A Ready To Use Web-Application Providing a Personalized Biopsy Schedule for Men With Low-Risk PCa Under Active Surveillance*

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Abstract

Background: Prostate cancer active surveillance (AS) patients undergo repeat biopsies. Active treatment is advised when biopsy Gleason grade group ≥ 2 (upgrading). Many patients never experience upgrading, yet undergo biopsies frequently. Personalized biopsy decisions based on upgrading-risk may

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^{*}Word count abstract (headings excluded): 300; Word count text: 2551

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reduce patient burden.

Objective: Develop a risk prediction model and web-application to assist patients/doctors in personalized biopsy decisions.

Design, Setting, and Participants: Model development: world's largest AS study PRIAS, 7813 patients, 1134 experienced upgrading; External validation: largest six cohorts of Movember Foundation's GAP3 database (> 20,000 patients, 27 centers worldwide); Data: repeat prostate-specific antigen (PSA) and biopsy Gleason grade.

Outcome Measurements, and Statistical Analysis: A Bayesian joint model fitted to the PRIAS dataset. This model was validated in GAP3 cohorts using risk prediction error, calibration, area under ROC (AUC). Model and personalized biopsy schedules based on predicted risks were implemented in a web-application.

Results and Limitations: Cause-specific cumulative upgrading-risk at year five of follow-up: 35% in PRIAS, at most 50% in GAP3 cohorts. PRIAS based model: PSA velocity was stronger predictor of upgrading (Hazard Ratio: 2.47, 95%CI: 1.93–2.99) than PSA value (Hazard Ratio: 0.99, 95%CI: 0.89–1.11). Validation: Moderate AUC (0.55–0.75) in PRIAS and GAP3 cohorts. Moderate prediction error (0.1–0.3) in GAP3 cohorts where impact of PSA value and velocity on upgrading-risk was similar to PRIAS, but large (0.3–0.45) otherwise. Recalibration of baseline risk required for external cohorts.

Conclusions: We successfully developed and validated a model for predicting upgrading-risk, and providing risk-based personalized biopsy decisions, in prostate cancer AS. The model made available via a web-application enables shared decision making of biopsy schedules by comparing fixed and personalized schedules on total biopsies and expected time delay in detecting upgrading.

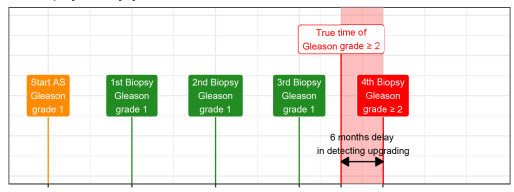
Patient Summary: Personalized prostate biopsies are a novel alternative to fixed one-size-fits-all schedules. The underlying statistical models are made available through a user-friendly web-application and may help to reduce unnecessary prostate biopsies while maintaining cancer control.

Keywords: Active Surveillance, Biopsies, Personalized Medicine, Prostate Cancer, Shared Decision Making

1. Introduction

- Patients with low- and very low-risk screening-detected localized prostate
- cancer are usually recommended active surveillance (AS) instead of immedi-
- 4 ate radical treatment [1]. In AS, cancer progression is routinely monitored
- ⁵ via prostate-specific antigen (PSA), digital rectal examination, and repeat
- 6 biopsies. Among these, the strongest indicator of cancer-related outcomes is
- the biopsy Gleason grade group [2]. When it increases from group 1 (Glea-
- s son 3+3) to 2 (Gleason 3+4) or higher, called upgrading [3], patients are
- 9 commonly advised curative treatment [4].
- Usually, AS protocols schedule biopsies periodically. Consequently, up-

