NAG Fortran Library Routine Document

F02FJF

Note: before using this routine, please read the Users' Note for your implementation to check the interpretation of **bold italicised** terms and other implementation-dependent details.

1 Purpose

To find eigenvalues and eigenvectors of a real sparse symmetric or generalized symmetric eigenvalue problem.

2 Specification

3 Description

F02FJF finds the m eigenvalues of largest absolute value and the corresponding eigenvectors for the real eigenvalue problem

$$
Cx = \lambda x \tag{1}
$$

where C is an n by n matrix such that

$$
BC = C^T B \tag{2}
$$

for a given positive-definite matrix B . C is said to be B -symmetric. Different specifications of C allow for the solution of a variety of eigenvalue problems. For example, when

 $C = A$ and $B = I$ where $A = A^T$

the routine finds the m eigenvalues of largest absolute magnitude for the standard symmetric eigenvalue problem

$$
Ax = \lambda x.\tag{3}
$$

The routine is intended for the case where A is sparse.

As a second example, when

 $C = B^{-1}A$

where

$$
A = A^T
$$

the routine finds the m eigenvalues of largest absolute magnitude for the generalized symmetric eigenvalue problem

$$
Ax = \lambda Bx.\tag{4}
$$

The routine is intended for the case where A and B are sparse.

The routine does not require C explicitly, but C is specified via a user-suppli[ed routine IMAGE which,](#page-3-0) given an n element vector z , computes the image w given by

$$
w=Cz.
$$

For instance, in the above example, where $C = B^{-1}A$ [, routine IMAGE will nee](#page-3-0)d to solve the positivedefinite system of equations $Bw = Az$ for w.

To find the m eigenvalues of smallest absolute magnitu[de of \(3\) we](#page-0-0) can choose $C = A^{-1}$ and hence find the reciprocals of the required eigenvalues[, so that IMAGE will nee](#page-3-0)d to solve $Aw = z$ for w, and correspondingl[y for \(4\) we](#page-0-0) can choose $C = A^{-1}B$ and solve $Aw = Bz$ for w.

A table of examples of [choice of IMAGE is given](#page-3-0) in Table 1. It should be remembered that the routine also returns the corresponding eigenvectors and that B is positive-definite. Throughout A is assumed to be symmetric and, where necessary, non-singularity is also assumed.

Eigenvalues	Problem		
Required	$Ax = \lambda x$ $(B = I)$	$Ax = \lambda Bx$	$ABx = \lambda x$
Largest	Compute $w = Az$	Solve $Bw = Az$	Compute $w = ABz$
Smallest (Find $1/\lambda$)	Solve $Aw = z$	Solve $Aw = Bz$	Solve $Av = z$, $Bw = v$
Furthest from σ (Find $\lambda - \sigma$)	Compute $w = (A - \sigma I)z$	Solve $Bw = (A - \sigma B)z$	Compute $w = (AB - \sigma I)z$
Closest to σ (Find $1/(\lambda - \sigma)$)	Solve $(A - \sigma I)w = z$	Solve $(A - \sigma B)w = Bz$	Solve $(AB - \sigma I)w = z$

Table 1 The Requi[rement of IMAGE for Vario](#page-3-0)us Problems.

The matrix B also need not be supplied explicitly, but is specified via a user-supplied [routine DOT which](#page-2-0), given *n* element vectors *z* and *w*, computes the generalized dot product $w^T B z$.

F02FJF is based upon routine SIMITZ (see Nikolai (1979)), which is itself a derivative of the Algol procedure ritzit (see Rutishauser (1970)), and uses the method of simultaneous (subspace) iteration. (See Parlett (1980) for description, analysis and advice on the use of the method.)

The routine performs simultaneous iteration on $k > m$ vectors. Initial estimates to $p \leq k$ eigenvectors, corresponding to the p eigenvalues of C of largest absolute value, may be supplied by the user to F02FJF. When possible k should be chosen so that the kth eigenvalue is not too close to the m required eigenvalues, but if k is initially chosen too small then F02FJF may be re-entered, supplying approximations to the k eigenvectors found so far and with k then increased.

