pdInd: G matrix with pattern of zeros

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pdInd is a constructor for pdClasses that define G matrices to model the variance of random effects for models in the nlme package.

Mixed models in which many predictors have random slopes often fail to converge in part because of the large number of parameters in the full covariance (G) matrix for random effects. One way of fitting a more parsimonious model that includes random slopes is to use pdDiag with zeros off the diagonal. However, this also forces zero covariances between random slopes and and the random intercept, resulting in a model that is not equivariant with respect to location transformations of the predictors with random slopes. The alternative remedy of omitting random slopes for some predictors can lead to biased estimates and incorrect standard errors of regression coefficients.

The default covariance pattern for pdInd produces a G matrix with zero covariances except in the first row and column. If the first random effect is the intercept, the resulting model assumes independence between random slopes without imposing minimality of variance over the possibly arbitrary origin. This imposition is the reason that having all covariances equal to zero results in a model that fails to be equivariant under location transformations.

The optional cov parameter can be used to allow selected non-zero covariance between random slopes.

For example, if two variables, X1 and X2 have random effects, the random effects model would be specified in a call to lme as random = ~1 + X1 + X2.

The default G matrix has the form:

$$G = \begin{pmatrix} g_{00} & g_{01} & g_{02} \\ g_{10} & g_{11} & g_{12} \\ g_{20} & g_{21} & g_{22} \end{pmatrix}$$

With pdDiag, all the off-diagonal elements of G are constrained to 0. Forcing g_{01} and g_{02} to be 0 produces a model that is not equivariant with respect to location changes in X1 and X2. The value at which the variance of Y given X1 and X2 is minimized is forced to be 0 for both variables.

However, constraining $g_{12} = 0$ produces a model that is equivariant with respect to location-scale transformation of X1 and X2 and in which the random between cluster values of regression slopes for each variable are independent of each other.

The pdInd class of positive-definite matrices creates, by default, a matrix with arbitrary values along the diagonal and in the first row and column, but zeros elsewhere. In the case of a 4×4 matrix, this produces:

$$G = \begin{pmatrix} g_{00} & g_{01} & g_{02} & g_{03} \\ g_{10} & g_{11} & 0 & 0 \\ g_{20} & 0 & g_{22} & 0 \\ g_{30} & 0 & 0 & g_{33} \end{pmatrix}$$

The challenge in parametrizing the G matrix is finding an unconstrained parametrization that results in a positive-definite matrix with selected covariances constrained to 0.

We consider a right-Cholesky decomposition noting that the diagonal component in the following factorization results in a diagonal component in the variance matrix.

$$\begin{pmatrix} G_{11} & G_{12} \\ G_{12}' & G_{22} \end{pmatrix} = \begin{pmatrix} R_{11} & R_{12} \\ 0 & D_{22} \end{pmatrix} \begin{pmatrix} R_{11}' & 0 \\ R_{12}' & D_{22} \end{pmatrix} = \begin{pmatrix} R_{11}R_{11}' + R_{12}R_{12}' & R_{12}D_{22} \\ D_{22}R_{12}' & D_{22}' \end{pmatrix}$$

With R_{11} square upper-triangular and D_{22} square diagonal, G_{22} must be diagonal. In addition, patterns of zeros in the R_{12} matrix, lying above the diagonal D_{22} matrix, are preserved in G_{12} .

Note that the unconstrained Cholesky parametrization uses the log of the diagonal elements of the triangular factor.

$$(fac \leftarrow cbind(c(1,0,0,0,0), c(1,2,0,0,0), c(0,1,3,0,0), c(1,0,0,4,0), c(0,1,0,0,5)))$$

fac %*% t(fac)

Note how the pattern of zeros in the last three columns of the first two rows is preserved in the cross product since the lower 3×3 diagonal block matrix is itself diagonal.

However, if the lower 3×3 block diagonal matrix is not diagonal, then the pattern of zeros in the top two rows above it is not necessarily preserved.

$$(fac \leftarrow cbind(c(1,0,0,0,0), c(1,2,0,0,0), c(0,1,3,0,0), c(1,0,0,4,0), c(0,1,0,-1,5)))$$

```
[,1] [,2] [,3] [,4] [,5]
[1,] 1 1 0 1 0
[2,] 0 2 1 0 1
[3,] 0 0 3 0 0
[4,] 0 0 0 4 -1
[5,] 0 0 0 5
```

fac %*% t(fac)

[,1] [,2] [,3] [,4] [,5]

But patterns of zeros above non-diagonal blocks are not necessarily preserved.

The cov parameter allows the user to specify a pattern of zeros in the upper triangle of the upper-triangular 'R' factor of the G matrix. As observed above, in some cases this will result in the same pattern in the G matrix. Even if the pattern in the R factor does not create a similar pattern in the G matrix, the model for the G matrix will nevertheless have the number of additional parameters for covariance as given by the TRUE entries in the upper diagonal of the cov matrix.

Predictor transformations to improve convergence

We will show how to use lchol(getG(fit)) to suggest transformations of predictors that appear in random effects models to help improve the convergence of mixed models fits.