A Biopsy every year



B Biopsy every 2 years

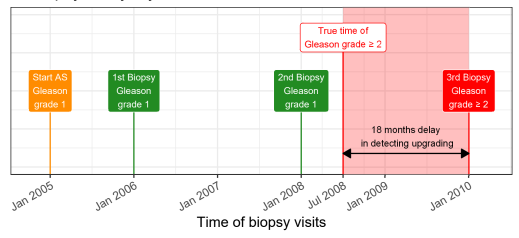


Figure 1: Trade-off between the number of biopsies and time delay in detecting upgrading (Increase in Gleason grade group from 1 to 2 or higher): The true time of upgrading for the patient in this figure is July 2008. When biopsies are scheduled annually (Panel A), upgrading is detected in January 2009 with a time delay of six months, and a total of four biopsies are scheduled. When biopsies are scheduled biennially (Panel B), upgrading is detected in January 2010 with a time delay of 18 months, and a total of three biopsies are scheduled. Since biopsies are conducted periodically, the time of upgrading is observed as an interval. For example, between Jan 2008–Jan 2009 in Panel A and between Jan 2008–Jan 2010 in Panel B. The phrase 'Gleason grade group' is shortened to 'Gleason grade' for brevity.

grading is always detected with a time delay (Figure 1). For detecting upgrading timely, many AS programs schedule fixed and frequent biopsies (e.g., annually) for all patients [5, 6]. However, this leads to unnecessary biop-13 sies in slow/non-progressing patients. Biopsies are invasive, may be painful, and are prone to medical complications such as bleeding and septicemia[7]. Biopsy burden and patient non-compliance to frequent biopsies [8] have raised concerns regarding the optimal biopsy schedule [9, 10]. In this regard, in some cohorts, magnetic resonance imaging (MRI) is employed for targeted biopsies and to study the value for tumor monitoring. Although, due to currently limited AS data, MRI's value is not clear. Others have proposed the option of scheduling biopsies infrequently (e.g., biennially) [9, 11]. However, due to differences in baseline upgrading-risk across cohorts [9], fixed biopsy schemes can still lead to many unnecessary biopsies. A promising alternative to fixed and frequent biopsies is personalized biopsy schedules based on the patient-specific upgrading-risk (Figure 2).

The first challenge in creating personalized biopsy schedules is developing a statistical model to consolidate accumulated patient data (e.g., PSA, previous biopsy results) into estimates for upgrading-risk. Existing upgrading-risk [12, 13] calculators use only the latest PSA measurement of a patient. Comparatively, more information is captured by considering all repeatedly measured PSA, previous biopsy results, and baseline characteristics of a patient. To this end, a suitable model is the joint model for time-to-event and longitudinal data [14, 15, 16]. A joint model predicts the upgrading-risk in a personalized manner. However, a subsequent challenge is translating predicted risks into clinical decisions. For example, a 10% upgrading-risk can

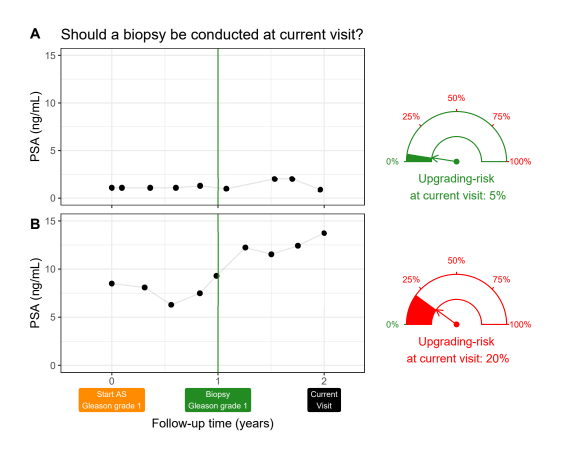


Figure 2: Motivation for personalized upgrading-risk based decisions of biopsy: Patient A (Panel A) and B (Panel B) had their latest biopsy at year one of follow-up (green vertical line). Patient A's prostate-specific antigen (PSA) profile remained stable until his current visit at year two, whereas patient B's profile has shown a rise. Consequently, patient B's upgrading-risk at the current visit (year two) is higher than that of patient A. This makes patient B a more suitable candidate for biopsy than Patient A. Risk estimates in this figure are only illustrative.

