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Complete List of Authors	Xiang Li Quefeng Li Donglin Zeng Karen Marder Jane Paulsen and Yuanjia Wang
Corresponding Author	Quefeng Li
E-mail	quefeng@email.unc.edu
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Time-varying Hazards Model for Incorporating Irregularly Measured, High-Dimensional Biomarkers

Xiang Li¹, Qiefeng Li², Donglin Zeng³,
Karen Marder⁴, Jane Paulsen⁵, and Yuanjia Wang⁶

^{1,4,6} Columbia University

^{2,3} University of North Carolina, Chapel Hill

⁵ University of Iowa

Abstract

Clinical studies with time-to-event outcomes often collect measurements of a large number of time-varying covariates over time (e.g., clinical assessments or neuroimaging biomarkers) to build time-sensitive prognostic model. An emerging challenge is that due to resource-intensive or invasive (e.g., lumbar puncture) data collection process, biomarkers may be measured infrequently and thus not available at every observed event time point. Leveraging all available, infrequently measured time-varying biomarkers to improve prognostic model of event occurrence is an important and challenging problem. In this paper, we pro-

pose a kernel-smoothing based approach to borrow information across subjects to remedy infrequent and unbalanced biomarker measurements under a time-varying hazards model. A penalized pseudo-likelihood function is proposed for estimation, and an efficient augmented penalization minimization algorithm related to the alternating direction method of multipliers (ADMM) is adopted for computation. Under some regularity conditions to carefully control approximation bias and stochastic variability, we show that even in the presence of ultra-high dimensionality, the proposed method selects important biomarkers with high probability. Through extensive simulation studies, we demonstrate superior performance in terms of estimation and selection performance compared to alternative methods. Finally, we apply the proposed method to analyze a recently completed real world study to model time to disease conversion using longitudinal, whole brain structural magnetic resonance imaging (MRI) biomarkers, and show a substantial improvement in performance over current standards including using baseline measures only.

Keywords: Biomarker studies; High-dimensional covariates; Irregular measurements; Kernel-weighted estimation; Neurological disorders; Time-varying hazards model.

1 Introduction

Time-varying biomarkers are often collected to assist studying disease mechanism and building time-varying prognostic models for time-to-event outcomes such as disease onset. With rapid advancements in technologies, high-dimensional time-varying biomarkers are repeatedly measured over time for each individual. While potentially useful to improve the power of prediction, there are also emerging challenges, statistically and computationally. First, since collecting certain biomarkers may be resource-intensive or invasive (e.g., neuroimaging measures involving radiation exposure) the measurements are infrequent and irregularly time-spaced for each subject. Thus, the biomarkers as covariates may not be available at every observed event time point. Second, there is an extensive body of literature (Desikan et al., 2006; Chen et al., 2014; Paulsen et al., 2014a; Ryan et al., 2015) suggesting that biomarker effects on neurological disorders vary with age, time, or an individual's disease progression. Specifically, a large natural history study of a neurological disorder (Paulsen et al., 2014a), Huntington's disease (HD), showed that regional brain atrophy measures, considered as important HD biomarkers, manifest differential rates of decline at distinct disease stages; another work on neurobiological processes revealed an age-dependent effect pattern for brain activation as measured by neuroimaging biomarkers (Ryan et al., 2015). Third, biomarkers often exhibit some biological network structure (e.g., structural covariation network, He et al. (2008); structural and functional brain networks, Bullmore and Bassett (2011); gene co-expression network, Stuart et al. (2003)), where linked biomarkers may indicate similar likelihood of disease diagnosis or prognosis due to sharing disease pathways. Given all these challenges, a valid statistical method to include the biomarkers for prediction should take into account all available, non-frequent biomarker measurements, time-varying ef-

fects, high dimensionality, and informative network structure among covariates.

Usual methods associating time-dependent biomarkers with hazards of time-to-event outcomes requires biomarkers to be completely observed at each event time (Honda and Härdle, 2014; Honda and Yabe, 2017), so they cannot be applied when the biomarkers are infrequently and irregularly measured. An easy solution is to treat this issue as a missing covariates problem, and impute missing biomarkers at an event time using last value carried forward (LVCF; Andersen and Liestol, 2003). While straightforward to carry out, this approach may induce bias and lead to incorrect inference especially when the biomarkers show a substantial change (Prentice, 1982; Tsiatis and Davidian, 2001). Several sophisticated approaches (Tsiatis and Davidian, 2004; Gould et al., 2014; Taylor et al., 2013) are proposed as alternatives, where they link group-average biomarker trajectories with time-to-event outcome models instead of using individual trajectories. However, predictive performance of models obtained from these approaches may not reflect the true predictivity of the actual biomarker measures collected on each individual. Other methods link unobserved random effects to time-to-event models through a measurement error model and joint modeling (e.g., Rizopoulos, 2011). But, measurement error models are not practically relevant when biomarker variation is due to true biological variability instead of random errors. Thus, it is desirable to directly associate observed biomarker values on each individual (instead of group-average or random effects) with the event times. Several recent work along this line includes Cao et al. (2014, 2015). However, they neither handle time-varying effects nor deal with high-dimensional biomarkers.

In addition, the estimation of high-dimensional, time-dependent effect profile functions for biomarkers on event outcomes with limited data is an ambitious goal. Thus, incorporating biological information is crucial to reduce model space complexity and stabilize the estimation.

Strong biological evidence observed for neurodegenerative disorders include: (1) signals are clustered in networks (He et al., 2008; Eidelberg and Surmeier, 2011; Parikshak et al., 2015); (2) biomarker signals evolve with disease progression and/or age (Desikan et al., 2006; Chen et al., 2014; Paulsen et al., 2014a); (3) signals are expected to be sparse (e.g., Liu et al., 2014). Existing methods on selecting functions for hazards models (e.g., Yan and Huang, 2012; Liu et al., 2013) or transformation models (Liu and Zeng, 2013) do not simultaneously handle irregularly measured time-dependent covariates and incorporate biological information. In our motivating study and other applications, subject's biomarker assessments were less frequently scheduled than clinical visits and thus did not necessarily coincide, rendering existing work referenced above non-applicable.

In this article, we propose a unified method to estimate time-varying effects in a hazards model using high-dimensional time-varying biomarkers measured irregularly. Our first contribution is to handle the complication of unavailable biomarkers at some event times without excluding subjects with missing biomarkers. To this end, we adopt local kernel smoothing in order to pool observations across event times and subjects. Secondly, to facilitate selection of the entire profile function, after approximating each function using B-splines, we incorporate a group sparsity penalty on the spline coefficients, inspired by the work of Huang et al. (2010). Furthermore, to incorporate the available biological network structure among the biomarkers, we include an additional regularization to encourage the strongly linked biomarkers to yield similar prognosis effects. It is challenging to control selection and estimation accuracy when dealing with high-dimensional functions because estimation noise at any given time point may cause a biomarker to enter the model. Thus, our third contribution is to propose an efficient computational algorithm to achieve ℓ_0 -penalty like sparsity of the functions by modifying the popular augmented

penalization methods, including the alternating direction method of multipliers (ADMM; Boyd et al. (2011)) algorithm. Fourthly, our examination of theoretical properties involves establishing high-dimensional oracle selection for functions (instead of scalar parameters) in the presence of kernel approximation, which requires techniques to appropriately control for approximation bias and stochastic variability that is not available in the existing theories.

The remainder of the paper is organized as follows. In Section 2, we describe a time-varying hazards model with time-varying biomarkers, an approximated likelihood function to borrow information, and an efficient algorithm for implementation. In Section 3, we provide theories showing that, under a general class of penalty functions, our method admits the “oracle property” in terms of selecting and estimating the true time-dependent effects. In Section 4, we extend the proposed algorithm to further incorporate biological network structure in the model regularization. In Section 5, we present extensive simulation studies to examine finite-sample performance of the proposed method and show much improved performance compared to alternative approaches. In Section 6, we apply our method to a recently completed real world study (Paulsen et al., 2014b), where the whole brain structural magnetic resonance imaging (MRI) data are used to estimate a network-regularized biomarker signature for predicting time-to-onset of HD. We show that the predictive performance of the proposed model substantially outperform LVCF, baseline-only analyses, and current standards developed independently in recent literature (Long et al., 2016). Finally, we conclude the article with some discussions in Section 7.

2 Methodologies

2.1 Model and estimation

Assume that data are collected from n i.i.d subjects. For subject i , let $\mathbf{X}_i(t)$ denote a p_n -dimensional vector of covariates including time-dependent biomarkers and let T_i denote time-to-event of interest (e.g., age-at-onset of a disease). To model the time-varying effects of covariates on the event, we propose a time-varying hazards model by assuming that the conditional hazard rate function of $T_i = t$ given the covariate history by time t is

$$\lambda(t|\mathbf{X}_i(s), s \leq t) = \lambda_0(t) \exp \{ \boldsymbol{\beta}^T(t) \mathbf{X}_i(t) \}, \quad (1)$$

where $\lambda_0(t)$ is an unspecified baseline hazard function, and $\boldsymbol{\beta}(t)$ is a vector of covariate effects at time t . Note that (1) can also include additional components of the covariate history, for instance, lagging effects, by expanding $\mathbf{X}_i(t)$ to include the covariate history.

Assume that $\mathbf{X}_i(t)$ is only measured at n_i discrete time points: t_{i1}, \dots, t_{in_i} . The observed data consist of $\{ \tilde{T}_i = \min(T_i, C_i), \Delta_i = I(T_i \leq C_i), \mathbf{X}_i(t_{i1}), \dots, \mathbf{X}_i(t_{in_i}) \}$, $i = 1, \dots, n$, where C_i denotes the censoring time assumed to be conditionally independent of T_i given $\mathbf{X}_i(t)$, \tilde{T}_i is the observed event time or censoring time, and Δ_i the censoring indicator. Furthermore, let $Y_i(t) = I(\tilde{T}_i \geq t)$ and $N_i(t) = I(\tilde{T}_i \leq t, \Delta_i = 1)$ denote the at-risk process and the observed counting process, respectively.

