

```
c=====
c      Test program for LAPACK "driver" routine 'dgesv'
c      which computes the solution of a real system
c      of linear equations:  A x = b
c
c      This version uses fixed-size 2-d arrays (size fixed at
c      some maximum value commensurate with needs and/or
c      available memory), illustrating another commonly used
c      Fortran technique to implement run-time dimensioning,
c      PARTICULARLY FOR RANK-2 ARRAYS.
c
c      This time the rules are as follows: All subroutines and
c      functions which manipulate the array must be passed:
c
c      (1) The array itself.
c      (2) The "true" or "physical" dimensions;
c           i.e. the dimensions in MAIN (*).
c      (3) The "run-time" or "logical" dimensions (*).
c
c      (*) More precisely, due to the nature of the FORTRAN
c      subscripting computation, the leading d-1 dimensions
c      must be passed for a rank-d array. In particular,
c      for rank-2 array (matrices), THE leading physical
c      dimension (often denoted 'LDA' in LAPACK code), and
c      THE leading logical dimension (often denoted 'N')
c      must BOTH be passed.
c
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c      Passing the physical dimensions ensures that the
c      linearization/subscripting calculation is identical
c      in all program units INCLUDING MAIN---so that, e.g.,
c      one can safely and conveniently use a(i,j) etc. in
c      MAIN.
c
c      Passing the logical dimensions allows us to write
c      routines which function for a general case (here,
c      typically for N x N matrices).
c
c      Passing BOTH sets of dimensions is slightly cumbersome,
c      but is the price we pay in this case for convenience
c      and generality.
c=====
program          tdgesv1

      implicit      none
c-----
c      Maximum size of linear system.
c-----
      integer        maxn
      parameter      ( maxn = 100 )

c-----
c      Storage for arrays.
c-----
      real*8         a(maxn,maxn),
      &                  b(maxn)
      integer        ipiv(maxn)

      integer        i,           nrhs,
      &                  n,           info

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c-----
c      Set up sample 3 x 3 system ...
c-----
a(1,1) =  1.23d0
a(1,2) =  0.24d0
a(1,3) = -0.45d0

a(2,1) = -0.43d0
a(2,2) =  2.45d0
a(2,3) =  0.78d0

a(3,1) =  0.51d0
a(3,2) = -0.68d0
a(3,3) =  3.23d0

b(1)   =  6.78d0
b(2)   = -3.45d0
b(3)   =  1.67d0

c-----
c      ... and solve it. Note that 'dgsev' is general
c      enough to solve A x_i = b_i for multiple right-hand-
c      sides b_i. Here we have only one right-hand-side.
c      Also note that the procedure performs the LU
c      decomposition in place, thus destroying the
c      input-matrix, it also overwrites the right-hand-side(s)
c      with the solution(s). Finally, observe that we
c      pass the "leading dimension" (maxn) of both 'a' and
c      'b' to the routine. Again, this allows us to load array
c      elements in the main program as we have just done,
c      without running into troubles due to the fact that
c      these elements ARE NOT, in general all contiguous in
c      memory. This certainly includes the current 3 x 3 case.
c-----

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n      = 3
nrhs = 1

call dgesv( n, nrhs, a, maxn, ipiv, b, maxn, info )

if(      info .eq. 0 ) then
c-----
c      Solution successful, write soln to stdout.
c      Note the use of "implied-do-loop" to write a
c      sequence of elements: the enclosing parenthesis
c      around the "loop" are required.
c-----
c-----      write(*,*) ( b(i) , i = 1 , n )
else if( info .lt. 0 ) then
c-----
c      Bad argument detected.
c-----
c-----      write(0,*) 'tdgesv1: Argument ', abs(info),
&                  ' to dgesv() is invalid'
else
c-----
c      Matrix is singular.
c-----
c-----      write(0,*) 'tdgesv1: dgesv() detected singular ',
&                  'matrix'
end if

stop

end

```

```

SUBROUTINE DGESV( N, NRHS, A, LDA, IPIV, B, LDB, INFO )
*
* -- LAPACK driver routine (version 2.0) --
* Univ. of Tennessee, Univ. of California Berkeley, NAG Ltd.,
* Courant Institute, Argonne National Lab, and Rice University
* March 31, 1993
*
* .. Scalar Arguments ..
