SLCARE: An R Package for Semiparametric Latent Class Analysis of Recurrent Events

by Qi Yu and Limin Peng

Abstract Recurrent event data frequently arise in biomedical follow-up studies. The concept of latent classes enables researchers to characterize complex population heterogeneity in a plausible and parsimonious way. This article introduces the R package SLCARE, which implements a robust and flexible algorithm to carry out Zhao, Peng, and Hanfelt (2022)'s latent class analysis method for recurrent event data, where semiparametric multiplicative intensity modeling is adopted. SLCARE returns estimates for non-functional model parameters along with the associated variance estimates and p values. Visualization tools are provided to depict the estimated functional model parameters and related functional quantities of interest. SLCARE also delivers a model checking plot to help assess the adequacy of the fitted model.

1 Introduction

Recurrent event data frequently arise in biomedical follow-up studies where the event of interest, such as infection or hospitalization, occurs repeatedly over time. The event recurrence often demonstrates different patterns across individuals. To accommodate such heterogeneity, many regression methods have been developed with implementation available as R packages. To name a few, analyses based on the proportional intensity model (Andersen and Gill 1982) can be carried out with function coxph() from R package survival (Therneau 2023) or cph() from R package rms (Harrell Jr 2023). Fitting Wang, Qin, and Chiang (2001) 's method based on a multiplicative intensity model is implemented by function reReg() from R package reReg (Chiou et al. 2023). Modeling the gap time between recurrent events, Clement and Strawderman (2009) proposed conditional generalized estimating equation, and developed R package condGEE implementing this approach. Fine, Yan, and Kosorok (2004) 's temporal regression strategy can be applied to regress the mean function of recurrent events through using function tpr() from R package tpr (Yan and Fine 2004).

More recently, viewing the observed data as a manifestation of latent classes or subgroups, Zhao, Peng, and Hanfelt (2022) proposed a semiparametric latent class analysis (LCA) method to help reveal more realistic heterogeneity structure of recurrent event data that may not be adequately captured by a single regression model. Zhao, Peng, and Hanfelt (2022) adopted latent variable mixture modeling (LVMM) while avoiding stringent parametric assumptions commonly employed in LCA literature. Note that several R packages are available for fitting LVMM for data types other than recurrent event data. For example, R package flexmix (Grün and Leisch 2008) delivers a general tool for tackling finite mixtures of regression models, such as the standard linear model and the generalized linear model, with the expectation-maximization (EM) algorithms. R package lcmm (Proust-Lima, Philipps, and Liquet 2017) allows for fitting joint mixture models with longitudinal and single time-to-event outcomes based on the Newton-Raphson algorithm. However, none of these packages offer options to conduct LCA oriented to recurrent event outcomes.

In this paper, we introduce the R package SLCARE, which delivers robust and flexible implementation of Zhao, Peng, and Hanfelt (2022) 's LCA method for recurrent event data and is available from the Comprehensive R Archive Network (CRAN) at https://CRAN.R-project.org/package=SLCARE and Github at https://github.com/qyxxx/SLCARE. The package SLCARE offers a user-friendly software for conducting LCA of recurrent event data in R with the core function SLCARE(). The function SLCARE() enables a variety of options for specifying and estimating the flexible semiparametric latent class model of Zhao, Peng, and Hanfelt (2022). For example, the number of latent classes can be pre-specified or chosen based on an entropy-based measure. Users can opt to built-in or customized initializer. Algorithm convergence criterion can be set through specifying the maximum number of iterations or the minimum change in parameter estimates. The function SLCARE() also provides visual tools to aid in result presentation and interpretation as well as model checking upon ggplot2 plotting environment (Wickham 2016).

The remainder of this paper is organized as follows. Next section describes the methodological background for the R package SLCARE, including model assumptions, estimation procedure and algorithm, inference, and model evaluation. In the third section, we elaborate the structure of the package. The utility of the package is illustrated via simulated data examples in Section 4. A real application is presented in Section 5. We conclude with a few remarks in the last section.

2 Methodological background

Recurrent event data and notation

For subject i, let $T_i^{(j)}$ denote time to the j^{th} recurrent event $(i=1,\ldots,n,j=1,2,\ldots)$. The underlying counting process of recurrent events is defined as $N_i^*(t) = \sum_{j=1}^\infty I(T_i^{(j)} \leq t)$, representing the number of events occurred before or at time t for subject i, where $I(\cdot)$ denotes the indicator function. Suppose the observation of subject i is ended at time C_i . Then the observed counting process is given by $N_i(t) = N_i^*(\min(t,C_i)) = \sum_{j=1}^\infty I(T_i^{(j)} \leq \min(t,C_i))$ $(i=1,\ldots,n)$. Let \tilde{Z}_i be a p-dimensional vector of time-independent covariates for subject i. The observed data consist of $\{N_i(t),C_i,\tilde{Z}_i\}_{i=1}^n$.

Models and assumptions

Suppose the whole population consists of K latent classes, and assume that how covariates are related to the intensity function of recurrent events are the same within each latent class while being allowed to vary across different latent classes. Zhao, Peng, and Hanfelt (2022) proposed a latent class multiplicative intensity model, which assumes that $N_i^*(t)$ is a nonstationary Poisson process with the intensity function,

$$\lambda_i(t) = \sum_{k=1}^K I(\xi_i = k) \times \lambda_0(t) \times W_i \times \eta_{0,k} \times \exp(\tilde{Z}_i^{\top} \tilde{\beta}_{0,k})$$
 (1)

where $I(\cdot)$ denotes the indicator function, ξ_i denotes the unobserved latent class membership, $\lambda_0(t)$ is an unspecified, continuous, nonnegative baseline intensity function shared by all latent classes, W_i is a positive subject-specific latent variable independent of $(\xi_i, \tilde{Z}_i, C_i)$. The latent variable W_i plays the same role as individual frailty reflecting more or less frequent occurrences of recurrent events. Here $\eta_{0,k}$ is a positive number that captures the class-k scale shift from the baseline intensity function, and $\tilde{\beta}_{0,k}$ is the regression coefficient that represents the class-k covariate effects on the intensity function. For the identifiability of $\lambda_0(t)$ and $\eta_{0,k}$, it is assumed that $E(W_i|\tilde{Z}_i,\xi_i=k)=1$ for $k=1,\ldots,K$ and $\int_0^{\nu^*} \lambda_0(u) du=1$, where ν^* is a predetermined constant. As commented in Zhao, Peng, and Hanfelt (2022), one may choose ν^* to be slightly smaller than the upper bound of C_i 's support. A different choice of ν^* only results in a scale shift to $\lambda_0(t)$ and has no influence on the regression coefficients, $\tilde{\beta}_{0,k}$'s.

