SOFTWARE

joineRML: A joint model and software package for time-to-event and multivariate longitudinal outcomes

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Abstract

Background: Joint modelling of longitudinal and time-to-event outcomes has received considerable attention over recent years. Commensurate with this has been a rise in statistical software options for fitting these models. However, these tools have generally been limited to a single longitudinal outcome. Here, we describe the classical joint model to the case of *multiple* longitudinal outcomes, propose a practical algorithm for fitting the models, and demonstrate how to fit the models using a new package for the statistical software platform R, joineRML.

Results: A multivariate linear mixed sub-model is specified for the longitudinal outcomes, and a Cox proportional hazards regression model with time-varying covariates is specified for the event time sub-model. The association between models is captured through a zero-mean multivariate latent Gaussian process. The models are fitted using a Monte Carlo Expectation-Maximisation algorithm, and inferences are based on approximate standard errors from the empirical profile information matrix, which are contrasted to an alternative bootstrap estimation approach. We illustrate the model and software on a real data example for patients with primary biliary cirrhosis with three repeatedly measured biomarkers.

Conclusions: An open-source software package capable of fitting multivariate joint models is available. The underlying algorithm and source code makes use of several methods to increase computational speed.

Keywords: Joint modelling; Longitudinal data; Multivariate data; Time-to-event data; Software

Background

1

- ⁴ In many clinical studies, subjects are followed-up repeatedly and response data col-
- 5 lected. For example, routine blood tests might be performed at each follow-up clinic
- ⁶ appointment for patients enrolled in a randomized drug trial, and biomarker mea-

surements recorded. An event time is also usually of interest, for example time of death or study drop-out. It has been repeatedly shown elsewhere that if the longi-8 tudinal and event-time outcomes are correlated, then modelling the two outcome processes separately, for example using linear mixed models and Cox regression 10 models, can lead to biased effect size estimates [1]. The same criticism has also 11 been levelled at the application of so-called two-stage models [2]. The motivation 12 for using joint models can be broadly separated into interest in drawing inference 13 about (1) the time-to-event process whilst adjusting for the intermittently measured 14 (and potentially error-prone) longitudinal outcomes, and (2) the longitudinal data 15 process whilst adjusting for a potentially informative drop-out mechanism [3]. The 16 literature on joint modelling is extensive, with excellent reviews given by Tsiatis 17 and Davidian [4], Gould *et al.* [5], and the monologue by Rizopoulos [6]. 18

Joint modelling has until recently been predominated by modelling a single lon-19 gitudinal outcome together with a solitary event time outcome; herein referred to 20 as univariate joint modelling. Commensurate with this methodological research has 21 been an increase in wide-ranging clinical applications (e.g. [7]). Recent innovations in 22 the field of joint models have included the incorporation of multivariate longitudinal 23 data [8], competing risks data [9, 10], recurrent events data [11], multivariate time-24 to-event data [12, 13], non-continuous repeated measurements (e.g. count, binary, 25 ordinal, and censored data) [14], non-normally and non-parametrically distributed 26 random effects [15], alternative estimation methodologies (e.g. Bayesian fitting and 27 conditional estimating equations) [16, 17], and different association structures [18]. 28 In this article, we specifically focus on the first innovation: multivariate longitudinal 29 data. In this situation, we assume that multiple longitudinal outcomes are measured 30 on each subject, which can be unbalanced and measured at different times for each 31 subject. 32

Despite the inherently obvious benefits of harnessing all data in a single model 33 or the published research on the topic of joint models for multivariate longitudinal 34 data, a recent literature review by Hickey et al. [19] identified that publicly avail-35 able software for fitting such models was lacking, which has translated into limited 36 uptake by biomedical researchers. In this article we present the classical joint model 37 described by Henderson et al. [3] extended to the case of multiple longitudinal out-38 comes. An algorithm proposed by Lin et al. [20] is used to fit the model, augmented by techniques to reduce the computational fitting time, including a quasi-Newton 40 update approach, variance reduction method, and dynamic Monte Carlo updates. 41 This algorithm is encoded into a R sofware package-joineRML. A simulation anal-42

43 ysis and real-world data example are used to demonstrate the accuracy of the algo-

⁴⁴ rithm and the software, respectively.

45 Implementation

⁴⁶ As a prelude to the introduction and demonstration of the newly introduced software

⁴⁷ package, in the following section we describe the underlying model formulation and
⁴⁸ model fitting methodology.

49 Model

For each subject i = 1, ..., n, $\boldsymbol{y}_i = (\boldsymbol{y}_{i1}^{\top}, ..., \boldsymbol{y}_{iK}^{\top})$ is the K-variate continuous out-50 come vector, where each y_{ik} denotes an $(n_{ik} \times 1)$ -vector of observed longitudinal 51 measurements for the k-th outcome type: $\boldsymbol{y}_{ik} = (y_{i1k}, \dots, y_{in_{ik}k})^{\top}$. Each outcome is 52 measured at observed (possibly pre-specified) times t_{ijk} for $j = 1, \ldots, n_{ik}$, which can 53 differ between subjects and outcomes. Additionally, for each subject there is an event 54 time T_i^* , which is subject to right censoring. Therefore, we observe $T_i = \min(T_i^*, C_i)$, 55 where C_i corresponds to a potential censoring time, and the failure indicator δ_i , 56 which is equal to 1 if the failure is observed $(T_i^* \leq C_i)$ and 0 otherwise. We assume 57 that both censoring and measurement times are non-informative. 58

The model we describe is the natural extension of the model proposed by Henderson *et al.* [3] to the case of multivariate longitudinal data. The model posits an unobserved or latent zero-mean (K+1)-variate Gaussian process that is realised independently for each subject, $W_i(t) = \left\{ W_{1i}^{(1)}(t), \ldots, W_{1i}^{(K)}(t), W_{2i}(t) \right\}$. This latent process subsequently links the separate sub-models via association parameters.

The k-th longitudinal data sub-model is given by

$$y_{ik}(t) = \mu_{ik}(t) + W_{1i}^{(k)}(t) + \varepsilon_{ik}(t), \tag{1}$$

where $\mu_{ik}(t)$ is the mean response, and $\varepsilon_{ik}(t)$ is the model error term, which we assume to be independent and identically distributed normal with mean 0 and variance σ_k^2 . The mean response is specified as a linear model

$$\mu_{ik}(t) = \boldsymbol{x}_{ik}^{\mathsf{T}}(t)\boldsymbol{\beta}_k,\tag{2}$$

where $\boldsymbol{x}_{ik}(t)$ is a p_k -vector of (possibly) time-varying covariates with corresponding fixed effect terms $\boldsymbol{\beta}_k$. $W_{1i}^{(k)}(t)$ is specified as

$$W_{1i}^{(k)}(t) = \boldsymbol{z}_{ik}^{\top}(t)\boldsymbol{b}_{ik},\tag{3}$$

where $z_{ik}(t)$ is an r_k -vector of (possibly) time-varying covariates with corresponding subject-and-outcome random effect terms b_{ik} , which follow a zero-mean mul-65 tivariate normal distribution with $(r_k \times r_k)$ -variance-covariance matrix D_{kk} . To 66 account for dependence between the different longitudinal outcome outcomes, we 67 let $cov(\mathbf{b}_{ik}, \mathbf{b}_{il}) = \mathbf{D}_{kl}$ for $k \neq l$. Furthermore, we assume $\varepsilon_{ik}(t)$ and \mathbf{b}_{ik} are uncor-68 related, and that the censoring times are independent of the random effects. These 69 distributional assumptions together with the model given by (1)-(3) are equivalent 70 to the multivariate extension of the Laird and Ware [21] linear mixed effects model. 71 More flexible specifications of $W_{1i}^{(k)}(t)$ can be used [3], including for example, sta-72 tionary Gaussian processes. However, we do not consider these cases here owing to 73 the increased computational burden it carries, even for the univariate case. 74