If the complete history of the covariate processes $\mathbf{X}_i(t)$ were observed for all $t < \tilde{T}_i$ and all

i , then the classic log-partial likelihood (Fleming and Harrington, 2011), which is defined as

$$n^{-1} \sum_{i=1}^n \int_0^{\tau} \left(\boldsymbol{\beta}^T(t) \mathbf{X}_i(t) - \log \left[n^{-1} \sum_{j=1}^n Y_j(t) \exp\{\boldsymbol{\beta}^T(t) \mathbf{X}_j(t)\} \right] \right) dN_i(t), \quad (2)$$

could be maximized to estimate all the coefficients, where τ is the duration of study. Since $\mathbf{X}_i(t)$ are only observable at some distinct time points, we need to approximate each term in the above log-partial likelihood function using the observed data. Note that the objective function (2) relies on some empirical average of functionals of $\mathbf{X}_i(t)$'s. Thus, the approximation does not need to be accurate for each subject's $\mathbf{X}_i(t)$; instead, an accurate approximation to this empirical average of $\mathbf{X}_i(t)$'s is sufficient. This motivates one to adopt kernel smoothing by pooling observations not only from the same trajectory on subject i , but also from the other subjects when approximating $\mathbf{X}_i(t)$. Specifically, consider kernel smoothing in Andersen and Liestol (2003) and weight subjects differently in the pooled data, where weights are based on the distance between the observed measurement times and t ; the resulting approximated objective function is given as

$$l_n^s(\boldsymbol{\beta}) = n^{-1} \sum_{i=1}^n \int_0^{\tau} \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) \left(\boldsymbol{\beta}^T(t) \mathbf{X}_i(t_{iv}) - \log \left[n^{-1} \sum_{j=1}^n \sum_{k=1}^{n_j} K_{h_n}(t - t_{jk}) Y_j(t) \exp\{\boldsymbol{\beta}^T(t) \mathbf{X}_j(t_{jk})\} \right] \right) dN_i(t),$$

where $K_{h_n}(t) = h_n^{-1} K(t/h_n)$ for some symmetric kernel function $K(\cdot)$ and a bandwidth h_n .

Since $\boldsymbol{\beta}(t)$'s are fully nonparametric, the maximization is not feasible. We further use B-spline approximation for each $\beta_j(t)$, $j = 1, \dots, p_n$. Specifically, let $\phi_{1,m}(t), \dots, \phi_{q_n,m}(t)$ be the B-spline basis functions of order m associated with $(q_n - m)$ equally-spaced interior knots $0 = t_0 < t_1 < \dots < t_{q_n-m} < t_{q_n-m+1} = \tau$, where τ is the study duration. The basis functions can

be generated by Cox-de Boor recursion formula $\phi_{\ell,0}(t) := 1$, if $t_\ell \leq t < t_{\ell+1}$ and $\phi_{\ell,0}(t) := 0$, otherwise. $\phi_{\ell,k}(t) := \frac{t-t_\ell}{t_{\ell+k-1}-t_\ell}\phi_{\ell,k-1}(t) + \frac{t_{\ell+k}-t}{t_{\ell+k}-t_{\ell+1}}\phi_{\ell+1,k-1}(t)$, for $1 \leq k \leq m$. To simplify notation, we write $\phi_{\ell,m}(t)$ as $\phi_\ell(t)$ in the following. Define $\phi(t) = (\phi_1(t), \dots, \phi_{q_n}(t))^T$. Then, for the j th component of $\beta(t)$, we approximate $\beta_j(t)$ by

$$\beta_j(t) \approx \gamma_j^T \phi(t), \quad j = 1, \dots, p_n, \quad (3)$$

where $\gamma_j = (\gamma_{j1}, \dots, \gamma_{jq_n})^T$ is a coefficient vector for the B-spline approximation. Consequently, define $\mathbf{Z}_i(t, u) = \mathbf{X}_i(t) \otimes \phi(u)$, where \otimes is the Kronecker product, and $\gamma = (\gamma_1^T, \dots, \gamma_{p_n}^T)^T$, we propose to maximize

$$\begin{aligned} l_n(\gamma) &= n^{-1} \sum_{i=1}^n \int \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) \left(\gamma^T \mathbf{Z}_i(t_{iv}, t) - \right. \\ &\quad \left. \log \left[n^{-1} \sum_{j=1}^n \sum_{k=1}^{n_j} I(\tilde{T}_j \geq t) K_{h_n}(t - t_{jk}) \exp \left\{ \gamma^T \mathbf{Z}_j(t_{jk}, t) \right\} \right] \right) dN_i(t) \\ &= n^{-1} \sum_{i=1}^n \sum_{v=1}^{n_i} \Delta_i K_{h_n}(\tilde{T}_i - t_{iv}) \left(\gamma^T \mathbf{Z}_i(t_{iv}, \tilde{T}_i) - \right. \\ &\quad \left. \log \left[n^{-1} \sum_{j=1}^n \sum_{k=1}^{n_j} I(\tilde{T}_j \geq \tilde{T}_i) K_{h_n}(\tilde{T}_i - t_{jk}) \exp \left\{ \gamma^T \mathbf{Z}_j(t_{jk}, \tilde{T}_i) \right\} \right] \right) \end{aligned} \quad (4)$$

to estimate γ and thus $\beta(t)$'s.

2.2 Sparsity regularization

With a large number of biomarkers, directly maximizing $l_n(\gamma)$ may lead to high variability of the time-varying effects and may even be infeasible. Furthermore, since most of the biomarkers are expected to be non-informative of disease prognosis, it is important to identify which ones

contribute to the underlying biological mechanism. Thus, we impose regularization both for the purpose of stabilizing computation and variable selection. Specifically, we propose to minimize the following penalized function:

$$\hat{\gamma} = \arg \min_{\gamma} \{-l_n(\gamma) + p(\gamma; \nu_n)\}, \quad (5)$$

where $p(\gamma; \nu_n)$ is a pre-specified penalty function with a tuning parameter ν_n .

Since we aim to select important $\beta_j(t)$'s as functions in $[0, \tau]$, or equivalently, γ_j as a vector, the penalty is imposed on the Euclidean norm of γ_j instead of each single component of γ_j . Furthermore, to encourage oracle selection, following Zhang and Zhang (2012), one may wish to choose a concave penalty function. Therefore, in this paper, we choose the penalty term $p(\gamma; \nu_n) = \nu_n \sum_{j=1}^{p_n} \sqrt{q_n} \rho(\|\gamma_j\|_2)$, where $\rho(\cdot)$ is a general penalty function imposed on $\|\gamma_j\|_2$, the Euclidean norm of γ_j . For example, $\rho(t) = t$ gives the LASSO penalty,

$$\rho(t) = \int_0^t I(u \leq \nu_n) + \frac{(a\nu_n - u)_+}{(a-1)\nu_n} I(u \geq \nu_n) du$$

gives the SCAD penalty (Fan and Li, 2001), and

$$\rho(t) = \int_0^t \frac{(a\nu_n - u)_+}{a\nu_n} du$$

yields the MCP penalty (Zhang, 2010). As an extreme case, we can choose $p(\gamma; \nu_n)$ to be the ℓ_0 -penalty, which is defined as $\nu_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\gamma_j\|_{G_0}$ with $\|\gamma_j\|_{G_0} = I(\|\gamma_j\|_2 \neq 0)$. A concave penalty may lead to non-convex minimization; however, in the next section, we will describe a unified algorithm to facilitate computation.

2.3 Computational algorithm

We propose a unified computational algorithm for the optimization (5). The algorithm is motivated by a class of proximal methods performing augmentation and splitting, including the ADMM algorithm (Boyd et al., 2011). Specifically, additional slack variables θ of the same dimension as the target variables γ are introduced to facilitate efficient computation and scale up.

Following Li et al. (2017), we first approximate (5) by the following constrained optimization problem

$$\arg \min_{\gamma, \theta} -l_n(\gamma) + p(\theta; \nu_n) \quad \text{subject to} \quad \sum_{j=1}^{p_n} \|\gamma_j - \theta_j\|_2 \leq c_n, \quad (6)$$

where c_n is some constant controlling the difference between γ and θ . Note that the ADMM algorithm is a special case when setting $c_n = 0$ in (6). We further propose to solve the equivalent

Lagrangian problem

$$\arg \min_{\gamma, \theta} -l_n(\gamma) + p(\theta; \nu_n) + \phi_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\gamma_j - \theta_j\|_2, \quad (7)$$

where ϕ_n is the Lagrangian multiplier. We delegate the proof of the equivalence between (6) and (7) to the supplementary material. Minimizing (7) for given ν_n and ϕ_n thus can be carried out by iteratively updating all parameters using the following algorithm: at the k th iteration,

$$\gamma^{k+1} = \arg \min_{\gamma} -l_n(\gamma) + \phi_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\gamma_j - \theta_j^k\|_2, \quad (8)$$

$$\theta^{k+1} = \arg \min_{\theta} p(\theta; \nu_n) + \phi_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\gamma_j^{k+1} - \theta_j\|_2. \quad (9)$$

Note that the above update (8) is similar to updating a regularized regression with group LASSO,

where the objective function is convex. For (9), when $p(\boldsymbol{\theta}; \nu_n) = \nu_n \sum_{j=1}^{p_n} \sqrt{q_n} \rho(\|\boldsymbol{\gamma}_j\|_2)$, the minimization can be performed group-wise which often results in explicit solution. Therefore, each iteration of the proposed algorithm only involves one step of convex minimization and one step of simple calculation. The tuning parameters ν_n and ϕ_n can be chosen using the likelihood-based cross-validation.