Zhao, Peng, and Hanfelt (2022) modeled the distribution of the latent class membership ξ_i by a logistic regression model:

$$P(\xi_i = k | \tilde{Z}_i) = p_k(\alpha_0, \tilde{Z}_i) \doteq \frac{\exp(\tilde{Z}_i^\top \alpha_{0,k})}{\sum_{k=1}^K \exp(\tilde{Z}_i^\top \alpha_{0,k})}, \quad k = 1, \dots, K$$
 (2)

where $\alpha_0 = (\alpha_{0,1}^\top, \dots, \alpha_{0,K}^\top)^\top$ with $\alpha_{0,1} = \mathbf{0}_{p \times 1}$.

Estimation and inference procedures

Estimation procedure

Let $Z_i = (1, \tilde{Z}_i^\top)^\top$ and $\beta_{0,k} = (\log \eta_{0,k}, \tilde{\beta}_{0,k}^\top)^\top$. By the model assumptions stated in Section 2, the following equalities hold:

$$E[I(\xi_i = k)Z_i \{ \frac{N_i^*(C_i)}{\mu_0(C_i)} - \exp(Z_i^\top \beta_{0,k}) \}] = 0, \ k = 1, \dots, K,$$
(3)

$$E[\tilde{Z}_{i}\{I(\xi_{i}=k) - \frac{\exp(\tilde{Z}_{i}^{\top}\alpha_{0,k})}{\sum_{k=1}^{K}\exp(\tilde{Z}_{i}^{\top}\alpha_{0,k})}\}] = 0, \ k = 1, \dots, K,$$
(4)

where $\mu_0(t) = \int_0^t \lambda_0(s) ds$, representing the cumulative baseline intensity function.

Equalities in (3) and (4) cannot be directly used to construct estimating equations for α_0 and $\beta_{0,k}$'s because ξ_i 's are not observable and $\mu_0(\cdot)$ is unknown. Adapting the principle of conditional score (Stefanski and Carroll 1987) for handling missing covariates, Zhao, Peng, and Hanfelt (2022) proposed to address the unobserved $I(\xi_i = k)$ by substituting it with $\tau_{ik} \doteq E[I(\xi_i = k)|Z_i, C_i, D_i]$,

where $D_i \doteq N_i(C_i)$, capturing the number of the observed recurrent events from subject i. First, it follows from the definition that

$$\tau_{ik} = \frac{P(D_i = d_i | \xi_i = k, Z_i, C_i) P(\xi_i = k | Z_i, C_i)}{\sum_{l=1}^{K} P(D_i = d_i | \xi_i = l, Z_i, C_i) P(\xi_i = l | Z_i, C_i)}.$$
(5)

As implied by model (2) and the assumption that $C_i \perp (N_i^*(\cdot), \xi_i)|Z_i$,

$$P(\xi_i = k | Z_i, C_i) = p_k(\alpha_0, \tilde{Z}_i) = \frac{\exp(\tilde{Z}_i^\top \alpha_{0,k})}{\sum_{l=1}^K \exp(\tilde{Z}_i^\top \alpha_{0,l})}$$
(6)

and

$$P(D_{i} = d_{i} | \xi_{i} = k, Z_{i}, C_{i})$$

$$= \int_{0}^{\infty} \frac{\{\exp(Z_{i}^{\top} \beta_{0,k}) w \cdot \mu_{0}(C_{i})\}^{d_{i}}}{d_{i}!} \exp\{-\exp(Z_{i}^{\top} \beta_{0,k}) w \cdot \mu_{0}(C_{i})\} \cdot f_{W}(w) dw,$$
(7)

where $f_W(\cdot)$ denotes the density function of frailty W. Combining the results in (5)-(7), τ_{ik} can be explicitly expressed as a function of $\alpha_0 \doteq (\alpha_{0,1}^\top \dots, \alpha_{0,K}^\top)^\top$, $\beta_0 \doteq (\beta_{0,1}^\top, \dots, \beta_{0,k}^\top)^\top$ and $\mu_0(\cdot)$, which is denoted by $\tau_{ik}(\alpha_0, \beta_0, \mu_0)$.

In addition, Zhao, Peng, and Hanfelt (2022) proposed a Nelson-Aalen type estimator of $\mu_0(\cdot)$, given by

$$\hat{\mu}(t) = \exp\{\hat{H}(t)\} \quad \text{with} \quad \hat{H}(t) = -\int_{t}^{\nu^{*}} \frac{\sum_{i=1}^{n} dN_{i}(s)}{\sum_{i=1}^{n} I(C_{i} \ge s)N_{i}(s)}.$$
 (8)

Replacing $I(\xi_i = k)$ and $\mu_0(\cdot)$ by $\tau_{ik}(\alpha, \beta, \hat{\mu})$ and $\hat{\mu}(\cdot)$ respectively in the empirical counterparts of (3)-(4) then leads to the following estimating equations:

$$n^{1/2}S_{1,n}(\alpha,\beta,\hat{\mu}) = 0,$$
 (9)

$$n^{1/2}S_{2,n}(\alpha,\beta,\hat{\mu}) = 0,$$
 (10)

where $S_{j,n}(\alpha,\beta,\hat{\mu}) = (S_{j,n,1}(\alpha,\beta,\hat{\mu})^{\top},\cdots,S_{j,n,K}(\alpha,\beta,\hat{\mu})^{\top})^{\top}$ (j=1,2) with

$$S_{1,n,k}(\alpha,\beta,\mu) = \frac{1}{n} \sum_{i=1}^{n} \tau_{ik}(\alpha,\beta,\mu) Z_{i} \{ \frac{N_{i}^{*}(C_{i})}{\hat{\mu}(C_{i})} - \exp(Z_{i}^{\top}\beta_{k}) \}, \quad k = 1, \dots, K;$$
 (11)

$$S_{2,n,k}(\alpha,\beta,\mu) = \frac{1}{n} \sum_{i=1}^{n} \tilde{Z}_i \{ \tau_{ik}(\alpha,\beta,\mu) - \frac{\exp(\tilde{Z}_i^{\top}\alpha_k)}{\sum_{j=1}^{K} \exp(\tilde{Z}_i^{\top}\alpha_j)} \}, \quad k = 1, \dots, K.$$
 (12)

