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Summary

In many observational longitudinal studies, the outcome of interest presents a skewed distribution, is subject to censoring due to detection limit or other reasons, and is observed at irregular times that may follow a outcome-dependent pattern. In this work, we consider quantile regression modeling of such longitudinal data, because quantile regression is generally robust in handling skewed and censored outcomes and is flexible to accommodate dynamic covariate-outcome relationships. Specifically, we study a longitudinal quantile regression model that specifies covariate effects on the marginal quantiles of the longitudinal outcome. Such a model is easy to interpret and can accommodate dynamic outcome profile changes over time. We propose estimation and inference procedures that can appropriately account for censoring and irregular outcome-dependent follow-up. Our proposals can be readily implemented based on existing software for quantile regression. We establish the asymptotic properties of the proposed estimator, including uniform consistency and weak convergence. Extensive simulations suggest good finite-sample performance of the new method. We also present an analysis of data from a long-term study of a population exposed to Polybrominated Biphenyls (PBB), which uncovers an inhomogeneous PBB elimination pattern that would not be detected by traditional longitudinal data analysis.

Keywords

Censored quantile regression; Irregular outcome-dependent follow-up; Longitudinal Data; Proportional intensity model; Recurrent Events

1. Introduction

Epidemiological follow-up studies often present various features that can complicate statistical analysis, such as censoring, skewness, and irregular outcome-dependent follow-up.
up. Our motivating example is the Michigan Long-Term Polybrominated Biphenyls (PBBs) Study, which was established following the PBB exposures of residents of Michigan farms and neighboring communities after their consumption of PBB contaminated food products in the early 1970s. With over 20 years of follow-up, PBB study provides a rich database for investigating the elimination of PBB from the human body. However, there exist several challenges with the analysis of the PBB data. First, due to laboratory assay detection limit, PBB concentration was not detectable when it was less than 1 part per billion (p.p.b.). This caused some left censored PBB measurements. Secondly, the distribution of PBB concentration is highly skewed, as evidenced by the histogram of log-transformed PBB measurements; see Figure 1. Thirdly, serum samples from each subject were not taken at a set of common time points or intervals. Furthermore, visit/sample-taking times may be outcome dependent. In Figure 2, we present the box-plots of log PBB levels measured at study entry by the number of visits. It is shown that subjects with high initial PBB levels contributed more serum samples, or equivalently speaking, made more follow-up visits, than those with low initial PBB levels. Similar data situations are encountered in many other epidemiological studies.

Many standard longitudinal methods (Diggle et al., 2002) assumed the follow-up visit times to be pre-determined or outcome-independent given covariates. When the follow-up visit times are irregular and are correlated with outcomes, one intuitive approach is to group irregular visit times into a common set of time intervals and formulate the data as longitudinal data with informative intermittent missing. Methods, such as Robins et al. (1995), can be applied; however they may be sensitive to arbitrary divisions of time intervals. Without involving discretizing the visit time scale, Lipsitz et al. (2002) developed a likelihood-based approach assuming the outcome process follows a Gaussian process with mean and covariance parametrically specified. Under reasonable conditions on the dependency between visit times and outcomes, they showed that the likelihood for the longitudinal outcomes is separable from that for the follow-up times, and thus the estimation of the outcome process parameters can be carried out without modeling the follow-up times. Fitzmaurice et al. (2006) extended Lipsitz et al. (2002)’s method to longitudinal binary data. Ryu et al. (2007) presented a Bayesian regression method, which jointly modeled the follow-up time process and the longitudinal outcome process through introducing a subject-specific latent variable.

Marginal semiparametric regression has also been studied for longitudinal data with irregular outcome-dependent follow-up. This type of approach avoids strong parametric assumptions on the joint distribution of longitudinal outcomes, but as a trade-off, requires a model for the follow-up time process to facilitate correcting the bias resulted from outcome-dependent follow-up. For example, Lin et al. (2004) investigated the marginal mean regression of the longitudinal outcome, while modeling the follow-up time process by a proportional intensity model (Andersen and Gill, 1982). They proposed an inverse intensity weighting strategy to adjust for the effect of outcome-dependent follow-up on the observed outcomes. To avoid directly estimating the baseline intensity function of the follow-up visit process, which usually requires smoothing, Buzkova and Lumley (2007) justified the use of intensity-ratio as the inverse weight in Lin et al. (2004)’s approach. The resulting estimator
does not require smoothing, and thus is simpler to compute. Furthermore, it can be applied under a mixture of continuous and discrete follow-up visit times.

With skewed and censored outcomes, quantile regression is naturally advantageous over either Gaussian regression modeling or marginal mean regression because quantiles are more robust and informative summary statistics for a skewed distribution and have better identifiability than mean when censoring is present. This motivates us to consider quantile regression modeling for longitudinal data with irregular outcome-dependent follow-up. With a proper formulation of outcome process and follow-up time process as in Lipsitz et al. (2002), our model specifies how the quantiles of the outcome at time \( t \) relates to the covariates observed up to time \( t \). This type of marginal quantile regression model has been exploited in literature by many authors in various aspects (Jung, 1996; Lipsitz et al., 1997; Koenker, 2004; Wang and Fygenson, 2009; Yi and He, 2009; Yuan and Yin, 2010; Lee and Kong, 2013, among others). For example, Lipsitz et al. (1997) proposed a GEE-type estimating equation and adopted the technique of inverse probability weighting to handle missing outcomes due to random dropouts. Koenker (2004) considered subject-specific fixed effects which are intended to capture unobserved individual heterogeneity, and proposed an \( \ell_1 \)-regularization estimating method to modify the inflation effect caused by the introduction of individual fixed effects. Wang and Fygenson (2009) investigated the case with longitudinal outcomes left censored by fixed constants and developed inference procedures that properly account for censoring and intra-subject dependency. Lee and Kong (2013) recently presented an adaptation of Wang and Fygenson (2009)'s method to handle longitudinal data subject to both left censoring and random dropouts. However, all these methods are generally focused on standard longitudinal settings with common visit times or outcome-independent follow-up times.

