An Introduction to **RcppEigen**

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Abstract

The **RcppEigen** package provides access to the **Eigen** C++ template library from R. **Rcpp** classes and instantiations of the C++ templated functions as and wrap from **Rcpp** provide the "glue" for passing objects from R to C++ and back.

1 Introduction

As stated in the **Rcpp** vignette, "Extending **Rcpp**"

Rcpp facilitates data interchange between R and C++ through the templated functions Rcpp::as (for conversion of objects from R to C++) and Rcpp::wrap (for conversion from C++ to R).

The RcppEigen package provides the header files composing the Eigen C++ template library and implementations of Rcpp::as and Rcpp::wrap for the C++ classes defined in Eigen.

The **Eigen** classes themselves provide high-performance, versatile and comprehensive representations of dense and sparse matrices and vectors, as well as decompositions and other functions to be applied to these objects. In the next section we introduce some of these classes and show how to interface to them from R.

2 Eigen classes

Eigen (http://eigen.tuxfamily.org) is a C++ template library providing classes for many forms of matrices, vectors, arrays and decompositions. These classes are flexible and comprehensive allowing for both high performance and well structured code representing high-level operations. C++ code based on Eigen is often more like R code, working on the "whole object", than compiled code in other languages where operations often must be coded in loops.

As in many C++ template libraries using template meta-programming (Abrahams and Gurtovoy, 2004), the templates themselves can be very complicated. However, **Eigen** provides typedef's for common classes that correspond to R matrices and vectors, as shown in Table 1. We will use these typedef's throughout this document.

The C++ classes shown in Table 1 are in the Eigen namespace, which means that they must be written as Eigen::MatrixXd. However, if we preface our use of these class names with a declaration like

using Eigen::MatrixXd;

we can use these names without the qualifier. I prefer this approach.

Table 1: Correspondence between R matrix and vector types and classes in the Eigen namespace.

R object type	Eigen class typedef		
numeric matrix	MatrixXd		
integer matrix	MatrixXi		
complex matrix	MatrixXcd		
numeric vector	VectorXd		
integer vector	VectorXi		
complex vector	VectorXcd		
Matrix::dgCMatrix	SparseMatrix <double></double>		

2.1 Mapped matrices in Eigen

Storage for the contents of matrices from the classes shown in Table 1 is allocated and controlled by the class constructors and destructors. Creating an instance of such a class from an R object involves copying its contents. An alternative is to have the contents of the R matrix or vector mapped to the contents of the object from the Eigen class. For dense matrices we use the Eigen templated class Map. For sparse matrices we use the Eigen templated class MappedSparseMatrix.

We must, of course, be careful not to modify the contents of the R object in the C++ code. A recommended practice is always to declare mapped objects as const.

2.2 Arrays in Eigen

For matrix and vector classes **Eigen** overloads the '*' operator to indicate matrix multiplication. Occasionally we want component-wise operations instead of matrix operations. The **Array** templated classes are used in **Eigen** for component-wise operations. Most often we use the **array** method for Matrix or Vector objects to create the array.

2.3 Structured matrices in Eigen

There are **Eigen** classes for matrices with special structure such as symmetric matrices, triangular matrices and banded matrices. For dense matrices, these special structures are described as "views", meaning that the full dense matrix is stored but only part of the matrix is used in operations. For a symmetric matrix we need to specify whether the lower triangle or the upper triangle is to be used as the contents, with the other triangle defined by the implicit symmetry.

3 Some simple examples

C++ functions to perform simple operations on matrices or vectors can follow a pattern of:

- 1. Map the R objects passed as arguments into Eigen objects.
- 2. Create the result.
- 3. Return Rcpp::wrap applied to the result.

An idiom for the first step is

```
using Eigen::Map;
using Eigen::MatrixXd;
using Rcpp::as;
const Map<MatrixXd> A(as<Map<MatrixXd> >(AA));
```

where AA is the name of the R object (called an SEXP in C and C++) passed to the C++ function.

