Improving communication about cancer screening: moving towards informed decision making

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Abstract

For decades, public communications about cancer screening have used persuasive techniques with the aim of maximising the number of people being screened. However, perspectives have changed more recently to acknowledge that screening can lead to harm as well as benefit, and that it is important for service users to consider both. For some types of cancer screening, there is professional contention about whether benefits clearly outweigh harms. In light of this, an emerging trend in cancer screening communication is to try to support informed decision making – that is, to help people understand both the advantages and disadvantages of screening, allowing them to make individual decisions about their screening participation that reflect their informed preferences. In this review, we provide an overview of key theoretical and practical aspects of improving communication and supporting informed decision making about cancer screening, highlight relevant research and discuss future implications.

Introduction

Cancer screening involves a complex array of potential consequences, including benefits (e.g. reduced disease burden, mortality and morbidity; improved disease outcomes) and harms (e.g. false positives, false negatives, overdiagnosis, other physical and psychological harms).\textsuperscript{1} There is variation across cancer types and screening modalities in the possible consequences for the individual screened, the probabilities of the various outcomes and the ratio of benefits to harms. These variations underpin differences in recommendations about whether and how screening should be offered and implemented. Some forms of screening are generally regarded as advantageous (e.g. bowel cancer screening with the faecal occult blood test); some are viewed more negatively (e.g. prostate cancer screening with prostate-specific antigen testing); and others are seen to have a finer balance between benefits and harms (e.g. breast cancer screening with mammography).\textsuperscript{2} Heterogeneity of study methodologies,
population characteristics and screening delivery can lead to conflicting results and uncertainty about estimates of benefits and harms, which complicate the task of making policy decisions about whether and how screening should occur.

Despite these complexities, policy makers are increasingly seeking to enable informed decisions by providing balanced information on harms and benefits to potential screening participants. In this review, we give an overview of current research and practice relating to improving communication about cancer screening, including relevant examples across a range of contexts. At times, we focus on breast cancer screening as a case where communication issues have received intense attention in recent years.

**Public perceptions of screening**

Attitudes about cancer screening are highly positive among the general public, and screening is often supported uncritically. Qualitative research suggests that key reasons motivating people to undergo screening include taking responsibility for one’s own health, feeling a social obligation to do so, and gaining reassurance. A landmark US survey documented widespread public enthusiasm for routine cancer screening, with 87% of respondents considering it “almost always a good idea”. Very similar results were reported a decade later in a large British survey. People’s commitment to screening was strong enough that 58% of women in the US survey said they would overrule a physician who suggested having less frequent cervical screening, whereas 77% of men would continue prostate screening, and 74% of men and women would continue bowel screening even if their physician recommended against it. These attitudes may be based on unrealistic expectations: a systematic review of studies that quantitatively assessed patient or public expectations of medical interventions, including 15 studies on cancer screening, found that participants tended to overestimate benefits and underestimate harms.

**Conventional approach to providing information**

Public health messaging about cancer screening for many decades has largely reflected the positive views held by public health organisations, professional associations, patient advocacy groups, academics and clinicians. According to these favourable perspectives on screening, authorities are justified in using persuasive communication tools with the goal of maximising uptake. These messages can take diverse forms. One frequently used approach is to induce feelings of fear and vulnerability, then offer hope by framing screening as a simple method of protection. A common feature of many conventional approaches to screening communication is to emphasise benefits and minimise harms. For example, several international reviews (most recently by Gummersbach et al.) have found that breast screening information materials overestimate benefits and underplay harms. Such communications have created highly positive community views of screening that engender normative expectations that screening is the ‘right’ thing to do.

**Need for informed (shared) decision making**

Messaging about screening has been criticised in recent years as simplistic and overly positive. It is argued that, rather than encouraging uptake, screening communication should aim to inform people about harms and benefits so that they can make informed decisions about participation. The ethical justification is that screening puts large numbers of healthy people at risk of possible harm by offering interventions they might not otherwise seek out. Advocates of this perspective note that screening can trigger a cascade of serious interventions with lasting consequences, and individuals who choose to screen (or not) must live with their decision and its repercussions. Individuals may weigh attributes differently depending on their personal circumstances and psychological characteristics – and some might reasonably decline screening. According to this perspective, screening decisions are a good example of the value of helping people be engaged and involved in decision making.

The terms ‘informed decision making’ and ‘shared decision making’ relate to public and patient involvement in healthcare decision making. Both terms encompass a process that enables a person to participate in making a healthcare decision, having learnt about the intervention and its likely consequences, and having considered their preferences. Shared decision making is conceptualised specifically in terms of patient–clinician consultations in which both parties express preferences and participate in making decisions. Shared decision making may therefore be more relevant to screening tests based in primary care (e.g. prostate cancer screening) than those provided outside a clinical encounter (e.g. breast and bowel cancer screening), where informed decision making might be the more appropriate term.

**Supporting informed (shared) decision making with decision aids**

Decision aids are designed to support people’s decision making about healthcare interventions in circumstances where more than one reasonable option is available, with benefits and harms that people value differently (i.e. where personal preferences are relevant). Decision aids make the decision explicit (e.g. to be screened or not), describe why the choice exists, provide information about options and their outcomes, and are intended to help
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people consider the options from their own perspective. Randomised controlled trials in a variety of settings demonstrate that decision aids improve knowledge of risk and accuracy of risk perceptions, stimulate people to take a more active decision making role, and can improve congruence between personal values and the chosen option, usually without lengthening consultations. Decision aids can also assist informed decision making about screening participation.

