome sets

Participants suggested creative and innovative ways of transferring information, beyond traditional peer-reviewed publications and presentations at professional meetings, were suggested by participants. Table 3 describes specific considerations for promoting uptake of core outcome sets. The potential factors that could influence the implementation of core outcome sets which were described to the workshop participants during the first part of the session are outlined in Table 4.

[INSERT TABLE 3]

[INSERT TABLE 4]
Workshop participants suggested the following approaches to promote the uptake of published core outcome sets and other key OMERACT findings to the broader community:

1. Revise the OMERACT website to make it easier for people to find key information; provide an RSS feed to deliver updated website content; actively use social media (e.g. Twitter, Facebook).

2. Highlight OMERACT achievements through an OMERACT newsletter to disseminate highlights from finished work; more lay publications (not just academic journals), e.g. ACR/EULAR highlights.

3. Hold an “OMERACT Findings Symposium” attached to ACR/EULAR or other locations where stakeholders such as payers, regulators, outcomes methodologists, and health technology assessment agencies hold their major conferences; send an OMERACT representative to major meetings.

4. Develop a toolkit using plain language to describe both the methods behind the development of core outcome sets and the resulting set of recommended outcomes; use short messages targeted to different stakeholders; develop an app for the toolkit.

5. Use standard-practice marketing strategies, e.g. consider the presence, profile, and penetration of OMERACT in the different stakeholder groups; use story-telling; evaluate marketing plans after implementation.

6. Conduct strength, weaknesses, opportunities, challenges/constraints (SWOC) analysis for plans to promote the uptake of recommendations.

7. Early in their research program, OMERACT working groups should develop a promotion/marketing strategy to implement the core outcome set; consider the relevance of OMERACT to each stakeholder group and prioritize groups.

8. Deposit OMERACT core outcome sets in outcome measurement repositories, e.g. EULAR Outcomes Measures Library [http://oml.eular.org/], FDA’s compendium of clinical outcome
assessments, Mapi Research Trust and Patient-Reported Outcome and Quality of Life Instruments Database (PROQOLID, https://eprovide.mapi-trust.org), and other relevant databases.

9. Engage patient research partners to work with the committees, associations, and arthritis communities with which they are involved to increase OMERACT’s profile at the grassroots level; link to patient organization websites (e.g. www.creakyjoints.org).

10. Continue the concept of “generosity of ideas and collaboration” to help spread information from OMERACT’s work.

Moving from dissemination to facilitating implementation: Engaging payers and other “post-regulatory decision makers”

A specific focus of the workshop was on the potential influence of “post-regulatory decision makers” in increasing the uptake of core outcome instrument sets. The importance of regulators such as the Food and Drug Administration (FDA) and European Medicines Agency (EMA) in promoting implementation of core outcome instrument sets has been well recognized by OMERACT and other developers of core outcome instrument sets for many years. Regulators have defined mechanisms to review, approve and communicate preferred outcomes through guidance documents, compendia, etc. Researchers from the life sciences industry and elsewhere are highly motivated to pay close attention to the health outcomes that are recognized by regulators, given the implied significance of those outcomes in regulatory decisions. Less well appreciated has been the potential impact of “post-regulatory decision makers” (Table 5) in creating strong incentives for researchers to use health outcomes recognized by these decision makers. The premise behind working with these groups is their explicit recognition of core outcome sets as influential in their decision making. This would create strong incentives for researchers
to use those core outcome sets, much as FDA recognition of core outcome sets is a strong motivator for their use (26).

[INSERT TABLE 5]

Because each of these organizations directly or indirectly influences the speed and extent of market uptake of new drugs, devices, diagnostics and procedures — and the prices paid for these products — the role of health outcomes in their decision making has important practical consequences for product developers and other researchers. For this reason, when and how to effectively and efficiently engage these stakeholders in core outcome instrument set development and strategies for promoting uptake is an important area for further exploration. It is unlikely that representatives of these groups will be available to actively participate in all phases of core outcome set development. Potential conflicts of interest should always be considered and made explicit.

Case studies: Strategies for promoting uptake

1. Rheumatoid Arthritis Flare Working Group

- Developed text and video stories about the importance of patient reported outcomes as outcomes from multiple perspectives (patients, clinicians, researchers), results from a research study, and the impact patient reported outcomes had on making health decisions [www.hopkinsarthritis.org/PCOR](http://www.hopkinsarthritis.org/PCOR).
- Made plans to further disseminate this information via social media to a larger community. Such efforts require additional expertise (for example, from medical writers and media professionals) to provide appropriate context and to make information accessible for stakeholders.

