# LAPACK Working Note 117 A Fortran 90 Interface for LAPACK: LAPACK90, version 1.0

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#### Abstract

The purpose of this report is to discuss the design of a Fortran 90 interface to LAPACK. Our emphasis at this stage is on the design of an improved user-interface to the package, taking advantage of the considerable simplifications which Fortran 90 allows. The proposed design makes use of assumed-shape arrays, optional arguments, and generic interfaces. The Fortran 90 interface can be implemented initially by writing Fortran 90 jackets to call the existing Fortran 77 code, and can persist unchanged even if the underlying Fortran 77 LAPACK code is rewritten to take advantage of the new features of Fortran 90. We aim to maintain a comparable level of performance as with the Fortran 77 code. In this paper we implement interfaces to the subset of LAPACK routines for solving systems of linear equations AX = B with a general matrix A, and for symmetric and Hermitian eigenproblems.

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# 1 Introduction

This paper is a follow on paper to LAPACK WN 101 [2] which proposed an interface to some of the LAPACK [1] linear equation routines. Following comments to that paper we have added some additional functionality to the interface. In this paper we also consider the symmetric eigenproblem. We welcome comments on the proposal given here. Our emphasis at this stage is on the design of an improved user-interface to the package, taking advantage of the considerable simplifications which Fortran 90 allows (see [3]).

The Fortran 90 interface can be implemented initially by writing Fortran 90 jackets to call the existing Fortran 77 code, and can persist unchanged even if the underlying Fortran 77 LAPACK code is rewritten to take advantage of the new features of Fortran 90.

Although we would like to maintain a comparable level of performance to the Fortran 77 LAPACK code, due to the immaturity of many of the current Fortran 90 compilers, we cannot usually at present achieve this same level of performance. We reiterate that our goal is to provide a true Fortran 90 interface to LAPACK. We are aware of certain modifications to the interface which could enhance performance, but these modifications complicate the interface by requiring Fortran 77-ish constructs. If performance is the main focus behind the user's application, we recommend that the user call the Fortran 77 interface directly.

For convenience we use the name "LAPACK 77" to denote the existing Fortran 77 package, and "LAPACK 90" to denote the new Fortran 90 interface which we are proposing.

# 2 LAPACK 77 and Fortran 90 Compilers

# 2.1 Linking LAPACK 77 to Fortran 90 programs

LAPACK 77 can be called from Fortran 90 programs in its present form — with some qualifications. The qualifications arise only because LAPACK 77 is not written entirely in *standard* Fortran 77; the exceptions are the use of the COMPLEX\*16 data type and related intrinsic functions, as listed in Section 6.1 of [1]; these facilities are provided as extensions to the standard language by many Fortran 77 and Fortran 90 compilers. Equivalent facilities are provided in standard Fortran 90, using the parameterized form of the COMPLEX data type (see below).

To link LAPACK 77 to a Fortran 90 program (which must of course be compiled by a Fortran 90 compiler), one of the following approaches will be necessary, depending on the compilers available.

1. Link the Fortran 90 program to an existing LAPACK 77 library, compiled by a Fortran 77 compiler. This approach can only work if the compilers

have been designed to allow cross-linking.

- If such cross-linking is not possible, recompile and archive the LAPACK 77 library with the Fortran 90 compiler, provided that the compiler accepts COMPLEX\*16 and related intrinsics as extensions.
- 3. If these extensions are not accepted and the user requires this data type, the LAPACK 77 code must be rewritten in standard Fortran 90 (see below).

Some conversions needed to use the double precision complex data type in standard Fortran 90 code from LAPACK 77 are:

One further obstacle may remain: it is possible that if LAPACK 77 has been recompiled with a Fortran 90 compiler, it may not link correctly to an optimized assembly-language BLAS library that has been designed to interface with Fortran 77. Until this is rectified by the vendor of the BLAS library, Fortran 77 code for the BLAS must be used.

# 2.2 Interface blocks for LAPACK 77

Fortran 90 allows one immediate extra benefit to be provided to Fortran 90 users of LAPACK 77, without making any further changes to the existing code: that is a module of explicit interfaces for the routines. If this module is accessed by a USE statement in any program unit which makes calls to LAPACK routines, then those calls can be checked by the compiler for errors in the numbers or types of arguments.

The module can be constructed by extracting the necessary specification statements from the Fortran 77 code with a little modification, as illustrated by the following example containing an interface for the single routine SSYEV:

```
MODULE LA_SF77MOD

INTERFACE

SUBROUTINE SSYEV( JOBZ, UPLO, N, A, LDA, W, & WORK, LWORK, INFO )

USE LA_PRECISION, ONLY: WP => SP

CHARACTER(LEN=1), INTENT(IN) :: JOBZ, UPLO

INTEGER, INTENT(IN) :: LDA, LWORK, N

INTEGER, INTENT(OUT) :: INFO

REAL(WP), INTENT(INOUT) :: A(LDA,*)

REAL(WP), INTENT(OUT) :: W(*), WORK(*)

END SUBROUTINE SSYEV

...

END INTERFACE

...

END MODULE LA SF77MOD
```

A module containing interfaces for all of the routines in LAPACK 77 will be required; here we provide one module per precision, for example LA\_SF77MOD, LA\_DF77MOD, LA\_CF77MOD, and LA\_ZF77MOD, and one module for auxiliary routines, LA\_AUXMOD.

# 3 Proposed Design of the LAPACK 90 interface

In the design of a Fortran 90 interface to LAPACK, we propose to take advantage of the features of the language listed below.

1. **Assumed-shape arrays:** All array arguments to LAPACK 90 routines will be assumed-shape arrays. Arguments to specify problem dimensions or array dimensions will not be required.

This implies that the actual arguments supplied to LAPACK routines must have the exact shape required by the problem. The most convenient ways to achieve this are:

• using allocatable arrays, for example:

```
REAL, ALLOCATABLE :: A(:,:), B(:)

. . .

ALLOCATE( A(N,N), B(N) )

. . .

CALL LA_GESV( A, B )
```

• passing array sections, for example:

```
REAL :: A(NMAX,NMAX), B(NMAX)
. . .
CALL LA_GESV( A(:N,:N), B(:N) )
```

Zero dimensions (empty arrays) will be allowed.

There are some grounds for concern about the effect of Fortran 77 assumedsize arrays on performance because compilers cannot assume that their storage is contiguous. The effect on performance will of course depend on the compiler, and may diminish in time as compilers become more effective in optimizing compiled code. See section 7.

- 2. Automatic allocation of work arrays: Workspace arguments and arguments to specify their dimensions will not be needed. In simple cases, automatic arrays of the required size can be declared internally. In other cases, allocatable arrays may need to be declared and explicitly allocated. Explicit allocation is needed in particular when the amount of workspace required depends on the block-size to be used (which is not passed as an argument).
- 3. Optional arguments: In LAPACK 77, character arguments are frequently used to specify some choice of options. In Fortran 90, a choice of options can sometimes be specified naturally by the presence or absence of optional arguments: for example, options to compute the left or right eigenvectors can be specified by the presence of arguments VL or VR, and the character arguments JOBVL and JOBVR which are required in the LAPACK 77 routine DGEEV, are not needed in LAPACK 90.

In other routines, a character argument to specify options may still be required, but can itself be made optional if there is a natural default value: for example, in DGESVX the argument TRANS can be made optional, with default value 'N'.

Optional arguments can also help to combine two or more routines into one: for example, the functionality provided by the routine DGECON can be made accessible by adding an optional argument RCOND to DGETRF.

4. **Generic Interfaces:** The systematic occurrence in LAPACK of analogous routines for real or complex data, and for single or double precision lends itself well to the definition of generic interfaces, allowing four different routines to be accessed through the same generic name.

Generic interfaces can also be used to cover routines whose arguments differ in *rank*, and thus provide a slight increase in flexibility over LA-PACK 77. For example, in LAPACK 77, routines for solving a system of linear equations (such as DGESV), allow for multiple right hand sides, and so the arrays which hold the right hand sides and solutions are always

of rank 2. In LAPACK 90, we can provide alternative versions of the routines (covered by a single generic interface) in which the arrays holding the right hand sides and solutions may *either* be of rank 1 (for a single right hand side) or be of rank 2 (for several right hand sides).

- 5. **Naming:** For the generic routine names, we propose:
  - the initial letter (S, C, D or Z) is simply omitted.
  - the letters LA\_ are prefixed to all names to identify them as names of LAPACK routines.

In other respects the naming scheme remains the same as described in Section 2.1.3 of [1]: for example, LA\_GESV.

It would also be possible to define longer, more meaningful names (which could co-exist with the shorter names), but we have not attempted this here.

We have *not* proposed the use of any *derived types* in this Fortran 90 interface. They could be considered — for example, to hold the details of an *LU* factorization and equilibration factors. However, since LAPACK routines are so frequently used as building blocks in larger algorithms or applications, we feel that there are advantages in keeping the interface simple, and avoiding possible loss of efficiency.

# 6. Error-handling:

In LAPACK 77, all documented routines have a diagnostic output argument INFO. Three types of exit from a routine are allowed:

**successful termination:** the routine returns to the calling program with INFO set to 0.

- illegal value of one or more arguments, or error in store allocation: the routine sets INFO < 0 and calls the auxiliary routine XERBLA; the standard version of XERBLA issues an error message identifying the first invalid argument, and stops execution.
- failure in the course of computation: the routine sets INFO > 0 and returns to the calling program without issuing any error message. Only some LAPACK 77 routines need to allow this type of errorexit; it is then the responsibility of a user to test INFO on return to the calling program.

For LAPACK 90 we propose that the argument INFO becomes optional: if it is not present and an error occurs, then the routine always issues an error message and stops execution, even when INFO > 0 (in which case the error message reports the value of INFO). If a user wishes to continue

execution after a failure in computation, then INFO must be supplied and tested on return.

This behaviour simplifies calls to LAPACK 90 routines when there is no need to test INFO on return, and makes it less likely that users will forget to test INFO when necessary.