At each major iteration F02FJF solves an r by r $(r \leq k)$ eigenvalue sub-problem in order to obtain an approximation to the eigenvalues for which convergence has not yet occurred. This approximation is refined by Chebyshev acceleration.

4 References

Nikolai P J (1979) Algorithm 538: Eigenvectors and eigenvalues of real generalized symmetric matrices by simultaneous iteration ACM Trans. Math. Software 5 118-125

Parlett B N (1980) The Symmetric Eigenvalue Problem Prentice-Hall

Rutishauser H (1969) Computational aspects of F L Bauer's simultaneous iteration method Numer. Math. 13 4–13

Rutishauser H (1970) Simultaneous iteration method for symmetric matrices Numer. Math. 16 205–223

5 Parameters

1: $N - INTEGR$ *Input*

On entry: n , the order of the matrix C .

Constraint: $N > 1$.

2: M – INTEGER *Input/Output*

On entry: m , the number of eigenvalues required.

Constraint: $M \geq 1$.

On exit: m', the number of eigenvalues actually found. It is equal to m if IF[AIL](#page-6-0) = 0 on exit, and is less than m if IF[AIL](#page-6-0) $= 2$, 3 o[r 4. See Section 6 and Sect](#page-6-0)[ion 8 for furthe](#page-7-0)r information.

3: K – INTEGER *Input*

On entry: the number of simultaneous iteration vectors to be used. Too small a value of K may inhibit convergence, while a larger value of K incurs additional storage and additional work per iteration.

Suggested value: $K = M + 4$ will often be a reasonable choice in the absence of better information. Constraint: $M < K \leq N$.

4: NOITS – INTEGER Input/Output

On entry: the maximum number of major iterations (eigenvalue sub-problems) to be performed. If NOITS \leq 0, then the value 100 is used in place of NOITS.

On exit: the number of iterations actually performed.

$$
5: \qquad \text{TOL} - \textit{real} \qquad \qquad \text{Input}
$$

On entry: a relative tolerance to be used in accepting eigenvalues and eigenvectors. If the eigenvalues are required to about t significant figures, then TOL should be set to about 10^{-t} . d_i is accepted as an eigenvalue as soon as two successive approximations to d_i differ by less than $(|\tilde{d}_i| \times \text{TOL})/10$, where \tilde{d}_i is the latest approximation to d_i . Once an eigenvalue has been accepted, then an eigenvector is accepted as soon as $(d_i f_i)/(d_i - d_k) < \text{TOL}$, where f_i is the normalised residual of the current approximation to the eigenv[ector \(see Section 8 for furthe](#page-7-0)r information). The values of the f_i and d_i can be printed fro[m routine MONIT. If TOL](#page-4-0) is supplied outside the range $(\epsilon, 1.0)$, where ϵ is the *machine precision*, then the value ϵ is used in place of TOL.

6: DOT – real FUNCTION, supplied by the user. External Procedure

DOT must return the value $w^T B z$ for given vectors w and z. For the standard eigenvalue problem, where $B = I$, DOT must return the dot product $w^T z$.

Its specification is:

real FUNCTION DOT(IFLAG, N, Z, W, RWORK, LRWORK, IWORK, LIWORK) INTEGER IFLAG, N, LRWORK, IWORK(LIWORK), LIWORK $real$ $Z(N), W(N), RWORK(LRWORK)$

1: IFLAG – INTEGER *Input/Output*

On entry: IFLAG is always non-negative.

On exit: IFLAG may be used as a flag to indicate a failure in the computation of $w^T B z$. If IFLAG is negative on exit from DOT, then F02FJF will exit immediate[ly with IFAIL set](#page-6-0) to IFLAG. Note that in this case DOT must still be assigned a value.

