

Cancer screening has never been shown to “save lives”

Vinay Prasad and colleagues argue that overall mortality rather than disease specific mortality should be the benchmark against which cancer screening is judged

Despite growing appreciation of the harms of cancer screening,¹⁻³ advocates still claim that it “saves lives.”⁴ This assertion rests, however, on reductions in disease specific mortality rather than overall mortality.

The burden falls on providers to provide clear information about both disease specific and overall mortality and to ensure that the overall goal of healthcare—to improve quantity and quality of life—is not undermined.⁷

In this article we argue that overall mortality should be the benchmark against which screening is judged and discuss how to improve the evidence upon which screening rests.

Why cancer screening might not reduce overall mortality

Discrepancies between disease specific and overall mortality were found in direction or magnitude in seven of 12 randomised trials of cancer screening.⁸ Despite reductions in disease specific mortality in the majority of studies, overall mortality was unchanged or increased.

There are two chief reasons why cancer screening might reduce disease specific mortality without significantly reducing overall mortality. Firstly, studies may be underpowered to detect a small overall mortality benefit. Secondly, disease specific mortality reductions may be offset by deaths due to the downstream effects of screening.

Underpowered studies lead to uncertainty and assumptions of benefit rather than scientific evidence of benefit. In the 30 year follow-up of the Minnesota Colon Cancer Control Study, which assessed annual fecal occult blood testing, there were 128 deaths from colon cancer per 10 000 participants in the screened group and 192 per 10 000 in the control arm—a statistically significant difference of 64 deaths per 10 000.¹⁰ But there was a difference of only two overall deaths between the screened arm (7111 deaths per

10 000) and the control arm (7109 deaths per 10 000; $P=0.97$). Hazard ratios and Kaplan Meier curves corroborate this finding of no mortality difference. For 80% power to detect a difference in overall mortality of 64 deaths per 10 000 (assuming the disease specific benefit was not offset by other deaths), the trial would have needed to be about five times as large.

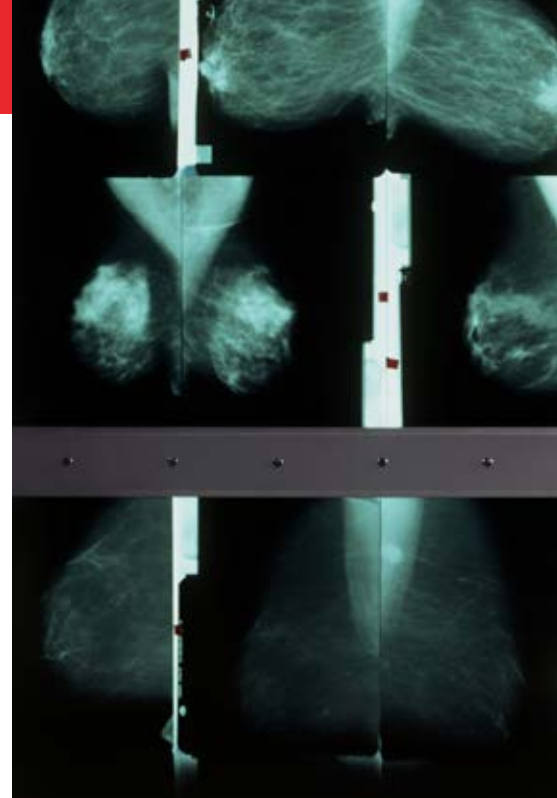
However, meta-analyses of fecal occult blood testing have shown a slight increase in deaths unrelated to colorectal cancer associated with screening, which implies that downstream effects of screening may partially or wholly negate any disease specific gains.¹¹

Such “off-target deaths” are particularly likely among screening tests associated with false positive results, overdiagnosis of non-harmful cancers, and detection of incidental findings. For example, prostate specific antigen (PSA) testing yields numerous false positive results, which contribute to over one million prostate biopsies a year.¹² Prostate biopsies are associated with serious harms, including admission to hospital and death.¹²⁻¹³ Moreover, men diagnosed with prostate cancer are more likely to have a heart attack or commit suicide in the year after diagnosis¹⁴ or to die of complications of treatment for cancers that may never have caused symptoms.¹²⁻¹³

The overall effect of cancer screening on mortality is more complex than a disease specific endpoint can capture, owing to the harms of further testing, overdiagnosis, and overtreatment. Realisation of this has led to reversal or abandonment of a number of screening campaigns, including chest radiography screening for lung cancer, urine testing for neuroblastoma, and PSA for prostate cancer.⁶⁻¹⁵⁻¹⁸

Mortality benefits of screening trial require close scrutiny

Arguably the strongest evidence that a single screening test can save lives comes from the National Lung Cancer Screening Trial (NLST), which randomised 53 454 heavy smokers to



receive either low dose computed tomography (CT) or chest radiography. CT was widely reported to show a 20% relative reduction in lung cancer deaths and a 6.7% relative reduction in overall mortality.¹⁹ However, the absolute risk reduction in overall mortality was only 0.46%, and several limitations undermine even this narrow margin.

