Systematic Review & Meta-Analysis of the Diagnostic Accuracy of Prostate Specific Antigen (PSA) for the Detection of Prostate Cancer in Symptomatic Patients

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Abstract

Background

Prostate Specific Antigen (PSA) is a commonly used test to detect prostate cancer. Attention has mostly focused on the use of PSA in screening asymptomatic patients, but the diagnostic accuracy of PSA for prostate cancer in patients with symptoms is less well understood.

Methods

A systematic database search was conducted of Medline, EMBASE, Web of Science, and the Cochrane library. Studies reporting the diagnostic accuracy of PSA for prostate cancer in patients with symptoms were included. Two investigators independently assessed the titles and abstracts of all database search hits and full texts of potentially relevant studies against the inclusion criteria, and data extracted into a proforma. Study quality was assessed using the QUADAS-2 tool by two investigators independently. Summary estimates of diagnostic accuracy were calculated with meta-analysis using bivariate mixed effects regression.

Results

563 search hits were assessed by title and abstract after de-duplication, with 75 full text papers reviewed. 19 studies met the inclusion criteria, 18 of which were conducted in secondary care settings (one from a screening study cohort). All studies used histology obtained by Transrectal Ultrasound guided biopsy (TRUS) as a reference test, usually only for patients with elevated PSA or abnormal prostate examination. Pooled data from 14,489 patients found estimated sensitivity of PSA for prostate cancer was 0.93 (95% CI 0.88, 0.96) and specificity was 0.20 (95% CI 0.12, 0.33). The area under the receiving-operator characteristic curve was 0.72 (95% CI 0.68, 0.76). All studies were assessed as having a high risk of bias in at least one QUADAS-2 domain.

Conclusions

Currently available evidence suggests PSA is highly sensitive but poorly specific for prostate cancer detection. However, significant limitations in study design and reference test reduces the certainty of this estimate. There is very limited evidence for the performance of PSA in primary care, the healthcare setting where most PSA testing is performed.

Introduction

Prostate specific antigen (PSA) is a commonly used test for the detection of prostate cancer(1). PSA testing is usually performed for one of two reasons: assessing a patient presenting to their GP or primary care physician with lower urinary tract symptoms (LUTS), or opportunistic screening for a patient who is asymptomatic but concerned about their risk of prostate cancer. Patients with an elevated PSA are usually referred to a Urologist for diagnostic testing, which may include magnetic resonance imaging (MRI) of the prostate and/or a prostate biopsy. Very large randomised controlled trials of PSA-based prostate cancer screening have been performed, summarised in a recent systematic review in 2018 that showed a small potential reduction in prostate cancer specific mortality with no change in all-cause mortality and an increased risk of complications from biopsy, overdiagnosis of clinically insignificant prostate cancer, and overtreatment(2). However, uncertainty remains about the diagnostic accuracy of PSA for prostate cancer in patients with LUTS(3).

The most recent systematic review of the diagnostic accuracy of PSA was published by Harvey et al in 2009(4). A range of estimates for the accuracy of PSA was found amongst the ten included studies. That review presented limited information on their methods; crucially, it was unclear whether the included studies were assessing PSA in symptomatic or asymptomatic patients, nor was it clear whether any were relevant to primary care populations. Just et al published a brief review of the literature in 2018, highlighting that the paucity of research in this area applicable to primary care, where a significant proportion of PSA testing is performed, still remains(3).
This systematic review aimed to determine the diagnostic accuracy of PSA for the detection of prostate cancer in patients, focusing on studies where the included patients (or a subset of included patients) had at least one symptom that could relate to an undiagnosed prostate cancer. Given the findings by Just et al, this review considered studies from primary and secondary care settings.

**Methods**

**Types of studies**

We included cross-sectional and cohort studies that reported paired data on the diagnostic accuracy of PSA for the detection of prostate cancer in symptomatic men, verified with the use of a reference test (prostate biopsy). We excluded studies if it was not possible to extract data for a complete two-by-two table for the target condition, or if the patient cohort was only asymptomatic patients (i.e. a screening cohort). We did not restrict studies by publication date, country, or clinical setting.

**Participants**

The study population of interest was any patient with symptoms of a possible prostate cancer, with no history of the disease. We defined symptoms of prostate cancer as at least one of: LUTS (nocturia, hesitancy, poor stream, incomplete voiding, double voiding, terminal dribbling, urgency, incontinence, frequency), haematuria, erectile dysfunction, or lower back pain. Symptoms may have been identified by a standardised tool, such as the International Prostate Symptom Score (IPSS), clinical coding, or through patient self-report. We did not exclude studies based on age of participants or study setting. Where studies included groups of both asymptomatic and symptomatic men we included men in the symptomatic group.