The cxxfunction from the inline package for R and its RcppEigen plugin provide a convenient way of developing and debugging the C++ code. For actual production code we generally incorporate the C++ source code files in a package and include the line LinkingTo: Rcpp, RcppEigen in the package's DESCRIPTION file.

The cxxfunction with the "Rcpp" or "RcppEigen" plugins has the as and wrap functions already defined as Rcpp::as and Rcpp::wrap. In the examples below we will omit these declarations. Do remember that you will need them in C++ source code for a package.

The first few examples are simply for illustration as the operations shown could be more effectively performed directly in R. We do compare the results from **Eigen** to those from the direct R results.

3.1 Transpose of an integer matrix

We create a simple matrix of integers

```
> (A <- matrix(1:6, ncol=2))
```

```
[,1] [,2]
[1,] 1 4
[2,] 2 5
[3,] 3 6
```

```
> str(A)
int [1:3, 1:2] 1 2 3 4 5 6
```

and use the transpose method for the Eigen::MatrixXi class to return its transpose.

For numeric or integer matrices the adjoint method is equivalent to the transpose method. For complex matrices, the adjoint is the conjugate of the transpose. In keeping with the conventions in the Eigen documentation we prefer the name adjoint with numeric or integer matrices.

3.2 Products and cross-products

As mentioned in Sec. 2.2, the '*' operator performs matrix multiplication on Matrix or Vector objects.

Notice that the create method for the Rcpp class List implicitly applies Rcpp::wrap to its arguments.

3.3 Crossproduct of a single matrix

As shown in the last example, the R function crossprod calculates the product of the transpose of its first argument with its second argument. The single argument form, crossprod(X), evaluates X'X. We could, of course, calculate this product as

```
> t(X) %*% X
```

but crossprod(X) is roughly twice as fast because the result is known to be symmetric and only half the result needs to be calculated.. The function tcrossprod evaluates crossprod(t(X)) without actually forming the transpose.

To express these calculations in Eigen we create a SelfAdjointView, which is a dense matrix of which only one triangle is used, the other triangle being inferred from the symmetry. ("self-adjoint" is equivalent to symmetric when applied to non-complex matrices.)

The **Eigen** class name is **SelfAdjointView**. The method for general matrices that produces such a view is called **selfadjointView**. Both require specification of either the **Lower** or **Upper** triangle.

For triangular matrices the class is TriangularView and the method is triangularView. The triangle can be specified as Lower, UnitLower, StrictlyLower, Upper, UnitUpper or StrictlyUpper.

For self-adjoint views the rankUpdate method adds a scalar multiple of AA' to the current symmetric matrix. The scalar multiple defaults to 1.

To some, the expressions to construct AtA and AAt in that code fragment are compact and elegant. To others they are hopelessly confusing. If you find yourself in the latter group, you just need to read the expression left to right. So, for example, we construct AAt by creating a general integer matrix of size $m \times m$ (where A is $m \times n$), ensure that all its elements are zero, regard it as a self-adjoint (i.e. symmetric) matrix using the elements in the lower triangle, then add AA' to it and convert back to a general matrix form (i.e.the strict lower triangle is copied into the strict upper triangle).

For these products we could use either the lower triangle or the upper triangle as the result will be symmetrized before it is returned.

3.4 Cholesky decomposition of the crossprod

The Cholesky decomposition of the positive-definite, symmetric matrix, \boldsymbol{A} , can be written in several forms. Numerical analysts define the "LLt" form as the lower triangular matrix, \boldsymbol{L} , such that $\boldsymbol{A} = \boldsymbol{L}\boldsymbol{L}'$ and the "LDLt" form as a unit lower triangular matrix \boldsymbol{L} and a diagonal matrix \boldsymbol{D} with positive diagonal elements such that $\boldsymbol{A} = \boldsymbol{L}\boldsymbol{D}\boldsymbol{L}'$. Statisticians often write the decomposition as $\boldsymbol{A} = \boldsymbol{R}'\boldsymbol{R}$ where \boldsymbol{R} is an upper triangular matrix. Of course, this \boldsymbol{R} is simply the transpose of \boldsymbol{L} from the "LLt" form.