The sub-model for the time-to-event outcome is given by the hazard model

$$\lambda_i(t) = \lambda_0(t) \exp\left\{\boldsymbol{v}_i^{\top}(t)\boldsymbol{\gamma}_v + W_{2i}(t)\right\},\,$$

where $\lambda_0(\cdot)$ is an unspecified baseline hazard, and $\boldsymbol{v}_i(t)$ is a *q*-vector of (possibly) time-varying covariates with corresponding fixed effect terms $\boldsymbol{\gamma}_v$. Conditional on $W_i(t)$ and the observed covariate data, the longitudinal and time-to-event data generating processes are conditionally independent. To establish a latent association, we specify $W_{2i}(t)$ as a linear combination of $\left\{W_{1i}^{(1)}(t), \ldots, W_{1i}^{(K)}(t)\right\}$:

$$W_{2i}(t) = \sum_{k=1}^{K} \gamma_{yk} W_{1i}^{(k)}(t),$$

where $\gamma_y = (\gamma_{y1}, \ldots, \gamma_{yK})$ are the corresponding association parameters. To emphasise the dependence of $W_{2i}(t)$ on the random effects, we explicitly write it as $W_{2i}(t, \mathbf{b}_i)$ from here onwards. As per $W_{1i}^{(k)}(t)$, $W_{2i}(t, \mathbf{b}_i)$ can also be flexibly extended, for example to include subject-specific frailty effects [3].

79 Estimation

80 Likelihood

For each subject *i*, let $\mathbf{X}_i = \bigoplus_{k=1}^K \mathbf{X}_{ik}$ and $\mathbf{Z}_i = \bigoplus_{k=1}^K \mathbf{Z}_{ik}$ be block-diagonal matrices, where $\mathbf{X}_{ik} = (\mathbf{x}_{i1k}^\top, \dots, \mathbf{x}_{in_{ik}k}^\top)$ is an $(n_{ik} \times p_k)$ -design matrix, with the *j*-th row corresponding to the p_k -vector of covariates measured at time t_{ijk} , and \bigoplus denotes the direct matrix sum. The notation similarly follows for the random effects design matrices, \mathbf{Z}_{ik} . We denote the error terms by a diagonal matrix $\mathbf{\Sigma}_i = \bigoplus_{k=1}^K \sigma_k^2 \mathbf{I}_{n_{ik}}$ and write the overall variance-covariance matrix for the random effects

as

$$oldsymbol{D} = \left(egin{array}{cccc} oldsymbol{D}_{11} & \cdots & oldsymbol{D}_{1K} \ dots & \ddots & dots \ oldsymbol{D}_{1K}^{ op} & \cdots & oldsymbol{D}_{KK} \end{array}
ight),$$

where I_n denotes an $n \times n$ identity matrix. We further define $\boldsymbol{\beta} = (\boldsymbol{\beta}_1^{\top}, \dots, \boldsymbol{\beta}_K^{\top})^{\top}$ and $\boldsymbol{b}_i = (\boldsymbol{b}_{i1}^{\top}, \dots, \boldsymbol{b}_{iK}^{\top})^{\top}$. Hence, we can then rewrite the longitudinal outcome sub-model as

For the estimation, we will assume that the covariates in the time-to-event submodel are time-independent and known at baseline, i.e. $v_i \equiv v_i(0)$. Extensions of the estimation procedure for time-varying covariates are outlined elsewhere [6, p. 115]. The *observed* data likelihood for the joint outcome is given by

$$\prod_{i=1}^{n} \left(\int_{-\infty}^{\infty} f(\boldsymbol{y}_{i} | \boldsymbol{b}_{i}, \boldsymbol{\theta}) f(T_{i}, \delta_{i} | \boldsymbol{b}_{i}, \boldsymbol{\theta}) f(\boldsymbol{b}_{i} | \boldsymbol{\theta}) d\boldsymbol{b}_{i} \right),$$
(4)

where $\boldsymbol{\theta} = (\boldsymbol{\beta}^{\top}, \operatorname{vech}(\boldsymbol{D}), \sigma_1^2, \dots, \sigma_K^2, \lambda_0(t), \boldsymbol{\gamma}_v^{\top}, \boldsymbol{\gamma}_y^{\top})$ is the collection of unknown parameters that we want to estimate, with $\operatorname{vech}(\boldsymbol{D})$ denoting the half-vectorisation operator that returns the vector of lower-triangular elements of matrix \boldsymbol{D} .

As noted by Henderson *et al.* [3], the observed data likelihood can be calculated by rewriting it as

$$\prod_{i=1}^{n} f(\boldsymbol{y}_{i} | \boldsymbol{\theta}) \left(\int_{-\infty}^{\infty} f(T_{i}, \delta_{i} | \boldsymbol{b}_{i}, \boldsymbol{\theta}) f(\boldsymbol{b}_{i} | \boldsymbol{y}_{i}, \boldsymbol{\theta}) d\boldsymbol{b}_{i} \right),$$

where the marginal distribution $f(\boldsymbol{y}_i | \boldsymbol{\theta})$ is a multivariate normal density with mean $X_i \boldsymbol{\beta}$ and variance-covariance matrix $\boldsymbol{\Sigma}_i + \boldsymbol{Z}_i \boldsymbol{D} \boldsymbol{Z}_i^{\top}$, and $f(\boldsymbol{b}_i | \boldsymbol{y}_i, \boldsymbol{\theta})$ is given by (6).

89 MCEM algorithm

We determine maximum likelihood estimates of the parameters θ using the Monte Carlo Expectation Maximisation (MCEM) algorithm [22], by treating the random effects b_i as missing data. This is effectively the same as the conventional Expectation-Maximisation (EM) algorithm, as used by Wulfsohn and Tsiatis [23] and Ratcliffe *et al.* [24] in the context of fitting univariate data joint models, except

- ⁹⁵ the E-step exploits a Monte Carlo (MC) integration routine as opposed to Gaus-
- $_{\rm 96}$ $\,$ sian quadrature methods, which we expect to be beneficial when the dimension of

⁹⁷ random effects becomes large.

- Starting from an initial estimate of the parameters, $\hat{\theta}^{(0)}$, the procedure involves iterating between the following two steps until convergence is achieved.
- 1 E-step. At the (m+1)-th iteration, we compute the expected log-likelihood of the complete data conditional on the observed data and the current estimate of the parameters,

$$Q(\boldsymbol{\theta} \mid \hat{\boldsymbol{\theta}}^{(m)}) = \sum_{i=1}^{n} \mathbb{E} \Big\{ \log f(\boldsymbol{y}_{i}, T_{i}, \delta_{i}, \boldsymbol{b}_{i} \mid \boldsymbol{\theta}) \Big\}$$
$$= \sum_{i=1}^{n} \int_{-\infty}^{\infty} \Big\{ \log f(\boldsymbol{y}_{i}, T_{i}, \delta_{i}, \boldsymbol{b}_{i} \mid \boldsymbol{\theta}) \Big\} f(\boldsymbol{b}_{i} \mid T_{i}, \delta_{i}, \boldsymbol{y}_{i}; \hat{\boldsymbol{\theta}}^{(m)}) d\boldsymbol{b}_{i}$$

Here, the complete-data likelihood contribution for subject i is given by the integrand of (4).

