PSA testing for men at average risk of prostate cancer

Bruce K Armstrong\textsuperscript{a,b}, Michael J Barry\textsuperscript{c,d}, Mark Frydenberg\textsuperscript{e,f}, Robert A Gardiner\textsuperscript{g,h,i,j,k}, Ian Haines\textsuperscript{l,m} and Stacy M Carter\textsuperscript{n,o}

\textsuperscript{a} School of Population Health, University of Western Australia, Perth
\textsuperscript{b} Sydney School of Public Health, University of Sydney, NSW, Australia
\textsuperscript{c} General Medicine Division, Massachusetts General Hospital, Boston, United States
\textsuperscript{d} Department of Medicine, Harvard Medical School, Boston, MA, United States
\textsuperscript{e} Urological Society of Australia & New Zealand, Sydney, NSW, Australia
\textsuperscript{f} Department of Surgery, Faculty of Medicine, Monash University, Melbourne, Vic, Australia
\textsuperscript{g} Faculty of Medicine, University of Queensland, Brisbane, Australia
\textsuperscript{h} Centre for Clinical Research, University of Queensland, Brisbane, Australia
\textsuperscript{i} Royal Brisbane and Women’s Hospital, Qld, Australia
\textsuperscript{j} School of Medical and Health Sciences, Edith Cowan University, Perth, WA, Australia
\textsuperscript{k} Griffith University, Brisbane, Qld, Australia
\textsuperscript{l} Cabrini Haematology and Oncology Centre, Cabrini Hospital, Melbourne, Vic, Australia
\textsuperscript{m} AMREP Dept of Medicine, Faculty of Medicine, Monash University, Melbourne, Vic, Australia
\textsuperscript{n} Centre for Values, Ethics and the Law in Medicine, University of Sydney, NSW, Australia
\textsuperscript{o} Corresponding author: stacy.carter@sydney.edu.au

Abstract

Prostate-specific antigen (PSA) testing of men at normal risk of prostate cancer is one of the most contested issues in cancer screening. There is no formal screening program, but testing is common – arguably a practice that ran ahead of the evidence. Public and professional communication about PSA screening has been highly varied and potentially confusing for practitioners and patients alike.

There has been much research and policy activity relating to PSA in recent years. Landmark randomised controlled trials have reported; authorities – including the 2013 Prostate Cancer World Congress, the Prostate Cancer Foundation of Australia, Cancer Council Australia, and the National Health and Medical Research Council – have made or endorsed public statements and/or issued clinical practice guidelines; and the US Preventive Services Task Force is revising its recommendations. But disagreement continues.

The contention is partly over what the new evidence means. It is also a result of different valuing and prioritisation of outcomes that are hard to compare: prostate cancer deaths prevented (a small and disputed number); prevention of metastatic disease (somewhat more common); and side-effects of treatment such as incontinence, impotence and bowel trouble (more common again). A sizeable proportion of men diagnosed through PSA testing (somewhere between 20\% and 50\%) would never have had prostate cancer symptoms sufficient to prompt investigation; many of these men are older, with competing comorbidities. It is a complex picture.
Below are four viewpoints from expert participants in the evolving debate, commissioned for this cancer screening themed issue of Public Health Research & Practice. We asked the authors to respond to the challenge of PSA testing of asymptomatic, normal-risk men. They raise important considerations: uncertainty, harms, the trustworthiness and interpretation of the evidence, cost (e.g. of using multiparametric magnetic resonance imaging to triage patients with elevated PSA), a likely bias towards intervention (particularly for cancer), and the potential to limit harm by treating more conservatively (although this may not occur consistently). They provide important insights, and disagree on some issues, but generally concur that men should decide for themselves whether to be tested. It seems reasonable to support men’s autonomy to make their own decisions based on their own values. However, the support men might require to decide is likely to be considerable, and this needs to be taken seriously in policy making.

**View from a cancer epidemiologist – by Bruce K Armstrong**

That early detection of disease is good is held as true by most health professionals and most people. It is not surprising, therefore, that early detection action runs ahead of evidence of early detection benefit. This is true of all national screening programs for cancers in Australia – cervical cancer, breast cancer and colorectal cancer – and it is true of prostate cancer, among others, for which there is no formal national program. The urge of clinicians to find cancer early and of people to avoid death from the ‘big C’ are so great that it is also not surprising that cancer screening without evidence of benefit persists, despite evidence or a strong presumption of harm. The principal harms are investigation and sometimes treatment based on a false positive test; clinical delay and perhaps poorer outcomes after a false negative test; and detection, diagnosis and treatment of cancers that would otherwise have never affected life or health (overdiagnosis).