The templated **Eigen** classes for the LLt and LDLt forms are called LLT and LDLT. In general we would preserve the objects from these classes so that we could use them for solutions of linear systems. For illustration we simply return the matrix L from the "LLt" form.

Because the Cholesky decomposition involves taking square roots we switch to numeric matrices.

```
> storage.mode(A) <- "double"</pre>
```

3.5 Determinant of the cross-product matrix

The "D-optimal" criterion for experimental design chooses the design that maximizes the determinant, |X'X|, for the $n \times p$ model matrix (or Jacobian matrix), X. The determinant, |L|, of the $p \times p$ lower Cholesky factor L, defined so that LL' = X'X, is the product of its diagonal elements. (This is true for any triangular matrix.) By the properties of determinants,

$$|X'X| = |LL'| = |L||L'| = |L|^2$$

Alternatively, if we use the "LDLt" decomposition, LDL' = X'X where L is unit lower triangular and D is diagonal then |X'X| is the product of the diagonal elements of D. Because we know that the diagonals of D must be non-negative, we often evaluate the logarithm of the determinant as the sum of the logarithms of the diagonal elements of D.

```
using Eigen::Lower;
using Eigen::Map;
using Eigen::MatrixXd;
using Eigen::VectorXd;
const Map<MatrixXd>
                      A(as<Map<MatrixXd> >(AA));
const int
                      n(A.cols());
const MatrixXd
                    AtA(MatrixXd(n, n).setZero().
                        selfadjointView<Lower>().rankUpdate(A.adjoint()));
const MatrixXd
                   Lmat(AtA.llt().matrixL());
const double
                   detL(Lmat.diagonal().prod());
const VectorXd
                   Dvec(AtA.ldlt().vectorD());
return List::create(_["d1"] = detL * detL,
                    _["d2"] = Dvec.prod(),
                    _["ld"] = Dvec.array().log().sum());
```

Note the use of the array() method in the calculation of the log-determinant. Because the log() method applies to arrays, not to vectors or matrices, we must create an array from Dvec before applying the log() method.

4 Least squares solutions

A common operation in statistical computing is calculating a least squares solution, $\hat{\beta}$, defined as

$$\widehat{oldsymbol{eta}} = rg \min_{eta} \|oldsymbol{y} - oldsymbol{X} oldsymbol{eta}\|^2$$

where the model matrix, X, is $n \times p$ ($n \ge p$) and y is an n-dimensional response vector. There are several ways based on matrix decompositions, to determine such a solution. We have already seen two forms of the Cholesky decomposition: "LLt" and "LDLt", that can be used to solve for $\hat{\beta}$. Other decompositions that can be used are the QR decomposition, with or without column pivoting, the singular value decomposition and the eigendecomposition of a symmetric matrix.

Determining a least squares solution is relatively straightforward. However, in statistical computing we often require a bit more information, such as the standard errors of the coefficient estimates. Calculating these involves evaluating the diagonal elements of $(X'X)^{-1}$ and the residual sum of squares, $\|y - X\widehat{\beta}\|^2$.