If an invalid argument is detected, we propose that routines issue an error message and stop, as in LAPACK 77. Note however that in Fortran 90 there can be different reasons for an argument being invalid:

illegal value: as in LAPACK 77.

**invalid shape** (of an assumed-shape array): for example, a 2-dimensional array is not square when it is required to be.

inconsistent shapes (of two or more assumed-shape arrays): for example, arrays holding the right hand sides and solutions of a system of linear equations must have the same shape.

No more core allocation needed for the LAPACK 77.

The specification could be extended so that the error-message could distinguish between these cases. For more detail see in appendix section D.2.

# 4 Prototype LAPACK 90 Interfaces

We have implemented Fortran 90 jacket procedures to the subset of LAPACK 77 routines concerned with the solution of systems of linear equations AX = B for a general matrix A—that is, the driver routines xGESV and xGESVX, and the computational routines xGETRF, xGETRS, xGETRI, xGECON, xGERFS and xGEEQU. We also consider here the symmetric and Hermitian eigenproblem routines xSYTRD, xSYGV, xSYGST, xORGTR, xSYEV, xSYEVD, xSYEVX and the factor routine xPOTRF which is strongly connected with the xSYGST subroutines.

Here we present calling sequences for each of the proposed routines, the first without using any of the optional arguments, the second using all the arguments. For ease of comparison between LAPACK 77 and LAPACK 90, we have retained the same names for the corresponding arguments, although of course Fortran 90 offers the possibility of longer names (for example, IPIV could become PIVOT\_INDICES). In this prototype implementation, we have assumed that the code of LAPACK 77 is not modified.

Detailed documentation of the proposed interfaces can be found in Appendices A, B and C.

# 4.1 Solution of Systems of Linear Equations for a General Matrix A

LA\_GESV (simple driver):

```
CALL LA_GESV( A, B )

CALL LA_GESV( A, B, IPIV, INFO )
```

# Comments:

• The array B may have rank 1 (one right hand side) or rank 2 (several right hand sides).

LA\_GESVX (expert driver):

```
CALL LA_GESVX( A, B, X )

CALL LA_GESVX( A, B, X, AF, IPIV, FACT, TRANS, EQUED, R, C, & FERR, BERR, RCOND, RPVGRW, INFO )
```

## Comments:

- The arrays B and X may have rank 1 (in which case FERR and BERR are scalars) or rank 2 (in which case FERR and BERR are rank-1 arrays).
- RPVGRW returns the reciprocal pivot growth factor (returned in WORK(1) in LAPACK 77).
- the presence or absence of EQUED is used to specify whether or not equilibration is to be performed, instead of the option FACT = 'E'.

LA\_GETRF (LU factorization):

```
CALL LA_GETRF( A, IPIV )

CALL LA_GETRF( A, IPIV, RCOND, NORM, INFO )
```

# Comments:

• instead of a separate routine LA\_GECON, we propose that optional arguments RCOND and NORM are added to LA\_GETRF to provide the same functionality in a more convenient manner. The argument ANORM of xGECON is not needed, because LA\_GETRF can always compute the norm of A if required.

LA\_GETRS (solution of equations using LU factorization):

```
CALL LA_GETRS( A, IPIV, B )

CALL LA_GETRS( A, IPIV, B, TRANS, INFO )
```

## Comments:

• The array B may have rank 1 or 2.

LA\_GETRI (matrix inversion using LU factorization):

```
CALL LA_GETRI( A, IPIV )

CALL LA_GETRI( A, IPIV, INFO )
```

LA\_GERFS (refine solution of equations and optionally compute error bounds):

```
CALL LA_GERFS( A, AF, IPIV, B, X )

CALL LA_GERFS( A, AF, IPIV, B, X, TRANS, FERR, BERR, INFO )
```

## Comments:

• The arrays B and X may have rank 1 (in which case FERR and BERR are scalars) or rank 2 (in which case FERR and BERR are rank-1 arrays).

LA\_GEEQU (equilibration):

```
CALL LA_GEEQU(A, R, C)

CALL LA_GEEQU(A, R, C, ROWCND, COLCND, AMAX, INFO)
```

# 4.2 Symmetric and Hermitian Eigenproblem Routines

LA\_SYEV / LA\_HEEV (all eigenvalues and optionally eigenvectors):

```
CALL LA_SYEV / LA_HEEV( A, W )

CALL LA_SYEV / LA_HEEV( A, W, JOBZ, UPLO, INFO )
```

 ${\tt LA\_SYEVD}$  /  ${\tt LA\_HEEVD}$  (all eigenvalues and optionally eigenvectors using a divide and conquer algorithm):

```
CALL LA_SYEVD / LA_HEEVD( A, W )

CALL LA_SYEVD / LA_HEEVD( A, W, JOBZ, UPLO, INFO )
```

LA\_SYEVX / LA\_HEEVX (selected eigenvalues and optionally eigenvectors):

```
CALL LA_SYEVX / LA_HEEVD( A, W )
       CALL LA_SYEVX / LA_HEEVD( A, W, JOBZ, UPLO, VL, VU, IL, IU, &
                                   M, IFAIL, ABSTOL, INFO )
LA_SYGV / LA_HEGV (all eigenvalues and optionally eigenvectors of the form
     Ax = \lambda Bx, ABx = \lambda x, or BAx = \lambda x
       CALL LA_SYGV / LA_HEGV( A, B, W )
       CALL LA_SYGV / LA_HEGV( A, B, W, ITYPE, JOBZ, UPLO, INFO )
LA_SYGST / LA_HEGST (reduction to standard form):
       CALL LA_SYGST / LA_HEGST( A, B )
       CALL LA_SYGST / LA_HEGST( A, B, ITYPE, UPLO, INFO )
LA_SYTRD / LA_HETRD (reduction to tridiagonal form):
       CALL LA_SYTRD / LA_HETRD( A, TAU )
       CALL LA_SYTRD / LA_HETRD( A, TAU, UPLO, INFO )
LA_ORGTR / LA_UNGTR (generates an orthogonal matrix):
       CALL LA_ORGTR / LA_UNGTR( A, TAU )
       CALL LA_ORGTR / LA_UNGTR( A, TAU, UPLO, INFO )
LA_POTRF (generates an orthogonal matrix):
       CALL LA_POTRF( A )
       CALL LA_POTRF( A, UPLO, RCOND, NORM, INFO )
```

# 5 Documentation

In appendices A, B and C, we give a first attempt at draft documentation for these routines. The style is somewhat similar to that of the LAPACK Users' Guide, but with various obvious new conventions introduced to handle the generic nature of the interfaces.

# 6 Test Software

Additional test software is being developed to test the new interfaces. At present, the test software is a modified version of the LAPACK 77 Test Suite.

# 7 Performance Issues and Timings

The present FORTRAN 90 compilers do not assume that array storage is contiguous. The effect on performance will of course depend on their compiler. The contents of arrays are copied to the temporary storage when calling FORTRAN 77 subroutines (functions) from the FORTRAN 90 procedures. This data copy results in a degradation in performance. As previously mentioned, we are aware of performance enhancements to the interface which could be made available through alternative interfaces to the routines. However, these enhancements violate our goal to present a true Fortran 90 interface to the LAPACK library by complicating the design of the interface through the use of Fortran 77 constructs. Our Fortran 90 interface is provided as a simplified user interface to the LAPACK library. If performance is the main focus of the user's application, he should call the Fortran 77 interface directly.

We have performed timings to measure the extra overhead of the Fortran 90 interface. We timed LA\_GETRF on a single processor of an IBM SP-2 (in double precision) and a single processor of a Cray YMP C90A (in single precision). All timings are given in megaflops.

 Speed of LAPACK 90 calling LAPACK 77 and BLAS from the ESSL library.

2. Speed of LAPACK 77, using BLAS from the ESSL library.

Array size	600	700	800	900	1000	1100	1200	1300	1400	1500
LAPACK90	187	180	182	170	172	172	176	177	181	182
LAPACK77	191	181	182	171	172	173	176	179	180	182

Cray 1. Speed of LAPACK 90 calling LAPACK 77 as provided by CRAY in LIBSCL

2. Speed of LAPACK 77 as provided by CRAY in LIBSCI.

Array size	600	700	800	900	1000	1100	1200	1300	1400	1500
LAPACK90	723	828	646	841	822	855	789	857	846	868
LAPACK77	778	834	649	845	825	860	794	864	848	873

The above tables show the LAPACK 90 results are a little slower (1 or 2%) than the LAPACK 77 results.

# 8 Acknowledgments

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# References

- [1] E. Anderson, Z. Bai, C. H. Bischof, J. Demmel, J. J. Dongarra, J. Du Croz, A. Greenbaum, S. Hammarling, A. McKenney, S. Ostrouchov and D. C. Sorensen. LAPACK Users' Guide Release 2.0. SIAM, Philadelphia, 1995.
- [2] J.J. Dongarra, J. Du Croz, S. Hammarling, J. Waśniewski and A. Zemła. LAPACK Working Note 101, A Proposal for a Fortran 90 Interface for LA-PACK. Report UNIC-95-9, UNI•C, Lyngby, Denmark, 1995. Report ut-cs-95-295, University of Tennessee, Computer Science Department, Knoxville, July, 1995.
- [3] M. Metcalf and J. Reid. Fortran 90 Explained. Oxford, New York, Tokyo, Oxford University Press, 1990.

# A Solving Systems of Linear Equations AX = B with a General Matrix A, Documentation

# A.1 LA\_GESV

# A.1.1 Purpose

**LA\_GESV** computes the solution to either a real or complex system of linear equations AX = B, where A is a square matrix and B and X are either rectangular matrices or vectors.

The LU decomposition with partial pivoting and row interchanges is used to factor A as A = PLU, where P is a permutation matrix, L is unit lower triangular, and U is upper triangular. The factored form of A is then used to solve the system of equations AX = B.

