# PRIAS Personalized Biopsy Schedules

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#### Abstract

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## 1 Introduction

Prostate cancer is the development of cancer in the prostate gland. With increase in life expectancy and increase in number of screening tests, an increase in diagnosis of low grade prostate cancers has been observed. Majority of these cancers have good long-term survival and in many cases the prostate cancer is (over) diagnosed solely due of screening. i.e. it wouldn't have shown any malignant symptoms for a long time otherwise. To avoid overtreatment, patients diagnosed with prostate cancer are often motivated to join active surveillance (AS) programs instead of taking immediate treatment. The goal of AS programs is to routinely check the progression of prostate cancer and avoid serious treatments such as surgery or chemotherapy as long as they are not needed.

Currently the largest AS program worldwide is PRIAS (www.prias-project.org) (Bokhorst et al., 2015). Patients enrolled in PRIAS are closely monitored using serum prostate-specific antigen (PSA) levels, digital rectal examination (DRE) and repeat prostate biopsies. Biopsies are evaluated using the Gleason grading system. Gleason scores range between 2 and 10, with 10 corresponding to a very serious state of prostate cancer. Patients who join PRIAS have a Gleason score of 6 or less, DRE score of cT2c or less and a PSA of 10 ng/mL or less at the time of induction. Although a PSA doubling time(measured as the inverse of the slope of regression line through the base 2 logarithm of PSA values) of less than 3 years, DRE of cT3 or more, and a Gleason score more than 6 are indicators of prostate cancer progression, only DRE and Gleason scores are considered to be the gold standard in this regard (Bokhorst et al., 2016). If either the DRE or the Gleason score are found to be above the aforementioned threshold, then it is considered that the disease has progressed and the patient is removed from AS for further curative treatment. When the Gleason score becomes greater than 6, it is also known as Gleason reclassification (referred to as GR here onwards).

The reliability of Gleason score comes at a high cost. Biopsies are difficult to obtain, are painful and have serious side effects such as hematuria and sepsis for prostate cancer patients (Loeb et al., 2013). So much so, that PRIAS as well as majority of the AS programs around the world strongly adhere to the rule of not having more than 1 biopsy per year. Performing a biopsy every year has the advantage that it is possible to detect GR within 1 year since its occurrence. The drawbacks of this schedule though, are not only medical but also financial. Keegan et al., 2012 have shown that if a biopsy is performed every year then the costs of AS per head, at 10 years of follow-up

exceed the costs of treatment (brachytherapy or prostatectomy) at 6 and 8 years of follow-up, respectively. They also found that performing biopsy every other year led to 99% increase in savings (AS vs. primary treatment) per head over a period of 10 years compared to the scenario where biopsy is performed every year. Despite this, several AS studies schedule biopsies for every patient annually (Tosoian et al., 2011; Welty et al., 2015). For patients enrolled in PRIAS the schedule is comparatively less rigorous. One biopsy is performed at the time of induction, and the rest are scheduled at 1, 4, 7, 10 years and every 5 years thereafter. For patients who have a PSA doubling time (PSA-DT) less than 10 years, repeat biopsy every year is advised.

The biopsy schedule of PRIAS program is less rigorous than other programs, and yet PRIAS has a high non compliance rate for repeat biopsies. Bokhorst et al., 2015 reported that the percentage of men receiving repeat biopsies decreased from 81% at year 1 to 60% in year 4, 53% in year 7 and 33% in year 10 of follow up. Non compliance of biopsy schedule reduces the effectiveness of AS programs, as progression is detected late. On the other hand even if patients comply with the schedule, be it annually or the schedule of PRIAS, it may not be suitable for them. A patient whose cancer progresses slowly will often end up having biopsies when they are not needed. For a patient who has a faster progressing disease, crude measures such as PSA-DT are employed to decide if frequent biopsies are required. The fact that existing schedules require improvement is also evident in some of the reasons given by patients for non-compliance: 'patient does not want biopsy', 'PSA stable', 'complications on last biopsy'and 'no signs of disease progression on previous biopsy'.

Let us assume that we have a new patient j enrolled in the AS program. The most useful biopsy schedule for him will be the one with the least number of biopsies  $N_i^b$  and the smallest offset  $O_j = T_j^o - T_j^*$  possible, where  $T_j^o$  is the time at which GR is observed and  $T_j^*$  is the actual time at which GR occurred. The search for the most useful biopsy schedule is the motivation behind this work. To this end, we have proposed alternative biopsy schedules, belonging to a class of schedules called personalized schedules. Personalized schedules are tailored separately for every patient and every disease. A simple example is the PRIAS schedule, which is personalized since it depends on the PSA-DT of the patient, an indicator of the state of disease. More sophisticated personalized schedules have been developed in the past. For e.g. Bebu and Lachin, 2017 have proposed Markov models based cost optimized personalized schedules. O'Mahony et al., 2015 have proposed cost optimized personalized equi-spaced screening intervals, using Microsimulation Screening Analysis (MISCAN) models. Parmigiani, 1998 have used information theory to come up with schedules for detecting time to event in the smallest possible time interval. Most of these methods however create an entire schedule in advance. Rizopoulos et al., 2016 have proposed dynamic personalized schedules for longitudinal markers using joint models for time to event and longitudinal data(Rizopoulos, 2012; Tsiatis and Davidian, 2004).

The personalized schedules we have proposed in this paper, utilize joint models and are dynamic. i.e. at a time only one future visit is scheduled, based on all the information gathered up to that point in time. More specifically, we have proposed two types of personalized schedules. One based on expected time of GR of a patient and the second based on the risk of GR. We have also analyzed an approach where the two types of personalized schedules are combined. Both types of schedules not only consider a patient's measurable attributes such as age, but also latent patient to patient variations in health, which cannot be measured directly. Results from previous repeat biopsies of the patient and PSA measurements as well as the population level information about hazard of GR, are used by the personalized schedules that we have proposed. It is important to note that a schedule for DRE measurements is not of interest since it is a non invasive procedure and has no serious medical implications. Thus the only event of interest is GR and not disease progression or DRE crossing the threshold of cT2c.

Using joint models to model the PSA measurements and risk of GR has the advantage that the association between the two is also modeled. More importantly, the association is modeled via random effects, and therefore the models have an inherent patient specific nature. Secondly, joint models allow modeling the entire longitudinal history of PSA measurements, which is more sophisticated than PSA-DT. The use of PSA measurements in creating a personalized schedule is important because PSA is easy to measure, is cost effective and does not have any side effects. Secondly, in PRIAS Bokhorst et al., 2015 found that compliance rate for PSA measurements was

as high as 91%. They also showed that there were more men who had a Gleason score greater than 6 as well as PSA-DT less than 3 years compared to men who had Gleason > 6 as well as PSA-DT larger than 3 years. i.e. Information from PSA was found to be indicative of GR. Lastly, some patients/doctors in PRIAS did not comply with the biopsy schedule because they considered PSA to be stable. However, if information from PSA is used in a methodical manner, it can lead to a more informative medical decision making process.

The rest of the paper is organized as follows. Section 2.1 covers briefly the joint modeling framework in context of the problem at hand. Section 2.2 details the personalized scheduling approaches we have proposed in this paper. In section 3 we demonstrate the efficacy of personalized schedules in a real world scenario by employing them for the patients from the PRIAS study. Lastly, in section 4, we present the results from a simulation study we conducted, to compare personalized schedules with the schedule of PRIAS study, as well with the most aggressive biopsy schedule of doing annual biopsies.

## 2 A framework for personalized biopsy schedules

The first step in creating a personalized schedule for biopsies is to come up with a model for Gleason scores, PSA levels and other patient specific characteristics. In PRIAS, PSA levels are measured at the time of induction, every 3 months for the first 2 years in the study and then every 6 months thereafter. Thus PSA levels can be modeled as a longitudinal outcome. As mentioned earlier, patients in PRIAS have a Gleason score of 6 or less at the time of induction in the study, and patients are removed from AS the first time GR takes place. Since our interest lies in finding the time of GR, we model it as a time to event outcome. While univariate modeling of the two aforementioned outcomes can be done separately using longitudinal model and survival models, but they are not independent. The association between the two exists because both are affected by the state of prostate cancer. To model the association between the two types of outcomes we use a joint model for time to event and longitudinal outcomes.

