

## NAG Library Function Document

### nag\_real\_sparse\_eigensystem\_iter (f12abc)

**Note:** this function uses **optional arguments** to define choices in the problem specification. If you wish to use default settings for all of the optional arguments, then the option setting function `nag_real_sparse_eigensystem_option` (f12adc) need not be called. If, however, you wish to reset some or all of the settings please refer to Section 11 in `nag_real_sparse_eigensystem_option` (f12adc) for a detailed description of the specification of the optional arguments.

#### 1 Purpose

`nag_real_sparse_eigensystem_iter` (f12abc) is an iterative solver used to find some of the eigenvalues (and optionally the corresponding eigenvectors) of a standard or generalized eigenvalue problem defined by real nonsymmetric matrices. This is part of a suite of functions that also includes `nag_real_sparse_eigensystem_init` (f12aac), `nag_real_sparse_eigensystem_sol` (f12acc), `nag_real_sparse_eigensystem_option` (f12adc) and `nag_real_sparse_eigensystem_monit` (f12aec). It is

#### 2 Specification

```
#include <nag.h>
#include <nagf12.h>

void nag_real_sparse_eigensystem_iter (Integer *irevcm, double resid[],
    double v[], double **x, double **y, double **mx, Integer *nshift,
    double comm[], Integer icomm[], NagError *fail)
```

#### 3 Description

The suite of functions is designed to calculate some of the eigenvalues,  $\lambda$ , (and optionally the corresponding eigenvectors,  $x$ ) of a standard eigenvalue problem  $Ax = \lambda x$ , or of a generalized eigenvalue problem  $Ax = \lambda Bx$  of order  $n$ , where  $n$  is large and the coefficient matrices  $A$  and  $B$  are sparse, real and nonsymmetric. The suite can also be used to find selected eigenvalues/eigenvectors of smaller scale dense, real and nonsymmetric problems.

`nag_real_sparse_eigensystem_iter` (f12abc) is a **reverse communication** function, based on the ARPACK routine **dnaupd**, using the Implicitly Restarted Arnoldi iteration method. The method is described in Lehoucq and Sorensen (1996) and Lehoucq (2001) while its use within the ARPACK software is described in great detail in Lehoucq *et al.* (1998). An evaluation of software for computing eigenvalues of sparse nonsymmetric matrices is provided in Lehoucq and Scott (1996). This suite of functions offers the same functionality as the ARPACK software for real nonsymmetric problems, but the interface design is quite different in order to make the option setting clearer and to simplify the interface of `nag_real_sparse_eigensystem_iter` (f12abc).

The setup function `nag_real_sparse_eigensystem_init` (f12aac) must be called before `nag_real_sparse_eigensystem_iter` (f12abc), the reverse communication iterative solver. Options may be set for `nag_real_sparse_eigensystem_iter` (f12abc) by prior calls to the option setting function `nag_real_sparse_eigensystem_option` (f12adc) and a post-processing function `nag_real_sparse_eigensystem_sol` (f12acc) must be called following a successful final exit from `nag_real_sparse_eigensystem_iter` (f12abc). `nag_real_sparse_eigensystem_monit` (f12aec), may be called following certain flagged, intermediate exits from `nag_real_sparse_eigensystem_iter` (f12abc) to provide additional monitoring information about the computation.

`nag_real_sparse_eigensystem_iter` (f12abc) uses **reverse communication**, i.e., it returns repeatedly to the calling program with the argument **irevcm** (see Section 5) set to specified values which require the calling program to carry out one of the following tasks:

- compute the matrix-vector product  $y = OPx$ , where  $OP$  is defined by the computational mode;
- compute the matrix-vector product  $y = Bx$ ;

- notify the completion of the computation;
- allow the calling program to monitor the solution.

The problem type to be solved (standard or generalized), the spectrum of eigenvalues of interest, the mode used (regular, regular inverse, shifted inverse, shifted real or shifted imaginary) and other options can all be set using the option setting function `nag_real_sparse_eigensystem_option` (f12adc) (see Section 11.1 in `nag_real_sparse_eigensystem_option` (f12adc) for details on setting options and of the default settings).

