Prostate Cancer Screening: Canadian Guidelines 2011

The Canadian Urological Association

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Methods/ data collection

A systematic literature search was conducted in the following electronic bibliographic databases: MEDLINE including PreMedline (2004 to November 2010), EMBASE (2004 to Week 44, 2010), the Cochrane Central Register of Controlled Trials (2010, 4th Quarter) and was restricted to studies published in English language.

The search queries were based on combination of exploded and non-exploded subject headings and free-text keywords and included terms prostate cancer, prostatic neoplasms, prostate tumor, prostate-specific antigen, digital rectal examination, dre, mass screening, screening test, early detection of cancer, cancer screening, screening, psa, transrectal ultrasound, trus, randomized, false-negative, false-positive, using alternative word spellings and endings.

The search strategy was modified for each database using database-specific thesaurus terms, syntax and search fields. For all searches, case reports and editorials were excluded along with news and letters. To identify additional relevant studies we also examined bibliographies of the relevant articles and selected reviews.
We compiled 1,938 unique citations and after removing the duplicates 1,036 citations were assessed for relevance. The screening process yielded 49 articles for a full text review.

Introduction:

The goal of prostate cancer (PCa) screening is to reduce the morbidity and mortality from this disease by early detection. There has been a steady decline in PCa mortality following the introduction of PSA testing in the late 1980’s. However, phase III trials on PCa screening have demonstrated a high number needed to treat, suggesting many PCAs diagnosed were not destined to have an impact upon a patient’s overall survival. (Level 1) We herein review the literature on PCa screening and describe what contemporary screening entails. We offer guidelines to physicians to facilitate the discussion of the risks and benefits of PCa screening for appropriate men. These guidelines are recommendations, are not a standard of care for all patients and should not preempt a physician’s clinical judgment.

Epidemiology

PCa remains the third leading cause of cancer death in men in Canada. The lifetime risk of death from PCa is 3.7 percent and the lifetime risk of PCa diagnosis is 13.6 percent. A Canadian male has a 1 in 7 chance of being diagnosed with PCa and a 1 in 27 chance of dying of PCa. The incidence of PCa at autopsy in men over 50 years of age dying of other causes ranges from 33 to
The chance that a man over 55 years of age will be diagnosed with PCa with a standard biopsy is 25 percent. It is evident from the epidemiologic data alone that the majority of men with histologic PCa are not destined to die from PCa.

**Prostate Cancer Screening Tests:**

DRE and PSA are the first line PCa screening tests. *(Level 2, Grade A)*

a) Digital Rectal Exam

DRE and PSA have different sensitivities and specificities for PCa detection and DRE may identify PCas that would not necessarily be picked up on PSA alone. However, from the European Randomized Study of Screening for Prostate Cancer (ERSPC), DRE did not provide any further additive information beyond PSA. *(Level 1)* The DRE can be normal or show findings consistent with BPH, such as symmetrical enlargement. Findings associated with PCa include induration or hard nodules. Controversy exists regarding asymmetry alone as a predictor of PCa. The positive predictive value of the DRE increases as the PSA increases. PCas identified by DRE are pathologically advanced in over 50 percent of men. *(Level 3)*

b) Prostate Specific Antigen

PSA is a glycoprotein produced primarily by the prostate epithelial cells that line the ducts and acini of the prostate. It is thought that the disruption of the normal prostate glandular architecture facilitates PSA’s access to the
systemic circulation. Conditions that increase PSA include benign prostatic hyperplasia, prostatitis, urethral instrumentation, prostate biopsy and PCa. Inconsistent causes of PSA elevation include a vigorous DRE and recent ejaculation. 5-alpha reductase inhibitors (5aRIs) reduce the PSA by about 50% by 6 months and is non dose-dependent.\(^5\) Patients on 5aRIs that have reached their nadir should have a baseline level after 6 months of medication. Any sustained rise should raise concerns for biologically significant PCa.\(^6\)