How should public health policy tackle prostate-specific antigen (PSA) testing for the early detection of prostate cancer? How do we proceed when there is uncertain benefit and certain harm, when up to 21% of Australian men aged 45–74 participate in screening each year and may benefit, and 19% aged over 74 also participate and probably won’t? Ban it? Regulate it? Or do nothing? (Note: These percentages were calculated from data on claims for PSA tests [item number 66655] processed by Medicare Australia in 2015–16 [medicarestatistics.humanservices.gov.au/statistics] and Australian Bureau of Statistics tables of estimated resident population by age and sex at 30 June 2015 and 30 June 2016 [ABS report series 3101.0 Australian Demographic Statistics], with the value 7.5 added to each calculated percentage to correct for the effect of Medicare’s ‘episode coning’ policy on recording of pathology items claimed for.)

Faced with these choices and the lack of other likely action, Cancer Council Australia (CCA) and the Prostate Cancer Foundation of Australia (PCFA) chose to ‘regulate’ by developing clinical practice guidelines for PSA testing and early management of test-detected prostate cancer. These nonbinding guidelines aimed “to maximise the benefits, if there are benefits, and minimise the harms from PSA testing”¹. They were approved by Australia’s National Health and Medical Research Council on 2 November 2015.¹ Early management was included because early management decisions can influence the realisation of assumed benefits and known harms of PSA testing.

The PCFA and CCA expert advisory panel made two important early decisions: that a national PSA testing program akin to the national breast cancer screening program (for example) would not be recommended, and that the reported results of the European Randomized Study of Screening for Prostate Cancer (ERSPC) would provide the evidence base for its PSA testing recommendations. The ERSPC is the largest study addressing PSA test efficacy, and one of the two largest, and the most recent, to report prostate cancer mortality reduction in PSA-tested, average-risk men. This was a necessary decision because only the ERSPC or statistical models based on its results, rightly or wrongly, can inform recommendations about the age range for, and frequency of, PSA testing, and the PSA level above which further investigation is recommended.

The guidelines make no recommendation about whether men should be routinely offered PSA testing. Their key testing recommendation is: “For men at average risk of prostate cancer who have been informed of the benefits and harms of testing and who decide to undergo regular testing for prostate cancer, offer PSA testing every 2 years from age 50 to age 69, and offer further investigation if total PSA is greater than 3.0 ng/mL”. Very importantly, they also recommend that the responsible clinical practitioner “offer evidence-based decisional support to men considering whether or not to have a PSA test, including the opportunity to discuss the benefits and harms of PSA testing before making the decision”. Guidance is given as to how the PSA testing recommendation might be modified for men at higher than average risk of prostate cancer, and recommendations are made on management of men who have a PSA >3.0 ng/mL or a biopsy diagnosis of prostate cancer, aiming at all times to obtain a balance between the benefits hoped for and the possible harms.
There is, as yet, no formal plan to evaluate implementation of the guidelines in practice or whether the assumed benefits of PSA testing are occurring, or whether the benefits are even increasing and harms decreasing. We might reasonably hope to see a fall in Australia’s very high prostate cancer incidence rates following guideline implementation because of a fall in prostate cancer overdiagnosis due to the recommended narrower age range for, and lower frequency of, PSA testing than has been common in recent Australian practice. Because of the lack of any recommendation for or against PSA testing, and PSA’s uncertain screening efficacy, any prediction of the prostate cancer mortality trend would be pure guesswork.

View from the United States – by Michael J Barry

There is a lot not to like about screening for prostate cancer with the PSA test. In the ERSPC, arguably the highest quality trial, although not without its faults\(^2\), benefits were modest. After 13 years, 1.28 fewer prostate cancer deaths per 1000 men were observed with screening, with no decrease in overall mortality.\(^3\) Heterogeneity among ERSPC countries in screening protocols, treatments and outcomes has raised questions about the magnitude of benefit actually attributable to PSA testing.\(^4\) On the other hand, the harms of screening are relatively high, with 31 additional cases of prostate cancer needing to be detected per prostate cancer death prevented.\(^5\) Men are not used to thinking of PSA screening as ‘causing’ cancer, but, for practical purposes, the substantially higher incidence with screening means just that. The higher incidence reflects more prostate biopsies, with short-term complications of haematuria, haematochezia, haematospermia and infections. The main harms, however, come from the treatment of men found to have prostate cancer, including substantial risks of erectile dysfunction and incontinence. Most of these cancers are never destined to cause morbidity or mortality,\(^6\) and the men who harbour them can only be harmed by early detection. Finally, at least in relation to practices in the US, PSA screening is not cost-effective.\(^6\)

