NAG Library Routine Document F08GUF (ZUPMTR)

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

F08GUF (ZUPMTR) multiplies an arbitrary complex matrix C by the complex unitary matrix Q which was determined by F08GSF (ZHPTRD) when reducing a complex Hermitian matrix to tridiagonal form.

2 Specification

```
SUBROUTINE FO8GUF (SIDE, UPLO, TRANS, M, N, AP, TAU, C, LDC, WORK, INFO)

INTEGER
M, N, LDC, INFO

COMPLEX (KIND=nag_wp) AP(*), TAU(*), C(LDC,*), WORK(*)

CHARACTER(1) SIDE, UPLO, TRANS
```

The routine may be called by its LAPACK name *zupmtr*.

3 Description

F08GUF (ZUPMTR) is intended to be used after a call to F08GSF (ZHPTRD), which reduces a complex Hermitian matrix A to real symmetric tridiagonal form T by a unitary similarity transformation: $A = QTQ^{\rm H}$. F08GSF (ZHPTRD) represents the unitary matrix Q as a product of elementary reflectors.

This routine may be used to form one of the matrix products

$$QC, Q^{H}C, CQ$$
 or CQ^{H} ,

overwriting the result on C (which may be any complex rectangular matrix).

A common application of this routine is to transform a matrix Z of eigenvectors of T to the matrix QZ of eigenvectors of A.

4 References

Golub G H and Van Loan C F (1996) Matrix Computations (3rd Edition) Johns Hopkins University Press, Baltimore

5 Parameters

1: SIDE - CHARACTER(1)

Input

On entry: indicates how Q or Q^H is to be applied to C.

SIDE = 'L'

Q or Q^{H} is applied to C from the left.

SIDE = 'R'

Q or Q^{H} is applied to C from the right.

Constraint: SIDE = 'L' or 'R'.

2: UPLO - CHARACTER(1)

Input

On entry: this **must** be the same parameter UPLO as supplied to F08GSF (ZHPTRD).

Constraint: UPLO = 'U' or 'L'.

Mark 25 F08GUF.1

F08GUF NAG Library Manual

3: TRANS - CHARACTER(1)

Input

On entry: indicates whether Q or Q^{H} is to be applied to C.

TRANS = 'N'

Q is applied to C.

TRANS = 'C'

 $Q^{\rm H}$ is applied to C.

Constraint: TRANS = 'N' or 'C'.

4: M – INTEGER

Input

On entry: m, the number of rows of the matrix C; m is also the order of Q if SIDE = 'L'.

Constraint: $M \ge 0$.

5: N – INTEGER

Input

On entry: n, the number of columns of the matrix C; n is also the order of Q if SIDE = 'R'. Constraint: N > 0.

6: AP(*) - COMPLEX (KIND=nag wp) array

Input/Output

Note: the dimension of the array AP must be at least $max(1, M \times (M+1)/2)$ if SIDE = 'L' and at least $max(1, N \times (N+1)/2)$ if SIDE = 'R'.

On entry: details of the vectors which define the elementary reflectors, as returned by F08GSF (ZHPTRD).

On exit: is used as internal workspace prior to being restored and hence is unchanged.

7: TAU(*) – COMPLEX (KIND=nag_wp) array

Input

Note: the dimension of the array TAU must be at least max(1, M - 1) if SIDE = 'L' and at least max(1, N - 1) if SIDE = 'R'.

On entry: further details of the elementary reflectors, as returned by F08GSF (ZHPTRD).

8: $C(LDC,*) - COMPLEX (KIND=nag_wp) array$

Input/Output

Note: the second dimension of the array C must be at least max(1, N).

On entry: the m by n matrix C.

On exit: C is overwritten by QC or $Q^{H}C$ or CQ or CQ^{H} as specified by SIDE and TRANS.

9: LDC – INTEGER

Input

On entry: the first dimension of the array C as declared in the (sub)program from which F08GUF (ZUPMTR) is called.

Constraint: LDC > max(1, M).

10: WORK(*) - COMPLEX (KIND=nag wp) array

Workspace

Note: the dimension of the array WORK must be at least max(1, N) if SIDE = 'L' and at least max(1, M) if SIDE = 'R'.

11: INFO – INTEGER

Output

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

F08GUF.2 Mark 25

6 Error Indicators and Warnings

INFO < 0

If INFO = -i, argument i had an illegal value. An explanatory message is output, and execution of the program is terminated.

7 Accuracy

The computed result differs from the exact result by a matrix E such that

$$||E||_2 = O(\epsilon)||C||_2$$

where ϵ is the *machine precision*.

8 Parallelism and Performance

F08GUF (ZUPMTR) is not threaded by NAG in any implementation.

F08GUF (ZUPMTR) 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 X06 Chapter Introduction for information on how to control and interrogate the OpenMP environment used within this routine. Please also consult the Users' Note for your implementation for any additional implementation-specific information.