4.1 Least squares using the "LLt" Cholesky

```
using Eigen::LLT;
using Eigen::Lower;
using Eigen::Map;
using Eigen::MatrixXd;
using Eigen::VectorXd;
                      X(as<Map<MatrixXd> >(XX));
const Map<MatrixXd>
const Map<VectorXd>
                      y(as<Map<VectorXd> >(yy));
const int
                      n(X.rows()), p(X.cols());
const LLT<MatrixXd> llt(MatrixXd(p, p).setZero().
                        selfadjointView<Lower>().rankUpdate(X.adjoint()));
const VectorXd betahat(llt.solve(X.adjoint() * y));
const VectorXd fitted(X * betahat);
                resid(y - fitted);
const VectorXd
                     df(n - p);
const int
const double
                      s(resid.norm() / std::sqrt(double(df)));
const VectorXd
                     se(s * llt.matrixL().solve(MatrixXd::Identity(p, p)).colwise().norm());
           List::create(_["coefficients"]
                                             = betahat,
                        _["fitted.values"]
                                            = fitted,
                        ["residuals"]
                                            = resid,
                        _["s"]
                                            = s,
                        _["df.residual"]
                                            = df.
                        ["rank"]
                                            = p,
                        _["Std. Error"]
                                             = se);
```

```
List of 7
 $ coefficients : num [1:2] -2.35 2.2
 $ fitted.values: num [1:31] 2.3 2.38 2.43 2.82 2.86 ...
               : num [1:31] 0.0298 -0.0483 -0.1087 -0.0223 0.0727 ...
                : num 0.115
 $ df.residual : int 29
 $ rank
                : int 2
               : num [1:2] 0.2307 0.0898
 $ Std. Error
> str(lmFit <- with(trees, lm.fit(cbind(1, log(Girth)), log(Volume))))
List of 8
 $ coefficients : Named num [1:2] -2.35 2.2
  ..- attr(*, "names")= chr [1:2] "x1" "x2"
 $ residuals
              : num [1:31] 0.0298 -0.0483 -0.1087 -0.0223 0.0727 ...
 $ effects
                : Named num [1:31] -18.2218 2.8152 -0.1029 -0.0223 0.0721 ...
  ..- attr(*, "names")= chr [1:31] "x1" "x2" "" "" ...
               : int 2
 $ fitted.values: num [1:31] 2.3 2.38 2.43 2.82 2.86 ...
 $ assign
               : NULL
 $ qr
                :List of 5
  ..$ qr
          : num [1:31, 1:2] -5.57 0.18 0.18 0.18 0.18 ...
  ..$ qraux: num [1:2] 1.18 1.26
  ..$ pivot: int [1:2] 1 2
  ..$ tol : num 1e-07
  ..$ rank : int 2
  ..- attr(*, "class")= chr "qr"
 $ df.residual : int 29
> for (nm in c("coefficients", "residuals", "fitted.values", "rank"))
      stopifnot(all.equal(lltFit[[nm]], unname(lmFit[[nm]])))
> stopifnot(all.equal(lltFit[["Std. Error"]],
                      unname(coef(summary(lm(log(Volume) ~ log(Girth), trees)))[,2])))
```

There are several aspects of the C++ code worth mentioning. The solve method for the LLT object evaluates, in this case, $(X'X)^{-1}X'y$ but without actually evaluating the inverse. The calculation of the residuals, $y - \hat{y}$, can be written, as in R, as y - fitted. (But note that Eigen classes do not have a "recycling rule as in R. That is, the two vector operands must have the same length.) The norm method evaluates the square root of the sum of squares of the elements of a vector. Although we don't explicitly evaluate $(X'X)^{-1}$ we do evaluate L^{-1} to obtain the standard errors. Note also the use of the colwise method in the evaluation of the standard errors.

In the descriptions of other methods for solving least squares problems, much of the code parallels that shown here and we omit this. We show only the evaluation of the coefficients, the rank and the standard errors. Actually, we only calculate the standard errors up to the scalar multiple of s, the residual standard error, in these code fragments. The calculation of the residuals and s and the scaling of the coefficient standard errors is the same for all methods. (See the file fastLm.cpp of the RcppEigen source package for details.)

4.2 Least squares using the unpivoted QR decomposition

A QR decomposition has the form

$$X = QR = Q_1R_1$$

where Q is an $n \times n$ orthogonal matrix, which means that $Q'Q = QQ' = I_n$, and R is $n \times p$ and zero below the main diagonal. The $n \times p$ matrix Q_1 is the first p columns of Q and the $p \times p$ upper triangular matrix R_1 is the top p rows of R. There are three Eigen classes for the QR decomposition: HouseholderQR provides the basic QR decomposition using Householder transformations, ColPivHouseholderQR incorporates column pivots and FullPivHouseholderQR incorporates both row and column pivots.

For the unpivoted QR decomposition the code is of the form

We see that this code is very close to the code for the "LLt" case. In fact, if we had extracted the upper triangular factor (the matrixU method) from the LLT object, the code would be nearly identical.