We conjecture that the condition number of the Hessian matrix for the parameters in the G matrix may be approximately the square of the condition number for the G matrix. Thus a condition number for G in the vicinity of 10⁷ would effectively result in singularity of the Hessian.

Rescaling and relocating the predictors with random slopes can improve ill-conditioning of the G matrix.

Let

$$\mathbf{Z} = egin{pmatrix} 1 \ Z_1 \ dots \ Z_k \end{pmatrix}$$

represent the vector of variables with random effects \mathbf{u} . For the *i*th cluster the contribution from level-2 random effects is:

$$\mathbf{Z}_{i}'\mathbf{u}_{i}$$

Consider a location-scale tranformation of the variables in \mathbf{Z} . It has the form $\mathbf{Z}^* = T\mathbf{Z}$ where T is lower triangular with the form

$$T = \begin{bmatrix} 1 & 0 \\ a_1 \\ a_2 \\ \vdots \\ a_k \end{bmatrix}$$

with B a diagonal matrix containing scaling coefficients, $b_1, b_2, ..., b_k$, so that

$$Z_i^* = a_i + b_i Z_i$$

If $G = Var(\mathbf{u})$ then

$$G^* = T'^{-1}GT^{-1}$$

where $G^* = Var(\mathbf{u}^*)$ with

$$\mathbf{Z}_i'\mathbf{u}_i = {\mathbf{Z}_i^*}'\mathbf{u}_i^*$$

Therefore, if we factor

$$G = T'T$$

with T lower-triangular, $\frac{1}{t_{11}}T$ provides a transformation of $\mathbf Z$ that minimizes the condition number of G^* . In R, using spida2::lchol this is simply getG(fit) %>% lchol %>% {./.[1,1]}.

Improving convergence – IN PROGRESS

- 1. Location scale transformations of Z variables. Note: don't need to change to X variables but might like to for purposes of inference.
- 2. Parsimonious G matrices: Use pdInd and pdDiag.
- 3. Note that LRTs with anova are likely to be informative to compare nested RE models with singular Hessians provided differences are only in covariance structure, i.e. same non-zero diagonal elements.
- 4. STUDY: any effect on vcov of changing to equivalent Zs without changing X.

Nuts and Bolts of the G matrix in lme – IN PROGRESS

The initial call to pdConstruct.pdInd occurs in the initialization phase of lme which call reStruct which in turn calls pdMat that creates an empty pdInd object, sets value <- numeric(0) and returns:

```
pdConstuct(object, value, form, nam, data)
```

pdInd methods

```
methods(class='pdInd')
   [1] pdConstruct pdFactor
                                pdMatrix
   see '?methods' for accessing help and source code
gnew:::pdInd
   Running .onLoad
   function (value = numeric(0), form = NULL, nam = NULL,
               data = sys.parent(), cov = NULL)
       # unchanged
       object <- numeric(0)</pre>
       class(object) <- c("pdInd", "pdMat")</pre>
       pdConstruct(object, value, form, nam, data, cov)
   <environment: namespace:gnew>
gnew:::pdConstruct.pdInd
   function (object, value = numeric(0), form = formula(object),
               nam = Names(object), data = sys.parent(),
               cov = NULL,
                ...)
       # note that pdConstruct.pdMat return an upper-triangular R factor, === might not be correct
       if(!is.null(attr(object,'cov'))) cov <- attr(object,'cov')</pre>
       if(!is.null(attr(value,'cov'))) cov <- attr(value,'cov')</pre>
       val <- nlme:::pdConstruct.pdMat(object,</pre>
                                         value = value,
                                         form = form,
                                         nam = nam
                                         data = data)
```

```
attr(val,'cov') <- cov
       if (length(val) == 0) {
         class(val) <- c("pdInd", "pdMat")</pre>
         return(val)
       # mod 2015 07 04: added arbitrary cov structure of non zero
       isRmat \leftarrow function(x) all(x[row(x) > col(x)] == 0) # is lower triangle == 0?
       if (is.matrix(val)) {
         if(is.null(cov)) {
            if(!is.null(attr(val,'cov'))) cov <- attr(val,'cov')</pre>
           else cov <- (row(val) == 1) & (col(val) > 1)
               disp(cov)
         if(isRmat(val)){
           value <- c(log(diag(val)), val[cov])</pre>
           # keeping only the entries that should be non-zero
         } else stop("matrix should be an upper triangular matrix")
         attributes(value) <-
            attributes(val)[names(attributes(val)) != "dim"]
         attr(value,"cov") <- cov</pre>
         class(value) <- c("pdInd", "pdMat")</pre>
         attr(value,"invert") <- FALSE</pre>
         return(value)
       }
       stop("shouldn't get here in pdConstruct.pdInd")
       Ncol \leftarrow (length(val) + 1)/2
       if (length(val) != 2*round(Ncol) - 1) {
         stop(gettextf("an object of length %d does not match a pdInd factor (diagonal + covariances wi
                         length(val)), domain = NA)
       class(val) <- c("pdInd", "pdMat")</pre>
       val
     }
   <environment: namespace:gnew>
gnew:::pdFactor.pdInd
   function (object)
     {
       invert <- attr(object, "invert")</pre>
       cov <- attr(object,"cov")</pre>
       object <- as.vector(object)</pre>
       Ncov <- sum(cov)</pre>
       Ncol <- length(object) - Ncov
       # was:
           L <- matrix(0,Ncol,Ncol)</pre>
           diag(L) <- exp( object[1:Ncol])</pre>
           if ( Ncol > 1 ) L[row(L)>1 & col(L)==1] <-
              object[(Ncol+1):length(object)]
           if(invert) c(t(solve(L))) else c(L2R(L))
       R <- matrix(0,Ncol,Ncol)</pre>
       diag(R) <- exp( object[1:Ncol])</pre>
       if ( Ncol > 1 ) R[cov] <-
         object[(Ncol+1):length(object)]
```

```
if(invert) c(t(solve(f2L(R)))) else c(R)
   <environment: namespace:gnew>
gnew:::pdMatrix.pdInd
   function (object, factor = FALSE)
       if (!isInitialized(object)) {
         stop("cannot extract matrix from an uninitialized object")
       cov <- attr(object,"cov")</pre>
       Ncov <- sum(cov)</pre>
       Ncol <- length(object) - Ncov</pre>
       value <- array(pdFactor(object), c(Ncol, Ncol),</pre>
                       attr(object, "Dimnames"))
       ob <- as.vector(object) # subsetting object calls pdMatrix!</pre>
       attr(value, "logDet") <- 2*sum(ob[1:Ncol])</pre>
       if (factor) value else crossprod(value)
   <environment: namespace:gnew>
gnew:::solve.pdInd
   function (a, b, ...)
       if (!isInitialized(a)) {
         stop("cannot get the inverse of an uninitialized object")
       attr(a, 'invert') <- !attr(a, 'invert')</pre>
       a
            Ncol \leftarrow (length(a) + 1)/2
       #
            ob <- as.vector(a)</pre>
       #
             if( Ncol == 1) ret <- -ob[1]
            else ret <-
              c( -ob[1:Ncol] ,
       #
                 - exp(ob[1])*ob[(Ncol+1):length(ob)]/exp(ob[2:Ncol]))
            attributes(ret) <- attributes(a)
       #
            ret
   <environment: namespace:gnew>
```