### 2.1 Joint model for risk of GR and PSA levels

Let  $T_i^*$  denote the true GR time for the  $i^{th}$  patient enrolled in an AS program. Let the vector of times at which a total of  $g_i$  biopsies are conducted for the  $i^{th}$  patient be denoted by  $C_i = \{C_{i0}, C_{i1}, ... C_{ig_i}; C_{ij} < C_{ik}, \forall j < k\}$ .  $T_i^*$  cannot be observed directly and it is only known that it falls in an interval  $(l_i, r_i]$ , where  $l_i = C_{i(g_{i-1})}, r_i = C_{ig_i}$  if GR is observed, and  $l_i = C_{ig_i}, r_i = \infty$  if patient drops out of AS. The latter is also known as right censoring. Let  $\mathbf{y}_i$  denote the  $n_i \times 1$  longitudinal outcome vector for the PSA levels of the  $i^{th}$  patient. The population of interest is all the patients enrolled in AS. For a sample of n patients from this population the complete data is denoted by  $\mathcal{D}_n = \{T_i, l_i, r_i, \mathbf{y}_i; i = 1, ...n\}$ , where  $T_i \in (l_i, r_i]$  denotes the observed time of GR.

To model the evolution of the PSA measurements over time, the joint model utilizes a linear mixed effects model. The longitudinal outcome  $y_i(t)$  at a time t is modeled as:

$$y_i(t) = m_i(t) + \varepsilon_i(t),$$
  
=  $\boldsymbol{x}_i^T(t)\boldsymbol{\beta} + \boldsymbol{z}_i^T(t)\boldsymbol{b}_i + \varepsilon_i(t)$ 

where,  $m_i(t)$  denotes the true and unobserved value of the longitudinal outcome at time t. The measurement error  $\varepsilon_i(t) \sim N(0, \sigma^2)$  is assumed normally distributed with variance  $\sigma^2$ .  $\boldsymbol{\beta}$  denotes the vector of the unknown fixed-effects parameters.  $\boldsymbol{b}_i \sim N(0, \boldsymbol{D})$  denotes the  $q \times 1$  vector of random effects, assumed normally distributed with mean zero, and  $q \times q$  covariance matrix  $\boldsymbol{D}$ .  $\boldsymbol{b}_i$  and  $\varepsilon_i(t)$  are assumed independent.  $\boldsymbol{x}_i(t)$  and  $\boldsymbol{z}_i(t)$  denote row vectors of the design matrices for the fixed and random effects, respectively. For non continuous longitudinal outcomes, joint models utilize Generalized linear mixed models (Rizopoulos, 2012).

To model the effect of PSA measurements on risk of GR, joint models utilize a relative risk submodel where the hazard of GR at any time point t, denoted by  $h_i(t)$ , depends on the history of true and unobserved values of PSA levels  $\mathcal{M}_i(t) = \{m_i(v), 0 \le v \le t\}$  measured up to that time point.

Joint models offer flexibility in modeling this dependence. In its simplest form, the hazard may depend on instantaneous value of PSA  $m_i(t)$  at time t. More sophisticated ones are dependence of hazard at time t on PSA-DT, PSA velocity  $m_i'(t) = \frac{dm_i(t)}{dt}$ , or even on the cumulative effect of PSA  $\int_0^t m_i(s) ds$  up to t. The fact that any functional form of dependence is possible, is evident from the following expression for hazard at time t:

$$h_i(t \mid M_i(t), \boldsymbol{w}_i) = h_0(t) e^{\boldsymbol{\gamma}^T \boldsymbol{w}_i + f\{M_i(t), \boldsymbol{b}_i, \boldsymbol{\alpha}\}}$$

where  $h_0(t)$  is the baseline hazard at time t.  $\boldsymbol{w}_i$  is a vector of baseline covariates and  $\boldsymbol{\gamma}$  are the corresponding parameters. The function  $f(\cdot)$  parametrized by vector  $\boldsymbol{\alpha}$  specifies the function form of longitudinal outcome that is used in the linear predictor of the relative risk model.

While  $\alpha$  controls the strength of association between the hazard of GR and features of the PSA history, the fact that both Gleason scores and PSA levels are internally related to a patient's health, is manifested by the random effects  $b_i$  in the model. The joint model postulates that given the random effects, time to GR and the different PSA measurements taken over time are all mutually independent. As mentioned earlier, in PRIAS study PSA-DT is used to decide the schedule of biopsies. Although PSA-DT is computed using observed PSA values, dependence on observed longitudinal history  $\mathcal{Y}(t) = \{y_i(v), 0 \leq v \leq t\}$  at any time t, is not the same as dependence on patient's health. On the contrary dependence on patient's health, manifested by  $b_i$  is same as dependence on future unobserved values of PSA. Thus the inference for the model parameters  $\theta = \{\beta^T, \gamma^T, \alpha^T, \sigma^2, \{d_{jk} \mid j=k=1,...q\}\}^T$  doesn't change even if uncertainty in biopsy schedule  $C_i$  is not modeled. The kernel of the corresponding joint likelihood conditional on the random effects and the model parameters is given by:

$$p(T_i, l_i, r_i, \boldsymbol{y}_i \mid \boldsymbol{b}_i, \boldsymbol{\theta}) \propto p(T_i \mid l_i, r_i, \boldsymbol{b}_i, \boldsymbol{\theta}) p(\boldsymbol{y}_i \mid \boldsymbol{b}_i, \boldsymbol{\theta})$$

## 2.2 Personalized scheduling approaches

Once a joint model for GR and PSA levels is obtained, the next step is to use it to create personalized schedules for biopsies. In this section we present the various personalized biopsy scheduling approaches and their motivation. The personalized schedules that we propose are dynamic in nature and thus at any given time, only 1 future biopsy is scheduled. The age of the patient and entire PSA, repeat biopsy history up to that time point is considered while computing the time of next biopsy. To elucidate the scheduling methods, let us assume that the a personalized schedule is to be created a new patient enumerated j, who is not present in the original sample of patients  $\mathcal{D}_n$ . Further let us assume that this patient did not have a GR at their last biopsy performed at time t, and that the PSA measurements are available up to a time point s. Combining these two pieces of information, the predictive distribution  $g(T_j^*)$  for time to GR for this patient is given by (conditioning on baseline covariates  $\mathbf{w}_i$  is dropped for notational simplicity here onwards):

$$g(T_{j}^{*}) = p(T_{j}^{*} \mid T_{j}^{*} > t, \mathcal{Y}_{j}(s), \mathcal{D}_{n})$$

$$= \int p(T_{j}^{*} \mid T_{j}^{*} > t, \mathcal{Y}_{j}(s), \boldsymbol{\theta}) p(\boldsymbol{\theta} \mid \mathcal{D}_{n}) d\boldsymbol{\theta}$$

$$= \int \int p(T_{j}^{*} \mid T_{j}^{*} > t, \mathcal{Y}_{j}(s), \boldsymbol{b}_{j}, \boldsymbol{\theta}) p(\boldsymbol{b}_{j} \mid T_{j}^{*} > t, \mathcal{Y}_{j}(s), \boldsymbol{\theta}) p(\boldsymbol{\theta} \mid \mathcal{D}_{n}) d\boldsymbol{b}_{j} d\boldsymbol{\theta}$$

$$(1)$$

where  $\mathcal{Y}_j(s)$  denotes the history of PSA measurements done up to time s. It can be seen that the predictive distribution depends on the observed longitudinal history via the random effects  $b_j$ . The posterior distribution of the parameters  $\theta$ , denoted by  $p(\theta \mid \mathcal{D}_n)$  is obtained from the joint model fitted to the original data set of patients  $\mathcal{D}_n$ .