4.3 Handling the rank-deficient case

One important consideration when determining least squares solutions is whether $\operatorname{rank}(X)$ is p, a situation we describe by saying that X has "full column rank". When X does not have full column rank we say it is "rank deficient".

Although the theoretical rank of a matrix is well-defined, its evaluation in practice is not. At best we determine a computational rank according to some tolerance. We refer to decompositions that allow us to estimate the rank of the matrix in this way as "rank-revealing".

Because the model.matrix function in R does a considerable amount of symbolic analysis behind the scenes, we usually end up with full-rank model matrices. The common cases of rank-deficiency, such as incorporating both a constant term and a full set of indicators columns for the levels of a factor, are eliminated. Other, more subtle, situations will not be detected at this stage, however. A simple example occurs when there is a "missing cell" in a two-way layout and the interaction of the two factors is included in the model.

```
> dd <- data.frame(f1 = gl(4, 6, labels = LETTERS[1:4]),
                   f2 = g1(3, 2, labels = letters[1:3]))[-(7:8), ]
> xtabs(~ f2 + f1, dd)
                                           # one missing cell
  f1
f2 ABCD
 a 2 0 2 2
 b 2 2 2 2
 c 2 2 2 2
> mm <- model.matrix(~ f1 * f2, dd)
                    # large condition number, indicating rank deficiency
> kappa(mm)
[1] 4.309225e+16
> rcond(mm)
                    # alternative evaluation, the reciprocal condition number
[1] 2.320603e-17
> (c(rank=qr(mm)$rank, p=ncol(mm))) # rank as computed in R's qr function
rank
       12
 11
> set.seed(1)
> dd\$y <- mm %*% seq_len(ncol(mm)) + rnorm(nrow(mm), sd = 0.1)
                           # lm detects the rank deficiency
> fm1 <- lm(y ~f1 * f2, dd)
> writeLines(capture.output(print(summary(fm1), signif.stars=FALSE))[9:22])
Coefficients: (1 not defined because of singularities)
            Estimate Std. Error t value Pr(>|t|)
(Intercept)
           0.97786
                        0.05816
                                  16.81 3.41e-09
f1B
            12.03807
                        0.08226
                                 146.35 < 2e-16
                                  37.90 5.22e-13
f1C
             3.11722
                        0.08226
f<sub>1D</sub>
             4.06852
                        0.08226
                                  49.46 2.83e-14
             5.06012
                        0.08226
                                  61.52 2.59e-15
f2b
```

```
f2c
             5.99759
                        0.08226
                                  72.91 4.01e-16
f1B:f2b
            -3.01476
                        0.11633 -25.92 3.27e-11
             7.70300
                        0.11633
f1C:f2b
                                  66.22 1.16e-15
f1D:f2b
             8.96425
                        0.11633
                                  77.06 < 2e-16
f1B:f2c
                  NA
                             NA
                                     NA
f1C:f2c
            10.96133
                        0.11633
                                  94.23
                                        < 2e-16
            12.04108
                        0.11633
                                103.51
f1D:f2c
                                         < 2e-16
```

The \mbox{lm} function for fitting linear models in R uses a rank-revealing form of the QR decomposition. When the model matrix is determined to be rank deficient, according to the threshold used in R's QR decomposition, the model matrix is reduced to rank X columns by pivoting selected columns (those that are apparently linearly dependent on columns to their left) to the right hand side of the matrix. A solution for this reduced model matrix is determined and the coefficients and standard errors for the redundant columns are flagged as missing.

An alternative approach is to evaluate the "pseudo-inverse" of X from the singular value decomposition (SVD) of X or the eigendecomposition of X'X. The SVD is of the form

$$X = UDV' = U_1D_1V'$$

where U is an orthogonal $n \times n$ matrix and U_1 is its leftmost p columns, D is $n \times p$ and zero off the main diagonal so that D_1 is a $p \times p$ diagonal matrix with non-decreasing positive diagonal elements, and V is a $p \times p$ orthogonal matrix. The pseudo-inverse of D_1 , written D_1^+ is a $p \times p$ diagonal matrix whose first r = rank(X) diagonal elements are the inverses of the corresponding diagonal elements of D_1 and whose last p - r diagonal elements are zero.