pdMat methods

```
methods(class='pdMat')
    Γ11 Γ
                       [<-
                                      as.matrix
                                                                    coef<-
    [6] corMatrix
                                                     isInitialized logDet
                       \mathtt{Dim}
                                      formula
   [11] matrix<-
                       Names
                                      Names<-
                                                     pdConstruct
                                                                   pdFactor
   [16] pdMatrix
                       plot
                                      print
                                                                    summary
   [21] VarCorr
   see '?methods' for accessing help and source code
```

```
nlme:::pdMat
   function (value = numeric(0), form = NULL, nam = NULL, data = sys.frame(sys.parent()),
       pdClass = "pdSymm")
   {
       if (inherits(value, "pdMat")) {
           pdClass <- class(value)</pre>
       object <- numeric(0)</pre>
       class(object) <- unique(c(pdClass, "pdMat"))</pre>
       pdConstruct(object, value, form, nam, data)
   <bytecode: 0x0000000232d4d38>
   <environment: namespace:nlme>
nlme:::pdConstruct.pdMat
   function (object, value = numeric(0), form = formula(object),
       nam = Names(object), data = sys.frame(sys.parent()), ...)
   {
       if (inherits(value, "pdMat")) {
           if (length(form) == 0) {
               form <- formula(value)</pre>
           if (length(nam) == 0) {
               nam <- Names(value)</pre>
           }
           if (isInitialized(value)) {
                return(pdConstruct(object, as.matrix(value), form,
                    nam, data))
           }
           else {
               return(pdConstruct(object, form = form, nam = nam,
                    data = data))
       if (length(value) > 0) {
           if (inherits(value, "formula") || data.class(value) ==
                "call") {
                if (!is.null(form)) {
                    warning("ignoring argument 'form'")
               form <- formula(value)</pre>
                if (length(form) == 3) {
                    form <- list(form)</pre>
                }
           }
           else if (is.character(value)) {
               if (length(nam) > 0) {
                    warning("ignoring argument 'nam'")
               }
               nam <- value
           else if (is.matrix(value)) {
               vdim <- dim(value)</pre>
```

```
if (length(vdim) != 2 || diff(vdim) != 0) {
            stop("'value' must be a square matrix")
        if (length(unlist(vnam <- dimnames(value))) > 0) {
            vnam <- unique(unlist(vnam))</pre>
            if (length(vnam) != vdim[1]) {
              stop("dimnames of 'value' must match or be NULL")
            dimnames(value) <- list(vnam, vnam)</pre>
            if (length(nam) > 0) {
              if (any(is.na(match(nam, vnam))) || any(is.na(match(vnam,
                nam)))) {
                 stop("names of 'value' are not consistent with 'nam' argument")
              value <- value[nam, nam, drop = FALSE]</pre>
            }
            else {
              nam <- vnam
        }
        form <- form
        nam <- nam
        object <- chol((value + t(value))/2)
        attr(object, "dimnames") <- NULL
        attr(object, "rank") <- NULL
    else if (is.numeric(value)) {
        value <- as.numeric(value)</pre>
        attributes(value) <- attributes(object)</pre>
        object <- value
    else if (data.class(value) == "list") {
        if (!is.null(form)) {
            warning("ignoring argument 'form'")
        form <- value
    }
    else {
        stop(gettextf("%s is not a valid object for \"pdMat\"",
            sQuote(deparse(object))), domain = NA)
    }
}
if (!is.null(form)) {
    if (inherits(form, "formula") && length(form) == 3) {
        form <- list(form)</pre>
    if (is.list(form)) {
        if (any(!unlist(lapply(form, function(el) {
            inherits(el, "formula") && length(el) == 3
            stop("all elements of 'form' list must be two-sided formulas")
        val <- list()</pre>
        for (i in seq_along(form)) {
```

```
if (is.name(form[[i]][[2]])) {
               val <- c(val, list(form[[i]]))</pre>
             }
            else {
               val <- c(val, eval(parse(text = paste("list(",</pre>
                 paste(paste(all.vars(form[[i]][[2]]), deparse(form[[i]][[3]]),
                   sep = "~"), collapse = ","), ")"))))
             }
        }
        form <- val
        class(form) <- "listForm"</pre>
        namesForm <- Names(form, data)</pre>
    else {
        if (inherits(form, "formula")) {
             namesForm <- Names(asOneSidedFormula(form), data)</pre>
        }
        else {
             stop("'form' can only be a formula or a list of formulae")