For marginal quantile regression inference, ignoring the dependency between irregular follow-up times and outcomes can lead to biased estimation. This is well demonstrated by an exploratory analysis of the PBB example. In Figure 3, we plot the 25th, 50th, 75th empirical quantiles of PBB measurements collected in each of four follow-up time intervals. Each gray dot represents one PBB measurement. Without accounting for outcome-dependent follow-up, a naive interpretation of Figure 3 would be that the distribution of PBB would first shift up and then go down over the time course. This is not scientifically plausible, and in fact, manifests the data distortion resulted from outcome-dependent follow-up. A further data examination reveals that the cohort participants who had PBB levels measured 10 to 16 years after PBB exposure are mostly those who had high PBB levels at the initial study visit. Thus, the PBB samples collected during this time period are not representative of the PBB concentrations of the whole study cohort, but rather reflect the PBB distribution for a subcohort that is likely to have high PBB levels. This explains the unexpected rise in empirical PBB quantiles observed in Figure 3, and more importantly indicates the need to appropriately address outcome-dependent follow-up in quantile regression analysis of longitudinal data.

In this paper, we develop a marginal quantile regression approach to analyzing longitudinal data with censored and skewed outcomes as well as irregular outcome-dependent follow-up. We propose an estimation procedure that properly accommodates these realistic data
features. More specifically, we employ Powell (1986)’s censored quantile regression technique to handle fixed or known random left censoring to longitudinal outcomes. To address outcome-dependent follow-up, we model the follow-up time process via a proportional intensity model, viewing each follow-up visit as a recurrent event. As one of the most popular models for recurrent events, a proportional intensity model can be conveniently implemented by standard statistical software such as SAS and R. It also allows us to specify how the intensity for the counting process of visits at time \( t \) depends on the past observed data, including visit history, outcomes, and covariates; thus it is an appropriate device for characterizing outcome-dependent follow-up. The adopted proportional intensity model forms the basis to correct the bias induced by outcome-dependent follow-up through inverse intensity-ratio weighting. It can also help understand the factors influencing the follow-up behaviors. We properly design our estimation and inference procedures so that they can be implemented via existing statistical software for quantile regression. Algorithmic issues are carefully addressed.

The rest of this paper is organized as follows. In Section 2, we introduce models and present the proposed estimation procedure and algorithm. We outline asymptotic studies in Section 3, and develop bootstrap and sample-based inference procedures in Section 4. In Section 5, we evaluate the proposed method by simulation studies. In Section 6, we present an analysis of PBB data, which demonstrates the importance and practical utility of the new method. We conclude with some remarks in Section 7.

2. Methods

2.1 Data and Notation

Let \( Y_i(t) \) denote the outcome process of interest, namely the outcome at time \( t \), and likewise let \( Z_i(t) \) denote a vector of external covariate processes for the \( i \)th subject. Let \([L_i, R_i]\) be a time interval indicating when the \( i \)th subject is under study. We assume that \( L_i \) and \( R_i \) are independent of \( Y_i(t) \) given \( Z_i(t) \). Note that \( Y_i(t) \) and \( Z_i(t) \) are only observed at \( L_i \) when the subject enters the study, and at a sequence of follow-up visit times, \( \{t_i^{(j)}: j=1, 2, \ldots, m_i\} \) within \([L_i, R_i]\). Here \( m_i \) is the total number of follow-up visits for the \( i \)th subject.

Define a counting process for study entry as \( N_i^L(t) = I(L_i \leq t) \) and a counting process for follow-up visits as \( N_i(t) = \sum_{j=1}^{m_i} I(t_i^{(j)} \leq t) \). Outcome \( Y_i^*(t) \) is subject to left censoring at a fixed constant \( c \). As explained in a remark in Section 7, the constant \( c \) can be replaced by a random variable which is observed for all subjects. Define \( Y_i(t) = \max(c, Y_i^*(t)) \). The observed data consist of \( \{L_i, Z_i(L_i), Y_i(L_i), t_i^{(j)}, Z_i(t_i^{(j)}), Y_i(t_i^{(j)}), R_i, m_i: j=1, 2, \ldots, m_i\}_{i=1}^n \). Notation with subscript \( i \) removed stands for the corresponding population analogue.

2.2 Models

Define the \( \tau \)th conditional quantile of a random variable \( Y \) given \( Z \) as \( Q_Y(\tau|Z) = \inf\{y: \Pr(Y \leq y|Z) \geq \tau\} \). We consider a marginal quantile regression model that takes the form,
where \( X_i(t) = (1, Z_i(t)^\top)^\top \) and \( \beta_0(\tau) \) is a vector of unknown regression coefficients. This model marginally specifies the relationship between outcome quantiles and covariates at time \( t \). The coefficients in \( \beta_0(\tau) \) are formulated as functions of \( \tau \), thereby allowing for inhomogeneous covariate effects across different segments of the outcome distribution.

Model (1) covers some commonly used models for longitudinal data. For example, one special case is the linear random intercept model, \( Y_i^*(t) = \mu + b^\top Z_i(t) + a_i + \varepsilon_i(t) \), where for subject \( i \), \( a_i \) is the random intercept effect, and \( \varepsilon_i(t) \) is the random error term at time \( t \) that follows a common distribution over \( t \). With \((a_i, \varepsilon_i(t))\) assumed to be independent of \( Z_i(t) \), it can shown that \( Q_{Y^*(\tau)\mid Z_i(t)} = \{\mu + Q_{a+\varepsilon(\tau)}\} + b^\top Z_i(t) \), where \( Q_{a+\varepsilon(\tau)} \) denotes the \( \tau \)th quantile of \( a_i + \varepsilon_i(t) \). Thus model (1) holds in this special case.

