Modified J-CAPRA scoring system in predicting treatment outcomes of metastatic prostate cancer patients undergoing androgen deprivation therapy

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Abstract
The J-CAPRA score is an assessment tool which stratifies risk and predicts outcome of primary androgen deprivation therapy (ADT) using prostate-specific antigen, Gleason score, and clinical TNM staging. Here, we aimed to assess the generalisability of this tool in multi-ethnic Asians. Performance of J-CAPRA was evaluated in 782 Malaysian and 16,946 Japanese patients undergoing ADT from the Malaysian Study Group of Prostate Cancer (M-CaP) and Japan Study Group of Prostate Cancer (J-CaP) databases, respectively. Using the original J-CAPRA, 69.6% metastatic (M1) cases without T and/or N staging were stratified as intermediate-risk disease in the M-CaP database. To address this, we first omitted clinical T and N stage variables, and calculated the score on a 0–8 scale in the modified J-CAPRA scoring system for M1 patients. Notably, treatment decisions of M1 cases were not directly affected.

Jasmine Lim and Shiro Hinotsu contributed equally to this work.

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1 | INTRODUCTION

Prostate cancer incidence varies markedly across the globe, attributing to population genetics, dietary intake, access to healthcare, local screening program, and diagnostic practices. Similar trend was observed in Asia. For instance, the incidence rate of prostate cancer in Japan (age-standardised rate, ASR 35.4 per 100,000) was higher than Malaysia (ASR 12.4 per 100,000). However, Malaysian had a higher mortality rate (ASR 5.6 per 100,000) compared to Japanese (ASR 4.4 per 100,000), resulting from ~60% of patients diagnosed at locally advanced and metastatic stages in Malaysia as well as limited access to survival-prolonging treatments.

Androgen-deprivation therapy (ADT) is the mainstay of first-line treatment for metastatic prostate cancer. Combination of radical prostatectomy or radiotherapy with ADT improved the survival of patients with high-risk localised or locally advanced disease. Of note, high-risk localised prostate cancer includes patients with prostate-specific antigen (PSA) >20 ng/ml or Gleason score >7 (Gleason Grade Group 4/5) or cT2c, while those with cT3-4 or cN+ (any PSA and Gleason score) diseases are classified as locally advanced. High proportion of advanced disease entails special challenge in predicting the disease risk of patients receiving ADT.

The J-CAPRA (Japan Cancer of the Prostate Risk Assessment) score is a novel, validated risk assessment tool in predicting outcomes of ADT in prostate cancer patients. This multivariable risk assessment tool was constructed from Japan Study Group Prostate Cancer (J-CaP) 2001–2003 database, which is a national registry of prostate cancer patients undergoing ADT in Japan. It is applicable to the entire spectrum of risk and stage, both localised and advanced prostate cancer. Therefore, the objective of this study was to assess the generalisability of J-CAPRA scoring system in stratifying the risk of prostate cancer patients from a multi-ethnic Asian population.

2 | MATERIALS AND METHODS

The M-CaP comprised patients accrued from nine urology referral centers across Malaysia, with 15.2% of patients treated at academic center. A total of 1152 men were recruited into the cohort between 2016 and May 2018, of whom 578 underwent primary ADT and 204 received ADT combined with radiation, surgery or chemotherapy. Type of ADT was categorised into orchidectomy, luteinizing hormone-releasing hormone (LHRH) agonist monotherapy, or LHRH agonist with anti-androgen (bicalutamide 50 mg daily) [maximum androgen blockade (MAB)]. Men receiving anti-androgen monotherapy were excluded from this study. Data on initial and subsequent PSA levels, TNM staging, treatment patterns, disease progression, and mortality were documented prospectively with a written form (proforma) based on medical, radiological, and pathological records. Details from the proforma were then transformed into an electronic database for further analysis. Patients were followed under specialist review every 3 months.

