Reduction of MRI-targeted biopsies in men with low-risk prostate cancer on active surveillance by stratifying to PI-RADS and PSA-density, with different thresholds for significant disease

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Background: The fear of undergrading prostate cancer (PCa) in men on active surveillance (AS) have led to strict criteria for monitoring, which have resulted in good long-term cancer-specific survival, proving the safety of this approach. Reducing undergrading, MRI-targeted biopsies are increasingly used in men with low-risk disease despite their undefined role yet. The objective of this study is to investigate the rate of upgrading using MRI-targeted biopsies in men with low-risk disease on AS, stratified on the basis of PI-RADS and PSA-density, with the aim to reduce potential unnecessary repeat biopsy procedures.

Methods: A total of 331 men were prospectively enrolled following the MRI-PRIAS protocol. MR imaging was according to Prostate Imaging Reporting and Data System (PI-RADSv2) guidelines. Suspicious MRI lesions (PI-RADS 3–5) were additionally targeted by MRI-TRUS fusion biopsies. Outcome measure was upgrading to Gleason score (GS) ≥3+4 with MRI-targeted biopsies, stratified for PI-RADS and PSA-density.

Results: In total, 25% (82/331) of men on AS showed upgrading from GS 3+3. Only 3% (11/331) was upgraded to GS ≥8. In 60% (198/331) a suspicious MRI lesion was identified, but in only 41% (82/198) of men upgrading was confirmed. PI-RADS 3, 4 and 5 categorized index lesions, showed upgrading in 30%, 34% and 66% of men, respectively. Stratification to PI-RADS 4–5, instead of PI-RADS 3–5, would have missed a small number of high volume Gleason 4 PCa in PI-RADS 3 category. However, further stratification into PI-RADS 3 lesions and PSA-density <0.15 ng/mL² could result in a safe targeted biopsy reduction of 36% in this category, without missing any upgrades.

Conclusions: Stratification with the combination of PI-RADS and PSA-density may reduce unnecessary additional MRI biopsy testing. Overall, the high rate of detected upgrading in men on AS may result in an unintended tightening of continuing in AS. Since patients, included under current AS criteria showed extremely favorable outcome, there might be no need to further restrict continuing on AS with MRI and targeted biopsies.

Keywords: Prostate cancer (PCa); biopsy; magnetic resonance imaging (MRI); MRI-guided targeted biopsy; PSA density; PI-RADS; risk stratification

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Introduction

In the Western world, about one half of all patients diagnosed with prostate cancer (PCa) has low-risk disease (1). In low-risk disease, active treatment hardly yields survival benefit (2). Therefore, active surveillance (AS) is the recommended option for the initial management of low-risk disease (3,4).

Monitoring in AS is based on repeated PSA measurements, clinical T-staging based on digital rectal exams, and repeated random systematic transrectal ultrasound (TRUS)-guided biopsies (3). Repeated biopsies are cumbersome for patients and have severe complication risks. Moreover, sampling errors lead to underestimation of the Gleason grading (5).

The fear of undergrading PCa in men on AS has however led to strict criteria for monitoring, which has resulted in good long-term cancer-specific survival, proving the safety of this approach (6). To reduce the fear of undergrading, MRI and MRI-targeted biopsies are increasingly used in the management of patients with clinically low-risk PCa, despite their role has not yet been established definitively (7). The use of MR imaging in AS has improved the inclusion of true low-grade PCa; targeted biopsies of suspected lesions on MRI may result in excluding those men found with intermediate/high-grade PCa (8).

However, this additional testing by MRI and MRI targeted biopsy may result in an unintended reclassification to perceived higher risk. This is termed as ‘risk inflation’; a cancer that is stable may be more accurately sampled at MRI-targeted biopsy and found to include higher risk features than when it was sampled in a routine systematic manner (9). This lesion targeting results in an increase in risk attribution, if traditional criteria [i.e., Gleason score (GS), cancer core length and the proportion of positive cores on routine sampling] are still applied. It would therefore be wrong to falsely encourage men to cease AS because of an apparent increase in risk (reclassification) rather than a true change in their cancer.

