Inference for High Dimensional Censored Quantile Regression

Zhe Fei, Qi Zheng, Hyokyoung G. Hong, and Yi Li
1. Department of Biostatistics, University of California, Los Angeles
2. Department of Bioinformatics and Biostatistics, University of Louisville
3. Department of Statistics and Probability, Michigan State University
4. Department of Biostatistics, University of Michigan

Abstract

With the availability of high dimensional genetic biomarkers, it is of interest to identify heterogeneous effects of these predictors on patients’ survival, along with proper statistical inference. Censored quantile regression has emerged as a powerful tool for detecting heterogeneous effects of covariates on survival outcomes. To our knowledge, there is little work available to draw inference on the effects of high dimensional predictors for censored quantile regression. This paper proposes a novel procedure to draw inference on all predictors within the framework of global censored quantile regression, which investigates covariate-response associations over an interval of quantile levels, instead of a few discrete values. The proposed estimator combines a sequence of low dimensional model estimates that are based on multi-sample splittings and variable selection. We show that, under some regularity conditions, the estimator is consistent and asymptotically follows a Gaussian process indexed by the quantile level. Simulation studies indicate that our procedure can properly quantify the uncertainty of the estimates in high dimensional settings. We apply our method to analyze the heterogeneous effects of SNPs residing in lung cancer pathways on patients’ survival, using the Boston Lung Cancer Survival Cohort, a cancer epidemiology study on the molecular mechanism of lung cancer.

KEYWORDS: Conditional Quantiles; Fused-HDCQR; High Dimensional Predictors; Statistical Inference; Survival Analysis.

CONTACT: Yi Li, yili@umich.edu, Department of Biostatistics, University of Michigan, Ann Arbor, MI 48109. Supplementary materials for this article are available online. Please go to www.tandfonline.com/r/JASA.

ARTICLE HISTORY: Received December 2019; Accepted July 2021.
1 Introduction

Lung cancer presents much heterogeneity in etiology (McKay et al., 2017; Dong et al., 2012; Huang et al., 2009), and some genetic variants may insert different impacts on different quantile levels of survival time. For example, in the Boston Lung Cancer Survival Cohort (BLCSC) (Christiani, 2017), a cancer epidemiology cohort of over 11,000 lung cancer cases enrolled in the Boston area since 1992, it was found that SNP AX.37793583 (rs115952579), along with age, gender, cancer stage and smoking status, had heterogeneous effects on different quantiles of survival time. A total of 674 patients in the study were genotyped, with the goal of identifying lung cancer survival-predictive SNPs. Target gene approaches, which focus on SNPs residing in cancer-related gene pathways, are appealing for increased statistical power in detecting significant SNPs (Moon et al., 2003; Risch and Plass, 2008; Ho et al., 2019), and the investigators have identified SNPs residing in 14 well-known lung cancer-related genes (Zhu et al., 2017; Korpanty et al., 2014; Yamamoto et al., 2008; Kelley et al., 2001). One goal was to investigate whether and how each SNP might play a different role among the high-risk (i.e., lower quantiles of overall survival) and low-risk (i.e., higher quantiles of overall survival) cancer survivors.

Quantile regression (QR) (Koenker and Bassett Jr, 1978) is a significant extension of classic linear regression. By permitting the effects of active variables to vary across quantile levels, quantile regression can naturally accommodate and examine the heterogeneous impacts of biomarkers on different segments of the response variable’s conditional distribution. As survival data are subject to censoring and may be incomplete, QR methods developed for complete data may be unsuitable. Efforts have been devoted to developing censored quantile regression (CQR) (Powell, 1986; Portnoy, 2003; Peng and Huang, 2008, among others), which has become a useful alternative strategy to traditional survival models, such as the Cox model and the accelerated failure time model. QR has also been widely studied to accommodate high dimensional predictors. For example, Wang et al. (2012) dealt with
variable selection using non-convex penalization; Zheng et al. (2013) proposed an adaptive penalized quantile regression estimator that can select the true sparse model with high probability; and Fan et al. (2014) studied the penalized quantile regression with a weighted $L_1$ penalty in an ultra-high dimensional setting. As to high dimensional CQR (HDCQR), He et al. (2013) provided a model-free variable screening procedure for ultrahigh dimensional covariates, and Zheng et al. (2018) proposed a penalized HDCQR built upon a stochastic integral based estimating equation. However, most of the existing works in HDCQR were designed to select a subset of predictors and estimate the effects of the selected variables, instead of drawing inference on all predictors.

Progress in high dimensional inferences has been made for linear and non-linear models (Zhang and Zhang, 2014; Bühlmann et al., 2014; Javanmard and Montanari, 2014; Ning and Liu, 2017; Fei et al., 2019; Fei and Li, 2021). For example, Meinshausen et al. (2009) proposed to aggregate p-values from multi-sample splittings for high dimensional linear regression. Another line of works referred to as post-selection inference includes Berk et al. (2013), Lee et al. (2016), and Belloni et al. (2019), which provided post-selection inference at fixed quantiles for complete data. However, these methods may not handle censored outcomes. For censored median regression, Shows et al. (2010) provided sparse estimation and inference, but it cannot handle high dimensional data.

We propose to draw inference on high dimensional HDCQR based on a splitting and fusing scheme, termed Fused-HDCQR. Utilizing a variable selection procedure for HDCQR such as Zheng et al. (2018), our method operates partial regression followed by smoothing. Specifically, partial regression allows us to estimate the effect of each predictor, regardless of whether or not it is chosen by variable selection. The fused estimator aggregates the estimates based on multiple data-splittings and variable selection, with a variance estimator derived by the functional delta method (Efron, 2014; Wager and Athey, 2018). To comprehensively assess the covariate effects on the survival distribution, we adopt a “global”
quantile model (Zheng et al., 2015) with the quantile level varying over an interval, instead of a local CQR that focuses only on a few pre-specified quantile levels. The global quantile model can address the molecular mechanism of lung cancer, our motivating disease, that hypothesizes that some genetic variants may cause heterogeneous impacts on different but unspecified segments of survival distribution (McKay et al., 2017; Dong et al., 2012; Huang et al., 2009).

Our work presents several advantages. First, compared to high dimensional Cox models (Zhao and Li, 2012; Fang et al., 2017; Kong et al., 2021), the employed HDCQR stems from the accelerated failure time model (Wei, 1992) and offers straightforward interpretations (Hong et al., 2019). Second, utilizing the global conditional quantile regression, it uses various segments of the conditional survival distribution to improve the robustness of variable selection and capture global sparsity. Third, our splitting-and-averaging scheme avoids the technicalities of estimating the precision matrix by inverting a \( p \times p \) Hessian matrix of the log likelihood, which is a major challenge for debiased-LASSO type methods (Zhang and Zhang, 2014; Van de Geer et al., 2014) and is even more so if we apply debiased-LASSO to the CQR setting. Finally, as opposed to post-selection inferences (Belloni et al., 2019, among others), Fused-HDCQR accounts for variations in model selection and draws inference for all of the predictors.

The rest of the paper is organized as follows. Section 2 introduces the method, and Section 3 details the asymptotic properties. Section 4 derives a non-parametric variance estimator, Section 5 conducts simulation studies, and Section 6 applies the proposed method to analyze the BLCSC data. The technical details, such as proofs and additional lemmas, are relegated to the online Supplementary Materials.
2 Model and Method

2.1 High dimensional censored quantile regression

Let $T$ and $C$ denote the survival outcome and censoring time, respectively. We assume that $C$ is independent of $T$ given $\tilde{Z}$, a $(p-1)$-dimensional vector of covariates ($p > 1$). Let $X = \min\{T, C\}$, $\Delta = 1\{T \leq C\}$, and $Z = (1, \tilde{Z}^T)^T$, where $1\{\cdot\}$ is the binary indicator function.

The observed data, $D^{(n)} = \{(X_i, \Delta_i, Z_i), i = 1, \ldots, n\}$, are $n$ identical and independently distributed (i.i.d.) copies of $(X, \Delta, Z)$. With $Y = \log T$, let $Q_Y(\tau|Z) = \inf\{t: P(Y \leq t|Z) \geq \tau\}$ be the $\tau$-th conditional quantile of $Y$ given $Z$. A global censored quantile regression model stipulates $Q_Y(\tau|Z) = Z^T \beta^*(\tau)$, $\tau \in (0, 1)$, where $\beta^*(\tau)$ is a $p$-dimensional vector of coefficients at $\tau$. We aim to draw inference on $\beta^*_j(\tau)$ for each $\tau \in (0, \tau_U]$ and for all $j \in \{1, \ldots, p\}$, where $0 < \tau_U < 1$ is an upper bound for estimable quantiles subject to identifiability constraint caused by censoring (Peng and Huang, 2008).

Let $N(t) = 1\{\log X \leq t, \Delta = 1\}$, $\Lambda_T(t|Z) = -\log(1 - P(\log T \leq t|Z))$, and $H(u) = -\log(1 - u)$. Then, $M(t) = N(t) - \Lambda_T(t \wedge \log X|Z)$ is a martingale process under model (1) (Fleming and Harrington, 2011) and hence $E(M(t)|Z) = 0$. We use $N_i(t)$ and $M_i(t)$, $i = 1, \ldots, n$, to denote the sample analogs of $N(t)$ and $M(t)$. Let $\theta_i(\tau) = Z_i^T \beta(\tau)$ and $\theta_i^*(\tau) = Z_i^T \beta^*(\tau)$.

We denote by $u(\beta, \tau)$ the expectation of $U_n(\beta, \tau)$.

The martingale property implies $u(\beta^*, \tau) = 0$ with $\tau \in [0, \tau_U]$, entailing an estimating equation with $\tau \in (0, \tau_U]$:

$$n^{1/2} U_n(\beta, \tau) = n^{-1/2} \sum_{i=1}^n Z_i \left\{N_i(\theta_i(\tau)) - \int_0^\tau 1\{\log X_i \geq \theta_i(u)\} dH(u)\right\} = 0. \quad (2)$$
The stochastic integral in (2) naturally suggests sequential estimation with respect to \( \tau \). We define a grid of quantile values \( \Gamma_m = \{\tau_0, \tau_1, \ldots, \tau_m\} \) to cover the interval \([\nu, \tau_U]\), where \( \tau_0 = \nu \) and \( \tau_m = \tau_U \). The assumption on the lower bound \( \nu > 0 \) is made to circumvent the singularity problem with CQR at \( \tau = 0 \), as detailed in assumption (A1). In practice, \( \nu \) is chosen such that only a small proportion of observations are censored below the \( \nu \)-th quantile.