**Index test**

The index test was Prostate Specific Antigen (PSA) in a peripheral blood sample, measured in nanograms per millilitre (ng/mL). We did not set an *a priori* abnormal PSA level, but instead extracted data based on the definition of abnormal PSA used in each study.

**Target condition**

The target condition was prostate cancer, regardless of Gleason grade or clinicopathological stage.

**Reference test**

The reference test was a biopsy of the prostate with histological examination. We did not set an inclusion criteria on the basis of prostate biopsy approach used in studies, but this was recorded as part of the data extraction.

**Electronic searches**

Medline Ovid, EMBASE, the Cochrane Central Register of Controlled Trials (CENTRAL) and Web of Science databases were utilised to identify relevant studies. Key search terms, informed by the Scottish Intercollegiate Guidelines Network (SIGN) search strategies and pre-existing systematic reviews in the field of prostate cancer, were combined with MeSH terms for each database search. Hand-searching of reference lists from included studies and snowballing techniques were performed to locate any other possibly relevant studies.

**Data collection and analysis**

**Selection of studies**

Search hits from each database were downloaded and combined into a review database managed in Mendeley Desktop. Each search hit was screened against the inclusion/exclusion criteria by SM and a 2nd investigator (LP, SC, or EG) independently, based on title and abstract. Full text articles were reviewed if a reviewer was unclear on the basis of title and abstract. Any discrepancies of study inclusion were adjudicated by a third reviewer (WH or AS).
Data extraction

A pre-prepared proforma for data extraction was used to collate relevant data from each included study, including two by two tables for the index and reference tests. SM extracted the data from all included studies. A second investigator extracted data from a random sample of 10% of included studies for verification of accuracy of data extraction. Any discrepancies were adjudicated by a third reviewer (WH or AS).

Quality assessment

Risk of bias and applicability of all included studies was assessed by SM using the QUADAS-2 tool, with a second investigator independently assessing 10% of included studies and discussed any discrepancies with SM.

Meta-analysis

Raw data extracted from included papers on PSA result and prostate cancer diagnoses were extracted and combined into 2 x 2 tables to assess diagnostic accuracy. Measures of pooled diagnostic accuracy were determined for the following outcomes using bi-variate mixed effects regression:

- Any prostate cancer diagnosis
- Clinically significant prostate cancer diagnosis (Gleason Grade Group ≥ 2)

The majority of included studies used a fixed PSA threshold of 4ng/mL, and this was also used as the threshold for meta-analysis. No included studies reported sufficient information to Meta-analyse age-adjusted thresholds.

Heterogeneity

Heterogeneity was assessed for visually using Forest plots of sensitivity and specificity.

All analyses were performed using Stata Version 16 (StataCorp, http://www.stata.com)

Protocol publication

The protocol for this systematic review and meta-analysis was registered with PROSPERO (CRD42021257783).

PRISMA reporting guidelines

This systematic review was conducted following the PRISMA reporting guidelines for systematic reviews and meta-analyses(6).

Results

Database searching identified 631 potentially relevant studies, and a further 42 studies were identified through reference list checking and snowballing techniques from initial search hits and key papers. Following de-duplication, 563 search hits were assessed by two reviewers independently, and 75 papers selected for full text assessment. 19 papers were ultimately included. Details of full-text exclusions can be found in figure 1 below.

Risk of bias assessment using the QUADAS-2 tool demonstrated a number of potential areas of bias in the included studies (see table 2 and figure 2). None of the studies were assessed as having a low risk of bias with regards to the reference standard test, which was almost always a Transrectal Ultrasound-guided (TRUS) biopsy. TRUS biopsy suffers from a significant risk of false negative or misclassification of prostate cancer diagnosis owing to the random nature of sampling of the prostate(7). The reference standard was performed with knowledge of the index test (PSA) in 16 of 19 studies. Patient populations were drawn from hospital urology clinics in all but one study, affecting applicability to other clinical settings. Limited information with regards to patient selection was available in eight studies, and the majority had a low risk of bias with regards to the conduct of the index test.
| Study              | Risk of Bias | Applicability Concerns |
|-------------------|--------------|-----------------------|
|                   | Patient Selection | Index Test | Reference Standard | Flow and Timing | Patient Selection | Index Test | Reference Standard |
| Abirabbi et al    | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Aghbash et al     | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Jagoda et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Chang et al       | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Chavan et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Gales et al       | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Hofer et al       | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Lee et al         | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Magistro et al    | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Meigs et al       | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Nordstrom et al   | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Painel et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Pape et al        | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Rashid et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Richie et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Sore et al        | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Shahab et al      | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Teurs et al       | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |
| Wyngaard et al    | ?            | ?                     | ?                   | ?               | ?               | ?               | ?               |

