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Comparison of methods for estimating the effect of salvage therapy in prostate cancer when treatment is given by indication

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Abstract

For patients who were previously treated for prostate cancer, salvage hormone therapy is frequently given when the longitudinal marker PSA begins to rise during follow-up. Because the treatment is given by indication, estimating the effect of the hormone therapy is challenging. In a previous paper, Kennedy et al (2010), we described two methods for estimating the treatment effect, called two-stage and sequential stratification. The two-stage method involved modeling the longitudinal and survival data. The sequential stratification method involves contrasts within matched sets of people, where each matched sets includes people who did and did not receive hormone therapy. In this paper we evaluate the properties of these two methods and compare and contrast them with the marginal structural model methodology. The marginal structural model methodology involves a weighted survival analysis, where the weights are derived from models for the time of hormone therapy. We highlight the different conditional and marginal interpretations of the quantities being estimated by the three methods. Using simulations, which mimic the prostate cancer setting, we evaluate bias, efficiency, accuracy of estimated standard errors and robustness to modeling assumptions. The results show differences between the methods in terms of the quantities being estimated and in efficiency. We also demonstrate how the results of a randomized trial of salvage hormone therapy are strongly influenced by the design of the study, and discuss how the findings from using the three methodologies can be used to infer the results of a trial.

Keywords

treatment by indication; time-dependent confounder; proportional hazards model; causal effect; prostate cancer

1. Introduction

In this paper we consider observational data that might arise in a prostate cancer study in which there is longitudinal data, a treatment that may be assigned at some timepoint during the follow-up and an event time outcome variable that may be censored. The goal is to estimate the effect of the treatment on the outcome variable. The longitudinal data is assumed to arise from a stochastic process. If the longitudinal process affects both the outcome of interest and the assignment of the treatment, then the longitudinal process is a time-dependent confounder. If the treatment affects the ensuing longitudinal process, then

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the process is an intermediate variable as well as a time-dependent confounder. Standard naive covariate adjustment, that adjusts for the longitudinal data, will only yield an estimate of the treatment effect beyond that due to changes in the process itself. Hence, if the longitudinal process is both a time-dependent confounder and an intermediate variable then, to estimate the treatment effect, covariate adjustment is necessary but problematic using standard methods. This situation is sometimes called treatment by indication, and the goal in this paper is to evaluate and compare various approaches to estimating the treatment effect when the treatment is given by indication.

The motivating example for this research comes from the prostate cancer setting. After initial diagnosis of prostate cancer and subsequent treatment by radiation therapy, elevated levels of prostate-specific antigen (PSA) and rates of increase of PSA indicate an increased risk for clinical recurrence of the cancer [1]. In addition, because of the increased risk, those patients with elevated PSA are more likely to initiate salvage androgen deprivation therapy (SADT) in order to prevent or delay the recurrence of cancer. In this example PSA is the longitudinal variable, recurrence time is the outcome variable and SADT is the treatment. As explained above, PSA is a time-dependent confounder in the relation between SADT and recurrence. Furthermore, patients experience a marked decrease in PSA for at least the first few months after initiation of SADT. Therefore PSA is also intermediate variables in the relation between SADT and recurrence. A standard Cox regression analysis including covariates representing time-dependent PSA, along with a time-dependent treatment indicator and other covariates, would therefore estimate the benefit of SADT beyond that due to the decrease in PSA at the time of SADT, a relatively useless quantity.

In the last 15 years marginal structural models (MSM) and related methods have been developed [2, 3, 4, 5, 6] to estimate a causal treatment effect of such a time varying treatment when there exists confounding by time-dependent covariates affected by earlier treatment as described above. This approach has been rigorously developed with an elegant theory linked to counterfactual models and randomized trials. In its simplest form, the MSM methodology can be used to estimate, from observational data, a hazard ratio between two counterfactual scenarios, one in which subjects are all treated at time τ and another in which subjects are not treated. Specifically, denote the counterfactual hazard at time t when the treatment was not assigned as $\lambda_0(t)$, and if the treatment was assigned at time τ for all subjects, the counterfactual hazard would be $\lambda_0(t) \exp[\phi I(t > \tau)]$. Here, the quantity ϕ is the causal treatment effect, which is assumed not to depend on τ or $t - \tau$, and it matches the target quantity of interest in a randomized clinical trial for which half the patients are randomized to treatment at time τ and the other half do not receive treatment, provided the assumptions of the MSM hold. Note that ϕ is a marginal quantity since it averages over subjects with possibly different hazards due to different measured and unmeasured covariates and other unexplainable sources of variability. Note also that the model defining ϕ does not condition on any time-dependent covariates. Recent causal inference literature has tended to use the terms marginal and causal interchangeably, however in this paper we will keep them as distinct because we will also be considering conditional causal effects where we condition on covariates, including time-dependent covariates. The MSM methodology [2, 4] estimates ϕ from observational data by weighting the observations to "mimic" data which would have arisen had a randomized trial been conducted. Specifically, inverse-probability-of-treatment weighting (IPTW) is used in order to estimate ϕ the marginal causal effects, and the weights are derived from models for the probability of treatment.

The marginal structural model can be extended to include baseline (but not time-dependent) covariates in the hazard [3, 4], specifically to estimate ϕ from a marginal model of the form $\lambda_0(t) \exp[X\beta + \phi I(t > \tau)]$, where X are baseline covariates. The history-adjusted MSM (HA-

In recent work [9], we presented two different methods for estimating treatment effects using observational data in situations like those presented above, where a time-dependent confounder is also an intermediate variable in the relation between treatment and outcome. One method, which we called the two-stage method, specified for each subject a model for the hazard of recurrence in the absence of treatment, called the "natural hazard". This hazard, denoted by $\lambda_i^0(t)$, can also be thought of as the counterfactual hazard for that person if he never receives the treatment. In this model, the hazard for subject *i* at time *t* is given by $\lambda_i^0(t)$, so that if the subject were to be assigned treatment at time τ then the hazard for that subject would be $\lambda_i^0(t) \exp[\gamma I(t > \tau)]$. The method links $\lambda_i^0(t)$ to the process for the longitudinal data, and then jointly estimates γ and $\lambda_i^0(t)$. The other method, called Sequential Stratification ([10, 11]), matches those patients who received treatment (called index cases) to similar patients still at risk, thereby reorganizing observed data to mimic a sequence of conditionally randomized treatment assignments. The estimation then proceeds by fitting stratified models and comparing patients within strata. Both these methods can be thought of as estimating conditional treatment effects, since they condition on subject-specific factors that could be time-dependent.

The question of whether the quantity of interest should be a marginal or a conditional causal effect, as formulated here, depends on the clinical context in which it would be used. For health policy situations, one is often interested in making guidelines for groups of patients and results from randomized trials of groups of patients would be considered the gold standard; thus, in such cases, estimates from marginal models would be desirable. In clinical settings, where subject-specific decisions regarding treatment are paramount, conditional treatment effects may be more useful. In the context of prostate cancer recurrence, the patient will know his baseline covariates and his pattern of PSA up to the current time, and hence it would be more valuable from a clinical perspective for an individual patient to know, under multiple salvage treatment options, his risk of recurrence, as opposed to the risk of recurrence among a wide array of patients with varied PSA patterns. The randomized trial that would be relevant for this patient would be one which only enrolled patients who had similar amount of follow-up since the the initial therapy and also a similar pattern of PSA values.

In our previous work [9], we described the two-stage and sequential stratification methods, but we did not evaluate their properties via simulation. Similarly, simulation-based evaluations of the MSM, and comparisons of the MSM with other methods, are limited in the literature. Young et al. [12] compared two types of structural nested models (SNMs) with the MSM, finding that the MSM is advantageous with respect to bias, variance, and ease of computation. Xiao et al. [13] compared the Cox MSM to the pooled logistic MSM (commonly used as an approximation to the Cox MSM) across varying weighting schemes, reporting that the pooled logistic MSM yields estimators with larger variances than the Cox MSM, and that normalized and stabilized weights outperform weights which are either unstabilized or unnormalized or both. Westreich et al [14] found good bias and coverage rate properties of MSM methods but sometimes with less precision compared to simple methods depending on how the weights were implemented. Their work also demonstrated the benefit

of using stabilized weights. Ertefaie et al. [15] compare IPTW and propensity score methods, finding that propensity score methods surpass IPTW methods with respect to mean squared error in both point-treatment and longitudinal settings. In the current paper the design of the simulation is strongly linked to the motivating prostate cancer study.