    if (length(namesForm) > 0) {
        if (length(nam) == 0) {
             nam <- namesForm
        }
        else {
             if (any(noMatch <- is.na(match(nam, namesForm)))) {</pre>
               err <- TRUE
               namCopy <- nam
               indNoMatch <- seq_along(nam)[noMatch]</pre>
               if (any(wch1 <- (nchar(nam, "c") > 12))) {
                 wch1 <- substring(nam, nchar(nam, "c") -</pre>
                   10) == "(Intercept)"
                 if (any(wch1)) {
                   namCopy[indNoMatch[wch1]] <- substring(nam[wch1],</pre>
                     1, nchar(nam[wch1], "c") - 12)
                   noMatch[wch1] <- FALSE
                   indNoMatch <- indNoMatch[!wch1]</pre>
               }
               if (sum(noMatch) > 0) {
                 namCopy[indNoMatch] <- paste(namCopy[indNoMatch],</pre>
                   "(Intercept)", sep = ".")
               }
               if (!any(is.na(match(namCopy, namesForm)))) {
                 err <- FALSE
               if (err)
                 stop("'form' not consistent with 'nam'")
             }
        }
    }
}
if (is.matrix(object)) {
```

```
if (length(nam) > 0 && (length(nam) != dim(object)[2])) {
               stop("length of 'nam' not consistent with dimensions of initial value")
       }
       attr(object, "formula") <- form</pre>
       attr(object, "Dimnames") <- list(nam, nam)</pre>
       object
   <bytecode: 0x000000023341cd8>
   <environment: namespace:nlme>
nlme:::pdMatrix.pdMat
   function (object, factor = FALSE)
       if (!isInitialized(object)) {
           stop("cannot access the matrix of uninitialized objects")
       }
       if (factor) {
           stop("no default method for extracting the square root of a \"pdMat\" object")
       }
       else {
           crossprod(pdMatrix(object, factor = TRUE))
       }
   }
   <bytecode: 0x00000001e592f20>
   <environment: namespace:nlme>
nlme:::pdFactor.pdMat
   function (object)
       c(qr.R(qr(pdMatrix(object))))
   <bytecode: 0x00000001e6b6840>
   <environment: namespace:nlme>
nlme:::solve.pdMat
   function (a, b, ...)
   {
       if (!isInitialized(a)) {
           stop("cannot get the inverse of an uninitialized object")
       matrix(a) <- solve(as.matrix(a))</pre>
   }
   <bytecode: 0x00000001e5efe88>
   <environment: namespace:nlme>
nlme:::VarCorr.pdMat
   function (x, sigma = 1, rdig = 3, ...)
   {
       sx <- summary(x)</pre>
       sd <- sigma * attr(sx, "stdDev")</pre>
       var <- sd^2</pre>
```

```
p \leftarrow dim(sx)[2]
       v <- array(c(var, sd), c(p, 2), list(names(sd), c("Variance",</pre>
           "StdDev")))
       attr(v, "formStr") <- if (inherits(attr(x, "formula"), "listForm")) {</pre>
           paste(class(x)[[1]], "(list(", paste(sapply(attr(x, "formula"),
                function(x) as.character(deparse(x))), collapse = ","),
                "))", sep = "")
       }
       else {
           paste(class(x)[[1]], "(", substring(deparse(attr(x, "formula")),
                2), ")", sep = "")
       }
       if (attr(sx, "noCorrelation") || p <= 1)</pre>
           return(v)
       ll <- lower.tri(sx)</pre>
       sx[ll] <- format(round(sx[ll], digits = rdig))</pre>
       sx[!11] <- ""
       if (!is.null(colnames(sx))) {
           sx[1, ] <- abbreviate(colnames(sx), minlength = rdig +</pre>
       dimnames(sx) <- list(names(sd), c("Corr", rep("", p - 1)))</pre>
       attr(v, "corr") <- sx[, -p, drop = FALSE]</pre>
   }
   <bytecode: 0x00000001e4cd778>
   <environment: namespace:nlme>
nlme:::as.matrix.pdMat
   function (x, ...)
   pdMatrix(x)
   <bytecode: 0x00000001e399b90>
   <environment: namespace:nlme>
nlme:::`matrix<-.pdMat`</pre>
   function (object, value)
   {
       value <- as.matrix(value)</pre>
       if (isInitialized(object) && any(dim(value) != Dim(object))) {
           stop("cannot change dimensions on an initialized \"pdMat\" object")
       pdConstruct(object, value)
   <bytecode: 0x00000001e4b1950>
   <environment: namespace:nlme>
nlme:::coef.pdMat
   function (object, unconstrained = TRUE, ...)
       if (unconstrained || !isInitialized(object)) {
           as.vector(object)
       }
       else {
```

```
stop("do not know how to obtain constrained coefficients")
       }
   }
   <bytecode: 0x00000001e65e498>
   <environment: namespace:nlme>
nlme:::`coef<-.pdMat`</pre>
   function (object, ..., value)
       value <- as.numeric(value)</pre>
       if (isInitialized(object)) {
           if (length(value) != length(object)) {
                stop("cannot change the length of the parameter after initialization")
           }
       }
       else {
           return(pdConstruct(object, value))
       class(value) <- class(object)</pre>
       attributes(value) <- attributes(object)</pre>
       value
   <bytecode: 0x00000001e5aac68>
   <environment: namespace:nlme>
```

