

AN ALGORITHM FOR FITTING MIXTURES OF GOMPERTZ  
DISTRIBUTIONS TO CENSORED SURVIVAL DATA

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## PURPOSE AND DESCRIPTION

We consider the fitting of a mixture of two Gompertz distributions to censored survival data. This model is therefore applicable where there are two distinct causes for failure that act in a mutually exclusive manner, and the baseline failure time for each cause follows a Gompertz distribution. For example, in a study of a disease such as breast cancer, suppose that failure corresponds to death, whose cause is attributed either to breast cancer or some other cause. In this example, the mixing proportion for the component of the mixture representing time to death from a cause other than breast cancer may be interpreted to be the cure rate for breast cancer (Gordon, 1990a and 1990b). This Gompertz mixture model whose components are adjusted multiplicatively to reflect the age of the patient at the origin of the survival time, is fitted by maximum likelihood via the EM algorithm (Dempster, Laird, and Rubin, 1977). There is the provision to handle the case where the mixing proportions are formulated in terms of a logistic model to depend on a vector of covariates associated with each survival time. The algorithm can also handle the case where there is only one cause of failure, but which may happen at infinity for some patients with a nonzero probability (Farewell, 1982).

## METHOD

Let  $T$  be a random variable denoting the failure time for a patient with an associated  $p$ -dimensional covariate vector  $\mathbf{x}$  which contains at least the single covariate  $w$ . The density of  $T$  is postulated to be the two-component mixture

$$f(t; \mathbf{x}, \Psi) = \sum_{i=1}^2 \pi_i(\mathbf{x}; \boldsymbol{\alpha}) f_i(t; w, \boldsymbol{\theta}_i), \quad (1)$$

where

$$f_i(t; w, \boldsymbol{\theta}_i) = h_i(t; w, \boldsymbol{\theta}_i) \exp\{-\Lambda_i(t; w, \boldsymbol{\theta}_i)\}, \quad (2)$$

$$h_i(t; w, \boldsymbol{\theta}_i) = \exp(\gamma_i w + \lambda_i + \beta_i t), \quad (3)$$

$$\Lambda_i(t; w, \boldsymbol{\theta}_i) = e^{\gamma_i w} e^{\lambda_i} (e^{\beta_i t} - 1)/\beta_i, \quad (4)$$

and

$$\boldsymbol{\theta}_i = (\lambda_i, \beta_i, \gamma_i)'$$

for  $i = 1, 2$ , and where

$$\Psi = (\boldsymbol{\alpha}', \boldsymbol{\theta}'_1, \boldsymbol{\theta}'_2)'.$$

Here a prime denotes vector transpose.

The mixing proportion  $\pi_i(\mathbf{x}; \boldsymbol{\alpha})$  is modelled to depend on  $\mathbf{x}$  through the logistic model for which

$$\begin{aligned} \pi_1(\mathbf{x}; \boldsymbol{\alpha}) &= 1 - \pi_2(\mathbf{x}; \boldsymbol{\alpha}) \\ &= e^{a+\mathbf{b}'\mathbf{x}} / (1 + e^{a+\mathbf{b}'\mathbf{x}}), \end{aligned}$$

where  $\boldsymbol{\alpha} = (a, \mathbf{b}')'$  is the parameter vector.

It can be seen from (3) that the hazard function corresponding to the  $i$ th component of the mixture has the form

$$h_i(t; w, \boldsymbol{\theta}_i) = e^{\gamma_i w} h_{i,0}(t; \boldsymbol{\theta}_i),$$

where the baseline hazard function  $h_{i,0}(t; \boldsymbol{\theta}_i)$  is represented by the Gompertz model, for which

$$h_{i,0}(t; \boldsymbol{\theta}_i) = \exp(\lambda_i + \beta_i t) \quad (i = 1, 2).$$

Hence the covariate  $w$  is modelled to have an additive effect on the  $i$ th component hazard function on the log scale. Often in practice,  $w$  will be the age of the patient at the instance from which the time to failure is being measured.

For the  $j$ th entity ( $j = 1, \dots, n$ ), we observe

$$\mathbf{y}_j = (t_j, \mathbf{x}'_j, \delta_{1j}, \delta_{2j}, \delta_{3j})',$$

where  $\delta_{1j}$ ,  $\delta_{2j}$  and  $\delta_{3j}$  are zero-one indicator variables with  $\delta_{ij} = 1$  if entity  $j$  failed due to cause  $i$  and zero otherwise ( $i = 1, 2$ ), and  $\delta_{3j} = 1$  if entity  $j$  was censored at time  $t_j$ ;  $\mathbf{x}_j$  is the covariate vector recorded on the  $j$ th entity ( $j = 1, \dots, n$ ). Thus

$$\delta_{1j} + \delta_{2j} + \delta_{3j} = 1$$

for  $j = 1, \dots, n$ . As above, we let  $\boldsymbol{\Psi} = (\boldsymbol{\alpha}', \boldsymbol{\theta}'_1, \boldsymbol{\theta}'_2)'$  be the vector containing all the unknown parameters. Then the log likelihood for  $\boldsymbol{\Psi}$  formed on the basis of  $\mathbf{y}_1, \dots, \mathbf{y}_n$  is given by

$$\begin{aligned} \log L(\boldsymbol{\Psi}) &= \sum_{j=1}^n \left[ \sum_{i=1}^2 \delta_{ij} \log \{ \pi_i(\mathbf{x}_j; \boldsymbol{\alpha}) f_i(t_j; w_j, \boldsymbol{\theta}_i) \} \right. \\ &\quad \left. + \delta_{3j} \log \sum_{i=1}^2 \{ \pi_i(\mathbf{x}_j; \boldsymbol{\alpha}) S_i(t_j; w_j, \boldsymbol{\theta}_i) \} \right], \end{aligned}$$

where

$$S_i(t; w, \boldsymbol{\theta}_i) = \exp\{-\Lambda_i(t; w, \boldsymbol{\theta}_i)\}$$

is the  $i$ th component survival function.