DOT must be declared as EXTERNAL in the (sub)program from which F02FJF is called. Parameters denoted as Input must not be changed by this procedure.

7: IMAGE – SUBROUTINE, supplied by the user. External Procedure IMAGE must return the vector $w = Cz$ for a given vector z.

Its specification is:

SUBROUTINE IMAGE(IFLAG, N, Z, W, RWORK, LRWORK, IWORK, LIWORK) INTEGER IFLAG, N, LRWORK, IWORK(LIWORK), LIWORK $real$ $Z(N)$, $W(N)$, RWORK(LRWORK)

1: IFLAG – INTEGER *Input/Output*

On entry: IFLAG is always non-negative.

On exit: IFLAG may be used as a flag to indicate a failure in the computation of w . If IFLAG is negative on exit from IMAGE, then F02FJF will exit immediate[ly with IFAIL](#page-6-0) set to IFLAG.

2: $N - INTEGR$ Input

On entry: n, the number of elements in the vectors w and z, and the order of the matrix C .

- $3: Z(N)$ real array Input On entry: the vector z for which Cz is required.
- 4: W(N) real array Output

On exit: the vector $w = Cz$.

5: RWORK(LRWORK) – real array User Workspace 6: LRWORK – INTEGER *Input* 7: IWORK(LIWORK) – INTEGER array User Workspace 8: LIWORK – INTEGER *Input*

IMAGE is called from F02FJF with the parameters RWORK, LRWORK, IWORK and LIWORK as supplied to F02FJF. The user is free to use the arrays RWORK and IWORK to supply information to IMA[GE and DOT as an](#page-2-0) alternative to using COMMON.

IMAGE must be declared as EXTERNAL in the (sub)program from which F02FJF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

8: MONIT – SUBROUTINE, supplied by the user. External Procedure

MONIT is used to monitor the progress of F02FJF. MONIT may be the dummy subroutine F02FJZ if no monitoring is actually required. (F02FJZ is included in the NAG Fortran Library and so need not be supplied by the user. The routine name F02FJZ may be implementation dependent: see the Users' Note for your implementation for details.) MONIT is called after the solution of each eigenvalue sub-problem and also just prior to return from F02FJF. The parameters ISTATE and NEXTIT allow selective printing by MONIT.

Its specification is:

SUBROUTINE MONIT(ISTATE, NEXTIT, NEVALS, NEVECS, K, F, D) INTEGER ISTATE, NEXTIT, NEVALS, NEVECS, K

F(K), D(K) $F(K)$, $D(K)$ 1: ISTATE – INTEGER Input On entry: ISTATE specifies the state of F02FJF and will have values as follows: $ISTATE = 0$ No eigenvalue or eigenvector has just been accepted. $ISTATE = 1$ One or more eigenvalues have been accepted since the last call to MONIT. $ISTATE = 2$ One or more eigenvectors have been accepted since the last call to MONIT. $ISTATE = 3$ One or more eigenvalues and eigenvectors have been accepted since the last call to MONIT. $ISTATE = 4$ Return from F02FJF is about to occur. 2: NEXTIT – INTEGER Input On entry: the number of the next iteration. 3: NEVALS – INTEGER *Input* On entry: the number of eigenvalues accepted so far. 4: NEVECS – INTEGER *Input* On entry: the number of eigenvectors accepted so far. 5: K – INTEGER *Input* On entry: k, the number of simultaneous iteration vectors. 6: $F(K)$ – real array Input On entry: a vector of error quantities measuring the state of convergence of the simultaneous iteration vectors. See the par[ameter TOL of F0](#page-2-0)2FJF [above and Section 8 for](#page-7-0) further details. Each element of F is initially set to the value 4.0 and an element remains at 4.0 until the corresponding vector is tested.

7: $D(K)$ – real array Input On entry: $D(i)$ contains the latest approximation to the absolute value of the *i*th eigenvalue of C.

