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Pathological findings and PSA outcomes after radical prostatectomy in men eligible for active surveillance: does the risk of misclassification vary according to the different biopsy criteria?

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ABSTRACT

Purpose:
To evaluate and compare the pathological findings and the PSA outcomes after radical prostatectomy (RP) in men eligible for AS, according to 3 different biopsy inclusion criteria.

Materials and Methods:
The study population included 177 men eligible for AS who fulfilled clinico-biological criteria (Gleason score ≤6, PSA<10, clinical stage T1c) and biopsy criteria as follows: (1) <3 positive cores and <3 mm of total tumour length; (2) and/or <3 positive cores with a cancer involvement <50% in any core; (3) and/or <33% of positive cores. PSA density cut-offs of 15 and 20 ng/ml/gr were also studied among these groups. Pathological findings on RP specimens and biochemical recurrence-free survival (RFS) were studied. Median follow-up was 34 months.

Results:
Cancers were graded Gleason 7 on RP specimens from 48.3% to 55.4% of cases. The rates of extracapsular extension (ECE) and vesicle seminal invasion (SVI) ranged from 11.2% to 17.5%, and from 1.1% to 1.8%, respectively, regardless of PSA density. The use of PSA density as AS criterion decreased for AS by 1.4-fold and by 2.3-fold the number of men eligible according to the cut-off used. The risk of unfavourable disease (defined as pT3-4 stage and/or a Gleason score≥8) remained between 15% to 19.2% when a PSA density cut-off of 15 ng/ml/gr was used. The risk of overall unfavourable disease was significantly higher in men with a cancer involvement ≥3 mm in initial biopsy compared with men who fulfilled these most stringent biopsy criteria (27.3%, vs 13.5%, respectively; p=0.023). The 3-year biochemical RFS was 91.5% and was not affected by the 3 different AS definitions.

Conclusions:
Even with the use of a 21-core biopsy protocol, the rate of unfavourable disease on RP specimens remains still elevated in men eligible for AS. Patients must be informed of this risk of misclassification which is about 20% in men who fulfil the less stringent biopsy criteria.
INTRODUCTION

The prostate cancer (PCa) detection rate has strongly increased since the PSA screening and the use of extended prostate biopsy protocol. Active surveillance entails a strategy by which selected men are managed expectantly with the intention to apply potentially curative treatment in case of progression signs [4]. Cancers that are amenable to AS usually are identified on favourable preoperative parameters and the risk is estimated by integrating Gleason score, pretreatment PSA, clinical stage, prostate volume and the extent of biopsy involvement with tumour [6-10]. Published AS series use different criteria largely based on centre experiences and preferences with no hard data. The most common clinical data used to define AS criteria are a Gleason score ≤6, PSA≤10 ng/ml and a clinical stage T1c disease. The PSA density is noted as inclusion criteria in some studies with different reported cut-offs for AS inclusion [6,13-14, Carter]. Other characteristics to consider include pathological biopsy parameters with a wide variation concerning the AS inclusion criteria. A cancer involvement of <33% of biopsy cores was retained in the UCSF cohort [16]. Other series included cancers involving <3 cores only [14] and with an extent of cancer in any core <50% [13,17]. Recently, a prospective trial of AS (SURACaP trial) has been opened in France including patients with a tumour length <3 mm in <3 cores [18].

Studies comparing entry criteria for AS protocols, especially in terms of biopsy parameters, are needed to clarify the best candidates for active surveillance. Recent studies emphasized the risk of under-diagnosis, adverse pathological findings and thus, missing window of curability if AS is preferred [Suardi, Conti]. Based on these limitations and the wide variation of existing pathological criteria for AS inclusion, we decided to compare pathological findings and PSA outcome after radical prostatectomy (RP) in men eligible for AS, according to 3 different biopsy parameters reported in the literature. PSA density cut-offs which have been published in recent AS studies were also tested among each group in analyses.
MATERIALS AND METHODS