The parameter vector  $\boldsymbol{\Psi}$  is estimated by maximum likelihood. The likelihood equation

$$\partial \log L(\boldsymbol{\Psi}) / \partial \boldsymbol{\Psi} = \mathbf{0}$$

is solved via the EM algorithm of Dempster et al. (1977). For those entities with censored observations (that is,  $\delta_{3j} = 1$ ), we introduce the zero-one indicator vector,  $\mathbf{z}_j = (z_{1j}, z_{2j})'$ , where  $z_{ij} = 1$  or 0 according as entity  $j$  would have failed from cause  $i$  or not ( $i = 1, 2$ ). The EM algorithm is applied with the  $\mathbf{y}_j$  and the  $\mathbf{z}_j$  viewed as the complete-data. The actual time to failure for those entities with  $\delta_{3j} = 1$  was not

introduced as an incomplete variable in the complete-data framework, as it did not simplify the calculations.

It follows on application of the EM algorithm in the aforementioned framework that on the  $(k + 1)$ th iteration, the estimate of  $\Psi$ ,  $\Psi^{(k+1)}$ , is the maximizer of the function

$$Q(\Psi; \Psi^{(k)}) = \sum_{j=1}^n \sum_{i=1}^2 [\delta_{ij} \log\{\pi_i(\mathbf{x}_j; \boldsymbol{\alpha}) f_i(t_j; w_j, \boldsymbol{\theta}_i)\} \\ + \delta_{3j} \tau_i(t_j; \mathbf{x}_j, \Psi^{(k)}) \log\{\pi_i(\mathbf{x}_j; \boldsymbol{\alpha}) S_i(t_j; w_j, \boldsymbol{\theta}_i)\}], \quad (5)$$

where  $\Psi^{(k)}$  is the estimate of  $\Psi$  on the previous  $k$ th iteration and

$$\tau_i(t_j; \mathbf{x}_j, \Psi) = \pi_i(\mathbf{x}_j; \boldsymbol{\alpha}) S_i(t_j; w_j, \boldsymbol{\theta}_i) / \sum_{h=1}^2 \pi_h(\mathbf{x}_j; \boldsymbol{\alpha}) S_h(t_j; w_j, \boldsymbol{\theta}_h)$$

is the posterior probability that the  $j$ th entity with censored survival time  $t_j$  would have failed due to cause  $i$  ( $i = 1, 2$ ).

Unfortunately, for Gompertz component distributions,  $\Psi^{(k+1)}$  does not exist in closed form, as discussed in McLachlan and Krishnan (1997). In our algorithm, it is computed iteratively, commencing with  $\boldsymbol{\alpha}^{(k+1)}$ , since its calculation does not involve  $\boldsymbol{\theta}_1^{(k+1)}$  and  $\boldsymbol{\theta}_2^{(k+1)}$ . The estimate  $\boldsymbol{\alpha}^{(k+1)}$  is computed iteratively by Newton-Raphson. The computation of  $\boldsymbol{\theta}_1^{(k+1)}$  and  $\boldsymbol{\theta}_2^{(k+1)}$  is then undertaken iteratively using a quasi-Newton method. As noted by Dempster et al. (1977), it is not essential that  $\Psi^{(k+1)}$  globally maximizes  $Q(\Psi; \Psi^{(k)})$  for the monotonicity property of the sequence of iterates  $\{\Psi^{(k)}\}$  to hold. A sufficient condition is

$$Q(\Psi^{(k+1)}, \Psi^{(k)}) \geq Q(\Psi^{(k)}, \Psi^{(k)}). \quad (6)$$

The use of (6) as a means of choosing  $\Psi^{(k+1)}$  corresponds to using the generalized EM algorithm (Dempster et al. 1977).

In order to improve the speed of convergence, an E-step is performed in our program after the computation of  $\boldsymbol{\alpha}^{(k+1)}$  and before the computation of the other sub-vectors  $\boldsymbol{\theta}_1^{(k+1)}$  and  $\boldsymbol{\theta}_2^{(k+1)}$  in  $\Psi^{(k+1)}$ . In the terminology of Meng and Rubin (1993), it can be viewed as applying a multicycle ECM (expectation conditional-maximization) algorithm, where a cycle is defined to be an E-step followed by a CM-step. This multicycle E-step is effected here by updating  $\boldsymbol{\alpha}^{(k)}$  by  $\boldsymbol{\alpha}^{(k+1)}$  in  $\Psi^{(k)}$  in the right-hand side of (5). A multicycle ECM algorithm is not necessarily a GEM algorithm. However, it does increase the likelihood function at each cycle, and hence at each iteration (Meng and Rubin, 1993). To further enhance convergence in our program, another E-step is performed after the computation of  $\boldsymbol{\alpha}^{(k+1,m+1)}$  for each  $m$ . This may affect the monotone convergence of the consequent sequence of likelihood values. In our particular applications the monotone convergence was preserved.

The standard errors of the maximum likelihood estimator  $\hat{\Psi}$  of  $\Psi$  is assessed using the bootstrap methodology of Efron (1979, 1982). A number  $K$  (as specified by

the user) of independent bootstrap samples are obtained with each being randomly drawn with replacement from the observed data  $\mathbf{y}_1, \dots, \mathbf{y}_n$ .

The program can also handle the case where there is only one cause of failure, but which may not happen for some entities; that is, there is a nonzero probability that the time to failure is infinity. In this case,  $f_1(t_j; w_j, \boldsymbol{\theta}_1)$  and  $S_1(t_j; w_j, \boldsymbol{\theta}_1)$  are each set identically equal to 1. The mixing proportion  $\pi_1(\mathbf{x}_j; \boldsymbol{\alpha})$  now represents the prior probability that an entity with covariate vector  $\mathbf{x}_j$  will not fail (that is, will have an infinite failure time).