## 4 References

Lehoucq R B (2001) Implicitly restarted Arnoldi methods and subspace iteration *SIAM Journal on Matrix Analysis and Applications* **23** 551–562

Lehoucq R B and Scott J A (1996) An evaluation of software for computing eigenvalues of sparse nonsymmetric matrices *Preprint MCS-P547-1195* Argonne National Laboratory

Lehoucq R B and Sorensen D C (1996) Deflation techniques for an implicitly restarted Arnoldi iteration *SIAM Journal on Matrix Analysis and Applications* **17** 789–821

Lehoucq R B, Sorensen D C and Yang C (1998) *ARPACK Users' Guide: Solution of Large-scale Eigenvalue Problems with Implicitly Restarted Arnoldi Methods* SIAM, Philadelphia

## 5 Arguments

**Note:** this function uses **reverse communication**. Its use involves an initial entry, intermediate exits and re-entries, and a final exit, as indicated by the argument `irevcm`. Between intermediate exits and re-entries, **all arguments other than `x` and `y` must remain unchanged**.

1: `irevcm` – Integer \* *Input/Output*

*On initial entry:* `irevcm` = 0, otherwise an error condition will be raised.

*On intermediate re-entry:* must be unchanged from its previous exit value. Changing `irevcm` to any other value between calls will result in an error.

*On intermediate exit:* has the following meanings.

`irevcm` = -1

The calling program must compute the matrix-vector product  $y = OPx$ , where  $x$  is stored in `x` and the result  $y$  is placed in `y`.

`irevcm` = 1

The calling program must compute the matrix-vector product  $y = OPx$ . This is similar to the case `irevcm` = -1 except that the result of the matrix-vector product  $Bx$  (as required in some computational modes) has already been computed and is available in `mx`.

`irevcm` = 2

The calling program must compute the matrix-vector product  $y = Bx$ , where  $x$  is stored as described in the case `irevcm` = -1 and  $y$  is placed in `y`.

`irevcm` = 3

Compute the `nshift` real and imaginary parts of the shifts where the real parts are to be placed in the first `nshift` locations of the array `y` and the imaginary parts are to be placed in the first `nshift` locations of the array `mx`. Only complex conjugate pairs of shifts may be applied and the pairs must be placed in consecutive locations. This value of `irevcm` will only arise if the optional argument **Supplied Shifts** is set in a prior call to `nag_real_sparse_eigensystem_option` (f12adc) which is intended for experienced users only; the default and recommended option is to use exact shifts (see Lehoucq *et al.* (1998) for details).

**irevcn** = 4

Monitoring step: a call to `nag_real_sparse_eigensystem_monit` (f12aec) can now be made to return the number of Arnoldi iterations, the number of converged Ritz values, their real and imaginary parts, and the corresponding Ritz estimates.

*On final exit:* **irevcn** = 5: `nag_real_sparse_eigensystem_iter` (f12abc) has completed its tasks. The value of **fail** determines whether the iteration has been successfully completed, or whether errors have been detected. On successful completion `nag_real_sparse_eigensystem_sol` (f12acc) must be called to return the requested eigenvalues and eigenvectors (and/or Schur vectors).

*Constraint:* on initial entry, **irevcn** = 0; on re-entry **irevcn** must remain unchanged.

2: **resid**[*dim*] – double *Input/Output*

**Note:** the dimension, *dim*, of the array **resid** must be at least **n** (see `nag_real_sparse_eigensystem_init` (f12aac)).

*On initial entry:* need not be set unless the option **Initial Residual** has been set in a prior call to `nag_real_sparse_eigensystem_option` (f12adc) in which case **resid** should contain an initial residual vector, possibly from a previous run.

*On intermediate re-entry:* must be unchanged from its previous exit. Changing **resid** to any other value between calls may result in an error exit.

*On intermediate exit:* contains the current residual vector.

*On final exit:* contains the final residual vector.

3: **v**[**n** × **ncv**] – double *Input/Output*

The *i*th element of the *j*th basis vector is stored in location  $\mathbf{v}[\mathbf{n} \times (i - 1) + j - 1]$ , for  $i = 1, 2, \dots, \mathbf{n}$  and  $j = 1, 2, \dots, \mathbf{ncv}$ .

