Prostate-specific antigen 10–20 ng/mL: A predictor of degree of upgrading to ≥8 among patients with biopsy Gleason score 6

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Purpose: This study aimed to identify the predictors of upgrading and degree of upgrading among patients who have initial Gleason score (GS) 6 treated with robot-assisted radical prostatectomy (RARP).

Materials and Methods: A retrospective review of the data of 359 men with an initial biopsy GS 6, localized prostate cancer who underwent RARP between July 2005 to June 2010 was performed. They were grouped into group 1 (nonupgrade) and group 2 (upgraded) based on their prostatectomy specimen GS. Logistic regression analysis of studied cases identified significant predictors of upgrading and the degree of upgrading after RARP.

Results: The mean age and prostate-specific antigen (PSA) was 63±7.5 years, 8.9±8.77 ng/mL, respectively. Median follow-up was 59 months (interquartile range, 47–70 months). On multivariable analysis, age, PSA, PSA density and ≥2 cores positive were predictors of upgrading with (odds ratio [OR], 1.03; 95% confidence interval [CI], 1.01–1.06; p=0.003; OR, 1.006; 95% CI, 1.01–1.11; p=0.018; OR, 0.65; 95% CI, 0.43–0.98, p=0.04), respectively. On subanalysis, only PSA level of 10–20 ng/mL is associated with upgrading into GS ≥8. They also had lower biochemical recurrence free survival, cancer specific survival, and overall survival (p≤0.001, p=0.003, and p=0.01, respectively).

Conclusions: Gleason score 6 patients with PSA (10–20 ng/mL) have an increased risk of upgrading to pathologic GS (≥8), subsequently poorer oncological outcome thus require a stricter follow-up. These patients should be carefully counseled in making an optimal treatment decision.

Keywords: Prognosis; Prostate-specific antigen; Prostatic neoplasms; Radical prostatectomy

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INTRODUCTION

Due to protracted clinical course of prostate cancer (PCa), clinical guidelines recommend treatment in men with life expectancy of more than 10 years. [1] Appropriate treatment of PCAs is largely dependent on the preoperative risk of disease progression among patients. Based on patient's risk stratification, patients may be offered with
different treatment modalities. All of these factors should be considered in offering any intervention among patients with localized disease. Nowadays, due to the upstream availability of minimally invasive options low risk PCa are likewise being offered with minimally invasive approach with the aid of robotic system.

In counseling, physicians rely mostly on biopsy Gleason score (GS) of their patients. Discordance between prostate biopsy and prostatectomy GS is not uncommon. Albertsen et al. [2] showed that men 55 to 74 years old with biopsy GS 6 have 18% to 30% risk of dying from PCa in the next 15 years of their lives. It is estimated in previous literatures that 50% of needle biopsy score will be upgraded on final pathology specimen [3]. In our previous study, our results showed that patients treated with radical prostatectomy for clinically low risk disease has 129% to 36.4% chance of upgrading [4]. This inaccuracy has serious impact on pretreatment decision-making, predominantly in patients who choose active surveillance or nonsurgical options. Increase in biopsy GS in men with pathological GS 8 to 10 put them on higher risk of death from PCa [5]. Likewise, upgrade in GS put them in increased chances of biochemical recurrence (BCR) [6,7]. Several articles involving patients treated with open radical prostatectomy have discussed the predictors of GS upgrading [6,8-10]. Nevertheless, no one among them reported the predictor of the degree of upgrading into higher pathologic GS (≥8). To the best of our knowledge this is the first study to report on the predictors of upgrading and the degree of upgrade exclusively on GS 6 patients treated with RARP.

In this study, we report predictors of GS upgrading and degree of upgrading among patients who have an initial GS 6 biopsy treated with RARP.

**MATERIALS AND METHODS**

We retrospectively reviewed records from our prospectively maintained Institutional Review Board-database (approval number: 2014-009-001) of patients who underwent RARP for PCa between July 2005 to June 2010. Patients with clinically localized PCa based on magnetic resonance imaging, digital rectal examination, are included in the study. Twelve-core transrectal ultrasound guided needle biopsy was done on all patients. After excluding patients with nonorgan confined PCa, Prostate-specific antigen (PSA)>20 ng/mL and patients with missing information needed to classify disease risk, 359 men with an initial biopsy GS of 6 were included in the study.

RARP was performed via a transperitoneal approach as described in our previous study [11], using the daVinci Surgical System (Intuitive Surgical, Sunnyvale, CA, USA) with the pelvic lymph node dissection being performed when predicted probability of lymph node metastasis based on clinical information is more than 3% [12].