## ABOUT THE SOFTWARE

The software is written in FORTRAN77. To compile the algorithms, several steps are required to be carried out which are listed in the file “makefile.txt”. The main algorithm is named MGOMP in which the subroutine CHOL (algorithm AS6) and SYMINV (algorithm AS7) are called to invert a positive definite symmetric matrix. To generate random numbers for the bootstrap application, the subroutine RANDOM (algorithm AS183) is called to return a pseudo-random integer taken from a uniform distribution between 0 and 1, excluding the end points. In addition, the MINPACK routine HYBRD1 (Moré et al., 1980) is called to find a solution of a system of two non-linear equations. Thus, there are five FORTRAN files named MGOMP.f, CHOL.f, SYMINV.f, RANDOM.f, and HYBRID.f in the package.

To run the software, two input files are required. The first file “para.dat” provides the initial estimates for the parameters  $\lambda_i, \beta_i, \gamma_i$  ( $i = 1, 2$ ),  $a$  and the coefficient  $b_1$  of  $w$ . The other elements in the coefficient  $\mathbf{b}$  of  $\mathbf{x}$  are set initially to zero. The second file “surv.dat” contains the data  $\mathbf{y}_j = (t_j, w_j, \delta_{1j}, \delta_{2j}, \mathbf{x}'_j)'$  of each observation, where  $t_j$ ,  $w_j$ ,  $\mathbf{x}_j$ , and  $\delta_{ij}$  ( $i = 1, 2$ ) are defined as in the previous section.

For the sake of achieving satisfactory accuracy on the parameter estimates, double-precision is used in the algorithm. The stopping criterion for convergence is based on the absolute relative changes in the parameters, and a value of  $10^{-4}$  is used for the cumulative (absolute) relative change. The final estimates of the parameters are given in the output file “fort.25”.

The structure of the subroutine GETEST is given below. It defines the input and output variables in the algorithms.

SUBROUTINE GETEST(NBOOT,MODEL,PI2EQ1,SIZE,FTIME,AGE,DELTA,  
NUMCOV,COV,ESTIN,ESTOUT,LLOUT,COEOUT,ITER,IFAIL,CODE)

NBOOT	Integer	input:	bootstrap iterations indicator: = 1 indicate the complete data is used; ≠ 1 indicate the number of bootstrap iterations needed.
MODEL	Integer	input:	model indicator: = 1 if model with two distinct causes of failure; = 2 if only one cause of failure is considered.
PI2EQ1	Integer	input:	mixing proportion indicator: = 0 if fitting two components of Gompertz distributions; = 1 if fitting single Gompertz distribution ( $\pi_{2j} = 1 \forall j$ ).
SIZE	Integer	input:	the number of entities in the data set.
FTIME	Double precision array (SIZE)	input:	the failure (or censoring) time of each of the entities.
AGE	Double precision array (SIZE)	input:	the age of each of the entities at the time of operation, that is $w_j$
DELTA	Integer array (3,SIZE)	input:	zero-one indicator: $d(1,j) = 1$ if entity $j$ failed due to Cause 1; $d(2,j) = 1$ if entity $j$ failed due to Cause 2; $d(3,j) = 1$ if entity $j$ was censored.
NUMCOV	Integer	input:	the number of additional covariates besides AGE to be included in the logistic model.
COV	Double precision array (MAXCOV,SIZE)	input:	the entries $(l,j)(l = 1, \dots, NUMCOV)$ give the values for the $l$ th covariate of the $j$ th entity, MAXCOV=20 is the maximum number of covariates.
ESTIN	Double precision array (8)	input:	the initial estimates for the parameters: $\lambda_1, \beta_1, \gamma_1, \lambda_2, \beta_2, \gamma_2, a$ and $b_1$ .
ESTOUT	Double precision array (MB,8)	output:	the final estimates for the parameters for each bootstrap iteration $k$ ( $k = 1, \dots, NBOOT$ ), MB=100 is the maximum number of bootstraps.
LLOUT	Double precision array (MB)	output:	the final estimates for each of calculated log-likelihoods for each bootstrap iteration.
COEOUT	Double precision array (MB,MAXCOV)	output:	the entries $(k,l)(k = 1, \dots, NBOOT)$ give the final estimate of the coefficient for the $l$ th extra covariate in the $k$ th bootstrap.
ITER	Integer	output:	the number of iterations made in estimation.
IFAIL	Integer	output:	fault indicator: = 0 if no error is detected; = 1 if input or model-specification error is detected; = 2 if error is detected in estimation of $a$ & $b_1$ ; = 3 if error is detected in estimation of $\theta_1$ & $\theta_2$ .
CODE	Integer	output:	failure code for fault indicator IFAIL=2 or 3: = 0 if no error is detected; = 1 if error is detected in matrix inversion (for IFAIL=2); = 2 if suspected non-termination of $a$ & $b_1$ (for IFAIL=2); = IFAULT (for IFAIL=3); see the description in the subroutine HYBRD1 of file "HYBRID.F".

## EXAMPLE

As an illustration, the Stanford heart transplant data (Crowley and Hu, 1977) are analysed, using this algorithm to fit a Gompertz mixture model. These data have been analysed also by Larson and Dinse (1985) and Kuk (1992). We consider only the subset of 65 patients who received a transplant and who had complete data on the covariates of interest: mismatch score and age at transplant. Deaths were attributed to transplant rejection (Group 1) or other causes (Group 2). Among the 65 transplant recipients, there were 29 rejection deaths, 12 deaths from other causes, and 24 censored observations. Here the covariates are transformed to have zero means and unit variances. We include both covariates in the logistic model and the age at transplant in the component Gompertz distributions. The standard errors of the maximum likelihood estimates are obtained by bootstrap resampling with  $K = 100$  replications. The results are presented in Table 1.