# A.1.2 Specification

```
SUBROUTINE LA_GESV( A, B, IPIV, INFO ) type(wp), INTENT(INOUT) :: A(:,:), rhs \\ INTEGER, INTENT(OUT), OPTIONAL :: IPIV(:) \\ INTEGER, INTENT(OUT), OPTIONAL :: INFO \\ where \\ type ::= REAL \mid COMPLEX \\ wp ::= KIND(1.0) \mid KIND(1.0D0) \\ rhs ::= B(:,:) \mid B(:)
```

# A.1.3 Arguments

- A (input/output) REAL or COMPLEX array, shape (:,:), size(A,1) = size(A,2).
  - On entry, the matrix A.
  - On exit, the factors L and U from the factorization A = PLU; the unit diagonal elements of L are not stored.
- **B** (input/output) **REAL** or **COMPLEX** array, shape (:, :) or (:), and size(B, 1) or size(B) = size(A, 1).
  - On entry, the right hand side vector(s) of matrix B for the system of equations AX = B.
  - On exit, if there is no error, the matrix of solution vector(s) X.
- IPIV Optional (output) INTEGER array, shape (:), <math>size(IPIV) = size(A, 1).

• If IPIV is present, it contains indices that define the permutation matrix P; row i of the matrix was interchanged with row IPIV(i).

# INFO - Optional (output) INTEGER.

- If *INFO* is present
  - = 0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - > 0: if INFO = i, U(i, i) is exactly zero. The factorization has been completed, but the factor U is exactly singular, so the solution could not be computed.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

#### A.2 LA\_GESVX

# A.2.1 Purpose

LA\_GESVX computes the solution to a real or complex system of linear equations AX = B, where A is a square matrix and B and X are either rectangular matrices or vectors.

LA\_GESVX is an expert driver routine, which can also optionally perform the following functions:

- solve  $A^T X = B$  or  $A^H X = B$ ,
- estimate the condition number of A
- return the pivot growth factor
- refine the solution and compute forward and backward error bounds
- equilibrate the system if A is poorly scaled.

# A.2.2 Specification

SUBROUTINE LA\_GESVX (A, B, X, AF, IPIV, FACT, TRANS, EQUED, & R, C, FERR, BERR, RCOND, RPVGRW, INFO) type(wp), INTENT(INOUT) :: A(:,:), rhs

type(wp), INTENT(OUT) :: sol

type(wp), INTENT(INOUT), OPTIONAL :: AF(:,:)

INTEGER, INTENT(INOUT), OPTIONAL :: IPIV(:)

CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: FACT, TRANS

CHARACTER(LEN=1), INTENT(INOUT), OPTIONAL :: EQUED

REAL(wp), INTENT(INOUT), OPTIONAL :: R(:), C(:)

REAL(wp), INTENT(OUT), OPTIONAL :: err, RCOND, RPVGRW

```
INTEGER, INTENT(OUT), OPTIONAL :: INFO where type ::= REAL \mid COMPLEX
wp ::= KIND(1.0) \mid KIND(1.0D0)
rhs ::= B(:,:) \mid B(:)
sol ::= X(:,:) \mid X(:)
err ::= FERR(:), BERR(:) \mid FERR, BERR
```

# A.2.3 Description

The following steps are performed:

1. If FACT is not present or FACT = 'N', and EQUED is present, real scaling factors are computed to equilibrate the system:

```
TRANS = 'N' : diag(R) \ A \ diag(C) \ (diag(C))^{-1} \ X = diag(R) \ B
TRANS = 'T' : (diag(R) \ A \ diag(C))^T \ (diag(R))^{-1} \ X = diag(C) \ B
TRANS = 'C' : (diag(R) \ A \ diag(C))^H \ (diag(R))^{-1} \ X = diag(C) \ B
```

Whether or not the system will be equilibrated depends on the scaling of the matrix A, but if equilibration is used, A is overwritten by diag(R) A diag(C) and B by diag(R) B (if TRANS = 'N') or diag(C) B (if TRANS = 'T' or 'C').

- 2. If FACT = 'N', the LU decomposition is used to factor the matrix A (after equilibration if EQUED is present) as A = PLU, where P is a permutation matrix, L is a unit lower triangular matrix, and U is upper triangular.
- 3. The factored form of A is used to estimate the condition number of the matrix A. If the reciprocal of the condition number is less than machine precision, steps 4-6 are skipped.
- 4. The system of equations is solved for X using the factored form of A.
- 5. Iterative refinement is applied to improve the computed solution matrix and calculate error bounds and backward error estimates for it.
- 6. If equilibration was used, the matrix X is premultiplied by diag(C) (if TRANS = 'N') or diag(R) (if TRANS = 'T' or 'C') so that it solves the original system before equilibration.

# A.2.4 Arguments

 $\mathbf{A} - (input/output) \ \mathbf{REAL} \ \text{or} \ \mathbf{COMPLEX} \ \text{array, shape} \ (:,:), \ size(A,1) = size(A,2).$ 

If FACT is not present or FACT = 'N',

- On entry, the matrix A.
- On exit, if *EQUED* is present, the matrix *A* may have been overwritten by the equilibrated matrix (see *EQUED*).

If FACT is present and FACT = 'F',

- On entry, the matrix A, possibly equilibrated in a previous call to LA\_GESVX (see EQUED).
- On exit, A is unchanged.
- **B** (input/output) **REAL** or **COMPLEX** array, shape (:, :) or (:), and size(B, 1) or size(B) = size(A, 1).
  - On entry, the right hand side vector(s) of matrix B for the system of equations AX = B.
  - On exit, if EQUED is present, B may have been scaled in accordance with the equilibration of A (see EQUED); otherwise, B is unchanged.
- $\mathbf{X} (output) \ \mathbf{REAL}$  or  $\mathbf{COMPLEX}$  array, shape (:,:) or (:), size(X,1) or size(X) = size(A,1).

  If INFO = 0, the solution matrix (vector) X to the original system of equations. Note that X always returns the solution to the original system of equations; if equilibration has been performed (EQUED is present and  $EQUED \neq 'N'$ ), this does not correspond to the scaled A and B.
- $\mathbf{AF} Optional\ (input/output)\ \mathbf{REAL}\ \text{or}\ \mathbf{COMPLEX}\ \text{array}, \text{shape}\ (:,:),\ size(AF,1) = size(AF,2) = size(A,1).$ 
  - If FACT is not present or FACT = 'N', then AF is an *output* argument and returns the factors L and U from the factorization A = PLU of the original matrix A, possibly equilibrated if EQUED is present.
  - If FACT is present and FACT = 'F', then AF is an input argument (and must be present); on entry, it must contain the factors L and U of A (possibly equilibrated if EQUED is present), returned by a previous call to  $\mathbf{LA\_GESVX}$ .
- **IPIV** Optional (input/output) **INTEGER** array, shape (:), size(IPIV) = size(A, 1).

- If FACT is not present or FACT = 'N', then IPIV is an output argument and returns the pivot indices from the factorization A = PLU of the original matrix A, possibly equilibrated if EQUED is present.
- If FACT is present and FACT = 'F', then IPIV is an input argument (and must be present); on entry, it must contain the pivot indices from the factorization of A (possibly equilibrated if EQUED is present), returned by a previous call to  $\mathbf{LA\_GESVX}$ .

# TRANS - Optional (input) CHARACTER\*1.

- If TRANS is present, it specifies the form of the system of equations:
  - = 'N' : AX = B (No transpose)
  - $= 'T' : A^T X = B \text{ (Transpose)}$
  - $= 'C' : A^H X = B$  (Conjugate transpose)
- otherwise TRANS = 'N' is assumed.

# FACT - Optional (input) CHARACTER\*1.

Specifies whether or not the factored form of the matrix A is supplied on entry.

- If FACT is present then:
  - ='N': the matrix A will be equilibrated if EQUED is present, then copied to AF and factored.
  - ='F': on entry, AF and IPIV must contain the factored form of A (possibly equilibrated if EQUED is present).
- otherwise FACT = 'N' is assumed.

# EQUED - Optional (input/output) CHARACTER\*1.

- If FACT is not present or FACT = 'N', then EQUED is an output argument. If it is present, then the matrix is equilibrated, and on exit EQUED specifies the scaling of A which has actually been performed:
  - = 'N' : No equilibration.
  - = 'R': Row equilibration, i.e., A has been premultiplied by diag(R); also B has been premultiplied by diag(R) if TRANS = 'N'.
  - ='C': Column equilibration, i.e., A has been postmultiplied by diag(C); also B has been premultiplied by diag(C) if TRANS ='T' or 'C'.
  - = 'B': Both row and column equilibration: combines the effects of EQUED = 'R' and EQUED = 'C'.

- If FACT is present and FACT = 'F', then EQUED is an input argument; if it is present, it specifies the equilibration of A which was performed in a previous call to  $\mathbf{LA\_GESVX}$  with FACT not present or FACT = 'N'.
- **R** Optional (input/output) **REAL** array, shape (:), size(R) = size(A, 1). R must be present if EQUED is present and EQUED = 'R' or 'B'; R is not referenced if EQUED = 'N' or 'C'.
  - If FACT is not present or FACT = 'N', then R is an output argument. If EQUED = 'R' or 'B', R returns the row scale factors for equilibrating A.
  - If FACT is present and FACT = 'F', then R is an input argument. If EQUED = 'R' or 'B', R must contain the row scale factors for equilibrating A, returned by a previous call to  $\mathbf{LA\_GESVX}$ ; each element of R must be positive.
- C Optional (input/output) **REAL** array, shape (:), size(C) = size(A, 1). C must be present if EQUED is present and EQUED = 'C' or 'B'; C is not referenced if EQUED = 'N' or 'R'.
  - If FACT is not present or FACT = 'N', then C is an output argument. If EQUED = 'C' or 'B', C returns the column scale factors for equilibrating A.
  - If FACT is present and FACT = 'F', then C is an input argument. If EQUED = 'C' or 'B', C must contain the column scale factors for equilibrating A, returned by a previous call to  $\mathbf{LA\_GESVX}$ ; each element of C must be positive.
- **FERR** Optional (output) **REAL** array of shape (:) or **REAL** scalar. If it is an array, size(FERR) = size(X,2). The estimated forward error bound for each solution vector X(j) (the j-th column of the solution matrix X). If XTRUE is the true solution corresponding to X(j), FERR(j) is an estimated upper bound for the magnitude of the largest element in (X(j) XTRUE) divided by the magnitude of the largest element in X(j). The estimate is as reliable as the estimate for RCOND, and is almost always a slight overestimate of the true error.
- **BERR** Optional (output) **REAL** array of shape (:) or **REAL** scalar. If it is an array, size(BERR) = size(X, 2). The componentwise relative backward error of each solution vector X(j) (i.e., the smallest relative change in any element of A or B that makes X(j) an exact solution).
- RCOND Optional (output) REAL.