The tolerance for determining if an element of the diagonal of D is considered to be (effectively) zero is a multiple of the largest singular value (i.e. the (1,1) element of D).

We define a utility function, Dplus to return the pseudo-inverse as a diagonal matrix, given the singular values (the diagonal of D) and the apparent rank.

```
using Eigen::DiagonalMatrix;
using Eigen::Dynamic;
inline DiagonalMatrix<double, Dynamic> Dplus(const ArrayXd& D, int r) {
    VectorXd    Di(VectorXd::Constant(D.size(), 0.));
    Di.head(r) = D.head(r).inverse();
    return DiagonalMatrix<double, Dynamic>(Di);
}
```

4.4 Least squares with the SVD

With these definitions the code for least squares using the singular value decomposition can be written

In the rank-deficient case this code will produce a complete set of coefficients and their standard errors. It is up to the user to note that the rank is less than p, the number of columns in X and hence that the estimated coefficients are just one of an infinite number of coefficient vectors that produce the same fitted values. It happens that this solution is the minimum norm solution.

The interpretation of the standard errors from this code is also problematic when X is rank-deficient.

4.5 Least squares with the eigendecomposition

The eigendecomposition of X'X is defined as

$$X'X = V\Lambda V'$$

where V, the matrix of eigenvectors, is a $p \times p$ orthogonal matrix and Λ is a $p \times p$ diagonal matrix with non-increasing, non-negative diagonal elements, called the eigenvalues of X'X. When the eigenvalues are distinct this V is the same as that in the SVD. Also the eigenvalues of X'X are the squares of the singular values of X.

With these definitions we can adapt much of the code from the SVD method for the eigendecomposition.

4.6 Least squares with the column-pivoted QR decomposition

The column-pivoted QR decomposition provides results similar to those from R in both the full-rank and the rank-deficient cases. The decomposition is of the form

$$XP = QR = Q_1R_1$$

where, as before, Q is $n \times n$ and orthogonal and R is $n \times p$ and upper triangular. The $p \times p$ matrix P is a permutation matrix. That is, its columns are a permutation of the columns of I_p . It serves to reorder the columns of X so that the diagonal elements of R are non-increasing in magnitude.

An instance of the class Eigen::ColPivHouseholderQR has a rank method returning the computational rank of the matrix. When X is of full rank we can use essentially the same code as in the unpivoted decomposition except that we must reorder the standard errors. When X is rank-deficient we evaluate the coefficients and standard errors for the leading r columns of XP only.

In the rank-deficient case the straightforward calculation of the fitted values, as $X\widehat{\beta}$ cannot be used. We could do some complicated rearrangement of the columns of X and the coefficient estimates but it is conceptually (and computationally) easier to employ the relationship

$$\widehat{m{y}} = m{Q}_1 m{Q}_1' m{y} = m{Q} egin{bmatrix} m{I}_r & m{0} & m{0} \end{bmatrix} m{Q}' m{y}$$

The vector Q'y is called the "effects" vector in R.

```
using Eigen::ColPivHouseholderQR;
typedef ColPivHouseholderQR<MatrixXd>::PermutationType Permutation;
const ColPivHouseholderQR<MatrixXd> PQR(X);
const Permutation
                                     Pmat(PQR.colsPermutation());
const int
                                        r(PQR.rank());
VectorXd
                               betahat, fitted, se;
if (r == X.cols()) { // full rank case
    betahat = PQR.solve(y);
    fitted = X * m_coef;
             = Pmat * PQR.matrixQR().topRows(m_p).triangularView<Upper>().
        solve(MatrixXd::Identity(m_p, m_p)).rowwise().norm();
} else {
    MatrixXd
                                  Rinv(PQR.matrixQR().topLeftCorner(r, r).
           triangularView<Upper>().
           solve(MatrixXd::Identity(r, r)));
                               effects(PQR.householderQ().adjoint() * y);
    VectorXd
    betahat.head(r)
                                        = Rinv * effects.head(r);
    betahat
                                        = Pmat * betahat;
   // create fitted values from effects
   // (cannot use X * m_coef when X is rank-deficient)
    effects.tail(X.rows() - r).setZero();
    fitted
                                        = PQR.householderQ() * effects;
    se.head(r)
                                        = Rinv.rowwise().norm();
    se
                                        = Pmat * se:
}
```