The J-CaP 2001–2003 database contained 26,272 prostate cancer patients diagnosed during 2001–2003 and received ADT, either as primary treatment or in combination with radiotherapy or surgery across 384 institutions in Japan. The database represented 95% of Japanese patients treated with primary ADT in the academic centers or community hospitals. Clinical stage, treatment patterns, disease progression, all-cause, and prostate cancer-specific mortality (CSM) were ascertained from participating urologists on a quarterly basis. In this study, 16,946 patients treated with primary ADT were included for analysis. Men receiving anti-androgen as monotherapy were excluded. Ethical approval for the M-CaP and J-CaP databases was granted by the Medical Research and Ethics Committee (MREC), Ministry of Health Malaysia, and regional Medical Research Review Board, respectively.
Clinical and demographic characteristics were compared between M-CaP and J-CaP 2001–2003 databases with Student’s t-test and chi-square test, as appropriate. Of note, the J-CaP 2001–2003 was used in the initial development of J-CAPRA score. The disease risk of each man in M-CaP was assessed based on the J-CAPRA. The validated J-CAPRA scoring system uses a 12-point scale based on five parameters including PSA level (up to 3 scores), biopsy Gleason score (up to 2 scores), clinical T stage (up to 3 scores), clinical N stage (up to 1 score), and clinical M stage (up to 3 scores). Validated score groups were introduced to stratify low-risk (J-CAPRA score 0–2), intermediate-risk (J-CAPRA score 3–7), and high-risk (J-CAPRA score ≥8) groups.

Performance of J-CAPRA was evaluated across all disease stages using M-CaP and J-CaP 2001–2003 databases. First, we examined the risk scores distribution of M-CaP and J-CaP 2001–2003 databases. Second, we assessed the differences in risk scores distribution between J-CaP and M-CaP across Union for International Cancer Control (UICC) stage I-IV groups. The J-CAPRA was further modified by omitting T and N stage score points for the M1 group. To validate the J-CAPRA modifications for the M1 group, overall survival, prostate cancer-specific survival, and progression-free survival of J-CaP 2001–2003 were compared between risk categories defined by original J-CAPRA and modified J-CAPRA, through Kaplan–Meier survival curves and Cohen’s coefficient of agreement. Overall survival, prostate cancer-specific survival, and progression-free survival were defined as time from initiation of ADT to death, to death from disease, and to the first event of radiological and/or biochemical progression or death, respectively. Statistical analysis was performed using SPSS for Windows version 21.0 (SPSS Inc.). Two-tailed p value <0.05 was termed as statistically significant.

### RESULTS

There were 16,946 patients in J-CaP and 782 patients in M-CaP receiving ADT, either as primary ADT or in combination with radiotherapy, surgery or chemotherapy. Summary of clinical characteristics of both J-CaP and M-CaP cohorts is presented in Table 1. Patients receiving ADT in J-CaP were older than those in M-CaP with an average age of 75 ± 7.2 years versus 69.6 ± 7.7 years (p < 0.01, Student’s t-test). In M-CaP, there was a higher burden of comorbidity compared to the J-CaP cohort (p < 0.01, chi-square test). Maximum androgen blockade (67%) was the most common treatment, followed by orchidectomy (23.2%) and LHRH agonist monotherapy (9.8%) in the J-CaP cohort. The M-CaP cohort had relatively higher-risk disease compared to J-CaP cohort, with a median PSA level of 99 ng/ml (interquartile range, IQR 31.1–352.5 ng/ml) versus 27.2 ng/ml (IQR 10.5–109.2 ng/ml), a median biopsy Gleason score of 8 versus 7 and a higher percentage of UICC stage IV cases at 74.1% versus 35.4%.