Then, \( \hat{\beta}(\tau_k) \)'s, the estimates of \( \beta(\tau_k) \)'s, \( \tau_k \in \Gamma_m \), can be sequentially obtained by solving

\[
\sum_{i=1}^{n} Z_i \left( N_i (\theta_i(\tau_k)) - \sum_{r=0}^{k-1} \int_{\tau_r}^{\tau_{r+1}} \{\log X_i \geq \hat{\theta}_i(\tau_r)\} dH(u) \right) = 0,
\]

where \( \hat{\theta}_i(\tau_k) = Z_i^T \hat{\beta}(\tau_k) \). Due to the monotonicity of \( \theta_i(\tau) \) in \( \tau \), \( \hat{\beta}(\tau_k) \) can be solved efficiently via \( L_1 \)-minimization. And \( \hat{\beta}(\tau), \tau \in [\nu, \tau_U] \), is defined as a right-continuous piece-wise constant function that only jumps at the grid points. It can be shown that \( \hat{\beta}(\tau) \) is uniformly consistent and converges weakly to a mean zero Gaussian process for \( \tau \in [\nu, \tau_U] \) when \( p = o(n) \). More importantly, \( \hat{\beta}(\tau) \) provides a comprehensive understanding of the covariate effects on the conditional survival distribution over the quantile interval \([\nu, \tau_U]\).

We incorporate this sequential estimating procedure for low dimensional CQR estimation in our proposed method.

In addition, our method requires dimension reduction, which can be accomplished by existing methods, including the screening method proposed by He et al. (2013) and the penalized estimation and selection procedure developed by Zheng et al. (2018). Specifically, Zheng et al. (2018) incorporated an \( L_1 \) penalty into the stochastic integral based estimating equation in (2) to obtain an L-HDCQR estimator, which achieves a uniform convergence rate of \( \sqrt{q \log(p \lor n)/n} \), and results in “sure screening” variable selection with high probability, where \( q \) is defined in condition (A4). Zheng et al. (2018) also proposed an AL-HDCQR estimator by employing the Adaptive Lasso penalties, which attains a uniform convergence rate of \( \sqrt{q \log(n)/n} \) and selection consistency.
2.2 Fused-HDCQR estimator

Our proposed Fused-HDCQR procedure consists of multiple data splitting, selecting variables, fitting low dimensional CQRs with partitioned data, applying append-and-estimate to all predictors, and aggregating those estimates.

1. With the full data $D^{(n)}$, determine via cross-validation the tuning parameter(s) $\lambda_n$ of $S$, an HDCQR variable selection method.

2. Let $B$ be a large positive integer. For each $b = 1, \ldots, B$,
   
   (i) randomly split the data into equal halves, $D^b_1$ and $D^b_2$;
   
   (ii) on $D^b_1$, apply $S$ with $\lambda_n$ on $[\nu, \tau_U]$, to select a subset of predictors, denoted by $\hat{S}^b_{\lambda_n}$, or $\hat{S}^b$ for short;
   
   (iii) on $D^b_2$, for each $j = 1, \ldots, p$, append $j$ to $\hat{S}^b$ such that $\hat{S}^b_{+j} = \{j\} \cup \hat{S}^b$, fit a partial CQR on the covariates indexed by $\hat{S}^b_{+j}$, and denote their coefficient estimates by $\tilde{\beta}_{\hat{S}^b_{+j}}(\tau)$, $\tau \in [\nu, \tau_U]$. Here, $\tilde{\beta}_{\hat{S}^b_{+j}}(\tau)$ is a right-continuous piecewise-constant function with jumps only at the grid points of $\tau_k \in \Gamma_m$;
   
   (iv) denote by $\tilde{\beta}_j^b(\tau) = \left(\tilde{\beta}_{\hat{S}^b_{+j}}(\tau)\right)_j$ the entry of $\tilde{\beta}_{\hat{S}^b_{+j}}(\tau)$ corresponding to $Z_j$.

3. Fusing: the final estimate of $\beta_j^*(\tau)$, $\tau \in [\nu, \tau_U]$, $j = 1, \ldots, p$ is

   $$\hat{\beta}_j(\tau) = \frac{1}{B} \sum_{b=1}^{B} \tilde{\beta}_j^b(\tau).$$

*Remark 1.* We could select different tuning parameters for $S$ in each data split, but with much added computation. Our numerical evidence suggests that a globally chosen $\lambda_n$ work well.

*Remark 2.* Our procedure needs a variable selection procedure to reduce dimension. For example, L-HDCQR selects a subset: $\{j \in \{2, \ldots, p\} : \max_k |\hat{\gamma}_j(\tau_k)| > a_0, \tau_k \in \Gamma_m\}$, where
\(\hat{\gamma}_j(\tau_k)\)'s are the L-HDCQR estimates, \(a_0 > 0\) is a predetermined threshold, and \(j\) starts with 2 as the intercept term (corresponding to \(j = 1\)) is always included in the model. For the choice of variable selection methods, our experience suggests that we adopt the screening method in He et al. (2013) for fast computation, use L-HDCQR for detecting any non-zero effects in the quantile interval \([\nu, \tau_U]\), and choose AL-HDCQR if we opt to select fewer predictors.

**Remark 3.** We select \(\lambda_n\) by minimizing a \(K\)-fold cross-validation error defined by deviance residuals in the presence of censored outcomes (Zheng et al., 2018). Specifically, we partition the data to \(K\) folds, and let \(\hat{\beta}_\lambda^{(-k)}(\tau)\) be the penalized estimate of \(\beta(\tau)\) using all of the data excluding the \(k\)-th fold with a tuning parameter \(\lambda\) and \(\tau \in [\nu, \tau_U]\), where \(k = 1, \ldots, K\).

Under the global CQR model (1), we define the cross-validation error as

\[
\text{CV Error}(\lambda) = \sum_{k=1}^{K} \sum_{i \in \text{fold } k} \int_{\nu}^{\tau_U} |D_i[\hat{\beta}_\lambda^{(-k)}(\tau)]| d\tau,
\]

where

\[
D_i[\beta(\tau)] = \text{sign}\{M_i(\beta(\tau))\} \sqrt{-2M_i(\beta(\tau)) + \Delta_i \log \{\Delta_i - M_i(\beta(\tau))\}}
\]

with \(M_i(\beta(\tau)) = N_i(\mathbf{Z}_i^T \beta(\tau)) - \int_{\nu}^{\tau} \{\log X_i \geq N_i(\mathbf{Z}_i^T \beta(u))\} dH(u) - \nu\). Here, \(H(u) = -\log(1 - u)\), \(N_i(\cdot)\) is the counting process, and \(M_i(\beta(\tau))\) is the martingale residual under model (1) (Zheng et al., 2018).

**Remark 4.** When carrying out quantile regression at each grid point, we formulate it as a linear programming problem (Koenker, 2005), which can be solved by a simplex algorithm with a computational complexity of \(O(n^2p)\) (Klee and Minty, 1972). Since our grid size is \(O(n)\) and the number of resampling, \(B\), is \(O(n)\), the computational complexity of our procedure is \(O(n^4p)\).
3 Theoretical Studies

3.1 Notation and regularity conditions

For any vector $\delta \in \mathbb{R}^p$ and a subset $S \subset \{1, \ldots, p\}$, denote by $S^C$ its complementary set, and define $\|\delta\|_{r,S} = \|\delta_S\|_r$, the $l_r$-norm of the sub-vector $\delta_S$, in which $\delta_j = \delta_j$ if $j \in S$ and $\delta_j = 0$ if $j \in S^C$. We set the following conditions.

(A1) There exist a quantile $\nu$ and a constant $c > 0$ such that

$$n^{-1} \sum_{i=1}^n \{ \log C_i \leq Z_i^T \beta^*(\nu) \} (1 - \Delta_i) \leq cn^{-1/2}$$

holds for sufficiently large $n$.

(A2) (Bounded observations) $\|Z\|_\infty \leq C_0$. Without loss of generality, we assume $C_0 = 1$. In addition, $E|\log X| < \infty$.

(A3) (Bounded densities) Let $F_T(t|Z) = P(\log T \leq t|Z)$, $\Lambda_T(t|Z) = -\log (1 - F_T(t|Z))$, $F(t|Z) = P(\log X \leq t|Z)$, and $G(t|Z) = P(\log X \leq t, \Delta = 1|Z)$. Also, define $f(t|Z) = dF(t|Z)/dt$, and $g(t|Z) = dG(t|Z)/dt$.

(a) There exist constants $\underline{f}$, $\bar{f}$, $\underline{g}$ and $\bar{g}$ such that

$$\underline{f} \leq \inf_{z,\tau \in [\nu, \tau_U]} f(z^T \beta^*(\tau)|z) \leq \sup_{z,\tau \in [\nu, \tau_U]} f(z^T \beta^*(\tau)|z) \leq \bar{f},$$

$$\underline{g} \leq \inf_{z,\tau \in [\nu, \tau_U]} g(z^T \beta^*(\tau)|z) \leq \sup_{z,\tau \in [\nu, \tau_U]} g(z^T \beta^*(\tau)|z) \leq \bar{g}.$$ 

(b) There exist constants $\kappa > 0$ and $A$ such that, when $|t| \leq \kappa$,

$$\sup_{z,\tau \in [\nu, \tau_U]} \left| f(z^T \beta^*(\tau) + t|z) - f(z^T \beta^*(\tau)|z) \right| \leq A|t|,$$

$$\sup_{z,\tau \in [\nu, \tau_U]} \left| g(z^T \beta^*(\tau) + t|z) - g(z^T \beta^*(\tau)|z) \right| \leq A|t|.$$
(A4) (Sparsity) Assume \( \log p = o(n^{1/2}) \), and let

\[
S_\tau = \{ j : \beta_j^*(\tau) \neq 0 \}, \quad S^* = \bigcup_{\tau \in [\nu, \tau_U]} S_\tau = \left\{ j : \sup_{\tau \in [\nu, \tau_U]} |\beta_j^*(\tau)| > 0 \right\}, \quad \text{and} \quad q = |S^*|.
\]

Let \( \hat{S} \) be the index set of covariates selected by \( S \) with a tuning parameter \( \lambda_n \). There exist constants \( 0 \leq c_1 < 1/3, c_2, K_1, K_2 > 0 \) such that

\[
q \leq K_1 n^{c_1}, \quad |\hat{S}| \leq K_2 n^{c_1}, \quad \text{and} \quad P \left( S^* \subseteq \hat{S} \right) \geq 1 - K_2 (p \vee n)^{-1-c_2}.
\]

(A5) Let \( \tilde{\mu}(\tau) = E \left[ 1 \{ \log X > Z^T \beta^*(\tau) \} \right] \). There exists a constant \( L > 0 \) such that

\[
|\beta_j^*(\tau_1) - \beta_j^*(\tau_2)| \leq L|\tau_1 - \tau_2| \quad \text{and} \quad |\tilde{\mu}(\tau_1) - \tilde{\mu}(\tau_2)| \leq L|\tau_1 - \tau_2|, \quad \text{for all} \quad \tau_1, \tau_2 \in (\nu, \tau_U) \quad \text{and} \quad 1 \leq j \leq p.
\]

(A6) (Bounded eigenvalues) \( \delta^T E[Z_i Z_i^T] \delta / \| \delta \|^2 \) is bounded below and above by \( \lambda_{\text{min}} \) and \( \lambda_{\text{max}} \), respectively, over \( \| \delta \|_0 \leq K_1 n^{c_1}, \delta \neq 0 \), where \( 0 < \lambda_{\text{min}} < \lambda_{\text{max}} \);

(Nonlinear impact) \( c_2 := \inf_{\| \delta \|_0 \leq K_1 n^{c_1}, \delta \neq 0} E[(Z_i^T \delta)^2]^{3/2} / E[|Z_i^T \delta|^3] > 0 \).

