Psychological consequences of false-positive screening mammograms in the UK

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Abstract

Objectives To identify the psychological effects of false-positive screening mammograms in the UK.

Methods Systematic review of all controlled studies and qualitative studies of women with a false-positive screening mammogram. The control group participants had normal mammograms. All psychological outcomes including returning for routine screening were permitted. All studies had a narrative synthesis.

Results The searches returned seven includable studies (7/4423). Heterogeneity was such that meta-analysis was not possible. Studies using disease-specific measures found that, compared to normal results, there could be enduring psychological distress that lasted up to 3 years; the level of distress was related to the degree of invasiveness of the assessment. At 3 years the relative risks were, further mammography, 1.28 (95% CI 0.82 to 2.00), fine needle aspiration 1.80 (95% CI 1.17 to 2.77), biopsy 2.07 (95% CI 1.22 to 3.52) and early recall 1.82 (95% CI 1.22 to 2.72). Studies that used generic measures of anxiety and depression found no such impact up to 3 months after screening. Evidence suggests that women with false-positive mammograms have an increased likelihood of failing to reattend for routine screening, relative risk 0.97 (95% CI 0.96 to 0.98) compared with women with normal mammograms.

Conclusions Having a false-positive screening mammogram can cause breast cancer-specific distress for up to 3 years. The degree of distress is related to the invasiveness of the assessment. Women with false-positive mammograms are less likely to return for routine assessment than those with normal ones.

Introduction

The benefits and harms arising from mammography screening are a matter of national debate in the UK.1 2 The number of lives saved, amount of over diagnosis and degree of distress caused by ‘false alarms’ are hotly contested.3 4 5 This debate has led to a review of UK breast cancer screening services currently being undertaken by Professor Sir Michael Richards.

The negative psychological impact of false-positive screening results has been documented in the fields of prenatal and cervical cancer screening.6 7 Their impact on the medium to long-term psychological well-being and behaviour of women who receive false-positive results from routine mammography has been less well researched and synthesised.

Systematic reviews and meta-analyses in Europe and America have found conflicting evidence about the psychological impact from receiving a false-positive mammogram and also on future attendance at routine screening.8 9 10–14 Most studies showed a negative impact from receiving a false-positive mammogram on measures of, well-being, depression and anxiety compared to women with normal screening results. The exception to this was the meta-analysis of generic psychological measures by Salz et al10 which showed that only anxiety was positively correlated with having a false-positive mammogram (0.03 (95% CI 0.00 to 0.07)). The evidence varied concerning whether psychological distress had a short-term (<1 month after assessment) or long-term impact. There was some evidence that the degree of impact varied with the severity of the reassessment test; with women undergoing biopsy showing greater psychological distress than those with a repeat mammogram.11

The results for the impact of receiving a false-positive mammogram on returning for the next routine screening mammogram give a more complex picture. Armstrong et al11 found there was no statistically significant difference between groups in the likelihood of returning for routine breast screening, although it is not clear whether the studies were reporting actual attendance or intention to attend. This may be important as Bankhead et al12 found that women were more likely to say that they had an intention to attend their next routine mammogram than actually do so. Other studies showed a variation in the effect of a false-positive mammogram on returning for screening according to location; with European women unaffected in this domain, Canadian women less likely to return and women from the USA more likely to return for routine mammography.7

These systematic reviews had a wider geographical scope than the UK and included studies of the short-term impact of a false-positive mammogram (less than 1 month from assessment). Many of the included studies were based on programmes with a different approach and periodicity to that of the UK where mammography screening is national, free, opt-out and uses double screening of mammograms. Additionally the UK service runs on a 3 years cycle of invitations to women aged 47–73. In particular, the US system differs as it comes from a mixture of public and private providers and is insurance-based, opt-in, uses single screening, which produces a higher number of false-positives and has until recently recommended screening annually from the age of 40. Most European countries offer screening for the 50–69 age group but generally every 2 years, and are provided by a mixture of public and private organisations, which may or may not be publically financed.

Therefore, the differences in the provision of screening services indicate that the psychological effects of this experience and the impact on returning for screening.
remains unclear in the particular health service and cultural context of the UK.

