#### Personalized Schedules for Burdensome Surveillance Tests

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Summary: Commonly used gold standard surveillance tests (biopsies, endoscopies, etc.) for confirming disease progression in early-stage chronic non-communicable disease patients (cancer, cardiovascular, etc.) are invasive. For detecting progression timely (benefit), patients are exposed to numerous invasive tests repeatedly (burden) over their lifetime as per a fixed one-size-fits-all schedule. Motivated by this problem in the world's largest prostate cancer surveillance study PRIAS, we present disease progression risk-based personalized test schedules, that aim to better balance the number of invasive tests (burden), and time delay in detection of progression (less is beneficial) than fixed schedules.

Using joint models for time-to-event and longitudinal data, we first consolidate auxiliary longitudinal data (e.g., biomarkers) and results of previous invasive tests, into individualized future cumulative-risk of progression. Under this risk profile, we optimize a utility function of the number of tests and expected time delay in detection of progression to obtain a personalized schedule of invasive tests. Personalized schedules are updated as more patient data becomes available over follow-up. We assist patients/doctors in comparing the consequences of opting for personalized versus fixed schedules objectively. For this, we exploit a patient's cumulative-risk profile to estimate the expected time delay in detection of progression for following both personalized and fixed schedules, in a patient-specific manner. We implement our methodology in a web-application for real prostate cancer patients of the PRIAS study.

KEY WORDS: Chronic diseases; Invasive medical tests; Joint models; Personalized schedules; Prostate biopsy; Surveillance

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## 1. Introduction

Chronic non-communicable diseases (e.g., cancer, renal, cardiovascular diseases, etc.) are the primary cause of human deaths worldwide (Alwan et al., 2010). In many patients diagnosed with an early stage disease, periodical surveillance tests are recommended to detect disease progression, a non-terminal event. Often the most accurate or gold standard surveillance tests are also invasive. For example, to confirm progression, biopsies are conducted repeatedly in low-grade prostate cancer (Bokhorst et al., 2015), endoscopies in Barrett's esophagus (Streitz et al., 1993), and colonoscopies in colorectal cancer (Krist et al., 2007) patients. Repeat biopsies are also utilized to detect allograft deterioration in lung (McWilliams et al., 2008) and kidney transplant (Henderson et al., 2011) patients.

Usually, invasive tests are scheduled in a fixed manner, e.g., every six months. The frequency of tests in these fixed schedules varies between diseases (Henderson et al., 2011; Bokhorst et al., 2015; Krist et al., 2007) and cohorts. Although, due to the periodical nature of test schedules, progression is always detected with a time delay (Figure 1). This time delay can be reduced by scheduling invasive tests frequently. However, invasive tests are difficult to conduct, can lead to severe complications (Loeb et al., 2013; Krist et al., 2007), cause patient discomfort, and sometimes patients may not comply with frequent tests (Bokhorst et al., 2015). In this regard, fixed test schedules ignore the differences in speed of progression between patients, and impose an equal medical burden on all. Hence, the frequency of invasive tests holds important implications for patients.

# [Figure 1 about here.]

In this paper, we aim to balance the number of invasive tests (burden) and the time delay in detecting disease progression (less is beneficial) better than fixed schedules. For this purpose, we intend to create personalized test schedules that exploit patient-specific clinical data accumulated during follow-up. In surveillance, this data includes baseline characteristics

of patients; results from previous invasive tests; and auxiliary longitudinal outcomes such as biomarkers, physical examination, and medical imaging measurements, etc. Previous approaches for personalized schedules can be divided into three categories. First, heuristic methods such as decision making flowcharts, e.g., Bokhorst et al. (2015). However, flowcharts discretize continuous clinical outcomes, often utilize only the last measurement, and ignore the measurement error in observed outcomes. The second method is employing partially observable Markov decision processes (Alagoz et al., 2010; Steimle and Denton, 2017) for personalized test decisions. Although, the curse of dimensionality limits their application with continuous longitudinal outcomes. Third, personalized schedules obtained by optimizing an explicit utility function of the clinical parameters of interest (Bebu and Lachin, 2017; Rizopoulos et al., 2015), including our previous work on scheduling biopsies in prostate cancer (Tomer et al., 2019). In this work, we will employ the third approach.

Our methodology is as follows. First, we develop a full specification of the joint distribution of disease and patient-specific longitudinal outcomes and the time of progression. We achieve this using joint models for time-to-event and longitudinal data (Tsiatis and Davidian, 2004; Rizopoulos, 2012). We use joint models because they are inherently personalized. Specifically, they exploit patient-specific random effects (Laird and Ware, 1982) to model longitudinal outcomes without discretizing them. We subsequently employ the fitted joint model for new patients to estimate their patient-specific cumulative-risk of progression over their current and future follow-up visits. These risk predictions utilize their clinical data accumulated until their latest follow-up. We schedule invasive tests on all those future time points where a patient's conditional cumulative-risk of progression is equal to a certain threshold (e.g., 10% risk). We then automate the choice of this threshold and the resulting schedule. More specifically, we optimize a function of the expected number of tests and the expected time delay in detecting progression (Figure 1) corresponding to different risk threshold based

personalized schedules. We estimate the expected number of tests and time delay in a patientspecific manner for both fixed and personalized schedules. This can help patients/doctors to evaluate the consequences of opting for personalized versus fixed schedules objectively.

This research is motivated by the problem of scheduling biopsies (Nieboer et al., 2018) in the world's largest prostate cancer active surveillance study PRIAS (Bokhorst et al., 2015). It has 7813 patients, 104,904 longitudinal measurements, and 1134 patients with cancer progression. These patients have low/very-low grade prostate cancer, often over-diagnosed due to prostate-specific antigen (PSA) based screening tests (Crawford, 2003). The goal of surveillance upon diagnosis is to delay serious treatments (e.g., surgery, chemotherapy, etc.) until cancer progresses further. For this purpose, patients are monitored continually via PSA (ng/mL) blood tests, digital rectal examination (DRE) for shape and size of the tumor, and biopsy Gleason grade group (Epstein et al., 2016). Since biopsy results are the strongest indicator of cancer-related outcomes, treatment is commonly advised upon observing an increase in a patient's biopsy Gleason grade group (cancer progression). The most commonly used schedule of biopsies in prostate cancer active surveillance is annual biopsies (Loeb et al., 2014). However, they lead to many unnecessary biopsies in slow/nonprogressing patients (50% proportion in some cohorts). Biopsy burden combined with patient non-compliance to frequent biopsies (Bokhorst et al., 2015) has raised concerns regarding the optimal biopsy schedule. Since prostate cancer has the second-highest incidence among all cancers in males (Torre et al., 2015), biopsy schedules tailored for individual patients can reduce the overall burden of biopsies in a large number of patients worldwide.