(A7) \( \Gamma_m \) is equally gridded with \( \tau_k - \tau_{k-1} = \epsilon_n = c_0 n^{-1} \) for \( \tau_k \in \Gamma_m (k = 1, \ldots, m) \) and a constant \( c_0 > 0 \).

Assumption (A1) requires the number of censored observations below the \( \nu \)-th quantile not to exceed \( cn^{1/2} \), which is satisfied if the lower bound of \( C \)'s support is greater than the lower bound of \( T \)'s support, a reasonable scenario in real applications. As recommended in Zheng et al. (2018), \( \nu \) is chosen such that only a small proportion of the observed survival times below the \( \nu \)-th quantile are censored. (A2) assumes that the covariates are uniformly bounded. As pointed out by Zheng et al. (2015), the global linear quantile regression model is most meaningful when the covariates are confined to a compact set to avoid crossing of the quantile functions. (A3) ensures the positiveness of \( f(t|Z) \) between \( Z^T \beta^*(\nu) \) and \( Z^T \beta^*(\tau_U) \), which is essential for the identifiability of \( \beta^*(\tau) \) for \( \tau < \tau_U \). (A4) restricts
the order of data dimensions, as well as the sparsity of $\beta^*(\tau)$, which is necessary for the convergence of the low dimensional estimator in (2) (Condition C4 in Wang et al. (2012)). (A4) also characterizes the “sure screening” property by $S$. This asymptotic property does not assess the variability of selection with a finite sample; it is crucial to account for such variability for high dimensional inference (Fei et al., 2019; Fei and Li, 2021). Also, several variable selection methods for high dimensional CQR satisfy the sure screening property in (A4) with additional mild conditions.

- L-HDCQR: by Corollary 4.1 of Zheng et al. (2018), a beta-min condition is required in addition to the set of conditions in this paper. Explicitly, there exist constants $C_1, C_2 > 0$, such that

$$\inf_{j \in S^*} \sup_{\tau \in [\tau_L, \tau_U]} |\beta_j^*(\tau)| > C_1 \exp(C_2 q\tau_U) \sqrt{q \log(p \vee n)/n} + L \sqrt{q \epsilon n}.$$ 

- AL-HDCQR: by Corollary 4.2 of Zheng et al. (2018), AL-HDCQR achieves the stronger selection consistency property, which implies the sure screening property.

- Quantile-adaptive Screening: by Theorem 3.3 of He et al. (2013), with a proper threshold value in their technical conditions, the screening procedure achieves the sure screening property.

(A5) characterizes the smoothness of $\beta^*(\tau)$. The eigenvalue condition in (A6) is the sparse Riesz condition in Zhang and Huang (2008), satisfied by many commonly used covariance structures, including the compound symmetry structure and the first order autoregressive structure (AR(1)) (Zhang and Huang, 2008). Also, the nonlinear impact condition controls the minoration of the quantile regression objective function by a quadratic function, as adopted in Zheng et al. (2018), for establishing the consistency of L-HDCQR estimator. The condition is satisfied when the covariates $Z_i$ have a log-concave density, which includes the commonly used normal distribution, Wishart distribution and Dirichlet distribution.
(Lovász and Vempala, 2007). (A7) details the fineness of $\Gamma_m$, which renders an adequate approximation to the stochastic integration in (2).

### 3.2 Theoretical properties of Fused-HDCQR

We first extend the results in Peng and Huang (2008) from a fixed $p$ to a $p$-diverges-but-less-than-$n$ case. The results are novel and critical since we allow the true model size $q = |S^*|$ to increase with $n$, while the selected $\tilde{S}^{b_S}$'s in the fused procedure vary around $S^*$. Specifically, we assume a subset $S \subset \{1, \ldots, p\}$ in Theorems 1 and 2, where $|S| \leq K_1 n^{c_1}$, $0 \leq c_1 < 1/3$ and $K_1 > 0$. Let $\hat{\beta}_S(\tau), \tau \in [\nu, \tau_U]$ be the estimator from Peng and Huang (2008) of fitting the CQR with $Z_S$ over the $\tau$-grid $\Gamma_m$.

**Theorem 1.** (Consistency with a diverging number of covariates) Under Conditions (A1)–(A7) and given a subset $S \subset \{1, \ldots, p\}$ such that $S^* \subset S$ and $|S| \leq K_1 n^{c_1}$, there exist positive constants $\zeta_1$ and $\zeta_2$ such that

$$
\sup_{\nu \leq \tau \leq \tau_U} \|\hat{\beta}_S(\tau) - \beta^*(\tau)\| \leq \zeta_1 \exp(\zeta_2)(K_1 n^{c_1-1} \log n)^{1/2}
$$

with probability at least $1 - 20\epsilon_0^{-2} K_1 n^{c_1-2}$.

**Remark 5.** From the proof of this theorem (in particular, the proofs of Propositions 1 and 2 in the Supplementary Materials that lead to this theorem), it can be seen that $\zeta_1$ and $\zeta_2$ do not depend on the choice of $S$ or $n$. Thus, $\zeta_1$ and $\zeta_2$ are universal for all possible $S$ satisfying $S^* \subset S$ and $|S| \leq K_1 n^{c_1}$.

Next, we derive the weak convergence of $\hat{\beta}_j$ for any $j \in S$.

**Theorem 2.** (Weak convergence with a diverging number of covariates) Under Conditions (A1)–(A7) and given a $S \subset \{1, \ldots, p\}$ such that $S^* \subset S$ and $|S| \leq K_1 n^{c_1}$, it holds that

$$
\sqrt{n} \left( \hat{\beta}_j(\tau) - \beta_j^*(\tau) \right)
$$

converges weakly to a mean zero Gaussian process for $\tau \in [\nu, \tau_U]$ and any $j \in S$. 

12
In high dimensional settings, the next theorem shows that the fused estimator enjoys desirable theoretical properties.

**Theorem 3.** Consider the Fused-HDCQR estimator in (3). Under assumptions (A1) – (A7), for any $j \in \{1, \ldots, p\}$,

$$\sqrt{n} \left( \hat{\beta}_j(\tau) - \beta_j^*(\tau) \right)$$

converges weakly to a mean zero Gaussian process for $\tau \in [\nu, \tau_U]$.

Our framework enables us to obtain the joint distribution of any $K$-dimensional estimated coefficients, where $K$ is a finite number. Let $K$ be the collection of the indices of $K$ covariates of interest. We can show that the weak convergence result of $\hat{\beta}_K(\tau)$, a $K$-dimensional subvector of the oracle estimator, still holds for $\tau \in [\nu, \tau_U]$, that is, $\sqrt{n}(\hat{\beta}_K(\tau) - \beta_K^*(\tau)), \tau \in [\nu, \tau_U]$ converges to a $K$-dimensional Gaussian distribution at any $\tau \in [\nu, \tau_U]$. We only need to replace $\hat{\beta}_j(\tau)$ by $\hat{\beta}_K(\tau)$ in the proof of Theorem 2 in the Appendix and slightly modify the arguments accordingly. Consequently, the term I in the proof of Theorem 3 still converges weakly to a mean zero Gaussian distribution, while the norms of items II and III are still $o_p(1)$. Therefore, Theorem 3 still holds for any $K$-dimensional subvector of $\hat{\beta}_K(\tau)$, i.e., $\sqrt{n}(\hat{\beta}_K(\tau) - \beta_K^*(\tau))$ converges to a mean zero $K$-dimensional Gaussian distribution at any $\tau \in [\nu, \tau_U]$.

As shown in the proof, the covariance function of $\hat{\beta}_j(\tau)$ depends on the unknown active set $S^*$, the unknown conditional density functions $f(t|Z)$ and $g(t|Z)$, and other unknown quantities. Thus, it is not calculable. The next section proposes an alternative model-free variance estimator based on the functional delta method and the multi-sample splitting properties (Efron, 2014; Fei and Li, 2021).
4 A Variance Estimator via the Functional Delta Method

Let $J_{bi} \in \{0, 1\}$ indicate whether the $i^{th}$ observation is in the $b^{th}$ sub-sample $D_b^2$, and $J_i = B^{-1} \sum_{b=1}^{B} J_{bi}$. For each $i = 1, \ldots, n$, we define the re-sampling covariance between $J_{bi}$ and $\tilde{\beta}_j^b(\tau_k)$ at $\tau_k \in \Gamma_m$ as

$$s_{ij}(\tau_k) = \frac{1}{B} \sum_{b=1}^{B} (J_{bi} - J_i) \left( \tilde{\beta}_j^b(\tau_k) - \hat{\beta}_j(\tau_k) \right).$$

Define $S_j(\tau_k) = (s_{1j}(\tau_k), s_{2j}(\tau_k), \ldots, s_{nj}(\tau_k))^T$ and let $n_1 = |D_2^b|$. It follows that the covariance between $\hat{\beta}_j(\tau_k)$ and $\hat{\beta}_j(\tau_\ell)$ can be consistently estimated by

$$\hat{\text{Cov}}_j(\tau_k, \tau_\ell) = \frac{n-1}{n} \left( \frac{n}{n-n_1} \right)^2 \sum_{i=1}^{n} s_{ij}(\tau_k)s_{ij}(\tau_\ell) = \frac{n(n-1)}{(n-n_1)^2} S_j^T(\tau_k)S_j(\tau_\ell),$$

where the multiplier $n(n-1)/(n-n_1)^2$ is a finite-sample correction for sub-sampling (Wager and Athey, 2018). In particular, by taking $\tau_\ell = \tau_k$, a variance estimator for $\hat{\beta}_j(\tau_k)$ is

$$\hat{V}_j(\tau_k) = \frac{n(n-1)}{(n-n_1)^2} S_j^T(\tau_k)S_j(\tau_k). \quad (5)$$

As in Wager and Athey (2018), it follows that $\hat{V}_j(\tau_k)/\text{Var}\left( \hat{\beta}_j(\tau_k) \right) \xrightarrow{p} 1$ with $n, B \to \infty$. Furthermore, for a finite $B$, we propose a bias corrected version of (5):

$$\hat{V}_j^B(\tau_k) = \hat{V}_j(\tau_k) - \frac{nn_1}{B(n-n_1)} \left\{ B^{-1} \sum_{b=1}^{B} \left( \tilde{\beta}_j^b(\tau_k) - \hat{\beta}_j(\tau_k) \right)^2 \right\}, \quad \tau_k \in \Gamma_m. \quad (6)$$