2. Worker Productivity Working Group
• Published and presented findings and involved working group members in promoting results within existing networks.

• Considered further dissemination of these findings on the measurement properties of instruments measuring work productivity to potential users. These may include health technology assessment agencies, “owners” of the instruments, work disability researchers, and policy decision makers.

DISCUSSION

Challenges

Workshop participants noted potential challenges in implementing the ideas outlined above. Core outcome instrument sets and the methodology behind developing them is complex, and clear communication of this information can be difficult. Many different stakeholders were identified and methods are needed to prioritize which stakeholders to target in which stages of the process. Leveraging networks of patient research partners is one essential strategy to pursue for improved uptake but may require developing training materials in lay language. Consistent with the recent recommendations for patient research partner involvement in OMERACT research projects (27), further work is needed to develop and standardize training. This may involve training of researchers in engagement strategies with patient research partners. The issue of weighting patient involvement — including weighted patient voting to ensure that they are not a minority likely to be outvoted by other stakeholder groups—was also raised.

OMERACT is an international organization and an ongoing challenge is to ensure geographical representation. Many major national and international conferences do not allow concurrent meetings by other organizations, which may limit the ability to engage with stakeholders at these venues.
Recognition of and explicit discussion about real and potential conflicts of interest is important and necessary to ensure the integrity of OMERACT’s program of work. Lastly, the strategies to facilitate implementation of core outcome sets should not be an afterthought; initiating these strategies is resource-intensive and it takes a substantial amount of time and energy, thus requiring planning and budgeting from the beginning of every initiative.

Limitations

This guidance is preliminary and is based on the views of the participants who attended OMERACT 2016. While there was good representation from the different stakeholder groups at OMERACT, it is necessary to include a larger number of stakeholders when evaluating the strategies discussed in this paper. The focus of discussion was within the field of rheumatology, and further work with core outcome set developers in other fields would be useful. The knowledge translation concepts were adapted from work that was focused on the dissemination and implementation of research findings; core outcome set development and promotion may not be precisely comparable.

Research agenda for promoting uptake

Next steps include prioritizing the approaches suggested above for promoting uptake of OMERACT Core Outcome Instrument Sets. We will also focus on clarifying how best to work with “post-regulatory decision makers” by collaborating with the Center for Medical Technology Policy to define a parallel set of mechanisms through which post-regulatory decision makers could recognize health outcomes that best inform their decision making (28). These mechanisms will be identified by conducting a series of interviews with key representatives and a meeting during which we will identify and evaluate potential mechanisms through which these organizations could encourage implementation of core outcome instrument sets. We will engage with OMERACT groups, stakeholders, as well as other core outcome set
developers to evaluate their experiences with implementing these knowledge translation approaches and revise our guidance as necessary.

CONCLUSIONS

OMERACT has developed an international reputation for high-quality, leading-edge methodology over the last twenty-five years, and we now recognize the need to strengthen our engagement with stakeholders as potential users of the products of our work, and to market the evidence-based, consensus-driven core outcome sets that we have established. Further work on promoting uptake of core outcome sets is now underway, through collaboration between the Center for Medical Technology Policy (CMTP) and OMERACT, with a focus on “post-regulatory decision makers”. The OMERACT Executive will engage with OMERACT Working groups to identify the most useful knowledge translation methods and processes. These will be used to inform recommendations in a chapter in the OMERACT Handbook on engaging stakeholders and strategies for promoting uptake of core outcome sets to support individual OMERACT Working Groups. We will undertake evaluation of our knowledge translation strategies on an ongoing basis.
References


17. de Wit MPT, Kirwan JR, Tugwell P, Beaton D, Boers M, Brooks P, et al. Successful Stepwise Development of Patient Research Partnership: 14 years’ experience of actions and

18. de Wit M, Abma T, Koelewijn-van Loon MS, Collins S, Kirwan J. Involving patient research partners has a significant impact on outcomes research: a responsive evaluation of the international OMERACT conferences. BMJ Open 2013;3:e002241

19. de Wit MP, Abma TA, Koelewijn-van Loon MS, Collins S, Kirwan J. What has been the effect on trial outcome assessments of a decade of patient participation in OMERACT? J Rheumatol 2014;41:177-184


