

```

C COMPOSED ENTIRELY OF ONES
  N=100
  X=1
  DO 30 K=1,3
    DO 10 I=1,N
      G(1,I)=2.0
      G(2,I)=-1.0
      B(I)=0.0
10    CONTINUE
      G(1,1)=1.0+X
      G(1,N)=1.0+X
      B(1)=X
      B(N)=X
C SOLVE THE SYSTEM
  MU=2
  CALL BPSS(N,MU,G,2,B,N,1,COND)
  IWRITE=IWMACH(2)
  WRITE(IWRITE,11)X
11  FORMAT(/5H X IS,F15.7)
  WRITE(IWRITE,12)COND
12  FORMAT(20H CONDITION NUMBER IS,1PE15.7)
C COMPUTE THE ERROR
  ERR=0.0
  DO 20 I=1,N
    ERR=AMAX1(ERR,ABS(B(I)-1.0))
20  CONTINUE
  WRITE(IWRITE,21)ERR
21  FORMAT(22H FOR BPSS THE ERROR IS,F16.8)
  X=X/100.
30  CONTINUE
  STOP
  END

```

When the above program was executed on the Honeywell 6000 machine at Bell Laboratories, the following was printed

```

X IS      1.0000000
CONDITION NUMBER IS  4.0807862E 03
FOR BPSS THE ERROR IS      0.00000431

X IS      0.0100000
CONDITION NUMBER IS  2.3329148E 04
FOR BPSS THE ERROR IS      0.00002055

X IS      0.0001000
CONDITION NUMBER IS  1.9933923E 06
FOR BPSS THE ERROR IS      0.00491761

```



use the routine BPLE, which is a little faster than BPSS. Ordinarily, however, the user is strongly urged to choose BPSS, and to follow it by a test of the condition estimate.

**Note 2:** Users who wish to solve a sequence of problems with the same coefficient matrix, but different right-hand sides *not all known in advance*, should not use BPSS, but should call subprograms BPCE, BPFS and BPBS. (See the example of BPFS.) BPCE is called once to get the  $LDL^T$  decomposition (see the introduction to this chapter) and then the pair, BPFS (forward solve) and BPBS (back solve), is called for each new right-hand side.

**Note 3:** The  $LDL^T$  decomposition of A satisfies the equation  $A = LDL^T$  where L is lower unit triangular (1's on the diagonal, 0's above the diagonal) and D is diagonal. On return from BPSS, the diagonal of D occupies the first row of G and  $G(i-j+1,i) = l_{ij}$  for  $i > j$ .

**Note 4:** For complex Hermitian matrices ( $A = A^*$ , where  $A^*$  represents the conjugate transpose of A), the complex version of this subroutine computes the  $LDL^*$  decomposition and returns the conjugate of L rather than L in G.

**Error situations:** \*(The user can elect to 'recover' from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$
4	$IB < N$
5	$NB < 1$
$10 + k^*$	singular matrix whose rank is at least k
$10 + N + k^*$	$k^{th}$ principal minor is not positive definite

**Double-precision version:** DBPSS, with G, B, and COND declared double precision.

**Complex Hermitian version:** CBPSS with G and B declared complex (see Note 4 above)

## BPSS — band positive definite linear system solution with condition estimation

**Purpose:** BPSS (Band Positive definite System Solution) solves the system  $AX = B$  where  $A$  is a band symmetric positive definite matrix. An estimate of the condition number of  $A$  is provided.

**Usage:** CALL BPSS (N, MU, G, IG, B, IB, NB, COND)

N → the number of equations

MU → the number of nonzero bands on and above the diagonal of  $A$

G → a matrix into which the upper triangular portion of the matrix  $A$  has been placed as follows:

$$G(j-i+1, i) = a_{ij} \text{ for } j \geq i$$

(See the introduction to this chapter.)

$G$  should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

← L and D from the factorization of  $A$  into  $LDL^T$   
(see **Note 3**)

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

B → the matrix of right-hand sides, dimensioned (IB, KB) in the calling program, where  $IB \geq N$  and  $KB \geq NB$ .

← the solution  $X$

IB → the row (leading) dimension of  $B$ , as dimensioned in the calling program

NB → the number of right-hand sides

COND ← an estimate of the condition number of  $A$  (See **Note 1**)

**Note 1:** The condition number measures the sensitivity of the solution of a linear system to errors in the matrix and in the right-hand side. If the elements of the matrix and the right-hand side(s) of your linear system have  $d$  decimal digits of precision, the solution might have as few as  $d - \log_{10}(\text{COND})$  correct decimal digits. Thus if  $\text{COND}$  is greater than  $10^{\text{Bd}P}$ , there may be no correct digits.

If the given matrix,  $A$ , is known in advance to be well-conditioned, then the user may wish to

**Double-precision version:** DBPNM with G and DBPNM declared double precision

**Complex version:** CBPNM with G declared complex

**Storage:** None

**Time:**  $N \times (2 \times MU - 1)$  additions

**See also:** BPDC, BPLD, BPLE, BPSS, BPCE

**Author:** Linda Kaufman

**Example:** Because of roundoff error it is often very difficult to decide whether a matrix is singular. One criteria often used for symmetric positive definite matrices is to compute the  $LDL^T$  decomposition of the matrix and declare the matrix singular if any element of the diagonal matrix D is less than  $\epsilon \|A\|$  where  $\epsilon$  is the machine precision and is computed by R1MACH(4).

The following program fragment might be used to indicate whether the symmetric positive definite banded matrix packed into G is singular or nearly singular. It uses the fact that the subroutine BPLD, which computes the  $LDL^T$  decomposition of a banded symmetric matrix, issues a recoverable error when it detects an element of D less than EPS, an input parameter to the subroutine.

```

                                IWRITE=I1MACH(2)
                                CALL ENTSRC(IROLD,1)
                                EPS=BPNM(N,MU,G,IG)*R1MACH(4)
                                CALL BPLD(N,MU,G,IG,EPS)
                                IF (NERROR(IERR).EQ.0) GO TO 10
                                    CALL ERROFF
                                    WRITE(IWRITE,1)
1                                FORMAT(16H SINGULAR MATRIX)
10                               CONTINUE

```

## BPNM — norm of a banded symmetric positive definite matrix

**Purpose:** BPNM (Banded Positive definite matrix NorM) computes the norm of a banded symmetric positive definite matrix A stored in packed form. The infinity norm is defined as

$$\max_{1 \leq i \leq n} \sum_{j=1}^n |a_{ij}|$$

**Type:** Real function

**Usage:** <answer> = BPNM (N, MU, G, IG)

N → the number of rows in A

MU → the number of nonzero bands in A on and above the diagonal

G → a matrix into which the upper triangular portion of the matrix A has been packed as follows:

$$G(1+j-i, i) = a_{ij} \text{ for } j \geq i$$

i.e. the main diagonal of A is in the first row of G

(See the introduction to this chapter.)

