sleev: Semiparametric Likelihood Estimation with Errors in Variables

Jiangmei Xiong Sarah C. Lotspeich Joey B. Sherrill Gustavo Amorim Bryan E. Shepherd Ran Tao

Data with measurement error in the outcome, covariates, or both are not uncommon, particularly with the increased use of routinely collected data for biomedical research. In settings with error-prone data, two-phase studies, where researchers validate a subsample of study data, can be used to obtain unbiased estimates. The sieve maximum likelihood estimator (SMLE), which combines the error-prone data on all records with the validated data on a subsample, is a highly efficient and robust estimator to analyze such two-phase data. However, given their complexity, a computationally efficient and user-friendly tool is needed to obtain SMLEs. This paper introduces the sleev package for making semiparametric likelihood-based inference using SMLEs for error-prone two-phase data in settings with binary and continuous outcomes. Functions from this package can be used to analyze data with error-prone responses and covariates. Various examples are presented to provide users with guidance in handling different types of variables. To demonstrate the use of the functions in practice, they are applied to a two-phase dataset simulated to represent data obtained from the electronic health records of an HIV clinic.

1. Introduction

Routinely collected data are being used more frequently in biomedical research. For example, data extracted from electronic health records have been used in numerous studies as a cost-effective resource to obtain information on a large number of people. However, these data tend to be error-prone, often across multiple variables, and careless use of these data could lead to biased estimates and misleading research findings (Duan et al. 2016). To avoid invalid study results, routinely collected data may undergo an audition or validation, in which trained experts carefully verify and extract data elements. However, it is usually only feasible to validate data for a subset of records or variables. After validation, researchers have two types of data: (i) error-prone pre-validation data for all records (phase one data) and (ii) error-free

validated data on a subset of records (phase two data). For analyses, the goal is then to combine the two types of data to obtain estimates that have low bias and are as efficient (i.e., have the smallest variance) as possible.

Building off of the measurement error and missing data literature, there are several types of approaches for combining such two-phase data with errors, including design-based methods (e.g., inverse-probability weighted estimators (Horvitz and Thompson 1952) and generalized raking estimators (Deville, Särndal, and Sautory 1993; Oh et al. 2021)) and model-based methods (e.g., maximum likelihood estimation (Carroll et al. 2006; Tang et al. 2015) and multiple imputation (Little and Rubin 1986; Cole, Chu, and Greenland 2006; Giganti et al. 2020)). Both design- and model-based estimators require the missing at random assumption for unbiased estimation, i.e., conditional on observed data, those records to be validated are selected through simple random sampling. Design-based estimators also require that the probability of being selected for validation is non-zero for all records, whereas no such positivity assumption is required for model-based estimators. Because they make no model assumptions on the error mechanism, design-based estimators tend to be more robust but less efficient than model-based estimators (Bang and Robins 2005; Amorim et al. 2021).

A robust class of model-based estimators, sieve maximum likelihood estimators (SMLEs), have recently been developed to analyze two-phase data with errors in both the outcome and covariates (Tao et al. 2021; Lotspeich et al. 2022). SMLEs are semiparametric and robust because they avoid making parametric assumptions on the nuisance models of the error terms, and, as full-likelihood estimators, they remain highly efficient. Hence, they provide a nice balance between robustness and efficiency. Still, in practice these estimators can be difficult to implement, as they involve approximating nuisance conditional densities using B-splines (Schumaker 2007) and then maximizing the semiparametric likelihood via a sophisticated EM algorithm (Tao, Zeng, and Lin 2017).

In this paper, we introduce the sleev package, which computes SMLEs for linear and logistic regressions using partially-validated, error-prone data. The sleev package incorporates error-prone data on all records plus validated data on a subset of records to obtain efficient and robust estimates of regression parameters in a user-friendly manner. This paper describes the SMLE method (Sections 2 and 3) and demonstrates the features of the sleev package and the application of functions in the package with a detailed illustration using simulated HIV data (Section 4).

2. Sieve maximum likelihood estimators for linear regression

Suppose that we want to fit a standard linear regression model for a continuous outcome Y and vector of covariates X: $Y = \alpha + \boldsymbol{\beta}^T \mathbf{X} + \epsilon$, where ϵ follows a normal distribution with mean zero and variance σ^2 . Our goal is to obtain estimates of $\boldsymbol{\theta} = (\alpha, \boldsymbol{\beta}^T, \sigma^2)^T$. When we have error-prone data, Y and \mathbf{X} are unobserved except for a subset of subjects whose records are validated. For the majority of subjects whose records are not validated, only the error-prone

outcome $Y^* = Y + W$ and covariates $X^* = X + U$ are observed in place of Y and X, where W and U are the additive errors for the outcome and covariates, respectively. It is assumed that the measurement errors W and U are independent of ϵ . However, W and U can be correlated. Note that X^* can also include error-free covariates Z, which can be expressed as $X^* = (X_0^{*T}, Z^T)^T$, where X_0^* denotes error-prone covariates. However, for simplicity, we do not include the expression of error-free covariates Z throughout Sections 2 and 3. With potential errors in our data, a naive regression analysis using error-prone variables Y^* and X^* could render misleading results.