The rest of the paper is as follows. Section 2 briefly introduces the joint modeling framework. In Section 3, we present the methodology for personalized schedules and then demonstrate them for biopsies in real PRIAS patients in Section 4. Lastly, in Section 5, we show the efficacy of personalized schedules via a realistic simulation study based on PRIAS patients.

### 2. Joint Model for Time-to-Progression and Longitudinal Outcomes

Let the true time of disease progression for the *i*-th patient be  $T_i^*$ . Progression is always observed with interval censoring  $l_i < T_i^* \le r_i$  (Figure 1). In patients who obtain progression,  $r_i$  and  $l_i$  denote the time of their latest and second latest invasive tests. Otherwise,  $l_i$  denotes the time of their latest test and  $r_i = \infty$ . Assuming K auxiliary longitudinal outcomes, let  $\mathbf{y}_{ki}$  denote the  $n_{ki} \times 1$  longitudinal response vector of the k-th outcome,  $k \in \{1, \ldots, K\}$ . The observed data of all n patients is given by  $\mathcal{A}_n = \{l_i, r_i, \mathbf{y}_{1i}, \ldots, \mathbf{y}_{Ki}; i = 1, \ldots, n\}$ .

To accommodate longitudinal outcomes of different types in a unified framework, the joint model consists of a generalized linear mixed-effects sub-model (McCulloch and Neuhaus, 2005). In particular, the conditional distribution of  $y_{ki}$  given a vector of patient-specific random effects  $b_{ki}$  is assumed to be a member of the exponential family, with linear predictor given by,

$$g_k \big[ E\{y_{ki}(t) \mid \boldsymbol{b}_{ki}\} \big] = m_{ki}(t) = \boldsymbol{x}_{ki}^{\top}(t)\boldsymbol{\beta}_k + \boldsymbol{z}_{ki}^{\top}(t)\boldsymbol{b}_{ki},$$

where  $g_k(\cdot)$  denotes a known one-to-one monotonic link function,  $y_{ki}(t)$  denotes the value of the k-th longitudinal outcome for the i-th patient at time t, and  $\boldsymbol{x}_{ki}(t)$  and  $\boldsymbol{z}_{ki}(t)$  denote the time-dependent design vectors for the fixed  $\boldsymbol{\beta}_k$  and random effects  $\boldsymbol{b}_{ki}$ , respectively. To account for the association between the different longitudinal outcomes, we link their corresponding random effects. More specifically, the complete vector of random effects  $\boldsymbol{b}_i = (\boldsymbol{b}_{1i}^\top, \dots, \boldsymbol{b}_{Ki}^\top)^\top$  is assumed to follow a multivariate normal distribution with mean zero and variance-covariance matrix W.

For the survival process, we assume that the hazard of progression  $h_i(t)$  at a time t depends on a function of the patient and outcome-specific linear predictors  $m_{ki}(t)$  and/or the random effects. More specifically,

$$h_i \left\{ t \mid \mathcal{M}_i(t), \boldsymbol{w}_i(t) \right\} = h_0(t) \exp \left[ \boldsymbol{\gamma}^\top \boldsymbol{w}_i(t) + \sum_{k=1}^K f_k \left\{ \mathcal{M}_{ki}(t), \boldsymbol{w}_i(t), \boldsymbol{b}_{ki}, \boldsymbol{\alpha}_k \right\} \right], \quad t > 0,$$

where  $h_0(\cdot)$  denotes the baseline hazard function,  $\mathcal{M}_{ki}(t) = \{m_{ki}(s) \mid 0 \leq s < t\}$  denotes the

history of the k-th longitudinal process up to t, and  $\mathbf{w}_i(t)$  is a vector of exogenous, possibly time-varying, covariates with corresponding regression coefficients  $\gamma$ . Functions  $f_k(\cdot)$ , parameterized by vector of coefficients  $\alpha_k$ , specify which features of each longitudinal outcome are included in the linear predictor of the relative-risk model (Brown, 2009; Rizopoulos, 2012; Taylor et al., 2013). Some examples, motivated by the literature (subscripts k dropped for brevity), are:

$$\begin{cases} f\{\mathcal{M}_i(t), \boldsymbol{w}_i(t), \boldsymbol{b}_i, \boldsymbol{\alpha}\} = \alpha m_i(t), \\ f\{\mathcal{M}_i(t), \boldsymbol{w}_i(t), \boldsymbol{b}_i, \boldsymbol{\alpha}\} = \alpha_1 m_i(t) + \alpha_2 m_i'(t), & \text{with } m_i'(t) = \frac{\mathrm{d}m_i(t)}{\mathrm{d}t}. \end{cases}$$

These formulations of  $f(\cdot)$  postulate that the hazard of progression at time t may be associated with the underlying level  $m_i(t)$  of the longitudinal outcome at t, or with both the level and velocity  $m'_i(t)$  (e.g., PSA value and velocity in prostate cancer) of the outcome at t. Lastly,  $h_0(t)$  is the baseline hazard at time t, and is modeled flexibly using P-splines (Eilers and Marx, 1996). The detailed specification of the baseline hazard  $h_0(t)$ , and the joint parameter estimation of the longitudinal and relative-risk sub-models using the Bayesian approach are presented in Web-Appendix A.

#### 3. Personalized Schedule of Invasive Tests for Detecting Progression

We intend to develop a personalized schedule of invasive tests for a new patient j, not present in training dataset  $\mathcal{A}_n$ . Tests are conducted only until progression is detected (Figure 1). Let  $T_j^*$  be the true time of progression, and  $t < T_j^*$  be the time of the last conducted test on which progression was not detected for the j-th patient. Lastly, v > t denotes the time of the current follow-up visit.

#### 3.1 Cumulative-risk of progression

First we combine the history of observed longitudinal outcomes  $\{\mathcal{Y}_{1j}(v), \dots, \mathcal{Y}_{Kj}(v)\}$  until the current visit time v, and the result of the last conducted test  $T_j^* > t$  to define the patient-

specific cumulative-risk of progression at future time u (Figure 2).

$$R_{j}(u \mid t, v) = \Pr\{T_{j}^{*} \leq u \mid T_{j}^{*} > t, \mathcal{Y}_{1j}(v), \dots, \mathcal{Y}_{Kj}(v), \mathcal{A}_{n}\}$$

$$= \int \int \Pr(T_{j}^{*} \leq u \mid T_{j}^{*} > t, \boldsymbol{b}_{j}, \boldsymbol{\theta}) p\{\boldsymbol{b}_{j} \mid T_{j}^{*} > t, \mathcal{Y}_{1j}(v), \dots, \mathcal{Y}_{Kj}(v), \boldsymbol{\theta}\}$$

$$\times p(\boldsymbol{\theta} \mid \mathcal{A}_{n}) d\boldsymbol{b}_{j} d\boldsymbol{\theta}, \quad u \geq t.$$

$$(1)$$

The personalized cumulative-risk function  $R_j(\cdot)$  depends on the observed longitudinal data  $\{\mathcal{Y}_{1j}(v), \ldots, \mathcal{Y}_{Kj}(v)\}$ , and the training dataset  $\mathcal{A}_n$  via the posterior distribution of patient-specific random effects  $\boldsymbol{b}_j$ , and posterior distribution of the vector of joint model parameters  $\boldsymbol{\theta}$ , respectively. The risk also dynamically updates as more longitudinal data becomes available over follow-up (Panel B and C, Figure 2).