Model (1) is also flexible in characterizing the profile change of the longitudinal outcome over time, which is of particular interest in the PBB study. For instance, with \( Z_i(t) = t \), model (1) becomes \( Q_{Y^*(\tau)\mid Z_i(t)} = \{\mu + Q_{a+\varepsilon(\tau)}\} + b_1(\tau) \). In this case, the coefficient \( b_1(\tau) \) represents the \( \tau \)th quantile of baseline outcome (i.e. \( Y_i(0) \)) and \( b_1(\tau) \) represents the change rate of the \( \tau \)th outcome quantile over time. Clearly, this model can be expanded to adjust for relevant baseline covariates or covariates collected during follow-up.

We further model follow-up visit times, and give some special attention to the subtle difference between the initial study visit and follow-up visits. This is motivated by the PBB study, in which, the commonly adopted time origin is the PBB exposure date set as July 1, 1973, rather than the date at study entry. Since study participants had little or no knowledge about how much they were exposed to PBB until they received results from their initial visit, we expect little dependency between the initial visit time and outcomes. In fact, we assume that \( L_i \perp Y_i^*(\cdot)\mid Z_i(\cdot) \) and \( L_i \) is not necessarily fixed. Consequently, our modeling of the follow-up process starts after the initial study visit. That is, defining a history function \( H_i(t) \) as all observed data before time \( t \) of the \( i \)th subject, we assume a proportional intensity model (Andersen and Gill, 1982) that takes the form,

\[
\lambda(t\mid H_i(t)) = I(L_i < t \leq R_i)\lambda_0(t)\exp(h_i(t)^\top a_0),
\]

where \( \lambda(t\mid H_i(t)) = \lim_{\Delta t \to 0} \frac{1}{\Delta t} P\{N_i(t+\Delta t) - N_i(t) = 1 \mid H_i(t)\} \) and \( a_0 \) is a vector of unknown coefficients. Here \( h_i(t) \) is a vector of time-dependent covariates belonging to \( H_i(t) \), which may be flexibly set to contain prior outcomes or covariates observed before time \( t \).

### 2.3 Estimation

The primary estimation goal is to estimate the \( \beta_0(\tau) \) in model (1), which captures the covariate effects on the \( \tau \)th marginal quantile of the outcome of interest, \( Y^*(t) \). When the follow-up process is independent of the outcome process, one may follow Wang and Fygenson (2009)'s method to estimate \( \beta_0(\tau) \) through minimizing the objective function,
\[
\frac{n^{-1/2}}{\sigma} \sum_{i=1}^{n} \left[ \int_{0}^{\infty} \rho_{\tau} \{ Y_i(t) - \max(c, X_i(t)^T \beta) \} \left( dN_{i}(t) + dN_{i}(t) \right) \right], \quad (3)
\]

where \( \rho_{\tau}(u) = u \cdot \{ \tau - U(u < 0) \} \). The basic idea underlying objective function (3) is derived from an application of the equivariance property of quantiles to monotone transformation (Koenker, 2005). A similar strategy was used by Powell (1986)'s method for censored quantile regression. More specifically, by the equivariance property, under model (1), quantiles of the observed outcome, \( Y_i(\tau) \), satisfy,

\[
Q_{Y_i(\tau)}(\tau | Z_i(t)) = \max(c, X_i(t)^T \beta_0(\tau)).
\]

Such a relationship between the observed outcomes and covariates lays the key justification for objective function (3).

However, in the presence of outcome-dependent follow-up, minimizing (3) does not render a valid approach because \( Y_i(\tau) \) and \( dN_{i}(t) \) are not independent given \( X_i(t) \). To correct the bias resulted from outcome-dependent follow-up, we adopt the strategy of inverse intensity-ratio weighting (Buzkova and Lumley, 2007) with separate handling of the initial study visit and follow-up visits. That is, we intend to weigh the outcomes observed at follow-up visits by the reciprocal of the intensity ratios, which, according to the follow-up time model (2), take the form \( w_i(t; \alpha_0) = I(L_i < t \leq R_i) \exp \{ h_i(t)^T \alpha_0 \} \). At the same time, we do not weigh the data collected at the initial study visit because the study entry time \( L_i \) is assumed to be conditionally independent of outcomes given covariates. Since \( \alpha_0 \) is usually unknown, we employ weights \( w_i(t; \hat{\alpha}) \) instead of \( w_i(t; \alpha_0) \), where \( \hat{\alpha} \) is a consistent estimate for \( \alpha_0 \) obtained by Andersen and Gill (1982)'s method, which maximizes a partial likelihood function of model (2).

We propose to estimate \( \beta_0(\tau) \) by the minimizer of \( \Psi_{\tau}(\beta, \alpha) \) with respect to \( \beta \), where

\[
\Psi_{\tau}(\beta, \alpha) = n^{-1/2} \sum_{i=1}^{n} \left[ \int_{0}^{\infty} \rho_{\tau} \{ Y_i(t) - \max(c, X_i(t)^T \beta) \} \left( dN_{i}(t) + \frac{1}{w_i(t; \hat{\alpha})} dN_{i}(t) \right) \right]. \quad (4)
\]

The resulting estimator is denoted by \( \hat{\beta}(\tau) \). We recommend calculating \( w_i(t; \hat{\alpha}) \) based on centered covariates. Doing so would lead to a better interpretation of the weight and also naturally confine the weight to take values in a reasonable range. For example, suppose \( g_i(t) \) is a covariate included in \( h_i(t) \). A centered covariate \( g_i^*(t) \) is defined as \( g_i(t) - \bar{g} \), where

\[
\bar{g} = \sum_{i=1}^{n} \sum_{j=1}^{m_i} g_i(t_j) / \sum_{i=1}^{n} m_i.
\]

To find the minimizer of \( \Psi_{\tau}(\beta, \alpha) \), we note that \( \Psi_{\tau}(\beta, \alpha) \) has an equivalent form of
This shows that, by treating observed outcomes as independent and properly assigning weights as one or the reciprocal of estimated intensity-ratio, we can solve the minimization problem for objective function (4) by using the existing crq() function in R package quantreg via the option for Powell’s censored regression quantiles.