- Low Risk
- High Risk
- Unclear Risk
| First Author          | Year | Country | Number of patients | Mean age (range)* | Setting | PSA range | Stage/Grade data | Reference test                  |
|-----------------------|------|---------|--------------------|-------------------|---------|-----------|------------------|--------------------------------|
| Abdrabo et al (8)     | 2011 | Sudan   | 118                | 70 years (56 – 83)| One hospital urology clinic | 2.5-10ng/mL | No               | TRUS biopsy                |
| Agnihotri et al (9)   | 2014 | India   | 875 biopsied (of 4,702 patients) | 66 years (50 – 75) | One hospital urology clinic | Any | No               | TRUS biopsy                |
| Aragona et al (10)    | 2005 | Italy   | 3,171 biopsied (of 16,298 patients) | 62 years (40 – 75) | 15 hospital urology clinics | Any | Clinical TNM staging | TRUS biopsy                |
| Chang et al (11)      | 2015 | Taiwan  | 225                | PCa 72 years; BPH 67 years | One hospital urology clinic | Any | TNM stage and Gleason Score | TRUS biopsy                |
| Chavan et al (12)     | 2009 | India   | 440 biopsied (of 922 patients) | 64 years (40 – 95) | One tertiary hospital urology clinic | Any | No               | TRUS biopsy                |
| Galic et al (13)      | 2003 | Croatia | 88 biopsied (of 944 patients) | ≥50 years | Recruited from two villages to attend hospital clinic | Not stated | No               | TRUS biopsy                |
| Hofer et al (14)      | 2000 | Germany | 188                | PCa 70 years; BPH 68 years | One hospital urology clinic | Any | No               | TRUS biopsy / TURP/ non-cancer surgery |
| Lee et al (15)        | 2006 | Korea   | 201                | 63 years | One hospital urology clinic | < 4ng/mL | No               | TRUS biopsy                |
| Magistro et al (16)   | 2020 | Germany | 1,125              | 70 years | One hospital urology clinic | Any | TNM stage and Gleason Score | HoLEP (+ mpMRI with targeted and systemic biopsy for some patients) |
| Meigs et al (17)      | 1996 | USA     | 1,524              | 50-79 years | One hospital urology clinic + two BPH study cohorts | Any | Clinical T stage | TRUS biopsy / TURP/ non-cancer surgery |
| Nordstrom et al (18)  | 2021 | Sweden  | 1,554              | 64 years (50 – 69) | Population-based screening study cohort | >3ng/mL | TNM stage and Gleason Score | TRUS biopsy                |
| First Author     | Year | Country   | Number of patients | Mean age (range)* | Setting                                      | PSA range | Stage/Grade data | Reference test          |
|------------------|------|-----------|--------------------|-------------------|----------------------------------------------|-----------|------------------|-------------------------|
| Patel et al (19) | 2009 | UK        | 647 biopsied (of 3,976 patients) | 65 years (15 – 91) | One hospital urology clinic                  | Any       | No               | TRUS biopsy              |
| Pepe et al (20)  | 2007 | Italy     | 403 biopsied (of 13,294 patients) | 62 years (40 – 75) | Two hospital urology clinics                | <4ng/mL   | Pathological T stage | TRUS biopsy              |
| Rashid et al (21)| 2012 | Bangladesh | 206               | >50 years         | One hospital urology clinic and one nursing home | >2.5ng/mL | No               | TRUS biopsy              |
| Richie et al (22)| 1993 | USA       | 1,167 biopsied (of 6,630 patients) | 63 years (50 – 96) | Six medical centres                         | Any       | TNM stage and Gleason Score | TRUS biopsy              |
| Seo et al (23)   | 2007 | Korea     | 4,967              | 66 years (40 – 96) | 25 hospital urology clinics                | Any       | No               | TRUS biopsy              |
| Shahab et al (24)| 2013 | Indonesia | 404                | 64 years (34 – 84) | One hospital urology clinic                | Any       | TNM stage and Gleason Score | TRUS biopsy              |
| Tauro et al (25) | 2009 | India     | 100                | 68 years          | One hospital urology clinic                | Any       | No               | TRUS biopsy              |
| Wymenga et al (26)| 2000| The Netherlands | 716              | Not reported      | Two hospital urology clinics                | Any       | Clinical T stage | TRUS biopsy / TURP / prostatectomy |