# [Figure 2 about here.]

# 3.2 Personalized Schedule of Tests Using a Cumulative-risk Threshold $\kappa^*$

Our aim is to employ the cumulative-risk function in (1) to develop a risk-based personalized schedule of invasive tests for the j-th patient. Typically an invasive test is decided on the same visit on which auxiliary data (e.g., biomarkers) is measured. Let  $U = u_1, \ldots, u_L$  represent a schedule of such visits (e.g., every six months in prostate cancer for PSA measurement), where  $u_1 = v$  is also the time of the current visit.

First, we make L successive decisions for conducting tests on each of the L future visit times  $u_l \in U$ . Specifically, we decide to conduct a test at time  $u_l$  if the conditional cumulative-risk of progression at  $u_l$  is larger than a certain risk threshold  $0 \le \kappa^* \le 1$  (e.g.,  $\kappa^* = 10\%$  risk). If a test gets planned at time  $u_l$ , then the successive test decision at time  $u_{l+1}$  is made using an updated cumulative-risk profile (Figure 3). This updated cumulative-risk profile accounts for the possibility that progression may occur after time  $u_l < T_j^*$ . The test decisions on each

future visit time  $u_l$  are defined as:

$$Q_j^{\kappa^*}(u_l \mid t_l, v) = I\{R_j(u_l \mid t_l, v) \ge \kappa^*\},$$

$$t_l = \begin{cases} t, & \text{if} & l = 1\\ t_{l-1}, & \text{if} & Q_j^{\kappa^*}(u_{l-1} \mid t_{l-1}, v) = 0, l \ge 2\\ u_{l-1}, & \text{if} & Q_j^{\kappa^*}(u_{l-1} \mid t_{l-1}, v) = 1, l \ge 2 \end{cases}.$$

The cumulative-risk  $R_j(u_l \mid t_l, v)$  at future visit time  $u_l$  utilizes the time  $t_l$  as the time of the last conducted test on which progression may not be observed (Equation 1). However, the contribution of the observed longitudinal data  $\{\mathcal{Y}_{1j}(v), \ldots, \mathcal{Y}_{Kj}(v)\}$  in the risk function remains the same over all time points in U. The test decision at time  $u_l$  is denoted by  $Q_j^{\kappa^*}(u_l \mid t_l, v)$ . Via the indicator function  $I(\cdot)$  it obtains a value 1 (or 0) when a test is to be conducted (or not conducted) at time  $u_l$ . The subset of future time points in U on which a test is to be performed results into a personalized schedule of planned future tests, given by:

$$S_i^{\kappa^*}(U \mid t, v) = \{ u_l \in U \mid Q_i^{\kappa^*}(u_l \mid t_l, v) = 1 \}.$$
 (2)

The personalized schedule in (2) is updated as more patient data becomes available over subsequent follow-up visits.

# 3.3 Expected Number of Tests and Time Delay in Detecting Progression

The schedule  $S_j^{\kappa^*}(U \mid t, v)$  manifests a personalized test plan for the j-the patient. However, the number of tests that will get conducted and the time delay in detecting progression that may subsequently be observed (Figure 1), depends on the true time of progression  $T_j^*$  of the patient. Since two different patients with the same timing of tests will expect different number of tests and time delay, we estimate these two quantities in a patient-specific manner as well. Although, this calculation is not limited to personalized schedules only, but can be done for any schedule S of tests with N time points  $S = \{s_n \mid n = 1, ..., N\}$ .

For each of the N planned tests there exist N time intervals  $s_{n-1} < T_j^* \leqslant s_n$  in which

progression may be observed. Correspondingly, there are N possible number of tests  $1, \ldots, N$ , and N possible time delays in detecting progression  $s_n - T_j^*$ . Given a schedule S, the true number of tests  $N_j$  and true time delay in detecting progression  $D_j$  that the patient will experience can be defined as:

$$N_{j}(S \mid t) = \begin{cases} 1, & \text{if} & t < T_{j}^{*} \leq s_{1} \\ \dots & \\ N - 1, & \text{if} & s_{N-2} < T_{j}^{*} \leq s_{N-1} \\ N, & \text{if} & s_{N-1} < T_{j}^{*} \end{cases} ,$$

$$D_{j}(S \mid t) = \begin{cases} s_{1} - T_{j}^{*}, & \text{if} & t < T_{j}^{*} \leq s_{1} \\ \dots & \\ s_{N} - T_{j}^{*}, & \text{if} & s_{N-1} < T_{j}^{*} \leq s_{N} \end{cases} .$$

$$(3)$$

The time delay is not defined for the scenario in which the patient obtains progression after the time of the last test in the schedule  $T_j^* > s_N$ . To estimate the expected values of both  $N_j(\cdot)$  and  $D_j(\cdot)$  in a patient-specific manner, we exploit the personalized cumulative-risk profile of the patient (Equation 1). Specifically, the expected number of tests  $E\{N_j(\cdot)\}$  and expected time delay  $E\{D_j(\cdot)\}$  can be calculated as the weighted sum of N possible number of tests and time delays defined in (3). The n-th weight is equal to the probability of the patient obtaining progression in the n-th interval  $s_{n-1} < T_j^* \leq s_n$ .

$$E\{N_{j}(S \mid t)\} = \sum_{n=1}^{N-1} n \times \Pr(s_{n-1} < T_{j}^{*} \leqslant s_{n}) + N \times \Pr(T_{j}^{*} > s_{N-1}), \quad s_{0} = t$$

$$E\{D_{j}(S \mid t)\} = \sum_{n=1}^{N} \left\{s_{n} - E(T_{j}^{*} \mid s_{n-1}, s_{n}, v)\right\} \times \Pr(s_{n-1} < T_{j}^{*} \leqslant s_{n}), \quad s_{0} = t$$

$$E(T_{j}^{*} \mid s_{n-1}, s_{n}, v) = s_{n-1} + \int_{s_{n-1}}^{s_{n}} \Pr\left\{T_{j}^{*} \geqslant u \mid s_{n-1} < T_{j}^{*} \leqslant s_{n}, \mathcal{Y}_{1j}(v), \dots, \mathcal{Y}_{Kj}(v), \mathcal{A}_{n}\right\} du$$

$$\Pr(s_{n-1} < T_{j}^{*} \leqslant s_{n}) = R_{j}(s_{n} \mid t, v) - R_{j}(s_{n-1} \mid t, v)$$

$$\Pr(T_{j}^{*} > s_{N-1}) = 1 - R_{j}(s_{N-1} \mid t, v),$$

(4)

where  $E(T_j^* \mid s_{n-1}, s_n, v)$  denotes the conditional expected time of progression for the scenario  $s_{n-1} < T_j^* \leqslant s_n$ , and is calculated as the area under the corresponding survival curve.