The correction term in (6) is a suitable multiplier of the re-sampling variance of $\tilde{\beta}_j^b(\tau_k)$’s, and converges to zero with $n \to \infty$ and $n_1 = O(n)$. Thus, the two variance estimators in (5) and (6) are asymptotically equal. However, $\hat{V}_j(\tau_k)$ in (5) requires $B$ to be of order $n^{3/2}$ to reduce the Monte Carlo noise below the sampling noise, while $\hat{V}_j^B(\tau_k)$ in (6) only requires $B$ to be of order $n$ to achieve the same (Wager et al., 2014).
Since \( \hat{\beta}_j(\tau) \) converges weakly to a Gaussian process by Theorem 3, and our variance estimators are consistent on the grid points, we define an asymptotic \( 100(1-\alpha)\% \) point-wise confidence interval for \( \beta^*_j(\tau_k) \) at any \( \tau_k \in \Gamma_m \) as

\[
\left( \hat{\beta}_j(\tau_k) - \Phi^{-1}(1-\alpha/2)\sqrt{\hat{V}_B^j(\tau_k)}, \hat{\beta}_j(\tau_k) + \Phi^{-1}(1-\alpha/2)\sqrt{\hat{V}_B^j(\tau_k)} \right),
\]

where \( \hat{V}_B^j(\tau_k) \) is as defined in (6), and \( \Phi \) is the standard normal cumulative distribution function. The \( p \)-value of testing \( H_0: \beta^*_j(\tau_k) = 0 \) for a \( \tau_k \in \Gamma_m \) is

\[
2 \times \left\{ 1 - \Phi \left( |\hat{\beta}_j(\tau_k)|/\sqrt{\hat{V}_B^j(\tau_k)} \right) \right\}.
\]

5 Simulation Studies

In various settings, we compare the proposed method, Fused-HDCQR (referred to as “Fused” in the tables and figures hereafter), with some competing methods in quantile regression or high dimensional inference. These methods include Wang et al. (2012) (“W12”) and Fan et al. (2014) (“F14”) for quantile regression; Zheng et al. (2018) (“Z18”) for censored quantile regression; and Meinshausen et al. (2009) (“M09”) for inference with aggregated \( p \)-values from multi-sample splittings.

In the simulations and the later data analysis, we choose L-HDCQR described in Section 3 as the variable selection tool for Fused-HDCQR. We also explore the feasibility of using other alternatives for variable selection, such as Fan et al. (2009) (“F09”) and M09.

When implementing Fused-HDCQR, we specify the number of splits as \( B = 300 \), the quantile interval as \([\nu, \tau_U] = [0.1, 0.8]\), and the grid length as \( m = n/\log p \). As regards the selection of tuning parameters, Theorems 1 and 2 suggest that our procedure not be sensitive to tuning parameters as long as they can ensure sure screening. In practical settings, we recommend to select tuning parameters by minimizing the 5-fold cross-validation error as in (4), which may help achieve sure screening and works well in our simulations. We study the following examples with sparse non-zero effects, some of which are heterogeneous.
**Example 1.** The event times are generated by

\[
\log T_i = \tilde{Z}_i^T b + \varepsilon_i, \quad i = 1, \ldots, n,
\]

where the coefficient vector \( b \) is sparse with \( b_{20} = 0.5, b_{40} = 1, b_{60} = 1.5, b_j = 0 \) for all other \( j \)'s, and \( \varepsilon_i \sim N(0,1) \). Therefore, the true coefficients satisfy \( \beta^*(\tau) = (Q_\varepsilon(\tau), b^T)^T \) for all \( \tau \in (0,1) \), where \( Q_\varepsilon(\tau) \), the \( \tau \)-th quantile of the distribution of \( \varepsilon \), is the intercept. The covariates \( \tilde{Z}_{j,i} \)'s are i.i.d. from Unif\((-1,1)\) and are independent across \( j \in \{1, \ldots, p\} \). The censoring times are generated independently as \( \log C_i \sim N(3, 17.25) \), giving a censoring rate around 25%.

**Example 2.** The event times follow

\[
\log T_i = \tilde{Z}_i^T b + 1.5 \tilde{Z}_{3,i} \varepsilon_i,
\]

where \( b_{20} = 1, b_{40} = 1.5, b_{60} = 2, b_j = 0 \) for all other \( j \)'s, and \( \varepsilon_i \sim N(0,1) \). We first generate \( \tilde{Z}_i \sim N_p(0, \Sigma) \) with an AR(1) \( \Sigma = (\sigma_{k\ell})_{p \times p} \), where \( \sigma_{k\ell} = 0.3^{|k-\ell|} \), and then let \( Z_i = \tilde{Z}_i \), except that the third covariate \( \tilde{Z}_{3,i} = |\tilde{Z}_{3,i}| + 0.5 \). Thus, \( \beta^*_1(\tau) = 0, \beta^*_4(\tau) = 1.5 Q_\varepsilon(\tau) \), and \( \beta^*_{j+1}(\tau) = b_j \) for all other \( j \)'s. The censoring times are generated independently as \( \log C_i \sim N(4, 17.25) \), giving a censoring rate around 23%.

**Example 3.** The event times follow

\[
\log T_i = \tilde{Z}_i^T b + \phi_1(\xi_i) \tilde{Z}_{1,i} + \phi_4(\xi_i) \tilde{Z}_{4,i},
\]

where \( b_{20} = 1, b_{40} = 1.5, b_{60} = 2, b_j = 0 \) for all other \( j \)'s, \( \xi_i \sim N(0,1) \), and \( \phi_1, \phi_4 \) are monotone functions as the dashed lines in Figure 1, both are continuous with zero and non-zero pieces over \( \tau \). We first generate \( \tilde{Z}_i \sim N_p(0, \Sigma) \) as in Example 2, and then let \( \tilde{Z}_i = \tilde{Z}_i \), except that \( \tilde{Z}_{1,i} = |\tilde{Z}_{1,i}| + 0.5 \) and \( \tilde{Z}_{4,i} = |\tilde{Z}_{4,i}| + 0.5 \). Therefore, \( \beta^*_1(\tau) = 0, \beta^*_2(\tau) = \phi_1(\tau), \beta^*_5(\tau) = \phi_4(\tau) \), and \( \beta^*_{j+1}(\tau) = b_j \) for all other \( j \)'s. The censoring times are generated independently as \( \log C_i \sim N(6, 17.25) \), which gives a censoring rate around 20%.
For each of these examples, we set \((n,p)=(300,1000)\) and \((700,1000)\) to study the impacts of the sample size and the number of variables on the performance, and, in particular, how the methods fare when \(p > n\). In Example 3, which mimics the real data example in Section 6 most closely, we have also explored \((n,p) = (700, 2000)\), which is roughly equal to the dimension of the real dataset. For every parameter configuration, a total of 100 independent datasets are generated, and we report the average results based on these replications. We choose 100 replications because the penalized methods for high dimensional CQR in general take much computing time (Table 5).

We first evaluate the feasibility of using various variable selection tools for our proposed method. Comparisons of true positives and false negatives among F09, M09, and L-HDCQR under Examples 1–3 are reported in Table 1. F09 presents a subpar performance because, by taking intersections of variables selected from different partitions of data, it tends to miss out some true signals and thus have fewer true positives. In contrast, L-HDCQR retains more true positives than both F09 and M09, while having more false positives. Because our method requires the variable selection step to include the true signals with high probability, even at the cost of some false positives, we use L-HDCQR as the screening tool for our method.

We next compare the performance of Fused-HDCQR with other high dimensional quantile regression methods at \(\tau = .25, .5, .75\) under Example 1. As a benchmark for comparisons, we also compute the oracle estimates based on the true model (with \(S^*\) known). Since W12, F14, and Z18 only provide coefficient estimates without standard errors (SEs), we only report the estimation biases for them, while reporting the average SEs, empirical standard deviations (SDs) and coverage probabilities of the confidence intervals for our method. Table 2 shows that Fused-HDCQR presents the smallest biases, which are comparable to those of the oracle estimates. In contrast, Z18 has smaller biases when the sample size is large, and larger biases otherwise, while W12 and F14 incur substantial biases since they are
not designed for censored data. Moreover, the average SEs based on Fused-HDCQR agree with the empirical SDs of the estimates. The consistent estimates of coefficients and SEs obtained by Fused-HDCQR lead to proper coverage probabilities around the 0.95 nominal level. In addition, the coverage probabilities improve as \( n \) increases.

Table 2 also concerns the power for detection of signals. Since W12, F14, and Z18 cannot draw inference and, in general, there is a lack of literature that deals with inference for HDCQR, we compare our method with the aggregated \( p \)-value approach (M09) in the quantile setting, though M09 originated from linear regression. The results indicate that Fused-HDCQR outperforms M09, presenting more power when the effect size is moderate or large.

Table 3 summarizes the results from Example 2 with the heterogeneous effect \( \beta_4 \) varying with \( \tau \). We compare the estimation accuracy between Fused-HDCQR and Z18, as well as the statistical power between Fused-HDCQR and M09. Again, Fused-HDCQR presents smaller biases than Z18 and a higher power than M09. To assess whether the tuning parameters selected as in Remark 3 help the variable selection method (L-HDCQR), used by Fused-HDCQR, satisfy assumption (A4) in Section 3, we report the selection frequency of each signal variable in Table 3 (and also in Table 4), and observe that the selection frequency increases as the sample size increases, hinting that assumption (A4) may be satisfied with these selected tuning parameters.

Table 4 summarizes the results based on Example 3. For the two heterogeneous effects \( \beta_2 \) and \( \beta_5 \) that vary with \( \tau \), their estimation biases of Fused-HDCQR become smaller and the estimated SEs are closer to the empirical ones as \( n \) increases. Figure 1 shows that the Fused-HDCQR estimates in general agree with the oracle estimates and the truth, except at the non-smooth change points, and have narrower confidence intervals with a larger \( n \), where the vertical bars are the average confidence intervals of the \( \tau \) grid.

In regards to the choice of \( B \) in the variance computation, our numerical experience
suggests that it may be sufficient to use a $B$ that is of the same order of the sample size, even when $n$ is less than $p$. This coincides with the note under (6) that $B$ is only required to be of order $n$ to reduce the Monte Carlo noise below the sampling noise.

Finally, we compare the computation intensity among Z18, M09, W12, F14, and Fused-HDCQR under Example 1 and report in Table 5 the computing time on average per dataset. Our method is the most computationally intensive, because it involves multiple data-splittings and draws inferences on all of the $p$ coefficients. However, by utilizing parallel computing, we have managed to reduce the computational time to the same order of Z18, W12, and F14 that are based on penalized regression. The R code used for generating the simulation results can be accessed via https://github.com/feizhe/HDCQR_Paper.

