Women’s views on overdiagnosis in breast cancer screening: a qualitative study

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Abstract

Objective To elicit women’s responses to information about the nature and extent of overdiagnosis in mammography screening (detecting disease that would not present clinically during the woman’s lifetime) and explore how awareness of overdiagnosis might influence attitudes and intentions about screening.

Design Qualitative study using focus groups that included a presentation explaining overdiagnosis, incorporating different published estimates of its rate (1–10%, 30%, 50%) and information on the mortality benefit of screening, with guided group discussions

Setting Sydney, Australia

Participants Fifty women aged 40–79 years with no personal history of breast cancer and with varying levels of education and participation in screening.

Results Prior awareness of breast cancer overdiagnosis was minimal. Women generally reacted with surprise, but most came to understand the issue. Responses to overdiagnosis and the different estimates of its magnitude were diverse. The highest estimate (50%) made some women perceive a need for more careful personal decision making about screening. In contrast, the lower and intermediate estimates (1–10% and 30%) had limited impact on attitudes and intentions, with many women remaining committed to screening. For some women, the information raised concerns, not about whether to screen but whether to treat a screen detected cancer or consider alternative approaches (such as watchful waiting). Information preferences varied: many women considered it important to take overdiagnosis into account and make informed choices about whether to have screening, but many wanted to be encouraged to be screened.

Conclusions Women from a range of socioeconomic backgrounds could comprehend the issue of overdiagnosis in mammography screening, and they generally valued information about it. Effects on screening intentions may depend heavily on the rate of overdiagnosis. Overdiagnosis will be new and counterintuitive for many people and may influence screening and treatment decisions in unintended ways, underscoring the need for careful communication.

Introduction

Mammography screening is meant to reduce breast cancer mortality through early diagnosis and treatment of disease that would otherwise threaten life. However, in some cases a screening mammogram detects a cancer that would have neither caused death nor presented clinically during the woman’s lifetime,1 meaning she would never have acquired a cancer diagnosis had she not attended screening.2 This is termed overdiagnosis or overdetection. Overdiagnosis and the resulting overtreatment of inconsequential disease are considered the most serious and important harms associated with early detection of cancer through screening,1,3 generating morbidity that adds to the societal burden of cancer in medical, psychological, and economic terms.1 An overdiagnosed person is unlikely to benefit and may be harmed physically and emotionally by diagnostic procedures and treatments. The implications of the diagnosis...
may extend to female relatives, who are consequently classified as being at higher risk. 3,4

The problems of overdiagnosis and overtreatment, both in cancer screening5 and in the broader healthcare context,6 are receiving increasing attention, and an independent review took place in 2012 in the United Kingdom7 to assess whether the national breast screening programme needed to be changed in light of the latest evidence on benefits and harms, including overdiagnosis. Quantifying overdiagnosis in mammography screening remains a challenge—all estimates are subject to bias, and findings vary over a wide range amid continuing methodological debate. 8-10 Recent studies have suggested that the overdiagnosed proportion of breast cancers (invasive and non-invasive) diagnosed in women invited for screening is likely to be around 19% to 35%,8 12-15 whereas other estimates have been substantially lower16,17 or higher,18 19

Debate persists regarding the appropriateness of informing women about the risk of overdiagnosis before they decide to screen.20 Many argue that women considering screening should be aware of benefits and harms, including overdiagnosis, so they can make informed decisions about whether to participate.1 14 21-25 However, others are wary of the possibility that explicit information on the downsides of screening might dissuade people from participating.20 26 Materials distributed by breast screening programmes worldwide generally do not explain overdiagnosis, 3 5 27-29 and consequently the public is largely unaware of the issue.30 32 Two mammography decision aids (evidence based information resources to support decision making) have briefly introduced the concept of overdiagnosis via an optional hyperlink33 or appendix.34 However, the information was probably not sufficient to convey the full implications of overdiagnosis and overtreatment, including the number of women affected. In a New Zealand study,35 a citizens’ deliberation method was used to educate a panel of 11 women about mammography benefits and harms, of whom 10 then changed their minds to vote against government provision of screening for women aged 40–49. Although overdiagnosis was referred to repeatedly during group deliberations about screening policy,36 it remains unclear how women respond to overdiagnosis on an individual level and how they might integrate this information with their personal attitudes in deciding about their own screening participation. There is also a lack of evidence about how sensitive preferences might be to the varying estimates of overdiagnosis.3