The estimate of the reciprocal condition number of the matrix A after equilibration (if done). If RCOND is less than the machine precision (in

particular, if RCOND = 0), the matrix is singular to working precision. This condition is indicated by a return code of INFO > 0, and the solution and error bounds are not computed.

# RPVGRW - Optional (output) REAL.

The reciprocal pivot growth factor  $\|A\|_{\infty}/\|U\|_{\infty}$ . If RPVGRW is much less than 1, then the stability of the LU factorization of the (equilibrated) matrix A could be poor. This also means that the solution X, condition estimator RCOND, and forward error bound FERR could be unreliable. If factorization fails with  $0 < INFO \le size(A,1)$ , then RPVGRW contains the reciprocal pivot growth factor for the leading INFO columns of A.

# **INFO** – Optional (output) **INTEGER**.

• If INFO is present

= 0 : successful exit

< 0: if INFO = -i, the *i*-th argument had an illegal value

> 0: if INFO = i, and i is

 $\leq N: U(i,i)$  is exactly zero. The factorization has been completed, but the factor U is exactly singular, so the solution and error bounds could not be computed.

- =N+1: RCOND is less than machine precision. The factorization has been completed, but the matrix is singular to working precision, and the solution and error bounds have not been computed.
- ullet If INFO is not present and an error occurs, then the program is terminated with an error message.

# A.3 LA\_GETRF

# A.3.1 Purpose

**LA\_GETRF** computes an LU factorization of a general rectangular matrix A using partial pivoting with row interchanges.

The factorization has the form A = PLU where P is a permutation matrix, L is lower triangular with unit diagonal elements (lower trapezoidal if m > n), and U is upper triangular (upper trapezoidal if m < n), where m = size(A, 1) and n = size(A, 2).

When A is square (m=n), **LA\_GETRF** optionally estimates the reciprocal of the condition number of the matrix A, in either the 1-norm or the  $\infty$ -norm. An estimate is obtained for  $||A^{-1}||$ , and the reciprocal of the condition number is computed as  $RCOND = 1/(||A|| ||A^{-1}||)$ .

# A.3.2 Specification

```
SUBROUTINE LA_GETRF( A, IPIV, RCOND, NORM, INFO ) type(wp), INTENT(INOUT) :: A(:,:) INTEGER, INTENT(OUT) :: IPIV( : ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: NORM REAL(wp), INTENT(OUT), OPTIONAL :: RCOND INTEGER, INTENT(OUT), OPTIONAL :: INFO where type ::= REAL \mid COMPLEX \ wp ::= KIND(1.0) \mid KIND(1.0D0)
```

# A.3.3 Arguments

A - (input/output) **REAL** or **COMPLEX** array, shape (:,:).

- On entry, the matrix A.
- On exit, the factors L and U from the factorization A = PLU; the unit diagonal elements of L are not stored.
- **IPIV** (output) **INTEGER** array, shape (:), size(IPIV) = min(size(A, 1), size(A, 2)). Indices that define the permutation matrix P; row i of the matrix was interchanged with row IPIV(i).

# RCOND - Optional (output) REAL.

The reciprocal of the condition number of the matrix A for the case m=n, computed as  $RCOND=1/(\|A\| \|A^{-1}\|)$ . RCOND should be present if NORM is present. If  $m \neq n$  then RCOND is returned as zero.

# **NORM** – Optional (input) **CHARACTER\*1**.

Specifies whether the 1-norm condition number or the  $\infty$ -norm condition number is required:

- = '1', 'O' or 'o': 1-norm;
- = I', i':  $\infty$ -norm.

If NORM is not present, the 1-norm is used.

**INFO** – Optional (output) **INTEGER**.

- If *INFO* is present
  - = 0: successful exit
  - <0: if INFO=-i, the *i*-th argument had an illegal value

- > 0: if INFO = i, U(i,i)] is exactly zero. The factorization has been completed, but the factor U is exactly singular, so the solution could not be computed.
- If *INFO* is not present and an error occurs, then the program is terminated with an error message.

# A.4 LA\_GETRS

# A.4.1 Purpose

**LA\_GETRS** solves a system of linear equations AX = B,  $A^TX = B$  or  $A^HX = B$  with a general square matrix A, using the LU factorization computed by **LA\_GETRF**.

# A.4.2 Specification

```
SUBROUTINE LA_GETRS (A, IPIV, B, TRANS, INFO) type(wp), INTENT(IN) :: A(:,:) \\ INTEGER, INTENT(IN) :: IPIV(:) \\ type(wp), INTENT(INOUT) :: rhs \\ CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: TRANS \\ INTEGER, INTENT(OUT), OPTIONAL :: INFO \\ where \\ type ::= REAL \mid COMPLEX \\ wp ::= KIND(1.0) \mid KIND(1.0D0) \\ rhs ::= B(:,:) \mid B(:)
```

## A.4.3 Arguments

- **A** (input) **REAL** or **COMPLEX** array, shape (:,:), size(A,1) = size(A,2). The factors L and U from the factorization A = PLU as computed by **LA\_GETRF**.
- **IPIV** (input) **INTEGER** array, shape (:), size(IPIV) = size(A, 1). The pivot indices from **LA\_GETRF**; for  $1 \le i \le size(A, 1)$ , row i of the matrix was interchanged with row IPIV(i).
- **B** (input/output) **REAL** or **COMPLEX** array, shape (:, :) or (:), and size(B, 1) or size(B) = size(A, 1).
  - On entry, the right hand side vector(s) of matrix B for the system of equations AX = B.
  - On exit, if there is no error, the matrix of solution vector(s) X.

TRANS - Optional (input) CHARACTER\*1.

• If TRANS is present, it specifies the form of the system of equations:

```
= 'N': AX = B (No transpose)
= 'T': A^TX = B (Transpose)
= 'C': A^HX = B (Conjugate transpose)
```

• otherwise TRANS = 'N' is assumed.

# INFO - Optional (output) INTEGER.

 $\bullet$  If INFO is present

= 0: successful exit

<0: if INFO=-i, the *i*-th argument had an illegal value

 $\bullet$  If INFO is not present and an error occurs, then the program is terminated with an error message.

# A.5 LA\_GETRI

# A.5.1 Purpose

**LA\_GETRI** computes the inverse of a matrix using the LU factorization computed by **LA\_GETRF**.

# A.5.2 Specification

```
SUBROUTINE LA_GETRI (A, IPIV, INFO) type(wp), INTENT(INOUT) :: A(:,:) INTEGER, INTENT(IN) :: IPIV(:) INTEGER, INTENT(OUT), OPTIONAL :: INFO where type ::= \text{REAL} \mid \text{COMPLEX} wp ::= \text{KIND}(1.0) \mid \text{KIND}(1.0D0)
```

# A.5.3 Arguments

- A (input/output) REAL or COMPLEX array, shape (:,:), size(A,1) = size(A,2).
  - On entry contains the factors L and U from the factorization A = PLU as computed by **LA\_GETRF**.
  - On exit, if INFO = 0, the inverse of the original matrix A.

**IPIV** – (input) **INTEGER** array, shape (:), size(IPIV) = size(A, 1). The pivot indices from **LA\_GETRF**; for  $1 \le i \le size(A, 1)$ , row i of the matrix was interchanged with row IPIV(i).

# INFO - Optional (output) INTEGER.

- If INFO is present
  - = 0: successful exit
  - < 0: if INFO = -i, the *i*-th argument had an illegal value
  - > 0: if INFO = i,  $\mathrm{U}(\mathrm{i},\mathrm{i})$  is exactly zero; the matrix is singular and its inverse could not be computed.
- If *INFO* is not present and an error occurs, then the program is terminated with an error message.

# A.6 LA\_GERFS

# A.6.1 Purpose

**LA\_GERFS** improves the computed solution X of a system of linear equations AX = B or  $A^TX = B$  and provides error bounds and backward error estimates for the solution. **LA\_GERFS** uses the LU factors computed by **LA\_GETRF**.