6 Application to the Boston Lung Cancer Survival Cohort (BLCSC)

Detection of molecular profiles related to cancer survival can aid personalized treatment in prolonging patients’ survival and improving their quality of life. In a subset of BLCSC samples, 674 lung cancer patients were measured with survival times, along with 40,000 SNPs and clinical indicators, such as lung cancer subtypes (adenocarcinoma, squamous cell carcinoma, or others), cancer stages (1-4), age, gender, education level (≤ high school or > high school) and smoking status (active or non-active smokers); see Table 6 for patients’ characteristics. The censoring rate was 23% and a total of 518 deaths were observed during the followup period, with the observed followup time varying from 13 to 8,584 days.

We could have included all 40,000 SNPs in our analysis. However, for more statistical power, we opt for the targeted gene approach by focusing on 2,002 SNPs residing in 14 genes identified to be cancer related, namely, ALK, BRAF, BRCA1, EGFR, ERBB2, ERCC1, KRAS, MET, PIK3CA, RET, ROS1, RRM1, TP53, and TYMS (Brose et al., 2002; Toyooka
et al., 2003; Paez et al., 2004; Soda et al., 2007). Pinpointing the effects of individual loci
within the targeted genes is helpful for understanding disease mechanisms (Evans et al.,
2011; D’Antonio et al., 2019) and designing gene therapies (Pâques and Duchateau, 2007;
Hanawa et al., 2004). We also adjust for patients’ clinical and environmental characteristics
listed in Table 6, which gives a total of $p = 2,011$ predictors.

We apply Fused-HDCQR to compute the point estimates (3) and the variance estimates
(6). We set the quantile interval to be $[0.2, 0.7]$, which is wide enough to cover high- and
low-risk groups and, in the meantime, ensures the quantile parameters be estimable in the
presence of censoring (Zheng et al., 2015). We choose the lower bound $\tau_0 = \nu = 0.1$ to
circumvent the singularity problem with CQR at $\tau = 0$, because few ($< 2\%$) observations
are censored below the 0.1-th quantile. With $\epsilon_n = 0.01$, we form the $\tau$-grid $\Gamma_m$ of length
$m = 61$. We set $B = 750$ as the number of re-samples, which is sufficiently large and
comparable to the sample size. To determine the tuning parameter $\lambda_n$ in L-HDCQR for
selection, we use 5-fold cross-validation as specified in Remark 3.

For ease of presentation, we summarize the results evaluated at 6 quantile levels, $\tau =
0.2, 0.3, \ldots, 0.7$, instead of the whole grid $\Gamma_m$. To highlight the findings of the high-risk
group, we rank all SNPs based on their $p$-values at $\tau = 0.2$. In particular, after Bonferroni
correction for multiple testing, there are 83 significant SNPs for $\tau = 0.2$ with the overall
type I error of $\alpha = 0.05$. Our method estimates the coefficients and the $p$-values for
all predictors, and we only present the results for the patient characteristics, the top 10
significant SNPs, and the 3 least significant SNPs in Figure 2 and Table 7. The estimated
coefficient of active smoking drops from $-0.42$ ($p = 0.0011$) to $-0.53$ ($p = 0.0005$) as $\tau$
changes from 0.2 to 0.5, and then increases to $-0.31$ ($p = 0.038$) as $\tau$ changes to 0.7,
suggesting that active smoking might be more harmful to the high- and median-risk groups
than the low-risk group of patients. The most significant SNP at $\tau = 0.2$ is AX.37793583.T,
which remains significant throughout $\tau = 0.2$ to $\tau = 0.7$. However, its estimated coefficient

20
decreases from 2.75 ($\tau = 0.2$) to 1.39 ($\tau = 0.7$), indicating its heterogeneous impacts on survival, i.e., stronger protective effect at lower quantiles and vice versa.

The effects of some SNPs are nearly zero for higher quantiles. For example, the estimated coefficient of AX.15207405.G decreases from 2.03 ($\tau = 0.2$; $p = 10^{-24}$) to $-0.05$ ($\tau = 0.7$; $p = 0.92$), with the estimated standard error increasing from 0.20 to 0.48. Similarly, the estimated coefficient of AX.40182999.A decreases from 1.5 ($\tau = 0.2$; $p = 9.6 \times 10^{-13}$) to $-0.01$ ($\tau = 0.7$; $p = 0.96$). The results again hint at heterogeneous SNP effects in various risk groups, which cannot be detected using traditional Cox models.

Finally, our results shed light on the roles of SNPs in the high-risk group (i.e., lower quantiles). Specifically, we map the 83 SNPs with significant effects at the 0.2-th quantile by Fused-HDCQR to the corresponding genes and rank the genes by the number of significant SNPs (over total number of SNPs for each gene in the parenthesis), which are TP53 (14/321), RRM1 (14/174), ERCC1 (10/167), BRCA1 (10/114), ALK (8/163), ROS1 (5/294), EGFR (5/261), ERBB2 (4/167), and 6 other genes with numbers of significant SNPs less than 4. While these genes were reported to be associated with lung cancer (Toyooka et al., 2003; Takeuchi et al., 2012; Rosell et al., 2011; Lord et al., 2002; Zheng et al., 2007; Sasaki et al., 2006; Brose et al., 2002), our analysis provides more detailed information as to which SNPs and locations of the genes are jointly associated with the lung cancer survival, as well as the estimated effects and uncertainties. Analysis of heterogeneous SNP effects has been gaining increasing research attention in lung cancer research (McKay et al., 2017; Dong et al., 2012; Huang et al., 2009), and beyond it (Garcia-Closas et al., 2008; Cheng et al., 2010; Gulati et al., 2014).

7 Conclusions

Our proposed procedure involves repeated estimates from low dimensional CQRs, which are computationally straightforward and can be efficiently implemented with parallel comput-
ing. We require the variable selection to possess a sure screening property as in condition (A4). This seems to be supported by our simulations, which find our procedure works well when the variable selection method can select a superset of the true model with high probability. Our condition is much weaker than a condition of selection consistency as specified in Fei et al. (2019).

For the selection of $B$, we recommend $B$ to be in the same order of the sample size $n$. Smaller $B$ might not affect coefficient estimation much; but it might yield inaccurate estimated standard errors, leading to incorrect inferences. In addition, we opt to define $\Gamma_m$ by setting the grid as $n/\log p$ equally spaced points between $\tau_0$ and $\tau_U$. This may cover the quantile interval well, with reasonable computation efficiency.

There are open questions to be addressed. First, substantial work is needed for handling highly correlated predictors as the performance of our method, like the other competing methods, deteriorates when correlations among predictors become stronger. Second, it is of interest to investigate an alternative method when the sparsity condition fails. For example, it is challenging to find an effective strategy to draw inference when a non-negligible portion of predictors have small but non-zero effects. We will pursue them elsewhere.

**Acknowledgements**

We are deeply grateful toward the Editor, the AE and the two referees for their constructive comments and suggestions that have improved the manuscript substantially. We thank our long-term collaborator, David Christiani of Harvard Medical School, for providing the BLCSC data. The work is partially supported by grants from NIH (5R01CA249096 and U01CA209414).
References

Belloni, A., V. Chernozhukov, and K. Kato (2019). Valid post-selection inference in high-dimensional approximately sparse quantile regression models. *Journal of the American Statistical Association* 114(526), 749–758.

Berk, R., L. Brown, A. Buja, K. Zhang, and L. Zhao (2013). Valid post-selection inference. *The Annals of Statistics* 41(2), 802–837.

Brose, M. S., P. Volpe, M. Feldman, M. Kumar, I. Rishi, R. Gerrero, et al. (2002). BRAF and RAS mutations in human lung cancer and melanoma. *Cancer research* 62(23), 6997–7000.

Bühlmann, P., M. Kalisch, and L. Meier (2014). High-dimensional statistics with a view toward applications in biology. *Annual Review of Statistics and Its Application* 1, 255–278.

Cheng, I., S. J. Plummer, C. Neslund-Dudas, E. A. Klein, G. Casey, B. A. Rybicki, and J. S. Witte (2010). Prostate cancer susceptibility variants confer increased risk of disease progression. *Cancer Epidemiology and Prevention Biomarkers* 19(9), 2124–2132.

Christiani, D. C. (2017). The Boston lung cancer survival cohort. [http://grantome.com/grant/NIH/U01-CA209414-01A1](http://grantome.com/grant/NIH/U01-CA209414-01A1). [Online; accessed November 27, 2018].

D’Antonio, M., J. Reyna, D. Jakubosky, M. K. Donovan, M.-J. Bonder, et al. (2019). Systematic genetic analysis of the MHC region reveals mechanistic underpinnings of HLA type associations with disease. *eLife* 8, e48476.

Dong, J., Z. Hu, Y. Shu, S. Pan, W. Chen, Y. Wang, et al. (2012). Potentially functional polymorphisms in dna repair genes and non-small-cell lung cancer survival: A pathway-based analysis. *Molecular carcinogenesis* 51(7), 546–552.
Efron, B. (2014). Estimation and accuracy after model selection. *Journal of the American Statistical Association* 109(507), 991–1007.

Evans, D. M., C. C. Spencer, J. J. Pointon, Z. Su, D. Harvey, G. Kochan, et al. (2011). Interaction between ERAP1 and HLA-B27 in ankylosing spondylitis implicates peptide handling in the mechanism for HLA-B27 in disease susceptibility. *Nature genetics* 43(8), 761–767.

Fan, J., Y. Fan, and E. Barut (2014). Adaptive robust variable selection. *The Annals of Statistics* 42(1), 324–351.

Fan, J., R. Samworth, and Y. Wu (2009). Ultrahigh dimensional feature selection: beyond the linear model. *Journal of Machine Learning Research* 10, 2013–2038.

Fang, E. X., Y. Ning, and H. Liu (2017). Testing and confidence intervals for high dimensional proportional hazards models. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)* 79(5), 1415–1437.

Fei, Z. and Y. Li (2021). Estimation and inference for high dimensional generalized linear models: A splitting and smoothing approach. *Journal of Machine Learning Research* 22(58), 1–32.

Fei, Z., J. Zhu, M. Banerjee, and Y. Li (2019). Drawing inferences for high-dimensional linear models: A selection-assisted partial regression and smoothing approach. *Biometrics* 75(2), 551–561.

Fleming, T. R. and D. P. Harrington (2011). *Counting Processes and Survival Analysis*, Volume 169. John Wiley & Sons.

Garcia-Closas, M., P. Hall, H. Nevanlinna, K. Pooley, J. Morrison, D. A. Richesson, et al.
(2008). Heterogeneity of breast cancer associations with five susceptibility loci by clinical and pathological characteristics. *PLoS genetics* 4(4), e1000054.

Gulati, S., P. Martinez, T. Joshi, N. J. Birkbak, C. R. Santos, A. J. Rowan, et al. (2014). Systematic evaluation of the prognostic impact and intratumour heterogeneity of clear cell renal cell carcinoma biomarkers. *European urology* 66(5), 936–948.