We assume that the joint density of the complete data (Y^*, X^*, W, U) takes the form

$$P(Y^*, \boldsymbol{X^*}, W, \boldsymbol{U}) = P(Y^* | \boldsymbol{X^*}, W, \boldsymbol{U}) P(W, \boldsymbol{U} | \boldsymbol{X^*}) P(\boldsymbol{X^*})$$
$$= P_{\boldsymbol{\theta}}(Y | \boldsymbol{X}) P(W, \boldsymbol{U} | \boldsymbol{X^*}) P(\boldsymbol{X^*}),$$

where $P(\cdot)$ and $P(\cdot|\cdot)$ denote density and conditional density functions, respectively. $P_{\theta}(Y|X)$ then refers to the conditional density function of the linear regression model $Y = \alpha + \beta^{T}X + \epsilon$. Denote the validation indicator variable by V, with V = 1 indicating that a record was validated and V = 0 otherwise. For records that do not undergo validation, their measurement errors (W, \mathbf{U}) are missing. Therefore, the contributions of these unvalidated subjects to the log-likelihood can be obtained by integrating out W and \mathbf{U} . Let $(Y_i^*, \mathbf{X}_i^*, W_i, \mathbf{U}_i, V_i, Y_i, \mathbf{X}_i)$ for $i = 1, \ldots, n$ denote independent and identically distributed realizations of $(Y^*, \mathbf{X}^*, W, \mathbf{U}, V, Y, \mathbf{X})$ in a sample of n subjects. Then, the observed-data log-likelihood takes the form

$$\begin{split} &\sum_{i=1}^{n} V_{i} \{ \log P_{\pmb{\theta}}(Y_{i}|\pmb{X}_{i}) + \log P(W_{i},\pmb{U}_{i}|\pmb{X}_{i}^{*}) \} \\ &+ \sum_{i=1}^{n} (1-V_{i}) \log \left\{ \int \int P_{\pmb{\theta}}(Y_{i}^{*} - w|\pmb{X}_{i}^{*} - \pmb{u}) P(w,\pmb{u}|\pmb{X}_{i}^{*}) \mathrm{d}w \mathrm{d}\pmb{u} \right\}. \end{split} \tag{1}$$

Note that $P(X^*)$ is left out, because the error-prone covariates are fully observed and thus $P(X^*)$ can simply be estimated empirically.

Because the measurement error model, $P(W_i, U_i | X_i^*)$, is often unknown in practice, we prefer to leave it unspecified, and use nonparametric maximum likelihood estimation (NPMLE) to estimate it. NPMLE estimates $P(W, U | X^* = x^*)$ with the m distinct observed (W, U) values, $\{(w_1, u_1), ..., (w_m, u_m)\}$. Because NPMLE estimates $P(W, U | X^* = x^*)$ with the empirical density, it will not be applicable when X^* contains continuous elements, where only a small number of observations on (W, U) will be associated with each $X^* = x^*$. In this situation, we estimate $P(W_i, U_i | X_i^*)$ with B-spline sieves. Specifically, we approximate $P(w_i, u_i | X_i^*)$ and $\log P(W_i, U_i | X_i^*)$ by $\sum_{k=1}^m I(w_i = w_k, u_i = u_k) \sum_{j=1}^{s_n} B_j^q(X_i^*) p_{kj}$ and $\sum_{k=1}^m I(W_i = w_k, U_i = u_k) \sum_{j=1}^{s_n} B_j^q(X_i^*) \log p_{kj}$, respectively. $B_j^q(X_i^*)$ is the jth B-spline basis function of order q evaluated at X_i^* , s_n is the dimension of the B-spline basis, and p_{kj} is the coefficient associated with $B_j^q(X_i^*)$ and (w_k, u_k) . We note that the p_{kj} 's need to satisfy the constraints $\sum_{k=1}^m p_{kj} = 1$

and $p_{kj} \geq 0$ since they approximate conditional densities. The log-likelihood in expression (Equation 1) is now approximated by

$$\begin{split} &\sum_{i=1}^{n} V_{i} \{ \log P_{\pmb{\theta}}(Y_{i}|\pmb{X}_{i}) + \sum_{k=1}^{m} I[(W_{i} = w_{k}, \pmb{U}_{i} = \pmb{u}_{k}) \sum_{j=1}^{s_{n}} B_{j}^{q}(\pmb{X}_{i}^{*}) \log p_{kj}] \} \\ &+ \sum_{i=1}^{n} (1 - V_{i}) \log \left\{ \sum_{k=1}^{m} P_{\pmb{\theta}}(Y_{i}^{*} - w_{k}|\pmb{X}_{i}^{*} - \pmb{u}_{k}) \sum_{j=1}^{s_{n}} B_{j}^{q}(\pmb{X}_{i}^{*}) p_{kj} \right\}. \end{split} \tag{2}$$

The maximization of expression (Equation 2) is carried out through an EM algorithm to find the SMLEs $\hat{\boldsymbol{\theta}}$ and \hat{p}_{kj} . The covariance matrix of the SMLE $\hat{\boldsymbol{\theta}}$ is obtained through the method of profile likelihood (Murphy and Van der Vaart 2000). Full details on the SMLE method for linear regression with error-prone data, including its theoretical properties, can be found in Tao et al. (2021).

