Prostate health index is useful for prostate cancer detecting in Chinese people

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Background: Prostate cancer (PCa) incidence was increased substantially in the last two decades in China. We evaluated the performance of prostate health index (PHI) in predicting the presence of PCa in Chinese people.

Methods: A total of 108 consecutive patients were recruited to develop a PHI-based nomogram to predict PCa. Serum total prostate specific antigen (tPSA), free PSA (fPSA), and p2PSA were measured by Beckman Coulter’s DxI 800 Immunoassay System. PHI was calculated by \[\text{[p2PSA/fPSA]} \times \text{tPSA} \]. Performance of individual PSA and PHI measurements in discriminating clinical outcomes was measured using the detection rate of PCa, sensitivity, specificity, and area under the receiver operating characteristic curve (AUC).

Results: Among all patients, PHI was a better predictor of PCa compared with tPSA. PHI achieved the highest AUC of 76.6% in discriminating PCa from non-PCa compared to tPSA measurement, the difference of AUC area between PHI and PSA was about 6% in the entire cohort. The PHI-based nomogram reduced the number of biopsies. With the cutoff value (tPSA >4 ng/mL, PHI >40), there were 37 patients need to do biopsies in tPSA >4 ng/mL group, while with PHI >40, only 30 patients need to do. Seven patients could avoid unnecessary biopsies. When combined with the two values (tPSA >4 ng/mL, PHI >40) only 24 patients remained.

Conclusions: Our finding has implication PHI and provides added value over tPSA for PCa detecting in Chinese people.

Keywords: Prostate health index (PHI); prostate specific antigen (PSA); prostate cancer (PCa)

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(p2PSA), a truncated PSA isoform, was demonstrated better accuracy than serum tPSA and fPSA in detecting PCa. The prostate health index (PHI), derived from a combination of p2PSA, tPSA, and fPSA (free PSA) had been shown to be better predictor of PCa (9-14), had been approved by the United States Food and Drug Administration (FDA) and the European Medicines Agency (EMA). There were many papers reported the PHI is a better predictor of PCa in patients from Europe and American (1,6,10,12-14), but few papers in Chinese people. In this paper, we would using an observational prospective study design to test whether PHI could aid in clinical practice.

### Methods

**Patients and tissue samples**

A total of 108 consecutive patients which visited doctors between 2014 and 2016 in No. 1 affiliated Hospital of Soochow University were included in this study. After written informed consent, PHI and PSA was detected for each patient before any study-specific investigation was performed. Transrectal ultrasound (TRUS)-guided biopsy was performed using a 12-core scheme. All biopsy specimens were reviewed by two doctors at the Pathology Department. This study was approved by the ethic committees at local hospitals.

**Laboratory methods**

Beckman Coulter’s DxI800 Immunoassay System was using to detecting Serum p2PSA, tPSA and fPSA of each patient. PHI was calculated by \((\frac{\text{p2PSA}}{\text{fPSA}}) \times \sqrt{\text{tPSA}}\). Performance of individual PSA and PHI measurements in discriminating clinical outcomes was measured using the detection rate of PCa, sensitivity, specificity, and the area under the receiver operating characteristic curve (AUC) (1,6).

**Statistical analysis**

Statistical analyses were conducted using the Statistical Package of the Social Sciences software version 19.0 (SPSS, Inc.). A two-sided P<0.05 was considered to be statistically significant.

### Results

A total 108 patients were recruited in this research. Baseline characteristic of all patients were shown in Table 1. Most of the patients were aged between 60 and 70 years, mean of all patients age was 69.4 [47–87] years.

Among all people, 51 patients were diagnosed with PCa, 57 patients were absence of PCa, mean of age in PCa group is 69.9, while in absence of group it is 68.9. fPSA was 3.8 vs. 1.9 ng/mL. No different with age and fPSA. PHI and tPSA were higher in PCa group, in the group of absence of PCa, mean of tPSA was 10.3 ng/mL, in PCa group was 30.3 ng/mL (P<0.05). Similar as tPSA, PHI was higher in PCa group, in PCa group, mean of PHI was 233.2 ng/mL, while in absence of PCa, there was 46.5 ng/mL only (P<0.05).

To get the more effect marker in predicted PCa, the PHI, PHI + tPSA and tPSA were selected. When we choose 60% sensitivity, the specificity for PSA to detect PCa was 68%, while the PHI and tPSA + PHI was 86%, 87.7% respect. When sensitivity was 70%, the specificity of PSA was 56.1%, PHI and tPSA + PHI were decrease to 75.4% and 77.2%. With 80% sensitivity, the specificity of PSA was 38.6%, PHI and PHI + PSA were 45.6% (Table 2). All that means the specificity of PHI or PHI + PSA were higher than PSA.

Compared with tPSA measurement, AUC was always higher for PHI. In detecting PCa, AUC of PHI was 76.6%, tPSA + PHI was 76.7%, while tPSA was only 70.8% (compare with 50%, P<0.05), the difference of AUC between PHI and

### Table 1 Characteristics of study objects

| Characteristics | Overall, n=108 | Absence of PCa, n=57 (52.8%) | Presence of PCa, n=51 (47.2%) | P value |
|-----------------|----------------|-----------------------------|-----------------------------|---------|
| Age, yr         | 69.4±6.8 [47–87] | 68.9±7.4                    | 69.9±6.2                    | >0.05   |
| tPSA, ng/mL     | 20.6±4.24 (2.23–149) | 10.3±8.4                    | 30.3±38.4                   | <0.05   |
| fPSA, ng/mL     | 2.88±4.2 (0.03–20)  | 1.9±1.6                     | 3.8±5.6                     | >0.05   |
| PHI             | 142.8±340.8 (6.85–2,986.2) | 46.5±17.5                  | 233.2±458.2                 | <0.05   |

PCa, prostate cancer; tPSA, total prostate specific antigen; fPSA, free prostate specific antigen; PHI, prostate health index.
We also evaluated the benefit of PHI over tPSA in reducing the number of biopsies. As Table 3 show, in 57 PCa negative patients, there were 37 patients need to do biopsies with the cutoff value of tPSA >4 ng/mL; while with the value of PHI >40, only 30 patients need do. Seven patients could avoid unnecessary biopsies. When combined with the two values (tPSA >4 ng/mL, PHI >40) only 24 patients remained (P<0.05).

**Discussion**

Although reasons were unclear, the PCa incidence and its trend were differences in Western countries and China. The incidence in China was lower than Western counties. More PCa patients in first diagnosed were high-grade PCa in China compared with Western country. Furthermore, the gray zone is different in Chinese patients from Western country (2,6,15). So some useful marker in detecting PCa in Western country need to validated in Chinese people, such as PHI. Derived from a combination of p2PSA, tPSA, and fPSA, PHI had been shown to be a better predictor of PCa in Western country. Similar as the results of Western country, we find that PHI performed better than tPSA in PCa detecting. The specificity of PHI and PHI + tPSA is higher than tPSA. The PHI achieved an AUC 76.7% while tPSA was 70.8%. All those date meanings that PHI-based strategy can reduce the number of unnecessary biopsies while maintaining the same detection rate of PCa.

Although additional muti-center with larger sample size need, our results identified the PHI can be used as a new reference standard for PCa detecting in China.

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**Footnote**

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at http://dx.doi.org/10.21037/tcr.2019.05.08). The authors have no conflicts...
of interest to declare.

**Ethical Statement:** The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The institutional research ethics committee approved the experimental protocols. Informed consent was obtained from all patients.

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