Mammography screening involves large numbers of women worldwide, with participants annually numbering over 840 000 in Australia18 and 2.1 million in the UK.37 The Australian mammography programme is similar to those in other developed countries: women are invited to a screening service whose leaflets emphasise the benefit without mentioning overdiagnosis (such as www.bsnsw.org.au/publications/early-detection-is-vital). To develop optimal strategies for communicating about overdiagnosis to women considering screening, better understanding of women’s perspectives on the issue is needed,38 including how views may vary according to the magnitude of overdiagnosis. We conducted a qualitative study to elicit women’s responses to information about overdiagnosis in mammography screening. We employed qualitative methods, which have been used successfully in previous research into women’s views of screening, their reasoning about participation, and the ideas shaping these decisions.39 40 A qualitative approach was well suited to the goals of this study which aimed to generate insights into women’s conceptual understanding and interpretation of information on overdiagnosis; to explore women’s evaluation of overdiagnosis including whether and how different estimates influenced screening attitudes and intentions; and to investigate views about provision of such information to the community.

Method

Design

We used focus groups to explore how women integrated information about overdiagnosis (a topic likely to be new to them) with their existing understanding of breast screening (or “schema”)14 and how they might use the information when making their personal screening decisions. Choosing a qualitative design involving face to face contact with participants ensured we could be confident that we communicated the information effectively. Women were able to ask questions to seek clarification, meaning that we could check their understanding and identify sources of confusion as part of the research process. Collecting data through focus groups rather than individual interviews gave more control of conversation topics to the participants because it allowed participants to bring up and discuss among themselves, in their own language, relevant points beyond those anticipated by the researchers.42 Another useful feature of focus groups for this research was the opportunity for participants to hear each other’s views and exchange ideas, thereby helping to clarify individual understanding and attitudes.43 Furthermore, the presence of group members with contrasting opinions often led participants to elaborate their own point of view in greater depth.44 All study procedures were approved by the University of Sydney Human Research Ethics Committee. Participants gave written consent prior to participation.

Participant selection and recruitment

We recruited a community sample of women for whom making decisions about participating in mammography screening was relevant. Eligible women were aged 40–79 years, spanning the broad age range in which women commonly have screening. In Australia, women aged 50–69 are routinely invited for free biennial mammograms through the national screening programme, while women aged 40–49 or over 70 are eligible for screening on request but not actively invited.

We selected Sydney suburbs that varied in socioeconomic status and approached potential participants by randomly sampling households in these areas using the electronic telephone directory. Recruitment was carried out via telephone by the Hunter Valley Research Foundation (HVRF), an independent non-profit organisation with extensive experience running community surveys and recruiting people into health research. To gain a diverse range of perspectives, we used purposive sampling to ensure inclusion of women with varying levels of education (that is, with and without post-school qualifications) and prior participation in screening (including low and high screening women in all age groups). We excluded women not fluent in English and women who had ever personally been diagnosed with breast cancer. Trained HVRF interviewers used a short series of questions to determine eligibility and compiled a database of women who had consented to contact by the researchers. We mailed study information to these women and then telephoned them to address any questions and invite them to participate.

Of 118 women who were contacted and eligible, 78 (66%) consented to participate. We scheduled focus group sessions to best accommodate women’s availability. Recruitment to new groups continued until the team decided we had achieved
Data capture, coding, and analysis of qualitative data

Focus group discussions were audio recorded and transcribed verbatim. Data were managed using NVivo 9 software. We conducted a thematic analysis, aiming to identify a set of main themes that captured the diverse views and feelings expressed. After the first few focus groups, the group moderators (JH, JJ, and KM) began the analysis by independently documenting and then discussing together the most salient observations from the group discussions. These impressions, together with the research questions that shaped the group discussion guide, formed the basis of the initial coding framework.

Two researchers (JH and JJ) independently reviewed all transcripts, developed codes in an iterative process, and applied these to the data to identify patterns of ideas. Using constant comparison,44,45 we continually looked for similarity and differences in the data and our coding within and across transcripts. By comparing each part of the data with the rest of the data, we established analytical categories and selected key overarching concepts. Throughout this process we met regularly with KM and the research team to discuss the framework with which to interpret the data.

Quantitative measures

We administered brief written questionnaires at the start and end of each focus group, including demographics, preferred involvement in decision making about screening (control preferences scale),46 and items to assess basic background knowledge and understanding of content presented (see appendix 3 on bmj.com for key items from questionnaires 1 and 2).

Follow-up telephone interviews

JH conducted a brief telephone interview with a randomly selected third of the focus group participants in each age category (n=18) at one to four weeks after their session to identify whether views about overdiagnosis had changed and to elicit any responses that women might prefer to express privately. We asked whether the women had any further thoughts or feelings, since their focus group, about the information presented and their screening intentions. As the views expressed during follow-up telephone interviews presented no new data further to the focus group discussions, they were not analysed further.