For J-CAPRA scoring, a total of 734 men in M-CaP with complete risk stratification data including PSA level, biopsy Gleason Score, and disease staging were recruited into the analysis. The J-CAPRA score distribution of M-CaP is summarised in Figure 1. Most (60.9%) had a J-CAPRA score of 3–7, while 31.7% had a score greater than 8 and 7.4% had a score less than 2. Conversely, 81.2% were with J-CAPRA score ≤7 and 18.8% had a J-CAPRA score ≥8 in the J-CaP cohort.
database (Figure 1). Risk stratification of J-CAPRA scores showed that 36/97 (37.1%) Stage I and II cases were low-risk disease (J-CAPRA score 0–2), while 87/101 (86.1%) Stage III cases were classified as intermediate-risk (J-CAPRA score 3–7) (Table 2A). Interestingly, only 43.5% (233/536) of Stage IV cases were of high risk (J-CAPRA score ≥8) (Table 2A). Comparing to the J-CAP 2001–2003 database, a relatively higher proportion (57%; 2604/4572) of Stage IV cases were stratified into the high-risk group (J-CAPRA score ≥8) (Table 2B). Majority (82.8%; 5427/6552) of the Stage I and II cases were classified as low risk, while 75.4% (2071/2754) Stage III cases were intermediate-risk disease in the J-CaP 2001–2003 database.

Further analysis revealed that 208/299 (69.6%) of stage IV metastatic (M1) cases without T and/or N staging were grouped into intermediate-risk (J-CAPRA score 3–7) category in the M-CaP database. Of note, these M1 patients did not undergo T and/or N staging as it did not affect treatment decisions directly. All M1 patients in the J-CaP 2001–2003 database had complete clinical T, N, and M staging. To address this, we first omitted clinical T and N stage variables, and calculated the score on a 0–8 rather than 0–12 scale in the modified J-CAPRA scoring system for M1 patients. The J-CAPRA score threshold was further adjusted for intermediate (modified J-CAPRA score 3–5) and high-risk (modified J-CAPRA score ≥6) groups for M1 patients, based on biopsy Gleason score, PSA, and clinical M stage alone. Using the J-CaP 2001–2003 database, we performed sensitivity analyses to measure whether there were discrepancies of original J-CAPRA intermediate (3–7) and high-risk (≥8) groups against modified J-CAPRA intermediate (3–5) and high-risk (≥6) groups in the overall survival, prostate cancer-specific survival, and progression-free survival. We demonstrated that there were no significant difference in the median overall survival and prostate cancer-specific survival of M1 patients between original J-CAPRA intermediate and high-risk group and modified J-CAPRA intermediate and high-risk group (p > 0.05, Log-rank test) (Figure 2A,B). A comparable median progression-free survival (PFS) of M1 patients was observed in both intermediate (original J-CAPRA score 3–7; PFS 63.1 months, 95% CI 58.5–67.7 versus modified J-CAPRA 3–5; PFS 61.8 months, 95% CI 56.7–66.9) and high-risk (original J-CAPRA score ≥8; PFS 37.6 months, 95% CI 35.7–39.5 versus modified J-CAPRA score ≥6; PFS 39.6 months, 95% CI 37.7–41.5) stratified by original and modified J-CAPRA scoring systems (p > 0.05, Log-rank test) (Figure 2C). The agreement of J-CAPRA score and modified J-CAPRA score among M1 patients was further supported by Cohen’s coefficient of 0.65. Using the modified J-CAPRA scores, 227/299 (75.9%) of Stage IV cases in the M-CaP database were re-grouped from intermediate-risk category into high-risk category (Table 3); of which, 87.7% (199/227) were M1 patients.