PSA – Prostate Specific Antigen; TRUS – Transrectal Ultrasound guided biopsy; PCa – Prostate cancer; BPH – Benign Prostatic Hypertrophy; TNM – Tumour-node-metastasis; TURP – Transurethral resection of the prostate; HoLEP – Holmium laser enucleation of the prostate; mpMRI – multiparametric magnetic resonance imaging

* Age range and/or mean not present in table if not reported

Table 3 summarises the features of the included studies. There was a wide range of countries and study sizes. One study focused on a symptomatic cohort within a population screening study, and the remainder were set in hospital urology clinics. No study was performed in a primary care population. Five studies gathered stage and grade data. All but one study used TRUS biopsy as a reference test, with three studies also gathering diagnostic data from Transurethral Resection of the Prostate (TURP) or other urological surgical procedures involving the prostate.

Table 4 shows the measures of diagnostic accuracy calculated using reported data in 14 included studies featuring 14,489 patients that considered a PSA level of greater than or equal to 4ng/mL as abnormal. The remaining five studies focused on populations in a specific part of the PSA range – either a low or raised PSA level. Meta-analysis showed an estimated combined sensitivity of a PSA greater than or equal to 4ng/mL for any prostate cancer of 0.93 (95% CI 0.88, 0.96) and a
combined specificity of 0.20 (95% CI 0.12, 0.33) (See figure 3). There was significant heterogeneity between included studies (sensitivity $I^2$ 98.97, specificity $I^2$ 99.61). Receiver Operator Curve (ROC) analysis showed an AUC of 0.72 (95% CI 0.68, 0.76) (See figure 4).

Three studies included in the meta-analysis collected stage and grade data for prostate cancer cases; however, none of these studies reported data for clinically significant prostate cancer diagnoses at a PSA cut-off of $\geq 4$ng/mL. Chang et al(11) did not report the accuracy of PSA, but showed a statistically significant difference in free:total PSA ratio for a Gleason Score of seven or more compared to Gleason Score of six or lower ($11.69 +/- 0.98$ vs $16.47 +/- 2.25$, p = 0.029). Richie et al(22) did not report the Gleason Score data collected, but found higher PSA levels and increasing age were associated with a higher risk of metastatic prostate cancer. Shahab et al(24) identified a PSA cut-off of 6.95ng/mL for differentiating moderate versus high Gleason Score (which was not defined).

### Table 4

| Author  | Year | Sensitivity | Specificity | Positive Predictive Value | Negative Predictive Value |
|---------|------|-------------|-------------|---------------------------|---------------------------|
| Abdrabo | 2011 | 0.92        | 0.24        | 0.35                      | 0.87                      |
| Agnihotri| 2014 | 0.99        | 0.05        | 0.59                      | 0.80                      |
| Aragona | 2005 | 0.92        | 0.15        | 0.38                      | 0.76                      |
| Chang   | 2015 | 0.89        | 0.09        | 0.19                      | 0.76                      |
| Chavan  | 2009 | 0.96        | 0.03        | 0.18                      | 0.79                      |
| Galic   | 2003 | 0.91        | 0.32        | 0.47                      | 0.85                      |
| Hofer   | 2000 | 0.92        | 0.29        | 0.46                      | 0.85                      |
| Meigs   | 1996 | 0.61        | 0.74        | 0.34                      | 0.89                      |
| Rashid  | 2012 | 0.72        | 0.46        | 0.28                      | 0.85                      |
| Richie  | 1993 | 0.82        | 0.48        | 0.31                      | 0.90                      |
| Seo     | 2007 | 0.98        | 0.04        | 0.33                      | 0.87                      |
| Shahab  | 2013 | 0.98        | 0.19        | 0.13                      | 0.98                      |
| Tauro   | 2009 | 1.00        | 0.38        | 0.40                      | 1                         |
| Wymenga | 2000 | 0.95        | 0.16        | 0.44                      | 0.82                      |

### Discussion

#### Summary of findings

Published studies assessing the diagnostic accuracy of PSA in symptomatic patients reported high sensitivity and low specificity for the detection of prostate cancer. 18 of the 19 included studies were undertaken in hospital urology outpatient populations, with one study focused on a symptomatic cohort within a population screening study. Importantly, there were no studies assessing the performance of PSA in a primary care population. Insufficient data was available to assess the diagnostic accuracy of PSA for clinically significant prostate cancer. Furthermore, all included studies had a high risk of bias in at least one QUADAS domain.
Comparison to existing literature

Harvey et al.(4) published a systematic review of the diagnostic accuracy of PSA for prostate cancer in European populations, focused on studies published between 1998 and 2008. Individual study level data from 10 included papers was reported, though without estimating a combined level of accuracy. They considered the accuracy of PSA for all prostate cancer types overall, and showed a range of accuracy estimates similar to this study. Over half of the studies included in this review were published since the review by Harvey et al. A review of clinical features of prostate cancer in primary care by Young and colleagues(27) in 2015 identified one study from 1989 of 287 patients referred from primary care with bladder outlet obstruction, of whom 211 had a PSA test. High levels of sensitivity (89.5%) and specificity (90%) were reported, but Young and colleagues considered the true level of accuracy was likely to be lower given few patients with a normal PSA level had the reference test for prostate cancer.