G should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

IG → the row (leading) dimension of G, as dimensioned in the calling program

$$\text{<answer>} \leftarrow \max_{1 \leq i \leq n} \sum_{j=1}^n |a_{ij}|$$

**Error situations:** (All errors in this subprogram are fatal — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$

```

C
      CALL BPML(N,MU,G,IG,X,B)
C
C SOLVE THE SYSTEM AX=B
C
      CALL BPSS(N,MU,G,IG,B,N,1,COND)
C
C PRINT OUT THE TRUE SOLUTION AND THE COMPUTED SOLUTION
C
      IWRITE=I1MACH(2)
      WRITE(IWRITE,21)
21     FORMAT(34H TRUE SOLUTION   COMPUTED SOLUTION)
      WRITE(IWRITE,22)(X(I),B(I),I=1,N)
22     FORMAT(1H ,2E16.8)
      ERR=0.0
      DO 30 I=1,N
          ERR=ERR+ABS(B(I)-X(I))
30     CONTINUE
      ERR=ERR/SASUM(N,X,1)
      WRITE(IWRITE,31)ERR
31     FORMAT(19H RELATIVE ERROR IS ,1PE15.7)
      WRITE(IWRITE,32)COND
32     FORMAT(20H CONDITION NUMBER IS,1PE15.7)
      STOP
      END

```

When the above program was executed on the Honeywell 6000 machine at Bell Laboratories, which has a machine precision of  $1 \times 10^{-8}$ , the following was printed:

```

TRUE SOLUTION   COMPUTED SOLUTION
0.22925607E 00  0.22925608E 00
0.76687502E 00  0.76687504E 00
0.68317685E 00  0.68317687E 00
0.50919111E 00  0.50919112E 00
0.87455959E 00  0.87455962E 00
0.64464101E 00  0.64464103E 00
0.84746840E 00  0.84746842E 00
0.35396343E 00  0.35396345E 00
0.39889160E 00  0.39889160E 00
0.45709422E 00  0.45709422E 00
RELATIVE ERROR IS  3.1985797E-08
CONDITION NUMBER IS  2.1901962E 01

```

The condition number of the matrix and the precision of the Honeywell suggest that even in the absence of roundoff error in BPML, a relative error of  $2.2 \times 10^{-7}$  would not be surprising. The value computed above is quite reasonable.

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BPML

**Storage:** None**Time:** (2MU-1)×N additions  
(2MU-1)×N multiplications**See also:** BPBS, BPCE, BPDC, BPLU, BPLE, BPSS**Author:** Linda Kaufman

**Example:** This example checks the consistency of BPML and BPSS the banded system solver. First the example uses BPML to compute for a given vector  $x$  and a given matrix  $A$  the vector  $b = Ax$ . Then the problem is inverted, i.e., BPSS is used to find the vector  $x$  which satisfies  $Ax = b$ . This  $x$  is then compared with the original vector. The vector  $x$  is generated randomly and the 10×10 matrix  $A$  is given by

```

      4  -1  -1
     -1  4  -1  -1
     -1  -1  4  -1  -1
           -1  -1  4  -1  -1
                   .  .  .
                   -1 -1  4  -1  -1
                           -1 -1  4  -1
                                   -1 -1  4

```

```

      INTEGER IG, N, MU, I, IWRITE, IIMACH
      REAL G(3,20), X(20), B(20)
      REAL UNI, ERR, COND, SASUM, ABS
      IG=3
      N=10
      MU=3

C
C CONSTRUCT MATRIX A AND PACK IT INTO G
C
      DO 10 I=1,N
          G(1,I)=4.0
          G(2,I)=-1.0
          G(3,I)=-1.0
      10  CONTINUE
C
C CONSTRUCT A RANDOM VECTOR
C
      DO 20 I=1,N
          X(I)=UNI(0)
      20  CONTINUE
C
C CONSTRUCT B=AX

```



## BPML — banded positive definite matrix - vector multiplication

**Purpose:** BPML (Banded Positive definite matrix MuLtiplication) forms the product  $Ax$  where  $A$  is a symmetric banded positive matrix stored in packed form.

**Usage:** CALL BPML (N, MU, G, IG, X, B)

N → the length of  $x$

MU → the number of nonzero bands on and above the diagonal of  $A$

G → a matrix into which the upper triangular portion of the matrix  $A$  has been packed as follows:

$$G(1 + j - i, i) = a_{ij} \text{ for } j \geq i.$$

(See the introduction to this chapter.)  $G$  should be dimensioned (IG, KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

X → the vector  $x$  to be multiplied

B ← the vector  $Ax$

**Error situations:** (All errors in this subprogram are fatal — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$

**Double-precision version:** DBPML with  $G$ ,  $X$ , and  $B$  declared double precision.

**Complex Hermitian version:** CBPML with  $G$ ,  $X$ , and  $B$  declared complex

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BPLE

```

          G(MU,I)=-1.0
20      CONTINUE
          G(2,I)=0.0
30      CONTINUE
C
C SET UP RIGHT HAND SIDE SO SOLUTION IS ALL 1'S
C
          DO 40 I=1,N
            X(I)=1.0
40      CONTINUE
          CALL BPML(N,MU,G,IG,X,B)
C
C SOLVE THE SYSTEM
C
          CALL BPLE(N,MU,G,IG,B,100,1)
C
C COMPUTE THE ERROR
C
          ERR=0.0
          DO 50 I=1,N
            ERR=AMAX1(ERR,ABS(B(I)-1.0))
50      CONTINUE
          IWRITE=11MACH(2)
          WRITE(IWRITE,51)ERR
51      FORMAT(31H ERROR IN SOLUTION FROM BPLE IS,F15.8)
          STOP
          END