2 *M-step.* We maximise $Q(\boldsymbol{\theta} \mid \hat{\boldsymbol{\theta}}^{(m)})$ with respect to $\boldsymbol{\theta}$. Namely, we set

$$\hat{\boldsymbol{\theta}}^{(m+1)} = \operatorname*{argmax}_{\boldsymbol{\theta}} Q(\boldsymbol{\theta} \,|\, \hat{\boldsymbol{\theta}}^{(m)}).$$

The M-step estimators naturally follow from Wulfsohn and Tsiatis [23] and Lin *et al.* [20]. Maximizers for all parameters except γ_v and γ_y are available in closed-form; algebraic details are presented in **Additional file 1**. The parameters $\boldsymbol{\gamma} = (\boldsymbol{\gamma}_v^{\top}, \boldsymbol{\gamma}_y^{\top})^{\top}$ are jointly updated using a one-step Newton-Raphson algorithm as

$$\hat{\boldsymbol{\gamma}}^{(m+1)} = \hat{\boldsymbol{\gamma}}^{(m)} + I\left(\hat{\boldsymbol{\gamma}}^{(m)}\right)^{-1} S\left(\hat{\boldsymbol{\gamma}}^{(m)}\right),$$

where $\hat{\gamma}^{(m)}$ denotes the value of γ at the current iteration, $S(\hat{\gamma}^{(m)})$ is the corre-105 sponding score, and $I(\hat{\gamma}^{(m)})$ is the observed information matrix, which is equal to 106 the derivative of the negative score. Further details of this update are given in Addi-107 tional file 1. The M-step for γ is computationally expensive to evaluate. Therefore, 108 we also propose a quasi-Newton one-step update by approximating $I(\hat{\gamma}^{(m)})$ by an 109 empirical information matrix for γ , which can be considered an analogue of the 110 Gauss-Newton method [25, p. 8]. To further compensate for this approximation, 111 we also use a nominal step-size of 0.5 rather than 1, which is used when exactly 112 calculating $I(\boldsymbol{\gamma})$. 113

The M-step involves terms of the form $\mathbb{E}\left[h(\boldsymbol{b}_i) \mid T_i, \delta_i, \boldsymbol{y}_i; \hat{\boldsymbol{\theta}}\right]$, for known functions $h(\cdot)$. The conditional expectation of a function of the random effects can be written

as

$$\mathbb{E}\left[h(\boldsymbol{b}_{i}) \mid T_{i}, \delta_{i}, \boldsymbol{y}_{i}; \hat{\boldsymbol{\theta}}\right] = \frac{\int_{-\infty}^{\infty} h(\boldsymbol{b}_{i}) f(\boldsymbol{b}_{i} \mid \boldsymbol{y}_{i}; \hat{\boldsymbol{\theta}}) f(T_{i}, \delta_{i} \mid \boldsymbol{b}_{i}; \hat{\boldsymbol{\theta}}) d\boldsymbol{b}_{i}}{\int_{-\infty}^{\infty} f(\boldsymbol{b}_{i} \mid \boldsymbol{y}_{i}; \hat{\boldsymbol{\theta}}) f(T_{i}, \delta_{i} \mid \boldsymbol{b}_{i}; \hat{\boldsymbol{\theta}}) d\boldsymbol{b}_{i}},$$
(5)

where $f(T_i, \delta_i | \boldsymbol{b}_i; \hat{\boldsymbol{\theta}})$ is given by

$$f(T_i, \delta_i | \boldsymbol{b}_i; \boldsymbol{\theta}) = \left[\lambda_0(T_i) \exp\left\{\boldsymbol{v}_i^\top \boldsymbol{\gamma}_v + W_{2i}(T_i, \boldsymbol{b}_i)\right\}\right]^{\delta_i} \\ \times \exp\left\{-\int_0^{T_i} \lambda_0(u) \exp\left\{\boldsymbol{v}_i^\top \boldsymbol{\gamma}_v + W_{2i}(u, \boldsymbol{b}_i)\right\} du\right\}$$

and $f(\boldsymbol{b}_i | \boldsymbol{y}_i; \hat{\boldsymbol{\theta}})$ is calculated from multivariate normal distribution theory as

$$\boldsymbol{b}_{i} | \boldsymbol{y}_{i}, \boldsymbol{\theta} \sim N\left(\boldsymbol{A}_{i}\left\{\boldsymbol{Z}_{i}^{\top}\boldsymbol{\Sigma}_{i}^{-1}(\boldsymbol{y}_{i}-\boldsymbol{X}_{i}\boldsymbol{\beta})\right\}, \boldsymbol{A}_{i}\right),$$
(6)

with $\mathbf{A}_i = \left(\mathbf{Z}_i^{\top} \mathbf{\Sigma}_i^{-1} \mathbf{Z}_i + \mathbf{D}^{-1}\right)^{-1}$. As this becomes computationally expensive using Gaussian quadrature commensurate with increasing dimension of \mathbf{b}_i , we estimate the integrals by MC sampling such that the expectation is approximated by the ratio of the sample means for $h(\mathbf{b}_i)f(T_i, \delta_i | \mathbf{b}_i; \hat{\boldsymbol{\theta}})$ and $f(T_i, \delta_i | \mathbf{b}_i; \hat{\boldsymbol{\theta}})$ evaluated at each MC draw. Furthermore, we use antithetic simulation for variance reduction in the MC integration. Instead of directly sampling from (6), we sample $\mathbf{\Omega} \sim N(0, \mathbf{I}_r)$ and obtain the *pairs*

$$oldsymbol{A}_i\left\{oldsymbol{Z}_i^ op oldsymbol{\Sigma}_i^{-1}(oldsymbol{y}_i-oldsymbol{X}_ioldsymbol{eta})
ight\}\pmoldsymbol{C}_ioldsymbol{\Omega},$$

where C_i is the Cholesky decomposition of A_i such that $C_i C_i^{\top} = A_i$. Therefore we only need to draw N/2 samples using this approach, and by virtue of the negative correlation between the pairs, it leads to a smaller variance in the sample means taken in the approximation than would be obtained from N independent simulations. The choice of N is described below.

119 Initial values

The EM algorithm requires that initial parameters are specified, namely $\hat{\theta}^{(0)}$. By choosing values close to the maximizer, the number of iterations required to reach convergence should be reduced.