Just to check that this does indeed provide the desired answer

```
> print(summary(fmPQR <- fastLm(y ~ f1 * f2, dd)), signif.stars=FALSE)</pre>
Call:
fastLm.formula(formula = y ~ f1 * f2, data = dd)
           Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.977859 0.058165 16.812 3.413e-09
         12.038068 0.082258 146.346 < 2.2e-16
           3.117222 0.082258 37.896 5.221e-13
4.068523 0.082258 49.461 2.833e-14
f1C
f1D
f2b
           5.060123
                     0.082258 61.516 2.593e-15
                    0.082258 72.912 4.015e-16
f2c
           5.997592
           -3.014763 0.116330 -25.916 3.266e-11
f1B:f2b
           7.702999 0.116330 66.217 1.156e-15
f1C:f2b
          8.964251 0.116330 77.059 < 2.2e-16
f1D:f2b
f1B:f2c
                                  NA
                                           NΑ
                NA
                          NA
          f1C:f2c
         f1D:f2c
Residual standard error: 0.2868 on 11 degrees of freedom
Multiple R-squared: 0.9999,
                                Adjusted R-squared: 0.9999
> all.equal(coef(fm1), coef(fmPQR))
[1] TRUE
> all.equal(unname(fitted(fm1)), fitted(fmPQR))
[1] TRUE
> all.equal(unname(residuals(fm1)), residuals(fmPQR))
[1] TRUE
```

The rank-revealing SVD method produces the same fitted values but not the same coefficients.

```
> print(summary(fmSVD <- fastLm(y ~ f1 * f2, dd, method=4L)), signif.stars=FALSE)
Call:
fastLm.formula(formula = y ~ f1 * f2, data = dd, method = 4L)
             Estimate Std. Error t value Pr(>|t|)
(Intercept)
             0.977859
                        0.058165 16.812 3.413e-09
f1B
             7.020458
                        0.038777 181.049 < 2.2e-16
f1C
             3.117222
                        0.082258 37.896 5.221e-13
                        0.082258
f1D
             4.068523
                                  49.461 2.833e-14
f2h
             5.060123
                        0.082258
                                  61.516 2.593e-15
             5.997592
                        0.082258
                                  72.912 4.015e-16
f2c
f1B:f2b
             2.002847
                        0.061311
                                  32.667 2.638e-12
f1C:f2b
             7.702999
                        0.116330
                                  66.217 1.156e-15
f1D:f2b
             8.964251
                        0.116330
                                  77.059 < 2.2e-16
f1B:f2c
             5.017610
                        0.061311
                                  81.838 < 2.2e-16
f1C:f2c
            10.961326
                        0.116330 94.226 < 2.2e-16
f1D:f2c
            12.041081
                        0.116330 103.508 < 2.2e-16
Residual standard error: 0.2868 on 11 degrees of freedom
Multiple R-squared: 0.9999,
                                   Adjusted R-squared: 0.9999
> all.equal(coef(fm1), coef(fmSVD))
[1] "'is.NA' value mismatch: O in current 1 in target"
> all.equal(unname(fitted(fm1)), fitted(fmSVD))
[1] TRUE
> all.equal(unname(residuals(fm1)), residuals(fmSVD))
```

For this example the method based on the eigendecomposition gets the rank wrong

4.7 Comparative speed

In the RcppEigen package the R function to fit linear models using the methods described above is called fastLm. The natural question to ask is, "Is it indeed fast to use these methods based on Eigen?". We have provided a comparative benchmarking of these methods, plus the default method using R's lm function and the fastLm implementations in the RcppArmadillo and RcppGSL packages, if they are installed. The benchmark is a file called lmBenchmark.R and it uses the rbenchmark package.