The personalized expected number of tests and expected time delay in detecting progression have the advantage that they are updated over follow-up as more patient data becomes available. Since they can be calculated for any schedule, patients and doctors can utilize them to compare schedules before making a decision. Although, in order to have a fair comparison of time delays between different schedules for the same patient, a compulsory test at a common horizon time point should be planned in all schedules.

## 3.4 Choice of Risk Threshold $\kappa^*$

The risk threshold  $\kappa^*$  controls the timing and the total number of invasive tests in the personalized schedule  $S_j^{\kappa^*}(U \mid t, v)$ . Through the timing and the total number of planned tests,  $\kappa^*$  also indirectly affects the time delay (Figure 1) that may occur in detecting progression if this schedule is followed. Hence,  $\kappa^*$  should be chosen while balancing both the number of invasive tests (burden) and the time delay in detecting progression (less is beneficial).

Consider the bi-dimensional Euclidean space of the expected total number of invasive tests (x-axis) and the corresponding expected time delay in detecting progression (y-axis) for test schedules planned using various  $\kappa$  (Figure 4). An ideal schedule of tests will have only one test conducted exactly at the true time of progression  $T_j^*$  of a patient. In other words, it will lead to a zero time delay. This schedule is shown at the point of optimality (1, 0) in Figure 4. Subsequently, a risk threshold  $\kappa^*(v)$  specific to current visit time v, can be chosen by minimizing the Euclidean distance between the point (1, 0) and the set of points representing personalized schedules  $S_j^{\kappa}$  corresponding to various  $0 \leqslant \kappa \leqslant 1$ .

$$\kappa^*(v) = \arg\min_{\kappa} \sqrt{\left[ E\{N_j(S_j^{\kappa} \mid t)\} - 1\right]^2 + \left[ E\{D_j(S_j^{\kappa} \mid t)\} - 0\right]^2}.$$
 (5)

Additional consequences of following a particular schedule, such as (quality-adjusted) life-

years saved, can also be accommodated in (5). This can be achieved by first setting a point of optimality in a higher dimensional Euclidean space of the aforementioned consequences, and then minimizing the Euclidean distance to the point of optimality.

Certain patients may have preferences for the maximum number of invasive tests they will undergo. Others may be apprehensive about having an expected time delay higher than a certain number of months. In this regard, the Euclidean distance in (5) can be minimized under constraints on the expected number of tests and/or expected time delay (Figure 4). An additional benefit of this approach is that it alleviates the issue of time delay and the number of tests having different units of measurement (Cook and Wong, 1994).

[Figure 4 about here.]

# 4. Demonstration of Personalized Schedules

To demonstrate the application of personalized schedules on real patients, we return to the prostate cancer active surveillance dataset, PRIAS, described in Section 1. The current PRIAS protocol for biopsies is fixed biopsies at year one, four, seven, and ten of follow-up, and every five years after that. Additional annual biopsies are scheduled if a patient's PSA doubling-time (Bokhorst et al., 2015) is high. The PSA is measured as per a fixed schedule, quarterly for the first two years, and semi-annually after that. The DRE is also measured semi-annually. The dataset is summarized in Web-Appendix B.

The clinical data that we intend to use consists of longitudinal PSA (continuous: ng/mL) and DRE (binary: tumor palpable or not) measurements, patient age at baseline, history of biopsies, and interval-censored times of cancer progression. The event of interest is cancer progression. We aim to use the accumulated clinical data to build a joint model that can be utilized for creating personalized biopsy schedules in future PRIAS patients.

### 4.1 Fitting the Joint Model to the PRIAS Dataset

We fit a joint model with  $\log_2(\mathrm{PSA}+1)$  transformed PSA (Lin et al., 2000; Pearson et al., 1994), and DRE as longitudinal outcomes, and cancer progression as the event (Web-Appendix B.3 for exact specification). For PSA, we utilize a linear mixed-effects sub-model wherein PSA profiles are modeled non-linearly over follow-up using B-splines (De Boor, 1978). For DRE, we utilize a logistic mixed-effects sub-model. To link the longitudinal sub-models for the PSA and DRE with the relative-risk sub-model for cancer progression, we include three features of the longitudinal outcomes in the relative-risk sub-model. Specifically, the hazard of cancer progression at time t depends on the fitted instantaneous  $\log_2(\mathrm{PSA}+1)$  value at time t, the estimated instantaneous  $\log_2(\mathrm{PSA}+1)$  velocity at t, and fitted log-odds of having a DRE indicating a palpable tumor at t. We estimated the parameters of our model under the Bayesian framework using the R package JMbayes (Rizopoulos, 2016).

The follow-up period of PRIAS is currently limited. Hence, our joint model is able to predict the cumulative-risk of progression only until the year ten of follow-up. The cumulative-risk of progression at year ten in PRIAS is 50% (Web-Figure 1). We found that the strongest predictor for progression in our model is  $\log_2(\text{PSA}+1)$  velocity. Specifically, for an increase in fitted  $\log_2(\text{PSA}+1)$  velocity from -0.03 to 0.15 the adjusted hazard ratio of progression was 1.6 (95%CI: 1.45–1.78). Detailed parameter estimates are in Web-Appendix B.4. Since personalized schedules are risk-based, their overall performance is dependent on the predictive accuracy of the fitted model. In this regard, the PRIAS based model's time-dependent area under the receiver operating characteristic curve (?) was moderate (between 0.61 and 0.68) and time-dependent mean absolute prediction error (?) was moderate to large (between 0.08 and 0.24) over follow-up (Web-Appendix B.6).

### 4.2 Personalized Schedules for a Demonstration Patient

We utilized the joint model fitted to the PRIAS dataset to schedule biopsies in a real PRIAS patient (Figure 5), starting from his current visit at year five, until year ten of follow-up. The cumulative-risk of progression of this patient at his current visit is 6%, whereas, at ten years, it is 16.5%. Thus, the patient is predicted to progress slowly. Consequently, risk-based personalized schedules in Panel B of Figure 5 planned much fewer biopsies than the standard annual schedule. At the same time, the expected time delay in detection of progression in personalized schedules is also less than one year (maximum delay possible with annual schedule). It is important to note that for a fair comparison of expected time delay, we scheduled a compulsory biopsy at the horizon of ten years in all schedules. In addition, we maintained a recommended minimum gap of one year between consecutive biopsies (Bokhorst et al., 2016).