Appropriate risk thresholds are not yet understood when MRI and MRI-targeted biopsies are used. In this study we investigate the upgrading with MRI and MRI-targeted biopsies in men with low-risk disease on AS, and examine the potential reduction of targeted biopsies by stratifying to PI-RADS and PSA-density. We explore the possible risk inflation by MRI and targeted biopsies, and look into the potential extension of GS thresholds for continuing in AS when using MRI in strict monitoring.

Methods

The study was HIPAA compliant and was approved by the institutional ethical review board (NL45884.078.13/A301321). Written informed consent with guarantee of confidentiality was obtained from the participants. Men with low-risk PCa are prospectively enrolled in our in-house database as part of our AS protocol. From November 2013 until October 2017 a total of 347 consecutive men on AS for low-grade (GS 3+3) PCa detected by TRUS guided biopsy received a first multi-parametric MRI and targeted biopsies of visible suspicious MRI lesions at our tertiary referral center.

A total of 331 men were included in the current study. Men were excluded (16/347), as they did not undergo additional targeted biopsy, despite of having a positive MRI (n=8 PI-RADS 3 lesions, n=8 PI-RADS 4 lesions). Results of part of this prospective cohort have been previously published (10).

In 50% (166/331) men were participants of the PRIAS study (www.prias-project.org), an international web-based AS study with strict criteria for inclusion at diagnosis (GS 3+3, T-stage ≤ cT2, PSA ≤10 ng/mL, ≤2 positive cores, PSA density <0.2) and follow-up (11). Within the MRI-PRIAS side study protocol an MRI and targeted biopsies (if indicated) are performed at baseline (3 months after diagnosis) and during every repeat standard TRUS-guided biopsies, scheduled at 1, 4, 7 and 10 years after diagnosis. Inclusion in the MRI-PRIAS side study protocol is also possible after ≥1 repeat TRUS-guided biopsies. The only reclassification criterion in the MRI-PRIAS side study is the presence of high-grade PCa (GS ≥3+4) at MRI-targeted biopsy. A head-to-head comparison of MRI-targeted with standard TRUS-guided biopsies was only available in repeat biopsies, and was not further investigated in this study.

The remaining 165/331 (50%) men in the present study had low-grade PCa based on standard TRUS-guided biopsy findings, but were followed-up outside of the PRIAS protocol as they did not meet the strict PRIAS inclusion criteria or were referred from a center not participating in PRIAS. All men were included in our prospective, institutional review board approved database, which is HIPPA compliant.

Multi-parametric MRI

The institutional MRI protocol included T2-weighted imaging (T2w), diffusion-weighted imaging (DWI) with
apparent diffusion coefficient (ADC) reconstructions, and dynamic contrast enhanced (DCE) imaging, as previously described (12), according to the Prostate Imaging Reporting and Data System (PI-RADS) version 2 guidelines (13). MRI was performed on a 3-T system (Discovery MR750, General Electric Healthcare, USA) using a 32-channel pelvic phased-array coil. All MRIs were reviewed by one urogenital radiologist (IGS) with over 5 years of prostate MRI experience. Individual lesions were scored according to the PI-RADSv2 5-point likelihood scale for high-grade PCa, and the index lesions were annotated and delineated (13). Visible MRI lesions with a PI-RADS score from 3 to 5 were defined as suspicious.

**MRI-targeted biopsy**

The MRI-TRUS fusion technique was used (UroStation™, Koelis, France) to perform the targeted biopsies of all suspicious lesions, identified on MRI. The suspicious MRI lesions, delineated on DICOM images, were targeted with 2–4 cores under ultrasound guidance. Experienced operators (FHD, DFO, JFV) performed the biopsy procedures.