3. Sieve maximum likelihood estimators for logistic regression

For a binary outcome Y, we fit a logistic regression model $P_{\boldsymbol{\theta}}(Y=1|\mathbf{X}) = \left[1 + \exp\{-(\alpha + \boldsymbol{\beta}^{\mathrm{T}}\mathbf{X})\}\right]^{-1}$ with parameters $\boldsymbol{\theta} = (\alpha, \boldsymbol{\beta}^{\mathrm{T}})^{\mathrm{T}}$. The joint density of $(Y^*, \boldsymbol{X}^*, Y, \boldsymbol{X})$ is

$$P(Y^*, X^*, Y, X) = P(Y^*|X^*, Y, X)P(Y|X, X^*)P(X|X^*)P(X^*)$$

= $P(Y^*|X^*, Y, X)P_{\theta}(Y|X)P(X|X^*)P(X^*),$

where $P(Y|X, X^*) = P_{\theta}(Y|X)$ follows from the assumption that Y and X^* are conditionally independent given X (i.e., X^* is a surrogate for X). Similar to the linear regression case, the observed-data log-likelihood takes the form

$$\sum_{i=1}^{n} V_{i} \{ \log P_{\boldsymbol{\theta}}(Y_{i}|\boldsymbol{X}_{i}) + \log P(Y_{i}^{*}|\boldsymbol{X}_{i}^{*}, Y_{i}, \boldsymbol{X}_{i}) + \log P(\boldsymbol{X}_{i}|\boldsymbol{X}_{i}^{*}) \}
+ \sum_{i=1}^{n} (1 - V_{i}) \log \left\{ \sum_{y=0}^{1} \int P_{\boldsymbol{\theta}}(y|\boldsymbol{x}) P(Y_{i}^{*}|\boldsymbol{X}_{i}^{*}, y, \boldsymbol{x}) P(\boldsymbol{x}|\boldsymbol{X}_{i}^{*}) d\boldsymbol{x} \right\}.$$
(3)

We fit $P(Y^*|X^*,Y,X)$ with an additional logistic regression model $P_{\gamma}(Y^*|X^*,Y,X)$ with γ denoting its parameters. We estimate $P(X|X^*)$ with NPMLE when X^* is discrete, and use B-spline approximation when X^* contains continuous components. Specifically, we approximate $P(x|X^*)$ and $\log P(X_i|X_i^*)$ in expression (Equation 3) by $\sum_{k=1}^m I(x=x_k) \sum_{j=1}^{s_n} B_j^q(X_i^*) p_{kj}$ and $\sum_{k=1}^m I(X_i=x_k) \sum_{j=1}^{s_n} B_j^q(X_i^*) \log p_{kj}$, respectively. Consequently, the approximated log-likelihood can be rewritten with the B-splines as

$$\sum_{i=1}^n V_i \bigg\{ \log P_{\pmb{\theta}}(Y_i | \pmb{X}_i) + \log P_{\pmb{\gamma}}(Y_i^* | \pmb{X}_i^*, Y_i, \pmb{X}_i) + \sum_{k=1}^m I(\pmb{X}_i = \pmb{x}_k) \sum_{j=1}^{s_n} B_j^q(\pmb{X}_i^*) \log p_{kj} \bigg\}$$

$$+ \sum_{i=1}^{n} (1 - V_i) \log \left\{ \sum_{y=0}^{1} \sum_{k=1}^{m} P_{\boldsymbol{\theta}}(y|\boldsymbol{x}_k) P_{\boldsymbol{\gamma}}(Y_i^*|\boldsymbol{X}_i^*, y, \boldsymbol{x}_k) \sum_{j=1}^{s_n} B_j^q(\boldsymbol{X}_i^*) p_{kj} \right\}. \tag{4}$$

Similar to the linear regression case, we maximize expression (Equation 4) through an EM algorithm to obtain the SMLEs $\hat{\boldsymbol{\theta}}$ and $\hat{\boldsymbol{p}}_{kj}$. Then, we use the method of profile likelihood to estimate the covariance of $\hat{\boldsymbol{\theta}}$. More details, including its theoretical properties, can be found in Lotspeich et al. (2022).

4. Case study with mock data

4.1 Overview and installation of the sleev R package

The sleev package provides a user-friendly way to obtain the SMLEs and their standard errors. The package can be installed through CRAN:

```
install.packages("sleev")
library(sleev)
```

The sleev package includes two main functions: linear2ph() and logistic2ph(), to fit linear and logistic regressions, respectively, under two-phase sampling with an error-prone outcome and covariates. The input arguments are similar for the two functions and listed in Table 1. From Table 1, we see that in addition to the arguments for error-prone and error-free outcome and covariates, the user needs to specify the B-spline matrix $B_j^q(\boldsymbol{X}_i^*)$ to be used in the estimation of the error densities.

We now illustrate how to obtain SMLEs using the sleev package. First, we briefly describe the data that will be used, and then we show how to fit a linear regression model in the presence of errors in both the outcome and covariates using the linear2ph() function. We will explain how to choose the dimension of the B-spline basis, s_n . We will also demonstrate two ways to handle the situation when there is more than one continuous covariate in the model, where the dimension of the B-spline basis increases exponentially with the number of continuous covariates. Finally, we will briefly demonstrate the use of logistic2ph() to fit a logistic regression model, which is largely analogous to the use of linear2ph().