# [Figure 5 about here.]

#### 5. Simulation Study

Although we demonstrated personalized schedules for a real patient, we also intend to analyze and compare personalized and fixed schedules in a full cohort. The criteria for comparison of schedules are the total number of invasive tests planned, and the actual time delay in detection of progression for each schedule. However, due to the periodical nature of schedules, the actual time delay in detection of progression cannot be observed in real-world surveillance. Hence, instead, we compare personalized versus fixed schedules via an extensive simulated randomized clinical trial in which each hypothetical patient undergoes each schedule. To keep our simulation study realistic, we employ the prostate cancer active surveillance scenario. More specifically, our simulated population is manifested by the fitted joint model obtained using the PRIAS cohort (Web-Appendix B.3).

#### 5.1 Simulation Setup

From the simulation population, we first sample 500 datasets, each representing a hypothetical prostate cancer active surveillance program with 1000 patients in it. We generate a true cancer progression time for each of the  $500 \times 1000$  patients and then sample a set of longitudinal DRE and PSA measurements at the same follow-up visit times as given in the PRIAS protocol. We then split each dataset into training (750 patients) and test (250 patients) parts, and generate a random and noninformative censoring time for the training patients. All test and training patients also observe Type-I censoring at year ten of follow-up (current study period of PRIAS). We next fit a joint model of the same specification as the model fitted to PRIAS (Web-Appendix B.3), to each of the 500 training datasets and obtain MCMC samples from the 500 sets of the posterior distribution of the parameters. In each of the 500 hypothetical surveillance programs, we utilize the corresponding fitted joint models to develop the profiles for cumulative-risk of progression in each of the  $500 \times 250$  test patients. This cumulative-risk is further used to create personalized biopsy schedules for the test patients. For each test patient, we conduct biopsies using personalized biopsy schedules based on two fixed risk thresholds, namely,  $\kappa^* = 5\%$  and  $\kappa^* = 10\%$ , and an automatically chosen visit-specific  $\kappa^*(v)$  (Equation 5) with a constraint of 6 months on expected time delay in detection of progression. We also conduct biopsies according to the currently practiced PRIAS and annual schedules. We decide biopsies only on the standard PSA follow-up visits (Section 4) utilizing clinical data accumulated only until that visit. Also, we maintain a minimum recommended gap of one year between consecutive prostate biopsies (Bokhorst et al., 2015). Biopsies are conducted until progression is detected, or the maximum follow-up period at year ten (horizon) is reached. The actual time delay in detection of progression is equal to the time at which progression is detected minus the actual (simulated) time of progression of a patient.

#### 5.2 Results

Since the simulated cohorts are based on PRIAS, roughly only 50% of the patients progress in the ten year study period. While we are able to calculate the total number of biopsies scheduled in all  $500 \times 250$  test patients, but the time delay in detection of progression is available only for those patients who progress in ten years (progressing). Hence, we show the simulation results separately for progressing and non-progressing patients in Panel A, and Panel B of Figure 6, respectively.

For progressing patients (Panel A, Figure 6), the annual schedule leads to the maximum number of biopsies (Median 3, Inter-quartile range or IQR: 1–6). However, it also guarantees a maximum time delay of one year for all patients. The PRIAS protocol schedules much fewer biopsies (Median 1, IQR: 2–4), but also has a higher time delay (Median 0.74, IQR: 0.38–1.00 years). The personalized schedule based on automatically chosen risk threshold  $\kappa^*(v)$  schedules fewer biopsies than PRIAS and has a delay slightly higher than PRIAS (Median 0.96, IQR:0.54–1.74). Unless the patient progresses within the first year of prostate cancer active surveillance, a delay of up to three years may not increase the risk of adverse downstream outcomes in (??).

The patients who are at the most advantage with the personalized schedules are the non-progressing patients (Panel B, Figure 6). For all of these patients, the annual schedule leads to 10 (unnecessary) biopsies. The schedule of the PRIAS program schedules a median of 6 (IQR: 4–8) biopsies. In comparison, the schedule based on automatically chosen risk threshold  $\kappa^*(v)$  schedules a median of 4 (IQR: 3–4) biopsies.

#### 6. Discussion

In this paper, we presented a methodology to create personalized schedules for burdensome surveillance tests, to detect disease progression in early-stage chronic non-communicable diseases. To this end, we utilized the framework of joint models for time-to-event and longitudinal data. Our approach first combines a patient's auxiliary longitudinal data (e.g., biomarkers) and results from previous invasive tests to estimate the patient-specific cumulative-risk of disease progression over his current and future follow-up time period. Then, using this risk profile, we schedule future invasive tests whenever the patient's conditional cumulative-risk of progression is predicted to be above a certain threshold. We select this risk threshold automatically in a personalized manner, by optimizing a utility function of the patient-specific consequences of choosing a particular risk threshold based schedule. These consequences are, namely, the number of invasive tests for a particular schedule, and the expected time delay in detection of progression if that schedule is followed. Last, we calculate this expected time delay in a personalized manner for both personalized and fixed schedules to assist patients/doctors in making a more informed decision of choosing a test schedule.

The use of joint models gives our schedules certain advantages. First, joint models utilize individualized random-effects, making our schedules inherently personalized. Second, the patient-specific risk of progression employed by the proposed personalized schedules is estimated by utilizing all observed longitudinal and clinical data of a patient. In addition, the continuous longitudinal outcomes are not discretized, which is commonly a case in Markov Decision Process based (Alagoz et al., 2010; Steimle and Denton, 2017), and flowchart-based test schedules. The third and most significant advantage of our schedules, however, is that they update as more patient data becomes available over follow-up. Last, although this work concerns with the use of personalized schedules in disease surveillance, the methodology is generic for use under a screening setting as well.

Since our schedules are risk-based, we proposed a utility function to automate the choice of a risk threshold based schedule. The utility function that we proposed focused only on two aspects of a schedule, namely the burden and the benefit. In this regard, we chose the number of invasive tests in a schedule (burden) and expected time delay in detection of progression (less is beneficial) because they are easy to interpret and are critical in making the decision of an invasive test. Since we calculate the expected time delay in a patient-specific manner for both personalized and fixed schedules, patients/doctors can compare and choose various risk-based and fixed schedules according to their preferences for the expected burden-benefit ratio. Additional measures such as (quality-adjusted) life-years saved can also be easily added in our utility function.