# A.6.2 Specification

```
SUBROUTINE LA_GERFS (A, AF, IPIV, B, X, & TRANS, FERR, BERR, INFO)

type(wp), INTENT(IN) :: A(:,:), AF(:,:), rhs

INTEGER, INTENT(IN) :: IPIV(:)

type(wp), INTENT(INOUT) :: sol

CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: TRANS

REAL(wp), INTENT(OUT), OPTIONAL :: err

INTEGER, INTENT(OUT), OPTIONAL :: INFO

where

type ::= REAL \mid COMPLEX

wp ::= REAL \mid COMPLEX

wp ::= KIND(1.0) \mid KIND(1.0D0)

rhs ::= B(:,:) \mid B(:)

sol ::= X(:,:) \mid X(:)

err ::= FERR(:), BERR(:) | FERR, BERR
```

# A.6.3 Arguments

**A** – (input) **REAL** or **COMPLEX** array, shape (:,:), size(A, 1) = size(A, 2). The original matrix A.

- $\mathbf{AF} (input) \mathbf{REAL} \text{ or } \mathbf{COMPLEX} \text{ array, shape } (:,:), size(AF,1) = size(AF,2) = size(A,1).$ 
  - The factors L and U from the factorization A = PLU as computed by  $\mathbf{LA\_GETRF}$ .
- **IPIV** (*input*) **INTEGER** array, shape (:), size(IPIV) = size(A, 1). The pivot indices from **LA\_GETRF**; for  $1 \le i \le size(A, 1)$ , row i of the matrix was interchanged with row IPIV(i).
- **B** (input) **REAL** or **COMPLEX** array, shape (:,:) or (:), size(B, 1) or size(B) = size(A, 1). The right hand side vector(s) of matrix B for the system of equations AX = B.
- $\mathbf{X} (input/output) \mathbf{REAL}$  or  $\mathbf{COMPLEX}$  array, shape (:,:) or (:), size(X, 1) or size(X) = size(A, 1).
  - On entry, the solution matrix X, as computed by **LA\_GETRS**.
  - On exit, the improved solution matrix X.
- TRANS Optional (input) CHARACTER\*1.
  - If TRANS is present, it specifies the form of the system of equations:
    - = 'N' : AX = B (No transpose)
    - $= 'T' : A^T X = B \text{ (Transpose)}$
    - $= 'C' : A^H X = B$  (Conjugate transpose)
  - otherwise TRANS = 'N' is assumed.
- **FERR** Optional (output) **REAL** array of shape (:) or **REAL** scalar.
  - If it is an array, size(FERR) = size(X, 2). The estimated forward error bound for each solution vector X(j) (the j-th column of the solution matrix X). If XTRUE is the true solution corresponding to X(j), FERR(j) is an estimated upper bound for the magnitude of the largest element in (X(j) XTRUE) divided by the magnitude of the largest element in X(j). The estimate is as reliable as the estimate for RCOND, and is almost always a slight overestimate of the true error.
- **BERR** Optional (output) **REAL** array of shape (:) or **REAL** scalar. If it is an array, size(BERR) = size(X, 2). The componentwise relative backward error of each solution vector X(j) (i.e., the smallest relative change in any element of A or B that makes X(j) an exact solution).
- INFO Optional (output) INTEGER.
  - If INFO is present

= 0: successful exit

< 0: if INFO = -i, the i-th argument had an illegal value

• If *INFO* is not present and an error occurs, then the program is terminated with an error message.

## A.6.4 Internal Parameters

ITMAX – is the maximum number of steps of iterative refinement. It is set to 5 in the LAPACK 77 subroutines (see [1]).

# A.7 LA\_GEEQU

# A.7.1 Purpose

**LA\_GEEQU** computes row and column scalings intended to equilibrate a rectangle matrix A and reduce its condition number. R returns the row scale factors and C the column scale factors, chosen to try to make the largest entry in each row and column of the matrix B with elements  $B_{ij} = R_i A_{ij} C_j$  have absolute value 1.

 $R_i$  and  $C_j$  are restricted to be between SMLNUM = smallest safe number and BIGNUM = largest safe number. Use of these scaling factors is not guaranteed to reduce the condition number of A but works well in practice.

# A.7.2 Specification

```
SUBROUTINE LA_GEEQU ( A, R, C, ROWCND, COLCND, & AMAX, INFO ) type(wp), \ INTENT(IN) :: \ A(:,:) \\ REAL(wp), \ INTENT(OUT) :: \ R(:), \ C(:) \\ REAL(wp), \ INTENT(OUT), \ OPTIONAL :: \ ROWCND, & COLCND, \ AMAX \\ INTEGER, \ INTENT(OUT), \ OPTIONAL :: \ INFO \\ where <math display="block">type ::= REAL \mid COMPLEX \\ wp ::= KIND(1.0) \mid KIND(1.0D0)
```

# A.7.3 Arguments

```
A – (input) REAL or COMPLEX array, shape (:,:). The matrix A, whose equilibration factors are to be computed.
```

- **R** (output) **REAL** array, shape (:), size(R) = size(A, 1). If INFO = 0 or INFO > size(A, 1), R contains the row scale factors for A.
- C (output) REAL array, shape (:), size(C) = size(A, 2). If INFO = 0, C contains the column scale factors for A.
- ROWCND Optional (output) REAL.

If INFO = 0 or INFO > size(A, 1), ROWCND contains the ratio of the smallest R(i) to the largest R(i). If  $ROWCND \ge 0.1$  and AMAX is neither too large nor too small, it is not worth scaling by R.

COLCND - Optional (output) REAL.

If INFO = 0, COLCND contains the ratio of the smallest C(i) to the largest C(i). If COLCND > 0.1, it is not worth scaling by C.

AMAX - Optional (output) REAL.

Absolute value of largest matrix element. If AMAX is very close to over-flow or very close to underflow, the matrix should be scaled.

- INFO Optional (output) INTEGER.
  - If *INFO* is present

= 0: successful exit

 $<0\,:$  if INFO=-i, the i-th argument had an illegal value

> 0: if INFO = i, and i is

 $\leq m$ : the *i*-th row of A is exactly zero

>m : the (i-m)-th column of A is exactly zero

where m = size(A, 1).

• If *INFO* is not present and an error occurs, then the program is terminated with an error message.

# B Symmetric and Hermitian Eigenvalue and Eigenvector Procedures, Documentation

# B.1 LA\_SYEV / LA\_HEEV

# **B.1.1** Purpose

 ${f LA\_SYEV}$  /  ${f LA\_HEEV}$  computes all eigenvalues and, optionally, eigenvectors of a real symmetric or complex Hermitian matrix A.

# **B.1.2** Specification

```
SUBROUTINE LA_SYEV / LA_HEEV( A, W, JOBZ, UPLO, INFO ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: JOBZ, UPLO INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), \ INTENT(INOUT) :: \ A(:,:) \\ type(wp), \ INTENT(OUT) :: \ W(:) \\ where \\ type ::= REAL \mid COMPLEX \\ wp ::= KIND(1.0) \mid KIND(1.0D0)
```

#### B.1.3 Defaults

- If JOBZ is not present then JOBZ = 'N' is assumed.
- If UPLO is not present then UPLO = 'U' is assumed.

# **B.1.4** Arguments

A - (input/output) REAL or COMPLEX array, shape (:,:)

- On entry, the symmetric (Hermitian) matrix A.
  - If UPLO = 'U', the upper triangular part of A contains the upper triangular part of the matrix A.
  - If UPLO = L', the lower triangular part of A contains
  - the lower triangular part of the matrix A.
- On exit:
  - If JOBZ = V', then if INFO = 0, A contains the orthonormal eigenvectors of the matrix A.
  - If JOBZ = 'N', then on exit the lower triangle (if UPLO='L') or the upper triangle (if UPLO='U') of A, including the diagonal, is destroyed.

 $\mathbf{W} - (output) \mathbf{REAL} \text{ array, shape (:), size(W)} = \text{size(A,1)} > 0.$ 

• If INFO = 0, the eigenvalues in ascending order.

# JOBZ - Optional, (input) CHARACTER\*1

- If JOBZ is present then:
  - = 'N': Compute eigenvalues only;
  - = 'V': Compute eigenvalues and eigenvectors.
- otherwise JOBZ = 'N' is assumed.

# **UPLO** – Optional, (input) **CHARACTER\*1**

- If UPLO is present then:
  - = 'U': Upper triangle of A is stored;
  - = 'L': Lower triangle of A is stored.
- otherwise UPLO = 'U' is assumed.

# INFO - Optional, (output) INTEGER

- If INFO is present:
  - -=0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - $->0\colon$  if INFO=i, the algorithm failed to converge; i form did not converge to zero.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# B.2 LA\_SYEVD / LA\_HEEVD

# **B.2.1** Purpose

**LA\_SYEVD** / **LA\_HEEVD** computes all eigenvalues and, optionally, eigenvectors of a real symmetric or complex Hermitian matrix A. If eigenvectors are desired, it uses a divide and conquer algorithm.

The divide and conquer algorithm makes very mild assumptions about floating point arithmetic. It will work on machines with a guard digit in add/subtract, or on those binary machines without guard digits which subtract like the Cray X-MP, Cray Y-MP, Cray C-90, or Cray-2. It could conceivably fail on hexadecimal or decimal machines without guard digits, but we know of none.

# **B.2.2** Specification

```
SUBROUTINE LA_SYEVD / LA_HEEVD( A, W, JOBZ, UPLO, INFO ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: JOBZ, UPLO INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), INTENT(INOUT) :: A(:,:) type(wp), INTENT(OUT) :: W(:) where type ::= REAL \mid COMPLEX wp ::= KIND(1.0) \mid KIND(1.0D0)
```

#### B.2.3 Defaults

- If JOBZ is not present then JOBZ = 'N' is assumed.
- If UPLO is not present then UPLO = 'U' is assumed.

# B.2.4 Arguments

- **A** (input/output) **REAL** or **COMPLEX** array, shape (:,:).
  - On entry, the symmetric (Hermitian) matrix A.
    - If UPLO = 'U', the upper triangular part of A contains the upper triangular part of the matrix A.
    - If UPLO = L', the lower triangular part of A contains
    - the lower triangular part of the matrix A.
  - On exit:
    - If JOBZ = V', then if INFO = 0, A contains the orthonormal eigenvectors of the matrix A.
    - If JOBZ = 'N', then on exit the lower triangle (if UPLO='L') or the upper triangle (if UPLO='U') of A, including the diagonal, is destroyed.
- $\mathbf{W} (output) \mathbf{REAL} \text{ array, shape (:), size(W)} = \text{size(A,1)} \ge 0.$ 
  - If INFO = 0, the eigenvalues in ascending order.
- JOBZ Optional, (input) CHARACTER\*1
  - If JOBZ is present then:
    - = 'N': Compute eigenvalues only;
    - = 'V': Compute eigenvalues and eigenvectors.
  - otherwise JOBZ = 'N' is assumed.