```

When the above code was run on the Honeywell 6000 machine at Bell Laboratories, the following was printed:

```
ERROR IN SOLUTION FROM BPLE IS  0.00000012
```

**See also:** BPBS, BPCE, BPDC, BPFS, BPLD, BPSS

**Author:** Linda Kaufman

**Example:** The matrix in this example is derived from the usual five-point approximation to the Laplace operator on the unit square with an 11×11 mesh. The 100×100 matrix A has the form

$$\begin{array}{ccccccc}
 C & -I & & & & & \\
 -I & C & -I & & & & \\
 & -I & C & -I & & & \\
 & & & \dots\dots & & & \\
 & & & & -I & C & -I \\
 & & & & & -I & C
 \end{array}$$

where I is the identity matrix of order 10 and C is the matrix

$$\begin{array}{ccccccc}
 4 & -1 & & & & & \\
 -1 & 4 & -1 & & & & \\
 & -1 & 4 & -1 & & & \\
 & & & \dots & & & \\
 & & & & -1 & 4 & -1 \\
 & & & & & -1 & 4
 \end{array}$$

To make it easy to detect errors in the example, the right-hand side has been chosen to make the solution a vector of all 1's. To construct the right-hand side the subroutine BPML is used which produces b=Ax where A is a band symmetric positive definite matrix packed into the matrix G.

```

      INTEGER IG, N, MU, MLM1, I, KBLOK, KK, J
      INTEGER IWRITE, ILMACH
      REAL G(11,100), B(100), X(100)
      REAL ERR, AMAX1
      IG=11
      N=100
      MU=11
C
C SET UP MATRIX FOR ELLIPTIC PDE IN 2 DIMENSIONS
C
      MLM1=MU-1
      I=0
      DO 30 KBLOK=1,MLM1
        DO 20 KK=1,MLM1
          I=I+1
          G(1,I)=4.0
          G(2,I)=-1.0
          DO 10 J=3,MLM1
            G(J,I)=0.0
          10 CONTINUE
        20 CONTINUE
      30 CONTINUE
  
```

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BPLE

**Note 3:** The  $LDL^T$  decomposition of  $A$  satisfies the equation  $A = LDL^T$  where  $L$  is lower unit triangular (1's on the diagonal, 0's above the diagonal) and  $D$  is diagonal. On return from BPLE, the diagonal of  $D$  occupies the first row of  $G$  and  $G(i-j+1, i) = l_{ij}$  for  $i > j$ .

**Note 4:** For complex Hermitian matrices ( $A = A^*$ , where  $A^*$  represents the conjugate transpose of  $A$ ), the complex version of this subroutine computes the  $LDL^*$  decomposition and returns the conjugate of  $L$  rather than  $L$  in  $G$ .

**Error situations:** \*(The user can elect to 'recover' from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$
4	$IB < N$
5	$NB < 1$
$10 + k^*$	singular matrix whose rank is at least $k$
$10 + N + k^*$	$k^{th}$ principal minor is not positive definite

**Double-precision version:** DBPLE with  $G$  and  $B$  declared double precision

**Complex Hermitian version:** CBPLE with  $G$  and  $B$  declared complex (see **Note 4**).

**Storage:** None

**Time:**  $N \times ((MU-1) \times (MU/2 + 2 \times (NB+1)) + 1)$  additions  
 $N \times (MU-1) \times (MU/2 + 2 \times NB)$  multiplications  
 $N \times (MU-1 + NB)$  divisions

**Method:** BPLE calls BPDC to form the  $LDL^T$  decomposition, and then calls BPFS for the forward solution, and BPBS for the back solution.

## BPLE — band symmetric positive definite linear system solution

**Purpose:** BPLE (Banded symmetric Positive definite Linear Equation solution) solves the system  $AX = B$  where  $A$  is a banded symmetric positive definite matrix

**Usage:** CALL BPLE (N, MU, G, IG, B, IB, NB)

N → the number of equations

MU → the number of nonzero bands on and below the diagonal of  $A$

G → a matrix into which the upper triangular portion of the matrix  $A$  has been packed as follows:

$$G(j-i + 1, i) = a_{ij} \text{ for } j \geq i$$

(See the introduction to this chapter.)

$G$  should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

←  $L$  and  $D$  from the factorization of  $A$  into  $LDL^T$  (see **Note 3**)

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

B → the matrix of right-hand sides, dimensioned (IB, KB) in the calling program, where  $IB \geq N$  and  $KB \geq NB$ .

← the solution  $X$

IB → the row (leading) dimension of  $B$ , as dimensioned in the calling program

NB → the number of right-hand sides

**Note 1:** Unless the given matrix  $A$  is known in advance to be well-conditioned, the user should use BPSS instead of BPLE.

**Note 2:** Users who wish to solve a sequence of problems with the same coefficient matrix, but different right-hand sides *not all known in advance*, should call subprograms BPLE, BPFS and BPBS. (See the example in BPFS.) BPLE is called once to get the  $LDL^T$  decomposition (see the introduction to this chapter) and to solve for the first solution and then the pair, BPFS (forward solve) and BPBS (back solve), is called for each additional right-hand side.