*On initial entry:* need not be set.

*On intermediate re-entry:* must be unchanged from its previous exit.

*On intermediate exit:* contains the current set of Arnoldi basis vectors.

*On final exit:* contains the final set of Arnoldi basis vectors.

4: **x** – double \*\* *Input/Output*

*On initial entry:* need not be set, it is used as a convenient mechanism for accessing elements of **comm**.

*On intermediate re-entry:* is not normally changed.

*On intermediate exit:* contains the vector *x* when **irevcn** returns the value -1, +1 or 2.

*On final exit:* does not contain useful data.

5: **y** – double \*\* *Input/Output*

*On initial entry:* need not be set, it is used as a convenient mechanism for accessing elements of **comm**.

*On intermediate re-entry:* must contain the result of  $y = OPx$  when **irevcn** returns the value -1 or +1. It must contain the real parts of the computed shifts when **irevcn** returns the value 3.

*On intermediate exit:* does not contain useful data.

*On final exit:* does not contain useful data.

6: **mx** – double \*\* *Input/Output*

*On initial entry:* need not be set, it is used as a convenient mechanism for accessing elements of **comm**.

*On intermediate re-entry:* must contain the result of  $y = Bx$  when **irevcn** returns the value 2. It must contain the imaginary parts of the computed shifts when **irevcn** returns the value 3.

*On intermediate exit:* contains the vector  $Bx$  when **irevcn** returns the value +1.

*On final exit:* does not contain any useful data.

- 7: **nshift** – Integer \* *Output*  
*On intermediate exit:* if the option **Supplied Shifts** is set and **irevcn** returns a value of 3, **nshift** returns the number of complex shifts required.
- 8: **comm**[*dim*] – double *Communication Array*  
**Note:** the dimension, *dim*, of the array **comm** must be at least  $\max(1, \mathbf{lcomm})$  (see `nag_real_sparse_eigensystem_init` (f12aac)).  
*On initial entry:* must remain unchanged following a call to the setup function `nag_real_sparse_eigensystem_init` (f12aac).  
*On exit:* contains data defining the current state of the iterative process.
- 9: **icomm**[*dim*] – Integer *Communication Array*  
**Note:** the dimension, *dim*, of the array **icomm** must be at least  $\max(1, \mathbf{licomm})$  (see `nag_real_sparse_eigensystem_init` (f12aac)).  
*On initial entry:* must remain unchanged following a call to the setup function `nag_real_sparse_eigensystem_init` (f12aac).  
*On exit:* contains data defining the current state of the iterative process.
- 10: **fail** – NagError \* *Input/Output*  
The NAG error argument (see Section 3.6 in the Essential Introduction).

## 6 Error Indicators and Warnings

### NE\_ALLOC\_FAIL

Dynamic memory allocation failed.

### NE\_BAD\_PARAM

On entry, argument  $\langle value \rangle$  had an illegal value.

### NE\_INITIALIZATION

Either the initialization function has not been called prior to the first call of this function or a communication array has become corrupted.

### NE\_INT

The maximum number of iterations  $\leq 0$ , the option **Iteration Limit** has been set to  $\langle value \rangle$ .

### NE\_INTERNAL\_EIGVAL\_FAIL

Error in internal call to compute eigenvalues and corresponding error bounds of the current upper Hessenberg matrix. Please contact NAG.

### NE\_INTERNAL\_ERROR

An internal error has occurred in this function. Check the function call and any array sizes. If the call is correct then please contact NAG for assistance.

**NE\_MAX\_ITER**

The maximum number of iterations has been reached. The maximum number of iterations =  $\langle value \rangle$ . The number of converged eigenvalues =  $\langle value \rangle$ . The post-processing function `nag_real_sparse_eigensystem_sol` (f12acc) may be called to recover the converged eigenvalues at this point. Alternatively, the maximum number of iterations may be increased by a call to the option setting function `nag_real_sparse_eigensystem_option` (f12adc) and the reverse communication loop restarted. A large number of iterations may indicate a poor choice for the values of **nev** and **ncv**; it is advisable to experiment with these values to reduce the number of iterations (see `nag_real_sparse_eigensystem_init` (f12aac)).