**Pathological review of biopsy specimens**

One expert uropathologist (GJvL) reviewed all biopsy specimens according to the International Society of Urological Pathology (ISUP) 2014 modified Gleason Score (14). GS upgrading was defined as any GS \( \geq 3+4 \) PCa found by MRI-targeted biopsies.

**Study objectives**

The primary objective of this study was to identify upgrading with MRI and MRI-targeted biopsies (benefit) in men on AS with GS 3+3. In line with this objective, the absence of upgrading (harm) despite of additional testing with MRI and targeted biopsies, was also assessed. The outcome measure is the presence/absence of upgrading to GS \( \geq 3+4 \).

The secondary objective was to assess the value of risk-stratification based on MRI (PI-RADS) and PSA-density for the presence/absence of additional upgrading. Additional analysis was performed with the outcome measure of GS \( \geq 4+3 \), focussing on risk inflation with MRI.

**Statistical analysis**

In accordance with the START recommendations, the outcome measure of clinically significant PCa for MRI and targeted biopsies is the biopsy result of GS 3+4 and higher (15). The PSA density was calculated using the MRI-measured prostate volume. The MRI-measured volume was calculated by the prolate ellipsoid formula (length x width x height x \( \pi / 6 \)). The PSA density cut-off point of 0.15 and 0.20 ng/mL\(^2\) was used for stratification (16–19). Histograms of the stratified PI-RADS and GS biopsy outcomes were constructed to visualize in which men GS upgrading did or did not occur.

Statistical tests were two-sided with the criterion of significance set at P<0.05. Statistically significant differences in continuous non-parametric patient characteristics were assessed with the Mann-Whitney U test, while the \( \chi^2 \) test for trend was used to test for differences in categorical patient characteristics; in case of small numbers the Fischer's exact test was used. Statistical analyses were performed with SPSS for Windows (Version 21.0. Armonk, NY: IBM Corp.). In addition, R version 3.4.2 and R-package ggplot2 (20) were used for visualization. Gleason scores were dichotomized using cut-off score Gleason \( \geq 3+4 \), and Gleason \( \geq 4+3 \), in which a zero indicated a GS below the cut-off and a one indicated a GS above the cut-off score.

**Results**

In 331 men on AS for GS 3+3 PCa, the median (interquartile range) age and PSA level at diagnosis were respectively 67 (range, 62–72) years and 8.0 (range, 5.6–12.0) ng/mL (Table 1). A total of 66/331 (20%) men had more than two positive systematic biopsy cores at diagnosis. A total of 155/331 (47%) men received their first MRI at baseline, while 176/331 (53%) men received their first MRI at confirmatory biopsy or at surveillance biopsy. In these men no previous MRI was performed. Men included in PRIAS did not significantly differ from men not included in PRIAS, accept for a small PSA and PSA-density difference (Table 1), reflecting PRIAS inclusion criteria.

**Benefit of additional testing with MRI and targeted biopsies**

**Upgrading in all men on AS**

In total, 25% (82/331) of men on AS showed upgrading from GS 3+3, due to additional testing by MRI and if indicated targeted biopsies. The majority (71%) was upgraded to GS 3+4, only 16% (13/82) and 13% (11/82) to GS 4+3 and GS \( \geq 4+4 \), respectively. Most of the upgraded
Table 1 Patient characteristics

| Patient characteristics of men on active surveillance | Median [IQR] or n (%) | P value* |
|-------------------------------------------------------|-----------------------|----------|
| Men on active surveillance [based on standard TRUS-guided biopsy findings (GS 3+3)] | | |
| Age (years) | 67 [62–72] | 0.574 |
| Time since diagnosis (months) | 11.5 [4–33] | 0.124 |
| PSA (ng/mL) | 8.0 [5.6–12] | 0.001 |
| Prostate volume on MRI (mL) | 46 [32–65] | 0.272 |
| PSA-density (ng/mL²) | 0.17 [0.11–0.28] | 0.001 |
| No. of positive cores at diagnosis | 1 [1–2] | 0.112 |
| No. of suspicious MRI lesions | 1 [0–1] | 0.591 |
| DRE | | |
| cT1 | 224 (68%) | 0.142 |
| cT2 | 62 (19%) | 0.074 |
| cT3 | 5 (2%) | 0.192 |
| n.a. | 40 (11%) | 0.763 |
| Total | 331 (100%) | |