Between January 2001 and July 2008, we identified patients who were diagnosed with PCa in a set of 21 biopsies and who have undergone a RP for clinically localized and low-risk prostate cancer according to preoperative data. Inclusion criteria for low-risk cancers were PSA level ≤10ng/ml, a clinical stage T1c disease, a Gleason score ≤6 and a life-expectancy >10 years. All patients underwent clinical evaluations, including digital rectal examinations, serum PSA, and transrectal ultrasound. All the patients had undergone a 21-core biopsy protocol as previously described for abnormal digital rectal findings or elevated PSA [19]. All the biopsies and radical prostatectomies were performed in our department and specimens were evaluated by senior uropathologists. Tumour volume was not measured routinely. Data from clinical evaluation, biopsy and RP specimens, and follow-up were recorded in a prospective database. PSA recurrence was defined as PSA >0.2 ng/ml after RP.

Of all the patients in the database, we identified 177 patients who met the main favourable criteria (PSA ≤10ng/ml, T1c disease, Gleason score ≤6, life-expectancy >10 years) and who were diagnosed to have a cancer involvement of <33% of biopsy cores (group 3). Of these 177 men, 112 met the following criteria: <3 positive cores with a cancer involvement <50% in any core (group 2). Of these 112 men, 89 met the following criteria: <3 positive cores with total tumour length <3 mm (group 1).

We studied the pathological findings on RP specimens, such as Gleason score, extracapsular extension (ECE), seminal vesicle invasion (SVI), positive surgical margins, and the PSA outcomes in the 3 groups during the follow-up. Statistical analyses were conducted between the 3 groups regardless of PSA density, and then, with the use of 2 PSA density cut-offs: 15 and 20 ng/ml/gr. Secondly, we compared the results between men of group 1 (cancer involvement <3 mm) and men who did not fulfil this most stringent biopsy criteria (not group 1). The qualitative data were tested using a chi-square test or Fisher’s exact test as appropriate and the quantitative date were tested using Student’s t-test. The Mann-Whitney’s test was used in case of no normal distribution. Biochemical recurrence-free survival was established using the Kaplan-Meier method. Curves were tested by log-rank test. The limit of statistical significance was defined as p<0.05. The SPSS 13.0 (Chicago, Illinois) software was used for analysis.
RESULTS

Of the 468 patients with PCa in a 21-core biopsy scheme and who have undergone a RP, 177, 112 and 89 men fulfilled preoperatively the AS criteria of group 3, 2 and 1, respectively. Table 1 shows the patients’ characteristics. The 3 groups were comparable and there were no significant baseline differences between the 3 groups in terms of PSA level, age, DRE findings, prostate volume and PSA density.

Table 2 summarizes the pathological findings at RP and the rates of unfavourable disease according to each AS criteria. The use of PSA density as AS criterion decreased by 1.4-fold and by 2.3-fold the number of men eligible for AS, if a cut-off of 20 and 15 ng/ml/gr was used, respectively. A majority of Gleason score 6 was observed in group 1 (51.7%), whereas a majority of Gleason score 7 or more was reported in groups 2 (53.6%) and 3 (55.4%). The rates of unfavourable pathological findings increased in groups 2 and 3 compared with the rates in group 1. Primary Gleason score 4 was reported in 12.4% of men of group 3 (versus 8.9% in group 1). Extracapsular extension was noted in 17.5% of RP specimens in group 3, versus 11.2% in group 1. The rate of positive surgical margins reached 20.9% in group 3 compared with 18.0% in group 1. Globally, when unfavourable was defined as ECE, SVI and/or Gleason score 8 or more, the overall rate was 20.3% in group 3 and 13.5% in group 1. T3 or T4 cancers were reported in 12.4% of cases in group 1 and 19.2% of cases in group 3. No differences reached significance when comparing these rates between groups.

Table 3 lists pathological features when PSA density was also used as AS criterion (cut-offs of 20 and 15 ng/ml/gr). Risks of unfavourable disease were then slightly lower but differences did not reach significance.