To address this knowledge gap we conducted a systematic review of studies in the UK population of the medium to long-term psychological consequences of experiencing a false-positive screening mammogram and whether these affect future attendance at mammography screening. In addition qualitative studies, an important underpinning to the understanding of psychological consequences, did not appear to have been searched for in previous reviews, so we particularly targeted these in our study.

The definition of a false-positive mammogram used in this study is that given by the WHO: ‘an abnormal mammogram (one requiring further assessment) in a woman ultimately found to have no evidence of cancer’ and the American College of Radiology categories of mammographs considered as abnormal are detailed in http://www.imaginis.com/mammography/mammogram-interpretation-categories-and-the-acr-bi-rads.

Methods
This systematic review was carried out following the principles published by the NHS Centre for Reviews and Dissemination. The study protocol can be found in appendix 1.

Inclusion criteria
Studies were included if based in the UK, populated by women whose experience met the above definition of a false-positive screening mammogram, the comparator group were those with a normal screening mammogram, the outcomes were psychological, behavioural or those from qualitative studies and follow-up was at least 1 month from the ‘all clear’. All controlled studies and qualitative designs were included. Case studies were excluded.

Search strategy
The search strategy consisted of searching of electronic bibliographic databases, internet searches, scrutiny of references of included studies and contacting experts in the field.

The following electronic databases were searched in December 2010: Medline, Medline in Process and other Non-Indexed Citations, EMBASE, HMIC, Cochrane Central, Cochrane CDSR, CRD Dare, CRD HTA, Cochrane Methodology, Web of Science, Psychinfo, Cinahl, Sociological Abstracts, the International Bibliography of the Social Sciences and Zetoc. Ongoing trials were searched for at: UKCRN, Controlled Trials.com, Clinical Trials.gov, ICTRP (WHO International Clinical Trials Registry Platform), UK Database of Uncertainties about the Effects of Treatments (DUETs), a filter was applied to capture qualitative research as well as quantitative designs. Further searches for more qualitative and grey literature were run in January 2011 on the following databases: Medline in Process and other Non-Indexed Citations, Embase Classic and Embase, British Nursing Index and Archive, Social Policy and Practice, Cinahl plus, Cochrane Library, HMIC, PsycINFO, Assia, Sociological Abstracts, Web of Science, CRD and IBSS. All searches were run from inception to the then present date. Bibliographies of included studies were searched for further relevant studies. An update search was carried out on 26 November 2011. The MEDLINE search strategy is available in appendix 2.

Papers were selected for review from the titles and abstracts generated by the search strategy. This was done independently by two reviewers (MB and TP); discrepancies were resolved by discussion. Retrieved papers were again reviewed and selected against the inclusion criteria by the same independent process. Data were extracted from included studies by one reviewer using standardised data extraction forms and checked by another reviewer. Attempts were made to contact authors to provide missing information. Data were gathered on the design, participants, methods, outcomes, baseline characteristics and results of the studies.

Quality assessment
Studies were assessed for internal validity according to criteria suggested by the updated NHS CRD Report No. 4, according to study type. Randomised controlled trials (RCTs) were appraised with the CONSORT statement and observational studies with STROBE guidelines. External validity was assessed according to the applicability of findings to a relevant patient group and service setting.

Analysis
Analysis was carried out using StatSev12 software. The principle summary measures were relative risks with 95% CI. All study designs had a narrative synthesis. Observational studies were considered for possible meta-analysis. Overall, they had considerable amounts of missing information so that it was difficult to judge heterogeneity. Therefore, a meta-analysis was not carried out.

Results
Search results
Our searches retrieved 4423 titles and abstracts after deduplication. When screening was complete we found seven primary studies (nine papers) that met the inclusion criteria. Four of the studies were prospective cohorts; four were retrospective cohorts and one was an RCT of an intervention to improve reattendance.