Results

Sample characteristics

Fifty women participated in eight focus groups. Participants had diverse cultural and education backgrounds (table 1). Each age stratum included women who had participated regularly in screening and others who had not. Three quarters of women in the target 50–69 year old cohort and 60% of older women knew they were eligible for free screening, whereas two thirds of women in their 40s either believed they were too young to access the programme or were unsure. About 80% of respondents desired active involvement in decisions about their breast screening participation.

To illuminate the study findings presented below, we have selected representative participant quotations that illustrate both typical responses and the diversity of views expressed. The examples are labelled by participant number, age, and screening history, and are drawn from women with a variety of education levels.
The concept of overdiagnosis: women’s understanding and reactions

Understanding of the concept of overdiagnosis

Prior awareness of overdiagnosis was limited to only a few women who had heard of it in the context of prostate cancer. The idea of overdiagnosis occurring in breast cancer screening was surprising and challenged women’s beliefs about breast cancer generally being a serious and dangerous disease:

“In your brain, breast cancer—if it isn’t treated—is fatal. So I’ve never ever heard of that before, and I think most women wouldn’t have.” (Participant 26, age 55 years, 3 screens)

“That there are okay cancers, it’s something that we haven’t heard of before. It’s very new to me... here it is... in the statistics, but we don’t hear about it, not even from our doctors.” (Participant 15, age 47, 1 screen).

By the end of the session, most women gained a reasonable understanding of the concept of overdiagnosis, as shown by these examples of women paraphrasing the concept of indolent or “slow growing” cancers that do not threaten life:

“Sometimes it doesn’t matter if you’ve got [breast cancer], it wouldn’t make any difference, you’re still going to live as long.” (Participant 27, age 57, 0 screens)

“It could be dormant for years and years, and you wouldn’t know.” (Participant 49, age 78, 4 screens).

We also assessed participants’ conceptual knowledge about overdiagnosis via two items in the written questionnaire administered at the end of each focus group. As shown in table 2, most of the participants’ written responses indicated understanding of the key point that screening increases breast cancer incidence through the detection of non-threatening cancers. We marked women’s definitions of overdiagnosis strictly according to the following scoring criteria. To score full marks, the response had to indicate that (a) a person has cancer, and (b) it may not cause harm or treatment may not be necessary. The “partially correct” scoring category covered responses that ranged from those indicating some limited understanding (such as “finding a lump that is benign”) to those that were close to achieving full marks (such as “when there is a cancer that is detected, but may have not changed if it was not detected”).

The complete set of written definitions given by participants is provided in appendix 4 on bmj.com.

There were many questions around the methods used to identify overdiagnosis, Women often had trouble grasping how it is even possible to know that overdiagnosis is occurring:

“How do we know that things are being overdetected? How do we know that there are some cancers that move more quickly or become more malignant than others...? Overdiagnosis assumes that these women, who have been overdiagnosed, have a cancer that is not necessarily aggressive... Who determines what’s an aggressive and a non-aggressive cancer?” (Participant 33, age 62, 5 or more screens).

Many women expressed surprise or disbelief at the current limitations we described in distinguishing between cancers that do require treatment and those that may not. There was some prior awareness that cancers are graded and staged after biopsy, and our assertion that this process still cannot accurately predict the future course of every diagnosed cancer (and the need for treatment) conflicted with women’s expectations:

“If you have a lump... don’t they do a biopsy to check whether it is cancerous or not first? Yes and if it is, they would be able to tell you whether they are aggressive or not, right?” (Participant 11, age 46, 2 screens).

Women also struggled with the fact that overdiagnosis cannot be identified at the level of individual patients—so that even if a given case is “slow growing,” there is no way to know for sure that it will remain non-threatening throughout the woman’s lifetime and thus represents an instance of overdiagnosis:

“How do they find out?... Have they studied a slow growing breast cancer to see how long it goes for or how do they know that?” (Participant 24, age 52, 1 screen).

Box 2 summarises the main issues around overdiagnosis that caused confusion or required clarification during the focus groups.

Reactions to learning about overdiagnosis

For some women, learning about overdiagnosis prompted concern about the implications in terms of the undesirable
Box 2: Summary of main points of confusion about overdiagnosis, based on women’s questions

- How do you know that there are cancers that will not progress fast enough to cause the woman any harm in her lifetime?
- Why is it not possible to distinguish these non-threatening cancers from the potentially dangerous ones that require treatment, even after biopsy or surgical removal of the cancer?
- How can overdiagnosis be quantified if we cannot identify the individual overdiagnosed cases?
- Why does overdiagnosis matter?