Table 1: Maximum likelihood estimates (standard errors) for two-component Gompertz mixture model

Covariate	Logistic model	Component parameters	
	MLE (S.E.)	Rejection death MLE (S.E.)	Other causes MLE (S.E.)
Constant	1.396 (0.499)		
Mismatch score	0.358 (0.746)		
Age at transplant	0.303 (0.556)	1.023 (0.346)	0.275 (0.688)
$\lambda$		-6.335 (0.428)	-3.927 (0.742)
$\beta$		-0.0015 (0.0007)	-0.0055 (0.0124)

## CODE OF THE ALGORITHM MGOMP.F

```

PROGRAM MGOMP
c ****
c ** Purpose: To fit a mixture of two Gompertz distributions **
c ** to censored survival data **
c **Input files: (1) data file "surv.dat"; **
c ** (2) initial estimates file "para.dat" **
c **Output file: final estimates file "fort.25" **
c ****
c integer MB,MAX,MAXCOV
c parameter(MB=100)
c parameter(MAX=5000)
c parameter(MAXCOV=20)
c integer nboot,model,pi2eq1,size,delta(3,MAX),numcov,ifail
c integer iter,code
c double precision ftime(MAX),age(MAX),cov(MAXCOV,MAX),estin(8)
c double precision estout(MB,8),llout(MB),cofout(MB,MAXCOV)
c
c CALL init(nboot,model,pi2eq1,size,ftime,age,delta,numcov,
c + cov,estin,ifail)
c if (ifail.ne.0) goto 200
c CALL getest(nboot,model,pi2eq1,size,ftime,age,delta,numcov,
c + cov,estin,estout,llout,cofout,iter,ifail,code)
200 CALL output(nboot,size,numcov,estin,estout,llout,cofout,
c + iter,ifail,code)
c stop
c end

SUBROUTINE init(nboot,model,pi2eq1,size,ftime,age,delta,numcov,
c + cov,estin,ifail)
c ****
c ** To get data, initial estimates and model specification **
c ****
c integer MB,MAX,MAXCOV
c parameter(MB=100)
c parameter(MAX=5000)
c parameter(MAXCOV=20)
c common/randc/ix,iy,iz
c integer ix,iy,iz
c integer nboot,model,pi2eq1,size,delta(3,MAX),totcov,ifail
c double precision ftime(MAX),age(MAX),cov(MAXCOV,MAX),estin(8)
c double precision tempc(MAXCOV,MAX)
c character*1 userin
c
c open(21,file='surv.dat')
c open(22,file='para.dat')
c ifail=1
c print *, 'Please select an option: [c] for continue '
c print *, '                                [q] for quit '
c read *,userin
c if (userin(1:1).eq.'q') goto 900
c print *, ' '

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print *, 'The program fits the model either to the complete ',
+ 'data, or with optional'
print *, 'bootstrapping. Select 1 to use the complete data, ',
+ 'or type the number of'
print *, 'bootstrap repetitions which you require. Your ',
+ 'choice?'
read *, nboot
if (nboot.gt.MB) then
    print *, 'The maximum number of bootstraps allowed is ',MB
    goto 900
endif
c   input random seeds for SUBROUTINE random()
if (nboot.gt.1) then
    print *, ''
    print *, 'Input the seeds for random number generation: ',
+ 'input 3 integers between 1 and 30,000, e.g. 1 328 70'
    read *, ix, iy, iz
endif
print *, ''
print *, 'Which model is chosen? Type 1 or 2 '
read *, model
print *, ''
print *, 'Type 1 for PI_2=1.0, 0 otherwise. Your choice? '
read *, pi2eq1
print *, ''
print *, 'How many observations are there? '
read *, size
if (size.gt.MAX) then
    print *, 'The maximum number of observations allowed is ',MAX
    goto 900
endif
print *, ''
print *, 'Please input the total number of additional covariates',
+ '(other than w) in'
print *, 'the survival data file. If you do not have any',
+ 'additional covariates'
print *, 'in the data file, input 0. (At the next step, you may',
+ 'choose some (or all)'
print *, 'of them to be included in the logistic model)'
read *, totcov
if (totcov.gt.MAXCOV) then
    print *, 'The maximum number of covariates allowed is ',MAXCOV
    goto 900
endif
do 40 j=1,size
    read (21,*) ftime(j),age(j),delta(1,j),delta(2,j),
+ (tempc(k,j),k=1,totcov)
    delta(3,j)=1-delta(1,j)-delta(2,j)
40 continue
if (totcov.ne.0) CALL getcov(size,totcov,tempc,numcov,cov)
print *, ''
print *, 'RUNNING PROGRAM. PLEASE WAIT!'
read (22,*) (estin(i),i=1,8)
if (pi2eq1.eq.1) then

```

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        estin(7)=0.0
        estin(8)=0.0
    endif
    ifail=0
900  return
end

SUBROUTINE getcov(size,totcov,tempc,numcov,cov)
c ****
c ** To include additional covariates in the logistic model      **
c ****
parameter(MAX=5000)
parameter(MAXCOV=20)
integer size,totcov,numcov,addcov(MAXCOV)
double precision tempc(MAXCOV,MAX),cov(MAXCOV,MAX)
c
numcov=0
do 10 l=1,MAXCOV
    addcov(l)=0
10 continue
print *,'
print *,'Your data set has ',totcov, ' additional covariates.',
+ ' Type the covariate'
print *,'number to be included in the logistic model. Type 0',
+ ' to stop, or a number'
print *,'in the range of 1 to ', totcov
read *, k
if (k.eq.0) then
    print *,'No additional covariate is included!'
    goto 900
endif
l=1
c WHILE (k.ne.0) DO
100 if (k.ne.0) then
    numcov=numcov+1
    addcov(l)=k
    l=l+1
    print *,'
    print *,'What is the next covariate? Type 0 to stop, or a '
    print *,'number in the range of 1 to ', totcov
    read *, k
    goto 100
endif
c ENDWHILE
print *,'
print *,'Include covariate numbers: '
print *,'      ',(addcov(l),l=1,numcov)
c user specifies additional covariates
do 300 j=1,size
    do 200 l=1,numcov
        cov(l,j)=tempc(addcov(l),j)
200    continue
300    continue
900    return

```