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BPLD

**Example:** As noted above, after execution of BPLD, the determinant of the matrix stored in G is the product of the elements stored in the first row of G. The subroutine computes this product taking care to avoid underflow and overflow. The subroutine UMKFL is used to decompose a floating-point number, F, into a mantissa, M, and an exponent E such that  $F = Mb^E$  where b is the base of the machine and  $1/b \leq |M| \leq 1$

```

      SUBROUTINE BPDET(N,MU,G,IG,DETMAN,IDEDEX)
      C
      C THIS SUBROUTINE COMPUTES THE DETERMINANT OF A
      C BAND SYMMETRIC POSITIVE DEFINITE MATRIX STORED IN G.
      C IT IS GIVEN BY DETMAN*BETA**IDEDEX
      C WHERE BETA IS THE BASE OF THE MACHINE
      C AND DETMAN IS BETWEEN 1/BETA AND 1
      C
      REAL G(IG,N),DETMAN
      REAL ONOVBE,M
      INTEGER E
      INTEGER IDEDEX
      CALL BPLD(N,MU,G,IG,0.0)
      C
      C THE DETERMINANT IS THE PRODUCT OF THE ELEMENTS OF ROW 1 OF G
      C WE TRY TO COMPUTE THIS PRODUCT IN A WAY THAT WILL
      C AVOID UNDERFLOW AND OVERFLOW
      C
      ONOVBE=1.0/FLOAT(I1MACH(10))
      DETMAN=ONOVBE
      BETA=FLOAT(I1MACH(10))
      IDEDEX=1
      DO 10 I=1,N
         CALL UMKFL(G(1,I),E,M)
         DETMAN=DETMAN*M
         IDEDEX=IDEDEX+E
         IF(DETMAN.GE.ONOVBE) GO TO 10
         IDEDEX=IDEDEX-1
         DETMAN=DETMAN*BETA
10    CONTINUE
      RETURN
      END

```

**Error situations:** \*(The user can elect to ‘recover’ from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$
$10 + k^*$	singular matrix whose rank is at least $k$
$10 + N + k^*$	the $k^{th}$ principal minor is not positive definite

**Double-precision version:** DBPLD with G and EPS declared double precision

**Complex Hermitian version:** CBPLD with G declared complex (see Note 3 above)

**Storage:** None

**Timing:**  $N \times (MU - 1) \times MU / 2$  additions  
 $N \times (MU - 1) \times MU / 2$  multiplications  
 $(MU - 1) \times N$  divisions

**Method:** Gaussian elimination without pivoting

**See also:** BPBS, BPCE, BPDC, BPFS, BPLE, BPSS

**Author:** Linda Kaufman

BPLD —  $LDL^T$  decomposition of a band symmetric positive definite matrix

**Purpose:** BPLD (Band Positive definite  $LDL^T$  decomposition) decomposes a banded symmetric positive definite matrix  $A$  into  $LDL^T$  where  $L$  is lower triangular and  $D$  is diagonal. It allows the user to provide a threshold for considering a matrix singular BPLD is called by the decomposition routines BPCE and BPDC.

**Usage:** CALL BPLD (N, MU, G, IG, EPS)

N → the number of equations

MU → the number of nonzero bands on and above the diagonal of  $A$

G → a matrix into which the upper triangular portion of the matrix  $A$  has been packed as follows:

$$G(j-i+1, i) = a_{ij} \text{ for } j \geq i$$

(See the introduction to this chapter.)

$G$  should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

← the  $LDL^T$  decomposition suitable for input into BPFS and BPBS (see Note 2)

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

EPS → if there exists an index  $k$  such that  $|d_{kk}| \leq EPS$  then  $A$  is considered singular

**Note 1:** After the execution of BPLD, (if the matrix has not been found singular), the determinant of  $A$  is the product of the elements of the first row of  $G$ .

**Note 2:** The  $LDL^T$  decomposition of  $A$  satisfies the equation  $A = LDL^T$  where  $L$  is lower unit triangular (1's on the diagonal, 0's above the diagonal) and  $D$  is diagonal. On return from BPLD, the diagonal of  $D$  occupies the first row of  $G$  and  $G(i-j+1, i) = l_{ij}$  for  $i > j$ .

**Note 3:** For complex Hermitian matrices ( $A = A^*$ , where  $A^*$  represents the conjugate transpose of  $A$ ), the complex version of this subroutine computes the  $LDL^*$  decomposition and returns the conjugate of  $L$  rather than  $L$  in  $G$ .



```

          DO 50 I=1,N
            B(I)=B(I)+R(I)
            RNORM=RNORM+ABS(R(I))
50        CONTINUE
          IF(RNORM.LT.R1MACH(4)*BNORM) GO TO 70
60        CONTINUE
          WRITE(IWRITE,61)
61        FORMAT(18H REFINEMENT FAILED)
C COMPUTE NEW ERROR
70        ERR=0.0
          DO 80 I=1,N
            ERR=AMAX1(ERR,ABS(B(I)-1.0))
80        CONTINUE
          WRITE(IWRITE,81)IT,ERR
81        FORMAT(24H ERROR AFTER REFINEMENT ,I4,3H IS,E14.7)
          X=X/100.0
90        CONTINUE
          STOP
          END

```

When the above program was executed on the Honeywell 6000 at Bell Laboratories, the following was printed

```

X IS      1.00000000
FOR BPLE THE ERROR IS      0.00000431
ERROR AFTER REFINEMENT    2 IS 0.

X IS      0.01000000
FOR BPLE THE ERROR IS      0.00002055
ERROR AFTER REFINEMENT    3 IS 0.

X IS      0.00010000
FOR BPLE THE ERROR IS      0.00491761
ERROR AFTER REFINEMENT    4 IS 0.

```

This problem was chosen because as  $x$  approaches 0, the matrix becomes more ill-conditioned, the error in the solution grows, and more steps of iterative refinement are required. The reader should be aware that in this example we were able to represent the matrix and the right-hand side precisely, but due to roundoff error this is not always the case. Often the iterative refinement algorithm produces an exact, but useless, solution to a slightly incorrect problem.