The impact of overdiagnosis information on decision making

Implications of overdiagnosis for decisions about screening

Exploring how awareness of overdiagnosis affected women’s feelings about screening was one of our key objectives for the focus group discussions. We found that a few participants, especially younger women with limited screening experience, quickly interpreted the possibility of overdiagnosis and overtreatment as a distinctly negative, off-putting factor in the screening “equation”:

- “Turns me off further... I’m just adding the things that we’ve learnt together... and all the doctors not knowing if it is malignant or not, you add them all together and you think, ‘Why bother?’” (Participant 09, age 45, 0 screens)
- “I don’t know, makes me a bit scared now to go for a mammogram.” (Participant 12, age 46, 0 screens)

After explaining and discussing the concept of overdiagnosis, we moved on to its rate, starting with the 30% estimate. Although this figure clearly surprised some women and raised a few momentary doubts, most women found this level of overdiagnosis acceptable and considered it to be outweighed by the possible benefits of early detection. Ultimately, the 30% estimate had limited impact on women’s existing views and screening intentions:

- “I’m really not sure if I would prefer to be screened than not.” (Participant 38, age 70, 5 or more screens)
- “I still prefer early detection, and so I prefer taking the risk of [being] ‘overdetected’ than... risk of dying.” (Participant 15, age 47, 1 screen)

The 1–10% overdiagnosis estimate was generally perceived as negligible and nothing to worry about. It did not seem to be sufficient to influence screening decisions but rather tended to confirm women’s confidence in screening:

- “The 1–10 is like nothing.” (Participant 24, age 52, 1 screen)
- “If it’s in the 10% range, then I will say, ‘Yeah, just go ahead and do it.’” (Participant 11, age 46, 2 screens)

By contrast, the 50% overdiagnosis estimate was seen as extremely high. Considering this rate elicited emotional reactions, some bordering on disbelief, and raised concerns about the potential burden of unnecessary treatment affecting so many women:

- “Fifty percent, that’s a huge number. ... I do voluntary work at [hospital], and [the patients having cancer treatment] look dreadful and they feel dreadful.... I hate to think of 50% of people going through all that awful treatment for no reason.” (Participant 42, age 72, 3 screens)

The figure illustrates a range of ways in which women responded to the 50% overdiagnosis scenario. Some women were prompted to reconsider whether screening was likely to be worthwhile for them personally—that is, taking into account their own perceived risk factors for breast cancer (such as family history, breast feeding, the pill, breast self examination, lifestyle, stress). The discussions suggested that if the rate of overdiagnosis was about 50%, some participants might decline screening altogether. More commonly, women suggested that they might delay screening (particularly younger women) or be less concerned or rigorous in ensuring they attended within the recommended two year interval. Notably, women often presented this as a relatively minor change in their screening intentions. For one woman, the 50% overdiagnosis estimate had the effect of undermining her confidence in mammography as...
a screening tool and stimulating a search for other screening modalities. However, some women indicated that the information did not at all affect their feelings that mammography screening was worthwhile. A few declared that they would still participate in screening regardless of an even higher overdetection risk, and even while interpreting 50% as a high rate in itself.

Across the sample, women made a clear distinction between the 30% and 50% estimates. Responses suggested that an overdetection rate of 50% might deter some women from screening at all, and for many others it would at least necessitate more careful thought about whether and when to screen. In other words, the higher rate would affect the decision making process even if it might not change women’s ultimate behaviour:

“If there’s more chance that you would have all the treatment and it was unnecessary, you’d be less likely to have the screening, I would say.” (Participant 30, age 61, 0 screens)

“It does make a difference. I’d probably do it more easily.” (Participant 24, age 52, 1 screen).

We also observed a relationship between previous screening attendance and the extent to which information about overdagnosis seemed to influence perceptions of screening. Regular screeners tended not to alter their positive screening intentions after learning about overdagnosis, whereas women with less screening experience seemed more concerned about the issue.

**Implications of overdagnosis beyond screening decisions**

Women valued screening as an opportunity to gain a sense of reassurance that everything was all right. They recognised that, even in light of overdagnosis, they still had good chances of going through screening over many years without having any cancer diagnosed. Therefore, many women did not seem motivated to make a well informed and carefully considered decision about whether to have screening. Instead they chose to defer engaging in thorough deliberation until they were faced with the relatively unlikely event of being diagnosed with cancer:

“I think screening is ... something you go through to have a peace of mind. But as to ... after that, if you are diagnosed, then you have to weigh and look at the various options that you have.” (Participant 11, age 46, 2 screens).