MONIT must be declared as EXTERNAL in the (sub)program from which F02FJF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

9: NOVECS – INTEGER *Input*

On entry: the number of approximate vectors that are being supplied in X. If NOVECS is outside the range $(0,K)$, then the value 0 is used in place of NOVECS.

10: $X(NRX,K)$ – real array Input/Output Input/Output

On entry: if $0 <$ NOVECS \leq [K, t](#page-2-0)he first NOVECS columns of X must contain approximations to the eigenvectors corresponding to the NOVECS eigenvalues of largest absolute value of C. Supplying approximate eigenvectors can be useful when reasonable approximations are known, or when the routine is being restarted with a larger valu[e of K. O](#page-2-0)therwise it is not necessary to supply approximate vectors, as simultaneous iteration vectors will be generated randomly by the routine.

On exit: if IF[AIL](#page-6-0) $= 0, 2, 3$ or 4, the first m' columns contain the eigenvectors corresponding to the eigenvalues returned in the first m' elements of D (see below); and the next $k - m' - 1$ columns contain approximations to the eigenvectors corresponding to the approximate eigenvalues returned in the next $k - m' - 1$ elements of D. Here m' is the value returne[d in M \(se](#page-2-0)e above), the number of eigenvalues actually found. The kth column is used as workspace.

11: NRX – INTEGER *Input*

On entry: the first dimension of the array X as declared in the (sub)program from which F02FJF is called.

Constraint: $NRX > N$.

12: $D(K)$ – real array Output

On exit: if I[FAIL](#page-6-0) = 0, 2, 3 or 4, the first m' elements contain the first m' eigenvalues in decreasing order of magnitude; and the next $k - m² - 1$ elements contain approximations to the next $k - m' - 1$ eigenvalues. Here m' is the value returne[d in M \(se](#page-2-0)e above), the number of eigenvalues actually found. D(k) contains the value e where $(-e, e)$ is the latest interval over which Chebyshev acceleration is performed.

- 13: WORK(LWORK) real array Workspace Workspace
- 14: LWORK INTEGER *Input*

On entry: the dimension of the array WORK as declared in the (sub)program from which F02FJF is called.

Constraint: LWOR[K](#page-2-0) $\geq 3 \times K + \max(K \times K, 2 \times N)$ $\geq 3 \times K + \max(K \times K, 2 \times N)$ $\geq 3 \times K + \max(K \times K, 2 \times N)$.

15: RWORK(LRWORK) – real array User Workspace

RWORK is not used by F02FJF, but is passed directly to r[outines DOT and I](#page-2-0)[MAGE and may](#page-3-0) be used to supply information to these routines.

16: LRWORK – INTEGER *Input*

On entry: the dimension of the array RWORK as declared in the (sub)program from which F02FJF is called.

Constraint: LRWORK ≥ 1 .

17: IWORK(LIWORK) – INTEGER array User Workspace

IWORK is not used by F02FJF, but is passed directly to r[outines DOT and](#page-2-0) [IMAGE and may](#page-3-0) be used to supply information to these routines.

18: LIWORK – INTEGER *Input*

On entry: the dimension of the array IWORK as declared in the (sub)program from which F02FJF is called.

Constraint: LIWORK > 1 .

19: IFAIL – INTEGER *Input/Output*

On entry: IFAIL must be set to $0, -1$ or 1. Users who are unfamiliar with this parameter should refer to Chapter P01 for details.

On exit: IFAIL $= 0$ unless the routine detects an error (see Section 6).

For environments where it might be inappropriate to halt program execution when an error is detected, the value -1 or 1 is recommended. If the output of error messages is undesirable, then the value 1 is recommended. Otherwise, because for this routine the values of the output parameters may be useful even if IFAIL $\neq 0$ on exit, the recommended value is -1 . When the value -1 or 1 is used it is essential to test the value of IFAIL on exit.