The following program has been specifically tailored to the three problems.

```

      INTEGER N, ML, IG, NM1, K, I, IWRITE, ILMACH, IT, IEND, ITER
      REAL G(2,100), B(200), R(200)
      REAL X, ERR, AMAX1, RNORM, BNORM, RLMACH, ABS
      DOUBLE PRECISION DBLE
C CONSTRUCT MATRIX AND RIGHT HAND SIDE SO TRUE SOLUTION IS
C COMPOSED ENTIRELY OF 1S
      N=100
      X=1
      ML=2
      IG=2
      NM1=N-1
      DO 90 K=1,3
        DO 10 I=1,N
          G(1,I)=2.0
          G(2,I)=-1.0
          B(I)=0.0
10      CONTINUE
          G(1,1)=1.0+X
          G(1,N)=1.0+X
          B(1)=X
          B(N)=X
C SOLVE THE SYSTEM
          CALL BPLE(N,ML,G,IG,B,N,1)
          IWRITE=ILMACH(2)
          WRITE(IWRITE,11)X
11      FORMAT(/5H X IS,F16.8)
C COMPUTE THE ERROR
          ERR=0.0
          DO 20 I=1,N
            ERR=AMAX1(ERR,ABS(B(I)-1.0))
20      CONTINUE
          WRITE(IWRITE,21)ERR
21      FORMAT(22H FOR BPLE THE ERROR IS,F16.8)
          IEND=ILMACH(11)*IFIX(RLMACH(5)/ALOG10(2.0)+1.0)
C FIND THE NORM OF THE SOLUTION
          BNORM=0.0
          DO 30 I=1,N
            BNORM=AMAX1(BNORM,ABS(B(I)))
30      CONTINUE
C REFINE THE SOLUTION
          DO 60 ITER=1,IEND
            IT=ITER
C COMPUTE THE RESIDUAL R=B-AX, IN DOUBLE PRECISION
            DO 40 I=2,NM1
              R(I)=DBLE(B(I-1))+DBLE(B(I+1))-2.0*DBLE(B(I))
40          CONTINUE
              R(1)=X+B(2)-DBLE(1.0+X)*DBLE(B(1))
              R(N)=X+B(N-1)-DBLE(1.+X)*DBLE(B(N))
C SOLVE A(DELTA X)=R
              CALL BPFS(N,ML,G,IG,R,N,1)
              CALL BPBS(N,ML,G,IG,R,N,1)
C DETERMINE NORM OF CORRECTION AND ADD IN CORRECTION
              RNORM=0.0

```

**Example:** In the following example we solve three problems that might arise from a discretization of a 1-dimensional differential equation. The coefficient matrix in each problem has the form

$$\begin{pmatrix}
 1+x & -1 & & & & \\
 -1 & 2 & -1 & & & \\
 & -1 & 2 & -1 & & \\
 & & -1 & 2 & -1 & \\
 & & & & & \ddots & \\
 & & & & & & & \ddots & \\
 & & & & & & & & & -1 & 2 & -1 \\
 & & & & & & & & & -1 & 1+x
 \end{pmatrix}$$

with  $x=1., .01,$  and  $.0001$  defining the three problems. To make it easy to detect errors in the solution, the right-hand side has been chosen to make the solution a vector of all 1's.

The program below is an encoding of an iterative refinement algorithm which may be used to obtain a highly accurate solution to a system of linear equations with an ill-conditioned matrix. When the condition number is not excessively high, the iterative refinement algorithm usually returns a solution that is accurate to the working precision of the machine. The iterative refinement algorithm is essentially:

- (1) Solve  $Ax = b$
- (2) Set  $tol = \epsilon \sum |x_i|$   
where  $\epsilon$  is the precision of the machine
- (3) Compute in double precision the residual,  $Ax-b$ , and put it in the real vector  $r$ .
- (4) Solve  $A \delta x = r$
- (5) Compute  $norm = \sum |\delta x_i|$
- (6) Set  $x$  to  $x + \delta x$
- (7) If  $norm \leq tol$  stop, else return to step 3

In our code, step (1) is accomplished using the the linear equation solver for band positive definite matrices, BPLE. The subroutine BPLE leaves in the  $G$  matrix a decomposition which may be used by BPFS and BPBS to solve problems with the same coefficient matrix but different right-hand sides as in step(4). Since it is possible that the matrix will be so ill-conditioned that the iterative refinement algorithm will diverge, steps (3) through (7) in our code are performed only a finite number of times. This number is chosen to be an upper bound on number of bits in the mantissa of the floating-point number supported by the machine.

This algorithm is not included in PORT because for double-precision matrices part of the computation would have to be done in extended precision.

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BPFS

**Note 2:** Users who have to solve a sequence of problems with the same coefficient matrix, but different right-hand sides, *not all known in advance*, should not call BPSS or BPLE repeatedly, but should use BPDC to find the  $LDL^T$  decomposition of the coefficient matrix, which can then be used repeatedly (and efficiently) for the sequence of forward solutions (using BPFS) and back solutions (using BPBS) for each set of right-hand sides.

**Error situations:** (All errors in this subprogram are fatal — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$ML < 1$
3	$IG < ML$
4	$IB < N$
5	$NB < 1$

**Double-precision version:** DBPFS, with G and B declared double precision

**Complex Hermitian version:** CBPFS with G and B declared complex  
G should contain the conjugate of L

**Storage:** None

**Time:**  $N \times (ML - 1) \times NB$  additions  
 $N \times (ML - 1) \times NB$  multiplications

**See also:** BPBS, BPCE, BPDC, BPLD, BPLE, BPSS

**Author:** Linda Kaufman

## BPFS — band symmetric lower (unit) triangular linear system solution

**Purpose:** BPFS (Banded symmetric Positive definite matrix Forward Solution) solves  $LX = B$  where  $L$  is a banded unit lower triangular matrix, (i.e. 1's on the diagonal, 0's above the diagonal). BPFS can be used for the forward solution phase of a band symmetric positive definite linear system. (It is used in this way by the routines BPSS and BPLE.)

**Usage:** CALL BPFS (N, ML, G, IG, B, IB, NB)

N → the number of equations

ML → the number of nonzero diagonals on and below the diagonal of  $L$

G → a matrix (which may contain the results obtained by the routines BPCE, BPDC, or BPLD) into which  $L$  is packed as follows:

$$G(1+i-j, i) = l_{ij} \text{ for } i > j$$

(See the introduction to this chapter.)

$G$  should be dimensioned (IG,KG) in the calling program, where  $IG \geq ML$  and  $KG \geq N$ .

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

B → the matrix of right-hand sides, dimensioned (IB, KB) in the calling program, where  $IB \geq N$  and  $KB \geq NB$ .

← the solution  $X$

IB → the row (leading) dimension of  $B$ , as dimensioned in the calling program

NB → the number of right-hand sides

**Note 1:** BPFS and BPBS can be used directly on the output matrix produced by BPDC, BPLD, or BPCE to solve a general linear system.