### 4 DISCUSSION

Findings from this study demonstrated that prostate cancer patients receiving ADT in Malaysia tend to be younger with higher risk and more advanced tumors than those treated in Japan. In the present study, we report for the first time the generalizability of the J-CAPRA scoring system in a multi-ethnic Asian cohort. We made two modifications to the J-CAPRA scoring system in assessing the risk of metastatic prostate cancer patients undergoing ADT by omitting clinical T and N staging.
Risk-stratification for prostate cancer is pivotal to guide appropriate treatment decision making at diagnosis and subsequent decision points. Using the Cancer of the Prostate Strategic Urologic Research Endeavor (CaPSURE) database in United States,11 the Cancer of the Prostate Risk Assessment (CAPRA) scoring system was established to incorporate simplicity and clinical applicability in the nomogram performance for assessing risk of radical prostatectomy patients.12 Multiple independent studies have been conducted for validating the CAPRA score.13

With high burden of advanced disease, risk stratification of the M-CaP population meets unusual challenges. Standard risk stratification schemata apply for localised disease only.14,15 The J-CAPRA score estimates risk of progression-free survival and prostate cancer-specific mortality for locally advanced and metastatic prostate cancer patients undergoing ADT. Validation of J-CAPRA score has been performed in numerous academic institutions worldwide.13 Based on the J-CAPRA score risk stratification, it also showed that men on ADT in J-CaP had a lower prostate cancer-specific mortality rate compared to those in the USA, although Japanese had higher disease risk profiles.16

The modified J-CAPRA scoring system is a good and simple model requiring PSA level, biopsy Gleason score, and clinical M staging alone, which are easily assessable from a daily consultation setting, for metastatic prostate cancer patients undergoing ADT. Based on the validation analysis in a large J-CaP cohort (>10,000 subjects), it appears to perform well in predicting disease progression in patients of different risk groups and achieve substantial Cohen’s coefficient agreement of 0.65 with J-CAPRA scoring system. A relatively much lower M1 case (15.2%) in the J-CaP database were re-grouped from J-CAPRA intermediate-risk group to modified J-CAPRA high-risk group compared to M1 cases (88%) in the M-CaP database as all J-CAPRA-scored M1 cases of J-CaP database had complete TNM staging. This finding also suggests the comparability of modified J-CAPRA to J-CAPRA scoring system without the presence of T and N staging. Modified J-CAPRA scoring system is applicable for risk stratification of metastatic prostate cancer patients from other low- and middle-income countries with limited healthcare resources. While cost is a major issue for prostate cancer patients from these countries, complete TNM staging for M1 cases is unlikely to be performed in these countries since both T and N staging do not affect treatment decisions directly in M1 cases. High-risk prostate cancer patients receiving primary ADT may develop rapid disease progression and benefit from early enrolment into clinical trials of novel androgen-receptor targeted drugs, chemotherapy, radiopharmaceutical agent, immunotherapy or combination therapies.

There are limitations to this analysis. First, we were unable to address the 5-years median overall survival, prostate cancer-specific survival, and progression-free survival of M-CaP database as its median follow-up time is currently less than 5 years; therefore, a comparison of outcomes between the J-CaP and M-CaP cohorts could not be established at this time. Second, it remains unknown whether these findings can be extrapolated to other populations. A multi-institutional analysis involving other developing nations is currently underway. Third, there could be a random error in the reporting of disease progression in the J-CaP study by clinicians owing to various definitions of progression.8 Fourth, addition of CHAARTED17–19 and LATITUDE20 criteria such as visceral metastasis and number of bone lesions may increase the accuracy of modified J-CAPRA scoring system in classifying the risk of prostate cancer. Nevertheless, our intention was to establish a risk assessment tool that is simple and quick to be utilised in a routine clinic consultation setting particularly in low- and middle-income countries with limited healthcare resources.

In summary, the J-CAPRA scoring system is developed to aid clinicians and patients in predicting ADT treatment outcomes of prostate cancer. We have modified and validated the J-CAPRA scoring system in order to stratify the risk of M1 patients without T and/or N staging. It is essential to confirm and validate these findings in other developing nations as healthcare cost remains a key element in deciding the management of prostate cancer.

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CONFLICT OF INTEREST
All authors declare no conflict of interest.

DATA AVAILABILITY STATEMENT
The data used to support the findings of this study are available from the corresponding author upon request.

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