Hanawa, H., P. W. Hargrove, S. Kepes, D. K. Srivastava, A. W. Nienhuis, and D. A. Persons (2004). Extended β-globin locus control region elements promote consistent therapeutic expression of a γ-globin lentiviral vector in murine β-thalassemia. *Blood* 104(8), 2281–2290.

He, X., L. Wang, and H. G. Hong (2013). Quantile-adaptive model-free variable screening for high-dimensional heterogeneous data. *The Annals of Statistics* 41(1), 342–369.

Ho, D. S. W., W. Schierding, M. Wake, R. Saffery, and J. O’Sullivan (2019). Machine learning SNP based prediction for precision medicine. *Frontiers in Genetics* 10, 267.

Hong, H. G., D. C. Christiani, and Y. Li (2019). Quantile regression for survival data in modern cancer research: expanding statistical tools for precision medicine. *Precision clinical medicine* 2(2), 90–99.

Huang, Y.-T., R. S. Heist, L. R. Chirieac, X. Lin, V. Skaug, S. Zienolddiny, et al. (2009). Genome-wide analysis of survival in early-stage non–small-cell lung cancer. *Journal of clinical oncology* 27(16), 2660–2667.

Javanmard, A. and A. Montanari (2014). Confidence intervals and hypothesis testing for high-dimensional regression. *Journal of Machine Learning Research* 15(1), 2869–2909.

Kelley, M. J., S. Li, and D. H. Harpole (2001). Genetic analysis of the β-tubulin gene,
tubb, in non-small-cell lung cancer. *Journal of the National Cancer Institute* 93(24), 1886–1888.

Klee, V. and G. J. Minty (1972). How good is the simplex algorithm. *Inequalities* 3(3), 159–175.

Koenker, R. (2005). *Quantile Regression*. Cambridge University Press.

Koenker, R. and G. Bassett Jr (1978). Regression quantiles. *Econometrica: Journal of the Econometric Society* 46(1), 33–50.

Kong, S., Z. Yu, X. Zhang, and G. Cheng (2021). High dimensional robust inference for cox regression models using de-sparsified lasso. *Scandinavian Journal of Statistics*. doi: 10.1111/sjos.12543.

Korpanty, G. J., D. M. Graham, M. D. Vincent, and N. B. Leighl (2014). Biomarkers that currently affect clinical practice in lung cancer: EGFR, ALK, MET, ROS-1, and KRAS. *Frontiers in oncology* 4, 204.

Lee, J. D., D. L. Sun, Y. Sun, and J. E. Taylor (2016). Exact post-selection inference, with application to the lasso. *The Annals of Statistics* 44(3), 907–927.

Lord, R. V., J. Brabender, D. Gandara, V. Alberola, C. Camps, M. Domine, et al. (2002). Low ERCC1 expression correlates with prolonged survival after cisplatin plus gemcitabine chemotherapy in non-small cell lung cancer. *Clinical Cancer Research* 8(7), 2286–2291.

Lovász, L. and S. Vempala (2007). The geometry of logconcave functions and sampling algorithms. *Random Structures & Algorithms* 30(3), 307–358.

McKay, J. D., R. J. Hung, Y. Han, X. Zong, R. Carreras-Torres, D. C. Christiani, et al. (2017). Large-scale association analysis identifies new lung cancer susceptibility loci and
heterogeneity in genetic susceptibility across histological subtypes. *Nature genetics* 49(7), 1126–1132.

Meinshausen, N., L. Meier, and P. Bühlmann (2009). P-values for high-dimensional regression. *Journal of the American Statistical Association* 104(488), 1671–1681.

Moon, C., Y. Oli, and J. A. Roth (2003). Current status of gene therapy for lung cancer and head and neck cancer. *Clinical cancer research* 9(14), 5055–5067.

Ning, Y. and H. Liu (2017). A general theory of hypothesis tests and confidence regions for sparse high dimensional models. *The Annals of Statistics* 45(1), 158–195.

Paez, J. G., P. A. Jänne, J. C. Lee, S. Tracy, H. Greulich, S. Gabriel, P. Herman, F. J. Kaye, N. Lindeman, T. J. Boggon, et al. (2004). Egfr mutations in lung cancer: correlation with clinical response to gefitinib therapy. *Science* 304(5676), 1497–1500.

Pâques, F. and P. Duchateau (2007). Meganucleases and dna double-strand break-induced recombination: perspectives for gene therapy. *Current gene therapy* 7(1), 49–66.

Peng, L. and Y. Huang (2008). Survival analysis with quantile regression models. *Journal of the American Statistical Association* 103(482), 637–649.

Portnoy, S. (2003). Censored regression quantiles. *Journal of the American Statistical Association* 98(464), 1001–1012.

Powell, J. L. (1986). Censored regression quantiles. *Journal of econometrics* 32(1), 143–155.

Risch, A. and C. Plass (2008). Lung cancer epigenetics and genetics. *International Journal of Cancer* 123(1), 1–7.

Rosell, R., M. A. Molina, C. Costa, S. Simonetti, A. Gimenez-Capitan, J. Bertran- Alamillo, et al. (2011). Pretreatment EGFR T790M mutation and BRCA1 mRNA expression in
erlotinib-treated advanced non–small-cell lung cancer patients with EGFR mutations. 
*Clinical Cancer Research* 17(5), 1160–1168.

Sasaki, H., S. Shimizu, K. Endo, M. Takada, M. Kawahara, H. Tanaka, et al. (2006). EGFR and erbB2 mutation status in japanese lung cancer patients. *International Journal of Cancer* 118(1), 180–184.

Shows, J. H., W. Lu, and H. H. Zhang (2010). Sparse estimation and inference for censored median regression. *Journal of Statistical Planning and Inference* 140(7), 1903–1917.

Soda, M., Y. L. Choi, M. Enomoto, S. Takada, Y. Yamashita, S. Ishikawa, et al. (2007). Identification of the transforming eml4–alk fusion gene in non-small-cell lung cancer. *Nature* 448(7153), 561–566.

Takeuchi, K., M. Soda, Y. Togashi, R. Suzuki, S. Sakata, S. Hatano, et al. (2012). RET, ROS1 and ALK fusions in lung cancer. *Nature Medicine* 18(3), 378–381.

Toyooka, S., T. Tsuda, and A. F. Gazdar (2003). The TP53 gene, tobacco exposure, and lung cancer. *Human Mutation* 21(3), 229–239.

Van de Geer, S., P. Bühlmann, Y. Ritov, and R. Dezeure (2014). On asymptotically optimal confidence regions and tests for high-dimensional models. *The Annals of Statistics* 42(3), 1166–1202.

Wager, S. and S. Athey (2018). Estimation and inference of heterogeneous treatment effects using random forests. *Journal of the American Statistical Association* 113(523), 1228–1242.

Wager, S., T. Hastie, and B. Efron (2014). Confidence intervals for random forests: The jackknife and the infinitesimal jackknife. *Journal of Machine Learning Research* 15(1), 1625–1651.
Wang, L., Y. Wu, and R. Li (2012). Quantile regression for analyzing heterogeneity in ultra-high dimension. *Journal of the American Statistical Association 107*(497), 214–222.

Wei, L.-J. (1992). The accelerated failure time model: a useful alternative to the Cox regression model in survival analysis. *Statistics in medicine 11*(14-15), 1871–1879.

Yamamoto, H., H. Shigematsu, M. Nomura, W. W. Lockwood, M. Sato, N. Okumura, et al. (2008). Pik3ca mutations and copy number gains in human lung cancers. *Cancer research 68*(17), 6913–6921.

Zhang, C.-H. and J. Huang (2008). The sparsity and bias of the Lasso selection in high-dimensional linear regression. *The Annals of Statistics 36*(4), 1567–1594.

Zhang, C.-H. and S. S. Zhang (2014). Confidence intervals for low dimensional parameters in high dimensional linear models. *Journal of the Royal Statistical Society: Series B (Statistical Methodology) 76*(1), 217–242.

Zhao, S. D. and Y. Li (2012). Principled sure independence screening for cox models with ultra-high-dimensional covariates. *Journal of Multivariate Analysis 105*(1), 397–411.

Zheng, Q., C. Gallagher, and K. Kulasekera (2013). Adaptive penalized quantile regression for high dimensional data. *Journal of Statistical Planning and Inference 143*(6), 1029–1038.

Zheng, Q., L. Peng, and X. He (2015). Globally adaptive quantile regression with ultra-high dimensional data. *The Annals of Statistics 43*(5), 2225–2258.

Zheng, Q., L. Peng, and X. He (2018). High dimensional censored quantile regression. *The Annals of Statistics 46*(1), 308–343.
Zheng, Z., T. Chen, X. Li, E. Haura, A. Sharma, and G. Bepler (2007). DNA synthesis and repair genes RRM1 and ERCC1 in lung cancer. *New England Journal of Medicine* 356(8), 800–808.

Zhu, Q.-G., S.-M. Zhang, X.-X. Ding, B. He, and H.-Q. Zhang (2017). Driver genes in non-small cell lung cancer: Characteristics, detection methods, and targeted therapies. *Oncotarget* 8(34), 57680–57692.
Table 1: Summary of variable selection results based on the simulated datasets.

| (n, p)     | CR  | q | TP     | FP     |
|------------|-----|---|--------|--------|
| Example 1  |     |   | L-HDCQR | M09    |
| (300,1000) | 0.25| 3 | 2.67   | 2.12   |
| (700,1000) | 0.25| 3 | 2.98   | 2.78   |
| Example 2  |     |   | L-HDCQR | M09    |
| (300,1000) | 0.22| 4 | 3.60   | 3.58   |
| (700,1000) | 0.23| 4 | 3.99   | 3.99   |
| (300,1000) | 0.20| 5 | 3.82   | 3.63   |
| Example 3  |     |   | L-HDCQR | M09    |
| (700,1000) | 0.20| 5 | 4.81   | 4.77   |
| (700,2000) | 0.19| 5 | 4.78   | 4.76   |