Two basic premises in this paper are that (i) there exists heterogeneity in the disease process among individuals, and (ii) subject-level data in observational studies arise from realizations of stochastic probability models. This matches in spirit the concepts of causality discussed in Aalen and Frigessi [16], Aalen et al [17] and Commenges and Gegout-Petit [18]. In the prostate cancer context there are four relevant linked stochastic processes, one for the longitudinal PSA data, one for the recurrence of the cancer, one for the assignment of treatment, and one for censoring. The three estimation methods we compare either make assumptions such that some of these stochastic processes can be ignored, or else require specification of models for one or more of these stochastic processes. The model for recurrence includes a parameter (γ) representing the multiplicative effect of treatment on the hazard of recurrence; this quantity is the conditional causal effect of treatment and is the quantity of interest when one is interested in subject-specific effects of treatment. The marginal causal effect of treatment for a heterogeneous group of patients is determined by the stochastic models for PSA and recurrence, along with the posited treatment assignment of interest, and may not equal γ .

The purpose of this paper is to evaluate, via simulation, the two-stage, sequential stratification and MSM approaches in the context of the prostate cancer example. The simulation scheme includes a longitudinal biomarker, a treatment process which may be predicted by the biomarker, and an event process which is related to values of the biomarker in addition to treatment status and a censoring process. In other words, we specify a true probability model for the biomarker, treatment, and recurrence, each defined at the subject-specific level. We will compare and contrast the methods themselves, along with the quantities they estimate, their properties, as measured by bias and efficiency, and their robustness to modeling assumptions, as well as to various types of censoring mechanisms.

2. Motivating Prostate Cancer Example

The prostate cancer datasets to which we applied the two-stage and sequential stratification methods in [9] have the following structure. All patients are diagnosed with localized prostate cancer and treated with external beam radiation therapy. Patients have pre-treatment characteristics, such as T-stage, which we denote by \mathbf{x}_i for subject *i*. Each patient has a sequence of values of PSA after the radiation therapy and these are used to monitor the patient. Time t is measured in years from the end of radiation therapy. The typical pattern of PSA after radiation therapy is well known, and associated with some of the pre-treatment variables. It decreases in everyone for about a year and then may or may not start to rise; if it does rise, it increases approximately exponentially with time. Rising values of PSA are indicative of tumor cells growing and dividing, but the tumor may not have yet grown to such a size that it is detectable. The time of clinical recurrence is the time at which the tumor is detected, which we call R_i , and that is the event of interest in our research. Let C_i denote the censoring time. If the values of PSA start to rise, the patient and their doctor may consider starting SADT prior to any recurrence; we denote the time of initiating SADT as S_i . While there are guidelines for when SADT should be initiated, in typical observational patient series there is considerable heterogeneity in the values of S_i , and SADT is not always initiated. SADT quickly reduces the values of PSA in just about all patients, and to near zero in most patients, but later PSA may rise and the patient may experience clinical recurrence. In none of the modeling or analysis we undertake do we consider the observed values of PSA after S_i . The data structure is depicted in figure 1. In this prostate cancer setting there is

very strong belief that SADT delays clinical recurrence, but the amount by which it delays recurrence or reduces the risk of recurrence is not well quantified.

Randomized clinical trials would be one way to investigate the effects of SADT. Given the uniform belief that SADT is effective at delaying clinical recurrence, it would be unethical to run a randomized trial in which SADT was withheld. Trials that would be interesting from a treatment policy perspective are ones that compare early to late SADT where early and late may be determined by the values of PSA, or ones that compare giving everyone SADT at the same time as radiation therapy with a strategy of giving SADT in follow-up as suggested by high or increasing values of PSA. While such trials would be ideal, they have not been undertaken. Thus the challenge is understanding what one might find from such trials by analyzing observational data. For an individual patient in active follow-up, with his sequence of PSA values, it would not be viable to run a randomized trial that exactly matches his situation. For him the relevant question is what is the future risk of recurrence if he does start SADT compared to not starting it.

3. Methods

Here we describe three potential methods to estimate the treatment effect from the type of observational data described above.

3.1. Two-Stage Method

The two-stage method, with full details available in [9], specifies a form for the 'natural hazard' (the hazard of recurrence in the absence of treatment by SADT) for subject *i*, given by $\lambda_i^0(t)$. At times after initiation of SADT, this hazard changes to:

 $\lambda_i^0(t) \mathrm{exp}(\gamma)$ (1)

The form of $\lambda_i^0(t)$ depends on baseline covariates \mathbf{x}_i and is linked to the PSA process for subject *i*. Since we will assume that the PSA process is determined by subject-specific random effects and \mathbf{x}_i in a mixed model, $\lambda_i^0(t)$ is also determined by the subject-specific random effects and \mathbf{x}_i . The two-stage method estimates both $\lambda_i^0(t)$ and γ . In the first stage we estimate the biomarker process for PSA for each subject in the absence of treatment by SADT (i.e., using only data prior to initation of SADT). Quantities estimated from the first stage are provided to the second stage. In the second stage we estimate the treatment effect γ using a Cox proportional hazards model. The models we will be assuming for the longitudinal PSA process and for $\lambda_i^0(t)$ have a similar form to those that were developed in [19], and are derived from analysis of the data described in that paper.

The assumed model for PSA in the absence of treatment by SADT is:

$$\log P_i(t) = \log PSA_i(t) + \varepsilon_{it} = (\alpha_0 + a_{i0}) + (\alpha_1^T \mathbf{x}_i + a_{i1})f(t) + (\alpha_2^T \mathbf{x}_i + a_{i2})t + \varepsilon_{it}$$
(2)

where $P_i(t)$ are the observed values of PSA for subject *i* at time *t*, (a_0, a_1, a_2) are fixed effect parameters, (a_{i0}, a_{i1}, a_{i2}) are subject-specific random effects, and \mathbf{x}_i is a covariate vector including an intercept term and baseline T-stage indicators (i.e., *I*(T-stage = 2) and *I*(T-stage 3)). $f(t) = (1 + t)^{-1.5} - 1$ captures the short-term evolution of PSA, while *t* captures the long-term evolution. We assume the measurement error $\varepsilon_{it} \sim N(0, \sigma^2)$, and the random effects $(a_{i0}, a_{i1}, a_{i2}) \sim \text{MVN}(0, \Sigma)$.

The resulting BLUP estimates for logPSA_{*i*}(*t*) and logPSA[']_{*i*}(*t*), where logPSA[']_{*i*}(*t*) denotes the slope of log PSA_{*i*}(*t*), are given by log PSA_{*i*}(*t*) and logPSA[']_{*i*}(*t*), respectively. The assumption regarding the natural hazard, (i.e., for recurrence in the absence of SADT) is that

$$\lambda_{i}^{0}(t) = \lambda_{0}(t) \exp\left[\theta_{0}^{T} \mathbf{x}_{i} + \theta_{1} \log \mathrm{PSA}_{i}(t) + \theta_{2} \log \mathrm{PSA}_{i}^{'}(t)\right] \quad (3)$$

Then combining with equation (1) the following time-dependent Cox model is then fit to estimate θ_0 , θ_1 , θ_2 and γ .

$$\lambda_{i}(t) = \lambda_{0}(t) \exp\left[\theta_{0}^{T} \mathbf{x}_{i} + \theta_{1} \log P\widehat{S}A_{i}(t) + \theta_{2} \log P\widehat{S}A_{i}^{'}(t) + \gamma I(t \ge S_{i})\right]$$
(4)

where S_i is the time of SADT and the BLUP estimates $\log P\hat{S}A_i(t)$ and $\log P\hat{S}A_i'(t)$ are calculated for times both before and after S_i . Note that in the estimation in equation (4) $\lambda_0(t)$ is not assumed to be constant with respect to time, and is treated non-parametrically in the usual Cox model fashion.