Men included in PRIAS protocol

| Patient characteristics of men on active surveillance | Median [IQR] or n (%) | P value* |
|-------------------------------------------------------|-----------------------|----------|
| Age (years) | 67 [62–72] | 0.574 |
| Time since diagnosis (months) | 12 [4–37] | 0.124 |
| PSA (ng/mL) | 7.6 [5.3–10] | 0.001 |
| Prostate volume on MRI (mL) | 47 [34–67] | 0.272 |
| PSA-density (ng/mL²) | 0.14 [0.09–0.26] | 0.001 |
| No. of positive cores at diagnosis | 1 [1–2] | 0.112 |
| No. of suspicious MRI lesions | 1 [0–1] | 0.591 |
| DRE | | |
| cT1 | 128 (77%) | 0.142 |
| cT2 | 27 (16%) | 0.074 |
| cT3 | 1 (1%) | 0.192 |
| n.a. | 10 (6%) | 0.763 |
| Total | 166 (100%) | |

* Mann-Whitney U, Chi-square, Fisher’s Exact. IQR, interquartile range; TRUS, transrectal ultrasound; PSA, prostate specific antigen; DRE, digital rectal examination; MRI, magnetic resonance imaging; n.a., not available.

index lesions (82%) were categorized into PI-RADS 4 and 5. Highest GS’s were associated with PI-RADS 4 and 5.

Upgrading in men with a suspicious MRI index lesion

In 60% (198/331) a suspicious lesion was identified on MRI, and was additionally biopsied (Figure 1). 41% (82/198) of these suspicious lesions showed upgrading from GS 3+3 to GS 3+4 or higher. In PI-RADS 3, 4 and 5 index lesions, the upgrading was 30% (15/50), 36% (36/101) and 66% (31/47), respectively.

Upgrading in men with a PIRADS 3 lesion

Of all suspicious index lesions on MRI, 25% (50/198) was categorized into PI-RADS 3 (Figure 1), meaning that the MRI abnormalities are equivocal to high- or low-grade PCa. An upgrade to GS 3+4 and GS 4+3 was proven in 22% (11/50) and 8% (4/50) respectively, showing the additional value of targeting these PI-RADS 3 lesions. None of these lesions however showed GS ≥4+4.

Harms of additional testing with MRI and targeted biopsies

Abundantly or unnecessary MRI and targeted biopsies

In 40% (133/331) of men, no suspicious lesions on MRI were identified. In these men MRI testing did not result in MRI-targeted biopsies, and no further harm was attributed.
In 59% (116/198) of men with a suspicious MRI, additional MRI-targeted biopsies did not result in upgrading: GS 3+3 PCa was confirmed in 44% (87/198), and no PCa was detected in 15% (29/198). These biopsies can be considered as harm and this was most prominent in the PI-RADS 3 category (70% GS 3+3 or no PCa) (Figure 2). Similar analysis for PI-RADS 4 and PI-RADS 5 category resulted respectively in 64% and 34% unnecessary targeted biopsies.

Abundantly or unnecessary MRI and targeted biopsies adjusted to risk inflation
When looking at the detection of GS ≥4+3 PCa additional testing did not result in upgrading in respectively 93% (307/331) of all MRIs, and in 88% (174/198) of all MRI-targeted biopsies (Figure 3).

Potential strategies to reduce further harm
Excluding PI-RADS 3 lesions from targeted biopsies
Excluding PI-RADS 3 index lesions from targeted biopsies would result in a reduction of 25% (50/198) targeted biopsies, however, still missing 18% (15/82) of all upgrades to GS ≥3+4. Missed upgrading to GS ≥4+3 would be 5% (4/82).