A PSA cut point of 4ng/ml was originally chosen to differentiate normal PSA levels from pathological elevation. In contemporary series, a PSA level of 4 ng/ml has a sensitivity of 20 percent in detecting PCa.\(^6\) To improve upon PSA sensitivity in younger men and increase specificity in older men, PSA age-specific reference ranges have been utilized.\(^6\) However, the Prostate Cancer Prevention Trial (PCPT) demonstrated that the relationship between PSA and PCa incidence is continuous (i.e. the higher the PSA, the greater the risk). (Level 1) There is no single justifiable cutpoint, regardless of age.\(^8\)

Since PSA levels may fluctuate from causes other than PCa, a single PSA value is not sufficient. Decisions regarding whether a prostate biopsy should be recommended should be based on more than a single PSA.\(^18\) Between laboratory variation is 20-25% and using the same laboratory may reduce variability.\(^19,20\) If the PSA level is below 1.0 ng/ml, biologically significant PCa is very unlikely to develop over the next 7 years.\(^12\) (Level 2)
PSA velocity (PSAV) may improve the sensitivity of PSA. To calculate PSAV, at least three PSA determinations must be used over a time period of 18 months.\textsuperscript{21, 22} A PSAV rise of greater than 0.75 ng/ml/year when the PSA is between 4-10 ng/ml may indicate a higher risk of PCa.\textsuperscript{22} However, for PSA levels less than 4 ng/ml, PSA velocity rises of 0.4 ng/ml/year appear to denote a higher risk of PCa.\textsuperscript{21, 23, 24} Age-specific PSAV with cut points of 0.25, 0.5 and 0.75 ng/ml/y in men aged 40-59, 60-69 and over 70 years, respectively has been described to improve the sensitivity of PCa detection.\textsuperscript{24} (Level 3) It is important to note that in phase III trials, the PSAV has not been shown to be an independent predictor of a positive biopsy over PSA alone.\textsuperscript{20, 25} PSAV alone should not be the basis for a decision to biopsy.

PSA densities (PSAD) may improve the specificity in detecting PCa.\textsuperscript{26} PSAD greater than 0.15 ng/ml may indicate a greater risk of PCa. However, not all studies confirm the utility of PSAD. Measuring the PSAD for the transition zone only may also be utilized, as the transition zone usually accounts for the PSA production by the BPH component.\textsuperscript{26} For these volume/density measurements, a transrectal ultrasound (TRUS) is required. This imposes additional costs, has not been shown to improve PCa screening and TRUS has variable interoperator reproducibility.\textsuperscript{27, 28} (Level 3)
PSA isoforms can also improve PCa detection, while improving sensitivity of prostate biopsies. The majority of circulating PSA bound to alpha-1 antichymotripsin (ACT) and alpha-2 macroglobulin (AMG), while the remainder is free in the serum. In normal prostatic acini, PSA is processed by enzymatic nicking of a 7 kD segment, followed by back diffusion into the serum. In PCa acini, loss of polarity results in direct secretion into the serum with no enzymatic nicking. This results in greater avidity between ACT and PSA, thus less free PSA. The lower the free to total PSA ratio, the greater the risk of PCa. There is no known optimal cut point. Like total PSA, the free to total ratio presents a continuum of PCa risk. Using the free to total PSA ratio improves specificity and will reduce the number of negative prostate biopsies. Hemodialysis and peritoneal dialysis do not affect the total serum PSA levels, but may affect the free PSA levels.

Measured complexed PSA levels may improve the specificity of PSA screening, but the data on complexed PSA is limited and this isoform is not commonly utilized. Biomarkers, such as PCA3, may play a more significant role in PCa screening, but the data supporting its use for routine screening is limited.