Note that $P\widehat{S}A_i(t)$ and $P\widehat{S}A_i'(t)$ are estimates assuming SADT is not given, this eliminates the concern described in the introduction about PSA being an intermediate variable. Thus the two-stage method essentially compares what happened to people who were treated with an estimate of what would have happened to them if they had not been treated. In this sense it has some similarity to what is modeled in Structure Nested Models (SNM), where the g-estimation algorithm is used to estimate the unknown parameters ([20]).

There are a number of issues and challenges associated with this two-stage approach. A basic assumption is that the quantity γ is the same for all people. The parameters in the model would still be identifiable if γ were allowed to depend on baseline or time-dependent covariates, however additional subject-specific values of γ are not estimable. The method requires fully specifying longitudinal and survival models, thus there are legitimate questions about the robustness of the estimates of γ to misspecification of these models. Finding a good model may be challenging; however, in the prostate cancer example, PSA and recurrence data of the type used here have been collected for many years in many different studies, giving good knowledge about the structure of these models. The expression for the hazard in Equation (3) has as covariates a smoothed version of PSA and its slope, and may require extrapolation of these values, thus this method would only be applicable in situations for which it would seem plausible to extrapolate the longitudinal variable into the future. An implicit assumption in this method is that the treatment assignment depends on PSA, and that there are no other unmeasured factors that may affect the treatment or that are associated with PSA or recurrence. While we perform the estimation in two stages, it is certainly possible to fit the longitudinal and survival models jointly ([21]). The joint estimation would likely lead to better estimates of γ in some situations. The joint estimation method is much more computationally intensive, so we will use the simpler two-stage estimation in our numerical work.

3.2. Sequential Stratification

The sequential stratification (SS) method [10, 11] reorganizes observed data in an attempt to mimic a sequence of conditionally randomized treatment assignments. At the time of each treatment initiation, similar patients at risk who have not initiated treatment are matched to the patient initiating treatment; this process generates one stratum for each treated subject in the data. Then, a stratified Cox proportional hazards model is fit in order to estimate the treatment effect, allowing for differing baseline hazards across strata.

Let $S_{(j)}$ be the *j*th ordered time of SADT initiation, $j = 1, ..., n_S$, where n_S is the total number of patients undergoing SADT. With respect to the *j*th patient to initiate SADT (index case (*j*)), we define $e_{ij} = 1$ if patient *i* is at risk at time $S_{(j)}$ and has a similar PSA pattern, and $e_{ij} = 0$ otherwise. Specifically, the stratum-inclusion indicator for patient *i* is given by:

$$e_{ij} = I[\min(S_i, R_i, C_i) \ge S_{(j)}, \|P_i(S_{(j)}) - P_{(j)}(S_{(j)})\| \le \delta_{jk}] \quad (5)$$

where $P_i(t) = (\log P\widehat{S}A_i^*(t), \log P\widehat{S}A_i^{'*}(t), \widehat{a}_{i2})$ is a vector of the BLUP estimates of log PSA, slope of log PSA at time *t*, and the random effect for time, standardized across *i* to have mean zero and variance 1, and R_i is the recurrence time and C_i is the censoring time. $||P_i(S_{(j)}) - P_{(j)}(S_{(j)})||$ indicates the Euclidean distance between the vectors of BLUP estimates for subject *i* and the index case at time $S_{(j)}$, and δ_{jk} is chosen so that exactly *k* patients have $||P_i(S_{(j)}) - P_{(j)}(S_{(j)})|| = \delta_{jk}$. Therefore each stratum consists of the index case (the patient undergoing SADT), along with the matched (with respect to standardized current logPSA, current slope of logPSA and long term slope of logPSA) *k*-nearest-neighbor patients still at risk at the time of initiation of SADT. We also only considered matches who had the same baseline T-stage as the index case. We used k = 3 if three or more potential matches were available, and all available matches if less than three were available.

Once strata are defined, we fit the following model, which assumes that for patient i in stratum (j) the hazard is given by:

$$\lambda_i(t) = \lambda_{0(j)}(t) \exp[\omega_0 \widehat{a}_{i2} + \omega_1 \log P\widehat{S}A_i^*(S_{(j)}) + \omega_2 \log P\widehat{S}A_i^{**}(S_{(j)}) + \eta I[i=(j)]] \quad (6)$$

where $(j) = 1, ..., n_S$, and I[i = (j)] is an indicator for patient *i* being the index case, and the estimate of η is the quantity of primary interest. The BLUP estimates of log PSA and slope of log PSA (at the respective times of SADT initiation) are used as adjustment covariates as well as matching criteria in order to account for any residual heterogeneity within strata. The estimate of the random effect a_{i2} is included because it can be viewed as a predictor of future PSA values. A robust variance estimator is used and matched patients (non-index-cases) who later undergo SADT are censored at the time of their SADT.

Additional comment on the use of the random effect, a_{i2} in the matching and in equation (3), is in order. In developing the sequential stratification method, Schaubel et al [10, 11] did not require modeling of the longitudinal process. However, the methods did require a modified version of Inverse Probability of Censoring Weighting (IPCW) in order to account for the dependent censoring of treatment-free recurrence caused by the receipt of SADT. The version of SS evaluated in this report does not involve inverse weighting. However, since the analysis is conditional on a_{i2} , which essentially accounts for future treatment propensity, bias due to dependent censoring should be minimal.

There are a number of issues and challenges associated with the sequential stratification approach. In this method we form strata of similar subjects, but there are choices to be made about the size of the strata and how the strata are formed. Some of these choices were investigated in Kennedy et al (2010), where we relied on the matching to achieve homogeneous strata, but in this paper we have also included adjustment covariates in equation (6). In general, decisions need to be made about which factors are used to define strata and which are incorporated as adjustment covariates in the model of interest. In the matching procedure, we match on the BLUP estimates of PSA, slope of PSA and \hat{a}_{i2} from the longitudinal model, but this was not strictly necessary: one could instead match on the observed values of PSA without the need to fit a longitudinal model. This method has some similarity to propensity score matching, but propensity score matching would aim to match

on subjects who had similar probability of obtaining treatment, whereas we aim to match on patients who have the same prognosis, similar to the idea of prognostic matching ([22]). In the prostate cancer example, these are thought to be similar. In principle we could refine the matching on prognosis, by including in the matching criteria quantities such as the projected PSA value 2 years into the future or an estimated probability of recurrence within, say, 3 years. These approaches would give more homogeneous strata with respect to prognosis.

A further challenge with respect to the SS method is variance estimation. The articles proposing the SS method both suggested the use of the bootstrap. Since estimating equation methods are used to derive the method, it is possible that a robust (sandwich) variance estimator could be used instead. Since use of the bootstrap is computationally demanding, we use a robust variance estimator in this paper.

3.3. Marginal Structural Model

In the context of survival analysis, inverse-probability-of-treatment weighted (IPTW) estimators for the parameters of a marginal structural model (MSM) [3, 4] can be obtained via a Cox model for which contributions to the partial likelihood are weighted differentially across subjects and across time, where the weights are first calculated at a discrete set of time points. In this paper, we closely follow the methods and code given in [4].

First, the time scale is discretized into many small intervals, with the interval endpoints denoted by t_0, t_1, t_2, \ldots . Then subject-specific time-varying weights are computed using estimated probabilities from two separate logistic regression models. The first model regresses the probability of not initiating treatment at time t_j (conditional on not having already initiated treatment by time t_{j-1}) on baseline covariates:

$$\operatorname{logit}[\operatorname{Pr}(S_i > t_j | S_i > t_{j-1})] = \tilde{\beta}_0(t_j) + \tilde{\beta}_1^T \mathbf{x}_i \quad (7)$$

The second model regresses this probability on both baseline and time-dependent covariates:

$$\operatorname{logit}[\operatorname{Pr}(S_i > t_j | S_i > t_{j-1})] = \beta_0(t_j) + \beta_1^T \mathbf{x}_i + \beta_2 \operatorname{logPSA}_i(t_j) + \beta_3 \operatorname{logPSA}_i(t_j) \quad (8)$$

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We also considered an alternative for the second model

$$\operatorname{logit}[\operatorname{Pr}(S_i > t_j | S_i > t_{j-1}|)] = \beta_0(t_j) + \beta_1^T \mathbf{x}_i + \beta_2 \operatorname{logP}_i(t_j) \quad (9)$$

Note that in equation (8) the initiation of SADT depends on estimates of the value and slope of PSA, which are both important variables for the hazard of recurence, while in equation (9) the initiation of SADT depends only on the observed PSA value, and matches exactly the way the data is generated in the simulation study.