Some women said they would still have screening but felt that, if they were subsequently diagnosed with breast cancer, treatment (or at least adjuvant therapies beyond surgery) may not be worth while if overdagnosis were 50%. Rather than seeing treatment as the immediate logical consequence of a diagnosis, they would want to consider “watchful waiting” or alternative therapies. At the very least, women anticipated having more thorough discussions and desired greater involvement with their doctors to explore all their management options:

“If you were diagnosed you’d really go into what treatment involves ... you’d want to make a really informed decision ... but I probably would still have the breast screens and then make the decision after.” (Participant 26, age 55, 3 screens)

“It’ll still go [for screening]. ... Surgery’s okay too, but ... anything else being needed, any chemicals to be put through my body ... then that’s where I stop and think and look at the statistics.” (Participant 15, age 47, 1 screen)

“It doesn’t turn me off the screening, but I would certainly, probably second-think the treatment. Not that I wouldn’t get it done, I probably may take a bit longer to think about it.” (Participant 16, age 47, 0 screens).

There were suggestions that awareness of overdagnosis (that is, knowing that not all cancers progress and need treatment) might help newly diagnosed women to feel less afraid and more hopeful about their prognosis. Some participants felt that this knowledge might also give such women more of a sense of control and reduce the pressure to follow a predetermined disease management pathway.

Importantly too, it was also suggested that awareness of overdagnosis could worsen the experience of having cancer by making women feel less able to cope with the challenges of unpleasant treatment. As such, it was considered perhaps better for the woman concerned to believe the cancer to be life threatening and treatment crucial:

“That 50% is not good information ... because if I was told ... I’ve got cancer, I won’t want to have in the back of my mind, ‘Oh well, it might not be, and I’m going through all this and my family’s all going bananas for nothing.’” (Participant 42, age 72, 3 screens).

**How to communicate overdagnosis information to women**

**Personal preferences regarding screening information**

Some respondents stated that the information about overdetection would not make any difference to their own screening views or behaviour and therefore it did not matter whether they had this information or not:

“For me ... if I’m going to do it, I don’t care about [the overdetection rate].” (Participant 35, age 69, 5 or more screens).

Others felt that it was good to know about overdagnosis in order to be able to make informed decisions about screening. This included women who felt that their own screening decision would change and those who thought that it would not:

“I really think it’s good to know the downside of overdetection. ... That information [is] not discouraging me to ... screen, but it just made me feel I have knowledge. ... I feel it’s really good.” (Participant 18, age 49, 0 screens).

Some participants thought that making decisions was more difficult when there was a lot of information to go through, or they lacked confidence in their ability to understand numerical information (such as the chances of experiencing benefit or harm from screening). These women often preferred to trust recommendations from their doctor or the government:

“It’s confusing, especially if you’re not mathematical, like me.” (Participant 34, age 67, 5 or more screens)

“I just want [my doctor] to say, ‘You should go, you must go ... this is where you go.’” (Participant 12, age 46, 0 screens).

Many respondents were in favour of being encouraged to screen, whereas some preferred to be given balanced information and left to make an individual choice. Some women simultaneously endorsed both of these approaches:

“I’d like to be encouraged to have screening. I’d also like to be armed with all the information.” (Participant 33, age 62, 5 or more screens).
Views on providing information to the community

Many women favoured full, balanced information as the most ethical approach to communicating about screening. Some were surprised that they had not heard about overdiagnosis before and argued that up-to-date information about it ought to be more widely available. Again, this included women who remained positive about undergoing screening:

“...I think if you’ve got the information, it should be provided.”
(Participant 16, age 47, 0 screens)

“I would actually say that women are entitled to know. ... How you act upon it is probably neither here nor there. It’s not a bad thing, perhaps, if that reminder [letter every two years] comes with this information attached to it—or any new information, for that matter, that might arise.”
(Participant 17, age 48, 1 screen).

On the other hand, women recognised the challenge posed by trying to communicate the complex information from the focus group presentation in an alternative format that would be accessible and more practical. Furthermore, some women felt that changing the conventional pro screening message to include more balanced information could unleash widespread public confusion:

“They’re going to confuse the people. ... All this hard work they’ve been doing for years, encouraging people to have mammograms ... now you are saying today, ‘If you want to you can do it, but see it’s your choice, see the percentages and you make your own decision.’... Years and years saying, drumming in, ‘You should have this every two years,’ ... and now you come out and say, ‘Look, they are the percentages, think about it if you’re going to go and do it.’ You are confusing people.”
(Participant 35, age 69, 5 or more screens).

Some women who said they personally were not put off by the overdiagnosis information nonetheless expressed concern that it would dissuade others from screening. In their view, this could lead to more breast cancer deaths and compromise the gains that screening had achieved for women’s health. A couple of participants suggested providing overdiagnosis information to women only if and when they were diagnosed with breast cancer rather than before screening. Some participants took an intermediate position, suggesting that the amount of information women required before screening would vary according to personal preferences and that this variation should be accommodated somehow:

“I don’t think a vast wad of information should be pushed on everyone willy-nilly, but people should be made aware that the information is there if they would like it ... this might help you make a decision. But I’d want to be made aware that there was information that I can and should look at, should I choose to.”
(Participant 29, age 59, 5 or more screens).