Studies covered two domains: (a) three studies looked at the psychological impact of false-positive mammograms in the normal risk population and (b) six looked at the impact of this experience on returning for routine mammogram screening (some studies looked at both domains). No studies were found that were either about or that had subgroups of women from different ethnic, socio-economic or other groups within the general screening population. No published qualitative studies were found. A flow chart of the selection process is in appendix 3.

Quality and characteristics
The quality of the research was variable; the RCT was poor quality with few methodological details given.
although some of the observational studies, notably those from the Oxford Primary Care Education Research Group, were reasonably well reported. However, the majority had a number of weaknesses, including a failure to consider the possible effects of bias and confounding on the results and a failure to report participants’ demographic and other characteristics. These quality indicators appear to have been overlooked as in most cases there was no consideration of the limitations of the methods or conduct of the study. There were insufficient studies in each domain to produce a meaningful assessment of publication bias with a funnel plot. Table 1 provides summary characteristics of the included studies by their outcome domain. Appendix 4 shows a summary assessment of quality indicators.

Study results
Psychological impact
The studies of the psychological impact of false-positive mammograms gave conflicting results. When disease specific measures were used that is, the Psychological Consequences Questionnaire, an enduring negative impact was found for those with false-positive results compared to those with normal mammograms that lasted until 35 months from the last assessment. The degree of distress found was related to the level of invasiveness of the method of assessment used; so that at 35 months, women who had a biopsy were more distressed (relative risk (RR) 95% CI 2.07 (1.22 to 3.52)) than women who had fine needle aspiration (RR 95% CI 1.80 (1.17 to 2.77)), and non-significantly; further mammography (RR 95% CI 1.28 (0.82 to 2.00)). Additionally women placed on early recall were also at a greater relative risk of distress (RR 95% CI 1.82 (1.22 to 2.72)). The greatest relative risk of distress was felt at 5 months after assessment and was significant for all assessment procedures (figure 1).

Conversely, when generic measures of general anxiety and depression were used, the Hospital Anxiety and Depression Scale and the General Health Questionnaire, no significant differences were found between the two groups at 6 weeks after assessment and 3 months after screening (figure 2).

Reattendance
The evidence for the impact of having a false-positive mammogram on returning for the next screening round is again conflicting. The forest plot below (figure 3) compares the relative risks of the reattendance studies. The evidence comes from four retrospective observational studies that collected data from registries and other NHS databases. The weight of evidence, in terms of the numbers of participants, is that women with false-positive mammograms are less likely to return for their next round of screening than women with normal mammograms, although the effect is small. The largest study with this finding (N=140 387) had a relative risk of returning of 0.97 (95% CI 0.96 to 0.98), Brett and Austoker came to the same conclusion, 0.92 (95% CI 0.86 to 0.98). Two studies with a combined population of N=7231 found that there was no such association but had wide 95% CIs consistent with both increased and decreased likelihood of return. Evidence from a poor-quality RCT by Meldrum et al (N=3083) suggests that this finding can be reversed if women are given screening invitation letters that are tailored to the outcome of their last screening (RR of returning (95% CI) 1.10 (1.00 to 1.21)).

Discussion
The benefits and harms accruing from breast cancer screening are a matter of current UK concern. Included studies, comparing psychological impact of false-positive mammograms to normal ones, gave conflicting results. When disease specific measures were used an enduring negative impact was found that lasted until 35 months with the degree of distress related to the invasiveness of the assessment. Conversely, when measures of general anxiety and depression were used no significant differences were found between the two groups. However, this could be explicable if we speculate that false-positive mammograms may lead to breast cancer-specific psychological distress, enduring for up to 3 years, but that it is unlikely that general anxiety or depression will occur.

Concerning reattendance, the weight of evidence is that women with false-positive mammograms are less likely to return for subsequent rounds of screening than women with normal mammograms.

No systematic reviews were found that are directly comparable to ours as they all include non-UK studies, may have populations younger than ours, measure outcomes at less than 1 month and have screening programmes based on opt-in or insurance payments. Nevertheless, our results agree with theirs that there can be negative psychological consequences from having a false-positive mammogram and that reattendance can fall. Greater clarity has emerged by removing the effect of variation in the specific nature of different national programmes.