Firstly, chest radiography for lung screening is not standard of care—it is well known not to improve disease specific or overall mortality.²⁰

Secondly, in the CT group the improvement in overall mortality exceeded the gains in lung cancer mortality by 36 deaths (87 fewer deaths from lung cancer and 123 fewer deaths overall). But CT screening did not seem to reduce deaths due to other cancers or improve cardiovascular survival to account for these 36 fewer deaths. If we assume that the improvement in non-lung cancer mortality was by chance and remove this difference, the overall mortality benefit disappears ($P=0.11$).

Thirdly, the benefit in lung cancer mortality of CT screening (estimated to avert over 12 000 lung cancer deaths in the US annually)²⁵ must be set against the 27 034 major complications (such as lung collapse, heart attack, stroke, and death) that follow a positive screening test (NLST investigators, personal communication, 2015).¹⁹

Public perception of screening

A systematic review has shown that the public has an inflated sense of the benefits and discounted sense of the harms of mammography screening, the cervical smear test, and PSA screening.²⁷ In one study 68% of women thought that mammography



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Such trials are worth the expense compared with the continued cost of supporting widespread screening campaigns without knowing whether they truly benefit society.⁵

The cost of adopting CT screening for lung cancer by the Medicare population has been estimated to surpass \$6bn a year.⁴⁶

To reduce costs, trials could target just the highest risk groups, with successful results prompting trials in lower risk groups. For example, the potential benefits of CT screening for lung cancer vary by age and smoking history of the participant.⁴⁷

Screening trials could also ascertain all causes of death among all participants to monitor any increase in off target deaths.⁴² This would be an improvement over current standards, but it would not overcome most of the concerns we have identified. Primary study data should be made available in a usable format for re-analysis.⁴⁸⁻⁵⁰

Barriers to trials powered for overall mortality

Political will, financial resources, and public perception are common hurdles in building support for resource intensive scientific endeavours, and developing consensus on these matters will take time and effort.

Conclusion

We encourage healthcare providers to be frank about the limitations of screening—the harms of screening are certain, but the benefits in overall mortality are not. Declining screening may be a reasonable and prudent choice for many people. Providers should also encourage participation in open studies.

We call for higher standards of evidence, not to satisfy an esoteric standard, but to enable rational, shared decision making between doctors and patients. As Otis Brawley, chief scientific and medical officer of the American Cancer Society, often states: “We must be honest about what we know, what we don’t know, and what we simply believe.”

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would lower their risk of getting breast cancer, 62% thought that screening at least halved the rate of breast cancer, and 75% thought that 10 years of screening would prevent 10 breast cancer deaths per 1000 women.²⁸ Even the most optimistic estimates of screening do not approach these numbers.^{29 30}

As long as we are unsure of the mortality benefits of screening we cannot provide people with the information they need to make an informed choice. We must be honest about this uncertainty.

A summary of the Swiss government’s decision not to recommend mammography shows that for every 1000 women who undergo screening one breast cancer death is averted (from five to four), while non-breast cancer deaths either remain at 39 or may increase to 40.³⁴ If non-breast cancer deaths remain the same, a woman must weigh net benefit against harms. If screening increases non-breast cancer deaths to 40, women would simply be trading one type of death for another, at the cost of serious morbidity, anxiety, and expense. Women should be told that to date, with over 600000 women studied, there is no clear evidence of a reduction in overall mortality with mammography screening.³⁰

Harms

Consideration of harms becomes more important in the absence of clear overall mortality benefit. Empirical analyses show that primary screening studies pay little attention to the harms of screening—of 57 studies only 7% quantified overdiagnosis and just 4% reported the rate of false positive

results.³⁵ When researchers do examine the harms of screening the results are typically sobering.

False positive results on breast cancer screening have been associated with psychosocial distress as great as a breast cancer diagnosis 6 months after the event.³⁶

False positive results affect over 60% of women undergoing screening mammography for a decade or more,³⁷ and 12-13% of all men who have undergone three or four screening rounds with PSA.³⁸ In the NLST 39.1% of people had at least one positive test result, of which 96.4% were false positives. One in 12 (9.4%) of those people had invasive procedures—such as needle biopsy, mediastinoscopy, and open thoracotomy—to learn that they didn’t have lung cancer.

What next?

How can we know whether screening saves lives? We need trials that are ten times larger and powered for overall mortality.^{5 6} Researchers have postulated, based on a colorectal cancer trial, that 4.1 million participants would be needed to demonstrate a reduction in overall death, compared with 150000 for disease specific death.⁴²

Studies of this size may be estimated to cost upwards of \$1bn (£0.7bn; €0.9bn), but conducting such trials in large national observational registries would dramatically reduce the cost. Large trials should be pragmatic, with inclusion criteria that mirror the real world population in which the intervention is used. The safest way to introduce or change screening programmes at the national level is by incorporating randomisation.⁴⁵