For the time-to-event sub-model, a quasi-two-stage model is fitted when the measurement times are balanced, i.e. when $t_{ijk} = t_{ij} \forall k$. That is, we fit *separate* LMMs for each longitudinal outcome as per (1), ignoring the correlation between different outcomes. This is straightforward to implement using standard software, in particular using lme() and coxph() from the R packages nlme [26] and survival [27], respectively. From the fitted models, the best linear unbiased predictions (BLUPs) of the separate model random effects are used to estimate each $W_{1i}^{(k)}(t)$ function. These estimates are then included as time-varying covariates in a Cox regression model, alongside any other fixed effect covariates, which can be straightforwardly fitted using standard software. In the situation that the data are not balanced, i.e. when $t_{ijk} \neq t_{ij} \forall k$, then we fit a standard Cox proportional hazards regression model to estimate γ_v and set $\gamma_{uk} = 0 \forall k$.

For the longitudinal data sub-model, when K > 1 we first find the maximum likelihood estimate of $\{\beta, \operatorname{vech}(D), \sigma_1^2, \ldots, \sigma_K^2\}$ by running a separate EM algorithm for the multivariate linear mixed model. Both the E- and M-step updates are available in closed form, and the initial parameters for this EM algorithm are available from the separate LMM fits, with D initialized as block-diagonal. As these are estimated using an EM rather than MCEM algorithm, we can specify a stricter convergence criterion on the estimates.

¹⁴² Convergence and stopping rules

¹⁴³ Two standard stopping rules for the deterministic EM algorithm used to declare ¹⁴⁴ convergence are the relative and absolute differences, defined as

$$\Delta_{\text{rel}}^{(m+1)} = \max\left\{\frac{|\hat{\boldsymbol{\theta}}^{(m+1)} - \hat{\boldsymbol{\theta}}^{(m)}|}{|\hat{\boldsymbol{\theta}}^{(m)}| + \epsilon_1}\right\} < \epsilon_0, \text{ and}$$
(7)

$$\Delta_{\text{abs}}^{(m+1)} = \max\left\{ \left| \hat{\boldsymbol{\theta}}^{(m+1)} - \hat{\boldsymbol{\theta}}^{(m)} \right| \right\} < \epsilon_2$$
(8)

respectively, for some appropriate choice of ϵ_0 , ϵ_1 , and ϵ_2 , where the maximum is 145 taken over the components of θ . For reference, the R package JM [28] implements 146 (7) (in combination with another rule based on relative change in the likelihood), 147 whereas the R package joineR [29] implements (8). The relative difference might 148 149 be unstable about parameters near zero that are subject to MC error. Therefore, the convergence criterion for each parameter might be chosen separately at each 150 EM iteration based on whether the absolute magnitude is below or above some 151 threshold. A similar approach is adopted in the EM algorithms employed by the 152 software package SAS [30, p. 330]. 153

The choice of N and the monitoring of convergence are conflated when applying a MCEM algorithm, and a dynamic approach is required. As noted by [22], it is computationally inefficient to use a large N in the early phase of the algorithm when the parameter estimates are likely to be far from the maximizer. On the flip side, as the parameter estimates approach the maximizer, the stopping rules will fail as the changes in parameter estimates will be swamped by MC error. Therefore, it has been recommended that one increase N as the estimate moves towards the maximizer. Although this might be done subjectively [31] or by pre-specified rules [32], an automated approach is preferable and necessary for a software implementation. Booth and Hobert [33] proposed an update rule based on a confidence ellipsoid for the maximizer at the (m + 1)-th iteration, calculated using an approximate sandwich estimator for the maximizer, which accounts for the MC error at each iteration. This approach requires additional variance estimation at each iteration, therefore we opt for a simpler approach described by Ripatti *et al.* [34]. Namely, we calculate a coefficient of variation at the (m + 1)-th iteration as

$$\mathrm{cv}(\Delta_{\mathrm{rel}}^{(m+1)}) = \frac{\mathrm{sd}(\Delta_{\mathrm{rel}}^{(m-1)}, \Delta_{\mathrm{rel}}^{(m)}, \Delta_{\mathrm{rel}}^{(m+1)})}{\mathrm{mean}(\Delta_{\mathrm{rel}}^{(m-1)}, \Delta_{\mathrm{rel}}^{(m)}, \Delta_{\mathrm{rel}}^{(m+1)})},$$

where $\Delta_{\text{rel}}^{(m+1)}$ is given by (7), and $\text{sd}(\cdot)$ and $\text{mean}(\cdot)$ are the sample standard de-154 viation and mean functions, respectively. If $\operatorname{cv}(\Delta_{\operatorname{rel}}^{(m+1)}) > \operatorname{cv}(\Delta_{\operatorname{rel}}^{(m)})$, then N :=155 $N + \lfloor N/\delta \rfloor$, for some small positive integer δ . Typically, we run the MCEM algo-156 rithm with a small N (for a fixed number of iterations—a *burn-in*) before imple-157 menting this update rule in order to get into the approximately correct parameter 158 region. Appropriate values for other parameters will be application specific, however 159 we have found $\delta = 3$, N = 100K (for 100K burn-in iterations), $\epsilon_1 = 0.001$, and 160 $\epsilon_0 = \epsilon_2 = 0.005$ delivers reasonably accurate estimates in many cases, where K was 161 earlier defined as the number of longitudinal outcomes. 162

As the EM monotonicity property is lost due to the MC integrations in the MCEM algorithm, convergence might be prematurely declared due to stochasticity if the ϵ -values are too large. To reduce the chance of this occurring, we require that the stopping rule is satisfied for 3 consecutive iterations [33, 34]. However, in any case, trace plots should be inspected to confirm convergence is appropriate.

168 Standard error estimation

Standard error (SE) estimation is usually based on inverting the observed infor-169 mation matrix. When the baseline hazard is unspecified, as is the case here, this 170 presents several challenges. First, $\hat{\lambda}_0(t)$ will generally be a high-dimensional vector, 171 which might lead to numerical difficulties in the inversion of the observed informa-172 tion matrix [6]. Second, the profile likelihood estimates based on the usual observed 173 information matrix approach are known to be underestimated [35]. The reason for 174 this is that the profile estimates are implicit, since the posterior expectations, given 175 by (5), depend on the parameters being estimated, including $\lambda_0(t)$ [6, p. 67]. 176

To overcome these challenges, Hsieh et al. [35] recommended to use bootstrap 177 methods to calculate the SEs. However, this approach is computationally expensive. 178 Moreover, despite the purported theoretical advantages, we also note that recently it 179 has been suggested that bootstrap estimators might actually *overestimate* the SEs; 180 e.g. [36, p. 740] and [35, p. 1041]. At the model development stage, it is often of 181 interest to gauge the strength of association of model covariates, which is not feasible 182 with repeated bootstrap implementations. Hence, an approximate SE estimator is 183 desirable. In either case, the theoretical properties will be contaminated by the 184 addition of MC error from the MCEM algorithm, and it is not yet fully understood 185 what the ramifications of this are. Hence, any standard errors must be interpreted 186 with a degree of caution. We consider two estimators below. 187