4.2 Overview of Data

We illustrate the usage of the functions in the sleev package with a dataset constructed to mimic data from the Vanderbilt Comprehensive Care Clinic (VCCC) patient records from Giganti et al. (2020). The VCCC collects data for people living with HIV who were admitted to the clinic between 1998 and 2011. The VCCC cohort records are fully validated. Thus, it is

Table 1: Main arguments for linear2ph() and logistic2ph()

Argument	Description						
Y_unval	Column name of unvalidated outcome in the input dataset.						
Υ	Column name of validated outcome in the input dataset. NAs in the input will be counted as individuals not selected in phase two.						
X_unval	Column names of unvalidated covariates in the input dataset.						
Х	Column names of validated covariates in the input dataset. NAs in the input will be counted as individuals not selected in phase two.						
Z	Column names of error-free covariates in the input dataset.						
Bspline	Column names of the B-spline basis in the input dataset.						
data	Dataset.						
hn_scale	Scale of the perturbation constant in the variance estimation via the method of profile likelihood. The default is 1.						
noSE	Standard errors of the parameter estimators will not be estimated when set to TRUE. The default is FALSE.						
TOL	Convergence criterion. The default is 0.0001.						
MAX_ITER	Maximum number of iterations in the EM algorithm. The default is 1000.						
verbose	Print analysis details when set to TRUE. The default is FALSE.						

an ideal dataset for illustrating the SMLEs. The VCCC dataset contains complete data for all 2087 patients; we use this number as the sample size for our simulated dataset. The simulated VCCC data were created by sampling from distributions that are similar to the original dataset. For our illustrations, we assume that 835 (40%) patient records were validated. We selected the 835 patients through simple random sampling and hid the validated values for the remaining 1252 patients by setting them as missing. Table 2 lists the variables to be used in subsequent analyses.

The dataset is included in the sleev package, and it can be loaded by

data(mock.vccc)

Table Table 3 displays the first six rows of the VCCC dataset: patients 1, 3, and 5 have NA listed for the VL_val, ADE_val, and CD4_val variables, which means that these patients were not selected for data validation; in contrast, patients 2, 4, and 6 had their data validated. For example, from the data validation, patient 2 had a viral load (VL) of 907 copies/mL³ and no AIDS-defining events within one year of antiretroviral therapy (ART) initiation confirmed. However, the patient's CD4 at ART initiation was found to be 114 cells/mm³, not 36.

Because of skewness, we often transform both CD4 and VL. In our analysis, CD4 was divided by 10 and square-root transformed and VL was \log_{10} transformed:

Table 2: Data dictionary for mock.vccc

Name	Status	Type	Description			
ID	error-free		Patient ID			
VL_unval	error-prone	continuous	Viral load (VL) at antiretroviral therapy (ART)			
VL_val	validated	continuous	initiation			
ADE_unval	error-prone	binary	Had an AIDS-defining event (ADE) within one			
ADE_val	validated	binary	year of ART initiation: 1 - yes, 0 - no			
CD4_unval	error-prone	continuous				
CD4_val	validated	continuous	CD4 count at ART initiation			
prior_ART	error-free	binary	Experienced ART before enrollment: 1 - yes, 0 -			
			no			
Sex	error-free	binary	Sex at birth of patient: 1 - male, 0 - female			
Age	error-free	continuous	Age of patient			

Table 3: First six patients in mock.vccc

[H]

ID	VL_unval	VL_val	ADE_unval	ADE_val	CD4_unval	CD4_val	prior_ART	Sex	Age
1	1358	NA	0	NA	465	NA	0	1	33
2	907	907	0	0	36	114	1	1	25
3	2284	NA	0	NA	263	NA	1	1	35
4	25473	25473	0	0	244	235	0	0	65
5	19	NA	0	NA	263	NA	1	1	37
6	36662	36662	0	0	30	30	1	0	47

```
mock.vccc$CD4_val_sq10 <- sqrt(mock.vccc$CD4_val/10)
mock.vccc$CD4_unval_sq10 <- sqrt(mock.vccc$CD4_unval/10)
mock.vccc$VL_val_l10 <- log10(mock.vccc$VL_val)
mock.vccc$VL_unval_l10 <- log10(mock.vccc$VL_unval)</pre>
```

4.3 Linear regression with mock data

We first illustrate the use of the linear2ph() function by fitting a linear regression model with CD4 count at ART (Y) regressed on VL at ART initiation (X), adjusting for sex at birth (Z). Both CD4 and VL are error-prone, partially-validated variables, whereas sex is error-free.

4.3.1 Setting up the B-spline basis for modeling the error mechanisms

To obtain the SMLEs, we first need to set up the B-spline basis for the covariates VL_unval_110 (the transformed error-prone VL variable from phase one) and Sex. We used the splines package to achieve this, which can be loaded with:

```
library(splines)
```

Here, we use a cubic B-spline basis with the $\mathtt{degree=3}$ argument in our call to the $\mathtt{bs}()$ function from the $\mathtt{splines}$ package. The size of the basis s_n is set to be 20; this was specified through the $\mathtt{df}=20$ argument. The B-spline basis is set up separately for the two \mathtt{Sex} groups, and the size of the B-spline basis is assigned in proportion to the relative size of the two \mathtt{Sex} groups. This allows the errors in $\mathtt{VL_unval_110}$ to be heterogeneous between males and females. The described B-spline basis is constructed as follows.

```
Bspline[mock.vccc$Sex==1,(sn_0+1):sn] <- Bspline_1
# name B-spline basis matrix columns properly
colnames(Bspline) <- paste0("bs", 1:sn)</pre>
```

Alternatively, if the investigator has prior knowledge that the errors in VL_unval_110 are likely to be homogeneous, one may fit a simpler model by not stratifying the B-spline basis by Sex. As a final step before running the analysis, we add the B-spline basis to the analysis dataset to create a single data frame:

```
data <- data.frame(cbind(mock.vccc, Bspline))</pre>
```