be perceived high/low depending upon the patient's age. Patients may also weigh risks of upgrading with the potential consequences of another biopsy. Two relevant consequences of biopsies (Figure 1) are the timing and the total number of planned biopsies (burden), and the time delay in detecting upgrading (smaller is beneficial). The relative importance of these consequences can vary between the patients, and also over the follow-up period for the same patient. The goal of this work is two-fold. First, to develop a robust, generalizable model that gives reliable estimates for individual upgrading-risk in AS. Second, to utilize the predicted upgrading-risks to create personalized biopsy schedules. To facilitate shared decision making of biopsy schedules, we also intend to provide quantitative estimates of the aforementioned consequences of opting for a personalized versus the standard fixed schedule. For developing our model, we will use the world's largest AS dataset PRIAS. Subsequently, we want to externally validate our model in the largest six AS cohorts from the Movember Foundation's GAP3 database [17]. Last, we intend to implement our model and methodology in a web-application.

2. Patients and Methods

54 2.1. Study Cohort

For developing a statistical model to predict upgrading-risk, we used the world's largest AS dataset, Prostate Cancer International Active Surveillance or PRIAS [4] available as of April 2019 (Table 1). In PRIAS, PSA was measured quarterly for the first two years of follow-up and semiannually thereafter. Biopsies were scheduled at year one, four, seven, and ten of

follow-up. Additional yearly biopsies were scheduled when PSA doubling time was between zero and ten years.

We selected all 7813 patients who had Gleason grade group 1 at the time of inclusion in PRIAS. Our primary event of interest is an increase in this Gleason grade group upon repeat biopsy, called *upgrading* (1134 patients). Upgrading is a trigger for treatment advice in PRIAS. Also, 2250 patients were provided treatment based on their PSA, number of biopsy cores with cancer, or anxiety/other reasons. Our reasons for focusing solely on upgrading are, namely, upgrading is strongly associated with cancer-related outcomes, and other triggers for treatment vary between cohorts [5].

For model validation, we selected the following largest (by number of repeated measurements) six cohorts from Movember Foundation's GAP3 database version 3.1 [17]: University of California San Francisco AS (UCSF, version 3.2), University of Toronto AS (Toronto), Johns Hopkins AS (Hopkins), Memorial Sloan Kettering Cancer Center AS (MSKCC), King's College London AS (KCL), and Michigan Urological Surgery Improvement Collaborative AS (MUSIC). Only patients with a Gleason grade group 1 at the time of inclusion in these cohorts were selected. Summary statistics are presented in Supplementary A.2.

79 2.2. Statistical Model

For developing an upgrading-risk prediction model, the available data in the PRIAS cohort was patient age at inclusion in AS, longitudinally measured PSA, timing of repeat biopsies and Gleason grades, and observed time of upgrading. Analysis of this data required modeling the within-patient correlation for PSA, the association between the Gleason grades and PSA

Table 1: **Summary of the PRIAS dataset as of April 2019**. The primary event of interest is upgrading, that is, increase in Gleason grade group from group 1 [2] to 2 or higher. IQR: interquartile range, PSA: prostate-specific antigen. Study protocol URL: https://www.prias-project.org

Characteristic	Value
Total centers	> 100
Total patients	7813
Upgrading (primary event)	1134
Treatment	2250
Watchful waiting	334
Loss to follow-up	249
Death (unrelated to prostate cancer)	95
Death (related to prostate cancer)	2
Median age at diagnosis (years)	66 (IQR: 61–71)
Median maximum follow-up per patient (years)	1.8 (IQR: 0.9-4.0)
Total PSA measurements	67578
Median number of PSA measurements per patient	6 (IQR: 4–12)
Median PSA value (ng/mL)	5.7 (IQR: 4.1–7.7)
Total biopsies	15686
Median number of biopsies per patient	2 (IQR: 1–2)

profiles of a patient, and handling missing PSA measurements after a patient experienced upgrading. In such situations, a commonly used model is the joint model for time-to-event and longitudinal data [14, 15, 16].