```

    end

    SUBROUTINE getest(nboot,model,pi2eq1,size,ftime,age,delta,
+ numcov,cov,estin,estout,llout,coeout,iter,ifail,code)
c ****
c ** To obtain the bootstrap samples and get the final estimates **
c ****
    integer MB,MAX,MAXCOV
    parameter(MB=100)
    parameter(MAX=5000)
    parameter(MAXCOV=20)
    integer nboot,model,pi2eq1,size,delta(3,MAX),bdelta(3,MAX)
    integer numcov,iter,ifail,code
    double precision ftime(MAX),age(MAX),cov(MAXCOV,MAX),estin(8)
    double precision bftime(MAX),bage(MAX),bcov(MAXCOV,MAX)
    double precision estout(MB,8),llout(MB),coeout(MB,MAXCOV)

c
    if (nboot.eq.1) then
c      for complete data (nboot.eq.1)
      ifail=2
      CALL mle(nboot,model,pi2eq1,size,ftime,age,delta,numcov,cov,
+ estin,1,estout,llout,coeout,iter,ifail,code)
    else
      for (nboot.gt.1)
      do 500 k=1,nboot
        ifail=2
c      WHILE (ifail.ne.0) DO
300      if (ifail.ne.0) then
        CALL boot(size,ftime,age,delta,numcov,cov,bftime,
+ bage,bdelta,bcov)
        CALL mle(nboot,model,pi2eq1,size,bftime,bage,bdelta,
+ numcov,bcov,estin,k,estout,llout,coeout,iter,ifail,code)
        goto 300
      endif
c      ENDWHILE
500      continue
    endif
    return
  end

  SUBROUTINE mle(nboot,model,pi2eq1,size,tj,xj,dj,numcov,cj,
+ estin,bnum,estout,llout,coeout,iter,ifail,code)
c ****
c ** To obtain the maximum likelihood estimates via EM algorithm **
c ****
    integer MB,MAX,MAXCOV
    parameter(MB=100)
    parameter(MAX=5000)
    parameter(MAXCOV=20)
    integer nboot,model,pi2eq1,size,dj(3,MAX),numcov,bnum
    integer iter,ifail,code,done
    double precision tj(MAX),xj(MAX),cj(MAXCOV,MAX),estin(8)
    double precision estout(MB,8),llout(MB),coeout(MB,MAXCOV)
    double precision est(8),newest(8),pi(2,MAX),s(2,MAX),tau(2,MAX)

```

```

double precision coe(MAXCOV),ll,ddiff
intrinsic abs
c
done=0
code=0
iter=0
do 10 i=1,8
    est(i)=estin(i)
    newest(i)=est(i)
10 continue
do 12 l=1,numcov
    coe(l)=0.0
12 continue
c WHILE (done.ne.1) AND (code.eq.0) DO
15 if (done.ne.1.and.code.eq.0) then
    CALL setpi(size,pi2eq1,numcov,xj,est(7),est(8),coe,cj,pi)
    CALL calcs(model,size,tj,xj,est,s)
c for mixture model of two components
    if (pi2eq1.eq.0) then
        CALL calcab(size,numcov,est,pi,xj,dj,cj,coe,s,
+       newest(7),newest(8),tau,ifail,code)
    else
        CALL caltau(size,pi,s,tau)
        ifail=0
    endif
c ifail.ne.0 implies code.ne.0
    if (ifail.ne.0) goto 30
    ifail=3
    CALL callhy(model,size,tj,xj,dj,est,newest,tau,ifail,code)
c ifail.ne.0 implies code.ne.0
    if (ifail.ne.0) goto 30
    CALL calc1(model,size,tj,xj,dj,tau,newest(2),newest(3),1,
+   newest(1))
    CALL calc1(model,size,tj,xj,dj,tau,newest(5),newest(6),2,
+   newest(4))
    ddiff=0.0
    do 20 i=1,8
        if (newest(i).ne.0.0) then
            ddiff=ddiff+abs((est(i)-newest(i))/newest(i))
        endif
        est(i)=newest(i)
20 continue
        if (nboot.eq.1) CALL findll(model,pi2eq1,size,tj,xj,
+       dj,est,pi,ll)
        if (ddiff.lt.0.0001) done=1
        iter=iter+1
30 continue
        goto 15
    endif
c ENDWHILE
    if (code.ne.0) goto 900
    llout(bnum)=ll
    do 100 i=1,8
        estout(bnum,i)=est(i)

```

```

100  continue
    do 200 l=1,numcov
        coeout(bnum,l)=coe(l)
200  continue
900  return
end

SUBROUTINE boot(size ,ftime ,age ,delta ,numcov ,cov ,bftime ,
+  bage ,bdelta ,bcov )
c ****
c ** To obtain a nonparametric bootstrap sample **
c ****
integer MAX ,MAXCOV
parameter(MAX=5000)
parameter(MAXCOV=20)
integer size ,delta(3 ,MAX) ,numcov ,bdelta(3 ,MAX) ,num
real rand ,random
double precision ftime(MAX) ,age(MAX) ,cov(MAXCOV ,MAX)
double precision bftime(MAX) ,bage(MAX) ,bcov(MAXCOV ,MAX)
intrinsic int
c
do 150 j=1,size
    rand=random()
    rand=1.+rand*float(size)
    num=int(rand)
    bftime(j)=ftime(num)
    bage(j)=age(num)
    bdelta(1,j)=delta(1,num)
    bdelta(2,j)=delta(2,num)
    bdelta(3,j)=delta(3,num)
    do 120 l=1,numcov
        bcov(l,j)=cov(l,num)
120  continue
150  continue
return
end

SUBROUTINE setpi(size ,pi2eq1 ,numcov ,xj ,a ,b ,coe ,cj ,pi )
c ****
c ** To evaluate the mixing proportion \pi_i (i=1,2) **
c ****
integer MAX ,MAXCOV
parameter(MAX=5000)
parameter(MAXCOV=20)
integer size ,pi2eq1 ,numcov
double precision xj(MAX) ,a ,b ,coe(MAXCOV) ,cj(MAXCOV ,MAX)
double precision pi(2 ,MAX) ,tmp
intrinsic exp
c
do 100 j=1,size
    if (pi2eq1.eq.1) then
        pi(1,j)=0.0
    else
        tmp=a+b*xj(j)

```