Stratifying to PSA-density before targeting suspicious MRI lesions
Figures 4 and 5 show the number of men with targeted biopsies, stratifying to PIRADS and to PSA-density.

**PSA-density cut-off ≥0.15 ng/mL²**
Upgrades to GS ≥3+4 in PI-RADS 3 lesions were all identified in men with a PSA-density of ≥0.15 ng/mL². Hence, when first stratifying according to a PSA-density cut-off ≥0.15 ng/mL² in men with a PI-RADS 3 lesion, would result in a MRI-targeted biopsy reduction of 36% (18/50) in this category, without missing any upgrade to GS 3+4 or higher. These results are plotted in Figure 5 to visualize the amount of additional MRI testing with targeted biopsies, in men with initially low-risk disease on AS. The PSA-density thresholds of 0.15 and 0.20 ng/mL² are depicted as dotted lines.

Even for PI-RADS 4 lesions, risk stratification by PSA-density could be beneficial if adjustment to potential risk
Figure 2  PI-RADS distribution of (A) all men in active surveillance with a suspicious initial MRI, (B) men upgraded following targeted biopsies (true-positives) using outcome measure Gleason score (GS) ≥3+4, and (C) men without upgrading (true-negatives) in comparison to men with upgrading (true-positives). The right graph (C) depicts the unnecessary targeted biopsies in PI-RADS assessment category 3, 4 and 5 (dotted areas). MRI, magnetic resonance imaging; PI-RADS, MRI suspicion score.

Figure 3  Unnecessary MRIs (blue in left graphs) (left) and unnecessary targeted biopsies (blue in right graphs) for Gleason cut-off score GS ≥3+4 (upper half) and GS ≥4+3 (lower half), for detecting clinically significant prostate cancer (red) in men with initially low-risk disease, based in traditional criteria. AS, active surveillance; GS, Gleason score; MRI, magnetic resonance imaging; GS, Gleason score; PI-RADS, MRI suspicion score.
Schoots et al. Biopsy reduction in PIRADS 3 lesions by PSA-density

Figure 4 Number of positive MRIs with Gleason score outcome of MRI-targeted biopsies, stratified to PI-RADS and PSA-density. Blue: beneficial outcome of the test results, stratified to PI-RADS and PSA-density; light red: low detection of clinically significant prostate cancer, stratified to PI-RADS and PSA-density; red: high detection of clinically significant prostate cancer, stratified to PI-RADS and PSA-density. MRI, MRI, magnetic resonance imaging; PSA, prostate specific antigen.

Figure 5 Dot-plots (and integrated box-plots) of men included in active surveillance. The PI-RADS assessment category 3, 4, and 5 (x-axis) are plotted against the PSA-density (y-axis). The outcome measure is Gleason score (GS), which is dichotomized using cut-off score GS ≥3+4 (A), and GS ≥4+3 (B) for clinically significant prostate cancer. Zero [0] indicated a Gleason score below the cut-off score (blue dots and boxplots), and a one [1] indicated a Gleason score above the cut-off score (red dots and boxplots). PI-RADS, MRI suspicion score; PSA, prostate specific antigen.

inflation is performed: only 1% (1/82) of all upgrades would have been missed, reducing 43% (43/101) of targeted biopsies in this category.

PSA-density cut-off ≥0.20 ng/mL

Stratifying according to the PSA-density cut-off ≥0.20 ng/mL would result in a targeted biopsy reduction of 44% (22/50) of all PI-RADS 3 index lesions, at the cost of missing 2% (2/82) upgrades; these 2 index lesions were classified as GS 3+4. When changing outcome to GS ≥4+3, the reduction of 44% in targeted biopsies of PI-RADS 3 lesions, coincides...
with not missing any upgrades to GS 4+3 or higher (Figures 4 and 5).