Note: CR, average censoring rate; \(q = |S^*|\); TP, average true positives; FP, average false positives; M09, Meinshausen et al. (2009); F09, Fan et al. (2009); L-HDCQR, Zheng et al. (2018).
|        | Bias | EmpSD | SE  | Cov  | Power |
|--------|------|-------|-----|------|-------|
|        | Oracle | Fused | Z18 | F14  | W12   | Fused | M09 |
|        |       |       |     |      |       |       |     |
| $n = 300, p = 1000$ |       |       |     |      |       |       |     |
| $\beta_{21} = 0.5$ | 0.02 | 0.02 | -0.38 | -0.50 | -0.48 | 0.14 | 0.13 | 0.93 | 0.97 | 0.06 |
|         | 0.01 | 0.01 | -0.24 | -0.49 | -0.48 | 0.12 | 0.13 | 0.95 | 0.98 | 0.04 |
|         | -0.01 | -0.01 | -0.13 | -0.50 | -0.48 | 0.12 | 0.13 | 0.96 | 1.00 | 0.02 |
| $\beta_{41} = 1$ | -0.00 | -0.00 | -0.03 | -0.68 | -0.32 | 0.14 | 0.12 | 0.92 | 1.00 | 0.98 |
|         | 0.02 | 0.01 | -0.01 | -0.70 | -0.30 | 0.17 | 0.14 | 0.93 | 1.00 | 0.94 |
|         | -0.00 | 0.01 | 0.00 | -0.92 | -0.24 | 0.12 | 0.13 | 0.92 | 1.00 | 1.00 |
| $\beta_{61} = 1.5$ | 0.00 | 0.01 | 0.01 | -0.64 | -0.25 | 0.11 | 0.13 | 0.97 | 1.00 | 1.00 |
|         | 0.02 | 0.01 | 0.02 | -0.70 | -0.25 | 0.13 | 0.14 | 0.95 | 1.00 | 1.00 |
| $n = 700, p = 1000$ |       |       |     |      |       |       |     |
| $\beta_{21} = 0.5$ | -0.02 | -0.01 | -0.01 | -0.47 | -0.23 | 0.09 | 0.08 | 0.92 | 1.00 | 0.56 |
|         | -0.01 | -0.01 | -0.01 | -0.39 | -0.22 | 0.08 | 0.08 | 0.89 | 1.00 | 0.65 |
|         | 0.00 | 0.00 | 0.04 | -0.53 | -0.17 | 0.09 | 0.08 | 0.91 | 1.00 | 1.00 |
| $\beta_{41} = 1$ | -0.00 | 0.00 | 0.03 | -0.49 | -0.19 | 0.09 | 0.08 | 0.90 | 1.00 | 1.00 |
|         | -0.01 | -0.01 | 0.01 | -0.53 | -0.18 | 0.08 | 0.10 | 0.87 | 1.00 | 1.00 |
|         | 0.01 | 0.01 | 0.06 | -0.54 | -0.21 | 0.10 | 0.09 | 0.93 | 1.00 | 1.00 |
| $\beta_{61} = 1.5$ | 0.01 | 0.01 | 0.03 | -0.62 | -0.21 | 0.08 | 0.08 | 0.93 | 1.00 | 1.00 |
|         | -0.00 | 0.00 | 0.03 | -0.71 | -0.21 | 0.07 | 0.09 | 0.94 | 1.00 | 1.00 |

Note: Each $\beta$ has three rows corresponding to $\tau = .25, .5, .75$ from the top to bottom; EmpSD, empirical standard deviation; SE, average standard error; Cov, coverage probability; Oracle, Oracle estimator; Z18, Zheng et al. (2018); F14, Fan et al. (2014); W12, Wang et al. (2012); M09, Meinshausen et al. (2009).
### Table 3: Results of Example 2 based on the simulated datasets.

|        | Bias | EmpSD | SE | Cov | Freq | Power |
|--------|------|-------|----|-----|------|-------|
|        | Oracle | Fused | Z18 | Fused |       | Fused | M09  |
|        |       |       |     |      |       |       |
| $n = 300, \ p = 1000$ |       |       |     |      |       |       |
| $\beta_4 = 1.5Q_\varepsilon(\tau)$ | 0.01 | 0.13 | 0.29 | 0.32 | 0.31 | 0.88 | 0.82 | 0.16 |
| | -0.05 | -0.07 | 0.06 | 0.33 | 0.29 | 0.90 | 0.73 | 0.11 | 0.00 |
| | 0.01 | -0.14 | -0.05 | 0.31 | 0.34 | 0.82 | 0.62 | 0.10 |
| | -0.01 | -0.01 | -0.01 | 0.14 | 0.13 | 0.90 | 1.00 | 0.88 |
| $\beta_{21} = 1$ | -0.03 | -0.01 | -0.05 | 0.12 | 0.12 | 0.91 | 0.69 | 1.00 | 0.92 |
| | -0.01 | -0.00 | -0.02 | 0.14 | 0.13 | 0.92 | 1.00 | 0.84 |
| | 0.01 | 0.01 | 0.03 | 0.13 | 0.13 | 0.90 | 1.00 | 1.00 |
| $\beta_{41} = 1.5$ | -0.01 | 0.01 | 0.03 | 0.12 | 0.13 | 0.93 | 0.99 | 1.00 | 1.00 |
| | -0.00 | 0.02 | -0.02 | 0.13 | 0.14 | 0.93 | 1.00 | 1.00 |
| | -0.03 | -0.03 | 0.04 | 0.13 | 0.13 | 0.91 | 1.00 | 1.00 |
| $\beta_{61} = 2$ | -0.03 | -0.02 | 0.03 | 0.11 | 0.13 | 0.92 | 1.00 | 1.00 |
| | -0.01 | -0.01 | -0.00 | 0.12 | 0.15 | 0.95 | 1.00 | 1.00 |
| $n = 700, \ p = 1000$ |       |       |     |      |       |       |
| $\beta_4 = 1.5Q_\varepsilon(\tau)$ | 0.03 | 0.08 | 0.19 | 0.19 | 0.21 | 0.92 | 0.99 | 0.61 |
| | 0.02 | 0.03 | 0.14 | 0.18 | 0.19 | 0.89 | 0.89 | 0.11 | 0.00 |
| | 0.04 | -0.03 | -0.01 | 0.21 | 0.23 | 0.92 | 0.97 | 0.56 |
| | 0.01 | 0.01 | 0.05 | 0.09 | 0.08 | 0.94 | 1.00 | 1.00 |
| $\beta_{21} = 1$ | 0.01 | 0.01 | 0.01 | 0.08 | 0.08 | 0.87 | 0.99 | 1.00 | 1.00 |
| | 0.01 | 0.01 | 0.05 | 0.10 | 0.09 | 0.89 | 1.00 | 1.00 |
| | -0.01 | 0.00 | 0.08 | 0.08 | 0.08 | 0.94 | 1.00 | 1.00 |
| $\beta_{41} = 1.5$ | -0.00 | 0.00 | 0.05 | 0.09 | 0.08 | 0.92 | 1.00 | 1.00 |
| | 0.00 | 0.01 | 0.04 | 0.09 | 0.09 | 0.95 | 1.00 | 1.00 |
| | -0.01 | -0.01 | 0.10 | 0.08 | 0.09 | 0.93 | 1.00 | 1.00 |
| $\beta_{61} = 2$ | -0.01 | -0.01 | 0.06 | 0.08 | 0.09 | 0.91 | 1.00 | 1.00 |
| | -0.00 | -0.00 | 0.07 | 0.09 | 0.10 | 0.90 | 1.00 | 1.00 |

Note: See the footnote of Table 2; Freq, average selection frequency in $B$ splits.
Table 4: Results of Example 3 based on the simulated datasets.

|                  | Bias | EmpSD | SE | Cov | Freq | Power |
|------------------|------|-------|----|-----|------|-------|
|                  | Oracle | Fused | Z18 | Fused | Fused | M09   |
| $n = 300, p = 1000$ |          |       |     |      |      |       |
| $\beta_2 = \phi_1(\tau)$ | 0.08 | 0.06 | 0.59 | 0.34 | 0.36 | 0.94 |
|                    | 0.34 | 0.37 | 1.01 | 0.52 | 0.51 | 0.89 |
|                    | 0.08 | -0.20 | -0.05 | 0.80 | 0.72 | 0.89 |
|                    | 0.10 | 0.14 | 0.27 | 0.65 | 0.50 | 0.90 |
| $\beta_5 = \phi_4(\tau)$ | -0.16 | -0.20 | -0.36 | 0.62 | 0.51 | 0.91 |
|                    | 0.02 | 0.06 | -0.03 | 0.56 | 0.52 | 0.90 |
| $\beta_{21} = 1.5$ | 0.02 | 0.03 | 0.04 | 0.25 | 0.23 | 0.95 |
| $\beta_{41} = 2$  | 0.01 | -0.00 | 0.02 | 0.23 | 0.25 | 0.93 |
| $\beta_{61} = 2.5$ | -0.02 | -0.07 | 0.19 | 0.21 | 0.26 | 0.94 |
| $n = 700, p = 1000$ |          |       |     |      |      |       |
| $\beta_2 = \phi_1(\tau)$ | 0.01 | 0.04 | 0.27 | 0.21 | 0.23 | 0.94 |
|                    | 0.13 | 0.30 | 0.79 | 0.37 | 0.40 | 0.88 |
|                    | 0.00 | 0.08 | 0.35 | 0.51 | 0.51 | 0.90 |
|                    | 0.06 | 0.09 | 0.18 | 0.33 | 0.33 | 0.91 |
| $\beta_5 = \phi_4(\tau)$ | -0.09 | -0.19 | -0.23 | 0.35 | 0.34 | 0.85 |
|                    | -0.01 | -0.04 | -0.08 | 0.37 | 0.31 | 0.94 |
| $\beta_{21} = 1.5$ | -0.00 | 0.00 | 0.04 | 0.16 | 0.17 | 0.97 |
| $\beta_{41} = 2$  | 0.01 | -0.02 | -0.01 | 0.15 | 0.18 | 0.95 |
| $\beta_{61} = 2.5$ | 0.01 | 0.00 | 0.07 | 0.18 | 0.18 | 0.94 |
| $n = 700, p = 2000$ |          |       |     |      |      |       |
| $\beta_2 = \phi_1(\tau)$ | -0.01 | 0.05 | 0.13 | 0.32 | 0.32 | 0.93 |
|                    | 0.10 | 0.26 | 0.59 | 0.46 | 0.44 | 0.91 |
|                    | 0.05 | -0.07 | 0.15 | 0.53 | 0.46 | 0.87 |
|                    | 0.10 | 0.10 | 0.25 | 0.45 | 0.35 | 0.84 |
| $\beta_5 = \phi_4(\tau)$ | -0.03 | -0.18 | -0.31 | 0.41 | 0.36 | 0.89 |
|                    | -0.00 | -0.01 | -0.13 | 0.36 | 0.34 | 0.85 |
| $\beta_{21} = 1.5$ | 0.01 | 0.01 | 0.03 | 0.18 | 0.21 | 0.98 |
| $\beta_{41} = 2$  | 0.01 | 0.02 | -0.07 | 0.22 | 0.20 | 0.91 |
| $\beta_{61} = 2.5$ | -0.01 | -0.01 | -0.05 | 0.25 | 0.20 | 0.94 |

Note: See the footnote of Tables 2 and 3: For $\beta_2$ and $\beta_5$, the numbers are shown at $\tau = .25, .5, .75$ from the top to the bottom and, for the other $\beta$'s, at $\tau = 0.5$. 