To justify the proposed weighting strategy, we examine the gradient of $\Psi_{\tau}(\beta, \alpha)$ with respect to $\beta$, denoted by $U_{\tau}(\beta, \alpha)$, which equals

$$n^{-1/2}\sum_{i=1}^{n} \left[ \int_0^{\infty} X_i(t) I(X_i(t) ^\top \beta > c) \{ I(Y_i(t) \leq X_i(t) ^\top \beta) - \tau \} \left( dN_i^L(t) + \frac{1}{w_i(t; \alpha)} dN_i(t) \right) \right].$$

Assuming $dN_i(t) \perp \{ Y_i^*(t), Z_i(t) \} | \mathcal{H}_i(t)$, or in words, $dN_i(t)$ is independent of current outcome and covariates given the history, we can show that under model assumptions (1) and (2),

$$E \left[ \int_0^{\infty} \frac{1}{w_i(t; \alpha_0)} X_i(t) I(X_i(t) ^\top \beta > c) \{ I(Y_i(t) \leq X_i(t) ^\top \beta) - \tau \} dN_i(t) \right] = 0.$$

At the same time, provided that $dN_i^L(t)$ is independent of $Y_i(t)$ given $X_i(t)$, it is easy to see that

$$E \left[ \int_0^{\infty} X_i(t) I(X_i(t) ^\top \beta > c) \{ I(Y_i(t) \leq X_i(t) ^\top \beta) - \tau \} dN_i^L(t) \right] = 0.$$

Combining these results immediately gives $E[U_{\tau}(\beta_0; \alpha_0)] = 0$, which endorses the proposed idea for estimating $\hat{\beta}(\tau)$. As commented by Lin et al. (2004), the assumption, $dN_i(t) \perp \{ Y_i^*(t), Z_i(t) \} | \mathcal{H}_i(t)$ is weaker than those imposed by Lipsitz et al. (2002), and is reasonable for the follow-up mechanisms of many real studies, including the PBB study.

3. Asymptotic Properties

We study the asymptotic properties of the proposed estimator, $\hat{\beta}(\tau)$. Due to space limit, we relegate regularity conditions and proofs of theorems to Web Appendices A–C.

The asymptotic properties of $\hat{\beta}(\tau)$ are stated in the following theorems.

Theorem 1: Under conditions C1-C4, $\sup_{\tau \in [\gamma, \gamma'] \| \hat{\beta}(\tau) - \beta_0(\tau) \rightarrow 0}$, a.s..
Theorem 2: Under conditions C1-C6, \( \{n^{1/2} [\hat{\beta}(\tau) - \beta_0(\tau)] : \tau \in [\gamma, \gamma'] \} \) converges weakly to a Gaussian process with mean 0 and covariance matrix \( \Sigma \), which is defined in (C.3) in Web Appendix C.

Note that Theorems 1 and 2 provide the asymptotic properties of \( \{\hat{\beta}(\tau) \}, \tau \in [\gamma, \gamma'] \), with \( 0 < \gamma < \gamma' < 1 \). In principle, the choice of \( [\gamma, \gamma'] \) should reflect the range of the quantile levels of interest. In theory, we assume that \( \gamma \) and \( \gamma' \) satisfy the technical constraints imposed by the regularity conditions. These constraints are necessitated by considerations relating to the tail identifiability and estimation stability. For example, with \( \gamma' < 1 \), we can avoid the complication with extreme quantile inference. When \( Y_i(t) \) is subject to left censoring by a positive constant, the lower tail quantiles may not be identifiable, and thus requiring \( \gamma > 0 \) may be necessary. In practice, there is usually no definite way to verify whether these technical constraints are met or not. Our recommendation is to first select \( \gamma \) and \( \gamma' \) based on the scientific interest and then adjust \( \gamma \) and \( \gamma' \) in an adaptive manner. That is, when \( \hat{\beta}(\tau) \) turns out to be an infeasible solution at \( \tau = \gamma \) (or \( \gamma' \)), we would reset \( \gamma \) (or \( \gamma' \)) to a larger (or smaller) value. Based on our numerical experience, such an empirical adaptive rule performances very well.

4. Inference

Bootstrap procedures can be used to make inference on \( \beta_0(\tau) \). For example, one may generate a bootstrap sample by randomly selecting \( n \) subjects with replacement. Based on each bootstrap sample, the proposed estimation procedure can be applied to obtain a bootstrap estimator, denoted by \( \hat{\beta}^* (\tau) \). With many bootstrap samples generated, the asymptotic distribution of \( \sqrt{n} \{ \hat{\beta}(\tau) - \beta_0(\tau) \} \) can be approximated by the empirical distribution of \( \sqrt{n} \{ \hat{\beta}^* (\tau) - \hat{\beta}(\tau) \} \).