# **UPLO** – Optional, (input) **CHARACTER\*1**

- If UPLO is present then:
  - = 'U': Upper triangle of A is stored;
  - = 'L': Lower triangle of A is stored.
- otherwise UPLO = 'U' is assumed.

# INFO - Optional, (output) INTEGER

- If INFO is present:
  - = 0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - ->0: if INFO=i, the algorithm failed to converge; i form did not converge to zero.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# B.3 LA\_SYEVX / LA\_HEEVX

# B.3.1 Purpose

**LA\_SYEVX** / **LA\_HEEVX** computes selected eigenvalues and, optionally, eigenvectors of a real symmetric or complex Hermitian matrix A. Eigenvalues and eigenvectors can be selected by specifying either a range of values or a range of indices for the desired eigenvalues.

# **B.3.2** Specification

```
SUBROUTINE LA_SYEVX / LA_HEEVX ( A, W, JOBZ, UPLO, VL, VU, & IL, IU, M, IFAIL, ABSTOL, INFO )

CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: JOBZ, UPLO INTEGER, INTENT(IN), OPTIONAL :: IL, IU INTEGER, INTENT(OUT), OPTIONAL :: INFO, M REAL(wp), INTENT(IN), OPTIONAL :: ABSTOL, VL, VU INTEGER, INTENT(OUT), OPTIONAL :: IFAIL(:) type(wp), INTENT(INOUT) :: A(:,:) REAL(wp), INTENT(OUT) :: W(:) where type ::= REAL \mid COMPLEX \ wp ::= KIND(1.0) \mid KIND(1.0D0)
```

## **B.3.3** Defaults

- If JOBZ and IFAIL are not present then JOBZ = 'N' is assumed.
- If JOBZ is not present and IFAIL is present then JOBZ = 'V' is assumed.
- If UPLO is not present then UPLO = 'U' is assumed.
- If IL or IU are not present and VL or VU are not present then all eigenvalues are computed ( M = size(A,1) ).
- If IL is present and IU is not present then IU = size(A,1) is assumed. If IL is not present and IU is present then IL = 1 is assumed.
- If VL is present and VU is not present then VU = +infinity is assumed. If VL is not present and VU is present then VL = -infinity is assumed.
- If ABSTOL is not present then ABSTOL = 2\*SLAMCH('S') is assumed. In this case the eigenvalues are computed most accurately.

# B.4 Argument dependency

- If either IL or IU are present then neither VL or VU are present.
- If either VL or VU are present then neither IL or IU are present.
- If JOBZ = 'N' then IFAIL must not be present.

# **B.4.1** Arguments

- A (input/output) REAL or COMPLEX array, shape (:,:),  $size(A,1) = size(A,2) \ge 0$ .
  - On entry, the symmetric (Hermitian) matrix A.
    - If UPLO = 'U', the upper triangular part of A contains the upper triangular part of the matrix A.
    - If UPLO = L, the lower triangular part of A contains the lower triangular part of the matrix A.
  - On exit:
    - If JOBZ = 'V', then if INFO = 0, A contains the orthonormal eigenvectors of the matrix A corresponding to the selected eigenvalues, with the i-th column of A holding the eigenvector associated with W(i). If an eigenvector fails to converge, then that column of A contains the latest approximation to the eigenvector, and the index of the eigenvector is returned in IFAIL.

- If JOBZ = 'N', then on exit the lower triangle (if UPLO='L') or the upper triangle (if UPLO='U') of A, including the diagonal, is destroyed.
- $\mathbf{W} (output) \mathbf{REAL} \text{ array, shape (:), size(W)} = \text{size(A,1)} \ge 0.$ 
  - On normal exit, the first M elements contain the selected eigenvalues in ascending order.

# JOBZ - Optional, (input) CHARACTER\*1

- If JOBZ is present then:
  - -='N': Compute eigenvalues only;
  - -='V': Compute eigenvalues and eigenvectors.
- otherwise JOBZ = 'N' is assumed.

# **UPLO** – Optional, (input) **CHARACTER\*1**

- If UPLO is present then:
  - -='U': Upper triangle of A is stored;
  - -='L': Lower triangle of A is stored.
- otherwise UPLO = 'U' is assumed.
- VL Optional, (input) REAL.
- VU Optional, (input) REAL.
  - If VL and VU are present (VL < VU) then the lower and upper bounds of the interval to be searched for eigenvalues. All eigenvalues in the half-open interval (VL, VU] will be found.
- IL Optional, (input) INTEGER.
- IU Optional, (input) INTEGER.
  - If IL and IU are present ( $1 \leq IL \leq IU \leq size(A,1)$ ) then the indices (in ascending order) of the smallest and largest eigenvalues to be returned. The  $IL^{th}$  through  $IU^{th}$  eigenvalues will be found.
- $\mathbf{M} \mathit{Optional}, (\mathit{output}) \ \mathbf{INTEGER}.$ 
  - The total number of eigenvalues found (  $0 \le M \le size(A, 1)$  ).
  - If IL and IU are present then M = IU IL + 1.
- **IFAIL** Optional, (output) **INTEGER** array, shape (:),  $size(IFAIL) = size(A, 1) \ge 0$ .

- If IFAIL is present then JOBZ='V' is assumed and eigenvectors are computed.
  - If INFO=0, the first M elements of IFAIL are zero.
  - If INFO>0, then IFAIL contains the indices of the eigenvectors that failed to converge.

# **ABSTOL** – Optional, (input) **REAL**.

• The absolute error tolerance for the eigenvalues. An approximate eigenvalue is accepted as converged when it is determined to lie in an interval [a,b] of width less than or equal to

$$ABSTOL + EPS \times \max(|a|, |b|)$$

where EPS is the machine precision. If ABSTOL is less than or equal to zero, then  $EPS \times |T|r$  will be used in its place, where |T| is the 1-norm of the tridiagonal matrix obtained by reducing A to tridiagonal form.

- Eigenvalues will be computed most accurately when ABSTOL is set to twice the underflow threshold  $2 \times SLAMCH('S')$ , not zero. If this routine returns with INFO > 0, indicating that some eigenvectors did not converge, try setting ABSTOL to  $2 \times SLAMCH('S')$ .
- See "Computing Small Singular Values of Bidiagonal Matrices with Guaranteed High Relative Accuracy," by Demmel and Kahan, LA-PACK Working Note #3.
- If ABSTOL is not present then  $ABSTOL = 2 \times SLAMCH('S')$  is assumed.

# INFO - Optional, (output) INTEGER

- If INFO is present:
  - -=0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - ->0: if INFO=i, then i eigenvectors failed to converge. Their indices are stored in array IFAIL.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# B.5 LA\_SYGST / LA\_HEGST

### **B.5.1** Purpose

LA\_SYGST / LA\_HEGST reduces a real symmetric-definite or complex Hermitian-definite generalized eigenproblem to standard form.

- If ITYPE=1, the problem is  $Ax=\lambda Bx$ , and A is overwritten by  $(U^H)^{-1}AU^{-1}$  or  $L^{-1}A(L^H)^{-1}$
- If ITYPE = 2 or 3, the problem is  $ABx = \lambda x$  or  $BAx = \lambda x$ , and A is overwritten by  $UAU^H$  or  $L^HAL$ .
- B must have been previously factorized as  $U^HU$  or  $LL^H$  by LA\_POTRF.

### **B.5.2** Specification

```
SUBROUTINE LA_SYGST / LA_HEGST( A, B, ITYPE, UPLO, INFO ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: UPLO INTEGER, INTENT(IN), OPTIONAL :: ITYPE INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), INTENT(IN) :: B(:,:) type(wp), INTENT(INOUT) :: A(:,:) where type ::= REAL \mid COMPLEX wp ::= KIND(1.0) \mid KIND(1.0D0)
```

#### B.5.3 Defaults

- If ITYPE is not present then ITYPE = 1 is assumed.
- If UPLO is not present then UPLO = 'U' is assumed.

### **B.5.4** Arguments

```
A – (input/output) REAL or COMPLEX array, shape (:,:), size(A, 1) = size(A, 2) \ge 0.
```

- On entry, the symmetric (Hermitian) matrix A.
  - If UPLO = U', the upper triangular part of A contains the upper triangular part of the matrix A, and the strictly lower triangular part of A is not referenced.
  - If UPLO = L', the lower triangular part of A contains the lower triangular part of the matrix A, and the strictly upper triangular part of A is not referenced.

• On exit, if INFO = 0, the transformed matrix, stored in the same format as A.

### $\mathbf{B} - (input) \mathbf{REAL} \text{ or } \mathbf{COMPLEX} \text{ array, shape } (:,:), \ size(B,1) = size(A,1).$

• The triangular factor from the Cholesky factorization of B, as returned by LA\_POTRF.

### ITYPE - Optional, (input) INTEGER

- If ITYPE is present then:
  - = 1: compute  $(U^H)^{-1}AU^{-1}$  or  $L^{-1}A(L^H)^{-1}$ ;
  - -=2 or 3: compute  $UAU^H$  or  $L^HAL$ .
- otherwise ITYPE = 1 is assumed.

### **UPLO** – Optional, (input) **CHARACTER\*1**

- If UPLO is present then:
  - -='U': Upper triangle of A is stored and B is factored as  $U^HU$ ;
  - -='L': Lower triangle of A is stored and B is factored as  $LL^H$ .
- otherwise UPLO = 'U' is assumed.

### INFO - Optional, (output) INTEGER

- If INFO is present:
  - = 0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# B.6 LA\_SYGV / LA\_HEGV

## B.6.1 Purpose

**LA\_SYGV** / **LA\_HEGV** computes all the eigenvalues, and optionally, the eigenvectors of a real generalized symmetric-definite or complex Hermitian-definite eigenproblem, of the form  $Ax = \lambda Bx$ ,  $ABx = \lambda x$ , or  $BAx = \lambda x$ . Here A and B are assumed to be symmetric (Hermitian) and B is also positive definite.