Table 5: Comparisons of computing time (on average per dataset in seconds) when performing Example 1.

|                  | Fused | Z18 | W12 | F14 | M09 |
|------------------|-------|-----|-----|-----|-----|
| \((n,p) = (300,1000)\) | 888   | 853 | 509 | 390 | 170 |
| \((n,p) = (700,1000)\) | 3,108 | 1,812 | 2,230 | 1,231 | 440 |

Note: see the footnote of Table 2.
Figure 1: Estimated heterogeneous effects and confidence intervals of Fused-HDCQR using Example 3: $\beta^*_2(\cdot)$ (left panel) and $\beta^*_5(\cdot)$ (right panel). From the top to the bottom are the plots for $(n, p) = (300, 1000), (700, 1000)$ and $(700, 2000)$, respectively.
Table 6: Patients’ characteristics in the BLCSC samples.

\( (n = 674) \)

|                     | Mean (SD) | Count (%) |
|---------------------|-----------|-----------|
| **Age**             | 60 (10.8) |           |
| **Female**          | 259 (38)  |           |
| **Education level** |           |           |
| \( \leq \) High school | 264 (39)  |           |
| > High school       | 410 (61)  |           |
| **Smoking**         |           |           |
| Non-active          | 418 (62)  |           |
| Active              | 256 (38)  |           |
| **Cancer type**     |           |           |
| Adenocarcinoma      | 283 (42)  |           |
| Squamous cell       | 110 (16)  |           |
| Other               | 281 (42)  |           |
| **Cancer stage**    |           |           |
| 1                   | 283 (42)  |           |
| 2                   | 110 (16)  |           |
| 3                   | 256 (38)  |           |
| 4                   | 25 (4)    |           |
Figure 2: Estimated quantile-specific coefficients of the predictors in Table 7.
Table 7: Analysis of the BLCSC data with Fused-HDCQR. The SNPs are sorted by their p-values at $\tau = 0.2$, corresponding to the high-risk groups. Results for the top 10 and the bottom 3 are presented.

| $\tau$  | Estimator | SE  | p-value | $\tau$  | Estimator | SE  | p-value | $\tau$  | Estimator | SE  | p-value |
|---------|-----------|-----|---------|---------|-----------|-----|---------|---------|-----------|-----|---------|
| 0.2     | Int       | 6.90| 0.25    | 1.4E-165| 0.3       | 7.48| 0.28    | 4.3E-157| 0.4       | 7.94| 0.24    | 3.2E-241|
|         | Adeno     | 0.20| 0.16    | 2.1E-01 | 0.14      | 0.18| 4.5E-01 | 0.02     | 0.13      | 8.7E-01 |
|         | Squamous  | -0.16| 0.16   | 3.0E-01 | -0.20     | 0.16| 2.1E-01 | -0.34    | 0.13      | 1.0E-02 |
|         | Stage2    | -0.82| 0.24   | 6.3E-04 | -0.99     | 0.25| 6.0E-05 | -0.98    | 0.24      | 3.2E-05 |
|         | Stage3    | -0.97| 0.17   | 1.6E-08 | -1.04     | 0.20| 2.0E-07 | -1.13    | 0.14      | 2.0E-15 |
|         | Stage4    | -1.54| 0.17   | 3.0E-20 | -1.77     | 0.20| 1.7E-19 | -1.86    | 0.14      | 2.2E-42 |
|         | Age       | -0.01| 0.01   | 1.5E-02 | -0.01     | 0.01| 3.0E-02 | -0.02    | 0.01      | 1.0E-02 |
|         | Edu       | 0.08 | 0.14   | 6.0E-01 | 0.06      | 0.15| 6.9E-01 | 0.07     | 0.13      | 5.8E-01 |
|         | Female    | -0.30| 0.13   | 2.2E-02 | -0.35     | 0.14| 1.0E-02 | -0.37    | 0.12      | 1.6E-03 |
|         | Smoke     | -0.42| 0.13   | 1.1E-03 | -0.48     | 0.14| 5.0E-04 | -0.52    | 0.11      | 3.4E-06 |
|         | AX.37793583_T | 2.75| 0.22   | 3.0E-36 | 2.61      | 0.20| 4.6E-39 | 2.39     | 0.20      | 3.7E-33 |
|         | AX.83104700_A | 2.32| 0.20   | 4.0E-31 | 1.91      | 0.19| 6.3E-24 | 1.54     | 0.19      | 1.5E-15 |
|         | AX.15207405_G | 2.03| 0.20 | 1.0E-24 | 1.59      | 0.22| 9.8E-13 | 1.17     | 0.21      | 3.7E-08 |
|         | AX.16619495_T | 1.79| 0.20   | 3.3E-19 | 1.36      | 0.20| 1.3E-11 | 0.97     | 0.20      | 1.2E-06 |
|         | AX.13920550_G | 1.93| 0.23   | 2.5E-17 | 1.41      | 0.28| 5.3E-07 | 0.87     | 0.27      | 1.6E-03 |
|         | AX.83444620_C | 1.39| 0.17   | 7.4E-17 | 1.05      | 0.19| 6.6E-08 | 0.71     | 0.21      | 8.8E-04 |
|         | AX.82902859_T | 1.58| 0.20   | 8.7E-16 | 1.19      | 0.18| 2.0E-11 | 0.90     | 0.12      | 3.4E-14 |
|         | AX.40182999_A | 1.50| 0.21   | 9.6E-13 | 1.01      | 0.25| 3.9E-05 | 0.64     | 0.14      | 6.5E-06 |
|         | AX.82976133_A | 2.32| 0.33   | 3.8E-12 | 2.02      | 0.35| 6.7E-09 | 1.58     | 0.35      | 6.1E-06 |
|         | AX.82900605_G | 2.21| 0.35   | 1.6E-10 | 1.91      | 0.29| 9.1E-11 | 1.54     | 0.33      | 2.9E-06 |
|         | ...       |      |         |          |           |     |         |          |           |     |         |
|         | AX.41828883_G | 1.4E-03| 0.34 | 1.00 | -3.2E-02| 0.42 | 0.94 | -5.7E-02| 0.54 | 0.92 |
|         | AX.11293250_T | -3.6E-04| 0.14 | 1.00 | 6.2E-02| 0.15 | 0.67 | 5.0E-02| 0.12 | 0.68 |
|         | AX.37863475_C | -3.1E-04| 0.26 | 1.00 | -1.1E-01| 0.25 | 0.68 | -1.8E-01| 0.24 | 0.46 |
|     | Estimate | SE   | p-value  |     | Estimate | SE   | p-value  |     | Estimate | SE   | p-value  |
|-----|----------|------|----------|-----|----------|------|----------|-----|----------|------|----------|
| \( \tau \) |          |      |          |     |          |      |          |     |          |      |          |
| Int  | 8.30     | 0.27 | 4.8E-214 |     | 8.55     | 0.30 | 4.9E-180 |     | 8.69     | 0.35 | 2.8E-132 |
| Adeno| -0.09    | 0.15 | 5.3E-01  |     | -0.09    | 0.13 | 4.8E-01  |     | -0.09    | 0.13 | 5.1E-01  |
| Squamous | -0.50   | 0.15 | 1.0E-03  |     | -0.60    | 0.16 | 2.1E-04  |     | -0.50    | 0.19 | 7.1E-03  |
| Stage2 | -0.88   | 0.25 | 5.0E-04  |     | -0.73    | 0.24 | 2.1E-03  |     | -0.57    | 0.19 | 2.8E-03  |
| Stage3 | -1.08   | 0.17 | 1.7E-10  |     | -0.91    | 0.15 | 6.4E-10  |     | -0.68    | 0.16 | 2.0E-05  |
| Stage4 | -1.91   | 0.15 | 7.0E-38  |     | -1.93    | 0.14 | 1.7E-44  |     | -1.69    | 0.16 | 2.1E-27  |
| Age  | -0.02   | 0.00 | 3.3E-05  |     | -0.02    | 0.01 | 1.3E-03  |     | -0.02    | 0.01 | 1.9E-03  |
| Edu  | 0.15    | 0.14 | 2.7E-01  |     | 0.16     | 0.13 | 2.2E-01  |     | 0.11     | 0.13 | 4.0E-01  |
| Female | -0.44  | 0.11 | 6.4E-05  |     | -0.47    | 0.12 | 1.6E-04  |     | -0.38    | 0.15 | 1.3E-02  |
| Smoke | -0.53   | 0.15 | 4.9E-04  |     | -0.36    | 0.16 | 2.4E-02  |     | -0.31    | 0.15 | 3.8E-02  |
| AX.37793583.T | 2.16 | 0.20 | 4.1E-28 |     | 1.84 | 0.28 | 2.8E-11 |     | 1.39 | 0.25 | 4.2E-08 |
| AX.83104700.A | 1.15 | 0.27 | 1.6E-05 |     | 0.58 | 0.27 | 3.5E-02 |     | 0.13 | 0.25 | 6.0E-01 |
| AX.15207405.G | 0.75 | 0.25 | 2.3E-03 |     | 0.34 | 0.37 | 3.5E-01 |     | -0.05 | 0.48 | 9.2E-01 |
| AX.16619495.T | 0.66 | 0.22 | 3.1E-03 |     | 0.44 | 0.31 | 1.5E-01 |     | 0.18 | 0.35 | 6.1E-01 |
| AX.13920550.G | 0.54 | 0.27 | 4.3E-02 |     | 0.26 | 0.60 | 6.7E-01 |     | 0.11 | 0.60 | 8.6E-01 |
| AX.83444620.C | 0.55 | 0.23 | 2.0E-02 |     | 0.29 | 0.22 | 1.8E-01 |     | 0.01 | 0.18 | 9.7E-01 |
| AX.82902859.T | 0.73 | 0.13 | 4.2E-08 |     | 0.51 | 0.32 | 1.1E-01 |     | 0.22 | 0.46 | 6.3E-01 |
| AX.40182999.A | 0.41 | 0.18 | 2.6E-02 |     | 0.22 | 0.27 | 4.1E-01 |     | -0.01 | 0.30 | 9.6E-01 |
| AX.82976133.A | 1.17 | 0.42 | 5.4E-03 |     | 0.61 | 0.52 | 2.4E-01 |     | 0.24 | 0.46 | 6.0E-01 |
| AX.82900605.G | 1.22 | 0.35 | 4.5E-04 |     | 0.86 | 0.34 | 1.1E-02 |     | 0.50 | 0.31 | 1.0E-01 |
| ...  |         |      |          |     |         |      |          |     |         |      |          |
| AX.41828883.G | 0.26 | 0.60 | 0.66 |     | 0.32 | 0.52 | 0.54 |     | 0.12 | 0.68 | 0.86 |
| AX.11293250.T | -0.00 | 0.12 | 1.00 |     | -0.09 | 0.12 | 0.44 |     | -0.09 | 0.15 | 0.56 |
| AX.37863475.C | -0.24 | 0.20 | 0.23 |     | -0.37 | 0.17 | 0.03 |     | -0.57 | 0.32 | 0.08 |