1. Bootstrap method. These are estimated by sampling n subjects with re-188 placement and re-labelling the subjects with indices $i' = 1, \ldots, n$. We then re-fit the 189 model to the bootstrap-sampled dataset. It is important to note that we re-sample 190 subjects, not individual data points. This is repeated B-times, for a sufficiently 191 large integer B. Since we already have the MLEs from the fitted model, we can use 192 these as initial values for each bootstrap model fit, thus reducing initial computa-193 tional overheads in calculating approximate initial parameters. For each iteration, 194 we extract the model parameter estimates for $(\boldsymbol{\beta}^{\top}, \operatorname{vech}(\boldsymbol{D}), \sigma_1^2, \ldots, \sigma_K^2, \boldsymbol{\gamma}_v^{\top}, \boldsymbol{\gamma}_y^{\top})$. 195 Note that we do not estimate SEs for $\lambda_0(t)$ using this approach. However, they are 196 generally not of inferential interest. When B is sufficiently large, the SEs can be 197 estimated from the estimated coefficients of the bootstrap samples. Alternatively, 198 $100(1 - \alpha)$ %-confidence intervals can be estimated from the $100\alpha/2$ -th and 199 $100(1 - \alpha/2)$ -th percentiles. 200

2. Empirical information matrix method. Using the Breslow estimator for $\int_0^t \lambda_0(u) du$, the profile score vector for $\boldsymbol{\theta}_{-\lambda} = (\boldsymbol{\beta}^\top, \operatorname{vech}(\boldsymbol{D}), \sigma_1^2, \dots, \sigma_K^2, \boldsymbol{\gamma}^\top)$ is calculated (see Additional file 1). We approximate the profile information for $\boldsymbol{\theta}_{-\lambda}$ by $I_e^{-1/2}(\hat{\boldsymbol{\theta}}_{-\lambda_0})$, where $I_e(\boldsymbol{\theta}_{-\lambda_0})$ is the observed empirical information [25] given by

$$I_e(\boldsymbol{\theta}_{-\lambda}) = \sum_{i=1}^n s_i(\boldsymbol{\theta}_{-\lambda})^{\otimes 2} - \frac{1}{n} S(\boldsymbol{\theta}_{-\lambda})^{\otimes 2},\tag{9}$$

 $s_i(\boldsymbol{\theta}_{-\lambda})$ is the conditional expectation of the complete-data profile score for subject *i*, $S(\boldsymbol{\theta}_{-\lambda})$ is the score defined by $S(\boldsymbol{\theta}_{-\lambda}) = \sum_{i=1}^n s_i(\boldsymbol{\theta}_{-\lambda})$, and $\boldsymbol{a}^{\otimes 2} = \boldsymbol{a}\boldsymbol{a}^\top$ is outer product for a vector \boldsymbol{a} . At the maximizer, $S(\hat{\boldsymbol{\theta}}) = 0$, meaning that the right handside of (9) is zero. Due to the MC error in the MCEM algorithm, this will not be exactly zero, and therefore we include it in the calculations. As per the bootstrap

- ²⁰⁶ approach, SEs for the baseline hazard are again not calculated. We note that this SE
- ²⁰⁷ estimator will be subject to the exact same theoretical limitation of underestimation
- described by Hsieh et al. [35], since the profiling was implicit; that is, because the
- ²⁰⁹ posterior expectations involve the parameters θ .

210 Software

The model described here is implemented in the R package joineRML, which is available on the The Comprehensive R Archive Network (CRAN) (https: //CRAN.R-project.org/package=joineRML). The principal function in joineRML is mjoint(). The primary arguments for implementing mjoint() are summarised in Table 1. To achieve computationally efficiency parts of the MCEM algorithm in joineRML are coded in C++ using the Armadillo linear algebra library and integrated using the R package RcppArmadillo [37].

A model fitted using the mjoint() function returns an object of class mjoint. By 218 default, approximate SE estimates are calculated. If one wishes to use bootstrap 219 standard error estimates, then the user can pass the model object to the bootSE() 220 function. Several generic functions (or rather, S3 methods) can also be applied to 221 mjoint objects, as described in Table 2. These generic functions include common 222 methods, for example coef(), which extracts the model coefficients; ranef(), which 223 extracts the BLUPs (and optional standard errors); and resid(), which extracts 224 the residuals from the linear mixed sub-model. The intention of these functions is to 225 have a common syntax with standard R packages for linear mixed models [26] and 226 survival analysis [27]. Additionally, plotting capabilities are included in joineRML. 227 These include trace plots for assessment of convergence of the MCEM algorithm, 228 and caterpillar plots for subject-specific random effects (Table 2). 229

The package also provides several datasets, and a function simData() that allows for simulation of data from joint models with multiple longitudinal outcomes. joineRML can also fit univariate joint models, however in this case we would currently recommend that the R packages joineR [29], JM [28], or frailtypack [38] are used, which are optimized for the univariate case and exploits Gaussian quadrature. In addition, these packages allow for extensions to more complex cases; for example, competing risks [29, 28] and recurrent events [38].

237 Results

238 Simulation analysis

A simulation study was conducted assuming two longitudinal outcomes and n = 200

²⁴⁰ subjects. Longitudinal data were simulated according to a follow-up schedule of 6

time points (at times $0, 1, \ldots, 5$), with each model including subject-and-outcome-241 specific random-intercepts and random-slopes: $\boldsymbol{b}_i = (b_{0i1}, b_{1i1}, b_{0i2}, b_{1i2})^{\top}$, Correla-242 tion was induced between the 2 outcomes by assuming correlation of -0.5 between 243 the random intercepts for each outcome. Event times were simulated from a Gom-244 pertz distribution with shape $\theta_1 = -3.5$ and scale $\exp(\theta_0) = \exp(0.25) \approx 1.28$, 245 following the methodology described by Austin [39]. Independent censoring times 246 were drawn from an exponential distribution with rate 0.05. Any subject where the 247 event and censoring time exceeded 5 was censored at the truncation time C = 5.1. 248 For all sub-models, we included a pair of covariates $X_i = (x_{i1}, x_{i2})^{\top}$, where x_{i1} is a 249 continuous covariate independently drawn from N(0,1) and x_{i2} is a binary covariate 250 independently drawn from Bin(1, 0.5). The sub-models are given as 251

$$\begin{aligned} y_{ijk} &= (\beta_{0,k} + b_{i0k}) + (\beta_{1,k} + b_{i1k})t_j + \beta_{2,k}x_{i1} + \beta_{3,k}x_{i2} + \varepsilon_{ijk}, \text{ for } k = 1,2; \\ \lambda_i(t) &= \exp\left\{(\theta_0 + \theta_1 t) + \gamma_{v1}x_{i1} + \gamma_{v2}x_{i2} + \gamma_{y1}(b_{i01} + b_{i11}t) + \gamma_{y2}(b_{i02} + b_{i12}t)\right\}; \\ \mathbf{b}_i &\sim N_4(0,D); \\ \varepsilon_{ijk} &\sim N(0,\sigma_k^2), \end{aligned}$$

where D is specified unstructured (4×4) -covariance matrix with 10 unique param-252 eters. Simulating datasets is straightforward using the joineRML package by means 253 of the simData() function. The true parameter values and results from 500 simu-254 lations are shown in Table 3. In particular, we display the mean estimate, the bias, 255 the empirical SE (= the standard deviation of the the parameter estimates); the 256 mean SE (= the mean SE of each parameter calculated for each fitted model); the 257 mean square error (MSE), and the coverage. The results confirm that the model 258 fitting algorithm generally performs well. 259

A second simulation analysis was conducted using the parameters above (with 260 n = 100 subjects per dataset). However, in this case we used a heavier-tailed distri-261 bution for the random effects: a multivariate t_5 distribution [40]. The bias for the 262 fixed effect coefficients was comparable to the multivariate normal random effects 263 simulation study (above). The empirical standard error was consistently smaller 264 than the mean standard error, resulting in coverage between 95% and 99% for the 265 coefficient parameters. Rizopoulos et al. [41] noted that the misspecification of the 266 random effects distributions was minimised as the number of longitudinal measure-267 ments per subject increased, but that the standard errors are generally affected. 268 These findings are broadly in agreement with the simulation study conducted here, 269

and other studies [42, 43]. Choi *et al.* [44] provide a review of existing research on the misspecification of random effects in joint modelling.