**Transrectal Ultrasound-Guided Prostate Biopsy:**

Once the PCa screening evaluation determines that a man is at increased risk of PCa, a TRUS guided prostate biopsy is required to obtain the histologic
diagnosis. The biopsy provides important data, including the grade and volume of PCa and the presence of extraprostatic disease. No single biopsy template is optimal for all men. A 10 to 12 core biopsy template is standard and should incorporate tissue from the peripheral zones and anterior horns.\textsuperscript{34,35} (Level 3, Grade B) Transition zone biopsies are not necessary with initial prostate biopsies unless there is a suspicious lesion identified on the TRUS.\textsuperscript{35} Nomograms incorporating prostate volume and age may be useful to determine the number of biopsies cores to optimize detection rates.\textsuperscript{36} Despite the critical data biopsies provide, morbidity and rising post-biopsy sepsis rates have to be taken into consideration as a significant risk of PCa screening.\textsuperscript{37}

**Biologically Significant Prostate Cancer:**

There are no clinical parameters that afford unequivocal prediction of the biologic significance of a patient’s detected PCa. At the extremes of disease where a man has metastatic PCa or low volume, low-grade disease, PCa biology is easier to predict. For men with organ-confined PCa in the absence of high grade and high volume disease, predicting the risk of morbidity and mortality relative to other medical comorbidities can be challenging. The clinical data most commonly used to predict disease biology include PSA, histologic grade, clinical or pathologic stage and cancer volume. The number of biopsy cores containing cancer and the extent of cancer involvement of each core provides an estimate of cancer volume.\textsuperscript{38-40}
Based on autopsy data, biologically insignificant PCa has been defined as tumour volume < 0.5 cc with a Gleason score 6 or less. Histologic criteria for biologically insignificant PCa is based on core biopsies and include PSA density < 0.15, no Gleason grade 4 or 5, involvement of < 3 needle biopsy cores (on sextant biopsies) and involvement of < 3 mm of tissue in any one biopsy core.\(^1\) It is important to note that a ‘biologically significant’ PCa may be present in up to 24 percent of patients using these criteria.\(^2\) A caveat is that the biologic significance of grade 3 cancer > 0.5cc has never been established.

Standard radiologic imaging has not allowed for better local tumour volume assessment.\(^3\) Multiparametric magnetic resonance imaging may have an emerging role in assessing tumour volume and improving biopsy accuracy.\(^4\) Nomograms are available that combine many clinical variables to try to determine which PCas are biologically insignificant.\(^5\) Inaccuracies range from 10 to 20 percent. \((\text{Level 3})\)

**Active Surveillance:**

Many men diagnosed with screen-detected PCa will have disease that is not destined to affect their overall survival. Watchful waiting is a strategy whereby there is no therapy until the patient develops symptoms secondary to advanced PCa. Treatment usually involves androgen deprivation for metastatic disease. This approach does not offer an opportunity for cure in men with more aggressive disease.\(^6\) Active surveillance entails close follow-up on patients
diagnosed with early stage, low-risk PCa. Therapy is recommended at a time when cure is deemed possible and when disease progression occurs as defined by a rapid PSA doubling time, clinical progression on DRE, grade progression or tumour volume progression on repeat TRUS guided prostate biopsies.

Seven studies on active surveillance indicate it appears safe.\(^ {45-51}\) (Level 3) The total number of patients in these studies is 2365 with a median follow-up ranging from 22 to 73 months. Overall survival ranges from 82 to 100 percent. PCa survival ranges from 97 to 100 percent. Patients remaining on active surveillance range from 50 to 92 percent. The main limitation to these studies is the lack of long-term follow-up. An international, prospective, randomized, phase III trial (Surveillance Therapy Against Radical Treatment – START) randomizing men with low risk PCa to active surveillance or immediate curative therapy is accruing patients. Until the START trial is completed, active surveillance with possible delayed curative intervention seems to be a reasonable option for men with favourable risk, screen-detected PCa.