Given the predictive distribution  $g(T_j^*)$ , our goal is find the optimal time  $u \ge \max(t, s)$  of the next biopsy. To this end, we use principles from statistical decision theory in a Bayesian setting (Berger, 1985; Robert, 2007). More specifically, we propose to choose future biopsy time u by minimizing the posterior expected loss  $E_g[L(T_j^*, u)]$ , where the expectation is taken w.r.t. the predictive distribution  $g(T_j^*)$ .

$$E_g[L(T_j^*, u)] = \int_t^{\infty} L(T_j^*, u) p(T_j^* \mid T_j^* > t, \mathcal{Y}_j(s), \mathcal{D}_n) dT_j^*$$

Various loss functions  $L(T_j^*, u)$  have been proposed in literature (Robert, 2007). The ones we utilize, and the corresponding motivations are presented next.

#### 2.2.1 Expected time of GR

One of the reasons, patients did not comply with the existing PRIAS schedule was 'complications on a previous biopsy'. Therefore, it makes sense to have as less biopsies as possible. In the ideal case only 1 biopsy, performed at the exact time of GR is sufficient. Hence, neither a time which overshoots the true GR time  $T_j^*$ , nor a time which undershoots is preferred. In this regard, the squared loss function  $L(T_j^*,u)=(T_j^*-u)^2$  and absolute loss function  $L(T_j^*,u)=|T_j^*-u|$  have the properties that the posterior expected loss is symmetric on both sides of  $T_j^*$ . Secondly, both loss functions have well known solutions available. We first discuss the posterior expected loss for the squared loss function, given by:

$$E_g[L(T_j^*, u)] = E_g[(T_j^* - u)^2]$$

$$= E_g[(T_j^*)^2] + u^2 - 2uE_g[T_j^*]$$
(2)

The posterior expected loss in equation 2 attains its minimum at  $u = E_g[T_j^*]$ , also known as expected time of GR.

#### Estimation

Since there is no closed form solution available for  $E_g[T_j^*]$ , for its estimation we introduce a construct called dynamic survival probability (Rizopoulos, 2011). The dynamic survival probability  $\pi_j(v \mid t, s)$  of patient j is the survival probability at time v, conditional on the observed PSA history  $\mathcal{Y}_j(s)$  and the fact that the patient did not have GR up to t. It is given by:

$$\pi_i(v \mid t, s) = Pr(T_i^* \ge v \mid T_i^* > t, \mathcal{Y}_i(s), D_n), v \ge t \tag{3}$$

The relationship between expected time of GR and dynamic survival probability is given by:

$$E_g[T_j^*] = t + \int_t^\infty \pi_j(v \mid t, s) \, dv$$

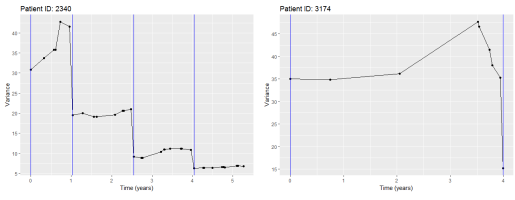
Since the R package JMbayes already provides an implementation of  $\pi_j(v \mid t, s)$ , we preferred this approach over Monte Carlo methods to estimate  $E_g[T_j^*]$  from the predictive distribution  $g(T_j^*)$ . There is no closed form solution available for the integral and hence we approximate it using Gauss-Kronrod quadrature. A limitation of 'Expected time of GR though', is that it is practically useful only when the variance of predictive distribution  $g(T_j^*)$  is small. The variance is given by:

$$Var_{g}[T_{j}^{*}] = E_{g}[T_{j}^{*2}] - E_{g}[T_{j}^{*}]^{2}$$

$$= 2 \int_{t}^{\infty} (v - t)\pi_{j}(v \mid t, s) dv - \left( \int_{t}^{\infty} \pi_{j}(v \mid t, s) dv \right)^{2}$$
(4)

Once again, a closed form solution is not available for the variance expression. The variance however depends on both t and  $\mathcal{Y}_j(s)$ . To elucidate the relationship between variance and observed information, we computed the variance at different time points for a couple of patients from the PRIAS study. The corresponding joint model fitted to the data set is discussed in section 3.1. The first patient in question is patient nr. 2340. He visited the hospital 34 times and was then right censored. 4 out those 34 visits were for biopsies. The variance of predictive distribution after each of those visits is shown in Figure 1. It is clear that the variance drops by a large margin each time a repeat biopsy is conducted. i.e. when more information about  $T_j^*$  is available. The effect of  $\mathcal{Y}_j(s)$  on variance can be seen in Figure 1b which corresponds to patient nr. 3174, who visited the hospital 9 times and was censored after his last biopsy. The observed PSA measurements for this patient are shown in Figure 2. It can be seen that the variance doesn't depend much on number of PSA measurements, but rather on the PSA values. For patient nr. 3174, variance drops quickly

when PSA measurements increase sharply. This becomes intuitive when we look at the model parameter estimates (section 3.1.1), where PSA velocity  $m'_i(t)$  is a strong predictor of the hazard of GR.



(a) Variance of the predictive distribution  $g(T_j^*)$  over a period of 9 years for patient 2340.  $g(T_j^*)$  over a period of 4 years for patient 3174. Blue vertical lines indicate biopsies.

Figure 1: Variance of the predictive distribution  $g(T_i^*)$ 

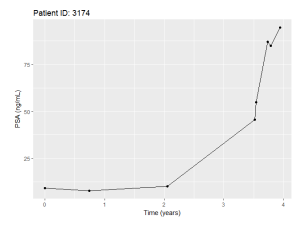


Figure 2: Observed evolution of PSA for patient 3174.

#### 2.2.2 Dynamic risk of GR

In a practical scenario it is possible that a doctor or a patient may not want to exceed a certain risk of GR  $1-\pi_j(u\mid t,s)$  since the last biopsy. This may be also useful in the cases where variance of  $g(T_j^*)$  is high, rendering expected time of GR, or any other measure of central tendency of  $g(T_j^*)$  unsuitable. The personalized scheduling approach based on dynamic risk of GR, schedules the next biopsy at a time point u such that the dynamic risk of GR is higher than a certain threshold  $1-\kappa$ ,  $\kappa\epsilon[0,1]$  beyond u. Or in other words the dynamic survival probability  $\pi_j(u\mid t,s)$  is below a threshold  $\kappa$  beyond u. To this end, the posterior expected loss for the following multilinear loss function can be minimized to find the most optimal u:

$$L_{k_1,k_2}(T_j^*, u) = \begin{cases} k_2(T_j^* - u) & if(T_j^* > u) \\ k_1(u - T_j^*) & \text{otherwise} \end{cases}$$
 (5)

where  $k_1 > 0$ ,  $k_2 > 0$  are constants parameterizing the loss function. The posterior expected loss function  $E_g[L_{k_1,k_2}(T_j^*,u)]$  obtains its minimum at  $u=\pi_j^{-1}\Big\{\frac{k_1}{k_1+k_2}\Big\}$  (Robert, 2007). The choice of  $k_1,k_2$  is equivalent to the choice of  $\kappa$ . More specifically,  $\kappa=\frac{k_1}{k_1+k_2}$ . When  $k_1=k_2=1$ , the

multilinear loss function is equal to the absolute loss function  $L(T_j^*, u) = |T_j^* - u|$ . Correspondingly,  $\kappa = 0.5$  for absolute loss and therefore time  $u = \pi_j^{-1}(0.5)$  is the median of  $g(T_j^*)$ .

#### Choice of $\kappa$

Since the value of  $\kappa$  dictates the biopsy schedule, its choice has important consequences. In certain cases it may be chosen on the basis of doctor's advice or the amount of risk that is acceptable to the patient. For e.g. if maximum acceptable risk is 75% then  $\kappa = 0.25$ , and correspondingly all  $k_1, k_2 \mid k_1 = \frac{k_2}{3}$  can be used in equation 5 to calculate u.