272 Example

We consider the primary biliary cirrhosis (PBC) data collected at the Mayo Clinic 273 between 1974 to 1984 [45]. This dataset has been widely analyzed using joint mod-274 elling methods [46, 47, 18]. PBC is a long-term liver disease in which the bile ducts 275 in the liver become damaged. Progressively, this leads to a build-up of bile in the 276 liver, which can damage it and eventually lead to cirrhosis. If PBC is not treated 277 or reaches an advanced stage, it can lead to several major complications, including 278 mortality. In this study, 312 patients were randomised to receive D-penicillamine 279 (n = 158) or placebo (n = 154). In this example we analyse the subset of patients 280 randomized to placebo. 281

Patients with PBC typically have abnormalities in several blood tests; hence, 282 during follow-up several biomarkers associated with liver function were serially 283 recorded for these patients. We consider three biomarkers: serum bilirunbin (de-284 noted serBilir in the model and data; measured in units of mg/dl), serum albumin 285 (albumin; mg/dl), and prothrombin time (prothrombin; seconds). Patients had a 286 mean 6.3 (SD = 3.7) visits (including baseline). The data can be accessed from the 287 joineRML package via the command data(pbc2). Profile plots for each biomarker 288 are shown in Figure 1, indicating distinct differences in trajectories between the 289 those who died during follow-up and those who did not (right-censored cases). A 290 Kaplan-Meier curve for overall survival is shown in Figure 2. There were a total of 291 69 (44.8%) deaths during follow-up in the placebo subset. 292

We fit a relatively simple joint model for the purposes of demonstration, which encompasses the following trivariate longitudinal data sub-model:

$$\begin{split} \log(\texttt{serBilir}) &= (\beta_{0,1} + b_{0i,1}) + (\beta_{1,1} + b_{1i,1})\texttt{year} + \varepsilon_{ij1}, \\ \texttt{albumin} &= (\beta_{0,2} + b_{0i,2}) + (\beta_{1,2} + b_{1i,2})\texttt{year} + \varepsilon_{ij2}, \\ (0.1 \times \texttt{prothrombin})^{-4} &= (\beta_{0,3} + b_{0i,3}) + (\beta_{1,3} + b_{1i,3})\texttt{year} + \varepsilon_{ij3}, \\ \textbf{b}_i &\sim N_6(0, \textbf{D}), \text{ and } \varepsilon_{ijk} \sim N(0, \sigma_k^2) \text{ for } k = 1, 2, 3; \end{split}$$

²⁹⁵ and a time-to-event sub-model for the study endpoint of death:

$$\begin{aligned} \lambda_i(t) &= \lambda_0(t) \exp\left\{\gamma_v \mathsf{age} + W_{2i}(t)\right\}, \\ W_{2i}(t) &= \gamma_{\mathsf{bil}}(b_{0i,1} + b_{1i,1}t) + \gamma_{\mathtt{alb}}(b_{0i,2} + b_{1i,2}t) + \gamma_{\mathtt{pro}}(b_{0i,3} + b_{1i,3}t) \end{aligned}$$

The log transformation of bilirubin is standard, and confirmed reasonable based 296 on inspection of Q-Q plots for residuals from a separate fitted linear mixed model 297 fitted using the lme() function from the R package nlme. Albumin did not require 298 transformation. Residuals were grossly non-normal for prothrombin time using both 299 untransformed and log-transformed outcomes. Therefore, a Box-Cox transformation 300 was applied, which suggested an inverse-quartic transform might be suitable, which 301 was confirmed by inspection of a Q-Q plot. The pairwise correlations for baseline 302 measurements between the three transformed markers were 0.19 (prothrombin time 303 vs. albumin), -0.30 (bilirubin vs. prothrombin time and albumin) The model is fit 304 using the joineRML R package (version 0.2.0) using the following code. 305

```
306
307
   # Get data
308
   data(pbc2)
309
    placebo <- subset(pbc2, drug == "placebo")</pre>
310
311
    # Fit model
312
    fit.pbc <- mjoint(</pre>
313
        formLongFixed = list(
314
            "bil" = log(serBilir) ~ year,
315
            "alb" = albumin ~ year,
316
            "pro" = (0.1 * \text{ prothrombin})^{-4} year),
317
        formLongRandom = list(
318
            "bil" = ~ year | id,
319
            "alb" = ~ year | id,
320
            "pro" = ~ year | id),
321
        formSurv = Surv(years, status2) ~ age,
322
        data = placebo,
323
        timeVar = "year",
324
        control = list(tol0 = 0.001, burnin = 400)
325
    )
326
327
```

Here, we have specified a more stringent tolerance value for ϵ_0 than the default setting in mjoint(). Additionally, the burn-in phase was increased to 400 iterations after inspection of convergence trace plots. The model fits in 3.1 minutes on a MacBook Air 1.6GHz Intel Core i5 with 8GB or RAM running R version 3.3.0, having completed 423 MCEM iterations (not including the EM algorithm iterations performed for determining the initial values of the separate multivariate linear mixed sub-model) with a final MC size of M = 3528. The fitted model results are shown in Table 4.

The fitted model indicated that an increase in the subject-specific random de-336 viation from the population trajectory of serum bilirubin was significantly associ-337 ated with increased hazard of death. A significant association was also detected for 338 subject-specific decreases in albumin from the population mean trajectory. However, 339 prothrombin time was not significantly associated with hazard of death, although 340 its direction is clinically consistent with PBC disease. Albert and Shih [46] anal-341 ysed the first 4-years follow-up from this dataset with the same 3 biomarkers and a 342 discrete event time distribution using a regression calibration model. Their results 343 were broadly consistent, although the effect of prothrombin time on the event time 344 sub-model was strongly significant. 345

We also fitted 3 univariate joint models to each of the biomarkers and the event 346 time sub-model using the R package joineR (version 1.2.0) owing to its optimization 347 for such models. The LMM parameter estimates were similar, although the absolute 348 magnitude of the slopes was smaller for the separate univariate models. Since 3 349 separate models were fitted, 3 estimates of γ_v were estimated, with the average 350 comparable to the multivariate model estimate. The multivariate model estimates 351 of $\gamma_{u} = (\gamma_{\text{bil}}, \gamma_{\text{alb}}, \gamma_{\text{pro}})^{\top}$ were substantially attenuated relative to the separate 352 model estimates, although the directions remained consistent. It is also interesting 353 to note that γ_{pro} was statistically significant in the univariate model. However, the 354 univariate models are not accounting for the correlation between different outcomes, 355 whereas the multivariate joint model does. 356

The model was refitted with the one-step Newton-Raphson update for γ replaced by a Gauss-Newton-like update in a time of 2.2 minutes for 419 MCEM iterations with a final MC size of M = 6272. This is easily achieved by running the following code.