### **B.6.2** Specification

```
SUBROUTINE LA_SYGV / LA_HEGV( A, B, W, ITYPE, JOBZ, UPLO, & INFO )

CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: JOBZ, UPLO INTEGER, INTENT(IN), OPTIONAL :: ITYPE INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), INTENT(INOUT) :: A(:,:), B(:,:) REAL(wp), INTENT(OUT) :: W(:) where type ::= REAL \mid COMPLEX \ wp ::= KIND(1.0) \mid KIND(1.0D0)
```

### B.6.3 Defaults

- If ITYPE is not present then ITYPE = 1 is assumed.
- If JOBZ is not present then JOBZ = 'N' is assumed.
- If UPLO is not present then UPLO =' U' is assumed.

### **B.6.4** Arguments

- **A** (input/output) **REAL** or **COMPLEX** array, shape (:,:),  $size(A, 1) = size(A, 2) \ge 0$ .
  - On entry, the symmetric (Hermitian) matrix A.
    - If UPLO = U', the upper triangular part of A contains the upper triangular part of the matrix A.
    - If UPLO = L', the lower triangular part of A contains the lower triangular part of the matrix A.
  - On exit:
    - If JOBZ = V', then if INFO = 0, A contains the matrix Z of eigenvectors. The eigenvectors are normalized as follows:
      - \* if ITYPE = 1 or 2,  $Z^HBZ = I$ ;
      - \* if ITYPE = 3,  $Z^{H}B^{-1}Z = I$ .
    - If JOBZ =' N', then on exit the upper triangle (if UPLO =' U') or the lower triangle (if UPLO =' L') of A, including the diagonal, is destroyed.
- $\mathbf{B} (input) \mathbf{REAL} \text{ or } \mathbf{COMPLEX} \text{ array, shape } (:,:), size(B,1) = size(A,1).$ 
  - On entry, the symmetric (Hermitian) matrix B.

- If UPLO = U', the upper triangular part of B contains the upper triangular part of the matrix B.
- If UPLO = L', the lower triangular part of B contains the lower triangular part of the matrix B.
- On exit, if  $INFO \leq size(A,1)$ , the part of B containing the matrix is overwritten by the triangular factor U or L from the Cholesky factorization  $B = U^H U$  or  $B = LL^H$ .
- $\mathbf{W} (output) \mathbf{REAL} \text{ array, shape (:), } size(W) = size(A, 1).$ 
  - If INFO = 0, the eigenvalues in ascending order.

# ITYPE - Optional, (input) INTEGER.

Specifies the problem type to be solved.

- If ITYPE is present then:
  - = 1:  $Ax = \lambda Bx$
  - -=2:  $ABx = \lambda x$
  - -=3:  $BAx=\lambda x$
- otherwise ITYPE = 1 is assumed.

### JOBZ - Optional, (input) CHARACTER\*1

- If JOBZ is present then:
  - = 'N': Compute eigenvalues only;
  - = 'V': Compute eigenvalues and eigenvectors.
- otherwise JOBZ =' N' is assumed.

### UPLO - Optional, (input) CHARACTER\*1

- If UPLO is present then:
  - = 'U': Upper triangle of A is stored and B is factored as  $U^HU$ ;
  - = 'L': Lower triangle of A is stored and B is factored as  $LL^{H}$ .
- otherwise UPLO = 'U' is assumed.

### INFO - Optional, (output) INTEGER

- If INFO is present:
  - = 0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - ->0: LA\_POTRF or LA\_SYEV / LA\_HEEV returned an error code:

- \*  $\leq size(A, 1)$ : if INFO = i, LA\_SYEV / LA\_HEEV failed to converge; i off-diagonal elements of an intermediate tridiagonal form did not converge to zero;
- \* > size(A, 1): if  $INFO = size(A, 1) + i \le 2size(A, 1)$ , then the leading minor of order i of B is not positive definite. The factorization of B could not be completed and no eigenvalues or eigenvectors were computed.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# B.7 LA\_SYTRD / LA\_HETRD

# B.7.1 Purpose

**LA\_SYTRD** / **LA\_HETRD** reduces a real symmetric or complex Hermitian matrix A to real symmetric tridiagonal form T by an orthogonal or unitary similarity transformation:  $Q^H A Q = T$ .

### **B.7.2** Specification

```
SUBROUTINE LA_SYTRD / LA_HETRD( A, TAU, UPLO, INFO ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: UPLO INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), INTENT(INOUT) :: A(:,:) type(wp), INTENT(OUT) :: TAU(:) where type ::= \text{REAL} \mid \text{COMPLEX}  wp ::= \text{KIND}(1.0) \mid \text{KIND}(1.000)
```

### B.7.3 Defaults

• If UPLO is not present then UPLO = 'U' is assumed.

### B.7.4 Arguments

```
A – (input/output) REAL or COMPLEX array, shape (:,:), size(A, 1) = size(A, 2) \ge 0.
```

- On entry, the symmetric (Hermitian) matrix A.
  - If UPLO = 'U', the upper triangular part of A contains the upper triangular part of the matrix A.
  - If UPLO = L, the lower triangular part of A contains the lower triangular part of the matrix A.

- On exit:
  - If UPLO = 'U', the diagonal and first superdiagonal of A are overwritten by the corresponding elements of the tridiagonal matrix T, and the elements above the first superdiagonal, with the array TAU, represent the unitary matrix Q as a product of elementary reflectors.
  - If UPLO = 'L', the diagonal and first subdiagonal of A are overwritten by the corresponding elements of the tridiagonal matrix T, and the elements below the first subdiagonal, with the array TAU, represent the unitary matrix Q as a product of elementary reflectors.
- See Further Details.

 $\mathbf{TAU} - (output) \mathbf{REAL} \text{ or } \mathbf{COMPLEX} \text{ array, shape (:)}, size(TAU) = size(A, 1) - 1.$ 

- The scalar factors of the elementary reflectors.
- See Further Details.

UPLO - Optional, (input) CHARACTER\*1

- If UPLO is present then:
  - = 'U': Upper triangle of A is stored
  - = 'L': Lower triangle of A is stored
- otherwise UPLO = 'U' is assumed.

INFO - Optional, (output) INTEGER

- If INFO is present:
  - -=0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
- If INFO is not present and an error occurs, then the program is terminated with an error message.

### B.7.5 Further Details

- If UPLO = 'U', the matrix Q is represented as a product of elementary reflectors  $Q = H_{n-1} \cdots H_2 H_1$ . Each  $H_i$  has the form  $H_i = I \tau v v'$ , where  $\tau$  is a complex scalar, and v is a complex vector with  $v_{i+1:n} = 0$  and  $v_i = 1$ ;  $v_{1:i-1}$  is stored on exit in A(1:i-1,i+1), and  $\tau$  in TAU(i).
- If UPLO = 'L', the matrix Q is represented as a product of elementary reflectors  $Q = H_1 H_2 \cdots H_{n-1}$ . Each  $H_i$  has the form  $H_i = I \tau v v'$ , where  $\tau$  is a complex scalar, and v is a complex vector with  $v_{1:i} = 0$  and  $v_{i+1} = 1$ ;  $v_{i+2:n}$  is stored on exit in A(i+2:n,i), and  $\tau$  in TAU(i).

The contents of A on exit are illustrated by the following examples with n=5:

```
if UPL0 = 'U':

( d e v2 v3 v4 )
 ( d e v3 v4 )
 ( e d )
 ( d e v4 )
 ( v1 e d )
 ( v1 v2 e d )
 ( v1 v2 v3 e d )
```

where d and e denote diagonal and off-diagonal elements of T, and  $v_i$  denotes an element of the vector defining  $H_i$ .

# B.8 LA\_ORGTR / LA\_UNGTR

### B.8.1 Purpose

**LA\_ORGTR** / **LA\_UNGTR** generates a real orthogonal / complex unitary matrix Q which is defined as the product of elementary reflectors, as returned by LA\_SYTRD / LA\_HETRD:

```
• if UPLO = 'U', Q = H_{n-1} \cdots H_2 H_1,
```

• if UPLO = 'L', 
$$Q = H_1 H_2 \cdots H_{n-1}$$
.

### B.8.2 Specification

```
SUBROUTINE LA_ORGTR / LA_UNGTR( A, TAU, UPLO, INFO ) CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: UPLO INTEGER, INTENT(OUT), OPTIONAL :: INFO type(wp), INTENT(IN) :: TAU(:) type(wp), INTENT(INOUT) :: A(:,:) where type:= \text{REAL} \mid \text{COMPLEX} wp:= \text{KIND}(1.0) \mid \text{KIND}(1.0\text{D0})
```

### B.8.3 Defaults

• If UPLO is not present then UPLO = 'U' is assumed.

### B.8.4 Arguments

- **A** (input/output) **REAL** or **COMPLEX** array, shape (:,:),  $size(A, 1) = size(A, 2) \ge 0$ .
  - On entry, the vectors which define the elementary reflectors, as returned by LA\_SYTRD or LA\_HETRD.
  - On exit the orthogonal or unitary matrix Q.
- TAU (input) REAL or COMPLEX array, shape (:), size(TAU) = size(A, 1) 1.
  - TAU(i) must contain the scalar factor of the elementary reflector  $H_i$ , as returned by LA\_SYTRD or LA\_HETRD.
- **UPLO** Optional, (input) **CHARACTER\*1** 
  - If UPLO is present then:
    - = 'U': Upper triangle of A is stored
    - = 'L': Lower triangle of A is stored
  - otherwise UPLO = 'U' is assumed.
- INFO Optional, (output) INTEGER
  - If INFO is present:
    - -=0: successful exit
    - -<0: if INFO=-i, the i-th argument had an illegal value
  - If INFO is not present and an error occurs, then the program is terminated with an error message.

# C Cholesky Factorization of a Real Symmetric or Complex Hermitian Positive Definite Matrix A, Documentation

# C.1 LA\_POTRF

### C.1.1 Purpose

**LA\_POTRF** computes the Cholesky factorization of a real symmetric or complex Hermitian positive definite matrix A.

The factorization has the form

- $A = U^H U$ , if UPLO = 'U', or
- $A = LL^H$ , if UPLO = 'L',

where U is an upper triangular matrix and L is lower triangular.