That all sounds like a pretty good argument for not screening, right? Perhaps. However, the absolute benefit from prostate cancer screening, if the ERSPC estimate is correct, is not dissimilar to the benefit of screening for breast cancer, and reducing overall mortality may be too high a bar for any screening test. Modelling strategies, and the reduction in metastases seen in the ERSPC, suggest that the absolute benefit may be greater over a longer time. Strategies being considered for maintaining most of the benefits while reducing the harms include testing less frequently (the screening interval in the ERSPC was 2–4 years), higher biopsy thresholds and, most importantly, active surveillance for men with low-risk cancers. These strategies to mitigate harm are not perfect and will come at the price of slightly higher risks of bad outcomes. For example, in the ProtecT study, active monitoring had the same cancer-specific mortality as surgery or radiation over 10 years (about 1%) but a higher risk of metastases (6% versus 3%).\(^7\) Whether these strategies, which may make screening cost-effective,\(^8\) will prove acceptable to most clinicians and patients remains to be seen. The most important reason for not rejecting PSA screening is that there is variation in how men see the trade-offs between possible benefits and harms. When men were fully informed about the trade-offs in a decision making process including a patient decision aid, and could answer key knowledge questions accurately, about a third still wanted a PSA test.\(^8\) Key points to cover in a conversation between a man and a clinician have been previously reviewed.\(^5\)

Some health systems will reasonably decide not to make PSA screening generally available because their limited resources can be used more effectively in other ways. However, if PSA screening is available, letting informed men help decide whether a PSA test is right for them seems the most patient-centred way forward. After all, they are the ones who must live with the consequences of the decision.

View from urologists – by Mark Frydenberg and Robert A Gardiner

Before considering selective screening for prostate cancer, the question to be asked is whether a diagnosis will benefit the patient. The answer can only be determined by establishing whether the patient accepts the risks and potential side effects of the diagnostic and treatment processes, appreciates the likelihood of cancer (if present) to affect his health and wellbeing, and can make a value judgement about what is really important to him. Consequently, there is no place for mass population or so-called opportunistic PSA screening.

The large majority of men investigated for prostate cancer are asymptomatic and do not have suspected malignancy based on a digital rectal examination. Because of the long natural history of most tumours, a 7–10-year life expectancy following treatment (and therefore diagnosis) is considered warranted before considering PSA testing. Pertinently, many patients have significant or latent comorbidities that will significantly affect their survival within a decade, so they will not live long enough to achieve a survival benefit.\(^9\)–\(^12\)

Serious health problems are not uncommon in middle-aged and elderly males, along with lifestyle-related factors such as smoking and obesity, which have yet to overly affect morbidity and mortality. A poor appreciation of individual life expectancy is not just limited to patients, as many clinicians are also overoptimistic and give patients ‘the benefit of the doubt’ when recommending investigations and treatments.\(^12\) To introduce some objectivity to overall patient prognosis, life expectancy tables\(^13\)–\(^14\) may be helpful, although they are population
based and do not take into account an individual’s comorbidities or sociodemographic factors.

Selective screening involves identifying men who are at risk of developing clinically significant prostate cancer, and are likely to benefit from a prostate cancer diagnosis and therefore PSA testing. As we have previously reported, a family history (particularly in first-degree relatives) is well recognised to predispose to a future diagnosis of prostate cancer, but a PSA –1.5, 12.4 ng/mL for men aged less than 50 years is regarded as even more predictive than either family history or ethnicity. Hereditary prostate cancers occur more commonly than any other tumour diagnosed – on average, detected 6 years earlier than for sporadic cancer. Patients with a family history of germline mutations in the family-susceptibility genes BRCA1 and BRCA2 have a significantly increased risk of developing this malignancy, tend to present at a younger age, and tend to have more aggressive disease and poorer survival outcomes.

Increasingly, multiparametric magnetic resonance imaging (mpMRI) is being used to triage patients with an elevated PSA. A combination of anatomical (T2-weighted) images with at least two of the three functional MRI parameters (diffusion-weighted imaging, dynamic contrast-enhanced imaging and spectroscopy) has been estimated to identify approximately 90% of moderate- to high-risk lesions, but is less reliable for detecting small (<0.5 cubic centimetres) and lower risk tumours. Limitations to using mpMRI are the level of accuracy of MRI interpretation and cost. However, until a cheaper and comparably accurate diagnostic test replaces PSA testing, the combination of PSA testing and mpMRI will remain the initial investigation of choice before biopsy.

It is easy to overlook or underestimate the quality-of-life impacts, which range from the anguish of possibly harbouring a malignancy to the uncommon but potentially devastating effects of infection associated with a biopsy, to the side-effects of the various treatments. Some men may rather risk a cancer spreading (especially in a cancer of lower histological grade) and remain untreated than risk losing their sexual or urinary abilities, in addition to other changes in bodily function that can affect social confidence and self-esteem. Consequently, it is imperative at the outset to evaluate and respect decisions made with respect to quality of life when deciding whether to test for prostate cancer.