**NE\_NO\_ARNOLDI\_FAC**

Could not build an Arnoldi factorization. The size of the current Arnoldi factorization =  $\langle value \rangle$ .

**NE\_NO\_SHIFTS\_APPLIED**

No shifts could be applied during a cycle of the implicitly restarted Arnoldi iteration.

**NE\_OPT\_INCOMPAT**

The options **Generalized** and **Regular** are incompatible.

**NE\_ZERO\_INIT\_RESID**

The option **Initial Residual** was selected but the starting vector held in **resid** is zero.

**7 Accuracy**

The relative accuracy of a Ritz value,  $\lambda$ , is considered acceptable if its Ritz estimate  $\leq \mathbf{Tolerance} \times |\lambda|$ . The default **Tolerance** used is the *machine precision* given by `nag_machine_precision` (X02AJC).

**8 Parallelism and Performance**

`nag_real_sparse_eigensystem_iter` (f12abc) is threaded by NAG for parallel execution in multithreaded implementations of the NAG Library.

`nag_real_sparse_eigensystem_iter` (f12abc) makes calls to BLAS and/or LAPACK routines, which may be threaded within the vendor library used by this implementation. Consult the documentation for the vendor library for further information.

Please consult the Users' Note for your implementation for any additional implementation-specific information.

**9 Further Comments**

None.

**10 Example**

This example solves  $Ax = \lambda x$  in shift-invert mode, where  $A$  is obtained from the standard central difference discretization of the convection-diffusion operator  $\frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \rho \frac{\partial u}{\partial x}$  on the unit square, with zero Dirichlet boundary conditions. The shift used is a real number.

## 10.1 Program Text

```

/* nag_real_sparse_eigensystem_iter (f12abc) Example Program.
 *
 * Copyright 2005 Numerical Algorithms Group.
 *
 * Mark 8, 2005.
 */

#include <math.h>
#include <nag.h>
#include <nag_stdlib.h>
#include <stdio.h>
#include <nagf12.h>
#include <nagf16.h>

static void my_dgtrrf(Integer, double *, double *, double *,
                    double *, Integer *, Integer *);
static void my_dgtrrs(Integer, double *, double *, double *,
                    double *, Integer *, double *, double *);

int main(void)
{
    /* Constants */
    Integer licomm = 140, imon = 0;
    /* Scalars */
    double h, rho, s, s1, s2, s3, estnrm, sigmai, sigmar;
    Integer exit_status, info, irevcm, j, lcomm, n, nconv, ncv;
    Integer nev, niter, nshift, nx;
    /* Nag types */
    NagError fail;
    /* Arrays */
    double *comm = 0, *dd = 0, *dl = 0, *du = 0, *du2 = 0, *eigvr = 0;
    double *eigvi = 0, *eigest = 0, *resid = 0, *v = 0;
    Integer *icomm = 0, *ipiv = 0;
    /* Pointers */
    double *mx = 0, *x = 0, *y = 0;

    exit_status = 0;
    INIT_FAIL(fail);

    printf("nag_real_sparse_eigensystem_iter (f12abc) Example Program "
           "Results\n");
    /* Skip heading in data file */
    scanf("%*[\n] ");

    /* Read problem parameter values from data file. */
    scanf("%ld%ld%ld%lf%lf%lf%*[\n] ", &nx, &nev, &ncv,
          &rho, &sigmar, &sigmai);
    n = nx * nx;
    lcomm = 3*n + 3*ncv*ncv + 6*ncv + 60;
    /* Allocate memory */
    if (!(comm = NAG_ALLOC(lcomm, double)) ||
        !(eigvr = NAG_ALLOC(ncv, double)) ||
        !(eigvi = NAG_ALLOC(ncv, double)) ||
        !(eigest = NAG_ALLOC(ncv, double)) ||
        !(dd = NAG_ALLOC(n, double)) ||
        !(dl = NAG_ALLOC(n, double)) ||
        !(du = NAG_ALLOC(n, double)) ||
        !(du2 = NAG_ALLOC(n, double)) ||
        !(resid = NAG_ALLOC(n, double)) ||
        !(v = NAG_ALLOC(n * ncv, double)) ||
        !(icomm = NAG_ALLOC(licomm, Integer)) ||
        !(ipiv = NAG_ALLOC(n, Integer)))
    {
        printf("Allocation failure\n");
        exit_status = -1;
        goto END;
    }
    /* Initialise communication arrays for problem using
    nag_real_sparse_eigensystem_init (f12aac). */