9 Further Comments

The total number of real floating-point operations is approximately $8m^2n$ if SIDE = 'L' and $8mn^2$ if SIDE = 'R'.

The real analogue of this routine is F08GGF (DOPMTR).

10 Example

This example computes the two smallest eigenvalues, and the associated eigenvectors, of the matrix A, where

$$A = \begin{pmatrix} -2.28 + 0.00i & 1.78 - 2.03i & 2.26 + 0.10i & -0.12 + 2.53i \\ 1.78 + 2.03i & -1.12 + 0.00i & 0.01 + 0.43i & -1.07 + 0.86i \\ 2.26 - 0.10i & 0.01 - 0.43i & -0.37 + 0.00i & 2.31 - 0.92i \\ -0.12 - 2.53i & -1.07 - 0.86i & 2.31 + 0.92i & -0.73 + 0.00i \end{pmatrix}$$

using packed storage. Here A is Hermitian and must first be reduced to tridiagonal form T by F08GSF (ZHPTRD). The program then calls F08JJF (DSTEBZ) to compute the requested eigenvalues and F08JXF (ZSTEIN) to compute the associated eigenvectors of T. Finally F08GUF (ZUPMTR) is called to transform the eigenvectors to those of A.

10.1 Program Text

```
Program f08gufe
! F08GUF Example Program Text
! Mark 25 Release. NAG Copyright 2014.
! .. Use Statements ..
    Use nag_library, Only: dstebz, nag_wp, x04dbf, zhptrd, zstein, zupmtr
! .. Implicit None Statement ..
    Implicit None
! .. Parameters ..
    Real (Kind=nag_wp), Parameter :: zero = 0.0E0_nag_wp
```

Mark 25 F08GUF.3

F08GUF NAG Library Manual

```
Integer, Parameter
                                        :: nin = 5, nout = 6
      .. Local Scalars ..
                                        :: v1, vu
      Real (Kind=nag_wp)
      Integer
                                         :: i, ifail, info, j, ldc, m, n, nsplit
      Character (1)
                                         :: uplo
      .. Local Arrays ..
     Complex (Kind=nag_wp), Allocatable :: ap(:), c(:,:), tau(:), work(:)
Real (Kind=nag_wp), Allocatable :: d(:), e(:), rwork(:), w(:)
                                         :: iblock(:), ifailv(:), isplit(:),
      Integer, Allocatable
                                            iwork(:)
     Character (1)
                                         :: clabs(1), rlabs(1)
      .. Executable Statements ..
!
      Write (nout,*) 'F08GUF Example Program Results'
!
      Skip heading in data file
      Read (nin,*)
      Read (nin,*) n
      ldc = n
     Allocate (ap(n*(n+1)/2),c(ldc,n),tau(n),work(n),d(n),e(n),rwork(5*n),w(n &
        ), iblock(n), ifailv(n), isplit(n), iwork(3*n))
     Read A from data file
     Read (nin,*) uplo
      If (uplo=='U') Then
        Read (nin,*)((ap(i+j*(j-1)/2),j=i,n),i=1,n)
      Else If (uplo=='L') Then
        Read (nin,*)((ap(i+(2*n-j)*(j-1)/2),j=1,i),i=1,n)
     End If
     Reduce A to tridiagonal form T = (Q**H)*A*Q
!
      The NAG name equivalent of zhptrd is f08gsf
      Call zhptrd(uplo,n,ap,d,e,tau,info)
     Calculate the two smallest eigenvalues of T (same as A)
1
      The NAG name equivalent of dstebz is f08jjf
1
      Call dstebz('I','B',n,vl,vu,1,2,zero,d,e,m,nsplit,w,iblock,isplit,rwork, &
        iwork,info)
     Write (nout,*)
      If (info>0) Then
        Write (nout,*) 'Failure to converge.'
        Write (nout,*) 'Eigenvalues'
        Write (nout,99999) w(1:m)
        Calculate the eigenvectors of {\tt T}, storing the result in {\tt C}
!
        The NAG name equivalent of zstein is f08jxf
!
        Call zstein(n,d,e,m,w,iblock,isplit,c,ldc,rwork,iwork,ifailv,info)
        If (info>0) Then
          Write (nout,*) 'Failure to converge.'
        Else
          Calculate the eigenvectors of A = Q * (eigenvectors of T)
          The NAG name equivalent of zupmtr is f08guf
!
          Call zupmtr('Left',uplo,'No transpose',n,m,ap,tau,c,ldc,work,info)
!
          Print eigenvectors
          Write (nout,*)
          Flush (nout)
          Normalize the eigenvectors
          Do i = 1, m
           c(1:n,i) = c(1:n,i)/c(1,i)
          End Do
!
          ifail: behaviour on error exit
                 =0 for hard exit, =1 for quiet-soft, =-1 for noisy-soft
          ifail = 0
```

F08GUF.4 Mark 25

10.2 Program Data

10.3 Program Results

Mark 25 F08GUF.5 (last)