Let $p_{1i}(t)$ be the predicted probability for subject *i* at time *t* estimated from model (7) and $p_{2i}(t)$ be the predicted probability for subject *i* at time *t* from models (8) or (9). Then the stabilized weight for subject *i* at time t_k is given by:

$$w_{i}(t_{k}) = \prod_{j=1}^{k} \left[\frac{\hat{p}_{1i}(t_{j})}{\hat{p}_{2i}(t_{j})} I(t_{j} < S_{i}) + \left(\frac{1 - \hat{p}_{1i}(t_{j})}{1 - \hat{p}_{2i}(t_{j})} \right) I(t_{j} = S_{i}) + I(t_{j} > S_{i}) \right]$$
(10)

This weight corresponds to the cumulative product (across time) of the ratio between two probabilities: in the numerator, the probability that the subject received his observed treatment given only baseline covariates, and in the denominator, the probability that the subject received his observed treatment given both baseline and time-dependent covariates. The numerator probability is used only for stabilization purposes, and although strictly not necessary, we include it here because it has been shown to improve the properties of IPTW estimators ([23, 14]). Note that, for a given subject, the weight is constant across time after initiation of SADT.

Finally, to estimate the quantity of interest ϕ , we fit the time-dependent Cox model:

$$\lambda_i(t) = \lambda_0(t) \exp[\theta_0^T \mathbf{x}_i + \varphi I(t \ge S_i)] \quad (11)$$

with subject-specific time-dependent weights $w_i(t)$. The SE of ϕ is obtained from a robust variance estimator.

There are a number of issues and challenges associated with this MSM approach. The quantity being estimated by the MSM method is a population average treatment effect, it is an identifiable quantity, and in contrast to the two-stage method which assumes the treatment effect is the same for all subjects the MSM method does not require this assumption, however it does assume that the treatment effect does not depend on the time of initiation of the treatment. In contrast to the two-stage method and the version of SS described above, the MSM method requires specifying and fitting models for the treatment assignment. This may or may not be easier than specifying a model for the outcome, depending on the context. The models fit in the MSM method are used to estimate the weights, and it has been observed that these weights can be quite unstable, negatively affecting properties of the estimated treatment effect ([24, 25, 23]). Various strategies to control this instability have been suggested, such as truncating very large weights or using stabilized weights (as is done in this paper). Xiao et al [13] suggested normalizing the weights, but we found that was not effective in our situation. In equations (7) and (8) or (9) we have assumed the intercepts $\beta_0(t)$ and $\beta_0(t)$ are time-dependent, where we use a B-spline estmator similar to Hernán et al [4]. For the way we generated data in the simulation study, assuming constants for $\beta_0(t)$ and $\beta_0(t)$ would have been adequate, but in general assuming smooth functions for $\beta_0(t)$ and $\beta_0(t)$ would be preferable. The choice of model to obtain $p_{2i}(t_i)$ may also be important. In the data generation scheme in the simulation study the initiation of SADT is determined by observed value of $\log PSA_i(t)$, corresponding to equation (9), while the recurrence event is determined by the true value and slope of log $PSA_i(t)$, hence consideration of equation (8). We will compare these two methods of obtaining the weights. As described in [4] more complex weights can be used that also take account of censoring, by developing an additional model for the censoring time. Although in practice it would usually be preferable to perform this extra modeling, we don't include this additional weight in this paper, because it wasn't necessary for nearly all the scenarios considered in the simulation study as we don't impose any censoring. Another practical issue when fitting the Cox model (equation (11)), using the weighted partial likelihood, is that it is necessary to have weights at the times of every event, whereas the weights are only calculated at a set of discrete times. To solve this problem, either the weights have to be interpolated to all times, or the data needs to be discretized so that events and initiation of SADT occur at the same set of times. An alternative to overcome this problem is to use survival models, instead of the logistic models in equations (7), (8) and (9), then the required weights could be calculated at any time.

3.4. Marginal vs. Conditional Causal Effects

In this paper we take the parameter γ to represent the relative decrease in the hazard for each subject when they receive SADT. It is a subject-specific effect that is assumed to be the same for every person. This assumption can be weakened; specifically, it would be possible to have γ depend on either baseline or time-dependent covariates. While this would be scientifically interesting we do not consider it in this paper.

The definition of γ from Equation (1) is conditional on the unknown natural hazard curve,

 $\lambda_i^0(t)$. In the two-stage method we parameterize the natural hazard to be a function of random effects. Because of its construction, the two-stage method is attempting to estimate the quantity γ . In contrast, the MSM method is trying to estimate a different quantity that is a marginal or population-averaged quantity; it is essentially averaging over the random effects. For non-linear mixed models it is well known that population-averaged estimates are different than subject-specific estimates and tend to be closer to zero, so we would expect population-averaged estimates of the treatment effect from the MSM method also to differ from those of the two stage method. This difference between subject-specific and population averaged quantities is also refered to as non-collapsibility of measures, such as hazard ratios, in non-linear models [26].

The MSM methodology is designed to estimate the ratio of two hazards, one being what the hazard would be if SADT is never given and the other being what the hazard would be if everyone who is at risk is given SADT at time τ . The form for both hazards can be directly derived from the subject-specific models for PSA and recurrence by integrating out the random effects. We note that these hazards are population quantities which do not depend on the details of the MSM methodology for estimating the weights. If SADT is never given, the marginal hazard at time t depends on $P(R \in (t, t + \delta)|R > t)$ for small δ , which can be written as:

$$\int_{a} P(R \in (t, t+\delta) | R > t, a) P(a | R > t) da \quad (12)$$

where *a* are the random effects. For simplicity of notation, assume there are no covariates \mathbf{x}_i ; then the term P(a|R > t) can be written as:

$$\exp\left[-\int_{s=0}^{t}\lambda_{0}(s)\exp(g(a,s,\omega))ds\right]f(a)/B \quad (13)$$

where f(a) is the distribution of the random effects, ω is the collection of parameters (*a*'s and θ 's) from equations (2) and (3), $g(\cdot)$ is the linear combination of PSA and slope of PSA obtained from plugging equation (2) into equation (3), and *B* is the integral of the numerator with respect to *a*. Thus the marginal hazard is:

$$\int_{a} \lambda_{0}(t) \exp(g(a, t, \omega)) \exp\left[-\int_{s=0}^{t} \lambda_{0}(s) \exp(g(a, s, \omega)) ds\right] f(a) da/B \quad (14)$$

For the group who received SADT at time τ , the marginal hazard at times $t > \tau$ is given by

$$\int_{a} \lambda_{0}(t) \exp(g(a,t,\omega) + \gamma) \exp\left[-\int_{s=0}^{t} \lambda_{0}(s) \exp[g(a,s,\omega) + \gamma I(s > \tau)] ds\right] f(a) da/B^{*}$$
(15)

where B^* is the normalizing constant.

The ratio of these hazards from equations (14) and (15) will be one at times prior to τ , but after τ it is a complicated expression, which certainly does not equal $\exp(\gamma)$. Furthermore,

the ratio of the hazards will depend on *t*, demonstrating that if the conditional model has proportional hazards, the marginal model will not be proportional hazards in this setting. This calls into question the merits of fitting a marginal proportional hazards model. Nevertheless, the estimate one obtains using the MSM methodology could still be considered a useful summary of the marginal effect of the treatment. If the treatment has no effect then there is no proportional hazards assumption, so in this case there is no violation of assumptions in fitting a marginal model.