Our joint model consisted of two sub-models. First, a linear mixed submodel [18] for longitudinally measured PSA (log-transformed). Second, a relative-risk sub-model (similar to the Cox model) for obtaining the causespecific upgrading-risk. We included patient age in both sub-models. In the PSA sub-model, we fitted a unique curve to the PSA measurements of each patient (Panel A, Figure 3). Subsequently, we calculated the mathematical derivative of the patient's fitted PSA profile (Equation 2, Supplementary A), to obtain his follow-up time specific instantaneous PSA velocity (Panel B. Figure 3). This instantaneous velocity is a stronger predictor of upgrading than the widely used average PSA velocity [19]. We modeled the impact of PSA on upgrading-risk by including fitted PSA value and instantaneous velocity as predictors in the relative-risk model. Last, the time of the latest negative biopsy was utilized in the relative-risk sub-model (Panel C, Figure 3). The parameters of the two sub-models were estimated jointly (Sup-101 plementary A) using the R package **JMbayes** [20]. 102

2.3. Risk Prediction and Model Validation

The predictions for upgrading-risk from our model are made for the entire future follow-up period of a patient. These predictions automatically
update over follow-up as more patient data becomes available (Figure 5,
Supplementary B). We validated our PRIAS based model internally in the
PRIAS cohort, and externally in the largest six GAP3 database cohorts. We
employed calibration plots [21, 22] and follow-up time-dependent mean ab-

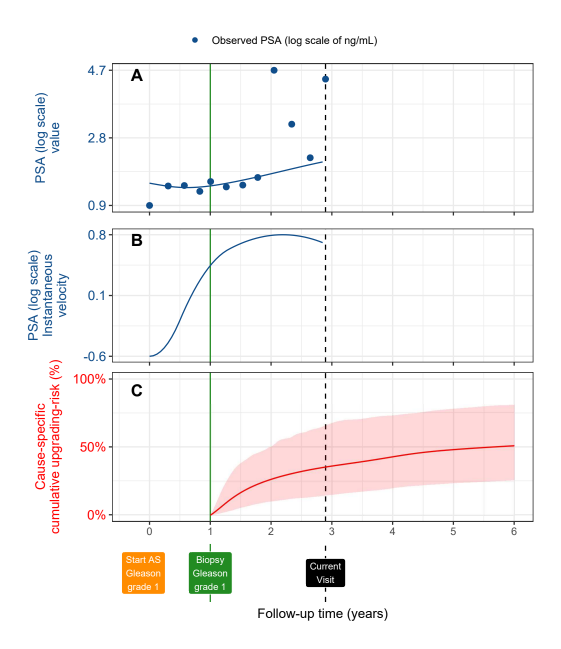


Figure 3: Illustration of the joint model on a real PRIAS patient. Panel A: Observed PSA (blue dots) and fitted PSA (solid blue line), log-transformed from ng/mL. Panel B: Estimated instantaneous velocity of PSA (log-transformed). Panel C: Predicted cause-specific cumulative upgrading-risk (95% credible interval shaded). Upgrading is defined as an increase in the Gleason grade group from group 1 [2] to 2 or higher. This upgrading-risk is calculated starting from the time of the latest negative biopsy (vertical green line at year 1 of follow-up). The joint model estimated it by combining the fitted PSA (log scale) value and instantaneous velocity, and time of the latest negative biopsy. Black dashed line at year 3 denotes the time of current visit.

solute risk prediction error or MAPE [23] to graphically and quantitatively evaluate our model's risk prediction accuracy, respectively. We assessed our model's ability to discriminate between patients who experience/do not experience upgrading via the time-dependent area under the receiver operating characteristic curve or AUC [23].