4.3.2 Model fitting and result interpretation

The SMLEs can be obtained by running

[1] "Run time: 2.789 min"

The fitted SMLEs are stored in a list object, denoted by res_linear in the code above. This list contains five slots: coefficients, covariance, sigma, converge, and converge_cov. We should first check if the EM algorithms for estimating the regression coefficients and their covariance matrix converged by checking if res_linear\$converge and res_linear\$converge_conv are TRUE.

```
c(res_linear$converge, res_linear$converge_cov)
```

[1] TRUE TRUE

The res_linear\$coefficient slot gives the regression coefficient estimates and their corresponding standard error estimates, z-statistics, and p-values.

res linear\$coefficients

```
Estimate SE Statistic p-value Intercept 4.8209166 0.15865204 30.386729 0.0000000000 VL_val_l10 -0.1413168 0.03983406 -3.547636 0.0003887047 Sex 0.2727984 0.10888178 2.505455 0.0122294098
```

Similar to interpreting the output from a standard linear model (i.e., fitted with lm()), the output here indicates that, after adjusting for sex, for every one-unit increase in the transformed viral load at ART initiation, there is expected to be 0.1413168 decrease in the transformed CD4 count at ART initiation. The transformed CD4 count can be transformed back to the original scale for interpretation. For example, a female patient with a VL of 1000 copies/ mL^3 is expected to have a CD4 count of approximately 193 cells/ mm^3 , which is lower than a female patient with a viral load of 100 copies/ mL^3 , whose expected CD4 count is approximately 206 cells/ mm^3 . It is expected that the average transformed CD4 count for males is 0.2727984 higher than that for females, adjusting for VL. Based on the p-values, both VL and sex are associated with CD4 count at the 0.05 significance level.

The res_linear\$covariance slot gives the covariance matrix, which can be used to calculate confidence intervals for the outcome variable for a subset of the patients. For example, suppose we want to know the 95% confidence interval of the expected CD4 count for male patients with VL of 1200 copies/mL³. First, we need to calculate the estimated mean transformed CD4 count for this patient group:

```
x.vec <- matrix(data = c(1,log10(1200),1), ncol=1) # set up data matrix
est.mean <- res_linear$coefficients[,1]%*%x.vec # calculate estimated mean
est.mean</pre>
```

```
[,1]
[1,] 4.658575
```

Then, we use the estimated covariance matrix to compute the variance of the linear combination est.mean:

res_linear\$covariance

```
[,1] [,2] [,3]
[1,] 0.025170471 -5.055755e-03 -8.873233e-03
[2,] -0.005055755 1.586752e-03 -5.841645e-05
[3,] -0.008873233 -5.841645e-05 1.185524e-02
```

```
est.cov <- t(x.vec)%*%res_linear$covariance%*%x.vec # covariance of est.mean
```

The 95% confidence interval of CD4 count $(cells/10mm^3)^{1/2}$ for this group is therefore

```
est.mean + c(-1, 1) * 1.96 * c(sqrt(est.cov))
```

Warning in est.mean + c(-1, 1) * 1.96 * c(sqrt(est.cov)): Recycling array of length 1 in array Use c() or as.vector() instead.

[1] 4.554328 4.762822

4.4 Choosing the B-spline basis through cross-validation

When constructing the B-spline basis in the estimation of the error models, one needs to specify the order of the B-spline functions q and the size of the B-spline basis s_n . It is customary to use cubic splines in practice. Quadratic and linear splines are also permissible, especially when the correlation between the covariates and their measurement errors are expected to be modest. The optimal size of the B-spline basis can be selected through k-fold cross-validation with the cv_linear2ph() function.

- 1. $cv_linear2ph()$ splits the data into k folds.
- 2. In each of k iterations, cv_linear2ph() holds out one fold and fits the model using the remaining k-1 folds, and then predicts the log-likelihood function in the hold-out fold.
- 3. cv linear2ph() calculates the average predicted log-likelihood across the k iterations.

The size of the B-spline basis that yields the largest average predicted log-likelihood will be chosen for subsequent analysis. The following code shows an example of using cv_linear2ph() to select the desirable size of the B-spline basis in the mock.vccc dataset. The number of folds is set to be five.

```
# set for reproducibility of fold assignment
set.seed(1)
# different B-spline sizes
sns <- c(15, 20, 25, 30, 35, 40)
# vector to hold mean log-likelihood for each sn
pred_loglike.1 <- rep(NA, length(sns))
# get number of rows of the dataset
n <- nrow(mock.vccc)
# specify number of folds in the cross validation
k <- 5
# calculate proportion of female patients in the dataset</pre>
```

```
sex_ratio <- sum(mock.vccc$Sex==0)/n</pre>
for (i in 1:length(sns)) {
  # constructing B-spline basis using the same process as in Section 4.3.1
  sn <- sns[i]
  sn 0 <- round(sn*sex ratio)</pre>
  sn_1 <- sn_sn_0
  Bspline_0 <- splines::bs(x=mock.vccc$VL_unval_110[mock.vccc$Sex==0],</pre>
                             df=sn_0, degree=3, intercept=TRUE)
  Bspline_1 <- splines::bs(x=mock.vccc$VL_unval_l10[mock.vccc$Sex==1],</pre>
                             df=sn_1, degree=3, intercept=TRUE)
  Bspline <- matrix(0, n, sn)</pre>
  Bspline[mock.vccc$Sex==0,1:sn_0] <- Bspline_0</pre>
  Bspline[mock.vccc$Sex==1,(sn_0+1):sn] <- Bspline_1</pre>
  colnames(Bspline) <- paste("bs", 1:sn, sep="")</pre>
  data.sieve <- data.frame(cbind(mock.vccc, Bspline))</pre>
  # cross validation, produce mean log-likelihood
  res.1 <- cv_linear2ph(Y="CD4_val_sq10", Y_unval="CD4_unval_sq10",
                         X="VL_val_110", X_unval="VL_unval_110", Z="Sex",
                         Bspline=colnames(Bspline), data=data.sieve,
                         nfolds=k, MAX_ITER = 2000, TOL = 1e-04,
                          verbose = FALSE)
  # save mean log-likelihood result
  pred_loglike.1[i] <- res.1$avg_pred_loglik</pre>
}
```