In the SS method strata are formed of similar subjects. If the matching was so successful that all subjects in each stratum had identical patterns of PSA and identical prognosis, then they would effectively have identical random effects and the quantity being estimated would be γ . In reality, there is some heterogeneity within each stratum, so everyone within strata would not have identical random effects. Thus the treatment effect being estimated by the sequential stratification method will be similar to but not the same as γ , because it involves averaging over the within-strata variation. If the stratification is quite coarse we might expect the estimate from sequential stratification to be closer to that from the MSM method than to γ . To minimize the impact of possibly coarse stratification we also adjusted for the stratification factors by including them as continuous covariates in the survival model. In work we do not present we found that not adjusting for PSA, slope of PSA and \hat{a}_{i2} in the stratified analysis in equation(6), gave estimated treatment effects further away from γ , compared to when we did adjust for PSA, slope of PSA and \hat{a}_{i2} . This demonstrates the value of more precise matching and adjustment. Another way to decrease the within-strata variation would be to increase the overall sample size, enabling more precise matching within strata.

The conditional treatment effect γ as defined by equation (1) is conditional on the person's random effects a_i . The treatment effect that the subject would be most interested in is one that conditions on his baseline covariates \mathbf{x}_i and his history of PSA up to the current time $\overline{P_i(t)}$. This will usually be well estimated by an estimate of γ obtained by effectively plugging in estimates of the random effects into equation (1). We contend that this would be a more useful measure of the treatment effect for patient *i* than one that conditions only on \mathbf{x}_i . There are calculators (for example, at psacalc.sph.umich.edu) that give the predicted probability of recurrence within three years for a patient who is in active follow-up given his history and pattern of PSA values. Such calculators could also give the probability of recurrence within three years, if the person were to start SADT immediately. For this calculation we contend, for the reasons given above, that γ is the more appropriate hazard ratio to consider than ϕ . Whether γ corresponds to anything that one would estimate from a clinical trial is less clear. The possible idealized trial would be one in which randomization happens at time τ , and the eligibility criteria for the trial would be people who had identical values of \mathbf{x}_i and random effects. This could be approximately achieved by enrolling subjects who had a specified value for \mathbf{x}_i and a specified path for PSA up to time τ . Having to specify \mathbf{x}_i and the path of PSA makes such a trial too restrictive and thus not feasible. However, it may be feasible to specify a set of possible values for \mathbf{x}_i and paths of PSA and then randomize within each set. If the analysis was also stratified, then this would be estimating a quantity that approximates γ . Because of its similarity to the SS method, the formulation of this clinical trial also makes it clear that SS is attempting to estimate the quantity γ .

The target quantities for the MSM method we use in this paper corresponds to a randomized trial in which subjects who are still at risk for recurrence at time τ are randomized to either SADT or no SADT. This in itself is not a very scientifically interesting or ethically plausible clinical trial, because withholding SADT until recurrence would not be allowed. The MSM methodology is flexible in that in principle, by using other weighting schemes for the final Cox model, the estimated parameter corresponds to randomized trials with different designs.

For example, if the trial design was to randomize people to either SADT now or no SADT until the first time PSA went above a certain threshold, then we would expect this trial to show a different and smaller marginal treatment effect than the simpler randomized one. However, the same conditional treatment effect γ would apply, and the marginal treatment effect from such a trial could be derived from γ by integrating out the random effects. Thus the conditional treatment effect γ can be regarded as a fixed inherent quantity that is not influenced by the design of the clinical trial, whereas the quantity being estimated by the MSM is a function of both γ and the design of the proposed clinical trial.

4. Data Simulation

Here we present simulation models used to generate realistic looking data for PSA, treatment by SADT, recurrence, and censoring. The generating models are designed to reflect the process by which data would arise in a clinical setting. Each model is a slightly simplified version of what is estimated from the real data.

We consider discrete and evenly distributed time points, with observation frequency f = 10 (number of evenly-spaced observations per year) and study duration K = 12 years. Let $T = \{0.1, 0.2, 0.3, ...\}$. PSA measurements, initiation of SADT, recurrence and censoring can only occur at this set of times. If more than one is simulated to occur at a specific time, then the sequence of them occuring is censored first, then PSA, then recurrence, then SADT.

4.1. Generating Model for PSA

Following [9, 19], for subject *i* at each time point after start of follow-up, we simulate observed PSA values (denoted by $P_i(t)$) from the following mixed model:

$$\log P_i(t) = \log PSA_i(t) + \varepsilon_{it} = (\alpha_0 + a_{i0}) + (\alpha_1^T \mathbf{x}_i + a_{i1})f(t) + (\alpha_2^T \mathbf{x}_i + a_{i2})t + a_{i3}t^2 + \varepsilon_{it}$$
(16)

where (a_0, a_1, a_2) are fixed effect parameters, $(a_{i0}, a_{i1}, a_{i2}, a_{i3})$ are subject-specific random effects, and \mathbf{x}_i is a covariate vector including an intercept term and baseline T-stage indicators. At a given time *t*, we assume the measurement error $\varepsilon_{it} \sim N(0, \sigma^2)$, and we assume the random effects $(a_{i0}, a_{i1}, a_{i2}) \sim \text{MVN}(0, \Sigma)$ and $a_{i3} \sim N(0, \tau^2)$. This model differs from that assumed for the two-stage estimation method only by the inclusion of a quadratic term t^2 . Note that, given the random effects, and in the absence of any treatment after time t = 0, $PSA_i(t)$ would be known and non-random for all *t*.

4.2. Generating Model for Treatment by SADT

For subject i we simulate the time of SADT by first calculating for each t in T a sequence of probabilities from the equation:

$$p_i(t_j) = \exp\left[\beta_0 + \beta_1^T \mathbf{x}_i + \beta_2 A_i(t_j) + \beta_3 \log P_i(t_j)\right] \quad (17)$$

where $(\beta_0, \beta_1, \beta_2, \beta_3)$ are fixed effect parameters for the intercept, baseline covariates \mathbf{x}_i , age $A_i(t)$, and observed time-dependent logPSA values. We then simulate $S_i(t_j) \sim$ Bernoulli $(p_i(t_j))$, and since subjects stay on treatment once treatment is initiated, the time to initiation of SADT for subject *i* is $S_i = \min\{t_j: S_i(t_j) = 1\}$.

4.3. Generating Model for Recurrence

For subject *i* we simulate the recurrence time given S_i by first calculating the hazard function at any time t from the following model:

$$\lambda_i(t) = \lambda_0 \exp[\theta_0^T \mathbf{x}_i + \theta_1 \log \text{PSA}_i(t) + \theta_2 \log \text{PSA}_i'(t) + \gamma I(t \ge S_i)] \quad (18)$$

where λ_0 is the constant baseline hazard. The survival function for subject *i* is:

$$S_i(t)\exp(-\int_0^t \lambda_i(u)du)$$

then the survival time for subject *i* is generated as $R_i^* = S_i^{-1}(V)$, where $V \sim Uniform(0, 1)$, and then R_i^* is rounded up to the closest visit time R_i , or censored at 12 years at the end of the study.

4.4. Generating Model for Censoring

For subject *i* at time *t*, we either assume no censoring or simulate censoring times from the following model for the probability of censoring:

$$\rho_i(t_k) = \operatorname{expit}[b_0 + b_1 t_k + b_2 A_i(t_k)] \quad (19)$$

where (b_0, b_1, b_2) are fixed effect parameters. We assume $C_i(t_k) \sim \text{Bernoulli}(\rho_i(t_k))$, and the censoring time for subject *i* is $C_i = \min\{K \land t_k: C_i(t_k) = 1\}$, where K = 12 is the maximum follow up time. Note that if $R_i = C_i$, then follow-up is stopped at R_i ; if $R_i > C_i$, then subject *i* does not experience a recurrence. Therefore $X_i = \min(R_i; C_i)$ is the observation time for subject *i*.