pdSymm methods

```
nlme:::pdSymm
   function (value = numeric(0), form = NULL, nam = NULL, data = parent.frame())
       object <- numeric(0)</pre>
       class(object) <- c("pdSymm", "pdMat")</pre>
       pdConstruct(object, value, form, nam, data)
   <bytecode: 0x00000001bae23e8>
   <environment: namespace:nlme>
methods(class='pdSymm')
   [1] coef
                   Dim
                                             pdConstruct pdFactor
                                logDet
                                                                      pdMatrix
   [7] solve
                    summary
   see '?methods' for accessing help and source code
nlme:::pdConstruct.pdSymm
   function (object, value = numeric(0), form = formula(object),
       nam = Names(object), data = sys.frame(sys.parent()), ...)
   {
       val <- NextMethod()</pre>
       if (length(val) == 0) {
           class(val) <- c("pdSymm", "pdMat")</pre>
           return(val)
       }
```

```
if (is.matrix(val)) {
           vald \leftarrow svd(val, nu = 0)
           object <- vald$v %*% (log(vald$d) * t(vald$v))</pre>
           value <- object[row(object) <= col(object)]</pre>
           attributes(value) <- attributes(val)[names(attributes(val)) !=</pre>
                "dim"l
           class(value) <- c("pdSymm", "pdMat")</pre>
           return(value)
       }
       Ncol \leftarrow round((sqrt(8 * length(val) + 1) - 1)/2)
       if (length(val) != round((Ncol * (Ncol + 1))/2)) {
           stop(gettextf("an object of length %d does not match the required parameter size",
               length(val)), domain = NA)
       class(val) <- c("pdSymm", "pdMat")</pre>
   <bytecode: 0x00000001b5ac978>
   <environment: namespace:nlme>
nlme:::pdMatrix.pdSymm
   function (object, factor = FALSE)
   {
       if (!isInitialized(object))
           stop("cannot extract matrix from an uninitialized object")
       if (factor) {
           Ncol <- Dim(object)[2]</pre>
           value <- array(pdFactor(object), c(Ncol, Ncol), attr(object,</pre>
                "Dimnames"))
           attr(value, "logDet") <- sum(log(abs(svd.d(value))))</pre>
           value
       }
       else {
           NextMethod()
       }
   <bytecode: 0x00000001b49ac58>
   <environment: namespace:nlme>
nlme:::pdFactor.pdSymm
   function (object)
   {
       Ncol \leftarrow round((-1 + sqrt(1 + 8 * length(object)))/2)
       .C(matrixLog_pd, Factor = double(Ncol * Ncol), as.integer(Ncol),
           as.double(object))$Factor
   }
   <bytecode: 0x00000001b4e1f20>
   <environment: namespace:nlme>
nlme:::solve.pdSymm
   function (a, b, ...)
       if (!isInitialized(a)) {
```

```
stop("cannot extract the inverse from an uninitialized object")
       }
       coef(a) <- -coef(a, TRUE)</pre>
   <bytecode: 0x00000001b144aa8>
   <environment: namespace:nlme>
nlme:::coef.pdSymm
   function (object, unconstrained = TRUE, ...)
       if (unconstrained || !isInitialized(object))
           NextMethod()
       else {
           val <- as.matrix(object)</pre>
           aN <- Names(object)
           aN1 <- paste("cov(", aN, sep = "")
           aN2 <- paste(aN, ")", sep = "")
           aNmat <- t(outer(aN1, aN2, paste, sep = ","))
           aNmat[row(aNmat) == col(aNmat)] <- paste("var(", aN,
               ")", sep = "")
           val <- val[row(val) <= col(val)]</pre>
           names(val) <- aNmat[row(aNmat) <= col(aNmat)]</pre>
           val
       }
   }
   <bytecode: 0x00000001a728b90>
   <environment: namespace:nlme>
```