Discussion

Principal findings

In this community sample of women eligible for mammography screening, awareness of overdiagnosis of breast cancer was minimal, consistent with earlier quantitative survey research.31 32 However, most participants were able to understand the issue after the explanation we presented and the opportunity to have their questions addressed. Although virtually everyone reacted to the information about overdiagnosis with surprise, effects on women’s screening attitudes were diverse. Some women expressed a commitment to screening that was unaffected by learning about the possibility of overdiagnosis, while others wanted to take this information into account and give further thought to their screening choices. The screening intentions expressed by these women depended heavily on the magnitude of overdiagnosis, with the highest estimate (50%) having a distinctly greater impact than both the lower and intermediate estimates (1–10% and 30%). Most participants felt that this information was important and should be available to enable women to make informed choices, although many also wanted to be encouraged to have screening.

Importantly, for some participants the information on overdiagnosis caused uncertainty not about whether to have screening but rather whether to have standard treatment for a screen detected cancer or consider alternative approaches (such as watchful waiting or unproven therapies). There was also the suggestion that, for women who were treated for breast cancer, being aware of the possibility of overdiagnosis might make it more difficult to endure unpleasant side effects because of uncertainty about whether treatment was really needed. Learning about overdiagnosis in mammography raised interest in seeking alternative forms of breast screening (such as ultrasound scanning) either instead of or in addition to mammography. Several participants expressed concern about other women inappropriately being deterred from screening altogether because of overdiagnosis. Moreover, some were suspicious about an agenda behind overdiagnosis research—including our own study—fearing a government plan to cut funding for mammography screening.

Strengths and limitations

The strategy for communicating about breast cancer screening in the UK is currently under review. By examining lay women’s perspectives on overdiagnosis, this study makes an important and timely contribution to the debate about the possible consequences and value of providing overdiagnosis information to women invited for screening.

The focus group presentation was carefully developed to be both accurate (that is, evidence based data) and understandable (such as innovative illustrations of the concepts), and was expert reviewed and extensively pilot tested. We presented the cumulative probabilities of different screening outcomes using natural frequencies, in line with evidence based recommendations.33 We explored responses to a diverse range of overdiagnosis estimates, reflecting the full body of existing literature, because the amount of overdiagnosis is contested and all estimation methods are subject to bias.3 The qualitative design is a strength of this exploratory study because it allowed unanticipated findings to emerge (such as the impact on decision making about accepting treatment for detected cancer).

Encouraging women to ask questions also provided new insights into points of confusion.

We tried to minimise volunteer bias by recruiting directly from the general community, and our provision of financial compensation helped us to achieve a broad sample including women from diverse socioeconomic backgrounds who varied in age, education, prior screening, and ethnicity. However, as the sample was not designed to be statistically representative, we cannot conclude that our findings reflect general population views. We stressed our neutrality at the start of each session to mitigate against “socially desirable” comments in favour of screening.

To avoid overloading women with information, we did not talk about the scientific uncertainty surrounding estimates of the mortality benefit, nor did we discuss in detail the chances of...
other relevant screening outcomes (such as false positives and false negatives). Discussing these additional points would have provided a more comprehensive picture of factors relevant to screening and the decision making process, and might have influenced the study results. However, further increasing the complexity of the information provided would have limited the study’s capacity to address the key research questions concerning overdiagnosis. It is possible that our choice of presenting the benefit information first may have created a “framing” effect resulting in more positive attitudes, but finding positive attitudes to cancer screening is consistent with previous literature.31 32 34

In comparison with earlier attempts to introduce the concept of breast cancer overdiagnosis to women,31 33 34 this study involved a much more thorough explanation of the issue and its magnitude, albeit in a more resource intensive manner. The purpose of the focus groups was not to imitate a “real life” setting but to allow us to explore the process through which women understood and responded to information about overdiagnosis, thereby providing valuable insight into women’s attitudes. In our diverse sample (including women with low education and non-English speaking backgrounds) most participants’ written definitions of overdetection showed full (60%) or partial (24%) understanding of the concept. However, some participants (several of whom, though reasonably fluent in English, were from other language backgrounds) did not demonstrate a clear understanding of overdiagnosis, which limited our ability to accurately gather their views about it. For such women, an individual counselling and interview session may be a more appropriate way to explain the issues and examine their responses. Importantly, in this study we asked women about their own screening intentions as opposed to their views on national screening policy.25 Earlier qualitative research with women diagnosed through screening found that they considered it important to take into account, when making screening decisions, the possibility of being diagnosed with an ultimately harmless condition.27 Our study showed that this view, expressed by breast cancer patients whose perspective on screening would have been coloured by their personal experience of cancer, was shared by many women in the general invited population.