We evaluated the efficacy of personalized schedules in a full cohort via a realistic simulation randomized clinical trial for prostate cancer active surveillance patients. We observed that the personalized schedule that used an automatically chosen risk threshold using (5) reduced unnecessary biopsies for patients who did not observe progression in the study period, compared to fixed schedules. In contrast, in patients who observed progression, the personalized schedule with automatically chosen risk threshold scheduled fewer biopsies at the cost of having a slightly more time delay in detecting progression than the PRIAS schedule. However, this by no means is the limit of the performance of the personalized schedules. In general, personalized schedules employing models with higher predictive accuracy (both discrimination and calibration) than the PRIAS based model may lead to an even better balance between the number of tests and the time delay in detecting progression.

There are certain limitations of our work. First, in practice, most cohorts observe Type-I right censoring. Hence, the cumulative-risk profiles of patients and the calculation of expected time delay in detection of progression is only possible up to the time of Type-I censoring. This problem can only be resolved as more follow-up data become available over

time. We proposed a joint model which assumes all events other than progression to be non-informative censoring. Alternative models that account for competing risks may lead to better results as they estimate absolute and not the cause-specific risk of progression. However, the methodology for scheduling biopsies need not change. Many surveillance tests are imperfect and prone to inter-observer variation (e.g., biopsy Gleason grade). Models that account for inter-observer variation in diagnostic tests (Balasubramanian and Lagakos, 2003) will be exciting to investigate further.

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#### SUPPORTING INFORMATION

Web Appendices referenced in this paper are available in the file titled 'supplementary.pdf'.

#### References

- Alagoz, O., Ayer, T., and Erenay, F. S. (2010). Operations research models for cancer screening. Wiley encyclopedia of operations research and management science.
- Alwan, A., MacLean, D. R., Riley, L. M., d'Espaignet, E. T., Mathers, C. D., Stevens, G. A., and Bettcher, D. (2010). Monitoring and surveillance of chronic non-communicable diseases: progress and capacity in high-burden countries. *The Lancet* **376**, 1861–1868.

- Balasubramanian, R. and Lagakos, S. W. (2003). Estimation of a failure time distribution based on imperfect diagnostic tests. *Biometrika* **90**, 171–182.
- Bebu, I. and Lachin, J. M. (2017). Optimal screening schedules for disease progression with application to diabetic retinopathy. *Biostatistics* **19**, 1–13.
- Bokhorst, L. P., Alberts, A. R., Rannikko, A., Valdagni, R., Pickles, T., Kakehi, Y., Bangma, C. H., Roobol, M. J., and PRIAS study group (2015). Compliance rates with the Prostate Cancer Research International Active Surveillance (PRIAS) protocol and disease reclassification in noncompliers. *European Urology* **68**, 814–821.
- Bokhorst, L. P., Valdagni, R., Rannikko, A., Kakehi, Y., Pickles, T., Bangma, C. H., Roobol, M. J., and PRIAS study group (2016). A decade of active surveillance in the PRIAS study: an update and evaluation of the criteria used to recommend a switch to active treatment. European Urology 70, 954–960.
- Brown, E. R. (2009). Assessing the association between trends in a biomarker and risk of event with an application in pediatric HIV/AIDS. The Annals of Applied Statistics 3, 1163–1182.
- Cook, R. D. and Wong, W. K. (1994). On the equivalence of constrained and compound optimal designs. *Journal of the American Statistical Association* **89**, 687–692.
- Crawford, E. D. (2003). Epidemiology of prostate cancer. *Urology* **62**, 3–12.
- De Boor, C. (1978). A practical guide to splines, volume 27. Springer-Verlag New York.
- Eilers, P. H. and Marx, B. D. (1996). Flexible smoothing with B-splines and penalties.

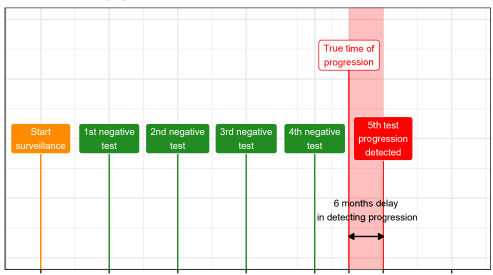
  Statistical Science 11, 89–121.
- Epstein, J. I., Egevad, L., Amin, M. B., Delahunt, B., Srigley, J. R., and Humphrey, P. A. (2016). The 2014 international society of urological pathology (isup) consensus conference on gleason grading of prostatic carcinoma. *The American journal of surgical pathology* **40**, 244–252.

- Henderson, L., Nankivell, B., and Chapman\*, J. (2011). Surveillance protocol kidney transplant biopsies: their evolving role in clinical practice. *American Journal of Transplantation* **11**, 1570–1575.
- Krist, A. H., Jones, R. M., Woolf, S. H., Woessner, S. E., Merenstein, D., Kerns, J. W., Foliaco, W., and Jackson, P. (2007). Timing of repeat colonoscopy: disparity between guidelines and endoscopists recommendation. *American journal of preventive medicine* 33, 471–478.
- Laird, N. M. and Ware, J. H. (1982). Random-effects models for longitudinal data. *Biometrics* pages 963–974.
- Lin, H., McCulloch, C. E., Turnbull, B. W., Slate, E. H., and Clark, L. C. (2000). A latent class mixed model for analysing biomarker trajectories with irregularly scheduled observations. *Statistics in Medicine* **19**, 1303–1318.
- Loeb, S., Carter, H. B., Schwartz, M., Fagerlin, A., Braithwaite, R. S., and Lepor, H. (2014). Heterogeneity in active surveillance protocols worldwide. *Reviews in urology* 16, 202–203.
- Loeb, S., Vellekoop, A., Ahmed, H. U., Catto, J., Emberton, M., Nam, R., Rosario, D. J., Scattoni, V., and Lotan, Y. (2013). Systematic review of complications of prostate biopsy. *European urology* **64**, 876–892.
- McCulloch, C. E. and Neuhaus, J. M. (2005). Generalized linear mixed models. *Encyclopedia* of biostatistics 4,.
- McWilliams, T. J., Williams, T. J., Whitford, H. M., and Snell, G. I. (2008). Surveillance bronchoscopy in lung transplant recipients: risk versus benefit. *The Journal of Heart and Lung Transplantation* 27, 1203–1209.
- Nieboer, D., Tomer, A., Rizopoulos, D., Roobol, M. J., and Steyerberg, E. W. (2018). Active surveillance: a review of risk-based, dynamic monitoring. *Translational andrology and urology* 7, 106–115.