Patients’ clinical and pathological variables: age, body mass index (BMI), prostate volume, PSA, number of core involvement, biopsy and prostatectomy GS were obtained. A dedicated pathologist evaluated the prostatectomy specimens with particular attention to pathologic stage, GS, surgical margins, perineural invasion and angiolymphatic invasion. GS upgrading is considered if initial biopsy GS≤6 increased on prostatectomy specimen to ≥GS 7 [13]. Patients were grouped into group 1 (nonupgrade GS) and group 2 (upgraded GS) and comparative analysis was performed. The primary aim was to evaluate the predictors of GS upgrading and its degree, meanwhile the secondary outcome was to evaluate the oncological outcomes of upgraded GS degrees 7 (3+4, 4+3), and ≥8.

Follow-up with physical examination and PSA quarterly for the first year, semiannually for the second year, and annually thereafter was done. BCR was defined as postoperative PSA≥02 ng/mL taken twice at least 6 weeks apart [14].

Statistical analysis was used to analyze demographic and pathological characteristics. Counts of frequencies were expressed as percentages, and continuous data were presented as mean±standard deviation (SD) or median and interquartile range (skewed data). Student t-test was used for continuous variables while chi-square test was used for categorical variables. The probability of BCR, Cancer specific survival (CSS) and overall survival (OS) among upgraded patients were calculated by the Kaplan-Meier analysis. The univariable and multivariable logistic regression models were used to estimate the impact of clinical features on upgrading and degree of upgrading. The IBM SPSS Statistics ver. 22.0 (IBM Co., Armonk, NY, USA) was used for all statistical analyses. A p-value of <0.05 was considered significant, and all p-values were 2-sided.

**RESULTS**

The mean age of the patient was 63 years old (SD, ±7.5). Mean BMI, prostate volume, and PSA were 24.2±2.6 kg/m² (interquartile range [IQR], 15.9–30.8 kg/m²), 39.2±20.9 (mL) (IQR, 10.5–164 mL), and 6.8 ng/mL (IQR, 5–10 ng/mL), respectively. Our median follow-up was 59 months (IQR, 47–70 months).

Table 1 summarized the comparative preoperative characteristics. BMI were almost the same for both groups.
Patients in the upgraded group were older (p=0.007) and had higher PSA and smaller prostate in volume; 7.7 ng/mL (IQR, 5.2–12.4 ng/mL) and 36.4±16.4 mL, respectively. The mean age of patients in the group 2 was 62.2±7.6 years while it was 64.3±7.3 years for group 1 (p=0.007). Likewise, PSA density is higher in group 2 compared to group 1 with 0.3±0.2 and 0.2±0.1, respectively. Among group 2 patients, 35.9% (52 of 145) had a PSA of 10–20 compared to group 1 with 17.3%. Moreover, 77.2% (112 of 145) of group 2 patients had ≥2 core involvements. More than 25% positive core involvement was seen in 43.4% of patients who upgraded compared to 26.2% who did not upgrade (p=0.007). Group 2 patients had higher pathologic T stage with 44.1% having >T3 (p≤0.001). Positive surgical margin (PSM) was higher in upgraded group with 44.1% having >T3 (p≤0.001). Significantly, perineural invasion was higher in upgraded group 51.7% versus 19.2% among the nonupgraded group (p≤0.001). Ten percent of group 2 patients showed angiolymphatic invasion compared to 2.3% of group 1 (p=0.002).

Table 2 summarized the comparison among upgraded patients. Patients who upgraded to Gleason score ≥8 had higher preoperative PSA (p=0.000). The results of univariable and multivariable logistic regression analyses were summarized in Table 3. Statistically significant preoperative variables were included in the regression model. Age, PSA, PSA density, clinical T stage, percentage of cores and
number of positive had significant result hence further analyzed on regression analysis. Moreover, on multivariable analysis age, PSA, PSA density and >2 cores positive were predictors of upgrading with (odds ratio [OR], 1.03; 95% confidence interval [CI], 1.01–1.06; p=0.005; OR, 1.006; 95% CI, 1.01–1.11; p=0.018; OR, 0.65; 95% CI, 0.43–0.98, p=0.04), respectively were strongest predictors of GS upgrade among clinicopathological variables.

On subanalysis of the predictors of degree of upgrading on Table 4, we found out that only PSA level of 10–20 ng/mL have increased likelihood of upgrading from GS 6 into pathologic GS≥8. Likewise, we report the predictors of upgrading like age, PSA, PSA density, and number of cores among patients who have solely GS 6 on needle biopsy. Knowledge on the impact of GS upgrading is of great importance especially at the point of shared decision-making between the physician and the patient prior to any planned intervention. In a study done by Carlsson et al. [15], they concluded that having surgery as primary treatment for low risk PCa might result in favorable oncologic outcome but may also jeopardize men's