lme

```
nlme:::lme.formula
   function (fixed, data = sys.frame(sys.parent()), random = pdSymm(eval(as.call(fixed[-2]))),
       correlation = NULL, weights = NULL, subset, method = c("REML",
            "ML"), na.action = na.fail, control = list(), contrasts = NULL,
       keep.data = TRUE)
   {
       Call <- match.call()</pre>
       miss.data <- missing(data) || !is.data.frame(data)</pre>
       controlvals <- lmeControl()</pre>
       if (!missing(control)) {
           controlvals[names(control)] <- control</pre>
       }
       fixedSigma <- controlvals$sigma > 0
       if (!inherits(fixed, "formula") || length(fixed) != 3) {
           stop("\nfixed-effects model must be a formula of the form \"resp ~ pred\"")
       }
       method <- match.arg(method)</pre>
       REML <- method == "REML"</pre>
       reSt <- reStruct(random, REML = REML, data = NULL)</pre>
       groups <- getGroupsFormula(reSt)</pre>
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if (is.null(groups)) {
    if (inherits(data, "groupedData")) {
        groups <- getGroupsFormula(data)</pre>
        namGrp <- rev(names(getGroupsFormula(data, asList = TRUE)))</pre>
        Q <- length(namGrp)</pre>
        if (length(reSt) != Q) {
             if (length(reSt) != 1) {
               stop("incompatible lengths for 'random' and grouping factors")
             randL <- vector("list", Q)</pre>
            names(randL) <- rev(namGrp)</pre>
             for (i in 1:Q) randL[[i]] <- random</pre>
            reSt <- reStruct(as.list(randL), REML = REML,
               data = NULL)
        }
        else {
             names(reSt) <- namGrp</pre>
        }
    }
    else {
        groups <- ~1
        names(reSt) <- "1"</pre>
    }
if (!is.null(correlation)) {
    add.form <- FALSE</pre>
    if (!is.null(corGrpsForm <- getGroupsFormula(correlation,</pre>
        asList = TRUE))) {
        corGrpsForm <- unlist(lapply(corGrpsForm, function(el) deparse(el[[2L]])))</pre>
        lmeGrpsForm <- unlist(lapply(splitFormula(groups),</pre>
             function(el) deparse(el[[2L]])))
        corQ <- length(corGrpsForm)</pre>
        lmeQ <- length(lmeGrpsForm)</pre>
        if (corQ <= lmeQ) {</pre>
             if (any(corGrpsForm != lmeGrpsForm[1:corQ])) {
               stop("incompatible formulas for groups in 'random' and 'correlation'")
             }
             if (corQ < lmeQ) {</pre>
               warning("cannot use smaller level of grouping for 'correlation' than for 'random'.
               add.form <- TRUE
             }
        else if (any(lmeGrpsForm != corGrpsForm[1:lmeQ])) {
             stop("incompatible formulas for groups in 'random' and 'correlation'")
        }
    }
    else {
        add.form <- TRUE
        corQ \leftarrow lmeQ \leftarrow 1
    if (add.form)
        attr(correlation, "formula") <- eval(substitute(~COV |</pre>
             GRP, list(COV = getCovariateFormula(formula(correlation))[[2L]],
             GRP = groups[[2L]]))
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}
else {
    corQ <- lmeQ <- 1
}
lmeSt <- lmeStruct(reStruct = reSt, corStruct = correlation,</pre>
    varStruct = varFunc(weights))
mfArgs <- list(formula = asOneFormula(formula(lmeSt), fixed,</pre>
    groups), data = data, na.action = na.action)
if (!missing(subset)) {
    mfArgs[["subset"]] <- asOneSidedFormula(Call[["subset"]])[[2L]]</pre>
}
mfArgs$drop.unused.levels <- TRUE
dataMix <- do.call(model.frame, mfArgs)</pre>
origOrder <- row.names(dataMix)</pre>
for (i in names(contrasts)) contrasts(dataMix[[i]]) = contrasts[[i]]
grps <- getGroups(dataMix, groups)</pre>
if (inherits(grps, "factor")) {
    ord <- order(grps)</pre>
    grps <- data.frame(grps)</pre>
    row.names(grps) <- origOrder</pre>
    names(grps) <- as.character(deparse((groups[[2L]])))</pre>
}
else {
    ord <- do.call(order, grps)</pre>
    for (i in 2:ncol(grps)) {
        grps[, i] <- as.factor(paste(as.character(grps[,</pre>
             i - 1]), as.character(grps[, i]), sep = "/"))
    }
}
if (corQ > lmeQ) {
    ord <- do.call(order, getGroups(dataMix, getGroupsFormula(correlation)))</pre>
}
grps <- grps[ord, , drop = FALSE]</pre>
dataMix <- dataMix[ord, , drop = FALSE]</pre>
revOrder <- match(origOrder, row.names(dataMix))</pre>
N <- nrow(grps)</pre>
Z <- model.matrix(reSt, dataMix)</pre>
ncols <- attr(Z, "ncols")</pre>
Names(lmeSt$reStruct) <- attr(Z, "nams")</pre>
contr <- attr(Z, "contr")</pre>
X <- model.frame(fixed, dataMix)</pre>
Terms <- attr(X, "terms")</pre>