4.5. Parameter Values and Simulation Conditions

Appropriate values for the parameters in models (16), (17), (18) and (19) are obtained by estimating the corresponding parameters from mixed-effects, logistic regression, and Cox proportional hazards models, respectively, fit to data for 2,781 patients with clinically localized prostate cancer, and all initially treated with radiation therapy. Baseline T-stage values are simulated from possible values (1,2,3,4) with probabilities corresponding to approximate proportions found in the real data. An older version of these data are described in [19]. When fitting the models to the simulated data T-stages 3 and 4 are combined into one category, to avoid problems with the very small numbers sometimes in T-stage 4. Ages are simulated from a $N(70, 6^2)$ distribution. We simulate 1000 datasets each with 1000 subjects, and PSA, SADT, recurrence, and censoring observations are generated. Unless otherwise stated, the true values of the PSA, SADT, recurrence, and censoring parameters are given in equations (20), (21), (22) and (23), respectively:

$$\alpha_0 = 1.635, \alpha_1 = \begin{pmatrix} 2.443\\ 0.217\\ 0.249 \end{pmatrix}, \alpha_2 = \begin{pmatrix} 0.242\\ 0.224\\ 0.547 \end{pmatrix}, \sigma^2 = 0.061, \sum = \begin{pmatrix} 1.084 & 1.065 & 0.148\\ 1.065 & 2.658 & 0.456\\ 0.148 & 0.456 & 0.322 \end{pmatrix}, \tau^2 = 0$$
(20)

$$\beta_0 = -7.258, \beta_1 = \begin{pmatrix} -0.036\\ -0.022 \end{pmatrix}, \beta_2 = 0, \beta_3 = 0.740 \quad (21)$$

$$\lambda_0 = 7.503 \times 10^{-3}, \theta_0 = \begin{pmatrix} 0.812\\ 0.918 \end{pmatrix}, \theta_1 = 0.050, \theta_2 = 2.018, \gamma = -1.5 \text{ or } 0.0 \tag{22}$$

$$b_0 = -\infty, b_1 = 0, b_2 = 0$$
 (23)

We generate two types of datasets. One type mimics an observational study in which there is variation in the time of SADT, using equations (16), (17), (18) and (19) to generate the data. With the specific parameter values as given above and with $\gamma = 0$ and 12 years of follow-up, on average 31% of people receive SADT and 38% of people experience a recurrence. The other type of dataset mimics what would arise in a randomized clinical trial with two groups, where the time of SADT differs between groups but is controlled within group according to a specified plan. The data was generated using the specified trial design and equations (16) and (18). For the observational studies, we obtain estimates of the treatment effects and their standard errors (SE) using the three methods. For the randomized trials, we simply fit a standard time-independent Cox model with treatment group as the covariate. We also fit Cox models that included both treatment group and \mathbf{x}_i as covariates, but the results are very similar and are not shown. We also fit Cox models that included treatment group, \mathbf{x}_i and the value of PSA at the time of randomization as covariates. We report the average of the 1000 estimated treatment effects, their standard deviation (SD), and the average of the 1000 SEs.

4.6. Fitting the Models

All three methods require fitting longitudinal and hazard models for which we use R. Program lmer is used for the longitudinal fitting and coxph is used for the hazard models. The R function bs() is used in the MSM methodology, where the degrees of freedom is set to 5 and all other parameters are set at default values. For the two-stage method and MSM method, the final Cox models have time-dependent covariates or time-dependent weights. For these we format the dataset for the function coxph(), such that for each subject, we discretized the time into intervals (0, 0.1], (0.1, 0.2], ..., and every time-dependent covariate or weight takes a constant value within each interval.

For the two-stage method, the time-dependent covariates (PSA and slope of PSA) take the value corresponding to the time at the end point of each interval. For the MSM method, the value of the weight in the interval $(t_{j-1}, t_j]$ is the value calculated from equation(10) at time t_{j-1} .

The actual weights used in the Cox model fit do vary between subjects and over time, but were not observed to be too extreme. Specifically for a random sample in the standard application of the method the 5th to 95th precentile range was approximately (0.38,1.29) and less than 0.2% of the time were they greater than 10.

5. Results

5.1. Evaluation of bias and efficiency of the three methods

Table 1 shows the results from simulated observational data, when there is a strong treatment effect and when there is no treatment effect. For standard application of the methods, in the case of a strong treatment effect, the two-stage method and SS give estimates which are moderately close to the true value of γ . As expected the MSM gives estimates closer to zero. The two-stage method is more efficient than the SS method, as measured by the SD, and both are more efficient than the MSM method. The SEs are close to the SD for the two-stage method, suggesting that the SEs do give appropriate measures of uncertainty. However, the SEs are somewhat too small for the SS method and the MSM method. When there is no treatment effect (i.e., when γ equals zero), the bias is small, but not zero. In this simulation design the MSM method has the largest bias, in other settings we observed the two-stage method to have larger bias. The results for the two MSM methods,

All the methods (except MSM(obsPSA)) utilize the longitudinal modeling of PSA in some way, specifically by using the BLUP estimates of $PSA_i(t)$. They are used either directly in the two-stage method, or to form matches for SS, or in the model for the probability of SADT in the MSM method. Compared to the true values of $PSA_i(t)$ these BLUP estimates will have some bias and uncertainty associated with them, and since the observations are generated based on the true values of PSA this uncertainty may lead to some bias in the estimates of the treatment effect. To investigate the impact of this uncertainty, we applied each estimation method using the true values of PSA and slope of PSA for the methods instead of the BLUP estimates. These results are presented in Table 1. This change appeared to have little impact on the bias and variability of the estimates from any of the methods.

The results for the larger sample size n=5000 show no real change in bias, but an expected reduction in precision for the two-stage and the SS method, but interestingly less gain in precision for the MSM method, for a five fold increase in the sample size compared to the standard case.

In Table 1 under "Analysis using Random Treatment Times", we show the results from simulated observational data in which the probability of receiving SADT is constant across time, and does not depend on the values of any covariates. Specifically, in Equation (17), $\beta_0 = -5.485$ is the intercept term, and we set $\beta_1 = \beta_2 = \beta_3 = 0$ for generating the data. Thus, in this case, the true weights for the MSM (at all times and for all subjects) should equal one. All methods now have small bias and except for Sequential Stratification have SEs that appropriately match the SD. There is less bias in the SS method as an estimate of γ , presumably because it is now easier to find similar people for each strata. The MSM, which gives appropriate estimates for both values of γ , matches the case when the weights are assumed to equal one. Also the MSM method is as efficient as the two-stage method in this case. We believe the reason that MSM gives better estimates of the SE is because the weights have less variability in this situation and estimation of them is less challenging.

To understand the target quantity for the MSM, we performed the numerical integrations as described in section 3.4. We found that the marginal hazard ratio was not constant and did become closer to one at longer times after treatment. Specifically at times right after the SADT treatment the log(hazard ratio) was close to -1.5, and then increased approximately linearly to be close to -1.0 twelve years later. Thus marginal proportional hazards does not hold. Hence if a constant hazard ratio is assumed we would expect the MSM method to estimate an intermediate value between -1.0 and -1.5, as it does. To further investigate this we simulated data from two randomized trials, where those at risk for recurrence at 3 or 6 years were randomized to either SADT or no SADT. The results shown in Table 2 for the estimated log hazard ratio are in the range of -1.1 to -1.3 when true γ is -1.5. The quantity derived from the MSM methodology (approximately -1.25) is also as expected in the -1.1 to -1.3 range. When the analysis of the randomized trial data also adjusted for the PSA value at the time of randomization, the estimated treatment effect is closer to the value of γ . The reason for this is because this analysis method more closely matches the conditional treatment effect, rather than the marginal treatment effect.

5.2. Robustness to misspecification of models

All three methods use models, and violation of the assumptions of these models could lead to poor properties of the methods.

To investigate the robustness of the methods to misspecifications of the correct structure of the longitudinal model for PSA, we simulated observational data in which individuals could have long-term quadratic trends, but fit a longitudinal model in which we assumed that the long-term trends were linear. The results in Table 3 show increasing bias for all methods of estimation with increasing τ . We speculate that for the two-stage method this is due to model misspecification, for SS it is due to the difficulty in finding matches who have similar prognosis and for MSM is due to the increased difficulty in estimating weights when there is more heterogeneity in the observed data. However, it should be noted that a value of 0.05 for τ is quite large and the lack of fit of equation (2) would likely be detectable from the observed data.