**View from an oncologist – by Ian Haines**

In 2012, the US Preventive Services Task Force (USPSTF) downgraded its recommendation for screening using PSA testing from C to D. Since that advice, the much-anticipated ProtecT study has been published. It randomised 1643 men equally between radical surgery, radical radiation and active monitoring, and reported results at 10 years. It revealed that only 1% of men with early-stage prostate cancer died of their disease in the first 10 years after diagnosis, irrespective of treatment and the usual prognostic factors.

Although treatment achieved a significant reduction in local disease progression and metastases, the lack of a significant survival benefit for radical surgery over active monitoring confirms the results of the Prostate Cancer Intervention Versus Observation Trial (PIVOT), the only other randomised trial done in the era of PSA screening tests. It found no significant prostate cancer–specific mortality or overall survival benefit for radical surgery over active monitoring at 12 years, confirmed at 19.5 years follow-up. PIVOT showed an absolute risk reduction of 5.5% (95% confidence interval [CI] –1.15, 12.4%; p = 0.06) for all-cause mortality (61.3% in patients treated with prostatectomy vs 66.8% in patients who had active monitoring) and an absolute risk reduction of 4% (95% CI –0.2%, 8.3%; p = 0.06) for prostate cancer–specific mortality (7.4% in patients treated with prostatectomy vs 11.4% in patients who had active monitoring). Although surgery reduced the need for treatment of progressive disease, there were increased treatment-related long-term complications such as urinary incontinence and erectile dysfunction. The dissemination of these results to every man over 40 will aid their informed decision making about PSA testing, and may reduce the overdiagnosis, overtreatment, and significant and immediate physical and psychological harms caused to many of them.

One of the reasons for this widespread testing has been that of the two large randomised studies of screening for prostate cancer with PSA, the ERSPC, showed a small survival benefit for screening. However, it has been suggested by the chief medical and scientific officer of the American Cancer Society that this study has flaws and unintended biases, such as a large disparity between primary treatment with androgen deprivation monotherapy given to similar risk patients in each arm. The call for an independent review of the mortality data was repeated in a recent review. Conversely, the other major randomised screening study, which showed no benefit for screening, was recently found to have major contamination of the control arm, in which 90% of men had a PSA test during the study period. This makes its results less reliable, although it is important to note that there were still 22% more cancers diagnosed in the screening arm after 2 years, with no associated survival benefit.

Despite the efforts of many urologists to reduce overdiagnosis and overtreatment, some urology groups suggest that all men over 40 should consider a screening PSA test, even when we know that 24.4% of men with a ‘normal’ PSA are diagnosed with prostate cancer when they have a biopsy and it is a common finding in this age group at autopsy. Many are advised to consider radical treatment, with all the possible adverse consequences, for a disease that kills 1 in 7 (2–3% of all men) at a median age of 82.4 and for which there is still no proven survival benefit for radical intervention.

This is poor public health policy. The USPSTF should not reverse its previous advice.
Competing interests

BA’s then-employer, the Sax Institute, was reimbursed by the PCFA for remunerated time spent advising the systematic evidence review team for the PSA testing guideline and on writing parts of the guideline document. BA chairs the PCFA’s Research Advisory Committee and is an ex officio member of its National Board. BA receives no remuneration for either of these memberships, but all expenses for attending meetings are paid for by the PCFA, and he attends Committee and Board dinners hosted by the PCFA. BA is an Associate Investigator of a PCFA randomised controlled trial of vitamin D in the prevention of progression of prostate cancer managed by active surveillance. He was one of the initiators of the process to develop an Australian guideline for PSA testing and the early management of test-detected prostate cancer, and was a member of the expert advisory panel that oversaw development of the guideline and agreed on its final text.

MB is a member of the USPSTF. His section does not necessarily represent the views and policies of the USPSTF. MB received salary support from the nonprofit organisation Healthwise as Chief Science Officer, and Healthwise provided grants to Massachusetts General Hospital for MB's prostate-related and shared decision making research.

RG is a member of the Andrology Australia Advisory Board. RG is also involved in NHMRC- and Movember-supported research into management of men with prostate cancer.

IH is a consultant oncologist who sees men in paid consultations who want second opinions about how to proceed with their PSA or biopsy results.

Author contributions

BA is the sole author of the section from a cancer epidemiologist. MB is the sole author of the section from the US. RG and MF worked together to produce the section from urologists. IH is the sole author of the section from an oncologist. SC wrote the abstract and key points, and collated and approved the final manuscript.

References