The average predicted log-likelihoods for the different s_n considered are:

```
out <- data.frame(sns, pred_loglike.1)
options(digits = 6)
print(out, row.names = FALSE)</pre>
```

It can be seen that the model with $s_n=20$ in the B-spline basis yields the highest average predicted log-likelihood, and is therefore chosen. We note that the average predicted log-likelihoods are fairly similar, indicating that the size of the B-spline basis does not impact the results very much in this dataset. This observation agrees with the results of Tao et al. (2021).

To confirm that there is negligible difference between the models fitted with different B-spline sizes, we can compare the SMLEs with $s_n = 20$ and $s_n = 35$.

```
# same process as in Section 4.3, fit with sn = 35
sn <- 35
sex_ratio <- sum(mock.vccc$Sex==0)/n
sn_0 <- round(sn*sex_ratio)</pre>
sn_1 <- sn_sn_0
Bspline_0 <- splines::bs(x=mock.vccc$VL_unval_110[mock.vccc$Sex==0], df=sn_0,</pre>
                          degree=3, intercept=TRUE)
Bspline_1 <- splines::bs(x=mock.vccc$VL_unval_110[mock.vccc$Sex==1], df=sn_1,
                           degree=3, intercept=TRUE)
Bspline <- matrix(0, n, sn)</pre>
Bspline[mock.vccc$Sex==0,1:sn_0] <- Bspline_0</pre>
Bspline[mock.vccc$Sex==1,(sn_0+1):sn] <- Bspline_1</pre>
colnames(Bspline) <- paste("bs", 1:sn, sep="")</pre>
data.sieve <- data.frame(cbind(mock.vccc, Bspline))</pre>
Bspline <- splines::bs(mock.vccc$VL_unval_110, df=sn, degree=3, intercept=TRUE)
colnames(Bspline) <- paste("bs", 1:sn, sep="")</pre>
data.sieve <- cbind(mock.vccc, Bspline)</pre>
fit.sn.35 <- linear2ph(Y="CD4_val_sq10", Y_unval="CD4_unval_sq10",
                        X="VL_val_110", X_unval="VL_unval_110", Z="Sex",
                        Bspline=colnames(Bspline), data=data.sieve,
                        hn_scale = 1, noSE = FALSE, TOL = 1e-04,
                        MAX_ITER = 1000, verbose = FALSE)
# compare the coefficients to those from Section 4.3.2
res_linear$coefficients
```

```
Estimate SE Statistic p-value Intercept 4.820917 0.1586520 30.38673 0.000000000 VL_val_l10 -0.141317 0.0398341 -3.54764 0.000388705 Sex 0.272798 0.1088818 2.50545 0.012229410
```

```
fit.sn.35$coefficients
```

```
Estimate SE Statistic p-value Intercept 4.846977 0.1592393 30.43832 0.000000000 VL_val_l10 -0.143348 0.0405102 -3.53856 0.000402319 Sex 0.250310 0.1058351 2.36509 0.018025598
```

The comparison shows that that the estimates, standard errors, z-statistics, and p-values of the parameters from the model with different B-spline sizes are very similar.

4.5 Example with two continuous covariates

In this section, we illustrate the use of the linear2ph() function with two continuous covariates using i) a bivariate B-spline basis and ii) a B-spline basis based on the first principle component (PC) of the covariates. Both approaches are reasonable, and they produce similar results in this example. The latter method is recommended for computational efficiency when there are more than two continuous covariates in the model. Suppose that we are fitting a model with CD4 count as the outcome and VL, age, and sex as covariates. This model is very similar to the model in Section 4.3, but with the addition of another error-free covariate age. Now, we have one binary and two continuous variables to be incorporated into the B-spline basis.

4.5.1 Bivariate B-spline

One approach to incorporate two continuous covariates is to use the tensor product of the one-dimensional B-spline bases for each variable. First, we construct the B-spline bases separately for the two continuous covariates, VL and age, stratified by sex. Due to the curse of dimensionality, the choice of number of knots has more restrictions than when there is only one continuous covariate. For instance, in this example the smallest size for the one-dimensional cubic B-spline basis is 4. If we set one-dimensional $s_n = 4$, the aggregate size of the multi-dimensional B-spline basis will be $4^2 + 4^2 = 32$, which is considered big with regard to the sample size we have available.

The first $4^2 = 16$ columns of the Bspline matrix will contain the bivariate B-spline basis for male patients, and the last 16 columns will contain the bivariate B-spline basis for female patients.