6 Error Indicators and Warnings

If on entry IFAIL $= 0$ or -1 , explanatory error messages are output on the current error message unit (as defined by X04AAF).

Errors or warnings detected by the routine:

IFAIL < 0

A negative value of IFAIL indicates an exit from F02FJF because the user has set IFLAG negative [in DOT or I](#page-2-0)[MAGE. The va](#page-3-0)lue of IFAIL will be the same as the user's setting of IFLAG.

$IFAIL = 1$

$IFAIL = 2$

Not all the requested eigenvalues and vectors have been obtained. Approximations to the rth eigenvalue are oscillating rapidly indicating that severe cancellation is occurring in the rth eigenvector and [so M is](#page-2-0) returned as $(r-1)$. A restart with a larger value [of K m](#page-2-0)ay permit convergence.

$IFAIL = 3$

Not all the requested eigenvalues and vectors have been obtained. The rate of convergence of the remaining eigenvectors suggests that [more than NOITS iteration](#page-2-0)s would be required and so the input valu[e of M has](#page-2-0) been reduced. A restart with a larger value [of K m](#page-2-0)ay permit convergence.

 $IFAIL = 4$

Not all the requested eigenvalues and vectors have been [obtained. NOITS iterations](#page-2-0) have been performed. A restart, possibly with a larger value [of K, m](#page-2-0)ay permit convergence.

 $IFAIL = 5$

This error is very unlikely to occur, but indicates that convergence of the eigenvalue sub-problem has not taken place. Restarting with a different set of approximate vectors may allow convergence. If this error occurs the user should check carefully that F02FJF is being called correctly.

7 Accuracy

Eigenvalues and eigenvectors will normally be computed to the accuracy requested by the par[ameter TOL,](#page-2-0) but eigenvectors corresponding to small or to close eigenvalues may not always be computed to the accuracy requested by the para[meter TOL. Us](#page-2-0)e of th[e routine MONIT to mon](#page-4-0)itor acceptance of eigenvalues and eigenvectors is recommended.

8 Further Comments

The time taken by the routine will be principally determined by the time taken to solve the eigenvalue subproblem and the time taken by the r[outines DOT and I](#page-2-0)[MAGE. The tim](#page-3-0)e taken to solve an eigenvalue subproblem is approximately proportional to nk^2 . It is important to be aware that several [calls to DOT and](#page-2-0) [IMAGE may oc](#page-3-0)cur on each major iteration.

As can be seen from Table 1, many applications of F02FJF will requi[re routine IMAGE to solve](#page-3-0) a system of linear equations. For example, to find the smallest eigenvalues of $Ax = \lambda Bx$ [, IMAGE needs to](#page-3-0) solve equations of the form $Aw = Bz$ for w and routines from Chapter F01 and Chapter F04 of the NAG Fortran Library will frequently be useful in this context. In particular, if A is a positive-definite variable band matrix, F04MCF may be used after A has been factorized by F01MCF. Thus factorization need be performed only once prior to calling F02FJF. An illustration of this type of use is given in the example program.

An approximation \tilde{d}_h , to the *i*th eigenvalue, is accepted as soon as \tilde{d}_h and the previous approximation differ by less than $|\tilde{d}_h| \times \text{TOL}/10$ $|\tilde{d}_h| \times \text{TOL}/10$ $|\tilde{d}_h| \times \text{TOL}/10$. Eigenvectors are accepted in groups corresponding to clusters of eigenvalues that are equal, or nearly equal, in absolute value and that have already been accepted. If d_r is the last eigenvalue in such a group and we define the residual r_i as

$$
r_j = Cx_j - y_r
$$

where y_r is the projection of Cx_j , with respect to B, onto the space spanned by x_1, x_2, \ldots, x_r , and x_j is the current approximation to the jth eigenvector, then the value f_i r[eturned in MONIT is given](#page-4-0) by

$$
f_i = \max ||r_j||_B / ||Cx_j||_B \quad ||x||_B^2 = x^T B x
$$

and each vector in the group is accepted as an eigenvector if

$$
(|d_r|f_r)/(|d_r|-e) < \text{TOL},
$$

where e is the current approximation to $|\tilde{d}_k|$. The values of the f_i are systematically increased if the convergence criteria appear t[o be too strict. See Rutishauser \(1970\) for further details.](#page-1-0)