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BPDC

```
TIME FOR BPDC  6450
TIME FOR BPCE  13170

N IS  48 ,NUMBER OF UPPER DIAGONALS IS 17
TIME FOR BPLD  7143
TIME FOR BPDC  8227
TIME FOR BPCE  12722

N IS  96 ,NUMBER OF UPPER DIAGONALS IS 17
TIME FOR BPLD  15001
TIME FOR BPDC  17160
TIME FOR BPCE  49809
```

As the example indicates, the cost of computing the condition estimate can be substantially greater than the cost of factoring the matrix when the width of the band is small. The ratio of BPCE to BPDC does not always decrease as  $N$  increases, as it does in the case of general linear systems, but depends rather on the scaling that is sometimes done by BPCE to prevent overflow during the calculation of the condition estimate. If no scaling is done, the ratio of the times for BPCE to BPDC remains constant, but for some examples most of the time is spent scaling and the time ratio increases as  $N$  increases.

```

30      CONTINUE
        DO 50 I=1,N
          DO 40 J=1,MU
            G2(J,I)=G(J,I)
            G3(J,I)=G(J,I)
40      CONTINUE
50      CONTINUE
        WRITE(IWRITE,51)N,MU
51      FORMAT(/6H N IS ,I4,30H ,NUMBER OF UPPER DIAGONALS IS,I3)
C TIME DECOMPOSITION BY BPLD
        IT=ILAPSZ(0)
        CALL BPLD(N,MU,G,IG,0.0)
        IT=ILAPSZ(0)-IT
        WRITE(IWRITE,52)IT
52      FORMAT(14H TIME FOR BPLD,I7)
C TIME DECOMPOSITION BY BPDC
        IT2=ILAPSZ(0)
        CALL BPDC(N,MU,G2,IG)
        IT2=ILAPSZ(0)-IT2
        WRITE(IWRITE,53)IT2
53      FORMAT(14H TIME FOR BPDC,I7)
C TIME DECOMPOSITION BY BPCE
        IT3=ILAPSZ(0)
        CALL BPCE(N,MU,G3,IG,COND)
        IT3=ILAPSZ(0)-IT3
        WRITE(IWRITE,54)IT3
54      FORMAT(14H TIME FOR BPCE,I7)
60      CONTINUE
        MLM1=MLM1*2
70     CONTINUE
        STOP
        END

```

When the above code was run on the Honeywell 6000 at Bell Laboratories with an optimizing compiler the following was printed

```

N IS   48 ,NUMBER OF UPPER DIAGONALS IS  5
TIME FOR BPLD   959
TIME FOR BPDC  1413
TIME FOR BPCE  3670

N IS   96 ,NUMBER OF UPPER DIAGONALS IS  5
TIME FOR BPLD  1868
TIME FOR BPDC  2673
TIME FOR BPCE  7965

N IS   48 ,NUMBER OF UPPER DIAGONALS IS  9
TIME FOR BPLD  2395
TIME FOR BPDC  3181
TIME FOR BPCE  6377

N IS   96 ,NUMBER OF UPPER DIAGONALS IS  9
TIME FOR BPLD  4854

```

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BPDC

The matrix in the example is derived from the traditional 5-point approximation to the Laplace operator on the unit square. The  $N \times N$  matrix A has the form

$$\begin{array}{ccccccc} C & -I & & & & & \\ -I & C & -I & & & & \\ & -I & C & -I & & & \\ & & & \dots\dots & & & \\ & & & & -I & C & -I \\ & & & & & -I & C \end{array}$$

where I is the identity matrix and C is the matrix

$$\begin{array}{cccc} 4 & -1 & & \\ -1 & 4 & -1 & \\ & -1 & 4 & -1 \\ & & & \dots \\ & & & -1 & 4 & -1 \\ & & & & -1 & 4 \end{array}$$

In this problem we vary the sizes of the blocks C and I which changes the bandwidth. For a given blocksize, the number of blocks is varied which changes N.

The subroutine ILAPSZ is a timer on the Honeywell 6000 with about 1% accuracy. It counts in 1/64 milliseconds.

```

INTEGER IG, MLM1, IWRITE, ILMACH, K, N, MU
INTEGER NBLOK, KBLOK, KK, I, J, IT, ILAPSZ, IT2
REAL G(17, 100), G2(17, 100), G3(17, 100)
REAL COND
IG=17
MLM1=4
IWRITE=IIMACH(2)
DO 70 K=1,3
  DO 60 N=48,96,48
    MU=MLM1+1
    I=0
    NBLOK=N/MLM1
C
C SET UP THREE MATRICES FOR ELLIPTIC PDE IN 2 DIMENSION
C
    DO 30 KBLOK=1,NBLOK
      DO 20 KK=1,MLM1
        I=I+1
        G(1,I)=4.0
        G(2,I)=-1.0
        G(MU,I)=-1.0
        DO 10 J=3,MLM1
          G(J,I)=0.0
10          CONTINUE
20          CONTINUE
          G(2,I)=0.0

```



**Error situations:** \*(The user can elect to ‘recover’ from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$
$10 + k^*$	singular matrix whose rank is at least $k$
$10 + N + k^*$	the $k^{\text{th}}$ principal minor is not positive definite

**Double-precision version:** DBPDC, with  $G$  declared double precision.

**Complex Hermitian version:** CBPDC with  $G$  declared complex (see *Note 2*).

**Storage:** None

**Time:**  $N \times (MU - 1) \times MU / 2 + N \times (2MU - 1)$  additions  
 $N \times (MU - 1) \times MU / 2$  multiplications  
 $N \times (MU - 1)$  divisions

**Method:** BPDC calls BPLD after setting  $EPS = \|A\| \epsilon$ , where  $\epsilon$  is machine precision, i.e. the value returned by R1MACH(4) (or, for double precision, by DIMACH(4)).

**See also:** BPBS, BPCE, BPFs, BPLD, BPLE, BPSS

**Author:** Linda Kaufman

**Example:** This example is designed to indicate the relative efficiency of BPDC, BPLD, and BPCE as a function of the width of the band. All three subroutines compute the same factorization, but the criterion for singularity is treated differently in each of the three. In all the subroutines the matrix is considered singular if for some  $i$ ,  $d_i \leq EPS$ . In BPLD the user provides EPS; in BPDC the subroutine computes EPS (see *Method*); in BPCE, 0.0 is used as EPS. (However, BPCE also provides an estimate of the condition number of the matrix.)