As an example, when $p(\boldsymbol{\gamma}; \nu_n)$ is chosen to be the ℓ_0 -penalty, the second step in each iteration of the above algorithm becomes

$$\boldsymbol{\theta}^{k+1} = \arg \min_{\boldsymbol{\theta}} \nu_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\boldsymbol{\theta}_j\|_{G_0} + \phi_n \sum_{j=1}^{p_n} \sqrt{q_n} \|\boldsymbol{\gamma}_j^{k+1} - \boldsymbol{\theta}_j\|_2.$$

It can be seen from Section 4 that the group-wise ℓ_0 -penalty acts as hard-thresholding the estimates obtained from the first step. Particularly, simple algebra gives for $j = 1, \dots, p_n$,

$$\boldsymbol{\theta}_j^{k+1} = \boldsymbol{\gamma}_j^{k+1} I(\|\boldsymbol{\gamma}_j^{k+1}\|_2 > \nu_n / \phi_n). \quad (10)$$

As a remark, the challenge in selecting informative biomarkers from a large candidate pool arises from that each component of the effect profiles $\boldsymbol{\beta}(t)$ is a function. Estimation noise on the entire range of t needs to be controlled and penalization needs to be imposed on its norm. Furthermore, when the dimension of $\mathbf{X}_i(t)$ is high, computation in (8) can still be intensive even if it is a convex minimization. In Section 4, we will suggest a coordinate-descent approach by first approximating $l_n(\boldsymbol{\gamma})$ by a summation of quadratic functions for each $\boldsymbol{\gamma}_j$. Thus, computation can easily scale up to high-dimensional scenarios.

3 Theoretical Properties

The main challenge in proving theoretical properties is to appropriately control for the approximation bias resulting from local kernel smoothing in the approximated likelihood (3) and stochastic variability. Let $\beta^*(t)$ denote the true value of $\beta(t)$ and let $\beta_j^*(t)$ denote its j -th element. We provide a non-asymptotic result showing that, with large probability, the nonzero $\beta_j^*(t)$ can be correctly selected and consistently estimated by $\hat{\beta}_j(t)$, where $\hat{\beta}_j(t) := \hat{\gamma}_j^T \phi(t)$ and $\hat{\gamma} = (\hat{\gamma}_1^T, \dots, \hat{\gamma}_{p_n}^T)^T$ is the estimator given by (5). In particular, we allow p_n and q_n to diverge to infinity and ν_n to converge to zero with n .

3.1 Technical notations

Before presenting regularity conditions, we need the following notations. Let

$$S_n^{(l)}(\gamma, t) = n^{-1} \sum_{i=1}^n \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) Y_i(t) \{Z_i(t_{iv}, t)\}^{\otimes l} \exp\{\gamma^T Z_i(t_{iv}, t)\}, \quad l = 0, 1, 2. \quad (11)$$

Then, the approximated log-partial likelihood for optimization can be rewritten as

$$l_n(\gamma) = \frac{1}{n} \sum_{i=1}^n \int_0^\tau \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) [\gamma^T Z_i(t_{iv}, t) - \log\{S_n^{(0)}(\gamma, t)\}] dN_i(t).$$

Denote $\mathbf{E}_n(\gamma, t) = S_n^{(1)}(\gamma, t)/S_n^{(0)}(\gamma, t)$. Then, the gradient vector, denoted by $\mathbf{U}_n(\gamma)$, and the negative Hessian matrix, denoted by $\mathbf{I}_n(\gamma)$, of $l_n(\gamma)$ are given by

$$\mathbf{U}_n(\gamma) = \frac{1}{n} \sum_{i=1}^n \int_0^\tau \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) \{Z_i(t_{iv}, t) - \mathbf{E}_n(\gamma, t)\} dN_i(t),$$

$$\mathbf{I}_n(\boldsymbol{\gamma}) = \frac{1}{n} \sum_{i=1}^n \int_0^\tau \sum_{v=1}^{n_i} K_{h_n}(t - t_{iv}) \left\{ \frac{S_n^{(2)}(\boldsymbol{\gamma}, t)}{S_n^{(0)}(\boldsymbol{\gamma}, t)} - \mathbf{E}_n(\boldsymbol{\gamma}, t)^{\otimes 2} \right\} dN_i(t).$$

In addition, we define

$$\bar{s}^{(l)}(\boldsymbol{\gamma}, t) = \mathbb{E}[Y(t)\{\mathbf{Z}(t, t)\}^{\otimes l} \exp\{\boldsymbol{\gamma}^T \mathbf{Z}(t, t)\}], \text{ for } l = 0, 1, 2;$$

$\mathbf{e}(\boldsymbol{\gamma}, t) = \bar{s}^{(1)}(\boldsymbol{\gamma}, t)/\bar{s}^{(0)}(\boldsymbol{\gamma}, t)$ and

$$\boldsymbol{\Sigma}(\boldsymbol{\gamma}, t) = \int_0^\tau \left\{ \frac{\bar{s}^{(2)}(\boldsymbol{\gamma}, t)}{\bar{s}^{(0)}(\boldsymbol{\gamma}, t)} - \mathbf{e}(\boldsymbol{\gamma}, t)^{\otimes 2} \right\} \bar{s}^{(0)}(\boldsymbol{\gamma}, t) \lambda_v(t) d\Lambda_0(t),$$

where we assume $\{t_{i1}, \dots, t_{in_i}\}$ follow an independent counting process with intensity function $\lambda_v(t)$.

In the penalized partial likelihood (5), we assume $\rho(t)$ belongs to a general class of folded-concave functions as discussed in Fan and Lv (2011). Such a class of functions will be characterized by Condition 11 in Section 3.2. In particular, denote $\kappa(\rho, \mathbf{u})$ as the ‘‘local concavity’’ of $\rho(\cdot)$ at a general vector $\mathbf{u} = (u_1, \dots, u_s)^T \in \mathcal{R}^s$ that

$$\kappa(\rho, \mathbf{u}) = \lim_{\varepsilon \rightarrow 0^+} \max_{1 \leq j \leq s} \sup_{t_1 < t_2 \in (|u_j| - \varepsilon, |u_j| + \varepsilon)} - \frac{\rho'(t_2) - \rho'(t_1)}{t_2 - t_1}.$$

For example, for the LASSO penalty, $\kappa(\rho, \mathbf{u}) = 0$, while for the SCAD penalty,

$$\kappa(\rho, \mathbf{u}) = \begin{cases} (a - 1)^{-1} \nu_n^{-1}, & \text{if there exists a } u_j \text{ such that } \nu_n \leq |u_j| \leq a\nu_n; \\ 0, & \text{otherwise.} \end{cases}$$

Our subsequent regularity condition for the penalty function will be written in terms of $\kappa(\rho, \mathbf{u})$.

We define additional notations. For a vector \mathbf{a} , let $\|\mathbf{a}\|_\infty = \max_j |a_j|$ denote its sup-norm. For a matrix \mathbf{A} , let $\|\mathbf{A}\|_\infty = \max_i \sum_j |a_{ij}|$ denote its matrix sup-norm, where a_{ij} is the (i, j) -th element of \mathbf{A} . Let $\lambda_{\min}(\mathbf{A})$ and $\lambda_{\max}(\mathbf{A})$ be the minimal and maximal eigenvalues of \mathbf{A} , respectively.

3.2 Regularity conditions

We define the unique projection of $\beta_j^*(t)$ on the sieve space consisting of $\phi(t)$ as $\gamma_j^{*T} \phi(t)$, where γ_j^* is a q_n -dimensional vector. Denote $\mathcal{M} = \{j : \beta_j^*(t) \neq 0, t \in [0, \tau]\}$ as the set of active $\beta_j^*(t)$'s. We assume that when q_n is large enough, \mathcal{M} is equivalent to the support of γ^* , i.e. $\mathcal{M} = \{j : \gamma_j^* \neq 0\}$. In other words, the important covariates with $\beta_j^*(t) \neq 0$ is fully characterized by the non-zero γ^* 's vectors, when we choose sufficient number of spline bases. Let $r_n = |\mathcal{M}|$ denote its cardinality. Denote $\mathcal{A} = \{j_l : j \in \mathcal{M} \text{ and } 1 \leq l \leq q_n\}$. Note that, $|\mathcal{A}| = r_n q_n$. Denote $d_n = \min_{j \in \mathcal{M}} \|\gamma_j^*\|_2$ as the minimal signal strength. For a set S , denote \mathbf{a}_S as the subvector of \mathbf{a} with indices in S and \mathbf{A}_{SS} as the submatrix with row and column indices in S . Let $\mathcal{B}_0 := \{\gamma \in \mathcal{R}^{p_n q_n} : \|\gamma_{\mathcal{A}} - \gamma_{\mathcal{A}}^*\|_\infty \leq d_n \text{ and } \gamma_{\mathcal{A}^c} = \mathbf{0}\}$. We assume that M is a universal positive constant and the following conditions hold.

Condition 1. $\Lambda_0(\tau) = \int_0^\tau \lambda_0(t) dt < \infty$ and $P\{C \geq \tau\} > 0$.

Condition 2. $\sup_{t \in [0, \tau]} |X_j(t)| \leq M$ for all $1 \leq j \leq p$; $\sup_{t \in [0, \tau]} |(\beta^*(t))^T \mathbf{X}(t)| \leq M$.

Condition 3. $\lambda_v(t)$ is bounded and twice continuously differentiable with bounded second derivative.

Condition 4. The kernel function $K(x)$ is symmetric and has finite second moment.

Condition 5. $|E[X_j(t)]| \leq M$ and $E[X_j(t)]$ is twice continuously differentiable and $|(E[X_j(t)])''| \leq M$. In addition, $E[X_j(t) - X_j(s)]^2 \leq M(t - s)^2$ holds almost everywhere for $t, s \in [0, \tau]$.