The algorithm implemented by F02FJF differs slightly fr[om SIMITZ \(Nikolai \(1979\)\) in that the](#page-1-0) eigenvalue sub-problem is solved using the singular value decomposition of the upper triangular matrix R of the Gram–Schmidt factorization of Cx_r , rather than forming $R^T R$.

9 Example

To find the four eigenvalues of smallest absolute value and corresponding eigenvectors for the generalized symmetric eigenvalue problem $Ax = \lambda Bx$, where A and B are the 16 by 16 matrices

[TOL is tak](#page-2-0)en as 0.0001 and 6 iteration vectors are used. F11JAF is used to factorize the matrix A, prior to calling F02FJF, and F11JCF is used [within IMAGE to solve](#page-3-0) the equations $Aw = Bz$ for w. Details of the factorization of A are passed from F11JAF to F11JCF by means of the COMMON block BLOCK1.

Ou[tput from MONIT occurs e](#page-4-0)ach time ISTATE is non-zero. Note that the required eigenvalues are the reciprocals of the eigenvalues returned by F02FJF.

9.1 Program Text

Note: the listing of the example program presented below uses **bold italicised** terms to denote precision-dependent details. Please read the Users' Note for your implementation to check the interpretation of these terms. As explained in the Essential Introduction to this manual, the results produced may not be identical for all implementations.

```
* F02FJF Example Program Text
* Mark 20 Revised. NAG Copyright 2001.
     .. Parameters ..<br>INTEGER
                      NMAX, LA, LRWORK, KMAX, LWORK, LIWORK, NRX
     PARAMETER (NMAX=16,LA=10*NMAX,LRWORK=1,KMAX=6,
    + LWORK=5*KMAX+2*NMAX,LIWORK=2*LA+7*NMAX+1,<br>+ NRX=NMAX)
                      NRX=NMAX)
     INTEGER NIN, NOUT
     PARAMETER (NIN=5, NOUT=6)
* .. Scalars in Common ..
     INTEGER
* .. Arrays in Common ..<br>
real A(LA)A(LA)INTEGER ICOL(LA), IPIV(NMAX), IROW(LA), ISTR(NMAX+1)
```

```
* .. Local Scalars ..
     real DSCALE, DTOL, TOL<br>INTEGER I TEATL J. K. L
                     I, IFAIL, J, K, L, LFILL, M, N, NNZC, NOITS,
    + NOVECS, NPIVM
     CHARACTER MIC, PSTRAT
     .. Local Arrays ..<br>real
     real<br>
D(NMAX), RWORK(LRWORK), WORK(LWORK), X(NRX,KMAX)
                     IWORK(LIWORK)
* .. External Functions ..<br>
real DOT
     real DOT
     EXTERNAL DOT
* .. External Subroutines ..
     EXTERNAL F02FJF, F02FJZ, F11JAF, IMAGE
* .. Common blocks ..
     COMMON /BLOCK1/A, IROW, ICOL, IPIV, ISTR, NNZ
* .. Executable Statements ..
     WRITE (NOUT,*) 'F02FJF Example Program Results'
     Skip heading in data file
     READ (NIN,*)
     READ (NIN,*) N, M, K, TOL
     WRITE (NOUT,*)
     IF (N.LT.5 .OR. N.GT.16) THEN
        WRITE (NOUT, 99999) 'N is out of range. N = ', N