Table 4 shows the statistical comparisons when analysis compared men who fulfilled the group 1 criteria with the remaining men (not group 1). Significant differences were noted. Men who did not fulfilled the group 1 criteria had statistically more frequently PCa with ECE (p=0.025), pT3-4 stage (p=0.020) and unfavourable disease (p=0.023). The use of a PSA density cut-off of 20 did not modify these statistical differences. Statistical significance was not found when the cut-off was decreased to 15 ng/ml/gr. These men had also more high-grade PCa and positive surgical margins, but difference did not reach significance.

No differences appeared between the groups in terms of seminal vesicle invasion and PSA failure after RP.

The overall 3-year recurrence-free survival rate was 91.5%. All the biochemical recurrences occurred during the 2 first years after RP. Figure 1 shows the biochemical recurrence-free survival curves according to the 3 groups. The survival rates between the men in group 1 (n=89) and the men who did not fulfilled the group 1 criteria (n=88) were not statistically different (log-rank test, p=0.571, see Figure 2). The 3-year recurrence-free survival rate was 92.7% for the men in group 1 and 89.8% for men who did not fulfilled the group 1 criteria.
DISCUSSION

Active surveillance is a treatment option for selected patients with low-risk PCAs. Data demonstrate that the proportion of low-risk men electing surveillance has risen in recent years [20]. Published AS series used different inclusion criteria largely based on centre experiences and preferences with no hard data. A wide variation concerning the biopsy inclusion criteria is also expressed in the literature. Many AS criteria have been based on series of patients diagnosed with much less than 21-core biopsies, sometimes even sextant biopsies only. The number of positive cores, the tumour length (total or at any core) and the percent of cancer involvement at any core are predictive factors of tumour volume in RP specimens or biochemical failure after RP [8,22-23].

The aim of our retrospective study was to compare the rate of misclassification associated with the use of 3 different biopsy criteria [13,16-17]. For each criterion, we also tested the impact of PSA density on the number of men eligible and the risk of unfavourable disease [van den Bergh, Carter]. A non organ-confined disease and/or a high-grade PCa defined a misclassification case. The 3 groups were comparable in terms of biopsy Gleason score (no grade 4 or 5), PSA level, age and DRE findings. Each patient underwent the same 21-core biopsy protocol under local anaesthesia. Thus, the different studied groups were strongly homogeneous.

The overall results of our series indicate that a cancer of low grade and small volume in biopsies is not necessarily indicative of a good pathological assessment on RP specimens. The Gleason score was upgraded from 48.3% to 55.4% according to the different biopsy criteria. The rate of misclassification, ie the presence of either non organ-confined disease or high grade PCAs was relatively high. Unfavourable disease defined as ECE, SVI or primary Gleason 4 was reported 27.7% of cases in group 3. The less stringent the biopsy criteria were, the higher the misclassification rates were. When unfavourable disease was defined as ECE, SVI and/or Gleason score 8 or more, this risk in group 3 was increased by 1.5-fold compared with the risk of group 1. Of the 97 men who fulfilled the group 2 and 3 criteria but not the stringent group 1 criteria (total tumour length >3 mm), about one third had unfavourable disease on RP specimens. This rate was only 13.5% when group 1 criteria were considered. Men who did not fulfil the group 1 criteria had statistically more frequently PCa with ECE, pT3-4 stage and unfavourable disease. However, concerning the biochemical recurrence after RP, the estimate risk of relapse was comparable and not significantly different in the 3 groups (about 5%).

Using the risk-stratification schemes based only on PSA level, DRE findings, biopsy Gleason score and extent of cancer involvement appears insufficient to identify cancers with a low risk of progression. Therefore, other factors, such as PSA density and PSA velocity, would provide additional significance [6,8]. In our cohort, patients who had a high PSA density at diagnosis were more likely to have unfavourable disease on RP specimens. Thus, the difference was slight and the use of PSA density as AS criterion did not strongly modify our results. Only 41% patients in our series had a PSA density lower than 0.15 ng/ml/gr and only 64 men fulfilled at once this measurement and the group 1 criteria. Ideally, the strictest criteria should be used in order to decrease the risk of unfavourable disease. However, very high selective and stringent AS inclusion criteria allows the inclusion of a relatively small proportion of patients from the pool of those with localized PCAs. Avoiding misclassification errors need to reduce severely the number of men eligible for AS. In our series, when the rate of misclassification was reduced from 27.7% to 21.3% (ie, by 1.3-fold), the number of men eligible for AS decreased by 2-fold (from 177 to 89 men).