Implications and future research

This study provides important insights into lay interpretations of information about overdiagnosis, with implications regarding how best to inform people about the issue. Firstly, the magnitude of overdiagnosis matters, with many women indicating that their screening behaviour may change depending on the trade-off between the chances of benefit and harm.30 The wide variation in published estimates (1% to 50%) is therefore problematic. Additionally, the suggestions from some women that overdiagnosis would be relevant to their decisions only if they were actually diagnosed with a screen detected cancer reveal a concerning misconception that a screening mammogram is a separable event from the cascade of investigation and intervention that may be triggered by an abnormal result. This highlights the need to explain clearly to women that once cancer is detected, evidence based treatment is virtually always indicated because potentially threatening cases cannot be differentiated from those representing overdiagnosis. Communicators must also take care not to create a situation in which women think they can avoid overdiagnosis by choosing a different screening method (for which evidence is limited) or by supplementing screening mammography with additional testing, which may increase the potential for overdiagnosis.

Thus the ethical imperative to provide information about overdiagnosis must be balanced with the responsibility to address misconceptions that may lead to problems in clinical practice.

Another important point to consider in developing strategies for communication about overdiagnosis relates to understanding how people interact with new information. A theoretical framework used in the field of education known as schema theory35 highlights that the way people engage with new material is influenced by their “schema” (existing topic knowledge and expectations). In the case of mammography screening, attitudes tend to be strongly favourable, which is partly a function of the historical and cultural context. Women’s attitudes will have been shaped through the information presented to them directly by screening services as well as their broader experience of many years of widespread public health campaigns focused on promoting the benefits of screening without explaining the harm of overdiagnosis.27 29 32 The very existence of overdiagnosis runs contrary to prevailing notions of cancer and screening. Therefore it is reasonable to expect that new messages about overdiagnosis will not necessarily be immediately understood by everyone and may provoke confusion or scepticism, as we found to some extent in our study. Rather than wishing to be informed in detail about the range of possible outcomes of screening, some participants expressed a preference to receive more basic information together with encouragement to have screening. Such preferences are consistent with the “consider an offer” approach,36 in which screening is offered with summarised key information and people are assisted to evaluate whether taking up the offer is right for them. At the same time, people can be encouraged to consider whether they want to access more detailed information that may help them make a choice.21 Public views and information needs may change over time as more balanced screening information and increased awareness of harms contribute to changing the schema of expectations people have about screening. Moreover, the diversity we found in women’s responses to overdiagnosis supports the need for information providers to tackle the challenge of developing ways to deliver screening information flexibly rather than assuming that women’s needs are homogenous.

Given that doctors’ recommendations are likely to influence screening behaviour,32 one avenue for future research may be to examine the perspectives of relevant clinicians, both in terms of their views on overdiagnosis and their responses to women’s attitudes as reported here. However, in countries such as Australia and the UK women are invited directly by the screening service and attend without necessarily having any discussion with a healthcare provider. Therefore, priorities for future research are to investigate whether a meaningful explanation of overdiagnosis can be condensed into a format that could be distributed with screening invitations, and to assess the impact of such information on screening intentions and participation. Also critical is clarification of the magnitude of overdiagnosis, though it is evident that such data cannot necessarily resolve whether screening benefits outweigh harms for any individual as this ultimately requires a value judgement.4 46 50 The question remains, how do we best inform women and support them to make informed decisions about mammography screening using the available evidence.21

Conclusions

Overdiagnosis was unfamiliar to a diverse sample of women eligible for mammography screening in Australia. None the less, we found it was possible to explain the issue in such a way that most participants understood the concept and its
implications. Women’s screening preferences were sensitive to the estimated level of overdiagnosis, underscoring the need for urgent efforts to clarify this issue. Critically, women differed in the way they evaluated the relevance of overdiagnosis in making their personal decisions about screening. Some felt that the new information was unlikely to change their existing attitudes and intentions; others considered it important to take the risk of overdiagnosis into account, particularly at higher estimates. Many women expressed a preference for this information to be more widely available. However, their responses to being informed about overdiagnosis also revealed a range of potentially unwanted consequences that communicators must address. Finally, the diversity of views elicited in this study highlights the inappropriateness of making screening policy and decision-making based on speculative assumptions about what women want and think regarding overdiagnosis. Instead, it is vital that research efforts continue to better understand women’s values about overdiagnosis, to clarify its influence on their behaviour, and to determine the optimal methods to support informed choice in mammography screening.