While a doctor's advice can be invaluable, it is also possible to automate the choice of  $\kappa$ . We propose to choose a  $\kappa$  for which a binary classification accuracy measure (López-Ratón et al., 2014; Sokolova and Lapalme, 2009), discriminating between cases and controls, is maximized. In PRIAS, cases are patients who experience GR and the rest are controls. However, a patient can be in control group at some time  $t_a$  and in the cases at some future time point  $t_b > t_a$ , and thus time dependent binary classification is more relevant. In joint models, a patient j is predicted to be a case if  $\pi_j(t + \Delta t \mid t, s) \leq \kappa$  and a control if  $\pi_j(t + \Delta t \mid t, s) > \kappa$  (Rizopoulos, 2014). The time window  $\Delta t$  can be either chosen on a clinical basis (such as 1 year in PRIAS WHY 1 YEAR) or it can be chosen at a point where  $AUC(t, \Delta t, s)$  (Rizopoulos, 2014) is largest. i.e.  $\Delta t$  for which the model has the most discriminative capability at time t. The binary classification accuracy measures we maximize to select the threshold  $\kappa$  are the following (the binary classification measures are functions of  $t, \Delta t, s$ , however the notation is dropped for readability):

- Accuracy:  $ACC = \frac{TP + TN}{TP + FP + TN + FN}$ , where TP, FP, TN and FN are the number of true positives, false positives, true negatives and false negatives at time point t. In this case if  $k_1 = TP + TN$  and  $k_2 = FP + FN$ , then  $\underset{k_1, k_2}{\operatorname{arg max}} ACC$  gives the optimal  $k_1, k_2$  or equivalently the  $\kappa$ .
- Youden's index: J = Sensitivity + Specificity 1, where sensitivity is defined as  $Pr(\pi_j(t + \Delta t \mid t, s) \leq \kappa \mid T_j^* \epsilon(t, t + \Delta t])$  and specificity is defined as  $Pr(\pi_j(t + \Delta t \mid t, s) > \kappa \mid T_j^* > t + \Delta t)$  (Rizopoulos, 2014). In this case if  $k_1 = FP \cdot TP FN \cdot TN$  and  $k_2 = (TP + FN)(FP + TN) k_1$ , then  $\underset{k_1, k_2}{\text{arg max }} J$  gives the optimal  $k_1, k_2$  or equivalently the  $\kappa$ .
- F1 Score:  $F1 = \frac{2TP}{2TP + FP + FN}$ . In this case if  $k_1 = 2TP$  and  $k_2 = FP + FN$ , then arg max F1 gives the optimal  $k_1, k_2$  or equivalently the  $\kappa$ .

### 2.2.3 A mixed approach between $E_g[T_i^*]$ and dynamic risk of GR

When the variance  $Var_g[T_j^*]$  is small, then  $E_g[T_j^*]$  is practically very useful. However when the variance is large, there may not be a clear central tendency of the distribution and  $E_g[T_j^*]$  will either overshoot  $T_j^*$  or undershoot it by a big margin. In the latter case more biopsies will be required until GR is detected at some time point  $T_j^o > T_j^*$ . The overshooting margin can be measured as an offset  $O_j = T_j^o - T_j^*$ . The maximum acceptable  $O_j$  in PRIAS is 3 years, which corresponds to the time gap between biopsies of the PRIAS fixed schedule. When  $Var_g[T_j^*]$  is large, the proposals based on  $E_g[T_j^*]$  can have a large  $O_j$ . Thus we propose that if the difference between the 0.025 quantile and  $E_g[T_j^*]$  is more than 3 years then proposals based on dynamic risk of GR be used instead. We call this approach a mixed approach.

#### 2.2.4 Scheduling multiple biopsies

Scheduling biopsies using personalized schedules is an iterative process. Only one biopsy is scheduled at once, using all the information available up to that point in time. The information is manifested via the predictive distribution  $g(T_j^*)$ . It is important to note that new biopsy times are only proposed when the patient visits the hospital for a PSA measurement or for biopsy, since  $g(T_j^*)$  is updated only at these time points. If a biopsy is conducted at time t, then scheduling the

next biopsy at time  $u \ge \max(t, s)$  for patient j is the goal. Although the predictive distribution  $g(T_j^*)$  is updated with the new information  $T_j^* > t$ , there are a couple of restrictions on time u, namely:

- 1. Two consecutive biopsies should have a gap of at least 1 year between them. i.e.  $u \ge t + 1$ . Given the medical side effects of biopsies, the 1 year gap is strongly advised for patients enrolled in PRIAS. However, the personalized scheduling methods do not take this rule into account.
- 2. Although it is required that  $u \ge \max(t, s)$ , the personalized scheduling methods may propose to perform a biopsy at a time  $u\epsilon(t, s]$ . The reason is that, GR is not a terminating event, and so the patients continue to visit for PSA measurements. The support of the predictive distribution  $g(T_j^*)$ , however does not depend on last time of PSA measurement s, and remains  $(t, \infty)$ .

Lastly, it is extremely likely that on consecutive visits to the hospital for PSA measurements, the personalized biopsy time is postponed or preponed. The choice of time of biopsy from consecutive visits is not clear. To resolve these issues, we propose to supplement the personalized scheduling methods with the following algorithm.

### Algorithm

Let us assume that patient j had their latest biopsy at time  $T_j^l$ , and the latest available PSA measurement is from the current visit to the hospital, at time  $T_j^{cv}$ . The predictive distribution for time of GR is given by  $p(T_j^*|T_j^* > T_j^l, \mathcal{Y}_j(T_j^{cv}), \mathcal{D}_n)$ . Further let,

- 1.  $T_i^p$  denote the time at which a biopsy is proposed by personalized scheduling methods.
- 2. Let  $T_j^s$  denote the time at which a biopsy is scheduled. This is not necessarily equal to the proposed time  $T_j^p$ , but rather a time devoid of the problems mentioned earlier. This time is generated from the algorithm.
- 3.  $T_i^{nv}$  denote the time of the next visit for PSA.
- 4.  $N_j^b$  denote the counter for number of biopsies conducted up to time  $T_j^l$ , excluding the biopsy done at the time of induction in AS.

Using these constructs, we have proposed an algorithm to schedule multiple biopsies. It is shown in Figure 3.

# 3 Personalized schedules for patients in PRIAS

To demonstrate how the personalized scheduling algorithm described in section 2.2.4 works, we apply it to the patients in the PRIAS dataset. To this end, we divide the PRIAS data set into training (5264 patients) and demonstration data sets (3 patients). We fit a joint model to the training data set and then use it to create a personalized schedule for patients in demonstration data set. We fit the joint model using the R package JMbayes (Rizopoulos, 2014), which uses the Bayesian methodology to estimate the model parameters.

#### 3.1 Fitting the joint model to PRIAS dataset

The training data set contains information about 5938 prostate cancer patients who satisfied the conditions for enrollment in AS. For every patient the age at the time of induction in AS was recorded. PSA was measured every 3 months for first 2 years and every 6 months thereafter. To detect GR, biopsies were conducted at different time points on the basis of a predetermined schedule as well as PSA-DT as described in section 1. For the longitudinal analysis of PSA measurements we used  $\log_2 PSA$  measurements instead of the raw data. The log transformation was done because the PSA scores took very large values around the time of disease progression. This indicated that the underlying distribution for PSA scores was right skewed. The longitudinal sub-model of the joint model we fit is given by:

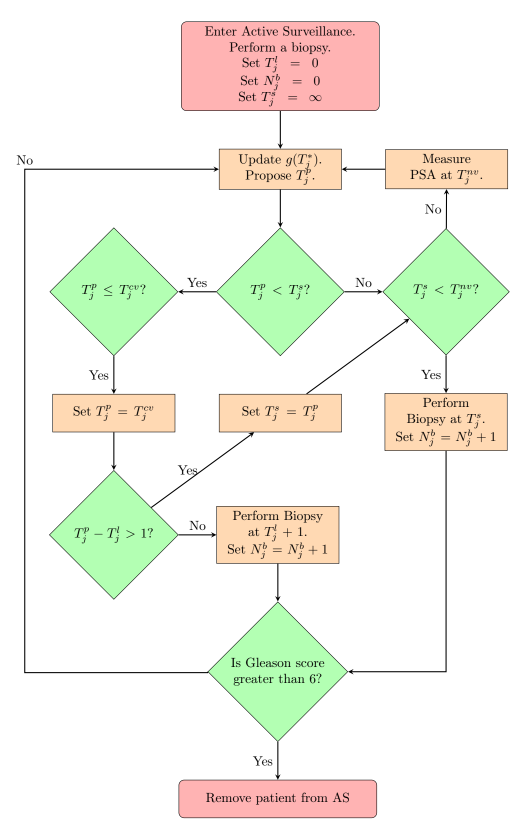


Figure 3: Algorithm for scheduling multiple biopsies using personalized schedules.

$$\log_{2} PSA(t) = m_{i}(t) + \varepsilon_{i}(t),$$

$$m_{i}(t) = (\beta_{0} + b_{i0}) + \beta_{1}(Age - 70) + \beta_{2}(Age - 70)^{2}$$

$$+ \sum_{k=1}^{4} \beta_{k+2}B_{k}(t, \mathcal{K}) + b_{i1}B_{7}(t, 0.5) + b_{i2}B_{8}(t, 0.5)$$

$$\varepsilon_{i}(t) \sim N(0, \sigma^{2}),$$

$$b_{i} \sim N(0, \mathbf{D})$$

The evolution of PSA levels over time is modeled flexibly using B-splines. For the fixed effects part the spline consists of 3 internal knots. The internal knots are at  $\mathcal{K} = \{0.5, 1.2, 2.5\}$  years, and boundary knots are at 0 and 7 years. For the random effects part there is only 1 internal knot at 0.5 years and the boundary knots are at 0 and 7 years. The choice of knots was based on exploratory analysis as well as on the basis of model selection criteria AIC and BIC. The variable Age was median centered to avoid numerical instabilities while estimating the parameters in the model. For the survival sub-model the hazard function we fitted is given by:

$$h_i(t) = h_0(t)e^{\gamma_1(Age-70) + \gamma_2(Age-70)^2\alpha_1 m_i(t) + \alpha_2 m_i'(t)}$$

where,  $\alpha_1$  and  $\alpha_2$  are measures of strength of association between hazard of GR and PSA value  $m_i(t)$  and PSA velocity  $m'_i(t)$ , respectively.  $h_0(t)$  is the baseline hazard at time t, and is modeled flexibly using P-splines(Eilers and Marx, 1996). Lastly, to fit the joint model we use the R package JMbayes Rizopoulos, 2014, which uses a Bayesian approach for parameter estimation.

#### 3.1.1 Parameter Estimates

The parameter estimates for the joint model we fitted to the PRIAS data set are shown in Table 1 and Table 2. Since the longitudinal evolution of  $\log_2 PSA$  is modeled with non-linear terms, the interpretation of the coefficients corresponding to time is not straightforward. In lieu of the interpretation we present the fitted evolution of PSA over a period of 10 years for a patient who is 70 years old in Figure 4. It can be seen that the after the first 6 months the PSA levels steadily increase over the follow up period. Since the model for PSA has only additive terms, this evolution remains same for all patients. The effect of Age only affects the baseline PSA score. However it is so small that it can be ignored for all practical purposes.

	Mean	Std. Dev	2.5%	97.5%	P
Intercept	2.455	0.012	2.433	2.480	< 0.000
(Age - 70)	0.003	0.001	$4.9 \times 10^{-4}$	0.006	0.032
$(Age - 70) \times (Age - 70)$	-0.001	$1.4 \times 10^{-4}$	-0.001	$-3.5 \times 10^{-4}$	< 0.000
Spline: visitTimeYears[0, 0.5]	-0.006	0.012	-0.031	0.017	0.674
Spline: visitTimeYears[0.5, 1.2]	0.228	0.019	0.192	0.265	< 0.000
Spline: visitTimeYears[1.2, 2.5]	0.140	0.029	0.088	0.197	< 0.000
Spline: visitTimeYears[2.5, 7]	0.303	0.039	0.227	0.379	< 0.000
$\sigma$	0.324	0.001	0.321	0.326	

Table 1: Longitudinal sub-model estimates for joint model.

For the survival sub-model, the parameter estimates in Table 2 show that only  $\log_2 PSA$  velocity is strongly associated with hazard of GR. For any patient, a unit increase in  $\log_2 PSA$  velocity corresponds to a 11 time increase in hazard of GR. The effect of  $\log_2 PSA$  value and effect of Age on hazard of GR are so small that they can be safely ignored for all practical purposes.

### 3.2 Demonstration of personalized schedules

In this section, we demonstrate how the personalized scheduling algorithm adapts the time of performing a biopsy according to the PSA history and results from repeat biopsies. The 3 patients

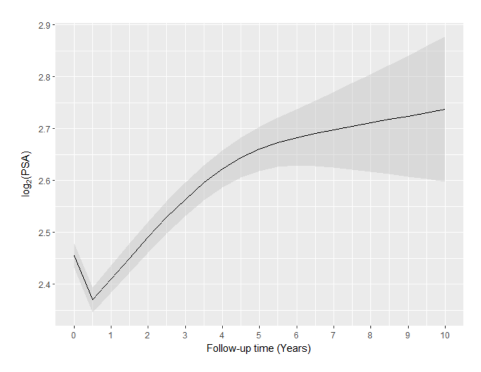


Figure 4: Fitted evolution of  $\log_2 PSA$  over a period of 10 years, for a patient who was inducted in AS at the Age of 70 years.

Variable	Mean	Std. Dev	2.5%	97.5%	P
Age - 70	0.037	0.006	0.025	0.0490	< 0.000
$(Age - 70) \times (Age - 70)$	-0.001	0.001	-0.003	$1.8 \times 10^{-4}$	0.104
$\log_2 PSA$	-0.049	0.064	-0.172	0.078	0.414
Slope: $\log_2 PSA$	2.407	0.319	1.791	3.069	< 0.000

Table 2: Survival sub-model estimates for joint model.

we have chosen for the demonstration data set are part of PRIAS program and they have had their repeat biopsies already. Hence a full scale comparison between PRIAS biopsy schedule and personalized scheduling algorithm's biopsy schedule is not possible.

The first patient of interest is patient nr. 3174 who was inducted in the PRIAS program at the age of 74 years. Between the last follow up and time of induction there was no repeat biopsy performed for this patient. Hence the predictive distribution  $g(T_j^*)$  for this patient depends only on the PSA measurements. For this patient, the evolution of PSA, time of last biopsy and proposed times of biopsies are shown in Figure 5. It can be seen that the PSA remains stable for the first 2 years of follow ups, but increases rapidly after that for the next 2 years of follow up. Since the hazard of GR depends on PSA velocity (Table2), the schedule of biopsy based on personalized scheduling algorithm (Section 3), adjust the times of biopsy according to the steep rise in PSA profile. At 2 years the expected and median time of GR are 12.5 years and 15.2 years respectively, whereas at 4 years, they are 5.3 and 4 years respectively. It is important to note that the estimates are year 2 are not as useful as the estimates at year 4, because the  $Var_g[T_j^*]$  is considerably lower at year 4, as shown in Figure 1b.