**Discussion**

In our data on men on AS, additional testing with MRI and targeted biopsies could be regarded as beneficial in 25% of men: additional testing resulted in an upgrade to GS ≥3+4 as compared to the GS 3+3 PCa based on systematic TRUS-guided biopsy findings. Our results matches well with upgrading data of other studies on AS, varying from 16% to 29% (21-27). This rate of upgrading was even higher (41%) as we consider only those men who had a positive MRI, defined as PI-RADS 3 to 5. However, the relevance of identifying and acting on this upgrading cannot be adequately interpreted without the presence of long-term data on cancer-specific survival. In fact, a similar cohort of men with low-risk disease, without additional testing by MRI and targeted biopsies, has a 15-year cancer-specific survival of 94.3% (6).

Autopsy data show that many men with intermediate-risk disease (Gleason 7) are never diagnosed and therefore have clinically ‘insignificant’ cancer (28,29). The Prostate Testing for Cancer and Treatment (ProtecT) trial, which compared in a randomized controlled manner three modalities of management (active monitoring, radical prostatectomy, and external beam radiotherapy) on patients with localized PCa (30), demonstrated no significant difference in the 10-year cancer-specific survival or overall survival. In the ProtecT trial there was a difference in metastasis rate, however, favouring radical treatment at 10 years follow-up. This difference in metastasis rate is considered to result from the 25% of men who had intermediate- or high-risk disease, for whom active monitoring is clearly associated with an increased risk of progression. Nonetheless, the lack of a mortality difference emphasizes that the majority of Gleason 7 patients are not at risk in the 10-year time frame.

In the Sunnybrook surveillance cohort the 15-year metastasis rate was at least 20% in Gleason 7 cancer at initial diagnosis (31). In a recent study, however, no increase in metastasis rate or progression of intermediate risk patients on surveillance was reported compared to low risk, with up to 10 year follow-up (32). This suggests that many intermediate risk patients may still be candidates for AS (33).

The experiences described were from the pre-MRI era: men with Gleason 7 PCA at standard systematic biopsy sampling might have consisted partially of men with higher Gleason grades. Today, such patients will have the benefit of an MRI and targeted biopsies, with a high likelihood of a more accurate biopsy and hence better representative histopathology results (34). This may result in an unintended tightening of inclusion for AS, as schematically depicted (Figure 6).

We detected 25% upgrading to Gleason 7 and higher, however, only 3% was Gleason 8 and higher, identical to other published reports (26,37). It is likely that only these men with Gleason 8 and higher disease significantly influenced cancer-specific survival in men classified as having low-risk disease, in which the overall prognosis showed to be excellent in the pre-MRI era (6). We may argue that low-risk patients upgraded with targeted biopsies to intermediate-risk disease should not be excluded from AS based on the Gleason criterion alone. The higher precision of MRI and targeted biopsies may create the opportunity to specify new risk thresholds that potentially could open AS to a larger group of patients. Treatment decisions should be based on multiple parameters next to patient age and co-morbidity, including percentage of Gleason 4, growth patterns (e.g., cribriform), PSA-density, and MRI findings.

Therefore, the major challenge is accurate patient selection for AS, without the burden of intensively additional testing. The additional testing by MRI and targeted biopsies, recommended in recent reviews on MRI in AS (8,36), comes with an extra invasive procedure, increasing the burden for patients staying in AS.

In this study, MRIs were abundantly or unnecessary performed in 75% of men in AS. Furthermore, unnecessary targeted biopsies were performed in 59% in men with a suspicious MRI (Figure 7), most prominent in PI-RADS 3 and 4 assessment category (Figure 2). Unnecessary testing would even be 93% for MRI and 88% for targeted biopsies, if we accept GS 3+4 as less significant disease. This critical evaluation of additional invasive testing is not to devaluate MRI, instead, this evaluation supports exploring refinements in the role of MRI at primary diagnosis and in AS, as proposed in Figure 6.