     ELSE IF (M.LT.1 .OR. M.GE.K .OR. K.GT.KMAX) THEN
        WRITE (NOUT, 99999) 'M or K out of range. M =', M, ' K =', K
     ELSE
*
* Set up the sparse symmetric coefficient matrix A.
\mathbf{r}L = 0DO 20 I = 1, N
           IF (I.GE.5) THEN
             L = L + 1A(L) = -0.25e0IROW(L) = IICOL(L) = I - 4END IF
           IF (I.GE.2) THEN
             L = L + 1A(L) = -0.25e0IROW(L) = IICOL(L) = I - 1END TF
           L = L + 1A(L) = 1.0e0IROW(L) = IICOL(L) = I20 CONTINUE
        NNZ = L*
        Call F11JAF to find an incomplete Cholesky factorisation of A.
*
        LFTLL = 2DTOL = 0.0e0MIC = 'M'DSCALE = 0.0e0PSTRAT = 'M'IFAIL = 1
*
        CALL F11JAF(N,NNZ,A,LA,IROW,ICOL,LFILL,DTOL,MIC,DSCALE,PSTRAT,
    + IPIV,ISTR,NNZC,NPIVM,IWORK,LIWORK,IFAIL)
*
        IF (IFAIL.NE.0) THEN
          WRITE (NOUT.99999) 'F11JAF fails. IFAIL =', IFAIL
        ELSE
*
* Call F02FJF to find eigenvalues and eigenvectors.
           IFAIL = 1
* * To obtain monitoring information from the supplied
* subroutine MONIT, replace the name F02FJZ by MONIT in
           the next statement, and declare MONIT as external *
```

```
*
            NOITS = 1000
            NOVECS = 0*
            CALL F02FJF(N,M,K,NOITS,TOL,DOT,IMAGE,F02FJZ,NOVECS,X,NRX,D,
     + WORK,LWORK,RWORK,LRWORK,IWORK,LIWORK,IFAIL)
\overline{u}IF (IFAIL.NE.0) THEN
                WRITE (NOUT, 99999) 'Warning - FO2FJF returns IFAIL =',
     + IFAIL
            END IF
            IF (IFAIL.GE.0 .AND. IFAIL.NE.1 .AND. IFAIL.LE.4 .AND. M.GE.
     + 1) THEN
               DO 40 I = 1, MD(I) = 1.0e0/D(I)40 CONTINUE
               WRITE (NOUT,*) 'Final results'
               WRITE (NOUT,*)
               WRITE (NOUT, *) ' Eigenvalues'
               WRITE (NOUT,99998) (D(I),I=1,M)
               WRITE (NOUT,*)
               WRITE (NOUT, *) ' Eigenvectors'
               WRITE (NOUT, 99998) (X(I,J),J=1,N),I=1,N)END IF
         END IF
      END IF
      STOP
*
99999 FORMAT (1X,A,I5,A,I5)
99998 FORMAT (1X,1P,4e12.3)
      END
*
      real FUNCTION DOT(IFLAG, N, Z, W, RWORK, LRWORK, IWORK, LIWORK)
* This function implements the dot product - transpose(W)*B*Z.<br>* DOT assumes that N is at least 3.
* DOT assumes that N is at least 3.<br>* Scalar Arguments
      .. Scalar Arguments ..<br>INTEGER IFLA
                         IFLAG, LIWORK, LRWORK, N
* .. Array Arguments ..<br>real RWC
                         RWORK(LRWORK), W(N), Z(N)
      INTEGER IWORK(LIWORK)
* .. Local Scalars ..
      real S<br>INTEGER T
      INTEGER I
      .. Executable Statements ..
      S = 0.0e0S = S + (Z(1)-0.5e0*Z(2))*W(1)S = S + (-0.5e0*Z(N-1)+Z(N))*W(N)DO 20 I = 2, N - 1S = S + (-0.5e0*Z(I-1)+Z(I)-0.5e0*Z(I+1))*W(I)20 CONTINUE