The second patient of interest is patient nr. 911. Figure 6 shows the evolution of PSA, time of last biopsy and proposed biopsy times for this patient. Firstly, it can be seen that at between year 1.5 and year 2, the PSA rises sharply, and accordingly the personalized schedules based on expected time of GR preponed the proposed biopsy time from 14.2 years to 13.8 years. The change in median time of GR is trivial. The second observation is that between year 2 and year 3 the PSA decreases sharply and accordingly, the proposed biopsy times are postponed. More specifically the median time of GR increased from 15.2 years at year 2 to 17 years at year 3. Whereas the expected

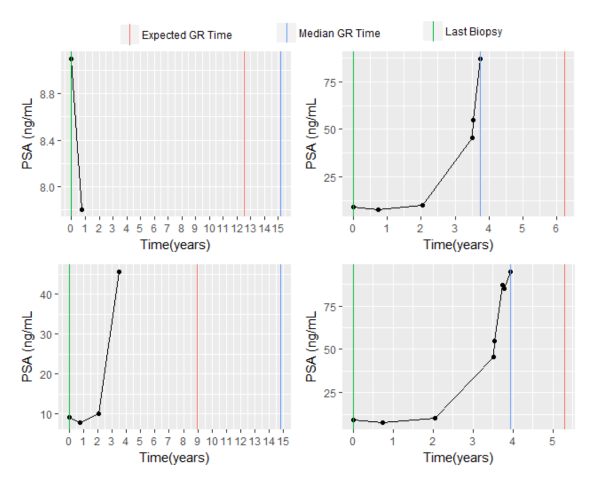


Figure 5: Proposed biopsy times for patient 3174 from PRIAS.

time of GR increased from 13.8 years to 16.6 years. It can also be seen that PSA remains stable up to to year 4. Lastly, because no GR is found at the repeat biopsy performed at 4.1 years, this further leads to postponing of the biopsy times, which become 18.7 and 19.9 for expected time of GR and median time of GR respectively.

The fact that PSA velocity affects the biopsy times is also evident in the case of patient 2340, whose PSA evolution is shown in Figure 7. We have purposefully ignored all the biopsies conducted for this patient in PRIAS, to isolate the effect of PSA history. It can be seen that PSA slowly rises to a value of 14 ng/mL over a period of 2.5 years. Correspondingly, there is little change in proposed biopsy times. On the other hand between 2.5 years and 3.2 years the PSA rises very sharply and correspondingly the expected time of GR decreases by an year from 12.2 years to 11.2 years in that period. The median time of GR remains the same though.

# 4 Simulation study

For contrasting the personalized scheduling approaches against the schedule of PRIAS and the schedule of doing biopsy every year, we performed a simulation study. We used the parameters from the joint model fitted to the original PRIAS data set to generate 60 data sets with 1000 patients each. We employed a Weibull baseline hazard while generating Gleason reclassification times for the simulated patients. We divided the 60 data sets into groups of 10 each, and within each group the parameter values for the Weibull baseline hazard were kept the same, while across the group they were different. The reason to create such groups was to ensure the consistency of results within in each group. Using different parameter values for the baseline hazard across the groups ensured that we tested not only the scenarios in which patients have early progression times but also those in which patients have late progression times. This is important because how late/early patients are observed to have Gleason reclassification, partially depends on the induction

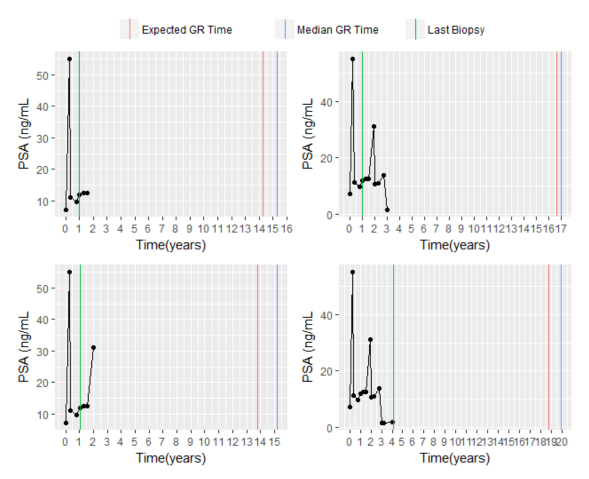


Figure 6: Proposed biopsy times for patient 911 from PRIAS.

criteria of the AS program.

#### 4.1 Simulation setup

We divided each of the 60 data sets into training (750 patients) and test (250 patients) parts. We fitted a joint model to the training data set and using it we generated biopsy schedules for patients in the test data set.

#### Evaluating performance of personalized biopsy schedules

Since we generate the true Gleason reclassification time for each of the patients in the test data set, we are able to calculate the offset for each patient and each scheduling method. While creating the personalized schedules we adhere to the PSA measurement schedule of PRIAS.

#### 4.2 Results

Since the Gleason reclassification times varied widely across the simulation data sets, we discuss the results separately for the different scenarios. In total we have 6 scenarios corresponding to the 6 groups of 10 data sets. For brevity, we only chose 3 most dissimilar scenarios and further we present one set of results for each of the 3 scenarios. The complete set of results can be found at <a href="https://goo.gl/u6dg8G">https://goo.gl/u6dg8G</a>. At this moment, we will not compare our results against the PRIAS schedule, since the PRIAS schedule we used is fixed for all patients, whereas in reality it can be dynamic if the PSA doubling time is less than 10 years.

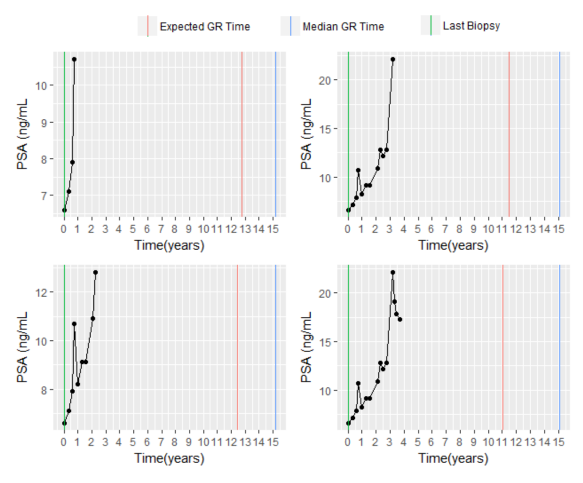


Figure 7: Proposed biopsy times for patient 2340 from PRIAS.

#### 4.2.1 Scenario 1

In the first scenario the Gleason reclassification times for the patients are mostly between 7 and 11 years. The Gleason reclassification times of test patients from one of the data sets is shown in Figure 8. Figure 11 shows that, for this data set the approaches with the least number of biopsies, 2 in number, are conditional expected failure time, and dynamic risk with a threshold  $\kappa$  chosen such that accuracy (section 2.2.2) is maximized. The mean and median offset for the two approaches is around 9 months. However for a few outlying patients the offset is more than 24 months. If we compare this to the schedule of performing biopsy every year the mean offset is around 6 months and mean number of biopsies is 8.

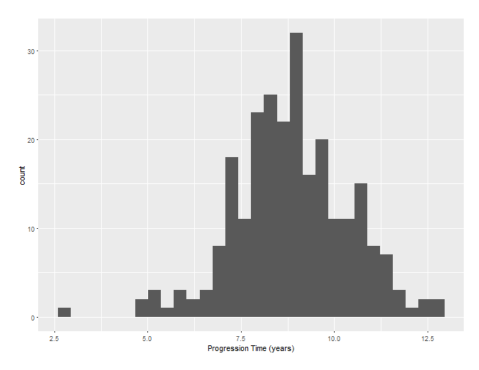


Figure 8: Gleason reclassification times for patients from the test data set of scenario 1.

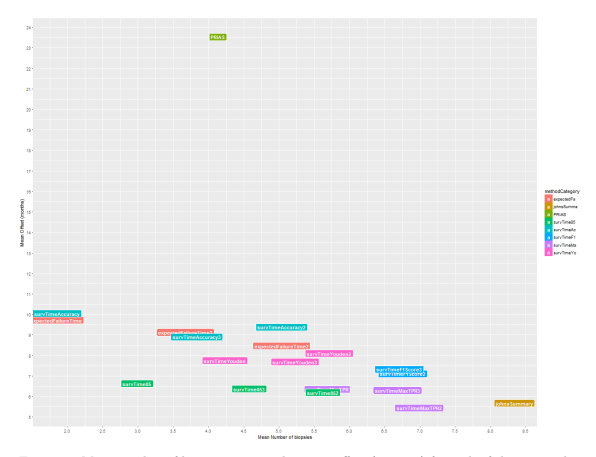


Figure 9: Mean number of biopsies against the mean offset (in years) for each of the approaches in scenario 1.