Uncertainties about the level of benefit and harm can make it difficult to determine what specific information to provide in decision support materials about cancer screening. Selecting the most appropriate data involves considering both the methodological quality of the evidence and its relevance to the local context. For transparency and to reduce bias, decision aids (or supplementary documents that describe their development) should cite the studies selected. When presenting quantitative information (e.g. frequencies of benefits and harms), risk communication research supports using either percentages (e.g. x%) or frequency formats (e.g. x in 1000) and specifying the denominator. Pictographs or icon arrays (e.g. as used by Smith et al. and Hersch et al.) improve comprehension, including among people with low literacy and numeracy. Given that there is rarely any single ‘right’ number for a specific outcome probability, decision aids should also try to communicate the scientific uncertainty in some way (e.g. as described by Forbes and Ramirez).

There are some concerns that encouraging laypeople to take a more active role in healthcare decisions might exacerbate inequalities. To examine this, a systematic review evaluated the impact of shared decision making interventions on health inequalities, including nine studies about cancer screening. The review concluded that such interventions significantly improve outcomes for disadvantaged groups, which suggests that a shared decision making approach for cancer screening is likely to be broadly beneficial, including for vulnerable populations. For example, an Australian trial of a bowel cancer screening decision aid for adults with low education achieved an 85% consent rate in a socio-economically disadvantaged population, with 93% of those randomised completing the trial. The decision aid increased the proportion of people making an informed choice by 22% (as per the study definition), providing evidence that informed decision making can be accessible to the wider community.

Communicating about overdiagnosis

One particularly contentious issue in some cancer screening contexts is overdiagnosis (or overdetection). A prominent concern in prostate, thyroid and breast cancer (with evidence emerging about lung cancer), overdiagnosis occurs when tumours are detected that would not become clinically apparent or cause death if left undetected and untreated. There is currently no way to reliably determine whether a given cancer will progress or remain indolent, so overdiagnosis typically leads to overtreatment, even among screen-detected cases that are recognised as low risk (e.g. low-grade prostate cancer).

Much of the shift in perspective towards promoting informed choice over increased uptake has been due to increasing concern about overdiagnosis, especially in breast cancer screening. Public awareness about overdiagnosis of slow-growing cancers is limited. An online survey of people aged 50–69 in the US who had been invited to participate in cancer screening by their doctor found that less than 10% of patients had been informed about the risk of overdiagnosis. Similarly, in an Australian telephone survey, 10% of women and 18%
of men who had been screened for breast and prostate cancer, respectively, said they were told about the risk of overdiagnosis.16

Communicating the concept of overdiagnosis is difficult, partly because it is undetectable on an individual level, which means that understanding it is not intuitive. A randomised controlled trial in Australia aimed to improve women’s understanding about overdiagnosis in breast cancer screening and assessed the effects of informing them.16 The decision aid explaining overdiagnosis increased knowledge and enabled more women to make an informed choice (as per the study definition), compared with a control decision aid omitting this information. Being better informed about potential consequences also made women feel slightly less positive about breast cancer screening and somewhat reduced intentions to be screened. This suggests that some women, when better informed, might make different decisions about participation.16 Similarly, a brief explanation of overdiagnosis reduced intentions to undergo thyroid cancer screening among Korean women.20

Communication in context

Crucially, evidence based information such as that presented in a decision aid is just one of many factors shaping healthcare choices. Decisions are influenced by emotions, not only scientific information. People’s sense of risk is often more about feelings than facts, which might limit the impact of new factual information.23 Breast cancer screening, in particular, has been highlighted as a decision context exemplifying the tension between powerful emotions and objective risk assessment.23 This tension makes it challenging to communicate potential harms of screening.

The source of information may shape how individuals interpret new information. Clinicians tend to be respected as credible sources of advice and can influence public perceptions of cancer screening.24 If future information materials encourage more deliberative decision making, clinicians might be called on to help people understand the evidence and guide them in clarifying their preferences. Current evidence suggests that many clinicians perform suboptimally in explaining the downside of screening to patients.25 Clinicians may not yet have the capacity or training to participate usefully in shared decision making about screening, and messages currently conveyed by clinicians may be inconsistent with recently developed screening information.17 An important future goal should therefore be to develop ways to support clinicians in communicating more effectively with patients about cancer screening, and its benefits and harms, including overdiagnosis and other challenging topics.10

Conclusion

When communicating with the public about cancer screening, promoting uptake by emphasising benefits was once considered justified. However, for some types of cancer screening, there is professional contention about whether the benefits of screening clearly outweigh the harms. Arguments have been made that individuals should be supported to make informed decisions about whether to undergo screening, and that they need accessible, high-quality information about screening outcomes to do so.

Despite challenges, research has shown that it is possible to help people become better informed about screening.10,12,16 Based on recent evidence16,20, it appears that most people would still prefer to be screened, even when given new information about risks such as overdiagnosis. Engaging and supporting individuals to construct and enact their own informed preferences in making decisions about their healthcare is vital if patients and citizens are really to be at the heart of the healthcare system.

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Competing interests

None declared

Author contributions

JH was responsible for the design, drafting and editing of the manuscript. BN, AG, JJ and KM were responsible for reviewing and editing the manuscript, and contributed to the design of the manuscript.

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