Is the tide turning against breast screening?

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See related research by Nederend et al., http://breast-cancer-research.com/content/14/1/R10

Abstract

Herein I argue that mammographic screening has not delivered on its fundamental premise: to reduce the incidence of advanced breast cancer. Indeed, achieving this goal is required if screening is to reduce breast cancer mortality or mastectomy use. Rather, screening has caused substantial increases in the incidence of in situ and early invasive cancers. Moreover, evidence indicates that these screen-detected cancers are unlikely to be cases that were ‘caught early’, but instead represent women who would not have been diagnosed in the absence of screening and who, as a result, have received harmful, unnecessary treatment. If true, these observations raise the specter that screening creates breast cancer patients and that this practice carries little or no benefit.

Introduction

The rationale for breast screening with mammography is deceptively simple: catch the cancer early and reduce mortality from the disease and the need for mastectomies. But breast cancer is a complex disease, and complex problems rarely have simple solutions.

A key question is if screening can prevent metastasis, as this would reduce breast cancer fatalities. Another is if the screen-detected tumours that previously required a mastectomy are now small enough to allow breast conserving surgery. The fundamental premise for both objectives is that screening must reduce the incidence of advanced breast cancer. Whether such a reduction occurs in a long-running, organized, population-based screening programme is what Joost Nederend and colleagues explore in their study from The Netherlands [1].

Current context

Other researchers have questioned whether breast screening reduce the number of advanced cancers [2,3].

A claimed reduction in breast cancer mortality [4-6] as well as a reduction in the use of mastectomies [7,8] have also been called into doubt in studies of population-based breast screening. In addition, the detection of cancers that would otherwise not have developed into clinical, symptomatic disease (overdiagnosis) is now recognised as an important harm, also for invasive breast cancer [9,10].

A recent systematic review of incidence trends in seven countries with at least seven years of screening [2] found that breast screening has not fulfilled its promise of fewer advanced breast cancers. It included The Netherlands, but not data from before organised screening was introduced in the late 1980s. Including data from 1980 to 2008 is a strength of the new study, as it allows reliable estimates of both pre- and post-screening incidence trends of advanced breast cancer. If the background incidence was increasing prior to screening, but stable during the screening period, screening may have prevented a further increase in advanced breast cancer. This seems not to have happened, either in The Netherlands or in other countries where data from the pre-screening period are available - for example, the United States and Norway [2]. A new study from Norway used the stepwise introduction of breast screening to compare the occurrence of stage 3 and 4 tumours in screened and non-screened areas, with data available also from the pre-screening era. The availability of a contemporary ‘control group’ circumvents the usual problems related to using stage distribution over time due to changing definitions and diagnostic methods. In Norway, there was a reduction in the occurrence of stage 3 and 4 tumours in screened areas in recent years compared to the pre-screening years, but an identical reduction was seen in the non-screened areas [11]. The authors also found evidence of substantial overdiagnosis of invasive breast cancers in Norway and ended their paper by raising the question of whether mammography screening programmes should exist.

Some studies have concluded that a reduction in advanced breast cancer has occurred, also in The Netherlands [12], but this is not correct [13]. The authors concluded that screening had reduced the incidence of cancers with metastases but they split these cancers into...
two groups: those above and those below 20 mm in diameter. When the data from the two groups were added, there was no reduction in incidence of cancers with metastases [13]. In fact, the combined incidence was identical for the first and last year reported (1989 and 1997) [13]. Another frequent fallacy is to compare the percentages of advanced breast cancers before and after screening. This neglects that screening finds many small, overdiagnosed cancers and will provide misleading results [13].

**Does breast screening ‘catch it early’?**

The stable incidence of advanced breast cancer is in stark contrast to the massive increases in the incidence of *in situ* and early breast cancer. When screening is introduced, the incidence of early stage breast cancer doubles, and it increases several-fold for *in situ* cases [8]. This has been seen everywhere screening has been introduced [9]. The increases do not disappear or diminish, even with follow-up over decades [9], and they coincide with screening even though its introduction is separated by many years between countries [9].

According to previous screening theory, these massive increases should prevent late stage disease, but despite more than 20 years of breast screening, this has not happened, which is not surprising if we consider the biology of breast cancer [3].

Breast screening brings forward the time of diagnosis only slightly compared to the fact that, on average, the woman has harboured the tumour for more than 20 years before it is detected [3,13]. When detected at screening, about one-third of the cancers have already metastasized [3,13]. In the randomised trials, tumours in the control group were 21 mm on average, and in the screened group they were 16 mm. Screening programmes today detect breast cancers that are a little over 10 mm on average, rather than 20 mm for clinically detected disease [14]. This represents a reduction of 1 to 2 volume doublings of the 32 necessary doublings to reach 20 mm [15], which is equivalent to a few months’ growth for aggressive, fast-growing cancers [14] that can therefore easily ‘slip through the screen.’

The true difference in size between screen-detected and clinically detected breast cancers in a screened population must be less than 10 mm, however. Overdiagnosis of small breast cancers artificially inflates the difference, and length bias means that screening preferentially detects small, slow-growing cancers, simply because there is more time to detect them. Similarly, the clinically detected cancers include the fast-growing interval cancers and attendees are those that already see their doctor when they notice something is wrong (selection bias, or the healthy screenee effect).

**What caused the large decline in breast cancer mortality?**

Despite high screening participation since the mid-1980s, breast cancer mortality in Sweden was reduced by only 16% from 1989 to 2006 in women aged 50 to 69 years, much less than in Denmark (26%) and Norway (23%), which had only limited screening - for example, 20% of women aged 50 to 69 years were offered screening in Denmark [16]. It is a general trait that the average reduction in breast cancer mortality in Europe has been almost twice as large in younger, non-screened age groups as in those screened, and equally large in countries with and without screening [16]. The laurels for this remarkable achievement should go to improved treatment of the sick, not screening of the healthy.

**Summing up breast screening**

Screening has not delivered on its promises. The risk of being diagnosed with advanced breast cancer today is the same as before screening, which is why screening has not reduced breast cancer mortality [5,6] or mastectomy use [7]. However, the risk of being diagnosed with breast cancer has increased by 50% in the screened age group due to overdiagnosis [9], with severe consequences for those who experience it. Screening for breast cancer has turned out to be very complex and hard to justify.

**Competing interests**

The author declares that they have no competing interests.

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