Strengths & weaknesses

This study benefited from a rigorous, focused, methodological approach in conducting the review. All clinical settings were eligible, ensuring we found as many relevant studies as possible. Most included studies employed PSA in a similar manner, using similar indications and diagnostic thresholds, allowing for cross-study comparisons.

The evidence for the association between lower urinary tract symptoms and prostate cancer, in particularly clinically significant prostate cancer, is equivocal. A number of secondary care studies suggest that symptoms do not discriminate well between prostate cancer and benign prostatic hypertrophy(28,29). This assumption is largely untested in primary care populations, and contrasts with studies showing that the majority of patients diagnosed with prostate cancer present to their GP with LUTS prior to diagnosis(30–33). This controversy also means that LUTS and other relevant symptoms may not be reported or be the focus of some potentially relevant studies of PSA for prostate cancer, and may have limited the sensitivity of the search strategy employed. However, key papers were picked up by the database searches and the majority of PSA studies will likely be focused on screening in asymptomatic populations.

All included studies employed TRUS biopsy as a reference test, with some also including pathological data obtained from urological procedures on the prostate. TRUS biopsy is recognised as having poor sensitivity as a diagnostic test(34), owing to the inability to visualise lesions within the prostate resulting in a random sampling of the gland, and thus misclassification bias. Most included studies only performed the reference test on patients with a raised PSA or abnormal prostate examination, introducing partial verification bias. Therefore, the true sensitivity of PSA in symptomatic patients is unknown and likely to be lower than reported.

Implications for research & practice

PSA is a commonly used test to assess for the presence of prostate cancer, mostly in a primary care setting, and is recommended as part of the assessment of patients with LUTS in national guidelines(35–37). The lack of primary care evidence for the use of PSA to detect prostate cancer is known, and is not the only condition for which secondary care evidence has been applied to primary care guidance(38). Even so, this is a major gap in knowledge, as spectrum bias means that secondary care data (or screening data) do not translate to primary care. High quality studies in primary care populations are needed to fill this gap, and future studies should report not just on prostate cancer per se, but on clinically significant cancer as well. The introduction of more accurate diagnostic tests for prostate cancer, including multiparametric magnetic resonance imaging (34), increases the need for better understanding of the role of PSA in the early detection of symptomatic prostate cancer.

Primary care clinicians are generally aware of the limitations of PSA testing(39), and clinical guidelines encourage a balanced discussion with patients of the potential benefits and harms of relying on PSA to detect prostate cancer(40,41). The findings of this review suggest this is a pragmatic approach in providing care to patients with LUTS. Alternative tests to PSA have been extensively researched(42,43), and some show promise of improving the level of confidence in detecting prostate cancer, though none has entered primary care practice as yet.
Conclusions

Published evidence from almost entirely secondary care based studies suggests that PSA has high sensitivity and low specificity for the diagnosis of prostate cancer in symptomatic patients. Published studies suffer from a number of biases, which probably overestimate the accuracy of PSA, and there were no included studies assessing the accuracy of PSA in a primary care population. The utility of PSA for the diagnosis of clinically significant prostate cancer in primary care remains unclear and needs urgent study. A major focus of such a study would be to identify patients with clinically significant cancer, warranting radical treatments, whilst avoiding exacerbating the issue of overdiagnosis of clinically insignificant prostate cancer.

Declarations

Conflict of interest statement – all authors confirm they have no relevant conflicts of interest to declare.

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Tables

Table 1 is not available with this version.

Figures

PRISMA 2020 flow diagram for new systematic reviews which included searches of databases and registers only

From: Page AU, Melcombe JE, Booth M, Baltzell G, Hofmann T, Vickers CD, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. BMJ 2021;372:n71. doi: 10.1136/bmj.n71

Figure 1

PRISMA 2020 flow diagram
Figure 2

Summary of QUADAS-2 risk of bias assessments

Figure 3

Forest plot of included studies using PSA cut-off of 4ng/mL
Figure 4

Receiver Operator Curve (ROC) of included studies using PSA cut-off of 4ng/mL

Supplementary Files

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