We also develop a sample-based inference procedure, which is expected to be more computationally efficient. One challenge is about how to estimate the asymptotic variance matrix of \( \sqrt{n} \{ \hat{\beta}(\tau) - \beta_0(\tau) \} \), which involves the unknown density function \( f_{Y_i(t)}(X_i(t)^\top \beta_0(\tau)) \) according to our asymptotic studies. To tackle this difficulty, we adopt the technique of Huang (2002) and Peng and Fine (2009). More specifically, we perturb \( U(\beta, \alpha) \) and then utilize the functional linearity of \( U(\beta) \) to derive a sample-based consistent estimate for \( B(\beta_0(\tau); \alpha_0) \) in condition C5. Then we can obtain a consistent estimate for the asymptotic variance of \( \sqrt{n} \{ \hat{\beta}(\tau) - \beta_0(\tau) \} \) based on its closed form derived in the proof of Theorem 2. The specific procedure is outlined as follows.

Step 1. Define

\[
\ell_j(\beta; \alpha) = \int_0^\infty X_j(t) I(X_j(t)^\top \beta > c) \{ I(Y_j(t) \leq X_j(t)^\top \beta) - \tau \} \left( dN_j(t)^L + \frac{1}{w_j(t; \alpha)} dN_j(t) \right)
\]

and \( \Omega(\tau) = n^{-1} \sum_{j=1}^n \left\{ \ell_j(\hat{\beta}(\tau); \hat{\alpha}) \right\} \otimes 2 \), where \( \mathbf{v} \otimes 2 = \mathbf{v}^\top \mathbf{v} \). Find a symmetric and nonsingular \((p + 1) \times (p + 1)\) matrix \( \mathbf{E}(\tau) \) such that \( \Omega(\tau) = \mathbf{E}^2(\tau) \). Let \( \mathbf{e}_j(\tau) \) denote the \( j \)th column of \( \mathbf{E}(\tau) \).

Step 2. Solve the equation

\[
\text{Biometrics. Author manuscript; available in PMC 2016 June 01.}
\]
for \( \mathbf{b} \), and denote the solution by \( \mathbf{\beta}^j(\tau) \) \((j = 1, \ldots, p + 1)\).

Step 3. Calculate \( \mathbf{D}(\tau) = (\mathbf{\beta}_1(\tau) - \mathbf{\beta}^1(\tau), \ldots, \mathbf{\beta}_{p+1}(\tau) - \mathbf{\beta}^{p+1}(\tau))\).

Step 4. Compute \( n^{1/2} \mathbf{E}(\tau) \mathbf{D}(\tau)^{-1} \), which provides a consistent estimate for \( \mathbf{B}_j(\mathbf{\beta}_0(\tau); \mathbf{\alpha}_0) \).

It is, however, not straightforward to employ the \text{crq()}\ function in R to solve equation (5) in Step 2. In order to take the advantage of existing software package, we propose an alternative solution-finding strategy for equation (5). That is, we first solve the equation,

\[
\sum_{i=1}^{n} \int_0^\infty \mathbf{X}_i(t) I(\mathbf{X}_i(t)^T \mathbf{b} > c) \left\{ I(Y_i(t) \leq \mathbf{X}_i(t)^T \mathbf{b}) - \tau \right\} \left( dN_i^0(t) + \frac{1}{w_i(t, \mathbf{\alpha}_0)} dN_i(t) \right) + I(\mathbf{X}_j^T \mathbf{b} > 0) \mathbf{X}_j^* \left\{ I(\mathbf{X}_j^T \mathbf{b} > 0) - \tau \right\} = 0,
\]

where \( \mathbf{X}_j^* = -n^{1/2} \mathbf{e}_j(\tau)/(1 - \tau) \). Mimicking the proposed algorithm for \( \mathbf{\beta}_j(\tau) \), we can solve equation (6) using the \text{crq()}\ function. It is easy to show that equation (6), coupled with condition \( \mathbf{X}_j^T \mathbf{b} > 0 \), is equivalent to equation (5). Therefore, we check whether the solution to equation (6), \( \mathbf{b}^\sim \), satisfies the condition \( \mathbf{X}_j^T \mathbf{b} > 0 \). If yes, we let \( \mathbf{\beta}^j(\tau) = \mathbf{b}^\sim \). Otherwise, we solve equation (6) switching the sign of \( \mathbf{e}_j(\tau) \). In this case, solving equation (6) would serve to locate a solution to a variant of equation (5), which is given by

\[
\mathbf{U}_\tau(\mathbf{b}, \mathbf{\alpha}) = -\mathbf{e}_j(\tau) \tag{5'}
\]

Note that the perturbations to \( \mathbf{U}_\tau(\mathbf{b}, \mathbf{\alpha}) \) posed by equation (5) and equation (5'), namely \( \mathbf{e}_j(\tau) \) and \(-\mathbf{e}_j(\tau)\), have the same asymptotic order. Following the theory presented in Huang (2002) and Peng and Fine (2009), we can show that the proposed inference procedure remains valid when one replaces equation (5) by (5'). When (5') is adopted for some \( j \), \( \mathbf{E}(\tau) \) in Step 4 needs to be updated by \( (\mathbf{e}_1(\tau), \ldots, -\mathbf{e}_j(\tau), \ldots, \mathbf{e}_{p+1}(\tau)) \) accordingly.