- Pearson, J. D., Morrell, C. H., Landis, P. K., Carter, H. B., and Brant, L. J. (1994). Mixed-effects regression models for studying the natural history of prostate disease. *Statistics in Medicine* **13**, 587–601.
- Rizopoulos, D. (2012). Joint Models for Longitudinal and Time-to-Event Data: With Applications in R. CRC Press.
- Rizopoulos, D. (2016). The R package JMbayes for fitting joint models for longitudinal and time-to-event data using MCMC. *Journal of Statistical Software* **72**, 1–46.
- Rizopoulos, D., Taylor, J. M., Van Rosmalen, J., Steyerberg, E. W., and Takkenberg, J. J. (2015). Personalized screening intervals for biomarkers using joint models for longitudinal and survival data. *Biostatistics* 17, 149–164.
- Steimle, L. N. and Denton, B. T. (2017). Markov decision processes for screening and treatment of chronic diseases. In *Markov Decision Processes in Practice*, pages 189–222. Springer.
- Streitz, J. J., Andrews, J. C., and Ellis, J. F. (1993). Endoscopic surveillance of barrett's esophagus. does it help? *The Journal of thoracic and cardiovascular surgery* **105**, 383–7.
- Taylor, J. M., Park, Y., Ankerst, D. P., Proust-Lima, C., Williams, S., Kestin, L., Bae, K., Pickles, T., and Sandler, H. (2013). Real-time individual predictions of prostate cancer recurrence using joint models. *Biometrics* 69, 206–213.
- Tomer, A., Nieboer, D., Roobol, M. J., Steyerberg, E. W., and Rizopoulos, D. (2019). Personalized schedules for surveillance of low-risk prostate cancer patients. *Biometrics* **75**, 153–162.
- Torre, L. A., Bray, F., Siegel, R. L., Ferlay, J., Lortet-Tieulent, J., and Jemal, A. (2015). Global cancer statistics, 2012. *CA: A Cancer Journal for Clinicians* **65**, 87–108.
- Tsiatis, A. A. and Davidian, M. (2004). Joint modeling of longitudinal and time-to-event data: an overview. *Statistica Sinica* **14**, 809–834.

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# **B** Test every 2 years

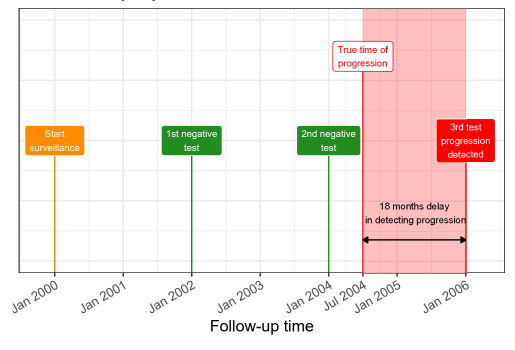


Figure 1. Trade-off between the number of invasive tests and time delay in detecting progression (non-terminal event of interest): The true time of progression for this patient July 2004. More frequent invasive tests in Panel A lead to a smaller time delay in detecting progression than less frequent invasive tests in Panel B. Since invasive tests are conducted periodically, the time of progression is observed as an interval. For example, between Jan 2004–Jan 2005 in Panel A and between Jan 2004–Jan 2006 in Panel B.

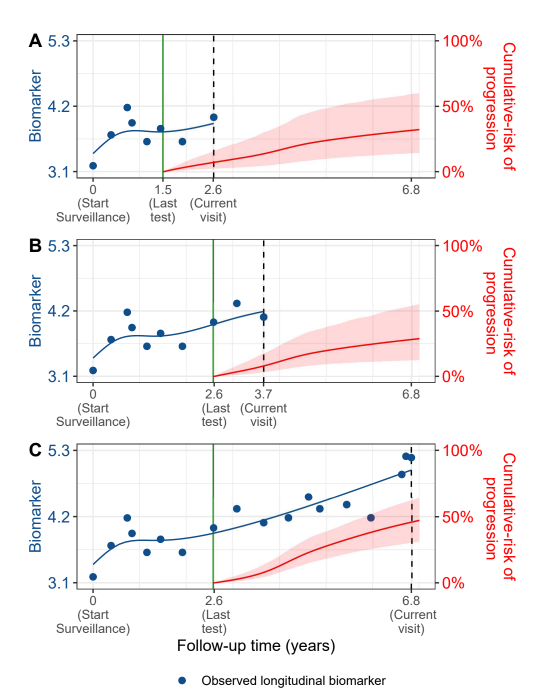


Figure 2. Cumulative-risk of progression changing dynamically over follow-up as more patient data is gathered. A single longitudinal outcome, namely, a continuous biomarker of disease progression, is used for illustration. Panels A, B and C: are ordered by the time of the current visit (dashed vertical black line) of a new patient. At each of these visits, we combine the accumulated longitudinal measurements (shown in blue), and last time of negative invasive test (solid vertical green line) to obtain the updated cumulative-risk profile (shown in red) of the patient. All values are illustrative.

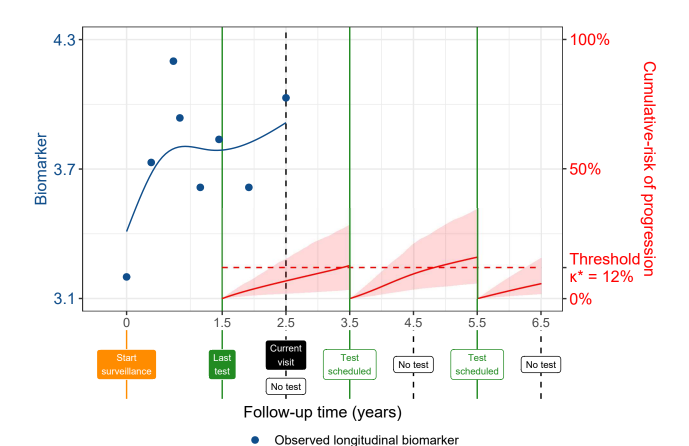


Figure 3. Personalized Invasive Test Schedule Using Patient-specific Conditional Cumulative-risk of Progression. A single longitudinal outcome, namely, a continuous biomarker (observed: blue dots, fitted: blue line) of disease progression is used for illustration. The last test on which progression was not observed was conducted at t=1.5 years. The current visit time of the patient is v=2.5 years. Decisions for invasive test need to be made at a gap of every one year starting from the current visit until a horizon of 6.5 years. That is,  $U=\{2.5,3.5,4.5,5.5,6.5\}$  years. Based on an example risk threshold of 12% ( $\kappa^*=0.12$ ) the future test decisions at time points in U lead to a personalized schedule  $S_j^{\kappa}(U \mid t=1.5, v=2.5)=\{3.5,5.5\}$  years. The conditional cumulative-risk profiles  $R_j(u_l \mid t_l, v)$  employed in (3.2) are shown with red line (confidence interval shaded). It is called 'conditional' because, for example, the second test at future time 5.5 years, is scheduled after accounting for the possibility that progression (true time  $T_j^*$ ) may not have occurred until the time of the previously scheduled test at time 3.5 <  $T_j^*$  years. All values are illustrative.