**Table 2. Comparison of upgraded Gleason score 6 patients**

| Variable                        | 3+4 (n=87) | 4+3 (n=37) | ≥4+4 (n=18) | p-value |
|---------------------------------|------------|------------|------------|---------|
| Age (y)                         | 63±7       | 65±7       | 65±9       | 0.26*   |
| Body mass index (kg/m²)         | 23.8±2.4   | 24.6±2.1   | 24.6±3.0   | 0.16*   |
| PSA (ng/mL)                     | 8.0±3.9    | 8.3±3.8    | 13.1±4.8   | 0.000*  |
| Prostate volume (mL)            | 36.0±17.2  | 36.3±15.0  | 36.0±16.0  | 0.19*   |
| PSA density                     | 0.25±0.13  | 0.25±0.18  | 0.23±0.15  | 0.72*   |
| Core number                     |            |            |            | 0.74*   |
| 1                               | 16 (19.8)  | 9 (25.7)   | 4 (23.5)   |         |
| ≥2                              | 65 (80.2)  | 26 (74.3)  | 14 (76.5)  |         |
| Percent positive core>25%       |            |            |            | 0.58*   |
| Yes                             | 35 (43.2)  | 13 (37.1)  | 9 (52.9)   |         |
| No                              | 46 (56.8)  | 22 (62.9)  | 8 (47.1)   |         |

Values are presented as mean±standard deviation or number (%).

PSA, prostate-specific antigen.

*Based on Student t-test. **Based on chi-square test.

**Table 3. Logistic regression analysis of predictive factors associated with overall upgrading**

| Variable                        | Univariable | Multivariable |
|---------------------------------|-------------|---------------|
|                                | OR         | 95% CI | p-value | OR         | 95% CI | p-value |
| Age                             | 1.03       | 1.01–1.06 | 0.005   | 1.03       | 1.01–1.06 | 0.003   |
| PSA                             | 1.09       | 1.05–1.13 | 0.000   | 1.006      | 1.01–1.11 | 0.018   |
| PSA density                     | 5.20       | 2.80–9.60 | 0.000   | 2.70       | 1.15–6.35 | 0.022   |
| Prostate volume                 | 0.99       | 0.98–1.00 | 0.212   | -          | -        | -       |
| Core number                     | -          | 0.020   | 0.04    | -          | -        | -       |
| 1                               | Reference  |         |         | Reference  | -        |         |
| ≥2                              | 0.62       | 0.41–0.92 | 0.017   | 0.65       | 0.43–0.98 | 0.51    |
| Positive cores>25%              |            |         |         |            |         |         |
| No                              | Reference  |         |         | Reference  | -        |         |
| Yes                             | 1.01       | 1.00–1.01 | 1.14    | 0.76–1.71  | 0.30    |
| Clinical T stage                |            |         |         |            |         |         |
| T1–T2a                          | Reference  | -       | 0.006   | Reference  | -        | 0.25    |
| T2b–T2c                         | 1.00       | 0.62–1.62 | 0.984   | 0.76       | 0.48–1.21 | 0.25    |
| >T3                             | 1.92       | 1.27–2.91 | 0.002   | 0.63       | 0.34–1.14 | 0.13    |

OR, odds ratio; CI, confidence interval; PSA, prostate-specific antigen.

**DISCUSSION**

In this study, we have shown that patients preoperative PSA level of 10–20 ng/mL have increased likelihood of upgrading from GS 6 into pathologic GS≥8. Likewise, we report the predictors of upgrading like age, PSA, PSA density, and number of cores among patients who have solely GS 6 on needle biopsy. Knowledge on the impact of GS upgrading is of great importance especially at the point of shared decision-making between the physician and the patient prior to any planned intervention. In a study done by Carlsson et al. [15], they concluded that having surgery as primary treatment for low risk PCa might result in favorable oncologic outcome but may also jeopardize men’s
potency and continent rate postoperatively. This scenario may have an impact on patients’ overall quality of life and great importance is often placed during counseling. At this point, preoperative predictors of upgrading may help urologist in proper counseling of their patients.

Previous reports have discussed that PSA level is a significant predictor of upgrading [16, 17]. At some point the level of PSA to which upgrading is most likely to occur have been discussed by Colleselli et al. [18]. They reported that patients with PSA range of 2.0–3.9 and 4.0–10.0 ng/mL had a 32.6% and 44% risk of upgrading, respectively. Among the previously mentioned studies, no one has investigated on the predictor of the degree of upgrading among their cohorts. To our knowledge, our study is the first to discuss the predictor of the degree of upgrading on exclusively GS 6 patients on needle biopsy.