The B-spline matrix is combined with the dataset and the corresponding column names are added as input arguments. Note that for the linear2ph() function, argument hn_scale is set to be 1/4 here. This is a parameter for step size in variance estimation using the method of profile likelihood (Murphy and Van der Vaart 2000). It tunes the numerical calculation of the profile log-likelihood and can trouble-shoot the issue of occasional NA values in the covariance matrix. In this case, there are NAs in the result when hn_scale is 1. We re-run the analysis with hn_scale set to 1/2, 1/4, 1/8, and the variance estimates are very similar among these hn_scale values (data not shown). Therefore, we choose 1/4 to be the hn_scale value.

res_linear\$coefficients

```
Estimate SE Statistic p-value
Intercept 5.02567 0.25515 19.70 0.000000
VL_val_l10 -0.13057 0.03938 -3.32 0.000913
Age -0.00575 0.00475 -1.21 0.225746
Sex 0.28170 0.10901 2.58 0.009764
```

4.5.2 Principal component analysis

When there are several continuous covariates in the model, it may be challenging to construct a multidimensional B-spline basis using the tensor product method. The challenge is due to the curse of dimensionality, and is especially true when there is a relatively small validation sample. One way around this is to use principal component analysis (PCA) to first reduce the dimension of the continuous covariates and then construct the B-spline basis based on the top principal component rather than the original covariates. Here we illustrate the use of this approach by using the top principle component to reduce the dimension from two to one. However, this approach is versatile (and probably more useful) when there are more than two continuous covariates. We use the prcomp() function in base R to perform PCA for the two continuous covariates. The two input variables are the error-prone unvalidated VL and error-free age:

The steps below are identical to what we did in Section 4.3, except that we construct the B-spline basis on the first PC of VL and age rather than the original covariates.

[1] "Run time: 10.722 min"

res_linear\$coefficients

```
Estimate SE Statistic p-value
Intercept 5.0657 0.2564 19.76 0.000000
VL_val_l10 -0.1374 0.0395 -3.48 0.000502
Age -0.0061 0.0047 -1.30 0.194326
Sex 0.2818 0.1082 2.60 0.009229
```

Note that these results are very close to those generated when using the bivariate B-spline basis.

4.6 Fitting a logistic regression model with logistic2ph()

We now illustrate fitting a logistic regression model using logistic2ph(). Suppose we are interested in fitting a logistic regression model of having an AIDS-defining event (ADE) within one year of ART initiation on CD4 count at ART initiation (CD4), adjusting for whether the patient is ART naive at enrollment. Among the three variables, both ADE and CD4 are error-prone and partially validated, and ART is error-free.

We set up the B-spline basis for estimating the error mechanisms in a similar way as in Section 4.3.1. That is, we set up different B-spline bases within each stratum of ART status at enrollment. This allows the errors in CD4 count to be heterogeneous between patients who are and are not ART naive at enrollment. Again, we assemble the variables and B-splines into one data frame prior to fitting the SMLE.

```
# same process as in Section 4.3.1
n <- nrow(mock.vccc)
sn=20
art_ratio <- sum(mock.vccc$prior_ART==0)/n
sn_0 <- round(sn*art_ratio)</pre>
```

Now, we obtain the SMLEs for the logistic regression model of interest by running the logistic2ph() function on our augmented dataset:

[1] "Run time: 2.703 min"

The arguments here are analogous to those of res_linear. Argument hn_scale is set to be 1/2, and it is set using the same method as in Section 4.5.2.

Like linear2ph(), the logistic2ph() function returns the results in a list, which we have stored in the object res_logistic. The coefficient slot gives the coefficient estimates and corresponding standard errors, z-statistics, and p-values.

res_logistic\$coefficients

```
Estimate SE Statistic p-value
Intercept -0.809 0.3770 -2.146 3.19e-02
CD4_val_sq10 -0.543 0.0872 -6.222 4.90e-10
prior_ART -0.254 0.3118 -0.815 4.15e-01
```

The coefficient estimate associated with CD4 indicates that the odds of having an ADE within one year of ART initiation decreases with increasing CD4. Specifically, adjusting for whether a

patient is ART naive at enrollment, a person with a CD4 count of 360 cells/mm^3 is estimated to have $\exp(-0.543) = 0.581$ times the odds of having an ADE within one year of ART initiation compared to a person with a CD4 count of 250 cells/mm^3 . A 95% confidence interval for this odds ratio, computed in the usual manner, is

```
exp(res_logistic\\coefficients[2,1]+c(-1,1)*1.96*res_logistic\\coefficients[2,2])
```

[1] 0.49 0.69

After adjusting for CD4 count, the estimated odds ratio for having an ADE within one year for ART naive patients versus patients not ART naive is exp(-0.254) = 0.776, and the 95% confidence interval is

```
\exp(\text{res\_logistic\$coefficients}[3,1]+c(-1,1)*1.96*\text{res\_logistic\$coefficients}[3,2])
```

[1] 0.421 1.429

Based on these results, the association between having ADE within one year of ART initiation and CD4 is significant at the 0.05 level. However, the association between having ADE within one year of ART initiation and whether the patient is ART naive at enrollment is not.

5. Summary and discussion

The sleev R package is a useful tool for analyzing two-phase validation studies with error-prone outcome and covariates. It empowers users to perform linear regression for continuous outcomes and logistic regression for binary outcomes. It allows the errors among variables to be correlated with each other and with additional error-free covariates. It also accommodates the conventional measurement error scenarios with errors in the outcome or covariates only. The resulting SMLEs are statistically efficient and numerically stable while making minimal assumptions on the error distributions.