It can be run as

```
> source(system.file("examples", "lmBenchmark.R", package="RcppEigen"))
```

Results will vary according to the speed of the processor and the number of cores and the implementation of the BLAS (Basic Linear Algebra Subroutines) that is being used. (**Eigen** methods do not use the BLAS but the other methods do.)

Results obtained on a desktop computer, circa 2010, are shown in Table 2

These results indicate that methods based on forming and decomposing X'X, (i.e. LDLt, LLt and SymmEig) are considerably faster than the others. The SymmEig method, using a rank-revealing decomposition, would be preferred, although the LDLt method could probably be modified to be rank-revealing. Do bear in mind that the dimensions of the problem will influence the comparative results. Because there are 100,000 rows in X methods that decompose the whole X matrix (all the methods except those named above) will be at a disadvantage.

The pivoted QR method is 1.6 times faster than R's lm.fit on this test and provides nearly the same information as lm.fit. Methods based on the singular value decomposition (SVD and GSL) are much slower but, as mentioned above, this is caused in part by \boldsymbol{X} having many more rows than columns. The GSL method from the GNU Scientific Library uses an older algorithm for the SVD and is clearly out of contention.

An SVD method using the Lapack SVD subroutine, dgesv, may be faster than the native Eigen implementation of the SVD, which is not a particularly fast method.

Table 2: lmBenchmark results on a desktop computer for the default size, $100,000 \times 40$, full-rank model matrix running 20 repetitions for each method. Times (Elapsed, User and Sys) are in seconds. The BLAS in use is a single-threaded version of Atlas (Automatically Tuned Linear Algebra System).

Method	Relative	Elapsed	User	Sys
LLt	1.000000	1.227	1.228	0.000
LDLt	1.037490	1.273	1.272	0.000
SymmEig	2.895681	3.553	2.972	0.572
QR	7.828036	9.605	8.968	0.620
PivQR	7.953545	9.759	9.120	0.624
arma	8.383048	10.286	10.277	0.000
$_{ m lm.fit}$	13.782396	16.911	15.521	1.368
SVD	54.829666	67.276	66.321	0.912
GSL	157.531377	193.291	192.568	0.640

5 Delayed evaluation

A form of delayed evaluation is used in **Eigen**. That is, many operators and methods do not force the evaluation of the object but instead return an "expression object" that is evaluated when needed. As an example, even though we write the X'X evaluation using <code>.rankUpdate(X.adjoint())</code> the <code>X.adjoint()</code> part is not evaluated immediately. The <code>.rankUpdate</code> method detects that it has been passed a matrix that is to be used in its transposed form and evaluates the update by taking inner products of columns of X instead of rows of X'.

Occasionally the method for Rcpp::wrap will not force an evaluation when it should. This is at least what Bill Venables calls an "infelicity" in the code, if not an outright bug. In the first example of the transpose of an integer matrix we assigned the transpose as a MatrixXi before returning it with wrap. The assignment forces the evaluation. If we skip this step we get an answer with the correct shape but the wrong contents.

```
using Eigen::Map;
using Eigen::MatrixXi;
const Map<MatrixXi> A(as<Map<MatrixXi> >(AA));
return wrap(A.transpose());
```

[1] "Mean relative difference: 0.4285714"

Another recommended practice is to assign objects before wrapping them for return to R.

6 Sparse matrices

Eigen provides sparse matrix classes. An R object of class dgCMatrix (from the Matrix package) can be mapped as shown below.

A sparse Cholesky decomposition is provided in **Eigen** as the SimplicialCholesky class. There are also linkages to the **CHOLMOD** code from the **Matrix** package. At present, both of these are regarded as experimental.

References

David Abrahams and Aleksey Gurtovoy. C++ Template Metaprogramming: Concepts, Tools and Techniques from Boost and Beyond. Addison-Wesley, Boston, 2004.