```

```

nag_real_sparse_eigensystem_init(n, nev, ncv, icomm, licomm,
                                comm, lcomm, &fail);
if (fail.code != NE_NOERROR)
{
    printf(
        "Error from nag_real_sparse_eigensystem_init (f12aac).\n%s\n",
        fail.message);
    exit_status = 1;
    goto END;
}
/* Select the required mode using
nag_real_sparse_eigensystem_option (f12adc). */
nag_real_sparse_eigensystem_option("SHIFTED INVERSE REAL", icomm,
                                comm, &fail);
/* Construct C = A - sigma*I, and factorize using my_dgttrf. */
h = 1.0 / (double)(n + 1);
s = rho * h / 2.0;
s1 = -1.0 - s;
s2 = 2.0 - sigmar;
s3 = s - 1.0;
for (j = 0; j <= n - 2; ++j)
{
    dl[j] = s1;
    dd[j] = s2;
    du[j] = s3;
}
dd[n - 1] = s2;

my_dgttrf(n, dl, dd, du, du2, ipiv, &info);

irevcm = 0;
REVCOMLOOP:
/* Repeated calls to reverse communication routine
nag_real_sparse_eigensystem_iter (f12abc). */
nag_real_sparse_eigensystem_iter(&irevcm, resid, v, &x, &y, &mx,
                                &nshift, comm, icomm, &fail);
if (irevcm != 5)
{
    if (irevcm == -1 || irevcm == 1)
    {
        /* Perform y <--- OP*x = inv[A-SIGMA*I]*x. */
        /* Use my_dgttrs, a cut down C version of Lapack's dgttrs. */
        my_dgttrs(n, dl, dd, du, du2, ipiv, x, y);
    }
    else if (irevcm == 4 && imon == 1)
    {
        /* If imon=1, get monitoring information using
nag_real_sparse_eigensystem_monit (f12aec). */
        nag_real_sparse_eigensystem_monit(&niter, &nconv, eigvr,
                                        eigvi, eigest, icomm, comm);
        /* Compute 2-norm of Ritz estimates using
nag_dge_norm (f16rac).*/
        nag_dge_norm(Nag_ColMajor, Nag_FrobeniusNorm, nev, 1, eigest,
                    nev, &estnrm, &fail);
        printf("Iteration %3ld, ", niter);
        printf(" No. converged = %3ld,", nconv);
        printf(" norm of estimates = %17.8e\n", estnrm);
    }
    goto REVCOMLOOP;
}
if (fail.code == NE_NOERROR)
{
    /* Post-Process using nag_real_sparse_eigensystem_sol
(f12acc) to compute eigenvalues/vectors. */
    nag_real_sparse_eigensystem_sol(&nconv, eigvr, eigvi, v, sigmar,
                                    sigmai, resid, v, comm, icomm,
                                    &fail);

    /* Print computed eigenvalues. */
    printf("\n");
    printf(" The %4ld Ritz values of closest", nconv);
    printf(" to unity are:\n\n");
}