auxContr <- lapply(X, function(el) if (inherits(el, "factor") &&
    length(levels(el)) > 1)
    contrasts(el))
contr <- c(contr, auxContr[is.na(match(names(auxContr), names(contr)))])</pre>
contr <- contr[!unlist(lapply(contr, is.null))]</pre>
X <- model.matrix(fixed, data = X)</pre>
y <- eval(fixed[[2L]], dataMix)</pre>
ncols <- c(ncols, dim(X)[2L], 1)</pre>
Q <- ncol(grps)
attr(lmeSt, "conLin") <- list(Xy = array(c(Z, X, y), c(N,
    sum(ncols)), list(row.names(dataMix), c(colnames(Z),
    colnames(X), deparse(fixed[[2L]])))), dims = MEdims(grps,
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ncols), logLik = 0, sigma = controlvals$sigma, auxSigma = 0)
tmpDims <- attr(lmeSt, "conLin")$dims</pre>
if (max(tmpDims$ZXlen[[1L]]) < tmpDims$qvec[1L]) {</pre>
    warning(gettextf("fewer observations than random effects in all level %s groups",
        Q), domain = NA)
fixDF <- getFixDF(X, grps, attr(lmeSt, "conLin")$dims$ngrps,</pre>
    terms = Terms)
lmeSt <- Initialize(lmeSt, dataMix, grps, control = controlvals)</pre>
parMap <- attr(lmeSt, "pmap")</pre>
if (length(lmeSt) == 1) {
    oldConLin <- attr(lmeSt, "conLin")</pre>
    decomp <- TRUE
    attr(lmeSt, "conLin") <- MEdecomp(attr(lmeSt, "conLin"))</pre>
}
else decomp <- FALSE
numIter <- 0
repeat {
    oldPars <- coef(lmeSt)</pre>
    optRes <- if (controlvals$opt == "nlminb") {</pre>
        control <- list(iter.max = controlvals$msMaxIter,</pre>
            eval.max = controlvals$msMaxEval, trace = controlvals$msVerbose)
        keep <- c("abs.tol", "rel.tol", "x.tol", "xf.tol",</pre>
             "step.min", "step.max", "sing.tol", "scale.init",
            "diff.g")
        control <- c(control, controlvals[names(controlvals) %in%</pre>
            keep])
        nlminb(c(coef(lmeSt)), function(lmePars) -logLik(lmeSt,
            lmePars), control = control)
    }
    else {
        reltol <- controlvals$reltol</pre>
        if (is.null(reltol))
            reltol <- 100 * .Machine$double.eps</pre>
        control <- list(trace = controlvals$msVerbose, maxit = controlvals$msMaxIter,</pre>
            reltol = if (numIter == 0) controlvals$msTol else reltol)
        keep <- c("fnscale", "parscale", "ndeps", "abstol",
            "alpha", "beta", "gamma", "REPORT", "type", "lmm",
             "factr", "pgtol", "temp", "tmax")
        control <- c(control, controlvals[names(controlvals) %in%</pre>
            keep])
        optim(c(coef(lmeSt)), function(lmePars) -logLik(lmeSt,
            lmePars), control = control, method = controlvals$optimMethod)
    }
    coef(lmeSt) <- optRes$par</pre>
    attr(lmeSt, "lmeFit") <- MEestimate(lmeSt, grps)</pre>
    if (!needUpdate(lmeSt)) {
        if (optRes$convergence) {
            msg <- gettextf("%s problem, convergence error code = %s\n message = %s",
              controlvals$opt, optRes$convergence, paste(optRes$message,
                 collapse = ""))
            if (!controlvals$returnObject)
               stop(msg, domain = NA)
            else warning(msg, domain = NA)
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}
        break
    }
    numIter <- numIter + 1L</pre>
    lmeSt <- update(lmeSt, dataMix)</pre>
    aConv <- coef(lmeSt)
    conv <- abs((oldPars - aConv)/ifelse(aConv == 0, 1, aConv))</pre>
    aConv <- NULL
    for (i in names(lmeSt)) {
        if (any(parMap[, i])) {
            aConv <- c(aConv, max(conv[parMap[, i]]))</pre>
            names(aConv)[length(aConv)] <- i</pre>
        }
    }
    if (max(aConv) <= controlvals$tolerance) {</pre>
        break
    if (numIter > controlvals$maxIter) {
        msg <- gettext("maximum number of iterations (lmeControl(maxIter)) reached without conve
        if (controlvals$returnObject) {
             warning(msg, domain = NA)
        else stop(msg, domain = NA)
    }
}
lmeFit <- attr(lmeSt, "lmeFit")</pre>
names(lmeFit$beta) <- namBeta <- colnames(X)</pre>
attr(fixDF, "varFixFact") <- varFix <- lmeFit$sigma * lmeFit$varFix
varFix <- crossprod(varFix)</pre>
dimnames(varFix) <- list(namBeta, namBeta)</pre>
Fitted <- fitted(lmeSt, level = 0:Q, conLin = if (decomp)
else attr(lmeSt, "conLin"))[revOrder, , drop = FALSE]
Resid <- y[revOrder] - Fitted</pre>
rownames(Resid) <- rownames(Fitted) <- origOrder</pre>
attr(Resid, "std") <- lmeFit$sigma/(varWeights(lmeSt)[revOrder])
grps <- grps[revOrder, , drop = FALSE]</pre>
lmeSt$reStruct <- solve(lmeSt$reStruct)</pre>
dims <- attr(lmeSt, "conLin")$dims[c("N", "Q", "qvec", "ngrps",</pre>
    "ncol")]