Condition 6. There exists some positive constant α such that $\sup_{t \in [0, \tau]} |\beta_j^*(t) - (\gamma_j^*)^T \phi(t)| \leq c_\alpha q_n^{-\alpha}$ for all $1 \leq j \leq p_n$, where c_α is some positive constant.

Condition 7. $(nh_n)^{-1/2} = O(1)$.

Condition 8. $r_n q_n c_n d_n^{1/2} = o(n)$, where $c_n := r_n q_n^2 h_n^{-1} \vee h_n^{-2}$.

Condition 9. $\sup_{\gamma \in \mathcal{B}_0} \|\Sigma_{\mathcal{A}\mathcal{A}}(\gamma)^{-1}\|_\infty \leq M$.

Condition 10. $\sup_{\gamma \in \mathcal{B}_0} \|\Sigma_{\mathcal{A}^c\mathcal{A}}(\gamma)\Sigma_{\mathcal{A}\mathcal{A}}(\gamma)^{-1}\|_\infty \leq (1 - \zeta)\rho'(0+)/\rho'(d_n/2)$ for some $\zeta \in (0, 1)$.

Condition 11. $\rho(t)$ is increasing and concave in $t \in [0, \infty)$ and has a continuous derivative $\rho'(t)$ with $0 < \rho'(0+) < M$. $\sup_{\gamma \in \mathcal{B}_0} |\nu_n \kappa(\rho, \gamma)| \leq M$.

Condition 1 is a common condition for Cox model. Condition 2 is used to establish the exponential type of concentration inequalities for the gradient vector. Condition 3 is a smoothness condition on $\lambda_v(t)$. Condition 4 is imposed on the kernel function and it is satisfied for common kernel functions such as Gaussian kernel and Epanechnikov kernel. Condition 5 is a smoothing assumption on $E[X_j(t)]$. Since this condition is only imposed on the population average, it allows the individual realization of $X_{ij}(t)$ to be non-smooth or even discontinuous. For example, the trajectory of $X_{ij}(t)$ may have a discontinuity point which is from a continuous distribution in $[0, \tau]$. Condition 6 requires $\beta_j^*(t)$ to be sufficiently smooth so that it can be well approximated by splines. If $\beta_j^*(t)$ has a bounded k -th derivatives, then $\alpha = k - 1$ (See Schumaker (2007)). Condition 9 and Condition 10 are imposed on the population information matrix $\Sigma(\gamma)$. Similar

conditions also appear in the derivation of oracle properties for parametric models (Fan and Lv, 2011). Condition 10 is an irrepresentable-type of condition. If some concave penalties (e.g., SCAD or MCP) are used, the upper bound in Condition 10 is allowed to diverge to infinity at a polynomial rate of n (Fan and Lv, 2011). If the ℓ_1 -penalty is used, the upper bound reduces to $1 - \zeta$, which is exactly the irrepresentable condition needed for LASSO (Zhao and Yu, 2006). Under Conditions 9 and 10, similar results for the sample information matrix $I_n(\gamma)$ can be shown to hold with high probability (See Lemma S5 in the Supplementary Material). Condition 11 is imposed on the penalty function, which is satisfied by commonly used penalty functions, such as LASSO, SCAD, and MCP.

3.3 Main results

We first give a concentration inequality for the gradient vector $U_n(\gamma^*)$, which will play a key role in establishing the main result.

Lemma 1. *Under conditions 1 to 8, there exist positive constants C_1, C_2, C_3, C_4 and D such that for any $x > 0$ and $\epsilon > 0$, it holds with probability no less than $1 - \epsilon - C_1 \exp(-C_2 n h_n^6 x^2) - C_3 \exp(-C_4 x)$ that*

$$|U_{n,j}(\gamma^*)| \leq D\{(n h_n^2)^{-1/2} x + \pi_n\}.$$

where $U_{n,j}(\gamma^*)$ is the j -th element of $U_n(\gamma^*)$ and $\pi_n = (r_n q_n c_n d_n^{1/2} / n)^{1/2} + h_n^2 + r_n q_n^{-\alpha}$.

Remark 1. *Different from the existing study of ordinary Cox model in the high dimensional setting (Bradic et al., 2011), the expectation of the gradient vector $U_n(\gamma^*)$ no longer equals to zero, due to the spline approximation to $\beta^*(t)$ and the local smoothing. The term π_n quantifies such a bias. In the expression of π_n , h_n^2 is due to the local smoothing, $r_n q_n^{-\alpha}$ is due to the*

approximation of $(\gamma_j^*)^T \phi(t)$ to $\beta_j^*(t)$, and $(r_n q_n c_n d_n^{1/2} / n)^{1/2}$ is introduced by $S_n^{(l)}(\gamma, t)$. However, due to the conditions of p_n, q_n and d_n , the bias π_n vanishes as sample size n grows so it will not affect variable selection as given in Theorem 1 below.

Remark 2. Under condition 3, $n_i = O_P(1)$, i.e. for any $\epsilon > 0$, there is a constant M_ϵ such that $P(n_i > M_\epsilon) \leq \epsilon$. Such an ϵ appears in Lemma 1. All other probabilities are calculated conditioning on the event $\{n_i \leq M_\epsilon\}$. We also condition on such an event when calculating the exception probabilities in Theorem 1 and additional lemmas in the supplementary material.

Our main theoretical result considers the variable selection property of our method. Specifically, we show that estimator $\hat{\beta}(t) = (\hat{\beta}_1(t), \dots, \hat{\beta}_p(t))^T$ possesses the “weak oracle property” as discussed in Fan and Lv (2011) for generalized linear model; that is, it is both variable selection consistent and consistently estimates the nonzero components of $\beta^*(t)$, i.e. the component functions which are not identically equal to zero.

Theorem 1. Suppose conditions 1 to 11 hold, and

$$\frac{n^2 h_n^8 (\nu_n \sqrt{q_n} - \pi_n)^2}{\log(p_n q_n)} \rightarrow \infty, \quad \frac{(n h_n^2)^{1/2} (\nu_n \sqrt{q_n} - \pi_n)}{\log(p_n q_n)} \rightarrow \infty, \quad (12)$$

$$\frac{n h_n^2 (r_n q_n)^{-1}}{\log(p_n r_n q_n^2)} \rightarrow \infty, \quad d_n > 2M \nu_n q_n. \quad (13)$$

There exist universal positive constants C_1, C_2, C_3, C_4, C_5 and C_6 such that, for any $\epsilon > 0$, with probability at least

$$1 - \epsilon - C_1 p_n q_n \exp\{-C_2 n^2 h_n^8 (\nu_n \sqrt{q_n} - \pi_n)^2\} \\ - C_3 p_n q_n \exp\{-C_4 (n h_n^2)^{1/2} (\nu_n \sqrt{q_n} - \pi_n)\} - C_5 p_n r_n q_n^2 \exp\{-C_6 n h_n^2 (r_n q_n)^{-1}\}, \quad (14)$$

there is a solution to (5) that yields $\hat{\beta}(t)$ satisfying

(a)(variable selection consistency): $\{j : \hat{\beta}_j(t) \neq 0\} = \{j : \beta_j^*(t) \neq 0\}$;

(b)(L_∞ error): $\max_{j \in \mathcal{M}} \sup_{0 \leq t \leq \tau} |\hat{\beta}_j(t) - \beta_j^*(t)| \leq M(\nu_n q_n^{3/2} + q_n^{-\alpha})$, where M is a positive constant.

The result in (a) guarantees the variable selection consistency with a high probability. The term $M\nu_n q_n^{3/2}$ in statement (b) gives the upper bound of $\max_{j_l \in \mathcal{A}} |\hat{\gamma}_{j_l} - \gamma_{j_l}^*|$, the other term $Mq_n^{-\alpha}$ corresponds to the approximation error of $\gamma_j^{*T} \phi(t)$ to $\beta_j^*(t)$. The first assumption in (13) gives the constraint on the divergence rates of p_n , r_n and q_n . They are allowed to diverge as long as $r_n q_n \log(p_n r_n q_n^2) = o(nh_n^2)$. The minimal signal strength d_n is allowed to converge to zero, given that the second term in (13) holds.

The conditions in Theorem 1 also restrict the choice of tuning parameter ν_n in the penalty function and bandwidth h_n in the kernel smoothing. Specifically, by (12) and (13), the tuning parameter ν_n needs to satisfy

$$\pi_n q_n^{-1/2} \vee \frac{\log(p_n q_n)}{n^{1/2} h_n q_n^{1/2}} \vee \frac{\sqrt{\log(p_n q_n)}}{n h_n^4 q_n} \ll \nu_n < \frac{d_n}{2M q_n}.$$

Moreover, the lower bound for h_n is given by

$$h_n \gg \frac{\log(p_n r_n q_n^2)}{n(r_n q_n)^{-1}} \vee \frac{\{\log(p_n q_n)\}^2}{n q_n \nu_n^2} \vee \frac{\{\log(p_n q_n)\}^{1/4}}{\sqrt{n q_n \nu_n}}.$$

To give an example, suppose that r_n and d_n are fixed and independent of n and $\alpha \geq 2$. Then, for $p_n = \exp\{O(n^{1/8})\}$, once we choose $q_n \asymp n^{1/8}$, $h_n \asymp n^{-1/8}$ and $n^{-5/16} \ll \nu_n \ll n^{-3/16}$, all requirements in Theorem 1 are met so that the probability in (14) becomes arbitrarily close to

one and the upper bound in Theorem 1(b) converges to zero.