- 361
- 362

363 364 fit.pbc.gn <- update(fit.pbc, gammaOpt = "GN")</pre>

In addition, we bootstrapped this model with B = 100 samples to estimate SEs and contrast them with the approximate estimates based on the inverse empirical profile information matrix. In practice, one should choose B > 100, particularly if using bootstrap percentile confidence intervals; however, we used a small value to reduce the computational burden on this process. In a similar spirit, we relaxed the

```
sro convergence criteria and lowered reduced the number of burn-in iterations. This is
easily implemented by running the following code, taking 1.8 hours to fit.
fit.pbc.gn.boot <- bootSE(fit.pbc.gn, nboot = 100, control = list(
    tol0 = 0.005, tol2 = 0.01, convCrit = "sas",
    burnin = 300, mcmaxIter = 350))</pre>
```

It was observed that the choice of gradient matrix in the γ -update led to virtually indistinguishable parameter estimates, although we note the same random seed was used in both cases. The bootstrap estimated SEs were broadly consistent with the approximate SEs, with no consistent pattern in underestimation observed.

382 Discussion

Multivariate joint models introduce three types of correlations: (1) within-subject 383 serial correlation for repeated measures; (2) between longitudinal outcomes corre-384 lation; and (3) correlation between the multivariate LMM and time-to-event sub-385 models. It is important to account for all of these types of correlations; however, 386 some authors have reported collapsing their multivariate data to permit univariate 387 joint models to be fitted. For example, Battes et al. [7] used an ad hoc approach 388 of either summing or multiplying the three repeated continuous measures (stan-380 dardized according to clinical upper reference limits of the biomarker assays), and 390 then applying standard univariate joint models. Wang et al. [48] fitted separate uni-391 variate joint models to each longitudinal outcome in turn. Neither approach takes 392 complete advantage of the correlation between the multiple longitudinal measures 393 and the time-to-event outcome. 394

Here, we described a new R package joineRML that can fit the models described 395 in this paper. This was demonstrated on a real-world dataset. Although in the fitted 396 model we assumed linear trajectories for the biomarkers, splines could be straight-397 forwardly employed, as have been used in other multivariate joint model applications 398 [15], albeit at the cost of additional computational time. Despite a growing availabil-399 ity of software for univariate joint models, Hickey et al. [19] noted that there were 400 very few options for fitting joint models involving multivariate longitudinal data. 401 To the best of our knowledge, options are limited to the R packages JMbayes [49], 402 rstanarm [50], and the Stata package stim [47]. Whilst all of these packages are 403 available, the extension to multivariate data remain features in the developmental 404 versions only. Moreover, none of these incorporates an unspecified baseline hazard. 405

 $_{406}$ $\,$ The first two packages use Markov chain Monte Carlo (MCMC) methods to fit the

⁴⁰⁷ joint models. Bayesian models are potentially very useful for fitting joint models,

and in particular for dynamic prediction; however, MCMC is also computationally

- demanding, especially in the case of multivariate models. Several other publications
- ⁴¹⁰ have made **BUGS** code available for use with WinBUGS and OpenBUGS (e.g. [51]),

⁴¹¹ but these are not easily modifiable and post-fit computations are cumbersome.

joineRML is a new software package developed to fill a void in the joint modelling 412 field, but is still in its infancy relative to highly developed univariate joint model 413 packages such as the R package JM [28] and Stata package st jm [47]. Future devel-414 opments of joineRML intend to cover several deficiencies. First, joineRML currently 415 only permits an association structure of the form $W_{2i}(t) = \sum_{k=1}^{K} \gamma_{yk} W_{1i}^{(k)}(t)$. As has 416 been demonstrated by others, the association might take different forms, including 417 random-slopes and cumulative effects or some combination of multiple structures, 418 and these may also be different for separate longitudinal outcomes [18]. Moreover, 419 it is conceivable that separate longitudinal outcomes may interact in the hazard 420 sub-model. Second, the use of MC integration provides a scalable solution to the 421 issue of increasing dimensionality in the random effects. However, for simpler cases, 422 e.g. bivariate models with random-intercepts and random-slopes (total of 4 random 423 effects), Gaussian quadrature might be computationally superior; this trade-off re-424 quires further investigation. Third, joineRML can currently only model a single 425 event time. However, there is a growing interest in competing risks [9] and recur-426 rent events data [11], which if incorporated into joineRML, would provide a flexible 427 all-round multivariate joint modelling platform. Competing risks [29, 28] and re-428 current events [38] have been incorporated into R packages already, but are limited 429 to the case of a solitary longitudinal outcome. Of note, the PBC trial dataset anal-430 ysed in this study includes times to the competing risk of kidney transplantation. 431 Fourth, with ever-increasing volumes of data collected during routine clinical vis-432 its, the need for software to fit joint models with very many longitudinal outcomes 433 is foreseeable [52]. This would likely require the use of approximate methods for 434 the numerical integration or data reduction methods. Fifth, additional residual di-435 agnostics are necessary for assessing possible violations of model assumptions. The 436 joineRML package has a resid() function for extracting the longitudinal sub-model 437 residuals; however, these are complex for diagnostic purposes due to the informative 438 dropout, hence the development of multiple-imputation based residuals [53]. 439

440 Conclusions

- ⁴⁴¹ In this paper we have presented an extension of the classical joint model proposed
- $_{442}$ by Henderson *et al.* [3] and an estimation procedure for fitting the models that
- $_{443}$ builds on the foundations laid by Lin *et al.* [20]. In addition, we described a new R
- ⁴⁴⁴ package joineRML that can fit the models described in this paper, which leverages
- the MCEM algorithm which should scale well for increasing number of longitudinal
- outcomes. This software is timely, as it has previously been highlighted that there
- 447 is a paucity of software available to fit such models [19]. The software is being
- ⁴⁴⁸ regularly updated and improved.

449 Availability and requirements

- 450 Project name: joineRML
- 451 Project home page: https://github.com/graemeleehickey/joineRML/
- 452 Operating system(s): platform independent
- 453 Programming language: R
- 454 Other requirements: none
- 455 License: GNU GPL-3
- 456 Any restrictions to use by non-academics: none

457 Abbreviations

- 458 MCEM Monte Carlo expectation maximisation; EM expectation maximisation; MC Monte Carlo; LMM -
- 459 linear mixed models; BLUP best linear unbiased prediction; SE standard error; MLE maximum likelihood
- 460 estimate; CRAN The Comprehensive R Archive Network; PBC primary biliary cirrhosis; SD standard deviation
- 461 Declarations
- 462 Ethics approval and consent to participate
- 463 Not applicable.
- 464 Consent for publication
- 465 Not applicable.

466 Availability of data and materials

- 467 The R package joineRML can be installed directly using install.packages("joineRML") in an R console. The
- 468 source code is available at https://github.com/graemeleehickey/joineRML. Archived versions are available from
- 469 the Comprehensive R Archive Network (CRAN) at https://cran.r-project.org/web/packages/joineRML/.
- 470 joineRML is platform independent, requiring R version \geq 3.3.0, and is published under a GNU GPL-3 license. The
- 471 dataset analysed during the current study is bundled with the R package joineRML, and can be accessed by running
- 472 the command data(pbc2, package = "joineRML").

473 Competing interests

474 The authors declare that they have no competing interests.

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- 478 manuscript.