```

        do 50 l=1,numcov
            tmp=tmp+coe(l)*cj(l,j)
50        continue
            pi(1,j)=exp(tmp)/(1.0+exp(tmp))
        endif
            pi(2,j)=1.0-pi(1,j)
100    continue
        return
    end

SUBROUTINE calcs(model,size,tj,xj,est,s)
c ****
c ** To evaluate the component survival functions s_i (i=1,2)      **
c ****
integer MAX
parameter(MAX=5000)
integer model,size,group,tmp
double precision tj(MAX),xj(MAX),est(8),s(2,MAX)
double precision lambda,beta,gamma,t1,t2,udrfl
intrinsic exp
c
udrfl=-75.
do 100 group=1,2
    tmp=(group-1)*3+1
    lambda=est(tmp)
    beta=est(tmp+1)
    gamma=est(tmp+2)
    do 50 j=1,size
        if (model.eq.2.and.group.eq.1) then
            s(group,j)=1.0
        else
            t1=exp(lambda+gamma*xj(j))
            t2=exp(beta*tj(j))-1.0
            if ((-t1*t2/beta).lt.udrfl) then
                s(group,j)=0.
                goto 50
            endif
            s(group,j)=exp(-t1*t2/beta)
        endif
50    continue
100    continue
        return
    end

SUBROUTINE calcab(size,numcov,est,pi,xj,dj,cj,coe,s,anew,bnew,
+ tau,ifail,code)
c ****
c ** To perform iterative steps for the computation of the          **
c ** coefficients \alpha of the covariates                         **
c ****
integer MAX,MAXCOV
parameter(MAX=5000)
parameter(MAXCOV=20)
integer size,dj(3,MAX),numcov,ifail,code,ctr,order,done

```

```

double precision est(8),pi(2,MAX),xj(MAX),cj(MAXCOV,MAX)
double precision coe(MAXCOV),s(2,MAX),anew,bnew,tau(2,MAX)
double precision a,b,newcoe(MAXCOV),ddiff
double precision arr(MAXCOV+2,MAXCOV+2),u(MAXCOV+2)
intrinsic abs

c
ctr=0
code=0
order=numcov+2
done=0
a=est(7)
b=est(8)
c WHILE (done.ne.1) AND (code.eq.0) DO
10 if (done.ne.1.and.code.eq.0) then
    ddiff=0.
    CALL caltau(size,pi,s,tau)
    CALL setarr(size,a,b,pi,xj,dj,cj,coe,numcov,tau,order,arr,u)
    CALL invert(arr,order,code)
    if (code.ne.0) goto 100
    anew=a
    bnew=b
    do 20 i=1,order
        anew=anew+arr(1,i)*u(i)
        bnew=bnew+arr(2,i)*u(i)
20 continue
    do 30 l=1,numcov
        newcoe(l)=coe(l)
30 continue
    do 50 i=1,order
        do 40 l=1,numcov
            newcoe(l)=newcoe(l)+arr(l+2,i)*u(i)
40 continue
50 continue
    ddiff=abs(a-anew)+abs(b-bnew)
    do 60 l=1,numcov
        ddiff=ddiff+abs(newcoe(l)-coe(l))
        coe(l)=newcoe(l)
60 continue
    CALL setpi(size,pi2eq1,numcov,xj,anew,bnew,newcoe,cj,pi)
    a=anew
    b=bnew
    if (ddiff.lt.0.0001) done=1
    ctr=ctr+1
    if (ctr.gt.800 .or. ddiff.gt.15.0) code=2
100 continue
    goto 10
endif
c ENDWHILE
if (code.eq.0) then
    ifail=0
    CALL caltau(size,pi,s,tau)
endif
return
end

```

```

SUBROUTINE caltau(size,pi,s,tau)
c **** To evaluate the posterior probability \tau, using ****
c ** pre-calculated values for \pi_i and s_i (i=1,2) ****
c ****
integer MAX
parameter(MAX=5000)
integer size
double precision pi(2,MAX),s(2,MAX),tau(2,MAX),temp
c
do 100 j=1,size
    temp=pi(1,j)*s(1,j)
    tau(1,j)=temp/(temp+pi(2,j)*s(2,j))
    tau(2,j)=1.0-tau(1,j)
100 continue
return
end

SUBROUTINE setarr(size,a,b,pi,xj,dj,cj,coe,numcov,tau,order,arr,u)
c **** To evaluate the matrix arr and the column vector u ****
c ****
integer MAX,MAXCOV
parameter(MAX=5000)
parameter(MAXCOV=20)
integer size,dj(3,MAX),numcov,order
double precision a,b,pi(2,MAX),xj(MAX),cj(MAXCOV,MAX)
double precision coe(MAXCOV),tau(2,MAX)
double precision arr(MAXCOV+2,MAXCOV+2),u(MAXCOV+2)
double precision varr(MAX),xarr(MAXCOV+2,MAX),tmp,yj,t
c
do 20 i=1,order
    u(i)=0.0
    do 10 k=1,order
        arr(i,k)=0.0
10    continue
20    continue
do 40 j=1,size
    varr(j)=pi(1,j)*pi(2,j)
    xarr(1,j)=1.0
    xarr(2,j)=xj(j)
    do 30 l=1,numcov
        xarr(l+2,j)=cj(l,j)
30    continue
40    continue
do 70 i=1,order
    do 60 k=1,order
        tmp=0.0
        do 50 j=1,size
            tmp=tmp+xarr(i,j)*xarr(k,j)*varr(j)
50    continue
        arr(i,k)=tmp
60    continue

```