4 Extension to Incorporate Network Structure

In many applications, biomarkers such as structural/functional brain measures (He et al., 2008; Alexander-Bloch et al., 2013) and gene co-expression (Stuart et al., 2003) exhibit network structures and hence can be naturally described by a graph $G = (V, E, \mathcal{W})$, where V is the set of vertices that correspond to biomarkers, $E = \{j \sim k\}$ is the set of edges that indicate connected vertices, and $\mathcal{W} = \{w_{jk} > 0 : (j, k) \in E\}$ is the set of edge weights. Note that \mathcal{W} is a $p_n \times p_n$ matrix with zero diagonal entries, $w_{jj} = 0$, $j = 1, \dots, p_n$.

When two biomarkers are highly linked, i.e., w_{jk} is large, they are likely to be involved in similar disease pathways, so the corresponding β_j and β_k are similar. Our proposed method can be easily extended to incorporate such network information to encourage this pattern in the analysis. Specifically, we add to (5) a Laplacian quadratic penalty on the effect size via a ℓ_2 -norm vector $|\gamma|_G = (\|\gamma_1\|_2, \dots, \|\gamma_{p_n}\|_2)^T$ to encourage smoothness of the functions over the network graph. Such a penalty takes a form of $|\gamma|_G^T \mathcal{L}_n^* |\gamma|_G$, where \mathcal{L}_n^* is a positive semi-definite matrix associated with the graph G . The choice of \mathcal{L}_n^* can be $\mathcal{L}_n^* = \mathbf{D} - \mathcal{W}$, where $\mathbf{D} = \text{diag}(d_1, \dots, d_{p_n})$ with $d_j = \sum_{l:(j,l) \in E} w_{jl}$, or its normalized version given by $\mathcal{L}_n^* = \mathbf{I} - \mathbf{D}^{-1/2} \mathcal{W} \mathbf{D}^{-1/2}$. The former choice of \mathcal{L}_n^* yields the penalty term

$$\sum_{(j,l) \in E} w_{jl} (\|\gamma_j\|_2 - \|\gamma_l\|_2)^2, \quad (15)$$

while the latter gives

$$\sum_{(j,l) \in E} w_{jl} \left(\frac{\|\gamma_j\|_2}{\sqrt{d_j}} - \frac{\|\gamma_l\|_2}{\sqrt{d_l}} \right)^2.$$

With this additional penalty, the previous algorithm in Section 2.2 is still applicable. In particular, the first step of each iteration solves

$$\begin{aligned} \hat{\gamma} = \arg \min_{\gamma} \left\{ -l_n(\gamma) + \lambda_1 \sum_{j=1}^{p_n} \sqrt{q_n} \|\gamma_j - \theta_j\|_2 \right. \\ \left. + (\lambda_2/2) \sum_{(j,l) \in E} w_{jl} \left(\frac{\|\gamma_j\|_2}{\sqrt{d_j}} - \frac{\|\gamma_l\|_2}{\sqrt{d_l}} \right)^2 \right\}. \end{aligned} \quad (16)$$

When the dimension p_n is large, it is not straightforward to directly find the minimum of the objective function (16), so we propose a majorization-minimization (MM) approach (Lange, 2013), and employ a group-wise descent algorithm to solve (16). The minimization is achieved by cyclic descent over each group at a time, where we choose a target group γ_j to minimize and consider other group coefficients $\gamma_k = \hat{\gamma}_k$, $k \neq j$, as fixed from the previous iteration. The details are given as follows.

Denote by $\nabla_j l_n(\gamma_j)$ the gradient taken over γ_j . By using a second-order Taylor expansion on γ_j centered at a point $\tilde{\gamma}_j$ and replacing the Hessian matrix by a suitable matrix \mathbf{H} , we first majorize $-l_n(\gamma_j)$ by a surrogate function $M(\gamma_j|\tilde{\gamma}_j)$,

$$M(\gamma_j|\tilde{\gamma}_j) = - \left\{ l_n(\tilde{\gamma}_j) + (\gamma_j - \tilde{\gamma}_j)^T \nabla_j l_n(\tilde{\gamma}_j) + \frac{1}{2} (\gamma_j - \tilde{\gamma}_j)^T \mathbf{H} (\gamma_j - \tilde{\gamma}_j) \right\}$$

with $-l_n(\gamma_j) \leq M(\gamma_j|\tilde{\gamma}_j)$ and $-l_n(\tilde{\gamma}_j) = M(\tilde{\gamma}_j|\tilde{\gamma}_j)$. We further assume that the matrix \mathbf{H} has the form $-\mathbf{H} = t^{-1} \mathbf{I}_{q_n}$, where \mathbf{I}_{q_n} is an identity matrix with dimension q_n and t is sufficiently

small such that the quadratic term $(2t)^{-1}\|\gamma_j - \tilde{\gamma}_j\|_2^2$ dominates the negative Hessian matrix of $l_n(\gamma_j)$. Thus, in each gradient step to update $\hat{\gamma}_j$, we solve

$$\arg \min_{\gamma_j} \left\{ \frac{1}{2t} \|\gamma_j - (\tilde{\gamma}_j + t\nabla_j l_n(\tilde{\gamma}_j))\|_2^2 + \lambda_1 \sqrt{q_n} \|\gamma_j - \theta_j\|_2 + \frac{\lambda_2}{2} \sum_{l:(j,l) \in E} w_{jl} \left(\frac{\|\gamma_j\|_2}{\sqrt{d_j}} - \frac{\|\hat{\gamma}_l\|_2}{\sqrt{d_l}} \right)^2 \right\}, \quad (17)$$

which can be carried out easily using Newton-Raphson method. Note that a similar approach was also used in Meier et al. (2008) and Simon et al. (2013).

Finally, we propose to use K -fold cross-validation to simultaneously choose all the tuning parameters. For each fixed bandwidth, the cross-validation criterion is the summation of the change in the log-partial likelihood function after we leave one-fold out. To choose the bandwidth, we let $h_n = Cn^{-1/8}$, start from a small positive constant C , and increase by a step size until the increment of the cross-validation is below a pre-specified threshold. All the algorithms have been implemented in an R package available upon request.

5 Simulation Studies

In this section, we conducted extensive simulations to evaluate the performance of the proposed method. We set the study duration $\tau = 1$. First, we considered p_n time-dependent covariates, each with piecewise constant trajectories given by

$$X_{ij}(t) = \sum_{l=1}^{20} I\{(l-1)/20 \leq t < l/20\} Z_{ijl},$$

where $\{Z_{ijl} : l = 1, \dots, 20\}$ were from a multivariate normal distribution with mean 0 and covariance $\text{Cov}(Z_{ijl}, Z_{ijl'}) = e^{-|l-l'|/20}$, $l, l' = 1, \dots, 20$. We also imposed a network structure to these covariates by assuming that there were links only within each block of four consecutive covariates, $\{X_{i1}, X_{i2}, X_{i3}, X_{i4}\}$, $\{X_{i5}, X_{i6}, X_{i7}, X_{i8}\}$, and so on. Furthermore, within each block of 4 covariates, the edge weight for each linked pair was set to be 0.5. Next, conditional on all covariates, the survival time T_i was generated from model (1) with $\lambda_0(t) = 2$ and $\beta(t)$'s were given for either of the following two scenarios:

(a) $\beta_1(t) = \dots = \beta_4(t) = 2 \exp\{-10(t - 0.1)^2\}$, $\beta_5(t) = \dots = \beta_8(t) = -1$, and $\beta_9(t) = \dots = \beta_{p_n}(t) = 0$;

(b) $\beta_i(t) = (-1)^{i+1} 2 \exp\{-10(t - i/10)^2\}$ for $i = 1, \dots, 4$, $\beta_i(t) = (-1)^{i+1}(i - 4)/2$ for $i = 5, \dots, 8$, and $\beta_9(t) = \dots = \beta_{p_n}(t) = 0$.

Therefore, for either scenario, only the first 8 covariates were informative. Additionally, in scenario (a), the linked important covariates had the same time-varying effects; while, this was not the case in scenario (b).

To simulate irregular measurements of the covariates, for each subject, we generated measurement times t_{i1}, \dots, t_{in_i} as the ordered uniform distributed times in $[0, \tau]$, where n_i was from a Poisson distribution with mean 8; thus the average number of the measurements per subject was 8. Furthermore, to generate right censored observations, we generated C_i from a uniform distribution in $[0, c]$ where c was chosen to yield about 30% censoring rate.

In the simulation studies, we varied $p_n = 20, 50, 1000$ and $n = 100, 200$. When applying the proposed method to each simulated data, we used Epanechnikov kernel function and quadratic B-splines with two interior knots fixed at sample quantiles of the observed failure times; therefore, $q_n = 5$. When using the penalty form in (16), we re-parameterized $\lambda_1 = \lambda_n \alpha$ and $\lambda_2 = \lambda_n(1 - \alpha)$.

We set $\alpha = 0.2, 0.5, 0.8, 1.0$, and for each α , we selected a path for λ_n as in Friedman et al. (2010). Specifically, λ_n starts from λ_{\max} which ensures all parameters are zero and decreases to a portion of λ_{\max} , i.e. $0.01 \times \lambda_{\max}$, with a length of 10 values. For the bandwidth, we chose from $h_n = 0.05, 0.1, 0.15, 0.2$. To select tuning parameters and bandwidth, 5-fold cross-validation was used. Simulations were repeated 100 times.

To evaluate the estimation performance, we computed the sum of squared errors (SSE) for the estimated β 's. We also calculated the number of true positive covariates (TP) and the number of false positive covariates (FP) as measures of the variable selection performance. Moreover, we compared the performance of the proposed method, time-varying biomarker hazard (DB-hazard) regression for irregularly measured covariates, with LVCF. We also compared different penalty functions, including group LASSO penalty (gLasso), group LASSO with network penalty (gNet) and ℓ_0 -regularization (10) with network penalty (ℓ_0 Net). We also compared with the LVCF (imputing missing covariates by the last observed values) under various penalty functions.