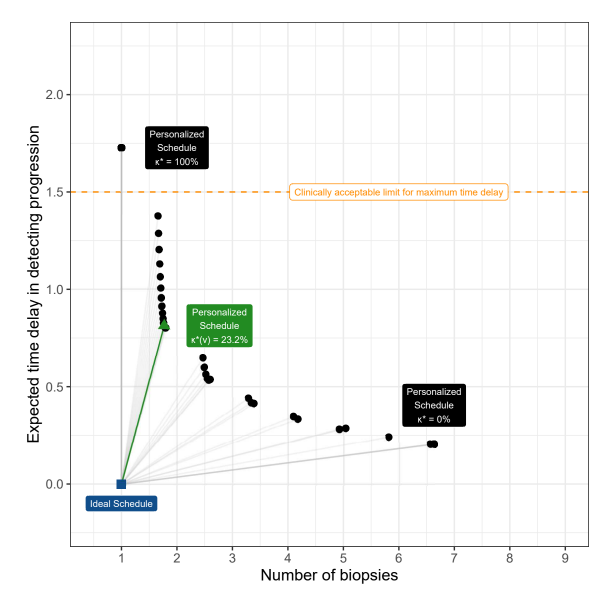


Figure 4. Automatic choice of risk threshold  $0 \le \kappa^* \le 1$  using (5). The ideal schedule of tests at point (1,0) is shown as a blue square. It plans exactly one invasive test at the true time of progression  $T_j^*$  of a patient and hence leads to a zero time delay in detecting progression. Personalized schedules based on a grid of thresholds chosen between  $0 \le \kappa^* \le 1$  are shown with black circles. Higher thresholds lead to fewer tests, but also higher expected time delay. We propose to choose the personalized schedule based on  $\kappa^*(v) = 21.2\%$  threshold (green triangle). This is because it has the least Euclidean distance (shown with a green line) to the ideal schedule. It is also possible to find the least distance under a certain clinically acceptable limit on time delay (orange dashed line), or number of tests.

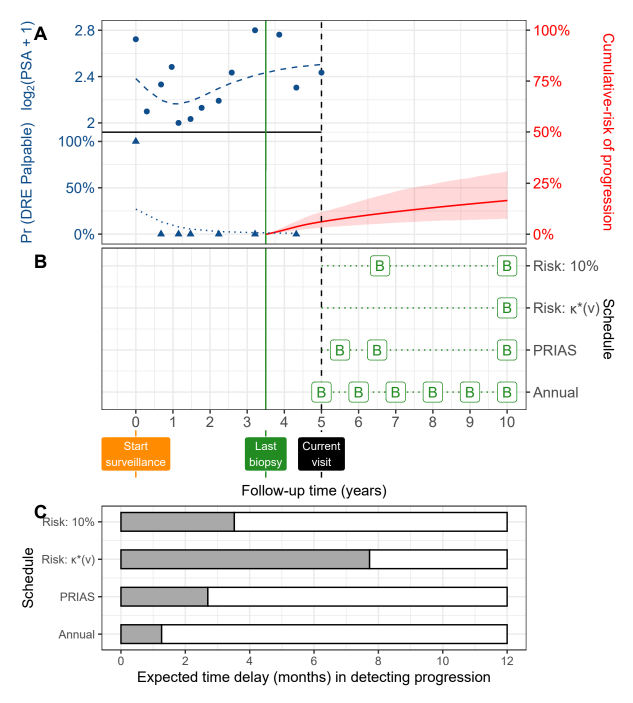


Figure 5. Demonstration of personalized schedules for a real PRIAS patient: In Panel A: Time of last negative biopsy is year 3.5 (vertical green solid line). Longitudinal data is repeated DRE (blue triangles) and PSA measurements (blue circles). The current visit is year five (vertical black dashed line). The estimated cumulative-risk profile is shown with a solid red line (95%CI is shaded). It is 6% at the current visit and 16.5% at year ten (horizon). In Panel B, we visualize different biopsy schedules, with a 'B' indicating a biopsy. Risk: 10% and Risk: Auto are personalized biopsy schedules using a fixed risk threshold  $\kappa^* = 10\%$ , and automatically chosen  $\kappa^*(v)$  (Equation 5), respectively. PRIAS and Annual denote the PRIAS biopsy schedule (paragraph 2 of Section 4) and annual biopsy schedule. Panel C: For personalized, PRIAS, and annual biopsy schedules, we calculate the corresponding expected time delays in detection of progression. A compulsory biopsy at year ten was planned in all schedules for a fair comparison of expected time delay.

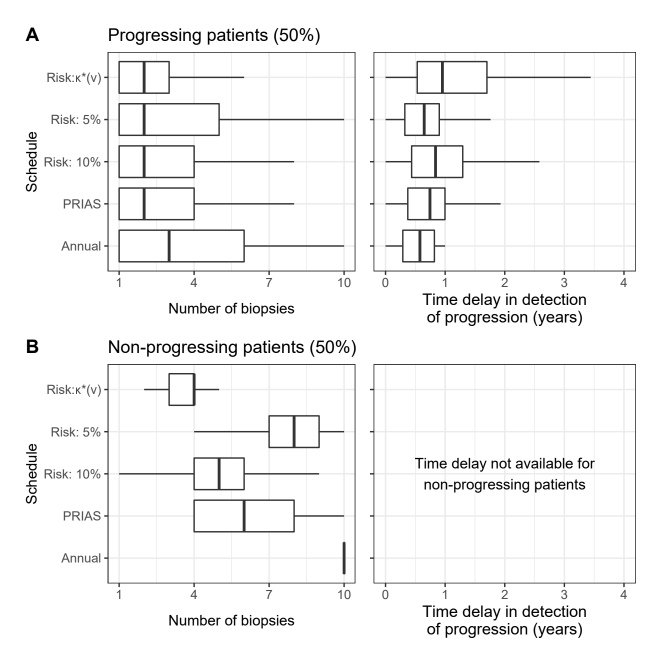


Figure 6. Boxplot showing variation in the number of biopsies, and the time delay in the detection of cancer progression for various biopsy schedules. Time delay (years) is calculated as (time of positive biopsy - the true time of cancer progression). Biopsies are conducted until cancer progression is detected. Panel A: results for simulated patients who obtained cancer progression in the ten year study period (progressing). Panel B: results for simulated patients who did not obtain cancer progression in the ten year study period (non-progressing). Types of personalized schedules: Risk: 10% and Risk: 5% approaches, schedule a biopsy if the cumulative-risk of cancer progression at a visit is more than 10% and 5%, respectively. Risk:  $\kappa^*(v)$  works similar as previous, except that a visit-specific risk threshold is chosen using (5). Annual corresponds to a schedule of yearly biopsies, and PRIAS corresponds to biopsies as per PRIAS protocol (Section 4).