This is the block version of the algorithm, calling Level 3 BLAS.

LA\_POTRF optionally estimates the reciprocal of the condition number (in the 1-norm) of a real symmetric or complex Hermitian positive definite matrix A. An estimate is obtained for  $||A^{-1}||$ , and the reciprocal of the condition number is computed as  $RCOND = 1/||A||||A^{-1}||$ .

### C.1.2 Specification

```
SUBROUTINE LA_POTRF( A, UPLO, RCOND, NORM, INFO )
   CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: NORM, UPLO
   INTEGER, INTENT(OUT), OPTIONAL :: INFO
   REAL(wp), INTENT(OUT), OPTIONAL :: RCOND
   type(wp), INTENT(INOUT) :: A(:,:)
   where
   type ::= REAL \mid COMPLEX
   wp ::= KIND(1.0) \mid KIND(1.0D0)
```

### C.1.3 Defaults

• If UPLO is not present then UPLO = 'U' is assumed.

### C.1.4 Arguments

```
A – (input/output) REAL or COMPLEX array, shape (:,:), size(A, 1) = size(A, 2) \ge 0.
```

• On entry, the symmetric (Hermitian) matrix A.

- If UPLO = 'U', the upper triangular part of A contains the upper triangular part of the matrix A, and the strictly lower triangular part of A is not referenced.
- If UPLO = 'L', the lower triangular part of A contains the lower triangular part of the matrix A, and the strictly upper triangular part of A is not referenced.
- On exit, if INFO=0, the factor U or L from the Cholesky factorization  $A=U^HU$  or  $A=LL^H$ .

# **UPLO** – Optional, (input) **CHARACTER\*1**

- If UPLO is present then:
  - = 'U': Upper triangle of A is stored;
  - = 'L': Lower triangle of A is stored.
- otherwise UPLO = 'U' is assumed.

### RCOND - Optional (output) REAL

• The reciprocal of the condition number of the matrix A computed as  $RCOND = 1/\|A\| \|A^{-1}\|$ .

### NORM - Optional (input) CHARACTER\*1

Specifies whether the 1-norm condition number or the infinity-norm condition number is required:

- $\bullet\,$  If NORM is present then:
  - = '1', 'O' or 'o': 1-norm;
  - = 'I' or 'i': infinity-norm.
- otherwise NORM = '1' is used.

# INFO - Optional, (output) INTEGER

- If INFO is present:
  - = 0: successful exit
  - < 0: if INFO = -i, the i-th argument had an illegal value
  - ->0: if INFO=i, the leading minor of order i is not positive definite, and the factorization could not be completed.
- If INFO is not present and an error occurs, then the program is terminated with an error message.

# D Code for One Version of LA\_SYEV

We illustrate here the sort of code that is needed to implement one of the Fortran 90 jacket procedures. The procedure shown is the real single precision version of LA\_SYEV.

### D.1 Precision-dependencies

To handle different precisions, we use a module LA\_PRECISION to define named constants SP and DP for the kind values of single and double precision, respectively.

```
MODULE LA_PRECISION
INTEGER, PARAMETER :: SP=KIND(1.0), DP=KIND(1.0D0)
END MODULE LA_PRECISION
```

Within the LAPACK 90 code, all real and complex constructs are expressed in terms of a symbolic kind value WP, which is defined by reference to the module LA\_PRECISION — in single precision:

```
USE LA_PRECISION :: WP => SP
and in double precision:
USE LA_PRECISION :: WP => DP
```

These are the only precision-dependent changes in the code, apart from changes to the procedure-names.

# D.2 Error-handling

To handle errors, as described in Section 4, we use a simple procedure ERINFO, which is assumed to be accessed from a module LA\_AUXMOD:

```
SUBROUTINE ERINFO (LINFO, SRNAME, INFO, ISTAT)
  .. Scalar Arguments ..
  CHARACTER( LEN = * ), INTENT(IN)
                                                  :: SRNAME
  INTEGER
                       , INTENT(IN)
                                                  :: LINFO
                       , INTENT(OUT), OPTIONAL
  INTEGER
                                                  :: INFO
  INTEGER
                       , INTENT(IN), OPTIONAL
                                                  :: ISTAT
   .. Executable Statements ..
  IF( ( LINFO < 0 .AND. LINFO > -200 ) .OR. &
       ( LINFO > O .AND. .NOT.PRESENT(INFO) ) )THEN
      WRITE (*,*) 'Program terminated in LAPACK_90 subroutine ',SRNAME
```

```
WRITE (*,*) 'Error indicator, INFO = ',LINFO
     IF ( PRESENT (ISTAT) ) THEN
       IF( ISTAT /= 0 ) THEN
          IF( LINFO == -100 )THEN
            WRITE (*,*) 'The statement ALLOCATE causes STATUS = ', ISTAT
          FLSE
            WRITE (*,*) 'LINFO = ', LINFO, ' not expected'
          END IF
       END IF
     END IF
     STOP
  ELSE IF ( LINFO <= -200 ) THEN
     WRITE(*,*) '*** WARNING, INFO = ', LINFO, ' WARNING ***'
     IF( LINFO == -200 THEN
       WRITE(*,*) 'Could not allocate sufficient workspace for the optimum'
       WRITE(*,*) 'blocksize, hence the routine may not have performed as'
       WRITE(*,*) 'efficiently as possible'
     ELSE
       WRITE(*.*) 'Unexpected warning'
     END IF
     END IF
  IF( PRESENT(INFO) ) INFO = LINFO
END SUBROUTINE ERINFO
```

A more elaborate error-handling mechanism could of course be devised.

### D.3 Accessing LAPACK 77 routines

We assume that interface-blocks for all the LAPACK 77 routines are accessible from modules La\_SF77MOD, La\_DF77MOD, La\_CF77MOD, and La\_ZF77MOD. Note that we do not use generic interfaces for the LAPACK 77 routines, since that would impose some restrictions on the way in which LAPACK 77 routines could be called.

However, we rename the routine in the USE statement, so that the precision-dependent name-change is localized in the USE statement.

## D.4 The code

```
MODULE LA_SSYEV
CONTAINS
!
SUBROUTINE SSYEV_F90( A, W, JOBZ, UPLO, INFO )
```

```
! .. Use Statements ..
  USE LA_PRECISION, ONLY: WP => SP
  USE LA_AUXMOD, ONLY: ERINFO, LSAME
  USE LA_AUF77MOD, ONLY: ILAENV_F77 => ILAENV
  USE LA_SF77MOD, ONLY: SYEV_F77 => SSYEV
! .. Implicit Statement ..
  IMPLICIT NONE
! .. Character Arguments ..
  CHARACTER(LEN=1), INTENT(IN), OPTIONAL :: JOBZ, UPLO
 .. Scalar Arguments ..
  INTEGER, INTENT(OUT), OPTIONAL :: INFO
  .. Array Arguments ..
  REAL(WP), INTENT(INOUT) :: A(:,:)
  REAL(WP), INTENT(OUT) :: W(:)
! .. Local Parameters ..
  CHARACTER(LEN=7), PARAMETER :: SRNAME = 'LA_SYEV'
  CHARACTER(LEN=6), PARAMETER :: BSNAME = 'SSYTRD'
! .. Local Scalars ..
  CHARACTER(LEN=1) :: LJOBZ, LUPLO
  INTEGER :: N, LINFO, LD, ISTAT, ISTAT1, LWORK, NB
! .. Local Arrays ..
  REAL(WP), POINTER :: WORK(:)
! .. Intrinsic Functions ..
  INTRINSIC MAX, PRESENT
! .. Executable Statements ...
  N = SIZE(A, 1); LINFO = 0; ISTAT = 0; LD = MAX(1,N)
  IF( PRESENT(JOBZ) ) THEN
     LJOBZ = JOBZ
  ELSE
     LJOBZ = 'N'
  END IF
  IF ( PRESENT (UPLO) ) THEN
     LUPLO = UPLO
  ELSE
     LUPLO = 'U'
  END IF
! .. Test the arguments
  IF( SIZE( A, 2 ) /= N .OR. N < O ) THEN
     LINFO = -1
  ELSE IF( SIZE( W ) /= N )THEN
     LINFO = -2
  ELSE IF ( .NOT.LSAME(LJOBZ,'N') .AND. .NOT.LSAME(LJOBZ,'V') )THEN
     LINFO = -3
  ELSE IF( .NOT.LSAME(LUPLO,'U') .AND. .NOT.LSAME(LUPLO,'L') )THEN
```

```
LINFO = -4
   ELSE IF( N > 0 )THEN
  .. Determine the workspace
      NB = ILAENV_F77(1, BSNAME, LUPLO, N, -1, -1, -1)
      IF ( NB \le 1 .0R. NB \ge N ) THEN
         NB = 1
      END IF
      LWORK = (2+NB)*N
      ALLOCATE(WORK(LWORK), STAT=ISTAT)
      IF( ISTAT /= 0 )THEN
         LWORK = 3*N-1
         ALLOCATE(WORK(LWORK), STAT=ISTAT)
         IF( ISTAT \neq 0 ) THEN
            LINFO = -100
         ELSE
            LINFO = -200
         ENDIF
      ENDIF
ļ
      IF( LINFO == 0 .OR. LINFO <= -200 )THEN
         .. Call LAPACK77 routine
!
         CALL SYEV_F77 ( LJOBZ, LUPLO, N, A, LD, W, WORK, LWORK, LINFO )
      DEALLOCATE(WORK, STAT=ISTAT1)
   ENDIF
   CALL ERINFO (LINFO, SRNAME, INFO, ISTAT)
END SUBROUTINE SSYEV_F90
END MODULE LA_SSYEV
```

# D.5 Accessing LAPACK 90 procedures

We assume that interface-blocks (module-procedures) for all the LAPACK 90 routines are accessible from module LA\_SCDZF90M0D.