Tables 1 and 2 summarize these simulation results for both settings of $\beta(t)$. It can be seen from both tables that in all cases DB-hazard using all available longitudinal measurements significantly improves the estimation relative to LVCF in terms of a much smaller SSE. It implies that the kernel smoothing method using all available measurements of covariates has less finite sample bias and is more efficient than using the last observed value to impute the covariate values at the observed event times. Using ℓ_0 -penalty in our method is superior to using either group LASSO or network regularization. The former always has a smaller SSE, much better FP and comparable TP. The results indicate the benefits by iteratively performing hard thresholding and considering network structure among variables. When comparing with gLasso and gNet without hard thresholding, gNet has a slightly better SSE and TP but much worse FP, which could be

explained by the grouping effect of using Laplacian penalty. By using Laplacian penalty, if any non-informative covariate is selected, other highly linked covariates would have more chance to be selected as well, which yields many more covariates being identified and poor performance of variable selection.

Table 1: Setting (a) comparison of estimation and selection performance of the proposed DB-hazard using all available longitudinal covariates with LVCF under various penalty functions.

	DB-hazard			LVCF		
	gLasso [†]	gNet [‡]	ℓ_0 Net [*]	gLasso	gNet	ℓ_0 Net
	$n = 100, p_n = 20$					
SSE ¹	4.38	3.76	3.06	5.43	5.19	4.30
TP ²	8.0	8.0	8.0	8.0	8.0	7.9
FP ³	6.3	9.2	1.1	5.9	8.3	0.8
	$n = 100, p_n = 50$					
SSE	5.19	4.30	3.12	6.58	5.74	4.28
TP	8.0	8.0	8.0	8.0	8.0	7.9
FP	14.2	23.9	1.5	11.4	21.9	1.4
	$n = 100, p_n = 1000$					
SSE	8.34	6.25	4.57	9.23	7.53	5.25
TP	7.7	8.0	8.0	7.7	8.0	7.8
FP	33.2	127.2	1.6	29.5	137.6	1.2
	$n = 200, p_n = 20$					
SSE	2.69	2.55	1.92	4.26	4.17	3.40
TP	8.0	8.0	8.0	8.0	8.0	8.0
FP	7.7	9.4	0.8	6.1	8.1	0.8
	$n = 200, p_n = 50$					
SSE	3.51	3.10	2.16	4.92	4.70	3.39
TP	8.0	8.0	8.0	8.0	8.0	8.0
FP	16.3	24.8	1.1	14.2	22.2	0.9
	$n = 200, p_n = 1000$					
SSE	5.04	4.17	2.83	6.52	5.73	4.06
TP	8.0	8.0	8.0	8.0	8.0	8.0
FP	57.1	149.0	1.7	41.8	137.6	1.4

[†]: group Lasso; [‡]: group Lasso with a Laplacian penalty; ^{*}: ℓ_0 -regularization penalty (10)
^[1]:sum of squared error; ^[2]:number of true positive; ^[3]:number of false positive.

Table 3 summarizes the performance of bandwidth selection. We specified a range of candidate bandwidths and performed five-fold cross-validation to select the optimal bandwidth. We compared our selection approach to the method with the smallest SSE among all candidates, denoted as “Best” in Table 3. It can be seen from the table that our selected bandwidths are very close to the “Best” bandwidth, indicating satisfactory performance of our data-driven procedure.

Table 2: Setting (b) comparison of estimation and selection performance of the proposed DB-hazard using all available longitudinal covariates with LVCF under various penalty functions.

	DB-hazard			LVCF		
	gLasso [†]	gNet [‡]	ℓ_0 Net [*]	gLasso	gNet	ℓ_0 Net
	$n = 100, p_n = 20$					
SSE ¹	6.12	5.96	4.96	8.93	8.65	7.49
TP ²	7.6	7.9	7.4	7.3	7.8	7.0
FP ³	7.5	9.2	0.8	6.0	8.0	0.8
	$n = 100, p_n = 50$					
SSE	8.67	8.00	6.08	10.76	10.33	8.26
TP	6.8	7.7	7.2	6.4	7.3	6.6
FP	14.4	24.2	1.2	11.3	18.9	1.1
	$n = 100, p_n = 1000$					
SSE	14.14	13.91	12.59	14.51	14.27	13.05
TP	2.1	3.3	3.5	1.9	3.4	3.4
FP	14.8	38.9	5.2	8.0	31.0	3.7
	$n = 200, p_n = 20$					
SSE	4.04	3.94	3.22	6.61	6.67	5.61
TP	7.9	8.0	7.8	7.9	7.9	7.6
FP	8.5	9.5	0.8	7.9	8.7	0.6
	$n = 200, p_n = 50$					
SSE	5.62	5.53	3.78	8.30	8.25	6.21
TP	7.8	7.9	7.7	7.7	7.8	7.4
FP	18.8	24.9	0.4	16.4	20.7	0.6
	$n = 200, p_n = 1000$					
SSE	10.43	10.02	8.06	12.33	11.77	9.21
TP	5.9	7.1	7.4	5.6	7.3	7.1
FP	48.2	133.6	1.0	26.8	75.3	0.7

[†]: group Lasso; [‡]: group Lasso with a Laplacian penalty; ^{*}: ℓ_0 -regularization penalty (10)
^[1]:sum of squared error; ^[2]:number of true positive; ^[3]:number of false positive.

Table 3: Performance of the bandwidth selection procedure for DB-hazard

	Setting (a)		Setting (b)		Setting (a)		Setting (b)	
	Selected	Best ^[1]	Selected	Best	Selected	Best	Selected	Best
	$n = 100, p_n = 20$				$n = 200, p_n = 20$			
Bandwidth	0.080	0.096	0.093	0.100	0.090	0.081	0.093	0.071
SSE ²	3.06	2.67	4.96	4.37	1.92	1.66	3.22	2.83
	$n = 100, p_n = 50$				$n = 200, p_n = 50$			
Bandwidth	0.080	0.089	0.081	0.095	0.089	0.080	0.088	0.084
SSE	3.12	2.65	6.08	5.24	2.16	1.73	3.78	3.25
	$n = 100, p_n = 1000$				$n = 200, p_n = 1000$			
Bandwidth	0.056	0.085	0.055	0.113	0.059	0.086	0.061	0.104
SSE	4.57	3.89	12.59	11.31	2.83	2.19	8.06	6.90

^[1]: defined as the bandwidth leading to the smallest SSE; ^[2]: sum of squared errors.

To relieve computational burden, we implemented several techniques to speed up our algorithms: we use warm starts for estimating $\beta(t)$ along a regularization path and use sparse data structure in the program to save memory and the time to search for the non-zero coefficients in a sparse β . As a result, the computing time for our method is highly manageable. Figure S1 in supplementary materials shows the running time of the proposed method with ℓ_0 -regularization penalty based on λ with length of 10 and fixed α and h . Overall, the computation time increased linearly with the number of covariates. When $p_n = 1000$ and $n = 200$, the running time is 634 seconds, with a total of $p_n q_n = 5000$ parameters.

We also evaluated the performance of the proposed method based on a different kernel function, i.e. Gaussian kernel. Similar results were obtained by using Gaussian kernel. In addition, the impact of various numbers of basis functions was considered by using quadratic B-splines with 5, 7 and 10 interior knots, corresponding to $q_n = 8, 10, 13$, respectively. We observed an increase in SSE and the number of identified variables as the number of basis functions increased. Note that $\beta_j(t)$ is a linear combination of basis functions. To obtain $\beta_j(t) = 0$, all the elements in the coefficient vector $\gamma_j = (\gamma_{j1}, \dots, \gamma_{jq_n})^T$ have to be zero. Thus, the trend is expected that it is more likely to obtain non-zero estimates with more basis functions. After increasing $n = 100$ to 200, the performance improved, which may suggest we need more sample sizes when describing a more complicated function $\beta_j(t)$ with more basis functions. Details of the above numerical studies are given in the supplementary material.

6 Application

Recent research has suggested that brain imaging biomarkers play an important role in predicting the onset of neurodegenerative disorders, in particular HD (Feigin et al., 2007; Paulsen, 2010, e.g.,). The diagnosis of HD is made based on neurological examination indicating 99% confidence that the extrapyramidal movement disorder is consistent with HD. By the time clinical symptoms are apparent, subjects may already be in an advanced disease stage. Therefore, identifying biomarkers informative for early prediction of disease onset preceding clinical diagnosis has important implications on recruiting pre-symptomatic subjects for clinical trials of early intervention (Paulsen, 2010). Current research indicates neuroimaging biomarkers to be among the most promising ones for predicting time to HD onset (Paulsen et al., 2014a,b). In this work, we analyze data collected from a newly completed, large natural history study of disease progression, PREDICT-HD (Paulsen et al., 2014b), in individuals who carry expanded CAG repeats and are destined to develop HD. CAG repeat length is inversely related to age at onset, but the exact onset age varies. We aim to predict time-to-onset of HD using structural MRI region of interest (ROI) volumetric measures for subjects without a diagnosis at the baseline but with expanded CAG repeats. The regional summary volumetric measures were created by a fully automated procedure and preprocessed using Freesurfer 5.2 (<http://surfer.nmr.mgh.harvard.edu>). Details on the imaging biomarker preprocessing can be found elsewhere (Paulsen et al., 2014a).