```

70    continue
do 100 j=1,size
    yj=float(dj(1,j))+dj(3,j)*tau(1,j)
    tmp=a+b*xarr(2,j)
    do 80 l=1,numcov
        tmp=tmp+coe(l)*cj(l,j)
80    continue
    t=exp(tmp)
    u(1)=u(1)+yj-t/(1.0+t)
    do 90 k=2,order
        u(k)=u(k)+yj*xarr(k,j)-xarr(k,j)*t/(1.0+t)
90    continue
100 continue
return
end

SUBROUTINE invert(arr,order,code)
c ****
c ** To return the inverse of the input matrix arr in the same      **
c ** matrix (so arr is over-written)                                     **
c ****
integer MAXCOV
parameter(MAXCOV=20)
integer order,code,ctr,nullty,ifault
real matrix((MAXCOV+2)*(MAXCOV+3))
real inv((MAXCOV+2)*(MAXCOV+3)),work(MAXCOV+2)
double precision arr(MAXCOV+2,MAXCOV+2)
c
ctr=0
do 20 i=1,order
    do 10 k=1,i
        ctr=ctr+1
        matrix(ctr)=arr(i,k)
10    continue
20    continue
nullty=0
ifault=0
CALL syminv(matrix,order,ctr,inv,work,nullty,ifault)
if (ifault.ne.0) then
    code=1
    goto 900
endif
ctr=0
do 40 i=1,order
    do 30 k=1,i
        ctr=ctr+1
        arr(i,k)=inv(ctr)
        arr(k,i)=inv(ctr)
30    continue
40    continue
900 return
end

SUBROUTINE callhy(model,size,tj,xj,dj,est,newest,tau,ifail,code)

```

```

c ****
c ** To call HYBRD1 for solving system of nonlinear equations    **
c ****
integer MAX
parameter(MAX=5000)
integer model,size,dj(3,MAX),ifail,code,group
double precision tj(MAX),xj(MAX),est(8),newest(8),tau(2,MAX)
external evalfn, HYBRD1
integer sizehy,n,indlo,tmp
double precision beta,gamma,x(2),xtol,fvec(2),work(100)
common/hybr1/hmodel,hsize,hdj,hgroup
integer hmodel,hsize,hdj(3,MAX),hgroup
common/hybr2/htj,hxj,htau
double precision htj(MAX),hxj(MAX),htau(2,MAX)

c
code=0
indlo=1
if (model.eq.2) indlo=2
c set values for the common statement
hmodel=model
hsize=size
do 10 j=1,hsize
  htj(j)=tj(j)
  hxj(j)=xj(j)
  htau(1,j)=tau(1,j)
  htau(2,j)=tau(2,j)
  hdj(1,j)=dj(1,j)
  hdj(2,j)=dj(2,j)
  hdj(3,j)=dj(3,j)
10 continue
do 20 group=indlo,2
  if (code.ne.0) goto 20
  hgroup=group
  tmp=(group-1)*3+1
  beta=est(tmp+1)
  gamma=est(tmp+2)
  ifault=0
  n=2
  x(1)=beta
  x(2)=gamma
  xtol=0.000000001
  sizehy=100
  CALL HYBRD1(evalfn,n,x,fvec,xtol,ifault,work,sizehy)
c ifault=1 implies no error is detected (see subroutine HYBRD1 in
c file "hybrid.f"); change code=1 for ifault=0
  if (ifaule.ne.1) then
    code=1
    if (ifaule.ne.0) code=ifaule
    goto 20
  endif
  newest(tmp+1)=x(1)
  newest(tmp+2)=x(2)
20 continue
  if (code.eq.0) ifail=

```

```

return
end

SUBROUTINE evalfn(n,x,fvec,iflag)
c ****
c ** To evaluate the system of functions **
c ****
integer MAX
parameter(MAX=5000)
integer n,iflag
double precision x(*),fvec(*)
double precision t1,t2,t3,dc,tc,hval
double precision tot1,tot2,beta,gamma,lambda
common/hybr1/hmodel,hsiz,hsize,hdj,hgroup
integer hmodel,hsiz,hdj(3,MAX),hgroup
common/hybr2/htj,hxj,htau
double precision htj(MAX),hxj(MAX),htau(2,MAX)
intrinsic exp
c
beta=x(1)
gamma=x(2)
CALL calcl(hmodel,hsiz,htj,hxj,hdj,htau,beta,gamma,
+ hgroup,lambda)
tot1=0.0
tot2=0.0
do 100 j=1,hsiz
    dc=hdj(hgroup,j)
    tc=htau(hgroup,j)
    t1=exp(lambda+gamma*hxj(j))
    t2=exp(beta*htj(j))-1.0
    hval=(t1*t2)
    t1=(dc+hdj(3,j)*tc)
    t2= -htj(j)/beta*exp(lambda+gamma*hxj(j)+beta*htj(j))
    t3=hval/(beta*beta)
    tot1=tot1+dc*htj(j)+t1*(t2+t3)
    t1=dc*hxj(j)
    t2= -(dc+hdj(3,j)*tc)
    t3=hxj(j)*hval/beta
    tot2=tot2+t1+t2*t3
100 continue
fvec(1)=tot1
fvec(2)=tot2
return
end

SUBROUTINE calcl(model,size,tj,xj,dj,tau,beta,gamma,ind,lambda)
c ****
c ** To solve for lambda given current estimates of beta & gamma **
c ****
integer MAX
parameter(MAX=5000)
integer model,size,dj(3,MAX),ind
double precision tj(MAX),xj(MAX),tau(2,MAX),beta,gamma,lambda
double precision t,t1,t2

```

```

intrinsic log
c
lambda=0.0
if (ind.eq.1.and.model.eq.2) then
  beta=0.0
  gamma=0.0
  goto 900
endif
t1=0.0
t2=0.0
do 10 j=1,size
  t1=t1+float(dj(ind,j))
  t=exp(gamma*xj(j))*(exp(beta*tj(j))-1.0)
  t2=t2+(float(dj(ind,j))+dj(3,j)*tau(ind,j))*t
10 continue
lambda=log(beta*t1/t2)
900 return
end