attr(lmeSt, "fixedSigma") <- fixedSigma</pre>
apVar <- if (controlvals$apVar) {</pre>
    lmeApVar(lmeSt, lmeFit$sigma, .relStep = controlvals[[".relStep"]],
        minAbsPar = controlvals[["minAbsParApVar"]], natural = controlvals[["natural"]])
}
else {
    "Approximate variance-covariance matrix not available"
attr(lmeSt, "conLin") <- NULL</pre>
attr(lmeSt, "lmeFit") <- NULL</pre>
grpDta <- inherits(data, "groupedData")</pre>
structure(class = "lme", list(modelStruct = lmeSt, dims = dims,
    contrasts = contr, coefficients = list(fixed = lmeFit$beta,
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random = lmeFit$b), varFix = varFix, sigma = lmeFit$sigma,
           apVar = apVar, logLik = lmeFit$logLik, numIter = if (needUpdate(lmeSt)) numIter,
           groups = grps, call = Call, terms = Terms, method = method,
           fitted = Fitted, residuals = Resid, fixDF = fixDF, na.action = attr(dataMix,
               "na.action"), data = if (keep.data && !miss.data) data),
           units = if (grpDta)
               attr(data, "units"), labels = if (grpDta)
               attr(data, "labels"))
   <bytecode: 0x000000022ca7698>
   <environment: namespace:nlme>
nlme:::reStruct
   function (object, pdClass = "pdLogChol", REML = FALSE, data = sys.frame(sys.parent()))
       if (inherits(object, "reStruct")) {
           if (!missing(REML))
               attr(object, "settings")[1] <- as.integer(REML)</pre>
           object[] <- lapply(object, function(el, data) {</pre>
               pdMat(el, data = data)
           }, data = data)
           return(object)
       }
       plen <- NULL
       if (inherits(object, "formula")) {
           if (is.null(grpForm <- getGroupsFormula(object, asList = TRUE))) {</pre>
               object <- list(object)</pre>
           }
           else {
               if (length(object) == 3) {
                    object <- eval(parse(text = paste(deparse(getResponseFormula(object)[[2]]),
                      deparse(getCovariateFormula(object)[[2]], width.cutoff = 500),
                      sep = "~")))
               }
               else {
                   object <- getCovariateFormula(object)</pre>
               object <- rep(list(object), length(grpForm))</pre>
               names(object) <- names(grpForm)</pre>
           }
       }
       else if (inherits(object, "pdMat")) {
           if (is.null(formula(object))) {
               stop("\"pdMat\" element must have a formula")
           object <- list(object)</pre>
       }
       else {
           if (data.class(object) != "list") {
               stop("'object' must be a list or a formula")
           if (is.null(names(object)) && all(unlist(lapply(object,
               function(el) {
                    inherits(el, "formula") && length(el) == 3
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})))) {
        object <- list(object)</pre>
    }
    else {
        object <- lapply(object, function(el) {</pre>
             if (inherits(el, "pdMat")) {
               if (is.null(formula(el))) {
                 stop("\"pdMat\" elements must have a formula")
               return(el)
             }
             if (inherits(el, "formula")) {
               grpForm <- getGroupsFormula(el)</pre>
               if (!is.null(grpForm)) {
                 el <- getCovariateFormula(el)</pre>
                 attr(el, "grpName") <- deparse(grpForm[[2]])</pre>
               return(el)
             }
             else {
               if (data.class(el) == "list" && all(unlist(lapply(el,
                 function(el1) {
                   inherits(el1, "formula") && length(el1) ==
                 })))) {
                 return(el)
               }
               else {
                 stop("elements in 'object' must be formulas or \"pdMat\" objects")
             }
        })
    }
    if (is.null(namObj <- names(object))) {</pre>
        namObj <- rep("", length(object))</pre>
    }
    aux <- unlist(lapply(object, function(el) {</pre>
        if (inherits(el, "formula") && !is.null(attr(el,
             "grpName"))) {
            attr(el, "grpName")
        }
        else ""
    }))
    auxNam <- namObj == ""
    if (any(auxNam)) {
        namObj[auxNam] <- aux[auxNam]</pre>
    names(object) <- namObj</pre>
object <- lapply(object, function(el, pdClass, data) {</pre>
    pdMat(el, pdClass = pdClass, data = data)
}, pdClass = pdClass, data = data)
object <- rev(object)</pre>
if (all(unlist(lapply(object, isInitialized)))) {
```

}