In this paper, we demonstrate the usage of functions in the sleev package under different scenarios. We showcase the selection process of the size of a B-spline basis, which is not frequently seen in papers that involve the use of B-splines. We also show the impact of curse of dimensionality on the construction of the B-spline basis, and recommend PCA as a dimension reduction technique that can circumvent this "challenge". We hope that users find the demonstrations in this paper useful for their applications.

In the future, we plan to extend the SMLE to address errors in outcomes and covariates for models with count and time-to-event outcomes. Additional functions will be added to the sleev package as methods are developed for these settings.

Acknowledgement

This research was supported by the National Institute of Health grants R01HL094786, R01AI131771, and P30AI110527 and the 2022 Biostatistics Faculty Development Award from the Department of Biostatistics at Vanderbilt University Medical Center.

- Amorim, G., R. Tao, S. Lotspeich, P. A. Shaw, T. Lumley, and B. E. Shepherd. 2021. "Two-Phase Sampling Designs for Data Validation in Settings with Covariate Measurement Error and Continuous Outcome." *Journal of the Royal Statistical Society. Series A, (Statistics in Society)*, 1368–89. https://doi.org/10.1111/rssa.12689.
- Bang, H., and J. M. Robins. 2005. "Doubly Robust Estimation in Missing Data and Causal Inference Models." *Biometrics* 61 (4): 962–73. https://doi.org/10.1111/j.1541-0420.2005. 00377.x.
- Carroll, R. J., D. Ruppert, L. A. Stefanski, and C. M. Crainiceanu. 2006. *Measurement Error in Nonlinear Models: A Modern Perspective*. Chapman; Hall/CRC.
- Cole, S. R., H. Chu, and S. Greenland. 2006. "Multiple-Imputation for Measurement-Error Correction." *International Journal of Epidemiology* 35 (4): 1074–81. https://doi.org/10.1093/ije/dyl097.
- Deville, J. C., C. E. Särndal, and O. Sautory. 1993. "Generalized Raking Procedures in Survey Sampling." *Journal of the American Statistical Association* 88 (423): 1013–20. https://doi.org/10.2307/2290793.
- Duan, R., M. Cao, Y. Wu, J. Huang, J. C. Denny, H. Xu, and Y. Chen. 2016. "An Empirical Study for Impacts of Measurement Errors on EHR Based Association Studies." In *AMIA Annual Symposium Proceedings*, 2016:1764. American Medical Informatics Association. https://pubmed.ncbi.nlm.nih.gov/28269935/.
- Giganti, M. J., P. A. Shaw, G. Chen, S. S. Bebawy, M. M. Turner, T. R. Sterling, and B. E. Shepherd. 2020. "Accounting for Dependent Errors in Predictors and Time-to-Event Outcomes Using Electronic Health Records, Validation Samples, and Multiple Imputation." The Annals of Applied Statistics 14 (2): 1045. https://doi.org/10.1214/20-aoas1343.
- Horvitz, D. G., and D. J. Thompson. 1952. "A Generalization of Sampling Without Replacement from a Finite Universe." *Journal of the American Statistical Association* 47 (260): 663–85. https://doi.org/10.2307/2280784.
- Little, R. J. A., and D. B. Rubin. 1986. Statistical Analysis with Missing Data. John Wiley & Sons.
- Lotspeich, S. C., B. E. Shepherd, G. Amorim, P. A. Shaw, and R. Tao. 2022. "Efficient Odds Ratio Estimation Under Two-Phase Sampling Using Error-Prone Data from a Multi-National HIV Research Cohort." *Biometrics* 78 (4): 1674–85. https://doi.org/10.1111/biom.13512.
- Murphy, S., and A. Van der Vaart. 2000. "On Profile Likelihood." *Journal of the American Statistical Association* 95 (450): 449–65. https://doi.org/10.2307/2669386.
- Oh, E. J., B. E. Shepherd, T. Lumley, and P. A. Shaw. 2021. "Raking and Regression Calibration: Methods to Address Bias from Correlated Covariate and Time-to-Event Error." Statistics in Medicine 40 (3): 631–49. https://doi.org/10.1002/sim.8793.

- Schumaker, L. 2007. Spline Functions: Basic Theory. 3rd ed. Cambridge Mathematical Library. Cambridge University Press. https://doi.org/10.1017/CBO9780511618994.
- Tang, L., R. H. Lyles, C. C. King, D. D. Celentano, and Y. Lo. 2015. "Binary Regression with Differentially Misclassified Response and Exposure Variables." *Statistics in Medicine* 34 (9): 1605–20. https://doi.org/10.1002/sim.6440.
- Tao, R., S. C. Lotspeich, G. Amorim, P. A. Shaw, and B. E. Shepherd. 2021. "Efficient Semiparametric Inference for Two-phase Studies with Outcome and Covariate Measurement Errors." *Statistics in Medicine* 40 (3): 725–38. https://doi.org/10.1002/sim.8799.
- Tao, R., D. Zeng, and D. Lin. 2017. "Efficient Semiparametric Inference Under Two-Phase Sampling, with Applications to Genetic Association Studies." *Journal of the American Statistical Association* 112 (520): 1468–76. https://doi.org/10.1080/01621459.2017.1295864.