BPDC —  $LDL^T$  decomposition of a band symmetric positive definite matrix A

**Purpose:** BPDC (Banded symmetric Positive definite matrix DeComposition) decomposes a banded symmetric positive definite matrix A into  $LDL^T$  where L is lower unit triangular (1's on the diagonal and 0's above the diagonal) and D is diagonal. It is called by BPLE as the first step of the solution of a banded symmetric positive definite linear system.

**Usage:** CALL BPDC (N, MU, G, IG)

N → the number of equations

MU → the number of nonzero bands on and above the diagonal of A

G → a matrix into which the upper triangular portion of the matrix A has been packed as follows:

$$G(j-i+1, i) = a_{ij} \text{ for } j \geq i$$

(See the introduction to the chapter.)

G should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

←  $LDL^T$  decomposition suitable for input into BPFS and BPBS (see Note 1)

IG → the row (leading) dimension of G, as dimensioned in the calling program

**Note 1:** The  $LDL^T$  decomposition of A satisfies the equation  $A = LDL^T$  where L is lower unit triangular (1's on the diagonal, 0's above the diagonal) and D is diagonal. On return from BPDC, the diagonal of D occupies the first row of G and  $G(i-j+1, i) = l_{ij}$  for  $i > j$ .

**Note 2:** For complex Hermitian matrices ( $A = A^*$ , where  $A^*$  represents the conjugate transpose of A), the complex version of this subroutine computes the  $LDL^*$  decomposition and returns the conjugate of L, rather than L, in G.

```

C SOLVE AX=B TO GET ITH COLUMN OF A (INVERSE)
      CALL BPFS(N,MU,G,IG,B,N,1)
      CALL BPBS(N,MU,G,IG,B,N,1)
C FIND NORM OF COLUMN
      AINORM=0.0
      DO 30 J=1,N
        AINORM=AINORM+ABS(B(J))
30      CONTINUE
      IF (AINVNO.LT.AINORM) AINVNO=AINORM
40      CONTINUE
      COND=4.0*AINVNO
      WRITE(IWRITE,41)COND
41      FORMAT(25H TRUE CONDITION NUMBER IS,E15.8)
      X=X/100.0
50      CONTINUE
      STOP
      END

```

When the above code was executed on the Honeywell 6000 machine at Bell Laboratories, the following was printed:

```

WHEN X IS 0.100000E 01
CONDITION ESTIMATE IS 0.40807862E 04
TRUE CONDITION NUMBER IS 0.50999824E 04

WHEN X IS 0.100000E-01
CONDITION ESTIMATE IS 0.23329148E 05
TRUE CONDITION NUMBER IS 0.24899523E 05

WHEN X IS 0.100000E-03
CONDITION ESTIMATE IS 0.19933923E 07
TRUE CONDITION NUMBER IS 0.19950487E 07

```

The comparison above of the condition number estimated by BPCE with the true condition number indicates that the order of magnitude (which is all one usually is interested in) of the estimated condition number is correct. Note that the inverse of a band matrix is usually a full  $n \times n$  matrix and should rarely be calculated.

**Example:** In the following example we obtain estimates of the condition numbers of the matrix

$$\begin{array}{ccccccc} 1+x & -1 & & & & & \\ -1 & 2 & -1 & & & & \\ & -1 & 2 & -1 & & & \\ & & -1 & 2 & -1 & & \\ & & & & & & \cdot \\ & & & & & & \cdot \\ & & & & & -1 & 2 & -1 \\ & & & & & -1 & & 1+x \end{array}$$

with  $x=1., .01, \text{ and } .0001$ . The matrix is singular when  $x$  is 0, and becomes more ill-conditioned as  $x$  approaches 0.

In this example we compare the condition number  $K = \|A\| \|A^{-1}\|$  with the estimate obtained from BPCE. In the code below  $A^{-1}$  is computed one column at a time and  $K$  is computed using the 1-norm. In the 1-norm,  $\|A\|$  is the maximum column sum in absolute value, which is obviously 4.

```

      INTEGER N, MU, IG, K, I, IWRITE, ILMACH, J
      REAL G(2,100), B(200)
      REAL X, COND, AINVNO, AINORM, ABS
      N=100
      X=1.0
      MU=2
      IG=2
      DO 50 K=1,3
C CONSTRUCT MATRIX
      DO 10 I=1,N
        G(1,I)=2.0
        G(2,I)=-1.0
10      CONTINUE
        G(1,1)=1.0+X
        G(1,N)=1.0+X
C GET ESTIMATE OF CONDITION NUMBER FROM BPCE
      CALL BPCE(N,MU,G,IG,COND)
      IWRITE=ILMACH(2)
      WRITE(IWRITE,11)X
11      FORMAT(/10H WHEN X IS,E14.6)
      WRITE(IWRITE,12)COND
12      FORMAT(25H CONDITION ESTIMATE IS ,E15.8)
C SINCE CONDITION NUMBER IS NORM(A)*NORM(INVERSE(A)),
C FIND THE NORM OF EACH COLUMN OF INVERSE(A). GENERATE
C THE COLUMNS ONE AT A TIME AND REUSE SPACE
      AINVNO=0.0
      DO 40 I=1,N
C GENERATE ITH COLUMN OF IDENTITY MATRIX AS RIGHT HAND SIDE
      DO 20 J=1,N
        B(J)=0.0
20      CONTINUE
        B(I)=1.0

```

**Error situations:** \*(The user can elect to ‘recover’ from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$MU < 1$
3	$IG < MU$
$10 + k^*$	singular matrix whose rank is at least $k$
$10 + N + k^*$	the $k^{\text{th}}$ principal minor is not positive definite

**Double-precision version:** DBPCE with  $G$  and  $COND$  declared double precision

**Complex Hermitian version:** CBPCE with  $G$  declared complex (see **Note 3** above).

**Storage:**  $N$  real (double precision for DBPCE, complex for CBPCE) locations of scratch storage in the dynamic storage stack

**Time:**  $N \times ((MU-1) \times (10+MU/2)+8)$  additions  
 $N \times ((MU-1) \times (6+MU/2)+4)$  multiplications  
 $N \times (MU+1)$  divisions

**Method:** Gaussian elimination without pivoting  
 See the reference below for the method used to estimate the condition number.  
 BPCE calls BPLD with  $EPS=0.0$

**See also:** BPBS, BPDC, BPFS, BPLD, BPLE, BPSS

**Authors:** Doris Ryan and Linda Kaufman

**Reference:** Cline, A. K., Moler, C. B., Stewart, G. W., and Wilkinson, J. H., An estimate for the condition number, *SIAM J. Numer. Anal.* 16 (1979), 368-375.