A necessary assumption for the interpretation of the treatment effect γ for the two-stage method is it that does not vary from one person to the next. In Table 4 we show results where there is heterogeneity in γ , specifically in generating the recurrence time using equation (18), we used $\gamma_i \sim U (\gamma \sim 0.75, \gamma + 0.75)$ instead of γ . The results show that there is little change in the quantities being estimated by all three methods compared to the standard application in Table 1.

To investigate the robustness of the MSM to misspecification of the model for SADT, and to investigate whether the two-stage or SS methods are sensitive to varying treatment assignment processes, we modified the model for simulating the initiation of SADT. In Table 4, the results are given in the case where in the simulated data we allow age to affect the probability of receiving SADT, but do not allow for this possibility in the model for SADT that gives the weights in the MSM method. Specifically, in Equation (17), we take $\beta_0 = -7.726$, $\beta_1 = (-0.086, -0.038)^T$, $\beta_2 = 0.20$, and $\beta_3 = 0.523$ for generating the data. The results from Table 4 are similar to those from Table 1, and thus the three methods are robust to misspecification of this type. The MSM is not affected in this case since age is a baseline covariate, and therefore, if age were included in both equations (7) and (8), the estimated weights in equation (10) would be approximately proportional to the weights that are computed without including age.

In another set of simulation results we investigate the effect of different censoring mechanisms. Results in Table 4 show that adding random censoring times (by taking $b_0 = -5.600$, $b_1 = 0.100$, and $b_2 = 0$ in Equation (19)) has little effect on the bias of any of the methods. Also, as expected more censoring does increase the SD and the SE.

Prostate cancer is a disease of older men; since the age of a subject will also affect the censoring rate, we simulated data with age-dependent censoring. It is also thought that older men are less likely to be given SADT, since such men could be more frail and therefore unable to tolerate potential side effects, or since SADT is thought to be less effective for older men, or since SADT could be considered less necessary for those with shorter life expectancies. Although we could not detect any age effect in real data, we include age in the simulation as a modifier of the probability of SADT. Specifically, we generate data using $\beta_0 = -7.726$, $\beta_1 = (-0.086, -0.038)^T$, $\beta_2 = 0.20$, and $\beta_3 = 0.523$ in Equation (17), and $b_0 = -8.03$, $b_1 = 0.25$, and $b_2 = 0.05$ in Equation (19). From the results in Table 4, we see that introducing age into the models that generated the data, but not the models that are used in the three estimation methods had little effect on the bias of any of the methods compared to what was seen in Table 1. Even though, this is exactly the situation in which an additional model for censoring would be considered necessary to correctly calculate the weights for the MSM method.

For all the above simulation scenarios we calculated the correlation between the estimates from the three methods, to assess whether for a particular dataset if one methods gives a

high value for the treatment effect, do the other methods tend also to give high values. The methods were correlated, the correlation between two-stage and SS was typically greater than 0.7 and the correlation of MSM with the other methods was typically greater than 0.4.

5.3. Results from Randomized Clinical Trials

As a last set of simulations we consider 4 different designs for randomized clinical trials. In all these simulations the true value of γ is –1.5. The results for the treatment effects are from simple analyses of the event times in the trial where a Cox model is fit and the only covariate is the treatment group indicator, and do not involve fitting any longitudinal models, or any time-dependent hazard models or calculating any weights.

In the Trial A (Simple Randomization at Baseline), there are three scenarios. In all we randomize 2,000 subjects at baseline and one arm never receives SADT prior to recurrence. The other arm is either to receive SADT immediately, to receive it at three years or to receive it at six years. While these trial are not ethically feasible, or even very scientifically interesting, they do demonstrate in Table 5 a decreasing marginal treatment effect in the trial if SADT is delayed.

In Trial B (SADT by Indication v's. no SADT), we randomize 2,000 subjects, at the first time PSA and slope of PSA rise above some threshold, into two arms: in the first arm, subjects receive SADT immediately, and in the other arm, subjects never receive SADT. Again, these trials are not ethically feasible; however, the estimates one would obtain from such trials are likely to correspond more closely with the quantity the two-stage and SS methods are estimating. The results in table 5 show that to be the case.

In Trial C (Early v's. Late SADT), we randomize 2,000 subjects, at the first time PSA and slope of PSA rise above some threshold, into two arms: here, in the first arm, subjects receive SADT immediately, whereas in the other arm, subjects receive SADT when their PSA and slope of PSA rise above some higher threshold. These trials would be regarded as clinically interesting and ethical. The results in Table 5 show that as expected, even though the true subject-specific treatment effect is -1.5 for all the trials, the estimated treatment effect from the trial is much smaller and depends on the design of the trial.

In Trial D (Immediate v's. SADT by Indication), at baseline we randomize 2,000 subjects to either receive SADT immediately, or else to receive SADT when PSA and slope of PSA rise above some threshold. Again, these trial are clinically interesting. The results in Table 5 show small treatment effects that depend on the design of the trial.

One conclusion from this exercise in simulating data from randomized clinical trials is that the target quantity for the trial will depend strongly on its design. The treatment effect being estimated by the two-stage and SS methods is most closely aligned with the target quantity in the trials in B. The treatment effect being estimated by the MSM method is most closely aligned with the target quantity in trial A with treatment assignment at baseline.

Another conclusion from these simulations is that even though both the conditional and marginal treatment effects are large (with a log hazard ratio of less than -1), the log hazard ratio in the clinically interesting trials is much smaller, which would clearly have implications for the sample size needed to detect an effect.

The estimated treatment effects for all four of the simulated randomized trial designs are totally determined by the structure of the models for PSA and recurrence, and by the value of γ , together with the trial design. If the results for Trials C and D are to be meaningful and useful, then these models would have to be accurate. A crucial assumption for the validity of

the efficacies in C and D is that γ does not depend on covariates. It would be possible to simulate observations for which γ depends on covariates in a number of different ways; for example, it could depend on baseline covariates, such as T-stage, or it could depend on time-dependent covariates, such as age or the value of PSA at S_i , or it could depend on time since baseline or on time after S_i . All of these variations, which can be thought of as interactions, would impact the efficacy in the trials. In Kennedy et al [9], we investigated estimating these interactions using the two-stage and sequential stratification methods, finding that the quantity of the data available to us was not sufficient to obtain accurate estimates of treatment covariate interactions.

6. Discussion

Estimating treatment effects from observational data in which there is treatment by indication is challenging, and none of the methods considered in this paper are without problems. All of the methods require building models for some aspects of the observed data, and any results are likely to be sensitive to the exact choice of these models. The two-stage method requires models for the disease process. To develop such models would generally require large datasets, and would likely benefit from subject matter knowledge as well. One of the models used in the MSM methodology is for the treatment initiation process and these gives weights that are used in another part of the MSM method. The sequential stratification method can be viewed as intermediate between the two-stage the MSM methods, it uses but does not rely as heavily on the disease process models as the two stage method.

A fundamental issue which we highlight in this paper is whether the desired quantity of interest is subject-specific or marginal. The context and intended use would dictate this. The development of the methods indicates that the two-stage and sequential stratification method are estimating subject-specific quantities, while the MSM is estimating a marginal quantity. This is supported by the simulation results. The MSM is designed to give estimates that correspond to a certain randomized trial, which may or may not be clinically relevant. However, in principle the methodology is flexible enough to allow different weighting schemes that may correspond to more relevant trials, for example by using the history-adjusted MSM method. The ability to obtain sufficiently accurate estimates of the weights may be a concern for more complex weighting schemes.

In the two-stage method, the subject specific treatment effect is defined conditional on latent variables; thus, it is not identifiable without distributional assumptions about the latent variable. Also, the methodology is only applicable in situations where the longitudinal process can be predicted into the future. The nature of changes in PSA, which mirror tumor growth, makes this possible in the prostate cancer example, but may not be possible in other examples.

While the MSM estimate derived from observational data is generally thought of as representing what the results of a randomized trial would be. However, for this to be reliable, it is necessary that the assumptions in the marginal model are appropriate, specifically the assumptions of proportional hazards, and that the hazard ratio does not depend on the starting time of the trial. In the prostate cancer example these assumptions would not be satisfied, so it is unclear in this case what the quantity being estimated from the observational data by the MSM represents. This suggests the need for some research into model checking procedures when fitting marginal models that involve estimated weights.