**Curative Therapies:**

For any cancer-screening program to be effective, there must be curative therapies. Of all the potentially curative therapies for organ-confined PCa, phase III data on 695 patients comparing curative therapy to watchful waiting exists only for radical prostatectomy.\(^ {52}\) It was shown that there was an overall and disease-specific survival advantage for those men undergoing radical
prostatectomy. (Level 1) However, one needed to treat 17 men to save 1 life from PCa and this study was in the pre-PSA era. In a separate evaluation of the same study with a longer median follow-up of 10.8 years, overall survival was not different, but PCa mortality and risk of metastases were reduced significantly by radical prostatectomy.\textsuperscript{53}

Phase III survival data also exists for post-operative radiotherapy conferring a survival benefit for men at higher risk for disease recurrence post-radical prostatectomy.\textsuperscript{54, 55} (Level 1) There are local therapies that do provide a survival benefit for patients, although given the early follow-up of these studies relative to the prolonged natural history of localized PCa, the overall benefit compared to harm is yet to be fully elucidated.

**Level 1 Evidence for Screening:**

The initial 2 multicenter, randomized, prospective PCa screening studies have been recently published.\textsuperscript{2, 3} The major similarities and differences of these studies are highlighted in Table 1. There were limitations associated with the Prostate, Lung, Colorectal, and Ovarian (PLCO) Cancer Screening Trial that had a significant impact upon the results and may reduce generalizability. Firstly, only a PSA cut point of 4 ng/ml was used to determine if a biopsy was required. Only 67 percent of the patients in this study had 10-year data, so the study had insufficient follow up given the prolonged natural history of screen detected PCa. The short follow-up period also had an impact upon the low level of events
or deaths from PCa in the trial. Contamination of PSA screening in the control arm was high at 52%. Thus, this study compared a group of men that were more closely screened to a group undergoing less screening, as opposed to a group that was unscreened.

A third phase III trial randomizing men to PSA screening versus no screening was recently published. In this study from Sweden, men were randomized prior to receiving information regarding the screening protocol. Results relative to the 2 aforementioned screening trials were significantly different in favour of PCa screening. The numbers needed to screen and numbers needed to be diagnosed to save 1 life from PCa were 293 and 12, respectively. Potential explanations for the differences were men were younger (median age 56 at baseline), the PSA cut point was lower (2.5 – 3.4 ng/ml), contamination was lower (3 percent received PSA testing prior to study), not all patients underwent immediate curative therapy and there was longer follow-up (median 14 years).

Although the ERSPC and Swedish studies did demonstrate a difference in favour of the more intensely screened population, the number of patients needed to screen and treat to save one life from PCa indicate that there are men that undergo treatment (and the associated morbidity) who were not destined to die from their PCa. These figures are similar to breast cancer; however, the quality of life issues specific to PCa therapies make comparisons difficult.
Contemporary Screening:

Contemporary PCa screening for men with at least a 10-year life expectancy now involves more than just a DRE and PSA. No single PSA value should be the only determinant of whether or not to biopsy a patient. PSAV, PSAD and PSA free to total ratio may improve upon the PSA sensitivity and specificity. (Level 2, Grade B) Furthermore, nomograms and mathematical models may guide a clinician by combining multiple clinical variables, such as DRE, PSA, PSAV, PSA isoforms, age, race, family history of PCa and genetic data to determine one’s risk of PCa and the risk of biologically significant disease.58-60 (Level 3, Grade B)

Guidelines exist that recommend PCa screening begin at age 40, particularly for those at higher risk.61 (Level 3, Grade B) There is evidence that a baseline PSA at such an early age may also be a predictor for future risk of PCa and allow a risk-stratified approach for timing and intensity of PCa screening.62-65 Men with a PSA lower than 0.5 ng/ml are at lower risk of PCa and may benefit from PSA screening every few years, as was standard in the ERSPC trial.2,62-65 It is important to realize that opening PSA testing to all men at age 40 does pose increased risks of further testing and therapy and the data supporting earlier testing is limited without any supportive outcome data.62-65
The benefits of screening also decline with age due to competing causes of death, the long natural history of PCa and the lead-time bias of PSA screening.\textsuperscript{9, 66, 67} Recommendations from the US Preventative Services Task Force recommends against screening once a man reaches age 75.\textsuperscript{68 (Level3, Grade C)} Strong consideration should be given to discontinuing PSA screening for Canadian men over 75 years of age.