Let \( \hat{\mathbf{B}}(\tau) \) denote the proposed sample-based estimate for \( \mathbf{B}_j(\mathbf{\beta}_0(\tau); \mathbf{\alpha}_0) \). A consistent sample-based covariance estimator for \( \hat{\mathbf{B}}(\tau) \) may be given by

\[
n^{-1} \sum_{i=0}^{n} \left[ -\hat{\mathbf{B}}(\tau)^{-1} \left\{ \mathbf{i}(\hat{\mathbf{\beta}}(\tau); \mathbf{\alpha}) - \hat{\mathbf{A}}_\tau \hat{\mathbf{J}}^{-1}(\hat{\mathbf{\alpha}}) \right\} \right]^{\otimes 2},
\]

where \( \mathbf{i}(\cdot) \) is an influence function defined in Web Appendix A (see (A.2)), and \( \hat{\mathbf{A}}_\tau \) and \( \hat{\mathbf{J}} \) are plug-in estimators of \( \mathbf{A}_j(\mathbf{\beta}_0; \mathbf{\alpha}_0) \) and \( \mathbf{J}(\mathbf{\alpha}_0) \) defined in (C.2) in Web Appendix C and (A.1) in Appendix A respectively.
5. Simulations

Simulation studies were conducted to assess finite-sample performance of the proposed method. We considered two time-independent covariates, \( Z_{i1} \sim \text{Uniform}(0, 1) \) and \( Z_{i2} \sim \text{Bernoulli}(0.5) \), and one time-dependent covariate \( Z_{i3}(t) = t \). For the study visit times, we generated the study entry time \( L_i \) from \( \text{Uniform}(0, 1) \), and the time at the end of follow-up \( R_i \) from \( \text{Uniform}(4, 5) \). Between \( L_i \) and \( R_i \), follow-up visit times were generated according to a proportional intensity model:

\[
P\{dN_i(t) = \mathcal{H}(t) = I(L_i < t \leq R_i)0.2\exp\{a_0 Y_{i}(t^-)\}dt, \quad (\tau)
\]

where \( Y(\tau) \) represents the last observed outcome before time \( t \) and \( a_0 = 0.2 \). A positive coefficient for \( Y(\tau) \) would indicate that subjects with larger previous outcomes have higher intensity of making subsequent visits.

We adopted Normal distribution and Gamma distribution for generating outcomes. More specifically,

Case 1: \( Y_i(t) = \max(0, 4.5 + d_i - Z_{i1} + Z_{i2} - t + \varepsilon_i(t)) \), where \( d_i \sim N(0, \frac{1}{4}((Z_{i1} + Z_{i2} + 1)^2 - \frac{1}{2})) \) and \( \varepsilon_i(t) \sim N(0, \frac{1}{8}) \) and they are independent. In this set-up, data follow a marginal quantile regression model,

\[
Q_{Y_i(\tau)}(\tau|Z_{i1}, Z_{i2}) = \max(0, 4.5 + \Phi^{-1}(\tau) + \{-1 + \Phi^{-1}(\tau)\} Z_{i1} + \{1 + \Phi^{-1}(\tau)\} Z_{i2} - t).
\]

Case 2: \( Y_i(t) = \max(0, 3.5 + d_i - 2Z_{i1} - t + \varepsilon_i(t)) \), where \( d_i \sim \text{Gamma}(\frac{3}{4}, \frac{1}{2}(Z_{i1} + Z_{i2} + 1)) \) and \( \varepsilon_i(t) \sim \text{Gamma}(1, \frac{1}{4}(Z_{i1} + Z_{i2} + 1)) \), and they are independent. In this set-up, data follow a marginal quantile regression model,

\[
Q_{Y_i(\tau)}(\tau|Z_{i1}, Z_{i2}) = \max\left(0, 3.5 + F^{-1}_{\text{Gamma}(\frac{3}{4}, t)}(\tau) + \{-2 + F^{-1}_{\text{Gamma}(\frac{1}{4}, t)}(\tau)\} Z_{i1} + F^{-1}_{\text{Gamma}(\frac{1}{4}, t)}(\tau) Z_{i2} - t\right).
\]

Under the set-ups described above, \( c \) is specified as 0, the average number of visits is 4.4, and the average left censoring rate is 10% in both case 1 and case 2. Note that fitting model (1) to a dataset with responses \( Y_i(t) \) subject to left censoring by \( c \) can be equivalently formulated as fitting model (1) to a transformed dataset with shifted responses, \( Y_i(t) - c \), subject to left censoring by the constant 0. Therefore, the cases we considered here with \( c = 0 \) are representative for the general scenarios with nonzero \( c \)’s.

For each set-up, we generated 1000 data sets of sample size \( n = 200 \). For each simulated dataset, we applied the proposed method to estimate covariate effects on the 25th, 50th, and 75th outcome quantiles. We also compared our method with a naive approach, which

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implements Wang and Fygenson (2009)'s method by obtaining the coefficient estimator as the minimizer of objective function (3). Empirical bias and standard deviations of estimators from both methods are presented in Table 1. It is shown that the proposed estimator is virtually unbiased. The bias from the naive method is quite evident; the magnitude of bias can be over half of the magnitude of standard deviation in some cases. The empirical standard deviations of the proposed estimator are reasonable for the sample size $n = 200$ and are fairly close to estimated standard deviations. The agreement between empirical and estimated standard deviations improves as the sample size is increased to $n = 400$. We also examined the estimates at higher quantiles, corresponding to $\tau = 0.85, 0.90, 0.95$. The results are presented in Table D.1 of Web Appendix D, indicating satisfactory performance of our proposals. The simulation results with $n = 400$ are presented Table D.2 of Web Appendix D.

In our simulations, we evaluated both bootstrap and sample-based inference procedures. The bootstrap size was chosen as 500. In Table 1, we report the averages of standard deviation (SD) estimates and the empirical coverage rates of 95% confidence intervals obtained from both inference approaches. Generally, both types of SD estimates are acceptably close to the empirical standard deviations. The bootstrap-based SD estimates are slightly better than the sample-based SD estimates. The computation of the sample-based approach is about 50 times faster than that of the bootstrapping procedure. Both bootstrap and sample-based inference procedures yield confidence intervals with accurate coverage rates.

We also investigated the robustness of the proposed estimation of model (1) to the potential mis-specification of the model for the follow-up time process. We consider three different scenarios of model mis-specification. The details about the set-ups and the results are relegated to Web Appendix D due to space limit.