479 Author's contributions

- 480 All authors collaborated in developing the model fitting algorithm reported. The programming and running of the
- 481 analysis was carried out by GLH. GLH wrote the first draft of the manuscript, with revisions provided by PP, AJ,
- 482 and RKD. All authors contributed to the manuscript revisions. All authors read and approved the final manuscript.

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607 Additional Files

- 608 Additional file 1
- An appendix (appendix.pdf) is available that includes details on the score vector and M-step estimators.
- 610 Figures

Figure 1 Longitudinal trajectory plots. The black lines show individual subject trajectories, and the coloured lines show smoothed (LOESS) curves stratified by whether the patient experienced the endpoint (blue) or not (red).

Figure 2 Kaplan-Meier curve for overall survival. A pointwise 95% band is shown (dashed lines). In total, 69 patients (of 154) died during follow-up.

611 Tables

 $\label{eq:table1} \begin{array}{l} \textbf{Table 1} \mbox{ The primary arguments}^{\dagger} \mbox{ with descriptions for the mjoint() function in the R package joineRML. \end{array}$

Argument	Description
formLongFixed	a list of formulae for the fixed effects component of each longitudinal outcome. The left hand-hand side defines the response, and the right-hand side specifies the fixed effect terms.
formLongRandom	a list of one-sided formulae specifying the model for the random effects effects of each longitudinal outcome.
formSurv	a formula specifying the proportional hazards regression model (not including the latent association structure).
data	a list of data.frame objects for each longitudinal outcome in which to interpret the variables named in the formLongFixed and formLongRandom. The list structure enables one to include multiple longitudinal outcomes with different measurement protocols. If the multiple longitudinal outcomes are measured at the same time points for each patient (i.e. $t_{ijk} = t_{ij} \forall k$), then a single data.frame object can be given instead of a list. It is assumed that each data frame is in <i>long format</i> .
survData	(optional) a data.frame in which to interpret the variables named in the formSurv. If survData is not given, then mjoint() looks for the time-to-event data in data.
timeVar	a character string indicating the time variable in the linear mixed effects model.
inits	(optional) a list of initial values for some or all of the parameters estimated in the model.
control	(optional) a list of control parameters. These allow for the control of $\epsilon_0, \epsilon_1,$ and ϵ_2 in (7) and (8); the choice of N, δ , and convergence criteria; the maximum number of MCEM iterations, and the minimum number of MCEM iterations during burn-in. Additionally, the control argument gammaOpt can be used to specify whether a one-step Newton-Raphson (="NR") or Gauss-Newton-like (="GN") update should be used for the M-step update of γ .

[†]mjoint() also takes the optional additional arguments verbose, which if TRUE allows for monitoring updates at each MCEM algorithm iteration, and pfs, which if FALSE can force the function not to calculate post-fit statistics such as the BLUPs and associated standard errors of the random effects and approximate standard errors of the model parameters. In general, these arguments are not required.

Function(s)	Returns
logLik, AIC, BIC	the log-likelihood, Akaike information criterion and Bayesian information
	criterion statistics, respectively.
coef, fixef	the fixed effects parameter estimates.
ranef	the BLUPs (and optional standard errors).
$print^{\dagger}$, summary*	short and long model summary outputs, respectively.
fitted, resid	the fitted and raw residuals from the multivariate LMM sub-model, re-
	spectively.
plot [‡]	the MCEM algorithm convergence trace plots.
sigma	the residual standard errors from the LMM sub-model.
vcov	the variance-covariance matrix of the main parameters of the fitted
	model (except the baseline hazard).
getVarCov	the random effects variance-covariance matrix.
confint	the confidence intervals based on asymptotic normality.
update	specific parts of a fitted model can be updated, e.g. by adding or re-
	moving terms from a sub-model, and then re-fitted.
sampleData	sample data (with or without replacement) from a joint model.

Table 2 Additional functions with descriptions that can be applied to objects of class \mathtt{mjoint}^\dagger .

[†]print() also applies to objects of class summary.mjoint and bootSE inheriting from the summary() and bootSE() functions, respectively. [‡]plot() also accepts objects of class ranef.mjoint inheriting from the ranef() function, which displays a caterpillar plot (with 95% prediction intervals) for each random effect. *summary() can also take the optional argument of an object of class bootSE inheriting from the function bootSE(), which overrides the approximate SEs and Cls with those from a bootstrap estimation routine.

Parameter	True value	Mean estimated value	Empirical SE	Mean SE	Bias	MSE	Coverage
D ₁₁	0.2500	0.2411	0.0435	_	-0.0089	0.0020	_
D_{21}	0.0000	0.0010	0.0136		0.0010	0.0002	_
D_{31}	-0.1250	-0.1212	0.0295		0.0038	0.0009	_
D_{41}	0.0000	-0.0006	0.0127	_	-0.0006	0.0002	_
D_{22}	0.0400	0.0396	0.0072		-0.0004	0.0001	—
D_{32}	0.0000	-0.0002	0.0138		-0.0002	0.0002	—
D_{42}	0.0000	-0.0001	0.0055		-0.0001	0.0000	—
D_{33}	0.2500	0.2420	0.0400		-0.0080	0.0017	—
D_{43}	0.0000	0.0007	0.0134		0.0007	0.0002	—
D_{44}	0.0400	0.0399	0.0075		-0.0001	0.0001	—
$\beta_{0,1}$	0.0000	0.0028	0.0612	0.0660	0.0028	0.0038	0.9660
$\beta_{1,1}$	1.0000	1.0012	0.0218	0.0229	0.0012	0.0005	0.9500
$\beta_{2,1}$	1.0000	1.0010	0.0449	0.0470	0.0010	0.0020	0.9540
$\beta_{3,1}$	1.0000	0.9932	0.0897	0.0925	-0.0068	0.0081	0.9440
σ_1^2	0.2500	0.2506	0.0165	0.0171	0.0006	0.0003	0.9560
$\beta_{0,2}$	0.0000	-0.0026	0.0637	0.0655	-0.0026	0.0041	0.9660
$\beta_{1,2}$	-1.0000	-1.0011	0.0229	0.0223	-0.0011	0.0005	0.9480
$\beta_{2,2}$	0.0000	0.0008	0.0399	0.0472	0.0008	0.0016	0.9700
$\beta_{3,2}$	0.5000	0.5061	0.0894	0.0923	0.0061	0.0080	0.9540
σ_2^2	0.2500	0.2501	0.0162	0.0171	0.0001	0.0003	0.9540
γ_{v1}	0.0000	0.0011	0.1243	0.1392	0.0011	0.0155	0.9720
γ_{v2}	1.0000	1.0487	0.2837	0.2750	0.0487	0.0829	0.9340
γ_{y1}	-0.5000	-0.5121	0.1936	0.2084	-0.0121	0.0376	0.9560
γ_{y2}	1.0000	1.0311	0.2220	0.2145	0.0311	0.0502	0.9400

Table 3 Results of simulation study.

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converge using joineRML within the maximum number of MC iterations, and so SEs and Cls are based on 99 bootstrap samples only. ^aSeparate model fit for serBilir. ^bSeparate model fit for serBilir. ^bSeparate model fit for serBilir. ^bSeparate model fit for albumin. ^cSeparate model fit for prothrombin. calculated from the maximum likelihood one model failed to