      DOT = SRETURN
      END
*
      SUBROUTINE IMAGE(IFLAG, N, Z, W, RWORK, LRWORK, IWORK, LIWORK)
* This routine solves A*W = B*Z for W.<br>* The routine assumes that N is at least
* The routine assumes that N is at least 3.
      A, IROW, ICOL, IPIV, ISTR and NNZ must be as returned by routine
* F11JAF.
* .. Parameters ..
                      NMAX, LA, LWORK
      PARAMETER (NMAX=16,LA=10*NMAX,LWORK=6*NMAX+120)
      .. Scalar Arguments ..<br>INTEGER IFLAG
                       IFLAG, LIWORK, LRWORK, N
* .. Array Arguments ..
      real RWORK(LRWORK), W(N), Z(N)<br>TNTEGER TWORK(LIWORK)
                        IWORK(LIWORK)
* .. Scalars in Common ..
      INTEGER
* .. Arrays in Common ..<br>real a(LA)
                        A(LA)
```

```
INTEGER ICOL(LA), IPIV(NMAX), IROW(LA), ISTR(NMAX+1)
* .. Local Scalars ..
     real RNORM, TOL<br>INTEGER IFAIL, ITN
                      IFAIL, ITN, J, MAXITN
     CHARACTER*2 METHOD
     .. Local Arrays ..<br>real
                      RHS(NMAX), WORK(LWORK)
     .. External Functions ..<br>real x02AJF
                      X02AJF<br>X02AJF
     EXTERNAL
* .. External Subroutines ..
     EXTERNAL
* .. Common blocks ..
                       /BLOCK1/A, IROW, ICOL, IPIV, ISTR, NNZ
* .. Executable Statements ..
*
     Form B*Z in RHS and initialize W to zero.
*
     RHS(1) = Z(1) - 0.5e^{0*Z(2)}W(1) = 0.0e0RHS(N) = -0.5e0*Z(N-1) + Z(N)W(N) = 0.0e0DO 20 J = 2, N - 1RHS(J) = -0.5e0 \times Z(J-1) + Z(J) - 0.5e0 \times Z(J+1)W(J) = 0.0e020 CONTINUE
*
* Call F11JCF to solve the equations A*W=B*Z.
\mathbf{r}METHOD = 'CG'TOL = X02AJF()MAXITN = 100IFAIL = 1
*
     CALL F11JCF(METHOD,N,NNZ,A,LA,IROW,ICOL,IPIV,ISTR,RHS,TOL,MAXITN,
     + W,RNORM,ITN,WORK,LWORK,IFAIL)
*
     IF (IFAIL.GT.0) IFLAG = -IFAIL
     RETURN
     END
*
     SUBROUTINE MONIT(ISTATE, NEXTIT, NEVALS, NEVECS, K, F, D)
* Monitoring routine for F02FJF.
      .. Parameters ..<br>INTEGER NOUT
     INTEGER
     PARAMETER (NOUT=6)
* .. Scalar Arguments ..
                      ISTATE, K, NEVALS, NEVECS, NEXTIT
* .. Array Arguments ..
                     D(K), F(K)* .. Local Scalars ..
     INTEGER I
      .. Executable Statements ..
     IF (ISTATE.NE.0) THEN
        WRITE (NOUT,*)
         WRITE (NOUT,99999) ' ISTATE = ', ISTATE, ' NEXTIT = ', NEXTIT
         WRITE (NOUT,99999) ' NEVALS = ', NEVALS, ' NEVECS = ', NEVECS
        WRITE (NOUT, \star) ' F D'
        WRITE (NOUT, 99998) (F(I),D(I),I=1,K)END IF
     RETURN
*
99999 FORMAT (1X,A,I4,A,I4)
99998 FORMAT (1X, 1P, e11.3, 3X, e11.3)
     END
```
9.2 Program Data

```
F02FJF Example Program Data
16 4 6 0.0001
```
9.3 Program Results

F02FJF Example Program Results

Final results