SUBROUTINE findll(model,pi2eq1,size,tj,xj,dj,est,pi,ll)
c ****
c ** To evaluate the maximum log likelihood **
c ****
integer MAX
parameter(MAX=5000)
integer model,pi2eq1,size,dj(3,MAX)
double precision tj(MAX),xj(MAX),est(8),pi(2,MAX),ll
double precision t1,t2,t3,te,tf,mul,udrfl
double precision l1,g1,b1,l2,g2,b2,p1,p2
intrinsic log
c
udrfl=-75.
t1=0.0
t2=0.0
t3=0.0
l1=est(1)
b1=est(2)
g1=est(3)
l2=est(4)
b2=est(5)
g2=est(6)
do 20 j=1,size
  p1=pi(1,j)
  p2=pi(2,j)
  te=exp(l2+g2*xj(j))*(exp(b2*tj(j))-1)/b2
  if (pi2eq1.eq.1) then
    t1=t1+dj(2,j)*log(p2)
  else
    t1=t1+dj(1,j)*log(p1)+dj(2,j)*log(p2)
  endif
  t2=t2+dj(2,j)*(l2+g2*xj(j)+b2*tj(j)-te)
  if (model.eq.2) then
    mul=1.0
  else

```

```

        tf=exp(l1+g1*xj(j))*(exp(b1*tj(j))-1)/b1
        t2=t2+dj(1,j)*(l1+g1*xj(j)+b1*tj(j)-tf)
        if (-tf.lt.udrf1) then
            mul=0.
        else
            mul=exp(-tf)
        endif
    endif
    if (-te.lt.udrf1) then
        t3=t3+dj(3,j)*log(p1*mul)
    else
        t3=t3+dj(3,j)*log(p1*mul+p2*exp(-te))
    endif
20    continue
    ll=t1+t2+t3
    return
end

SUBROUTINE output(nboot,size,numcov,estin,estout,llout,
+ coeout,iter,ifail,code)
c ****
c ** To write the result into output file "fort.25" **
c ****
integer MB,MAXCOV
parameter(MB=100)
parameter(MAXCOV=20)
integer nboot,size,numcov,iter,ifail,code
double precision estin(8),estout(MB,8),llout(MB)
double precision coeout(MB,MAXCOV),sum(8+MAXCOV),bar(8+MAXCOV)
double precision se(8+MAXCOV),sumsq(8+MAXCOV)
c
write (25,900) (estin(i),i=1,3)
write (25,905) (estin(i),i=4,6)
write (25,910) (estin(i),i=7,8)
write (25,915) size
if (ifail.ne.0) then
    print *, 'The program is terminated: ifail= ',ifail
    write (25,920) ifail
    if (ifail.eq.2.and.code.eq.1) write (25,922)
    if (ifail.eq.2.and.code.eq.2) write (25,925)
    if (ifail.eq.3) write (25,927) code
else
    print *
    print *, 'FINISH. THE RESULT IS AT THE OUTPUT FILE "fort.25".'
    do 100 l=1,8+numcov
        sum(l)=0.
        sumsq(l)=0.
100   continue
    write (25,930)
    do 500 k=1,nboot
        if (nboot.eq.1) then
            write (25,935) k,iter,llout(k)
            write (25,940) (estout(k,l),l=1,3)
            write (25,945) (estout(k,l),l=4,6)

```

```

        write (25,950) (estout(k,1),l=7,8)
        if (numcov.ne.0) write (25,955) (coeout(k,1),l=1,numcov)
        goto 500
    endif
c      if (nboot.eq.1) goto 500
    do 200 l=1,8
        sum(l)=sum(l)+estout(k,l)
        sumsq(l)=sumsq(l)+estout(k,l)*estout(k,l)
200   continue
    do 300 l=1,numcov
        sum(8+l)=sum(8+l)+coeout(k,l)
        sumsq(8+l)=sumsq(8+l)+coeout(k,l)*coeout(k,l)
300   continue
500   continue
        if (nboot.eq.1) goto 650
        do 550 l=1,8+numcov
            bar(l)=sum(l)/float(nboot)
            se(l)=sqrt(sumsq(l)/float(nboot)-bar(l)*bar(l))
550   continue
        do 600 l=1,8+numcov
            if (l.le.8) then
                write (25,960) l,bar(l),se(l)*float(nboot)/float(nboot-1)
            else
                write (25,965) l-8,bar(l),
+                se(l)*float(nboot)/float(nboot-1)
            endif
600   continue
650   endif
900   format('Initial Estimates:',/2x,'lambda1=',F9.5,',     beta1=',
+ F9.5,'     gamma1=',F9.5)
905   format(2x,'lambda2=',F9.5,',     beta2=',
+ F9.5,'     gamma2=',F9.5)
910   format(8x,'a=',F9.5,',     b_1=',F9.5)
915   format(/,'There are ',I6,', values in the data set',/)
920   format(/,'The program is terminated: ifail= ',I3)
922   format(/,'The inverion of matrix does not make good!!')
925   format(/,'Suspected non-termination of a and b_1!!')
927   format(/,'The failure code in HYBRD1 routine is ',I3)
930   format(/,'-----',
+ /,'Summary of results:')
935   format(/,'Bootstrap ',I3,: after iteration ',I3,
+ ', Log Likelihood= ',F15.8)
940   format('  Estimates:',/2x,'lambda1=',F11.7,',     beta1=',
+ F11.7,'     gamma1=',F11.7)
945   format(2x,'lambda2=',F11.7,',     beta2=',
+ F11.7,'     gamma2=',F11.7)
950   format(8x,'a=',F11.7,',     b_1=',F11.7)
955   format(2x,'Coefficients for additional covariates are: ',
+ /,20F9.5)
960   format('mean (s.e) of estimate ',I2,: ',F11.7,' (',F9.7,')')
965   format('mean (s.e) of additional covariate ',I2,: ',F11.7,
+ ' (',F9.7,')')
    return
end

```

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