The aforementioned time-dependent AUC and MAPE [23] are temporal extensions of their standard versions [22] in a longitudinal setting. More specifically, at every six months of follow-up we calculated a unique AUC and MAPE for predicting upgrading-risk in the subsequent one year (Supplementary B.1). For emulating a realistic situation, we calculated the AUC and MAPE at each follow-up using only the validation data available until that follow-up. Lastly, to resolve any potential model miscalibration in validation cohorts, we aimed to recalibrate our model's baseline hazard of upgrading (Supplementary A), individually for each cohort.

3. Results

The cause-specific cumulative upgrading-risk at year five of follow-up was 35% in PRIAS, and at most 50% in the six validation cohorts (Panel B, Figure 4). That is, many patients may not require all biopsies planned in the first five years of AS. In the PRIAS based fitted model, the adjusted hazard ratio (aHR) of upgrading for an increase in patient age from 61 to 71 years (25-th to 75-th percentile) was 1.45 (95%CI: 1.30–1.63). For an increase in fitted PSA value (log scale) from 2.36 to 3.07 (25-th to 75-th percentile), the aHR was 0.99 (95%CI: 0.89–1.11). The strongest predictor of upgrading-risk was instantaneous PSA velocity, with an increase from -0.09 to 0.31 (25-th to

75-th percentile), leading to an aHR of 2.47 (95%CI: 1.93–2.99). The aHR for PSA value and velocity varied between cohorts (Table 8, Supplementary A.2). 135 The time-dependent MAPE; time-dependent AUC; and calibration plot 136 of our model in different validation cohorts are shown in Panel B, Figure 8, Supplementary B; Panel A, Figure 4; and Panel B, Figure 4, respectively. In 138 all cohorts, time-dependent AUC was moderate (0.55 to 0.75) over the whole 139 follow-up period. Time-dependent MAPE was large (0.3 to 0.45) in those 140 cohorts where the impact of PSA value and velocity on upgrading-risk was 141 different from PRIAS (e.g., MUSIC cohort, Table 8, Supplementary A.2), and moderate (0.1 to 0.3) otherwise. In all cohorts, the MAPE decreased rapidly after year one of follow-up. Our model was miscalibrated for validation cohorts (Panel B, Figure 4). We resolved this by recalibrating the baseline hazard of upgrading in all cohorts (Figure 6, Supplementary B). We compared risk predictions from the recalibrated models with predictions from separately fitted joint models to each cohort (Figure 7, Supplementary B). 148 The difference in predictions was lowest in Johns Hopkins cohort (impact of PSA similar to PRIAS, Table 5, Supplementary A.2). Comprehensive results are in Supplementary A.2 and B. 151

3.1. Personalized Biopsy Schedules

We utilized the PRIAS based model to create personalized biopsy schedules. Specifically, first, we utilized the model to predict a patient's causespecific cumulative upgrading-risk on his future follow-up visits (usually every six months in PRIAS) based on his available data (Figure 5). We then
planned biopsies on those visits where the patient's conditional cumulative
upgrading-risk was more than a certain threshold (Supplementary C). Exam-