Our analysis data consist of 866 subjects who had expanded CAG repeats at the huntingtin gene (MacDonald et al., 1993) without a clinical diagnosis at the baseline. These subjects will develop HD during their lifetime due to the expanded repeat length at the HD gene, but the exact age of onset is unknown. The median length of follow-up time was 4.0 years with an

average of 1.9 follow-ups per subject. Imaging biomarkers were measured approximately bi-annually with some random variation, and thus obtained less frequently than the clinical measures of the time-to-diagnosis outcome (assessed annually). Figure S2 in supplementary materials displays the number of subjects with available clinical measures (time-to-diagnosis outcome) and longitudinal imaging measurements at follow ups, which shows sparse measurements of imaging biomarkers at times (e.g., 18 month after baseline). Biomarkers and clinical assessments included in the analyses are: baseline CAP score (scaled product of CAG repeats length at the HD gene and baseline age; Zhang et al. (2011)) 8 demographic or clinical measures (gender, baseline total motor score, TMS, from the United Huntington's Disease Rating Scale; and cognitive and functioning measures), and whole brain MRI volumetric biomarkers including 58 subcortical region of interest (ROI) measures and 68 cortical ROIs.

Figure S3 in supplementary materials shows the heatmaps of the 136 features measured at the baseline and at the last visit for 142 subjects who were diagnosed with HD during the study (converters) and 390 subjects who remained free of HD diagnosis (non-converters). The goal is to simultaneously select informative biomarkers, estimate their time-dependent effect profiles, and their combination that tracks with HD conversion. In Figure S3, no single feature can definitively distinguish converters from non-converters, suggesting a multi-dimensional approach considering all features will outperform univariate analyses. However, covariation among features is also prevalent and a multi-dimensional approach needs to account for high dimensionality and covariation patterns through regularization. Most biomarkers and subjects show a smooth trend between first and last visit. Appropriately smoothing over time and borrow information from the nearby measurements would be essential for predicting the conversion events, especially for the time points where the imaging measurements were sparse. Lastly, several features show subtle

differences in discriminating converters from non-converters at the first and last visit, suggesting their prognostic power may vary with time (i.e., more discriminant power when using most recent imaging biomarker measurements).

In the analysis, we applied DB-hazard with ℓ_0 -penalty and Laplacian network regularization to the data. We constructed the imaging biomarkers' co-variation network (He et al., 2008) based on an independent control group (no expanded repeat length at the huntingtin gene) in PREDICT-HD. The obtained co-variation pattern was introduced in the Laplacian regularization. For DB-hazard, the estimation follows the same procedure as described in Section 2 by using all longitudinal imaging measurements over time. We used five-fold cross-validation to select the bandwidth and tuning parameters. We compared DB-hazard with using baseline data alone ("Baseline"), and with the last value carried forward ("LVCF"). All features were standardized before fitting the model. The running time of the proposed DB-hazard with ℓ_0 -penalty for this analysis is 927 seconds.

Table S4 in supplementary materials summarizes the area under the ROC curve (AUC), time-dependent sensitivity (SEN), specificity (SPE), positive predictive value (PPV), and negative predictive value (NPV) at a given time where the threshold is obtained by optimizing Youden's index. The results show that overall DB-hazard performs better than the other two alternatives. For example, the AUC of DB-hazard is the highest among three methods at all time points. Comparing with LVCF, DB-hazard outperforms substantially, which may be due to the bias of LVCF. When compared with using only baseline measurements, DB-hazard performs better especially at later years (e.g., year 6), demonstrating the advantages of using current values of longitudinal biomarkers to update longer-term prediction. In a recent independent study, Long et al. (2016) compared Harrell's C-index (i.e., AUC) of ten models including data from four studies of progres-

sion of HD: PREDICT-HD, TRACK (Tabrizi et al., 2013), COHORT (Dorsey et al., 2012), and REGISTRY (Handley et al., 2012). The best model had a median AUC of 0.87, which is similar to our baseline-only analyses but lower than using DB-hazard to incorporate all available longitudinal measures. This comparison further supports the information gain due to incorporating all available time-dependent structural MRI biomarkers and clinical assessments. Our analysis is the first one to use longitudinal imaging biomarkers to track HD conversion.

Regarding other measures, the specificity for DB-hazard by year 6 is 0.873, while it is only 0.651 and 0.810 for Baseline and LVCF, respectively. Similarly, the PPV estimated by DB-hazard by year 6 (0.540) is higher than the other two methods (Baseline: 0.278; LVCF: 0.414). The high time-dependent sensitivity and specificity of the DB-hazard combined HD biomarker signature suggest that it is a valuable tool that tracks with clinically defined disease onset. The higher PPV at year 4 and 6 demonstrates valuable information gain by using the longitudinal imaging measures to improve prediction. However, the moderate magnitude of PPV implies that additional biomarkers (e.g., genomic or proteomic biomarkers; Langfelder et al. (2016)) may need to be identified to improve prospective prediction performance.

From 136 features, DB-hazard identified 6 non-imaging covariates (i.e. CAP, total functional capacity - TFC, baseline total motor score - baseline TMS, symbol digit modality test - SDMT, stroop color naming total and stroop word reading total) and 6 imaging biomarkers (i.e. left Caudate, left and right Putamen, left Pallidum, left Accumbens, left lateral occipital volume) as informative for predicting time-to-onset of HD, including 5 subcortical measures and 1 cortical measure (left lateral occipital volume). Figure S4 in supplementary materials shows the heatmaps of the selected features, where they are seen to better distinguish converters from non-converters than other non-selected noise features in Figure S3. In addition, through the use of network

regularization, redundancy among features was removed and a total of 12 features achieves a high AUC. Some features' discriminant power changes at the first and last visit (e.g., TFC). It is interesting that more subcortical ROIs were selected and only 1 cortical ROI was selected when both were included in DB-hazard as candidates. This result is consistent with clinical research suggesting regional atrophy of subcortical grey matter is an important biological feature of HD progression (Ross et al., 2014). All subcortical ROIs identified were ranked as top candidate biomarkers in existing clinical research (Paulsen et al., 2014a), while the cortical measure was not reported before. In Figure S5, we present the effects of top-ranking measures estimated by DB-hazard and the corresponding 95% confidence intervals obtained by bootstrap 100 times. The baseline TMS has the strongest effect and a similar shape as the baseline CAP. The effects of TMS, TFC and SDMT increases in the first two years and the largest effect is reached between year 2 and 3. The two imaging biomarkers, left Caudate and left Putamen, show similar effect size as baseline CAP, TFC and SDMT. The measure with the largest effect is the baseline TMS.

7 Discussion

In this article, we propose methods for fitting time-varying hazards model with sparsely measured time-dependent covariates. A contribution of our method compared to analyses in the existing literature (e.g., Paulsen et al., 2014b; Long et al., 2016) is that we used all longitudinal measures (both imaging biomarkers and clinical measures at the follow-ups) to perform analyses, which was not available previously due to imbalanced assessments of imaging measures. Our simulation studies show that smoothing over longitudinal measurements across subjects improve performance over the commonly used LVCF and baseline only analysis. In addition, the pro-

posed DB-hazard with ℓ_0 -penalty solved by a two-step procedure substantially outperform other methods without the hard-thresholding or using group LASSO alone, in term of both estimation and selection accuracy. We prove the theoretical oracle property under the local kernel smoothing, which has not been investigated in the literature previously. We also demonstrate substantial improvement on the applications to a real world study data (PREDICT-HD) compared to current clinical standards.

Here, we assume a constant network structure in equation (15). It would be interesting and challenging to explore time-varying network $\mathcal{L}_n^*(t)$. One method is to incorporate a time-varying Gaussian graphical model (Zhou et al., 2010; Razavian et al., 2010), where the time-varying network $\widehat{\mathcal{L}}_n^*(t)$ can be obtained from $\widehat{\mathcal{L}}_n^*(t)^{-1} = \arg \max_{\Theta} \log |\Theta(t)| - \text{tr}(\mathbf{R}(t)\Theta(t)) - \rho \|\Theta(t)\|_1$, where $\mathbf{R}(t)$ is the weighted correlation matrix and can be calculated from the weighted covariance matrix

$$\mathbf{R}'(t) = \frac{\sum_{i=1}^n \sum_{v=1}^{n_i} K_{h_n}(t_{iv} - t) \mathbf{X}_i(t_{iv}) \mathbf{X}_i(t_{iv})^T}{\sum_{i=1}^n \sum_{v=1}^{n_i} K_{h_n}(t_{iv} - t)}.$$

Another extension is to study the effect of time-varying network on the disease outcome, and to distinguish the effect from the longitudinal measurements themselves and their time-varying network.

Lastly, here we focus on time-to-event data. However, the proposed approach can be easily extended to other types of outcomes. One would replace the log-partial likelihood function with the least squares loss function for continuous outcome or appropriate likelihood for generalized outcomes.

Supplementary Material

Supplementary material available online includes the proofs of Lemma 1 and Theorem 1, and additional information for simulation and real data analysis.

References

- Alexander-Bloch, A., Giedd, J. N. et al. (2013) Imaging structural co-variance between human brain regions. *Nature Reviews Neuroscience*, **14**, 322–336.
- Andersen, P. K. and Liestol, K. (2003) Attenuation caused by infrequently updated covariates in survival analysis. *Biostatistics*, **4**, 633–649.
- Boyd, S., Parikh, N., Chu, E., Peleato, B. and Eckstein, J. (2011) Distributed optimization and statistical learning via the alternating direction method of multipliers. *Foundations and Trends in Machine Learning*, **3**, 1–122.
- Bradic, J., Fan, J. and Jiang, J. (2011) Regularization for coxs proportional hazards model with np-dimensionality. *Annals of statistics*, **39**, 3092–3120.
- Bullmore, E. T. and Bassett, D. S. (2011) Brain graphs: graphical models of the human brain connectome. *Annual review of clinical psychology*, **7**, 113–140.
- Cao, H., Churpek, M. M., Zeng, D. and Fine, J. P. (2015) Analysis of the proportional hazards model with sparse longitudinal covariates. *Journal of the American Statistical Association*, **110**, 1187–1196.