Solving equations (9) and (10) renders estimators of α_0 and β_0 , denoted by $\hat{\alpha}$ and $\hat{\beta}$. Detailed derivations of estimating equations as well as the asymptotic properties of $\hat{\alpha}$ and $\hat{\beta}$ can be found in Zhao, Peng, and Hanfelt (2022) .

Estimation algorithm

Based on the result in (8) and estimating equations in (9), and (10), we propose the following algorithm to obtain $\hat{\alpha}$ and $\hat{\beta}$, the parameter estimates for models (1) and (2).

Step 1: Compute $\hat{\mu}(\cdot)$ based on formula (8). Let $L_{iter}=0$ and set initial values of α_0 and β_0 as $\hat{\alpha}^*$ and $\hat{\beta}^*$ respectively. Calculate $\hat{\tau}^*_{ik} \doteq \tau_{ik}(\hat{\alpha}^*, \hat{\beta}^*, \hat{\mu})$ based on equations (5), (6) and (7).

Step2: Repeat in the loop:

- (i) Solve estimating equations (9) and (10) with $\tau_{ik}(\alpha, \beta, \hat{\mu})$ fixed as $\hat{\tau}_{ik}^*$. Denote the solutions by $\check{\alpha}$ and $\check{\beta}$.
- (ii) $\hat{\alpha}^* := \check{\alpha}$; $\hat{\beta}^* := \check{\beta}$; $\hat{\tau}_{ik}^* := \tau_{ik}(\hat{\alpha}^*, \hat{\beta}^*, \hat{\mu})$. Increase L_{iter} by 1.
- (iii) If $\max\left(\|\frac{\check{\beta}-\hat{\beta}^*}{\hat{\beta}^*}\|_{\infty}, \|\frac{\check{\alpha}-\hat{\alpha}^*}{\hat{\alpha}^*}\|_{\infty}\right)$ is smaller than a pre-specified threshold value or L_{iter} is greater than a pre-specified maximum number of iteration, then exit the loop. Here $\|\cdot\|_{\infty}$ denotes the L_{∞} norm and the fraction between vectors α and β is component-wise fraction.

End repeat loop.

Return $\check{\alpha}$ and $\check{\beta}$ as the final estimates for α_0 , and β_0 (i.e., $\hat{\alpha}$ and $\hat{\beta}$).

Algorithm implementation

In the following, we present some details related to the algorithm implementation, such as how to set initial values, $\hat{\alpha}^*$ and $\hat{\beta}^*$, and how to solve the estimating equations involved in the presented algorithm.

Setting initial values for the estimation algorithm The function SLCARE() in package **SLCARE** allows users to specify the initial values, $\hat{\alpha}^*$ and $\hat{\beta}^*$, by their own choice. Unless users manually specify the initial values, SLCARE() implements an automated initializer, which obtains $\hat{\alpha}^*$ and $\hat{\beta}^*$ through the following procedure:

Step 1: Perform K-means clustering (Lloyd 1982) of the observed covariates and number of recurrent events, i.e., $\{\tilde{Z}_i, N_i(C_i)\}_{i=1}^n$, with R function kmeans (), and divide all subjects into K group, where K stands for the number of latent classes. Denote the assigned group membership for subject i by $G_i (\in \{1, \ldots, K\})$.

Step 2: Obtain $\hat{\alpha}^*$, the initial estimate for α_0 , as the regression coefficients from fitting the multinomial regression of $\{G_i\}_{i=1}^n$ on covariates $\{\tilde{Z}_i\}_{i=1}^n$.

Step 3: Obtain $\hat{\beta}^*$, the initial estimate of β_0 , as $(\hat{\beta}_1^{*\top}, \dots, \hat{\beta}_K^{*\top})$, where $\hat{\beta}_k^{*\top}$ is the regression coefficient estimate from fitting Wang, Qin, and Chiang (2001) 's multiplicative intensity model to the subset with $G_i = k$ using reReg() from R package reReg (Chiou et al. 2023).

Specifying the distribution of frailty W The SLCARE() function allows users to specify the distribution of frailty W. While Zhao, Peng, and Hanfelt (2022) 's method allows the frailty distribution to be a general parametric distribution, the implementation by SLCARE() confines to settings where W=1 or W follows a distribution that is parameterized as Gamma(k,k). These choices of frailty distributions cover a variety of density forms.

Solving estimating equations In Step 2(i) of the presented algorithm, an updated estimate for β_0 , $\check{\beta}$, is obtained from solving equations, $\sum_{i=1}^n \hat{\tau}_{ik} \cdot Z_i \cdot \left\{ \frac{N_i^*(C_i)}{\hat{\mu}(C_i)} - \exp(Z_i^\top \beta_k) \right\} = 0$, $k = 1, \ldots, K$. As suggested by Zhao, Peng, and Hanfelt (2022), we solve these equations by fitting a 'pseudo' weighted Poisson regression model to an augmented dataset which includes responses, $\{\frac{N_i^*(C_i)}{\hat{\mu}(C_i)}\}_{i=1}^n$, and covariates, $\{Z_i\}_{i=1}^n$, along with weights $\hat{\tau}_{ik}$. **SLCARE** performs such Poisson regression with R function glm. fit() from R package **stats**.