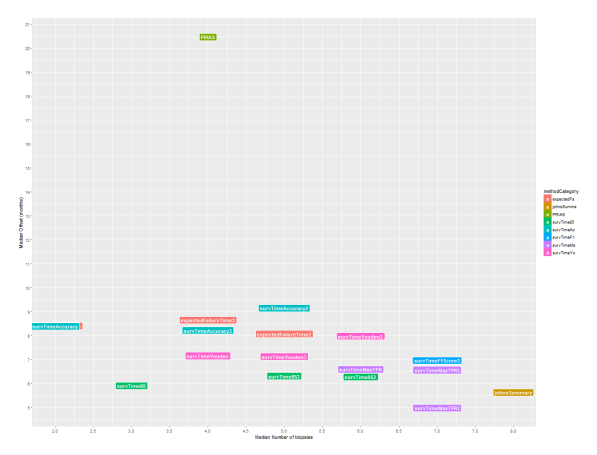


Figure 10: Median number of biopsies against the median offset (in years) for each of the approaches in scenario 1.

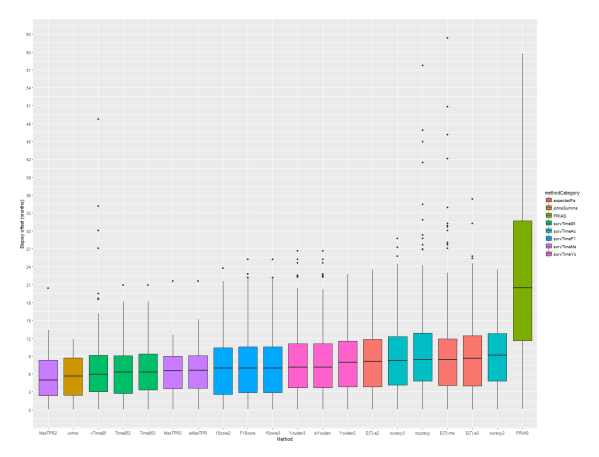


Figure 11: Boxplot for the offset corresponding to the various approaches in scenario 1.

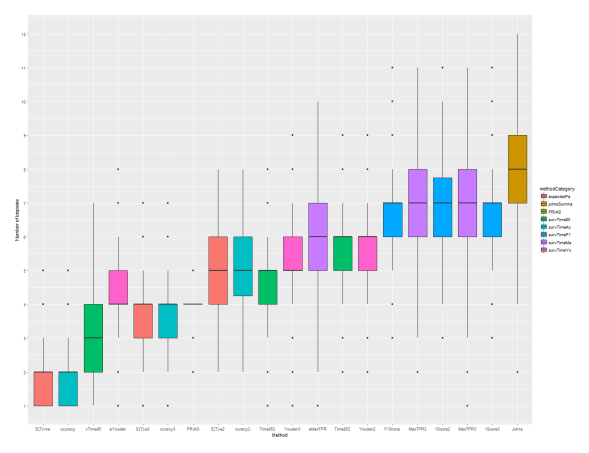


Figure 12: Boxplot for the number of biopsies corresponding to the various approaches in scenario 1.

#### 4.2.2 Scenario 2

Unlike the previous scenario, the Gleason reclassification times for the patients in Scenario 2 are mostly between 0 and 5 years. The Gleason reclassification times of test patients from one of the data sets is shown in Figure 13. Figure 16 shows that, for this data set the approach with the least number of biopsies is conditional expected failure time. The mean and median number of biopsies are 1.5 and 1 respectively. While the mean and median offset is around 18 months, there is considerable variation in the offset for the various patients. The first and third quartiles are at 12 months and 27 months respectively. The large variation in offset is due to the fact that time to Gleason reclassification has larger variance since it is based on a small history of the patient. In such a scenario one might be inclined towards doing a biopsy every year as they have the least offset. However that approach leads to a very large number of biopsies as shown in Figure 17. Certain approaches based on dynamic risk perform better here, both in terms of number of biopsies as well as the offset. In particular if the choice of  $\kappa$  can be based on maximization of the Youden index or a fixed  $\kappa$  can be chosen.

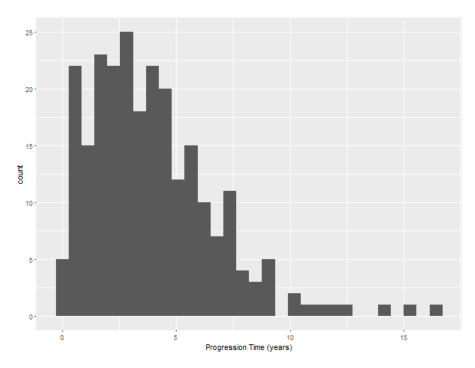


Figure 13: Gleason reclassification times for patients from the test data set of scenario 2.

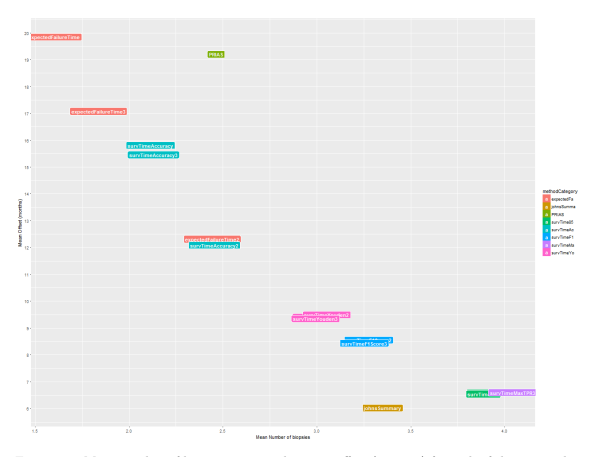


Figure 14: Mean number of biopsies against the mean offset (in years) for each of the approaches in scenario 2.

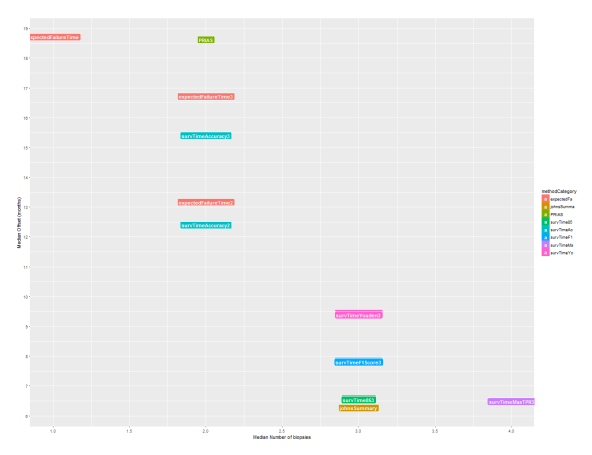


Figure 15: Median number of biopsies against the median offset (in years) for each of the approaches in scenario 2.

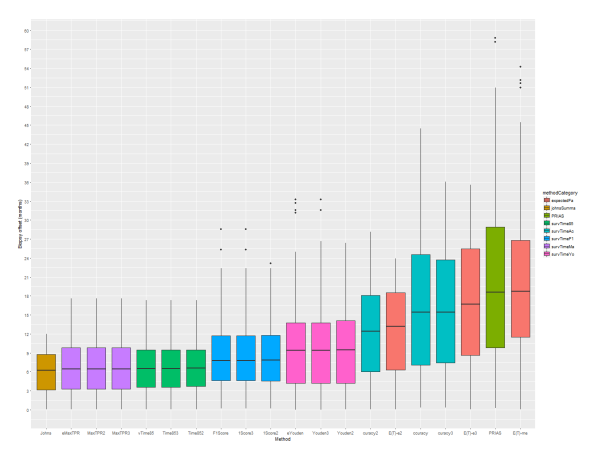


Figure 16: Boxplot for the offset corresponding to the various approaches in scenario 2.