- Cao, H., Zeng, D. and Fine, J. P. (2014) Regression analysis of sparse asynchronous longitudinal data. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **77**, 775–776.
- Chen, T., Wang, Y., Chen, H., Marder, K. and Zeng, D. (2014) Targeted local support vector machine for age-dependent classification. *Journal of the American Statistical Association*, **109**, 1174–1187.
- Desikan, R. S., Ségonne, F., Fischl, B., Quinn, B. T., Dickerson, B. C., Blacker, D., Buckner, R. L., Dale, A. M., Maguire, R. P., Hyman, B. T. et al. (2006) An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. *NeuroImage*, **31**, 968–980.
- Dorsey, E. R., Investigators, H. S. G. C. et al. (2012) Characterization of a large group of individuals with huntington disease and their relatives enrolled in the cohort study. *PLoS One*, **7**, e29522.
- Eidelberg, D. and Surmeier, D. J. (2011) Brain networks in Huntington disease. *Journal of Clinical Investigation*, **121**, 484–492.
- Fan, J. and Li, R. (2001) Variable selection via nonconcave penalized likelihood and its oracle properties. *Journal of the American Statistical Association*, **96**, 1348–1360.
- Fan, J. and Lv, J. (2011) Nonconcave penalized likelihood with np-dimensionality. *Information Theory, IEEE Transactions*, **57**, 5467–5484.
- Feigin, A., Tang, C., Ma, Y., Mattis, P., Zgaljardic, D., Guttman, M., Paulsen, J., Dhawan, V. and Eidelberg, D. (2007) Thalamic metabolism and symptom onset in preclinical huntington’s disease. *Brain*, **130**, 2858–2867.

- Fleming, T. R. and Harrington, D. P. (2011) *Counting processes and survival analysis*, vol. 169. John Wiley & Sons.
- Friedman, J., Hastie, T. and Tibshirani, R. (2010) Regularization paths for generalized linear models via coordinate descent. *Journal of statistical software*, **33**, 1–22.
- Gould, L. A., Boye, M. E., Crowther, M. J., Ibrahim, J. G., Quartey, G., Micallef, S. and Bois, F. Y. (2014) Joint modeling of survival and longitudinal non-survival data: current methods and issues. Report of the DIA Bayesian joint modeling working group. *Statistics in Medicine*, **34**, 2181–2195.
- Handley, O., Landwehrmeyer, B., Committee, R. S., Investigators, E. R. et al. (2012) European huntington’s disease network registry: current status. *Journal of Neurology, Neurosurgery & Psychiatry*, **83**, A47–A47.
- He, Y., Chen, Z. and Evans, A. (2008) Structural insights into aberrant topological patterns of large-scale cortical networks in alzheimer’s disease. *The Journal of neuroscience*, **28**, 4756–4766.
- Honda, T. and Härdle, W. K. (2014) Variable selection in cox regression models with varying coefficients. *Journal of Statistical Planning and Inference*, **148**, 67–81.
- Honda, T. and Yabe, R. (2017) Variable selection and structure identification for varying coefficient cox models. *Journal of Multivariate Analysis*, **161**, 103–122.
- Huang, J., Horowitz, J. L. and Wei, F. (2010) Variable selection in nonparametric additive models. *Annals of Statistics*, **38**, 2282–2313.

- Lange, K. (2013) The MM algorithm. In *Optimization*, 185–219. Springer.
- Langfelder, P., Cattle, J. P., Chatzopoulou, D., Wang, N., Gao, F., Al-Ramahi, I., Lu, X.-H., Ramos, E. M., El-Zein, K., Zhao, Y. et al. (2016) Integrated genomics and proteomics define huntingtin cag length-dependent networks in mice. *Nature neuroscience*, **19**, 623–633.
- Li, X., Xie, S., Zeng, D. and Wang, Y. (2017) Efficient 0-norm feature selection based on augmented and penalized minimization. *Statistics in medicine*, in press.
- Liu, J., Huang, J., Ma, S. and Wang, K. (2013) Incorporating group correlations in genome-wide association studies using smoothed group lasso. *Biostatistics*, **14**, 205–219.
- Liu, M., Zhang, D., Shen, D., Initiative, A. D. N. et al. (2014) Identifying informative imaging biomarkers via tree structured sparse learning for ad diagnosis. *Neuroinformatics*, **12**, 381–394.
- Liu, X. and Zeng, D. (2013) Variable selection in semiparametric transformation models for right-censored data. *Biometrika*, **100**, 859–876.
- Long, J. D., Langbehn, D. R., Tabrizi, S. J., Landwehrmeyer, B. G., Paulsen, J. S., Warner, J. and Sampaio, C. (2016) Validation of a prognostic index for huntington’s disease. *Movement Disorders*, **32**, 256–263.
- MacDonald, M. E., Ambrose, C. M., Duyao, M. P., Myers, R. H., Lin, C., Srinidhi, L., Barnes, G., Taylor, S. A., James, M., Groot, N. et al. (1993) A novel gene containing a trinucleotide repeat that is expanded and unstable on huntington’s disease chromosomes. *Cell*, **72**, 971–983.
- Meier, L., Van De Geer, S. and Bühlmann, P. (2008) The group lasso for logistic regression. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **70**, 53–71.

- Parikshak, N. N., Gandal, M. J. and Geschwind, D. H. (2015) Systems biology and gene networks in neurodevelopmental and neurodegenerative disorders. *Nature Reviews Genetics*, **16**, 441–458.
- Paulsen, J. S. (2010) Early detection of huntington’s disease. *Future Neurology*, **5**, 85–104.
- Paulsen, J. S., Long, J. D., Johnson, H. J., Aylward, E. H., Ross, C. A., Williams, J. K., Nance, M. A., Erwin, C. J., Westervelt, H. J., Harrington, D. L. et al. (2014a) Clinical and biomarker changes in premanifest huntington disease show trial feasibility: a decade of the predict-hd study. *Frontiers in Aging Neuroscience*, **6**, 78.
- Paulsen, J. S., Long, J. D., Ross, C. A., Harrington, D. L., Erwin, C. J., Williams, J. K., Westervelt, H. J., Johnson, H. J., Aylward, E. H., Zhang, Y. et al. (2014b) Prediction of manifest huntington’s disease with clinical and imaging measures: a prospective observational study. *The Lancet Neurology*, **13**, 1193–1201.
- Prentice, R. (1982) Covariate measurement errors and parameter estimation in a failure time regression model. *Biometrika*, **69**, 331–342.
- Razavian, N. S., Moitra, S., Kamisetty, H., Ramanathan, A. and Langmead, C. J. (2010) Time-varying gaussian graphical models of molecular dynamics data. *Technical Report*.
- Rizopoulos, D. (2011) Dynamic predictions and prospective accuracy in joint models for longitudinal and time-to-event data. *Biometrics*, **67**, 819–829.
- Ross, C. A., Aylward, E. H., Wild, E. J., Langbehn, D. R., Long, J. D., Warner, J. H., Scahill, R. I., Leavitt, B. R., Stout, J. C., Paulsen, J. S. et al. (2014) Huntington disease: natural history, biomarkers and prospects for therapeutics. *Nature Reviews Neurology*, **10**, 204–216.

- Ryan, N. P., Catroppa, C., Cooper, J. M., Beare, R., Ditchfield, M., Coleman, L., Silk, T., Crossley, L., Beauchamp, M. H. and Anderson, V. A. (2015) The emergence of age-dependent social cognitive deficits after generalized insult to the developing brain: A longitudinal prospective analysis using susceptibility-weighted imaging. *Human brain mapping*, **36**, 1677–1691.
- Schumaker, L. (2007) *Spline functions: basic theory*. Cambridge University Press.
- Simon, N., Friedman, J., Hastie, T. and Tibshirani, R. (2013) A sparse-group lasso. *Journal of Computational and Graphical Statistics*, **22**, 231–245.
- Stuart, J. M., Segal, E., Koller, D. and Kim, S. K. (2003) A gene-coexpression network for global discovery of conserved genetic modules. *Science*, **302**, 249–255.
- Tabrizi, S. J., Scahill, R. I., Owen, G., Durr, A., Leavitt, B. R., Roos, R. A., Borowsky, B., Landwehrmeyer, B., Frost, C., Johnson, H. et al. (2013) Predictors of phenotypic progression and disease onset in premanifest and early-stage huntington’s disease in the track-hd study: analysis of 36-month observational data. *The Lancet Neurology*, **12**, 637–649.
- 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.
- Tsiatis, A. A. and Davidian, M. (2001) A semiparametric estimator for the proportional hazards model with longitudinal covariates measured with error. *Biometrika*, **88**, 447–458.
- (2004) Joint modeling of longitudinal and time-to-event data: an overview. *Statistica Sinica*, **14**, 809–834.

Yan, J. and Huang, J. (2012) Model selection for cox models with time-varying coefficients.

Biometrics, **68**, 419–428.

Zhang, C.-H. (2010) Nearly unbiased variable selection under minimax concave penalty. *The*

Annals of Statistics, **38**, 894–942.

Zhang, C.-H. and Zhang, T. (2012) A general theory of concave regularization for high-

dimensional sparse estimation problems. *Statistical Science*, 576–593.

Zhang, Y., Long, J. D., Mills, J. A., Warner, J. H., Lu, W. and Paulsen, J. S. (2011) Indexing

disease progression at study entry with individuals at-risk for huntington disease. *American*

Journal of Medical Genetics Part B: Neuropsychiatric Genetics, **156**, 751–763.

Zhao, P. and Yu, B. (2006) On model selection consistency of lasso. *J. Mach. Learn. Res.*, **7**,

2541–2563.

Zhou, S., Lafferty, J. and Wasserman, L. (2010) Time varying undirected graphs. *Machine Learn-*

ing, **80**, 295–319.