```

```

    for (j = 0; j <= nconv-1; ++j)
    {
        printf("%8ld%5s( %12.4f ,%12.4f )\n", j+1, "",
            eigvr[j], eigvi[j]);
    }
else
    {
        printf(
            " Error from nag_real_sparse_eigensystem_iter (f12abc).\n%s\n",
            fail.message);
        exit_status = 1;
        goto END;
    }
END:
    NAG_FREE(comm);
    NAG_FREE(eigvr);
    NAG_FREE(eigvi);
    NAG_FREE(eigest);
    NAG_FREE(dd);
    NAG_FREE(dl);
    NAG_FREE(du);
    NAG_FREE(du2);
    NAG_FREE(resid);
    NAG_FREE(v);
    NAG_FREE(icom);
    NAG_FREE(ipiv);
    return exit_status;
}

static void my_dgtrf(Integer n, double dl[], double d[],
                    double du[], double du2[], Integer ipiv[],
                    Integer *info)
{
    /* A simple C version of the Lapack routine dgtrf with argument
       checking removed */
    /* Scalars */
    double temp, fact;
    Integer i;
    /* Function Body */
    *info = 0;
    for (i = 0; i < n; ++i)
    {
        ipiv[i] = i;
    }
    for (i = 0; i < n - 2; ++i)
    {
        du2[i] = 0.0;
    }
    for (i = 0; i < n - 2; i++)
    {
        if (fabs(d[i]) >= fabs(dl[i]))
        {
            /* No row interchange required, eliminate dl[i]. */
            if (d[i] != 0.0)
            {
                fact = dl[i] / d[i];
                dl[i] = fact;
                d[i+1] = d[i+1] - fact * du[i];
            }
        }
        else
        {
            /* Interchange rows I and I+1, eliminate dl[I] */
            fact = d[i] / dl[i];
            d[i] = dl[i];
            dl[i] = fact;
            temp = du[i];
            du[i] = d[i+1];
            d[i+1] = temp - fact*d[i+1];
            du2[i] = du[i+1];
        }
    }
}

```

```

        du[i+1] = -fact * du[i+1];
        ipiv[i] = i + 1;
    }
}
if (n > 1)
{
    i = n - 2;
    if (fabs(d[i]) >= fabs(dl[i]))
    {
        if (d[i] != 0.0)
        {
            fact = dl[i] / d[i];
            dl[i] = fact;
            d[i+1] = d[i+1] - fact * du[i];
        }
    }
    else
    {
        fact = d[i] / dl[i];
        d[i] = dl[i];
        dl[i] = fact;
        temp = du[i];
        du[i] = d[i+1];
        d[i+1] = temp - fact * d[i+1];
        ipiv[i] = i + 1;
    }
}
/* Check for a zero on the diagonal of U. */
for (i = 0; i < n; ++i)
{
    if (d[i] == 0.0)
    {
        *info = i;
        goto END;
    }
}
END:
return;
}

static void my_dgttrs(Integer n, double dl[], double d[],
                    double du[], double du2[], Integer ipiv[],
                    double b[], double y[])
{
    /* A simple C version of the Lapack routine dgttrs with argument
       checking removed, the number of right-hand-sides=1, Trans='N' */
    /* Scalars */
    Integer i, ip;
    double temp;
    /* Solve L*x = b. */
    for (i = 0; i <= n - 1; ++i)
    {
        y[i] = b[i];
    }
    for (i = 0; i < n - 1; ++i)
    {
        ip = ipiv[i];
        temp = y[i+1-ip+i] - dl[i]*y[ip];
        y[i] = y[ip];
        y[i+1] = temp;
    }
    /* Solve U*x = b. */
    y[n-1] = y[n-1] / d[n-1];
    if (n > 1)
    {
        y[n-2] = (y[n-2] - du[n-2]*y[n-1])/d[n-2];
    }
    for (i = n - 3; i >= 0; --i)
    {
        y[i] = (y[i]-du[i]*y[i+1]-du2[i]*y[i+2])/d[i];
    }
}

```

```
    return;  
}
```

## 10.2 Program Data

```
nag_real_sparse_eigensystem_iter (f12abc) Example Program Data  
10 4 20 10.0 1.0 0.0 : Values for nx, nev, ncv, rho, sigmar, sigmai
```

## 10.3 Program Results

```
nag_real_sparse_eigensystem_iter (f12abc) Example Program Results
```

The 4 Ritz values of closest to unity are:

1	(	1.0192	,	0.0000	)
2	(	0.9656	,	0.0000	)
3	(	1.0738	,	0.0000	)
4	(	0.9129	,	0.0000	)

---