The sequential stratification method has a number of features that can be optimized, these include the size of the strata and how much you adjust for other variables in the stratified analysis. In previous work ([9]) we investigated the strata size and did not adjust for other

variables. In this paper we found that adjusting for other variables was beneficial for estimating the subject-specific treatment effect. One feature of the SS method which still needs development is estimation of the standard errors. We used a sandwich estimator to account for the fact that subjects could be in more than one strata, yet the standard errors were still lower than the empirical SD's in the simulation studies. Previous articles proposed the bootstrap to estimate the variance of the SS treatment effect estimator ([10, 11]). Although this may indeed be a solution in certain settings, it should be noted that the variability associated with the matching process is not accurately captured by the bootstrap for several matching methods; most notably those using nearest-neighbor matching [29]. Other options for forming the strata include either random sampling (or perhaps selecting all subjects) within categories of a discrete covariate, or using caliper matching based on a risk score. It is possible that the accuracy of the robust variance estimator depends strongly on the number of patients treated (for whom matches can be found), and strata size. Neither methods of matching nor variance estimation have been fully explored in the context of sequential stratification.

MSM is a method that is designed to analyse observational data which contains treatment by indication and then infer the results of a randomized clinical trial. The simulated randomized trial section of this paper suggests a different possible approach to this problem of infering the results of a randomized trial. The disease progression processes and treatment effects are modelled and estimated from the observational data using subject-specific models, then these estimated models are used to simulate the clinical trial of interest. This is a micro-simulation approach, which is used in the health policy area, and also has some similarities to g-computation ([28]). Both approaches have challenges, but are worthy of further evaluation in specific contexts.

All three methods described in this paper can be generalized to allow for interactions or treatment effects that are modified by covariate values. The results from the simulated randomized trials A, B, C and D, assume there are no such interactions. If there were interactions then the estimates from the randomized trials would likely change. Thus accurate estimates of these interactions will be crucial in order for estimates from observational data to be used in the micro-simulation approach. Understanding these interactions would also be important for the patient and his doctor in helping them make a decision about initiating SADT.

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Table 1

Evaluation of bias and efficiency

		$\gamma = -1.5$			$\gamma = 0$	
Method	Mean Est	SD Est	Mean SE	Mean Est	SD Est	Mean SE
		Star	ıdard Applica	ttion of Meth	spo	
Two-stage	-1.526	0.239	0.230	-0.043	0.161	0.156
Seq Strat	-1.475	0.314	0.253	0.027	0.217	0.158
MSM	-1.259	0.358	0.318	0.060	0.296	0.254
MSM (obsPSA)	-1.228	0.385	0.314	0.065	0.291	0.243
			Analysis Usi	ng True PSA		
Two-stage	-1.584	0.238	0.230	-0.076	0.158	0.156
Seq Strat	-1.546	0.321	0.259	-0.002	0.213	0.158
MSM	-1.272	0.356	0.318	0.015	0.293	0.253
		Standar	d Application	: Results for	n=5000	
Two-stage	-1.513	0.104	0.102	-0.039	0.069	0.070
Seq Strat	-1.442	0.211	0.174	0.031	0.095	0.069
MSM	-1.246	0.259	0.185	0.014	0.238	0.157
		Analysi	s Using True	SADT Prob	abilities	
MSM	-1.290	0.434	0.327	0.014	0.335	0.256
		Analysis	s Using Rand	om Treatmen	t Times	
Two-stage	-1.411	0.226	0.209	0.022	0.140	0.132
Seq Strat	-1.387	0.252	0.213	0.057	0.166	0.131
MSM	-1.269	0.214	0.204	-0.028	0.130	0.131
Unweighted MSM	-1.271	0.218	0.204	-0.030	0.137	0.131

Table 2

Randomized Trial Simulation

		,			4	
		$\gamma = -1.5$			$\gamma = 0$	
Time of Randomization	Mean Est	SD Est	Mean SE	Mean Est	SD Est	Mean SE
3 years	-1.165	0.087	0.093	-0.005	0.067	0.069
6 years	-1.283	0.116	0.119	-0.002	0.081	0.082
	ad	justed for	T-stage and F	SA at randon	nization tim	Je
3 years	-1.441	0.099	0.095	-0.001	0.076	0.069
6 years	-1.517	0.128	0.121	-0.003	0.089	0.083

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Analysis

		$\gamma = -1.5$			$\gamma = 0$	
Method	Mean Est	SD Est	Mean SE	Mean Est	SD Est	Mean SE
			$\tau = 0$.025		
Two-stage	-1.454	0.223	0.217	0.002	0.156	0.152
Seq Strat	-1.327	0.308	0.234	0.138	0.209	0.155
MSM	-1.123	0.360	0.304	0.042	0.312	0.255
MSM (obsPSA)	-1.059	0.349	0.299	0.122	0.297	0.246
			$\tau = 1$	0.05		
Two-stage	-1.272	0.216	0.196	0.088	0.162	0.145
Seq Strat	-0.981	0.281	0.206	0.386	0.214	0.155
MSM	-0.861	0.327	0.279	0.080	0.312	0.254
MSM(obsPSA)	797.0-	0.335	0.281	0.156	0.293	0.248

Table 4

Impact of Model Misspecification

		$\gamma = -1.5$			$\gamma = 0$	
Method	Mean Est	SD Est	Mean SE	Mean Est	SD Est	Mean SE
	I	Misspecifie	ed treatment e	ffect:Heterog	ceneity in γ	
	$y_i \sim t$) (−2.25, –1	0.75)	$\gamma_i \sim \gamma_i$	U (-0.75, C	.75)
Two-stage	-1.454	0.228	0.226	0.011	0.154	0.155
Seq Strat	-1.388	0.321	0.244	0.086	0.209	0.157
MSM	-1.199	0.391	0.317	0.052	0.330	0.257
		Misspec	ified SADT N	Aodel: Includ	ing Age	
Two-stage	-1.480	0.223	0.211	-0.005	0.148	0.141
Seq Strat	-1.432	0.270	0.215	0.051	0.190	0.136
MSM	-1.246	0.283	0.246	0.005	0.220	0.190
			Random (Censoring		
Two-stage	-1.526	0.291	0.279	-0.029	0.192	0.186
Seq Strat	-1.475	0.415	0.318	0.056	0.265	0.193
MSM	-1.281	0.449	0.369	0.037	0.334	0.281
			Age Depende	nt Censoring		
Two-stage	-1.479	0.269	0.259	0.001	0.175	0.169
Seq Strat	-1.446	0.354	0.277	0.056	0.228	0.169
MSM	-1.283	0.311	0.291	-0.004	0.221	0.214

Table 5

			5	i i	;
Conditions for Treatment	Time of Randomization	Comparison	Mean Est	SD Est	Mean SE
	Randomized Trial A: S	imple Randomization at Baseline			
At risk at:					
baseline	baseline	No Treatment	-1.104	0.091	0.085
3 years	baseline	No Treatment	-0.869	0.085	0.081
6 years	baseline	No Treatment	-0.443	0.074	0.073
	Randomized Trial B: S	ADT by Indication v's no SADT			
Pos. slope & PSA above:	First time PSA above:				
1	1	No Treatment	-1.332	0.073	0.072
2	2	No Treatment	-1.374	0.071	0.073
ß	ß	No Treatment	-1.397	0.081	0.075
	Randomized Tri	al C: Early v's Late SADT			
Pos. slope & PSA above:	First time PSA above:	Treatment when PSA above:			
1	1	2	-0.340	0.081	0.080
1	1	c	-0.689	0.076	0.077
2	2	3	-0.520	0.076	0.078
2	2	4	-0.825	0.073	0.076
	Randomized Trial D: Ir	nmediate v's SADT by Indicatior			
At risk at:		Treatment when PSA above:			
baseline	baseline	1	-0.103	0.103	0.100
baseline	baseline	2	-0.374	0.099	0.095
baseline	baseline	3	-0.644	0.097	0.091