Meanwhile, in Step 2(i) of the presented algorithm, an updated estimate for α_0 , $\check{\alpha}$, is obtained by solving equations, $\frac{1}{\sqrt{n}}\sum_{i=1}^n \tilde{Z}_i \left\{ \hat{\tau}_{ik} - \frac{\exp(\tilde{Z}_i^{\top}\alpha_k)}{\sum_{j=1}^K \exp(\tilde{Z}_i^{\top}\alpha_j)} \right\} = 0$, $k=2,\cdots,K$. To solve these equations, we use an alternative and yet equivalent approach that fits weighted multinomial regression model on an augmented dataset, which consists of nK response-covariate pairs, $\{(1,\tilde{Z}_i),\ldots,(K,\tilde{Z}_i)\}_{i=1}^n$ with weight $\hat{\tau}_{ik}$ assigned to the pair (k,\tilde{Z}_i) $(k=1,\ldots,K)$. SLCARE implements the weighted multinomial regression with function multinom() from R package nnet (Venables and Ripley 2002).

Determination of the number of latent classes Zhao, Peng, and Hanfelt (2022) proposed to determine *K*, the number of latent classes, based on a relative entropy measure (Ramaswamy et al. 1993). That is, upon fitting models (1) and (2) that assume *M* latent classes, the corresponding relative entropy measure is defined as

$$Entropy(M) = 1 - \frac{\sum_{i=1}^{n} \sum_{k=1}^{M} -\hat{\tau}_{ik} \log(\hat{\tau}_{ik})}{n \log(M)}$$
 (13)

where $\hat{\tau}_{ik} = \tau_{ik}(\hat{x}, \hat{\beta}, \hat{\mu})$ (k = 1, ..., M). By definition, Entropy(M) is bounded between 0 and 1 with a larger value indicating a better fit to the data (Celeux and Soromenho 1996). Following the rationale, one may choose K as the maximizer of $Entropy(\cdot)$ over a sequence of candidate values for K. Given a pre-specified K, the value of Entropy(K) can be generated by S3 method print(x, type = "Entropy").

Model checking Zhao, Peng, and Hanfelt (2022) proposed a graphical method for checking the overall fit of models (1) and (2). The key idea is to compare the observed recurrent events $D_i \doteq N_i(C_i)$ versus the expected number of recurrent events under models (1)-(2). Specifically, models (1) and (2) imply $E\{N_i(C_i)|Z_i\} = E\{N_i^*(C_i)|Z_i\} = \sum_{k=1}^K \tau_{ik} \cdot \mu_0(C_i) \exp(Z_i^\top \beta_{0,k})$. Then the expected number of recurrent events under the assumed models can be approximated by

$$\hat{D}_i \doteq \sum_{i=1}^K \hat{\tau}_{ik} \cdot \hat{\mu}(C_i) \cdot \exp(Z_i^\top \hat{\beta}_k). \tag{14}$$

Therefore, in the scatter plot of \hat{D}_i versus D_i , a major departure from the identity line $D_i = \hat{D}_i$ may suggest a lack-of-fit of the assumed models. Model checking plot can be generated by S3 method plot(x, type = "ModelChecking").

Class-specific mean function of recurrent events To help illustrate heterogeneity in recurrent event occurrence across different latent classes, **SLCARE** computes crude estimates for the class-specific mean functions of recurrent events. Specifically, the crude estimate for the class-specific mean function of recurrent event is computed as

$$\frac{\sum_{i=1}^{n} I(\hat{\xi}_{i} = k) \sum_{j=1}^{K} \hat{\tau}_{ij} \cdot \hat{\mu}(t) \exp(Z_{i}^{\top} \hat{\beta}_{j})}{\sum_{i=1}^{n} I(\hat{\xi}_{i} = k)}, \quad k = 1, \dots, K,$$
(15)

where $\hat{\xi}_i = \arg\max_{1 \leq k \leq K} \hat{\tau}_{ik}$. A plot of the estimated mean functions stratified by latent groups can be generated by S3 method plot(x, type = "EstMeans").

3 Package structure

The main function of R package SLCARE is SLCARE(). The dataset imported to function SLCARE() should take the long format, where each row corresponds to one time point of a subject at which a recurrent event is observed or censored. With a dataset coded in the wide format, where each row contains the timing information of all recurrent events observed or censored within one subject, users may convert such a dataset into the required long format by using function pivot_longer() from package tidyr (Wickham, Vaughan, and Girlich 2023) or function melt() from package reshape2 (Wickham 2007). In addition to allowing users to specify initial parameter estimates manually, SLCARE() offers a default initializer that implements the informative selection of initial values described in Section **Estimation procedure**. The built-in initializer performs K-means clustering of the observed data using function kmeans() from package stats and fitting multiplicative intensity models using reReg() from package reReg. When solving the estimating equations involved in the iterative estimation algorithm, SLCARE() performs weighted Poisson regression using glm.fit() from package stats and weighted multinomial regression using multinom() from package nnet. After running SLCARE(), model fitting results can be easily extracted from the output with S3 methods summary(), print(), predict(), and plot(). Graphical results generated by SLCARE() are presented via ggplot2 environment and are fully customizable.

Table 1 lists the main function SLCARE(), corresponding S3 methods and other functions called by this function, along with brief descriptions of their purposes.

Table 1: An overview of the main function and imported functions in SLCARE

FUNCTION	PURPOSE
SLCARE()	Conduct LCA with recurrent events data based on semiparametric multiplicative intensity modeling.
summary()	Generic function; used to summarize estimates for model parameters along with the associated variance estimates and p values.
print()	Generic function; used to initial estimates for the estimation algorithm
predict() plot()	convergence criterion, latent class membership probability and predicted number of recurrent events. Generic function; used to predict the posterior number of recurrent events. Generic function; used to generate cumulative baseline intensity function estimated mean function, model checking plot.
reReg() multinom() kmeans() glm.fit()	Imported from package reReg; used to find an initial estimate for β_0 . Imported from package nnet; used to find an initial estimate for α_0 ; used to solve the estimating equation to update the estimate for α_0 . Imported from package stats; used to find initial estimates for α_0 and β_0 . Imported from package stats; used to solve the estimating equation to
S()	update the estimate for β_0 .

4 Illustrations

The package **SLCARE** is designed to conduct LCA of recurrent events data based on semiparametric multiplicative intensity modeling (Zhao, Peng, and Hanfelt 2022). The main function SLCARE() has function arguments that enable flexible model specification and method implementation. In the below, we illustrate the use of SLCARE() through simulated datasets.