From our simulations, we find that the proposed estimator always has much smaller bias compared to that of the naive estimator. When only a moderate model misspecification presents, the bias of the proposed estimator is only slightly larger than the empirical bias observed in the case with correctly specified follow-up model. The bias increases as the departure from the true model increases. Overall, the proposed method demonstrates quite robust performance when the follow-up time model is misspecified.

### 6. Application to PBB study

Polybrominated biphenyls (PBBs) are manufactured chemicals added as flame retardants to electrical devices, plastics, and various textiles. A widespread contamination with PBBs occurred in Michigan during 1973 - 1974 when PBB was accidently substituted for a nutritional supplement manufactured at the same chemical plant. The PBB was then mixed with animal feed. Farmers and Michigan residents throughout the state were exposed to PBB by consuming contaminated animal food products (e.g. milk, beef, pork, chicken, eggs). The Michigan Department of Public (now Community) Health (MDCH), in collaboration with the US Public Health Service, established a registry of individuals exposed to the contaminated food products. Since the initial enrollment period (1976 - 1978), the MDCH has periodically contacted cohort members to obtain additional serum samples. Serum samples from cohort members were collected from 1976–1993.
Our analysis was focused on understanding the elimination of PBB from the body by examining repeated measurements of PBB in serum. The current analysis is limited to women. PBBs are stable, persistent halogenated organic pollutants with extremely long half-lives. Participants may continue to have measurable PBB levels in serum after more than 20 years. We included females who were born before the contamination incident (July, 1973) if they had at least two serum PBB measurements at least 6 months apart, and if they had an initial serum PBB measurement greater than 2 parts per billion (p.p.b.) and taken after age 16. We required an initial serum PBB measurement of at least 2 p.p.b. to ensure that their levels were above the limit of detection of 1 p.p.b. This imposed “artificial” truncation which would limit the interpretation of our analysis results to the population with initial visit PBB measurements greater than 2 p.p.b. We excluded females who were younger than age 16 at initial measurement because childhood growth could potentially affect the compartment mobility and thus the equilibrium of serum PBB concentration levels. We also excluded measurements taken during pregnancy or during any period of breast-feeding because of the potential mobilization of PBB into the bloodstream during these times (Eyster et al., 1983; Kreuzer et al., 1997).

The final dataset used for our analysis included 364 women. There are between 2 and 7 serum measurements of PBB per woman. Initial PBB concentration level ranges from 1 to 559.80 p.p.b. (mean=11.44, median=2.40). Outcome \( Y^*(t) \) is defined as \( \log(\text{PBB}) \) at time \( t \), with the time origin set as July 1, 1973, the time when the PBB contamination started. Study entry time \( L \) is defined as the time from PBB exposure date to the first study visit. Similarly, \( R \) is defined as the time from PBB exposure to the most recent PBB serum measurements available, December 31, 1993. The initial visit time \( L \) ranges from 2.67 to 8.04 years with mean=3.74, median=3.71, and interquartile range=(3.38, 3.96). A histogram of the gap times between adjacent visits is presented in Web Appendix F.

When we modeled follow-up visit times, we excised separate attentions to the three time periods, 1976-1981, 1982-1989, and 1990-1993, to accommodate the special design of the PBB study. During 1982-1989, a substudy focused on those with high PBB levels to examine the health of those who had been more severely exposed. As a result, serum samples available during this time period were mostly contributed by participants of this substudy, who tended to have higher PBB levels than a member randomly selected from the whole study cohort. After 1990, all participants of the PBB study were aggressively contacted for serum samples. Giving a careful consideration of such a study design and conjecturing that a high initial PBB level may lead to more frequent follow-up visits, we assume the following model for the follow-up visit time process:

\[
P(dN_i(t)|\mathcal{H}(t))=I(L_i< t \leq R_i)\lambda_0(t)\times \exp\{\alpha_1 I(t \leq 8.5)Y_i(L_i)+\alpha_2 I(8.5< t \leq 16.5)Y_i(L_i)+\alpha_3 I(t>16.5)Y_i(L_i)\}. \tag{8}
\]

Note that in model (8), we convert the three calendar time intervals stated above into time intervals starting from the assigned time origin. The coefficients, \( \alpha_1, \alpha_2 \) and \( \alpha_3 \), represent the effects of the initial outcome on the follow-up time process in these three time intervals respectively.
Table 2 presents the coefficient estimates for the assumed proportional intensity model (8). All coefficient estimates are positive. The estimated $\alpha_2$ has the largest magnitude, 0.584, and is significantly different from zero with $p < 0.01$. This result indicates that one unit larger in the initial log(PBB) level may be associated with 79% higher intensity of making a follow-up visit during 1982-1989. This is consistent with our observation in Figure 2, which demonstrates participants with higher initial PBB levels have more follow-up visits. In contrast, the coefficient estimates for $\alpha_1$ and $\alpha_3$ are close to zero. This may reflect a rather uniform visit patterns across all study cohort member during the cohort recruiting and during the most recent time periods.

We modeled the outcome log(PBB) by marginal quantile regression models taking the form,

$$Q_{Y_i(t)}(\tau) = \beta_0(\tau) + \beta_1(\tau) \times t, \quad t > 0, \quad (9)$$

where $0 < \tau < 1$. Here, the intercept, $\beta_0(\tau)$, represents the $\tau$th quantile of log(PBB) level at time origin. The time effect, $\beta_1(\tau)$, represents the elimination rate of the population $\tau$th quantile of log(PBB) level over time, and thus is the key quantity of interest in this analysis. We applied the proposed method to estimate model (9) with $\tau = 0.25, 0.50, 0.75, 0.85, 0.90$ and 0.95. We also fit these models by naively applying Wang and Fygenson (2009)'s method, which would ignore the dependency between follow-up and outcome. For inference, we adopted the bootstrap procedure to obtain standard deviation estimates and 95% confidence intervals. The bootstrap size was chosen as 500.