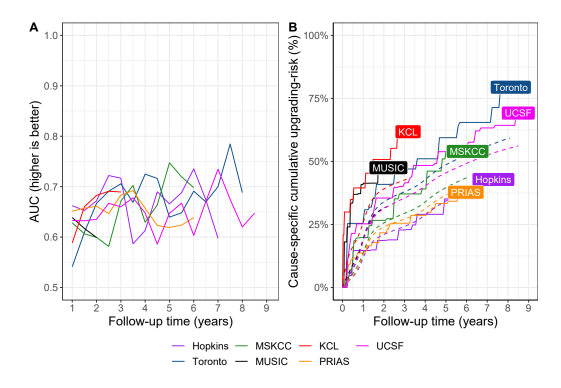


Figure 4: Model Validation Results. Panel A: time-dependent area under the receiver operating characteristic curve or AUC (measure of discrimination). Panel B: calibration-at-large indicates model miscalibration. This is because solid lines depicting the non-parameteric estimate of the cause-specific cumulative upgrading-risk [24], and dashed lines showing the average cause-specific cumulative upgrading-risk obtained using the joint model fitted to the PRIAS dataset, are not overlapping. Same plot after recalibration is shown in Figure 6, Supplementary B. Full names of Cohorts are PRIAS: Prostate Cancer International Active Surveillance, Toronto: University of Toronto Active Surveillance, Hopkins: Johns Hopkins Active Surveillance, MSKCC: Memorial Sloan Kettering Cancer Center Active Surveillance, KCL: King's College London Active Surveillance, MU-SIC: Michigan Urological Surgery Improvement Collaborative Active Surveillance, UCSF: University of California San Francisco AS.

ple personalized schedules based on 5% and 10% risk thresholds are shown in Figure 5, and in Figure 9–11, Supplementary C. For both personalized and fixed schedules, we estimated the expected time delay in detecting progression if a patient follows them (Panel C, Figure 5). The delay is calculated in a patient-specific manner (Supplementary C) and is updated as more data becomes available over follow-up. Using expected delay and schedule of biopsies as criteria, patients/doctors can compare fixed schedules with personalized schedules based on different risk thresholds.

3.2. Web-Application

We implemented our model and personalized schedules in a user-friendly web-application https://emcbiostatistics.shinyapps.io/prias_biopsy_
recommender/. Currently, the web-application supports PRIAS and the six validation cohorts. Patient data can be entered manually and in Microsoft Excel format. Predictions for upgrading-risk are available for a currently limited, cohort-specific, follow-up period (Table 7, Supplementary C). The web-application visualizes the timing of biopsies, and expected time delay in detecting upgrading, for personalized schedules based on 5%, 10%, and 15% risk threshold; annual biopsies; biennial biopsies; and PRIAS schedule.

77 4. Discussion

We successfully developed and externally validated a model for predicting upgrading-risk [3] in prostate cancer AS, and providing risk-based personalized biopsy decisions. Our work has four novel features over earlier
risk calculators [15, 25]. First, our model was fitted to the world's largest
AS dataset PRIAS and externally validated in the largest six cohorts of

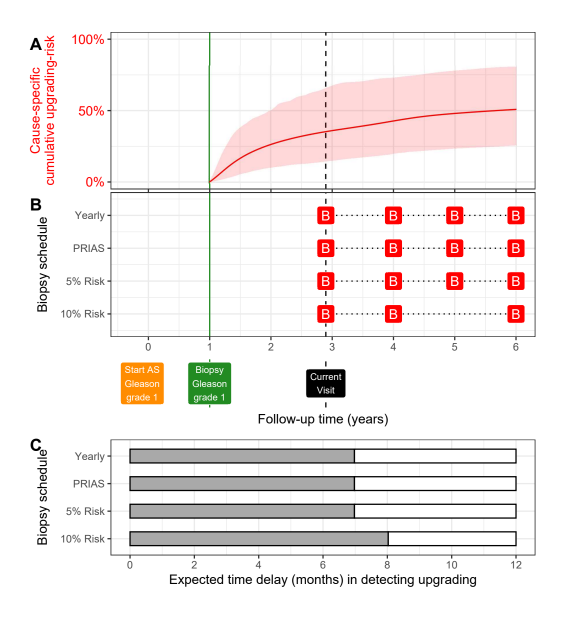


Figure 5: Illustration of personalized and fixed schedules of biopsies. Due to a lack of space, the PSA profile of this patient is shown in Figure 3. Panel A: Predicted cumulative upgrading-risk (95% credible interval shaded). Panel B: Personalized and fixed schedules of biopsies, with a red 'B' indicating a scheduled biopsy. The green vertical line at year 1 denotes the time of the latest negative biopsy. Black dashed line at year 3 denotes the time of the current visit. Panel C: Expected time delay in detecting upgrading (months) for different schedules. A compulsory biopsy was scheduled at year six (maximum biopsy scheduling time in PRIAS, Supplementary C) in all schedules for a meaningful comparison between them.