The PI-RADS steering committee recommended to biopsy lesions with PI-RADS category 4 and 5, and not lesions with PI-RADS category 1 and 2 (13). For findings with PI-RADS category 3, biopsy may or may not be appropriate, depending on factors other than MRI alone. In our study we biopsied all PI-RADS category 3 lesions, and detected high volume Gleason 4 pattern in 8% PI-RADS 3 lesions (all GS 4+3, no GS ≥4+4). We may argue that in some circumstances this might be acceptable, reducing the additional harms of additional biopsies. However, further
Figure 6 Active surveillance and initial MRI. Men included in active surveillance (circle 1), based on traditional criteria (PSA, clinical T-stage and Gleason score by systematic ultrasound-guided biopsies) have excellent prognosis as shown by long-term follow-up of several clinical trial (6,30,35). Nowadays, men undergo additional MRI, as suggested by recent reviews (8,36). These MRIs show in more than half at least one suspicious lesion. Subsequently, these lesions are biopsied by MRI-targeted approach (circle 2). A significant proportion shows upgrading (circle 3), of which the majority is Gleason score (GS) 3+4 (circle 4). In current clinical practice all upgraded men are advocated to cease active surveillance and change into active treatment, despite good prognosis (circle 5). This suggests 'risk inflation’. In the hypothesized clinical practice, only men with upgrading to primary Gleason 4 pattern and higher are excluded from active surveillance (circle 6), correcting the present initiated 'risk inflation' by MRI. AS, active surveillance; GS, Gleason score; MRI, magnetic resonance imaging.

stratification by PI-RADS, to biopsy only PI-RADS category 4 and 5 would results in missing high volume Gleason 4 pattern, as also confirmed by other studies (23,38).

We therefore additionally investigated the combination of PSA-density and MRI to further tailor the patient risk stratification in reducing unnecessary biopsies and improving the balance between the benefit and harms of additional testing. Using the PSA-density cut-off ≥0.15 ng/mL in men with a PI-RADS 3 lesion would result in a targeted biopsy reduction of 36% in this category, without missing any upgrade to GS 3+4 or higher (Figures 4 and 5). Even for PI-RADS 4 lesions, tailored risk stratification by PSA-density could be beneficial if adjustment to risk inflation is performed (Figures 4 and 5, Table 2).

Others have confirmed this correlation, however, data in men on AS is limited (37,39-41). In a multivariate cox-regression analysis, the PSA-density was shown to be a positive predictor to detect upgrading in men on AS, with a hazard ratio of 1.72 (40). In a receiver operating characteristic (ROC) analysis by Lai et al. the optimal PSA-density cut-off point was 0.18 ng/mL with an AUC of 0.77 (39). The optimal cut-off in men on AS should be further determined in larger cohorts. In a cohort of men with initially diagnosed low-risk disease by MRI/US fusion biopsy and monitoring with serial fusion biopsies, still PSA-density was an important predictor of subsequent upgrading (41). Our study clearly suggests that men on AS with a PI-RADS 3 index lesion and a PSA-density of <0.15 ng/mL may not benefit from a follow-up biopsy.

Incorporating prostate MRI at primary PCa diagnosis will result in better discrimination between true low-risk disease and intermediate-/high-risk disease. If TRUS-guided biopsies combined with MRI and targeted biopsies are able to minimise misclassification of PCa, we may
abandon the currently used confirmation biopsy testing at 1 year in AS management, as depicted in Figure 7. MRI and targeted biopsies are increasingly used in the surveillance management of patients with clinically low-risk PCa; however, their role has not yet been established definitively (7). We may hypothesize that surveillance management of men with low-risk PCa will incorporate results from MRI and targeted biopsies into multivariate risk models in nearby future (Figure 7B) (42).