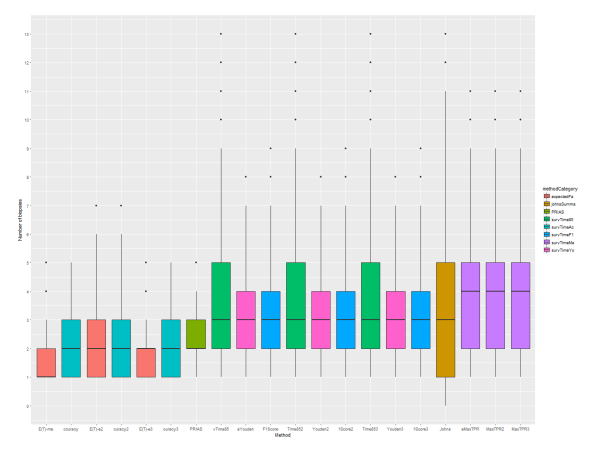


Figure 17: Boxplot for the number of biopsies corresponding to the various approaches in scenario 2.

#### 4.2.3 Scenario 3

In scenario 3, the Gleason reclassification times for the patients are widely spread, mostly between 2 and 9 years. i.e. patients fail late as well early. The Gleason reclassification times of test patients from one of the data sets is shown in Figure 18. Figure 21 shows that, for this data set the approach with the least number of biopsies is conditional expected failure time. The mean and median number of biopsies are 1.5 and 1 respectively. This method also has considerable variation in the offset for the various patients. The first and third quartiles are at 9 months and 27 months respectively. A more detailed analysis revealed that the variation in offset was higher for subjects with earlier failure times. For e.g. the mean and median offset in subjects with reclassification times more than 4 years was nearly 13.5 months. For others it was nearly 39.5 months. This is once again is due to the fact that time to Gleason reclassification has larger variance since it is based on a small history of the patient. And thus usefulness of conditional expected failure time is questionable.

Dynamic risk based methods once again provide an alternative. If a fixed  $\kappa$  of 0.15 is used then although the offset will be low, there is very high variation in number of biopsies. i.e. people who have reclassifications early will benefit but people who have reclassifications later will have too many biopsies. If  $\kappa$  is chosen such that accuracy is maximized, and a biopsy is done whenever there is a gap of 2 years then a reasonable offset is obtained. The first and third quartiles are almost 4.5 and 18 months. The first and third quartiles for number of biopsies are 3 and 4.

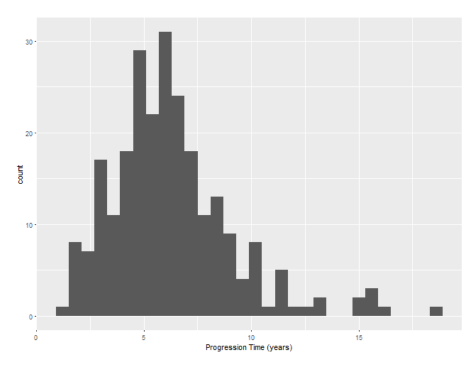


Figure 18: Gleason reclassification times for patients from the test data set of scenario 3.

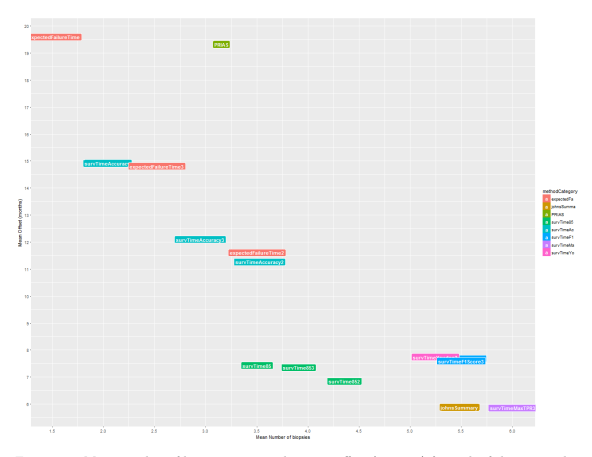


Figure 19: Mean number of biopsies against the mean offset (in years) for each of the approaches in scenario 3.

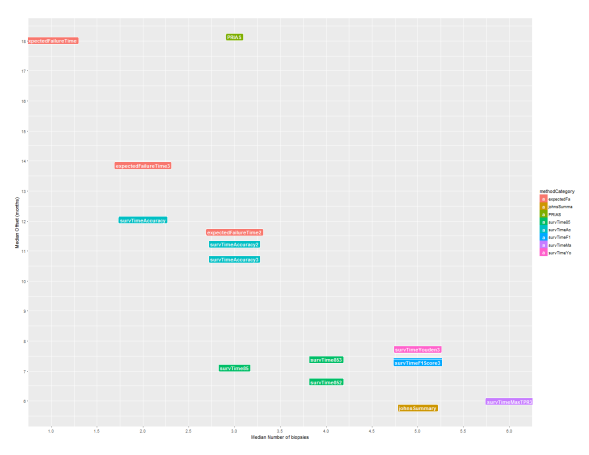


Figure 20: Median number of biopsies against the median offset (in years) for each of the approaches in scenario 3.

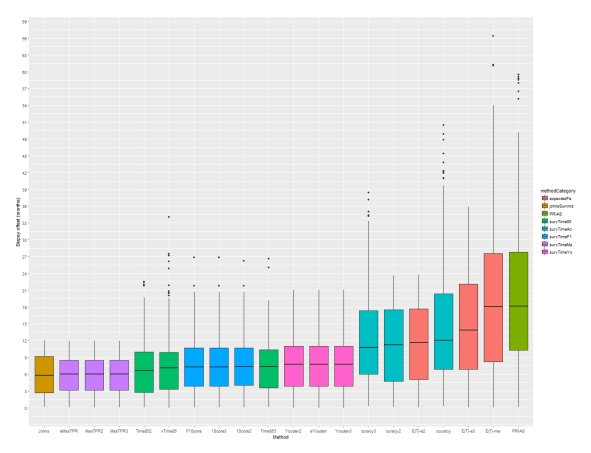


Figure 21: Boxplot for the offset corresponding to the various approaches in scenario 3.

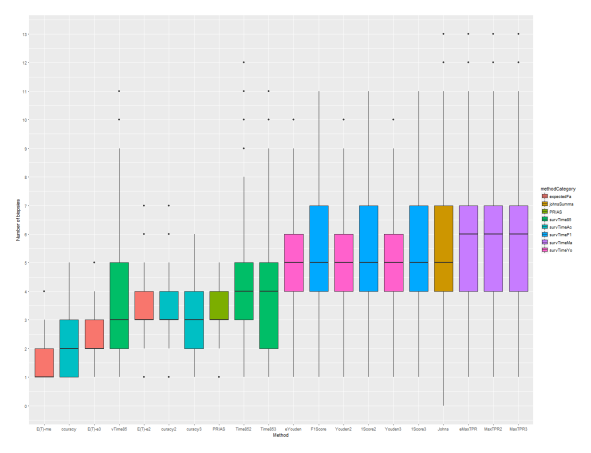


Figure 22: Boxplot for the number of biopsies corresponding to the various approaches in scenario 3.

## 4.3 A mixed approach: E(T) vs dynamic risk

While we did create biopsy schedules using conditional expected Gleason reclassification time, and dynamic risk of Gleason reclassification based approaches, we saw that no single approach works best in all scenarios. In particular, we observed that when the failure times are early dynamic risk based approaches perform better in terms of offset than conditional expected Gleason reclassification time. The latter works best when the Gleason reclassification times are later. This has motivated us to use a mixed approach, where we schedule biopsies using dynamic risk based approaches if the variance (Eq. ??) of time to Gleason reclassification is higher than a certain threshold, otherwise we use conditional expected Gleason reclassification time. This is still work in progress.

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# A Appendix Heading

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