In Table 3, we present the coefficient estimates and 95% confidence intervals obtained from the proposed method and the naive approach. Based on the naive approach, we obtain positive time coefficient estimates at all considered quantile levels. In particular, the 95% confidence interval for the time effect on the 75th percentile of PBB concentration is (0.001, 0.054). This result can lead to a conclusion that among individuals at the 75th percentile of PBB distribution the amount of PBB in their blood increases with time. It contradicts with the biologic fact that human bodies can not produce chemical PBB (and the exposure did not continue). As we explain in Section 1 on Figure 3, these positive time effect estimates are probably artifacts caused by ignoring the dependency between outcome and follow-up.

On the other hand, the estimates for $\beta_1(\tau)$ from the proposed approach are all negative except for the $\beta_1(\tau)$ estimate with $\tau = 0.25$, which is extremely close to zero. The negative estimates for the time effect are consistent with the biological evidence that PBB concentration in blood should decrease over time. By taking into account that subjects with lower initial PBB levels tend to contribute fewer serum samples, our inverse intensity-ratio weighting strategy assigned larger weights for these subjects and hence correct the bias due to outcome-dependent follow-up.

In addition, the estimated time coefficients demonstrate an overall trend of increasing with $\tau$. The time effect on lower percentiles, such as $\tau = 0.25, 0.50, 0.75$, are not significant while the time effect on the 95th quantile is significantly less than zero. Our estimates may be interpreted as that the 95th percentile of PBB distribution decreases 5.1% ($= 1 - \exp(-0.052)$) per year with 95% confidence interval between 0.7% and 9.2%. The faster
elimination rates of upper quantiles (compared to those of lower quantiles) provide some confirmation to the conjecture that subjects with higher PBB levels may demonstrate faster PBB elimination over time. The interesting and sensible varying pattern of time effect on different quantiles cannot be uncovered by traditional longitudinal models that are focused on modeling mean outcomes.

We also performed sensitivity analyses, in which we fit two different models for follow-up visit times. In one case (Case A), we assume that the proportional intensity model for visit times include one additional covariate, BMI, which represents the body mass index of study participants at study enrollment. In the other case (Case B), we fit the proportional intensity model (8) with \( Y_i(L_i) \) replaced by its discrete version with cutoff points chosen as \( \exp(1) \) and \( \exp(3) \). The results from adopting these different visit time models are presented in Web Appendix E; please see Table E.1–Table E.4. From Tables E.1 and E.3, we note that fitting the three different visit time models consistently evidences the presence of outcome-dependent following during 1982-1989, which conforms to the design of the PBB study. Moreover, it renders quite similar estimates for \( \beta_0(\tau) \). This demonstrates the robustness of the proposed method.

In summary, our analysis shows that the PBB concentration distribution shifts down slowly over time. Upper quantiles decrease faster than lower quantiles. A significant decreasing trend over time has been shown by the data for the 95th quantile of PBB distribution. Ignoring outcome-dependent follow-up would result in very biased estimates of \( \beta_1(\tau) \) leading to implausible scientific conclusions.

7. Remarks

Quantile regression offers a robust and flexible approach to analyzing longitudinal data with skewed outcomes. Irregular outcome-dependent follow-up is a common data feature but can be easily overlooked in practice. The proposed inverse intensity-ratio weighted estimator can effectively correct the bias due to irregular outcome-dependent follow-up, with reasonable modeling of the follow-up time process.

The current method exposition takes the assumption that the left censoring variable is a fixed constant. Nevertheless, the new method can be readily adapted to cases where the left censoring variable is random but always observed. Moreover, it is straightforward to extend the proposed method to deal with doubly censored longitudinal outcomes, for example, measurements subject to a upper detection limit as well as a lower detection limit.

Like Powell (1986)’s censored quantile regression method, our approach can be readily extended to cases where the left censoring variable is not just a fixed constant but a random variable which is always observed. More specifically, when the outcome \( Y_i(t) \) is subject to left censoring by an observed random variable \( C_i(t) \), the proposed method would remain valid if one replaces the constant \( c \) by \( C_i(\tau) \) and \( C_i(t) \) is independent of \( Y_i(t) \) given \( X_i(t) \). Such an extension of the proposed method can accommodate more general practical settings, for example, a long-term follow-up study where the assay detection limit changes over time.
The proposed method can be revised to accommodate other types of models for the follow-up time process, such as proportional mean/rate model (Lin et al., 2000). Our simulations suggest that the proposed estimation of the marginal quantile regression model is reasonably robust to mis-specifications of the follow-up time model. Even when the adopted proportional intensity model departs from the true model, our approach can still achieve considerable bias reductions compared to the naive approach that does not make any adjustment for outcome dependent follow-up.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**


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Figure 1.
Distribution of PBB concentration measurements after logarithm transformation.
Figure 2.
Distribution of observed log (PBB) at the first visit versus number of measurements.
Figure 3.
An intuitive data illustration of PBB study.
Simulation studies that compared the proposed method and the naive approach: EmpSD – empirical standard deviation; AvgSD – the average of standard deviation estimates; Cov95 – the coverage rate of a 95% confidence interval.

<table>
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<th>Proposed</th>
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<td>EmpSD</td>
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### Table 3
Parameter estimates and 95% confidence interval for PBB study

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<td>(0.609, 1.199)</td>
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<td>(1.246, 1.745)</td>
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<td>(1.763, 2.833)</td>
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<td>(3.112, 4.514)</td>
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<td>(3.379, 4.716)</td>
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