BPCE —  $LDL^T$  decomposition with condition estimation

**Purpose:** BPCE (Banded symmetric Positive definite matrix Condition Estimation) gives a lower bound for the condition number of a banded symmetric positive definite matrix A. It also returns the  $LDL^T$  decomposition of A and may be used in a linear equation package.

**Usage:** CALL BPCE (N, MU, G, IG, COND)

N → the order of the matrix A

MU → the number of nonzero bands on and above the diagonal of A

G → a matrix into which the upper triangular portion of the matrix A has been packed as follows:

$$G(j-i+1, i) = a_{ij} \text{ for } j \geq i$$

(See the introduction to this chapter.)

G should be dimensioned (IG,KG) in the calling program, where  $IG \geq MU$  and  $KG \geq N$ .

←  $LDL^T$  decomposition suitable for input into BPFS and BPBS (see Note 2)

IG → the row (leading) dimension of G, as dimensioned in the calling program

COND ← an estimate of the condition number of A (see Note 1)

**Note 1:** The condition number measures the sensitivity of the solution of a linear system to errors in the matrix and in the right-hand side. If the elements of the matrix and the right-hand side(s) of your linear system have **d** decimal digits of precision, the solution might have as few as  $\mathbf{d} - \log_{10}(\text{COND})$  correct decimal digits. Thus if COND is greater than  $10^{\text{BdP}}$ , there may be no correct digits.

**Note 2:** The  $LDL^T$  decomposition of A satisfies the equation  $A = LDL^T$  where L is lower unit triangular (1's on the diagonal, 0's above the diagonal) and D is diagonal. On return from BPCE, the diagonal of D occupies the first row of G and  $G(i-j+1, i) = l_{ij}$  for  $i > j$ .

**Note 3:** For complex Hermitian matrices ( $A = A^*$ , where  $A^*$  represents the conjugate transpose of A), the complex version of this subroutine computes the  $LDL^*$  decomposition and returns the conjugate of L rather than L in G.

**Example:** The program fragment below solves a linear system  $AX = B$ , where  $A$  is a symmetric positive definite band matrix. It is assumed that the  $A$  matrix has been packed into  $G$  according to the scheme  $G(j-i+1, i) = a_{ij}$ . The subroutine `BPCE` factors  $A$  into  $LDL^T$ , where  $L$  is unit lower triangular and  $D$  is diagonal. The factors are returned in  $G$  so that `BPFS` can forward solve (solve  $LY=B$ ) and `BPBS` can back solve (solve  $DL^TX=Y$ ).

The subroutine `BPCE` also provides an estimate of the condition number of  $A$ . In the code below if the condition number is larger than the reciprocal of the machine precision (given by `R1MACH(4)`), the matrix is considered too ill-conditioned and the system is not solved.

```

                IWRITE=I1MACH(2)
                CALL BPCE(N,ML,G,IG,COND)
                IF (COND .GT. 1.0/R1MACH(4)) GO TO 10
                CALL BPFS(N,ML,G,IG,B,IB,NB)
                CALL BPBS(N,ML,G,IG,B,IB,NB)
                GO TO 20
10              WRITE(IWRITE,11)
11              FORMAT(26H MATRIX TOO ILL-CONDITIONED)
20              CONTINUE

```

**Error situations:** \*(The user can elect to ‘recover’ from those errors marked with an asterisk — see *Error Handling*, Framework Chapter)

Number	Error
1	$N < 1$
2	$ML < 1$
3	$IG < ML$
4	$IB < N$
5	$NB < 1$
$10 + k^*$	singular D with kth diagonal element 0.0

**Double-precision version:** DBPBS, with G and B declared double precision

**Complex Hermitian version:** CBPBS with G and B declared complex

**Storage:** None

**Time:**  $NB \times (ML - 1) \times N$  additions  
 $NB \times (ML - 1) \times N$  multiplications  
 $NB \times N$  divisions

**See also:** BPCE, BPDC, BPFs, BPLD, BPLE, BPSS

**Author:** Linda Kaufman



**Purpose:** BPBS (Banded symmetric Positive definite matrix Back Solve) solves  $DRX = B$  where  $D$  is diagonal and  $R$  is banded unit upper triangular (1's on the diagonal and 0's below the diagonal). It can be used for the back solution phase of a banded linear system solution. (It is used in this way by the routines BPSS and BPLE.)

**Usage:** CALL BPBS (N, ML, G, IG, B, IB, NB)

N → the number of equations

ML → the number of nonzero diagonals of  $R$  (including the unit diagonal)

G → a matrix (which may contain results obtained by the routines BPLD, BPCE, or BPDC) into which  $D$  and  $R$  are packed as follows:

$$\begin{aligned} G(1, i) &= d_i \\ G(j-i+1, i) &= r_{ij} \text{ for } j > i \end{aligned}$$

(See the introduction to this chapter.)

$G$  should be dimensioned (IG,KG) in the calling program, where  $IG \geq ML$  and  $KG \geq N$ .

IG → the row (leading) dimension of  $G$ , as dimensioned in the calling program

B → the matrix of right-hand sides, dimensioned (IB, KB) in the calling program, where  $IB \geq N$  and  $KB \geq NB$ .

← the solution  $X$

IB → the row (leading) dimension of  $B$ , as dimensioned in the calling program

NB → the number of right-hand sides

**Note:** BPFS and BPBS can be used directly on the output matrix produced by BPDC, BPLD, or BPCE to solve a banded symmetric positive definite linear system.

## BANDED&lt; SYMMETRIC&lt; POSITIVE-DEFINITE MATRICES

BPBS - Back Solve  
BPCE - Condition Estimation  
BPDC - DeComposition  
BPFS - Forward Solve  
BPLE - Linear Equation solution  
BPLD -  $LDL^T$  decomposition  
BPML - MuLtiplication  
BPNM - NorM  
BPSS - System Solution