Rates of misclassification and non-organ confined disease were lower than those reported in recent studies [Conti, Suardi]. These discrepancies can be explained by non comparable cohorts, retrospective analyses or other biases. However, we think that the use of a 21-core biopsy scheme, compared with a 10-12 cores protocol, has a strong impact on this improvement. The counterpart is that the saturation biopsy strategies have been hypothesized to increase the potential risk of overtreating patients whose tumours induce very low risk to life. Nevertheless, the role of prostatic biopsies changed. The actual importance of prostate biopsies has evolved from purely cancer detection to investigating as to how biopsy results can assist clinical management
for patients. The inclusion of patients in AS protocols emphasizes the necessity of perfectly accurate staging strategies.

Biochemical recurrence rates were probably not a valid endpoint to address conclusions in men eligible for AS. The PSA failures can not reflect accurate measurement in men who did not undergo radical treatment. However, PSA failure is established to be associated with an increased risk of progression to metastatic disease and specific death [Freedland]. Few PSA failures occurred in our series and the biochemical recurrence-free survival was 91.5% at 3 years after surgery. None of the patients developed clinical recurrence with symptoms during the follow-up and no specific death appeared. These results may validate the use of AS as treatment option in selected patients who are informed of the need of close surveillance and the risk of deferred treatment. However, the number of events in all 3 groups was small and conclusions on differences in biochemical RFS may therefore be too premature. The lack of significant differences might be due to short follow-up since a large proportion of PSA failures occur beyond 3-year follow-up [Amling].

Our cohort can appear small, compared with the study of Suardi et al. [11]. However, Suardi et al. have considered only PSA, Gleason score and DRE as AS criteria. We added pathological indicators of cancer involvement (percent of cancer in any core, total tumor length) and the impact of PSA density in a very homogeneous and extensively biopsied cohort. One of the other limitations of our study was to consider for the PSA analyses only the men who have undergone RP. This may represent a bias, the target population for AS being larger than our restricted cohort. In this study, we tested current AS criteria which were internationally used or recently published [Carter, 13,16-18]. Our listing of AS criteria was not complete but provided a representative cohort of the men actually included in AS protocols. Recent AS criteria of the French protocol were interesting as it provided the most stringent definition in terms of biopsy parameters [18]. However, only prospective studies comparing biopsy inclusion criteria for AS protocols and comparing AS with immediate radical treatment would be able to clarify the best candidates for active surveillance according outcomes in terms of rising PSA and specific deaths.

Even with the use of a 21-core extended biopsy scheme, a number of PCs will be selected for AS, while these actually constitute more aggressive tumors. However, these results do not contraindicate the inclusions in AS protocol. AS protocols always included a close surveillance scheme, aiming to catch the aggressive tumors as soon as possible during follow-up. Nevertheless, our findings can help urologists to better inform the patients about the risk of misclassification and the need of close surveillance.
CONCLUSIONS

Our study demonstrates the limitations of AS inclusion criteria, even with the use of a 21-core biopsy protocol. The rate of unfavourable disease remains still elevated, even with the use of the most selective criteria. In the group of patients with the most stringent biopsy criteria, one fifth of patients have non organ-confined disease and/or primary Gleason score of grade 4/5. Thus, any AS selective model provides potential limitations with a risk of under-diagnosis, adverse pathological findings and potential hazards of missing window of curability if AS is preferred. These results can help urologists to better inform the men eligible for AS about this risk of misclassification.
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Table 1. Patients characteristics at baseline (n=194)

Table 2. Pathological findings and PSA outcomes after radical prostatectomy according to the 3 different biopsy criteria (groups 1, 2 and 3).

Table 3. Univariate analysis comparing pathological findings and PSA outcomes between group 1 (n=94) and group 3 excluding group 1 (n=94).

Figure 1. Biochemical recurrence-free survivals according the 3 different biopsy criteria (groups 1, 2 and 3).