Limitations
The robustness of the findings of this systematic review are limited by the reliability of the poorly reported observational studies. The degree of heterogeneity between these studies is subsequently unknown. This meant that we were unable to pool the data (without potentially reporting spurious relationships) and thus provide an overall estimate of distress and reattendance. Additionally, the evidence in this systematic review is at least 10 years old and may also have been influenced by publication bias.

Our decision to restrict included studies to those in the UK may also be seen as a limitation. However we felt that this was a reasonable approach given the variability in results worldwide, the most obvious explanation for which was likely to be variation in programme. Detailed investigation of qualitative research was also felt to be more appropriately conducted at national level. Also despite the UK-specific nature of this review we believe that generalisable messages remain, as indicated below.
<table>
<thead>
<tr>
<th>Study/author year (funding)</th>
<th>Design</th>
<th>N</th>
<th>Participants</th>
<th>Intervention group</th>
<th>Control group</th>
<th>Outcomes</th>
<th>Length of follow-up</th>
<th>Exclusion criteria</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brett and Austoker, 2001 (Cancer Research Campaign)</td>
<td>Prospective cohort multicentre Psychological impact</td>
<td>505</td>
<td>Women invited for routine screening by mammogram, already participating in the study at 5 months</td>
<td>Routine screening by mammogram with a false-positive result N=375</td>
<td>Routine screening by mammogram with a normal result N=130</td>
<td>PCQ intention to reattend and actual reattendance satisfaction with service ad hoc questionnaire</td>
<td>3 years (35 months) after assessment</td>
<td>Over 65 years, symptomatic referral, in another study, developed cancer</td>
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<tr>
<td>Brett et al, 1998 (Cancer Research Campaign)</td>
<td>Prospective cohort multicentre Psychological impact</td>
<td>284</td>
<td>Women invited for routine screening by mammogram, already participating in the study at 1 month</td>
<td>Routine screening by mammogram with a false-positive result N=163</td>
<td>Routine screening by mammogram with a normal result N=52</td>
<td>PCQ, intention to reattend, ad hoc questionnaire</td>
<td>5 months after assessment</td>
<td>As above</td>
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<tr>
<td>Ong et al, 1997a (Cancer Research Campaign, NHSBSP)</td>
<td>Prospective cohort multicentre Psychological impact</td>
<td>877</td>
<td>Women invited for routine screening by mammogram recalled for assessment</td>
<td>Women placed on early recall (3 years) N=182</td>
<td>Women placed on routine recall after mammography (N=173), further mammography assessment (N=166), FNA (N=109) or biopsy (N=31)</td>
<td>PCQ</td>
<td>1 month after assessment</td>
<td>Not reported</td>
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<td>Bull and Campbell, 1991 (Yorkshire Regional Health Authority)</td>
<td>Prospective cohort Psychological impact</td>
<td>750</td>
<td>Women invited for routine screening by mammogram recalled for assessment</td>
<td>Routine screening by mammogram with a false-positive result N=308</td>
<td>Routine screening by mammogram with a normal result N=420</td>
<td>Ad hoc questionnaire including frequency of breast self-examination HADS</td>
<td>6 weeks after the ‘all clear’</td>
<td>Not reported</td>
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<tr>
<td>Ellman et al, 1989 (DHSS Research Management Division)</td>
<td>Prospective cohort Psychological impact</td>
<td>752</td>
<td>Women invited for routine mammogram screening and those recalled for further assessment and those with symptoms being further investigated</td>
<td>Routine screening by mammogram with a false-positive result N=271</td>
<td>Routine screening by mammogram with a normal result N=295, symptomatic women who did not have cancer N=134, symptomatic or recalled screened women who did have cancer N=38, history of breast cancer with or without symptoms N=14</td>
<td>GHQ-28, ad hoc questionnaire</td>
<td>3 months after clinic attendance</td>
<td>Not reported</td>
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<tr>
<td>Impact on reattendance</td>
<td>Retrospective cohort Reattendance and interval cancer</td>
<td>140387</td>
<td>Women 49–63 years invited for routine breast screening by mammography</td>
<td>Routine screening by mammogram with a false-positive result N=4 792</td>
<td>Routine screening by mammogram with a normal result N=108 617</td>
<td>Subsequent attendance at routine screening after a false-positive result and rate of interval cancer—from records</td>
<td>3 years</td>
<td>Women who were older than 63 years at follow-up</td>
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Table 1  Continued