Meanwhile, several studies have reported different

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**Table 4. Univariable analysis of the predictor of degree of upgrading to Gleason score≥8**

| Variable                      | Univariable analysis | 95% CI          | p-value |
|-------------------------------|----------------------|-----------------|---------|
| Age                           | 1.01                 | 0.95–1.08       | 0.59    |
| Body mass index               | 1.09                 | 0.90–1.30       | 0.34    |
| PSA                           | 0.000                |                 |         |
| <10                           | Reference            |                 |         |
| 10–20                         | 6.65                 | 2.36–18.75      |         |
| PSA density                   | 0.48                 | 0.02–11.75      | 0.65    |
| Prostate volume               | 1.00                 | 0.97–1.03       | 0.80    |
| Core number                   | 0.77                 |                 |         |
| 1                             | Reference            |                 |         |
| ≥2                            | 1.17                 | 0.38–3.65       | 0.31    |
| Positive cores>25%            | No                   | Reference       |         |
|                               | Yes                  | 0.61            | 0.23–1.58 |

OR, odds ratio; CI, confidence interval; PSA, prostate-specific antigen.

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Fig. 1. Upgraded patients’ biochemical recurrence (BCR)-free survival rate. GS, Gleason score.

Fig. 2. Upgraded patients’ cancer specific survival rate. GS, Gleason score.

Fig. 3. Upgraded patients’ overall survival rate. GS, Gleason score.
Thus, even on patients with GS 6 on biopsy risk of upgrading is increased whenever multiple cores are present. In the current study, we found out that patients with more than or equal to 2 cores involved had higher chances of upgrading. This might be explained by the fact that presence of multiple cores increases the likelihood of lymph node invasion and less organ-confined disease (≥pT3). [19,20] Thus, even on patients with GS 6 on biopsy risk of upgrading is increased whenever multiple cores are present.

In a cohort of 2,771 men, the group of Serkin et al. [21] reported that patients with higher PSA density were likely to upgrade. And this could also happen even if they are eligible for active surveillance [22]. In support to their findings, PSA density was also a significant predictor of overall upgrading among our patients. However, when PSA density was entered on the model for predicting degree of upgrading it has lost its significance. This might be due to the small number of patients who upgraded to ≥8. Meanwhile, prostate volume has long been a point of debate; several reports have stated that prostate volume is a significant predictor of upgrading among patients who underwent radical prostatectomy [10,23,24]. There seems to be an inverse relationship between prostate size and chances of upgrading which was also evident in our current study.

On the other hand, patients’ age showed to have an impact on upgrading in our cohort. This may be explained by the fact that as man ages PSA level tends to increase stepwise. This was also evident in the cohort of 25,858 patients reported by Caster et al. [25], that older age was associated with GS upgrading and pathologically advanced disease.

Difference between biopsy and prostatectomy specimen GS has a significant impact on BCR. Corcoran et al. [6] reported that change in GS on prostatectomy was a predictor of BCR. Moreover, Boorjian et al. [5] also found out that upgrading in GS was associated with BCR and progression to systemic disease, though their study included all GS. Likewise, our findings suggest that BCR was notably higher among patients who upgraded into higher GS in support of the aforementioned studies. Badani et al. [26] reported in their series of 2,706 patients a BCR of 7.3% however its median follow-up was only 22 months. On the other hand, Menon et al. [7], did a study with a median follow-up of 602 months however, their study included all GS. In our current study, we reported the oncologic outcome of our cohort at a median follow-up of 59 months for specifically GS 6 patients alone after RARP. This reflects a more realistic oncologic behaviour of patients who were initially GS 6 on biopsy but subsequently changed on pathologic specimen.

Upgraded group of patients have had significant differences in PSM, extraprostatic extension, seminal vesicle invasion, and lymphovascular invasion [27]. Accordingly, the rate of extraprostatic extension, and PSM were noted to be higher among patients who have GS upgrading during radical prostatectomy [9]. In concordance, our results also showed that upgraded group had higher PSM, perineural invasion and angiolymphatic invasion compared to nonupgrade group depicting probably a more aggressive disease.

There are several limitations in our study. First of all, this study was analyzed in a retrospective manner. Small number of patients which represents a higher degree of upgrading from GS 6 is one of its main limitation. Comparative analysis of patients with originally GS ≥8 from biopsy and those who upgraded from GS 6 to ≥8 is likewise recommended. A longer follow-up maybe needed to portray satisfactory oncologic impact of upgrading to clinical course and disease progression of patients with initially GS 6 treated with radical prostatectomy. Despite these limitations, our study showed that patients with higher preoperative PSA level has increased risk of upgrading to higher degree of GS which should be discussed thoroughly before offering any treatment options to patients.

**CONCLUSIONS**

Patients with GS 6 on biopsy with PSA level of 10–20 ng/mL carries an increased risk of upgrading into higher pathologic GS (≥8) with poorer oncologic outcome. Presence of multiple cores, high PSA density increases the likelihood of upgrading. We recommend strict counseling especially in choosing treatment options between surgical and nonsurgical for patients with GS 6 on biopsy whenever these parameters are present.

**CONFLICTS OF INTEREST**

The authors have nothing to disclose.

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