Input dataset

An input dataset for SLCARE() is a data frame containing time to recurrent events or censoring along with time-independent covariates of interest. The input dataset should take the long format, where each row contains the information in one time interval in which a recurrent event is observed or censored. As such, multiple rows in the dataset may correspond to one subject. As mentioned in Section **Package structure**, a dataset in the wide format data can be readily converted into the long format required by SLCARE() with the use of function pivot_longer() from package tidyr or function melt() from package reshape2.

The package SLCARE has a build-in dataset, SimData, which is a simulated dataset perturbed from a real dataset. SimData contains 48 subjects. For each subject, SimData contains values for the following six variables: id, which is the unique identifier for each subject; start, which records the starting time in minutes of the counting process interval; stop, which records the ending time in minutes of the counting process interval. If the subject entered the study at time zero, this column represents the time in minutes from the baseline visit to the occurrence of event or censoring; event, which indicates whether a recurrent event is observed (event = 1) or not (event = θ) at the end of the counting process interval; x1, which is a binary covariate; x2, which is a continuous covariate. Below we display how the data from one example subject are recorded in SimData:

```
#> # A tibble: 3 x 6
#> # Groups: id [1]
#>
    id start stop event
                             х1
#>
    <chr> <dbl> <int> <int> <int> <dbl>
          0 1950
                      1
#> 1 G052
                           1 0 444
#> 2 G052
         1950 2580
                        1
                              1 0.444
#> 3 G052
          2580 7085
                        0
                              1 0.444
```

As shown above, subject 6052 experienced two current events at time 1950 and time 2580 before the censoring time 7085. The time-independent covariates, x1 and x2, remain unchanged across different time points within the same subject.

Command line and function arguments

The LCA of recurrent event data outlined in Section **Methodological background** can be carried out with a single command line SLCARE(). SLCARE() provides flexible function arguments, which are shown below:

```
#> function (formula = "x1 + x2", alpha = NULL, beta = NULL, data = data,
#> id_col = "id", start_col = "start", stop_col = "stop", event_col = "event",
#> K = NULL, gamma = 0, max_epochs = 500, conv_threshold = 0.01,
#> boot = NULL)
#> NULL
```

At minimum, SLCARE() requires the following three arguments:

data: a dataframe with the format similar to SimData.

K: pre-determined number of latent classes.

formula: a string containing the covariates of interest in data.

The optional arguments of SLCARE() are:

alpha: initial estimate for α_0 in the estimation procedure. The default is NULL, which represents the initial estimate for α_0 resulted from the automated initializer described in Section Estimation and inference procedures.

beta: initial estimate for β_0 in the estimation procedure. The default is NULL, which represents the initial estimate for α_0 resulted from the automated initializer described in Section Estimation and inference procedures.

id_col: name of the column that includes subject identifiers.

start_col: name of the column that records the start time of each at-risk time interval.

stop_col: name of the column that records the start time of each at-risk time interval.

event_col: name of the column that indicates whether a recurrent event is observed or not (i.e, 1=observed; 0=otherwise).

gamma: parameter that indicates the distribution of frailty W. The default is 0 which indicates model (1) holds without the subject-specific frailty (i.e., W=1); gamma = k indicates that W follows the Gamma(k,k) distribution.

max_epochs: maximum number of iterations for the estimation algorithm.

conv_threshold: convergence threshold for the estimation algorithm.

boot: number of bootstrap replicates used to obtain the standard error estimation. The default is NULL which indicates bootstrap is not conducted.

Output and illustration with a sample dataset

The following code is used to perform LCA of the recurrent event dataset SimData with two latent classes (i.e., K = 2), 20 bootstrap replicates, and covariates x1 and x2.

The default output/print of SLCARE() includes parameter estimates and the associated variance estimates and p-values, along with the model's relative entropy measure. The same results can also be obtained using the generic method summary(model1, digits = 3), which allows for manually setting the minimum number of significant digits.

```
model1
```

```
#> Call:
#> SLCARE(formula = "x1 + x2", data = SimData, id_col = "id", start_col = "start",
       stop_col = "stop", event_col = "event", K = 2, boot = 20)
#> Coefficients for Beta:
#>
       (intercept)
                          x1
                                 x2
#> class1
               3.21 -0.2350 -4.933
#> class2
                2.41 -0.0413 0.309
#>
#> Std. Errors for Beta:
#>
       (intercept) x1
               0.216 0.278 0.813
#> class1
               0.137 0.106 0.343
#> class2
#> P. Values for Beta:
         (intercept)
                        x1
#> class1
            4.81e-50 0.398 1.28e-09
#> class2
            8.97e-70 0.696 3.68e-01
#>
#> Coefficients for Alpha:
#>
           x1 x2
#> class1 0.00 0.00
#> class2 -0.71 4.81
#>
#> Std. Errors for Alpha:
#>
            x1 x2
#> class1 0.000 0.00
#> class2 0.928 1.88
#>
```

Recall that in model (2), $\alpha_{0,1} = 0$, indicating that class1 is used as the reference group for evaluating the odds ratio of being in another latent class. The numbers in the row corresponding to class2 represent covariate effects on the log odds ratio of belonging to class2 versus class1.

These results are also can be extracted separately with the generic method function print(model1, type = "Est"), print(model1, type = "PValue"), and print(model1, type = "Entropy").

Initial estimates for the estimation algorithm

Unless manually specifying the arguments alpha and beta, SLCARE() implements the automated initializer described in Section **Estimation and inference procedures**. The resulting initial estimates for β_0 and α_0 can be found through the following generic method function print():

```
print(model1, type = "Init")
#> $beta
#>
         (intercept)
                              x1
          2.01630 0.09073593 -0.45944133
#> class1
            2.56397 -0.24415627 0.06202144
#> class2
#>
#> $alpha
#>
                x1
                         x2
#> class1 0.0000000 0.000000
#> class2 0.5386871 2.985595
```

Convergence criterion

SLCARE() provides arguments to control the termination of the iterative estimation procedure. By default, max_epochs=500 and conv_threshold=0.01. This means that the iterations would stop when the number of iterations reaches 500 or the magnitude of the change in parameter estimates is below 0.01. In addition, users are allowed to check the change in parameter estimates in the last iteration using the following code:

```
print(model1, type = "ConvergeLoss")
#> [1] 0.00909132
```

If the output is greater than the pre-determined convergence threshold, users may consider increasing the maximum number of iterations and/or setting a larger convergence threshold.