Conclusions:

PCa screening allows the detection of potentially lethal cancer at a point in time when it is more likely to be curable. This comes at the expense of many patients being treated when their cancer poses no threat to their life. Therefore, the harms and benefits of PCa screening must be explained to each patient so they understand all the factors to be considered in the shared decision-making about screening. PCa screening should be offered to all men 50 years of age with at least a 10-year life expectancy. Annual screening has been the standard, however, the two screening studies that demonstrated screening was beneficial screened men every 2-4 years. If there is a higher risk of PCa, such as family history of PCa or if the patient is of African descent, screening should be offered at age 40 years. Furthermore, there may be benefit of offering a baseline PSA for men 40 to 49 years of age to establish future PCa risk. Initial screening should include DRE and PSA. PSA and PSA free/total ratio are currently the most reliable serum markers. Both present a continuum of PCa risk. No strict cut-point should be used for all patients. The lowest cut-point
used in phase III trials demonstrating a benefit to screening was 2.5 ng/ml. Many other factors may be utilized beyond the initial screening tests. If a biopsy is indicated, a 10 to 12-core TRUS guided peripheral zone prostate biopsy incorporating the anterior horn area should be performed. For men diagnosed with screen-detected PCa, tumour volume, grade, DRE and PSA results direct management. Selectively treating patients with favourable risk PCa may significantly improve screening outcomes.
Table 1. Comparisons between the PLCO, ERPSC and Goteborg studies.

<table>
<thead>
<tr>
<th></th>
<th>PLCO(^1)</th>
<th>ERSPC(^2)</th>
<th>Goteborg(^6)</th>
</tr>
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<tbody>
<tr>
<td>N</td>
<td>76,693</td>
<td>162,243</td>
<td>20,000</td>
</tr>
<tr>
<td>Age</td>
<td>55-74 (13% &gt; 70)</td>
<td>55-69</td>
<td>50-64</td>
</tr>
<tr>
<td>Site</td>
<td>Multiple centers USA</td>
<td>7 Countries</td>
<td>1 City (Goteborg, Sweden)</td>
</tr>
<tr>
<td>Methods</td>
<td>PSA &gt; 4 ng/ml Abnormal DRE</td>
<td>PSA &gt; 3 ng/ml Abnormal DRE</td>
<td>PSA &gt; 2.5 ng/ml (From 2005 on) PSA &gt;2.9 ng/ml (From 1999-2004) PSA &gt; 3.4 ng/ml (From 1995-98)</td>
</tr>
<tr>
<td>FUP</td>
<td>Every 1 y x 6 11 y median FUP</td>
<td>Every 4 y 9 y (complete)</td>
<td>Every 2 years 78% had 14 y FUP</td>
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<tr>
<td>Compliance</td>
<td>85%</td>
<td>82%</td>
<td>76%</td>
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<tr>
<td>Contamination</td>
<td>52%</td>
<td>Not known</td>
<td>3%</td>
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<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Screened</th>
<th>Control</th>
<th>Screened</th>
<th>Control</th>
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<tr>
<td>Prostate Cancers</td>
<td>7.3%</td>
<td>6%</td>
<td>8.2%</td>
<td>4.8%</td>
<td>7.2%</td>
<td>11.4%</td>
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<tr>
<td>Prostate Cancer Deaths</td>
<td>50</td>
<td>44</td>
<td>326</td>
<td>214</td>
<td>78</td>
<td>44</td>
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<tr>
<td>Risk Ratio</td>
<td>NS</td>
<td>20% (p = 0.04)</td>
<td>44% (p =0.002)</td>
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<tr>
<td>NNS</td>
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<td>1:293</td>
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References:


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