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What is already known on this topic

Mammography screening for breast cancer carries a risk of overdiagnosis (detection of disease that would not present clinically within the woman’s lifetime), but estimates of the magnitude of this risk vary widely. Public awareness about breast cancer overdiagnosis is low, and there is little evidence about how women respond to the concept and extent of overdiagnosis and how they might use this information in making personal choices about screening.

What this study adds

Evaluations of the relevance of overdiagnosis depended heavily on the different estimates of magnitude: the 50% estimate generated substantial concern, whereas the 1–10% and 30% estimates were considered more acceptable levels of risk. Women’s responses were diverse: some indicated that they would take the risk of overdiagnosis into account when making screening decisions, whereas others felt that their attitudes and intentions to screen were unaffected by the new information. Learning about overdiagnosis raised interest in alternative screening methods as well as concern about whether and how to treat screen detected cancers; hence great care is needed in communicating this information as it may influence screening and treatment decisions in unintended ways.

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### Table 1 | Characteristics of women in focus groups about mammography screening and overdiagnosis (n=50)

| Characteristics                                      | No (%) of women |
|------------------------------------------------------|-----------------|
| Age (years):                                          |                 |
| 40–49                                                 | 19 (38)         |
| 50–69                                                 | 16 (32)         |
| 70–79                                                 | 15 (30)         |
| Mammography screening history*:                       |                 |
| Screened at least once                                | 31 (62)         |
| Never screened (age <50 years)                       | 12 (24)         |
| Never screened (age >50 years)                       | 7 (14)          |
| Education†                                            |                 |
| No formal qualifications                              | 4 (8)           |
| Intermediate school certificate                       | 11 (22)         |
| Higher school certificate                             | 9 (18)          |
| Diploma or trade certificate                          | 12 (24)         |
| University degree                                     | 13 (26)         |
| Country of birth:                                    |                 |
| Australia                                             | 27 (54)         |
| England or Scotland                                   | 4 (8)           |
| China                                                 | 4 (8)           |
| Other                                                 | 15 (30)         |
| Marital status†                                       |                 |
| Married or living with partner                        | 36 (72)         |
| Single, separated, or widowed                         | 13 (26)         |
| No of children†                                       |                 |
| 0                                                     | 11 (22)         |
| 1                                                     | 7 (14)          |
| 2–6                                                   | 31 (62)         |
| Employment†                                           |                 |
| Retired                                               | 15 (30)         |
| Working full time                                     | 14 (28)         |
| Working part time                                     | 13 (26)         |
| Not in paid work                                      | 7 (14)          |
| Preferred role in screening decision†                 |                 |
| I decide alone                                        | 13 (26)         |
| I decide after consulting doctor                      | 12 (24)         |
| Doctor and I share equally in deciding                | 14 (28)         |
| Doctor decides after consulting or alone              | 8 (16)          |

*Although the Australian national mammography screening programme is available to women over 40 years old with no upper age limit (so all study participants were eligible), the programme actively recruits women for biennial screening between the ages of 50 and 69 in accordance with current recommendations.

†Missing data for between one and three participants.
Table 2 | Participants’ responses to written knowledge items completed at the end of focus group session about mammography screening and overdiagnosis (n=50)

| Item                                                                 | No (%) of women |
|----------------------------------------------------------------------|-----------------|
| Who is more likely to be diagnosed with breast cancer? (multiple choice): |                 |
| Women who have screening mammograms (correct answer)                 | 35 (70)         |
| Women who do not have screening mammograms                           | 10 (20)         |
| Unsure                                                               | 5 (10)          |
| What is meant by the term “overdetection”? (free response question): |                 |
| Full marks                                                           | 30 (60)         |
| Partially correct                                                    | 12 (24)         |
| Incorrect                                                            | 8 (16)          |
## Figure

| Response                                      | Example                                                                 | Participant Details     |
|-----------------------------------------------|------------------------------------------------------------------------|-------------------------|
| May decline breast screening entirely         | "I probably think, 'Oh yeah, 50% is a high number, and I won't do it!'" | Participant 01, age 40, 1 screen |
| Seek other forms of breast cancer screening   | "I may think about trying another sort of way of testing or checking, because I think, 50%, it's really unreliable" | Participant 18, age 49, 0 screens |
| Reconsider in light of personal risk factors  | "You'd have to look at your general health, your history, ... if I knew I had a good healthy lifestyle all along, I'd be inclined to say, 'No, I wouldn't do it!'" | Participant 22, age 50, 0 screens |
| Delay screening or rescreening               | "It may delay it a little bit more but probably not too much"          | Participant 13, age 47, 0 screens |
| Views unchanged - would still screen         | "Doesn't change anything for me. It wouldn't matter how high it was ... I'd rather be safe than sorry. I think any death is one death too many" | Participant 34, age 67, 5 or more screens |

Women's responses to a rate of 50% overdiagnosis for mammography screening