Latent class membership probability

SLCARE() generates output on the estimated probability that a subject belong to different latent classes. The following are the example code and output in the case where one is interested in the class membership probabilities of the 8^{th} to 10^{th} subjects in SimData.

Predicted number of recurrent events

SLCARE() computes the predicted number of recurrent events for each subject according to equation (14). The code to extract such a result for the 8^{th} to 10^{th} subjects in SimData is shown below, along with the corresponding output:

predict(model1)[8:10,]

#>		ID	PosteriorPrediction
#>	8	G052	2.231237
#>	9	U0M043	2.892228
#>	10	G064	13.333784

Note that SLCARE() returns the estimated numbers of recurrent events as float rather than integer. Users may use the argument integer = TRUE in the genetic function predict() to round those numbers to integers.

predict(model1, integer = TRUE)[8:10,]

#>		ID	PosteriorPrediction
#>	8	G052	2
#>	9	U0M043	2
#>	10	G064	13

Model checking plot

SLCARE() generates a model checking plot following the procedure described in Section **Methodological background** via ggplot2 environment. The following shows the S3 method for plotting a model checking plot:

plot(model1, type = "ModelChecking")

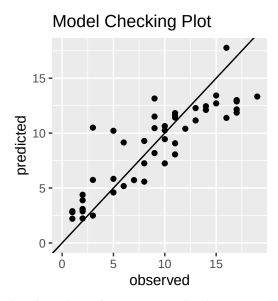


Figure 1: Predicted numbers of events versus the observe numbers of events.

Estimated cumulative baseline intensity function

By equation (8), SLCARE() calculates $\hat{\mu}(t)$, which is an estimates for $\mu_0(t)$, cumulative baseline intensity function. A plot of $\hat{\mu}(t)$ versus t can be generated with the following code:

```
plot(model1, type = "mu0")
```

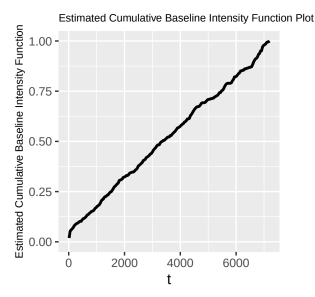


Figure 2: Estimated cumulative baseline intensity function.

Evaluation of $\hat{\mu}(t)$ at specific time points, for example, t=100,1000 and 5000, can be obtained by the following command:

```
model1$est_mu0(c(100, 1000, 5000))
#> [1] 0.06086907 0.17089670 0.70936436
```

Estimated mean function plot

Estimated mean function plots are useful for describing and comparing the expected number of recurrent events across different latent groups over time. SLCARE() computes the estimated mean function over time according to equation (15), which can be plotted with the following code:

plot(model1, type = "EstMeans")

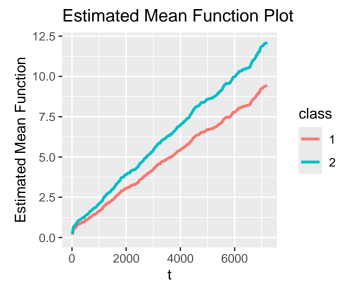


Figure 3: Estimated mean function for two latent classes.

5 A real application

We apply SLCARE to a dataset from a randomized phase III clinical trial FFCD 2000-05 conducted between 2002 and 2007 in patients diagnosed with advanced colorectal cancer (Ducreux et al. 2011).

The dataset is publicly available in **frailtypack** package (Rondeau, Marzroui, and Gonzalez 2012) and is also included in the **SLCARE** package. One recurrent event of interest is the appearance of new colorectal lesions. The colorectal lesion is the region in a colon that has suffered damage through colorectal cancer. The appearance of new colorectal lesions indicates colorectal cancer progression. The main interest of our analysis is to explore the heterogeneity in the recurrence patterns of colorectal lesions.

To address this interest, we extract the dataset colorectal SLCARE from the colorectal dataset. The dataset 'colorectal SLCARE is arranged in the long format, consisting of 150 patients and 289 records.

Each record stores the value of the six variables described as follows. Variable id is the unique patient identifier. Variable time0 is the start time (in years) of the recurrent event interval. Variable time1 records the interval end time (in years) of the detection of new colorectal lesions or terminal event (death or right-censoring). Variable new.lesions = 1 indicates that the appearance of a new colorectal lesion is observed and new.lesions = 0 indicates that a terminal event is observed. The other two variables represent covariates of interest: (1) treatment, which is coded as treatment = 1 if the patient received a combination chemotherapy and as treatment = 0 if the patient received sequential chemotherapy; (2) previous resection indicator, which is coded as prev.resection = 1 if the patient had a previous resection and as prev.resection = 0 otherwise.

The construction and the structure of the colorectal SLCARE dataset are presented below.

```
data("colorectal", package = "SLCARE")
colorectalSLCARE <- colorectal |>
 dplyr::select(id, time0, time1, new.lesions, treatment, prev.resection) |>
 dplyr::mutate(id = paste0("patient", id), treatment = as.numeric(treatment) -1,
               prev.resection = as.numeric(prev.resection) -1)
str(colorectalSLCARE, vec.len = 3)
#> 'data.frame':
                  289 obs. of 6 variables:
                 : chr "patient1" "patient2" "patient3" ...
#> $ id
#> $ time0
                 : num 0000.525 ...
#> $ time1
                 : num 0.71 1.282 0.525 0.921 ...
#> $ new.lesions : int 0 0 1 1 0 1 0 1 ...
#> $ treatment : num 0 1 0 0 0 1 1 0 ...
  $ prev.resection: num  0  0  0  0  1  1  0 ...
```

We fit models (1) and (2) to the dataset colorectal_SLCARE with $T_i^{(j)}$ corresponding to time to the j-th detection of new colorectal lesions. We consider three candidate distributions for W, the frailty term in model (1), which are W=1 and Gamma(r,r), r=1, 3. For each candidate $f_W(\cdot)$, we calculate the relative entropy measure Entropy as described in Section **Estimation and inference procedures** using the command illustrated in Section **Illustrations**. The results are presented in Table 2.