Our study comes with some limitations. First, our study is a retrospective design. Retrospective studies are known for their risk of selection bias. However, our study represents a prospectively monitored cohort of consecutive men on AS, with strict monitoring (within or without the PRIAS protocol). Second, we did not perform co-reading of MRIs, which likely would have increased detection sensitivity. However, even without co-reading the MRI detection rate in our cohort is comparable to those reported in recent publications on MRI in AS (21-27). Third, clinicians involved were not blinded to clinical data and MRI results. Hence, this process is daily clinical practice and therefore can be extrapolated to other hospitals. Fourth, the presence of standard systematic biopsy results in this study next to MRI-targeted biopsies would have shown the imperfection of MRI in detection all clinically significant PCa. Studies evaluating this added value are reporting up to 11% additionally found clinically significant PCa (8). As part of the monitoring protocol, the majority of men received their initial MRI without additional systematic biopsies. However, this group of men will decrease as a result of the increased introduction of MRI and targeted biopsies at the primary diagnostic work-up.

We acknowledge that the outcome measurement of our analysis was upgrading, based on MRI-revealed Gleason grading as recommended by the START consortium (15). Instead, the cancer-specific survival rate in long-term follow-up would have been more appropriate, especially in disputing the relevance of this high upgrading rate due to MRI targeted biopsies. However, this outcome may be debatable in a cohort of men with mostly low-risk disease that exhibits excellent long-term cancer-specific survival (6,35), and that furthermore experiences most shifts from AS to active treatment during the first 2 years of follow-up (43).

**Conclusions**

In this study on AS, we detected by MRI-targeted biopsies an upgrading to Gleason 7 and higher in 25%, however, only 3% was Gleason 8 and higher. This rate of upgrading was even higher (41%) as we consider only those men who had a suspicious finding on prostate MRI, defined as PI-RADS 3 to 5. Further stratification to PI-RADS 4–5 would have missed a small number of primary Gleason 4 PCa in the PI-RADS 3 category. Stratification with the combination of PI-RADS and PSA-density may reduce unnecessary additional MRI biopsy testing. We showed that men on AS with a PI-RADS 3 index lesion and a PSA-density of <0.15 ng/mL² will not benefit from a follow-up targeted biopsy.

The high rate of detected upgrading may result in an unintended tightening of continuing in AS. Since patients, included under current surveillance criteria showed

### Table 2

| MRI index lesions | Threshold csPCa: GS ≥3+4 | Threshold csPCa: GS ≥4+3 |
|-------------------|--------------------------|--------------------------|
|                   | No targeted biopsy | Targeted biopsy          | No targeted biopsy | Targeted biopsy          |
| Stratification into PSAD <0.15 and ≥0.15 ng/mL² | | | |
| PI-RADS 3         | P3 and PSAD <0.15      | P3 and PSAD ≥0.15       | P3 and PSAD <0.15  | P3 and PSAD ≥0.15       |
| PI-RADS 4         | P4 and any PSAD       |                           | P4 and any PSAD    |                           |
| PI-RADS 5         | P5 and any PSAD       |                           | P5 and any PSAD    |                           |
| Stratification into PSAD <0.20 and ≥0.20 ng/mL² | | | |
| PI-RADS 3         | P3 and PSAD <0.20      | P3 and PSAD ≥0.20       | P3 and PSAD <0.20  | P3 and PSAD ≥0.20       |
| PI-RADS 4         | P4 and any PSAD       |                           | P4 and any PSAD    |                           |
| PI-RADS 5         | P5 and any PSAD       |                           | P5 and any PSAD    |                           |

csPCa, clinically significant prostate cancer; GS, Gleason score; PSAD, prostate specific antigen-density; PI-RADS, MRI suspicion score.
extremely favorable outcome, there might be no need to further restrict continuing on AS with MRI and targeted biopsies. The higher precision of MRI and targeted biopsies may create the opportunity to specify new risk thresholds that potentially could open AS to a larger group of patients.

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Footnote
Conflicts of Interest: The authors have no conflicts of interest.
to declare.

**Ethical Statement:** The study was HIPAA compliant and was approved by the institutional ethical review board (NL45884.078.13/A301321). Written informed consent with guarantee of confidentiality was obtained from the participants.

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