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<td>O’Sullivan et al, 2001 (Cancer Research Campaign)</td>
<td>Retrospective cohort <em>Reattendance</em></td>
<td>5649</td>
<td>Women invited for mammography screening for the second or more time</td>
<td>Routine screening by mammogram with a false-positive result N=248</td>
<td>Routine screening by mammogram with a normal result N=5401</td>
<td>Subsequent attendance at routine screening after a false-positive result— from records</td>
<td>Unclear, probably from 1989 to 1997</td>
<td>Women invited for the first time and women who had been previously invited but had never attended</td>
<td>Effects of a false-positive result on reattendance for those on early recall and routine recall</td>
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<td>Brett and Austoker, 2001 (Cancer Research Campaign)</td>
<td>As above in psychological impact</td>
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<td>Meldrum, 1994 (Scottish Office Home and Health Department)</td>
<td>RCT-nested telephone interview study</td>
<td>3083</td>
<td>All women invited for second round routine mammography screening (50–65 years)</td>
<td>Tailored invitation with a false-positive result N=115 and with a normal result N=800</td>
<td>Standard invitation with a false-positive result N=112 and with a normal result N=791</td>
<td>Subsequent attendance at routine screening and effect of a tailored invitation on subgroups</td>
<td>Not reported</td>
<td>Women with breast cancer and those whose screening history was not available</td>
<td>Trial comparing the effect of a tailored invitation on second round screening attendance with a standard invitation</td>
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<tr>
<td>Orton, 1991 (funding not reported)</td>
<td>Retrospective cohort <em>Reattendance</em></td>
<td>1582</td>
<td>Women, aged 45–64, invited to attend for second round screening by mammography</td>
<td>Routine screening by mammogram with a false-positive result N=50</td>
<td>Routine screening by mammogram with a normal result N=1532</td>
<td>Reattendance acceptability of screening</td>
<td>NA</td>
<td>Not reported</td>
<td>Data are not available for the acceptability of screening for false-positive participants</td>
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FNA, fine needle aspiration; HADS, Hospital Anxiety and Depression Scale; NA, not applicable; PCQ, Psychological Consequences Questionnaire; RCT, randomized controlled trial.
Implications for policy and future research

Policy makers, particularly in the UK should consider the impact of false-positive mammograms when planning breast screening services. Measures need to be taken to reduce the distress caused; however, the evidence base for such measures is lacking. Internationally the need to take account of the context of the evaluations is illustrated and absence of supporting qualitative research to understand the underlying nature of psychological consequences in the UK is likely to have parallels in other countries.

Therefore, further research is needed to increase and update our understanding of the harms of breast cancer screening. In particular a qualitative interview study would further our appreciation of what this experience means to women, and principally shed light on how the subtleties of difference in response to assessment procedures relate to anxiety and probability of reattendance. Consequently, the authors are currently conducting such an interview study. Well-designed observational studies are also needed, that use disease specific and generic outcome measures in order to determine the degree and kind of negative psychological outcomes. Studies should include women from different ethnic and socio-economic groups and routinely collect demographic information so that future systematic reviews may be able to judge whether the pooling of data is legitimate.

Conclusions

We conclude that the experience of having a false-positive screening mammogram can cause breast cancer-specific psychological distress that may endure for 3 years. However, it is less likely that there will be pathological general anxiety and depression. The distress caused by a false-positive mammogram may be
sufficient to deter some women from attending their next breast cancer screening appointment.

▸ Additional data is published online only. To view this file please visit the journal online (http://ebm.bmj.com).

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**Disclaimer** The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the Department of Health.
Competing interests None.

Correction notice This article has been corrected since it was published Online First. Reference 33 was a repeat of reference 27 and so has been deleted from the article.

References