Table 2: Real data example: Relative entropy calculated with different choices of K and $f_W(w)$.

	K = 2	K = 3
Gamma(1,1)	0.785	0.788
Gamma(3,3)	0.802	0.793
W = 1	0.462	0.459

As shown in Table 2, the maximum relative entropy is achieved with the combination of K=2 and Gamma(3,3). Therefore, we set K=2 and select $f_W(\cdot)$ as the density of Gamma(3,3) for the rest of the analysis.

```
set.seed(66)
finalmodel <- SLCARE(formula = "treatment + prev.resection", data = colorectalSLCARE,
        id_col = "id", start_col = "time0", stop_col = "time1", event_col = "new.lesions",
        K = 2, gamma = 3, boot = 200)
summary(finalmodel, digits = 3)

#> Call:
#> SLCARE(formula = "treatment + prev.resection", data = colorectalSLCARE,
#> id_col = "id", start_col = "time0", stop_col = "time1", event_col = "new.lesions",
```

```
#>
      K = 2, gamma = 3, boot = 200)
#>
#> Coefficients for Beta:
         (intercept) treatment prev.resection
#>
             1.696
                      -0.415
                                      -0.493
#> class1
#> class2
               0.811
                         0.691
                                        0.494
#>
#> Std. Errors for Beta:
         (intercept) treatment prev.resection
             0.34
                      0.281
#> class2
                0.48
                         0.887
                                        0.703
#>
#> P. Values for Beta:
#>
         (intercept) treatment prev.resection
#> class1
            6.26e-07
                      0.139 0.0751
#> class2
            9.11e-02
                         0.436
                                       0.4823
#> Coefficients for Alpha:
     treatment prev.resection
#> class1
             0.0
#> class2
             -12.3
                            -11.7
#>
#> Std. Errors for Alpha:
#>
       treatment prev.resection
              0.00
#> class1
                             0.00
                             1.26
#> class2
              3.96
#>
#> P. Values for Alpha:
         treatment prev.resection
#> class1
              NaN
#> class2 0.00195
                         1.49e-20
#>
#> Relative Entropy: 0.802
```

Based on the results from fitting models (1) and (2), we classify patients into two groups according to the modal class assignment rule, i.e., $\hat{\xi}_i = \arg\max_{1 \leq k \leq K} \hat{\tau}_{ik}$. Table 3 (generated by R package table1 (Rich 2023)) summarizes the characteristics of the resulting two groups, Class 1 of size 127 and Class 2 of size 23. Comparing the total number of the observed colorectal lesions, captured by $N_i(C_i)$, we note that patients belong to Class 1 tend to experience more colorectal lesions as compared to those belong to Class 2. There is also notable separation in covariates treatment and prev. resection. That is, over 50% patients in Class 1 received combination chemotherapy in contrast to 0% patients in Class 2 on combination chemotherapy, and over 70% patients in Class 1 has previous resection, while all patients in Class 2 did not have resection in the past. These observations are consistent with the estimation results for α_0 (see model summary), which suggest that patients receiving combination chemotherapy or with previous resection are much less likely to be classified to Class 2. These observations entail a plausible characterization of the two latent classes. That is, Class 1 consists of patients with more aggressive progression that incurs more intensive past or ongoing treatment, while Class 2 is featured by relatively mild disease that requires less complicated treatment regimens.

We next examine the covariate effects within each latent class. The coefficient estimates presented in model summary indicate a reasonable direction of how treatment and prev.resection influence the intensity of lesion recurrence. That is, combination chemotherapy or previous resection may help reduce the risk of lesion recurrence. However, the estimated effects which conform to our intuition are associated with p values that do not reach statistical significance.

Table 3: Real data example	: Characteristics of the two latent classes.
----------------------------	----------------------------------------------

	Class 1	Class 2	P-value
	(N=127)	(N=23)	
observed.events			
Mean (SD)	1.02 (0.988)	0.391 (0.583)	< 0.001
Median [Min, Max]	1.00 [0, 4.00]	0 [0, 2.00]	
treatment			
sequential chemotherapy	54 (42.5%)	23 (100%)	< 0.001
combination chemotherapy	73 (57.5%)	0 (0%)	
prev.resection			
no	37 (29.1%)	23 (100%)	< 0.001
yes	90 (70.9%)	0 (0%)	

In Figure 4, we plot the average estimated mean function of experiencing new colorectal lesions for the two latent classes. Figure 4 shows that Class 1 is associated with a lower frequency of experiencing new colorectal lesions than Class 2. Such a distinction may be contributed by the underlying difference in disease severity between the two latent classes, which is also manifested by the different distributions of receiving combination chemotherapy and having previous resection between these two classes demonstrated in Table 3.

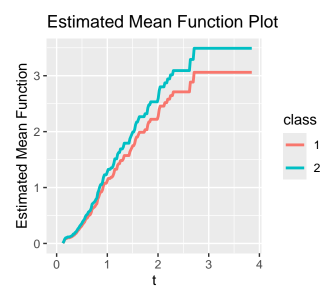


Figure 4: Real data example: Estimated mean function of experiencing new colorectal lesions for two classes.

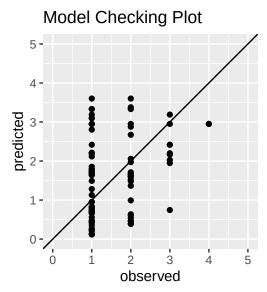


Figure 5: Real data example: Predicted numbers of new colorectal lesions versus the observed numbers of new colorectal lesions.

We further check the overall fit of the latent class models adopted by SLCARE() to the colorectalSLCARE dataset using the model checking plot described in Section **Methodological background**. As shown in Figure 5, the dots representing the predicted and the observed numbers of new colorectal lesions are generally balanced around the 45-degree line when there are two observed colorectal lesions. However, over-prediction and under-prediction are noted when the observed number of of colorectal lesions is one or three respectively. These observations may suggest a moderate lack-of-fit of the fitted model to this colorectal dataset.