the Movember Foundation's GAP3 database [17]. Second, the model predicts a patient's current and future upgrading-risk in a personalized manner. Third, we use the risks to make a personalized schedule, and to also calculate expected time delay in detecting upgrading (less is beneficial) if that schedule is followed. Thus, patients/doctors can compare schedules before making a choice. Fourth, we implemented our methodology in a user-friendly web-application (https://emcbiostatistics.shinyapps.io/prias_biopsy_recommender/) for PRIAS and validated cohorts.

Our model is useful for a large number of patients from PRIAS and 191 validated cohorts. Across these cohorts, our model had a moderate time-192 dependent AUC (0.55–0.75), a measure of discrimination. The moderate 193 AUC can be explained by the fact that, unlike the standard AUC [22], the time-dependent AUC utilizes only the validation data available until the time at which it is calculated. The same holds for the time-dependent MAPE (mean absolute prediction error), although it varied much more between cohorts than AUC. It was moderate in cohorts where the effect size for impact of PSA value and velocity on upgrading-risk was similar to that for PRIAS 199 (e.g., Hopkins cohort). Otherwise, as in the case of KCL or MUSIC cohorts, 200 the MAPE was large. In all cohorts, MAPE decreased rapidly after year one of follow-up. This may be explained by the fact at year one the validation 202 data also consists of those patients who may have been misclassified incorrectly as Gleason grade group 1 at the time of inclusion in AS. The currently practiced compulsory biopsy at year one of follow-up for all patients may obviate this issue. Last, we required recalibration of our model's baseline hazard of upgrading for all validation cohorts.

The clinical implications of our work are as follows. First, the cause-208 specific cumulative upgrading-risk at year five of follow-up was at most 50% 209 in all cohorts (Panel B, Figure 4). That is, many patients may not require 210 all biopsies planned in the first five years of AS. Given the non-compliance and burden of frequent biopsies [8], the availability of our methodology as a 212 web-application may encourage patients/doctors to consider upgrading-risk 213 based personalized schedules instead. An additional advantage of personal-214 ized schedules is that they update as more patient data becomes available 215 over follow-up. We have shown via a simulation study [26] that personalized schedules may reduce on average six biopsies compared to annual schedule and two biopsies compared to PRIAS schedule in slow/non-progressing 218 AS patients, while maintaining almost the same time delay in detecting progression as PRIAS schedule. Personalized schedules with different risk thresholds indeed have different performance. In this regard, to assist patients/doctors in choosing between various personalized, and fixed schedules, 222 the web-application provides a patient-specific estimate of expected time delay in detecting upgrading, for both personalized and fixed schedules. We hope that this will objectively address patient apprehensions regarding adverse outcomes in AS.

This work has certain limitations. Predictions for upgrading-risk and personalized schedules are available only for a currently limited, cohort-specific, follow-up period (Table 7, Supplementary C). This problem can be mitigated by refitting the model with new follow-up data in the future. Recently, some cohorts utilize MRI to explore the possibility of targeting visible lesions by biopsy. Currently GAP3 database has limited MRI follow-up data available.

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As more such data becomes available, the current model can be extended to include MRI based predictors. We scheduled biopsies using cause-specific cumulative upgrading-risk. Accounting for competing events, such as treatment based on the number of positive biopsy cores, may lead to improved personalized biopsy decisions. Although, in this work, we did not consider such additional triggers for treatment because, unlike upgrading, they differ between cohorts [5]. Upgrading is susceptible to inter-observer variation too. Models which account for this variation [15, 27] will be interesting to investigate further. However, the methodology for personalized scheduling, and for comparison of various schedules need not change.