6 Discussion

The R package SLCARE provides an easy-to-use software to conduct latent class analysis of recurrent events based on a flexible semiparametric multiplicative intensity modeling proposed by Zhao, Peng, and Hanfelt (2022). A practical automated initializer is embedded in SLCARE to help users set initial estimates in an informative way. SLCARE provides a single command line function to implement full analysis with optional arguments for model customization. SLCARE also provides visualization tools for result summary and model evaluation with ggplot2 plotting environment.

The R package **SLCARE** is maintained in both Comprehensive R Archive Network (CRAN) at https://CRAN.R-project.org/package=SLCARE and Github at https://github.com/qyxxx/SLCARE. There are several interesting directions to further improve this package. For example, computational speed may be improved with more efficient coding. A new option may be added to perform automated selection of *K* and frailty distribution. Such updates of package **SLCARE** will be made available at CRAN and Github.

References

Andersen, Per Kragh, and Richard D Gill. 1982. "Cox's Regression Model for Counting Processes: A Large Sample Study." *The Annals of Statistics*. https://www.jstor.org/stable/2240714.

Celeux, Gilles, and Gilda Soromenho. 1996. "An Entropy Criterion for Assessing the Number of Clusters in a Mixture Model." *Journal of Classification*. https://doi.org/10.1007/BF01246098.

Chiou, Sy Han, Gongjun Xu, Jun Yan, and Chiung-Yu Huang. 2023. "Regression Modeling for Recurrent Events Possibly with an Informative Terminal Event Using R Package reReg." *Journal of Statistical Software*. https://doi.org/10.18637/jss.v105.i05.

Clement, David Y, and Robert L Strawderman. 2009. "Conditional GEE for Recurrent Event Gap Times." *Biostatistics*. https://doi.org/10.1093/biostatistics/kxp004.

Ducreux, Michel, David Malka, Jean Mendiboure, Pierre-Luc Etienne, Patrick Texereau, Dominique Auby, Philippe Rougier, et al. 2011. "Sequential Versus Combination Chemotherapy for the Treatment of Advanced Colorectal Cancer (FFCD 2000–05): An Open-Label, Randomised, Phase 3 Trial." *The Lancet Oncology*. https://doi.org/10.1016/S1470-2045(11)70199-1.

- Fine, JP, J Yan, and MR Kosorok. 2004. "Temporal Process Regression." *Biometrika*. https://doi.org/10.1093/biomet/91.3.683.
- Grün, Bettina, and Friedrich Leisch. 2008. "FlexMix Version 2: Finite Mixtures with Concomitant Variables and Varying and Constant Parameters." *Journal of Statistical Software*. https://doi.org/10.18637/jss.v028.i04.
- Harrell Jr, Frank E. 2023. rms: Regression Modeling Strategies. https://CRAN.R-project.org/package=rms.
- Lloyd, Stuart. 1982. "Least Squares Quantization in PCM." *IEEE Transactions on Information Theory*. https://doi.org/10.1109/TIT.1982.1056489.
- Proust-Lima, Cécile, Viviane Philipps, and Benoit Liquet. 2017. "Estimation of Extended Mixed Models Using Latent Classes and Latent Processes: The R Package lcmm." *Journal of Statistical Software*. https://doi.org/10.18637/jss.v078.i02.
- Ramaswamy, Venkatram, Wayne S DeSarbo, David J Reibstein, and William T Robinson. 1993. "An Empirical Pooling Approach for Estimating Marketing Mix Elasticities with PIMS Data." *Marketing Science*. https://doi.org/10.1287/mksc.12.1.103.
- Rich, Benjamin. 2023. table1: Tables of Descriptive Statistics in HTML. https://CRAN.R-project.org/package=table1.
- Rondeau, Virginie, Yassin Marzroui, and Juan R Gonzalez. 2012. "frailtypack: An R Package for the Analysis of Correlated Survival Data with Frailty Models Using Penalized Likelihood Estimation or Parametrical Estimation." *Journal of Statistical Software*. https://doi.org/10.18637/jss.v047.i04.
- Stefanski, Leonard A, and Raymond J Carroll. 1987. "Conditional Scores and Optimal Scores for Generalized Linear Measurement-Error Models." *Biometrika*. https://doi.org/10.1093/biomet/74.4.703.
- Therneau, Terry M. 2023. A Package for Survival Analysis in R. https://CRAN.R-project.org/package=survival.
- Venables, W. N., and B. D. Ripley. 2002. *Modern Applied Statistics with S.* https://www.stats.ox.ac.uk/pub/MASS4/.
- Wang, Mei-Cheng, Jing Qin, and Chin-Tsang Chiang. 2001. "Analyzing Recurrent Event Data with Informative Censoring." *Journal of the American Statistical Association*. https://doi.org/10.1198/016214501753209031.
- Wickham, Hadley. 2007. "Reshaping Data with the reshape Package." *Journal of Statistical Software*. http://www.jstatsoft.org/v21/i12/.
- . 2016. *ggplot2: Elegant Graphics for Data Analysis*. Springer-Verlag New York. https://ggplot2.tidyverse.org.
- Wickham, Hadley, Davis Vaughan, and Maximilian Girlich. 2023. *tidyr: Tidy Messy Data*. https://CRAN.R-project.org/package=tidyr.
- Yan, Jun, and Jason Fine. 2004. "Estimating Equations for Association Structures." *Statistics in Medicine*. https://doi.org/10.1002/sim.1650.
- Zhao, Wei, Limin Peng, and John Hanfelt. 2022. "Semiparametric Latent Class Analysis of Recurrent Event Data." *Journal of the Royal Statistical Society Series B: Statistical Methodology.* https://doi.org/10.1111/rssb.12499.

Qi Yu Emory University Department of Biostatistics and Bioinformatics 1518 Clifton Rd. NE, Atlanta, GA 30322, USA ORCiD: 0009-0007-7585-1183 qi.yu2@emory.edu

Limin Peng
Emory University
Department of Biostatistics and Bioinformatics
1518 Clifton Rd. NE, Atlanta, GA 30322, USA
ORCiD: 0000-0002-8042-3112
lpeng@emory.edu