5. Conclusions

We successfully developed and validated a model for predicting upgradingrisk, and providing risk-based personalized biopsy decisions, in prostate cancer AS. The model made available via a user-friendly web-application (https:
//emcbiostatistics.shinyapps.io/prias_biopsy_recommender/) enables
shared decision making of biopsy schedules by comparing fixed and personalized schedules on total biopsies and expected time delay in detecting upgrading. Novel biomarkers and MRI data can be added as predictors in the model
to improve predictions in the future. Recalibration of baseline upgrading-risk
is advised for external cohorts.

3 Author Contributions

Anirudh Tomer had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data anal-

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   zopoulos
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    Obtaining funding: Roobol, Steverberg, and Rizopoulos
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    Supervision: Roobol, and Rizopoulos
    Other: none
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69 Acknowledgments

The first and last authors would like to acknowledge support by Nederlandse Organisatie voor Wetenschappelijk Onderzoek (the national research
council of the Netherlands) VIDI grant nr. 016.146.301, and Erasmus University Medical Center funding. Part of this work was carried out on the
Dutch national e-infrastructure with the support of SURF Cooperative. The
authors also thank the Erasmus University Medical Center's Cancer Computational Biology Center for giving access to their IT-infrastructure and
software that was used for the computations and data analysis in this study.

The PRIAS website is funded by the Rotterdam Prostate Cancer Research
Foundation Rotterdam (SWOP). This work was supported by the Movember

Foundation. The funder did not play any role in the study design, collection, analysis or interpretation of data, or in the drafting of this paper.

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don Hospital Trust, London, UK), Jenna Kimberly-Duffell (University of Cambridge & Cambridge University Hospitals NHS Foundation Trust, Cam-355 bridge, UK), Aida Santaolalla (King's College London, London, UK & Guys 356 and St Thomas NHS Foundation Trust, London, UK), Daan Nieboer (Erasmus Medical Center, Rotterdam, The Netherlands), Jonathan Olivier (Lille 358 University Hospital Center, Lille, France), Tiziana Rancati (Fondazione IR-350 CCS Istituto Nazionale dei Tumori di Milano, Milan, Italy), Heln Ahlgren 360 (Sahlgrenska University Hospital, Gteborg, Sweden), Juanma Mascars (Insti-361 tuto Valenciano de Oncologa, Valencia, Spain), Annica Lfgren (Skne University Hospital, Malm, Sweden), Kurt Lehmann (Kantonsspital Baden, Baden, 363 Switzerland), Catherine Han Lin (Monash University and Epworth Health-Care, Melbourne, Australia), Hiromi Hirama (Kagawa University, Kagawa, Japan), Kwang Suk Lee (Yonsei University College of Medicine, Gangnam Severance Hospital, Seoul, Korea). Research Advisory Committee: Guido Jenster (Erasmus MC, Rotterdam, 368 the Netherlands), Anssi Auvinen (University of Tampere, Tampere, Finland), Anders Bjartell (Skne University Hospital, Malm, Sweden), Masoom Haider (University of Toronto, Toronto, Canada), Kees van Bochove (The Hyve B.V. Utrecht, Utrecht, the Netherlands), Ballentine Carter (Johns Hopkins University, Baltimore, USA until 2018). 373 Management team: Sam Gledhill (Movember Foundation, Melbourne, 374 Australia), Mark Buzza / Michelle Kouspou (Movember Foundation, Mel-375 bourne, Australia), Chris Bangma (Erasmus Medical Center, Rotterdam, 376 The Netherlands), Monique Roobol (Erasmus Medical Center, Rotterdam, The Netherlands), Sophie Bruinsma / Jozien Helleman (Erasmus Medical

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