INTEGER           INFO, LDA, LDB, N, NRHS
*
* ..
* .. Array Arguments ..
INTEGER           IPIV( * )
DOUBLE PRECISION  A( LDA, * ), B( LDB, * )
*
* ..
*
* Purpose
* ======
*
* DGESV computes the solution to a real system of linear equations
* 
$$A * X = B,$$

* where A is an N-by-N matrix and X and B are N-by-NRHS matrices.
*
* The LU decomposition with partial pivoting and row interchanges is
* used to factor A as
* 
$$A = P * L * U,$$

* where P is a permutation matrix, L is unit lower triangular, and U is
* upper triangular. The factored form of A is then used to solve the
* system of equations  $A * X = B$ .
*
* Arguments
* ======
*
* N      (input) INTEGER
*        The number of linear equations, i.e., the order of the

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*
*      matrix A.  N >= 0.
*
*      NRHS  (input) INTEGER
*              The number of right hand sides, i.e., the number of columns
*              of the matrix B.  NRHS >= 0.
*
*      A      (input/output) DOUBLE PRECISION array, dimension (LDA,N)
*              On entry, the N-by-N coefficient matrix A.
*              On exit, the factors L and U from the factorization
*              A = P*L*U; the unit diagonal elements of L are not stored.
*
*      LDA   (input) INTEGER
*              The leading dimension of the array A.  LDA >= max(1,N).
*
*      IPIV  (output) INTEGER array, dimension (N)
*              The pivot indices that define the permutation matrix P;
*              row i of the matrix was interchanged with row IPIV(i).
*
*      B      (input/output) DOUBLE PRECISION array, dimension (LDB,NRHS)
*              On entry, the N-by-NRHS matrix of right hand side matrix B.
*              On exit, if INFO = 0, the N-by-NRHS solution matrix X.
*
*      LDB   (input) INTEGER
*              The leading dimension of the array B.  LDB >= max(1,N).
*
*      INFO  (output) INTEGER
*              = 0:  successful exit
*              < 0:  if INFO = -i, the i-th argument had an illegal value
*              > 0:  if INFO = i, U(i,i) is exactly zero.  The factorization
*                    has been completed, but the factor U is exactly
*                    singular, so the solution could not be computed.
*
* =====

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*     .. External Subroutines ..
      EXTERNAL          DGETRF, DGETRS, XERBLA
*
*     ..
*     .. Intrinsic Functions ..
      INTRINSIC        MAX
*
*     ..
*     .. Executable Statements ..
*
*     Test the input parameters.
*
      INFO = 0
      IF( N.LT.0 ) THEN
         INFO = -1
      ELSE IF( NRHS.LT.0 ) THEN
         INFO = -2
      ELSE IF( LDA.LT.MAX( 1, N ) ) THEN
         INFO = -4
      ELSE IF( LDB.LT.MAX( 1, N ) ) THEN
         INFO = -7
      END IF
      IF( INFO.NE.0 ) THEN
         CALL XERBLA( 'DGESV ', -INFO )
         RETURN
      END IF
*
*     Compute the LU factorization of A.
*
      CALL DGETRF( N, N, A, LDA, IPIV, INFO )
      IF( INFO.EQ.0 ) THEN
*
*     Solve the system A*X = B, overwriting B with X.
*
      CALL DGETRS( 'No transpose', N, NRHS, A, LDA, IPIV, B, LDB,
$                           INFO )

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END IF
RETURN
*
*      End of DGESV
*
END
```



```
#####
# Building 'tdgesv' and sample output on physics, a Sun 4
# Ultra-Enterprise running SunOS 5.6
#####
physics% pwd; ls
/export/ugrad/phys410/linsys/ex1
Makefile    tdgesv.f

physics% make
f77 -O -c tdgesv.f
tdgesv.f:
MAIN tdgesv1:
f77 -O -L/home5/choptuik/lib tdgesv.o -llapack -o tdgesv
Undefined first referenced
symbol      in file
dscal_          /usr/local/lib/liblapack.a(dgetf2.o)
dswap_          /usr/local/lib/liblapack.a(dlaswp.o)
dtrsm_          /usr/local/lib/liblapack.a(dgetrf.o)
idamax_         /usr/local/lib/liblapack.a(dgetf2.o)
dgemm_          /usr/local/lib/liblapack.a(dgetrf.o)
dger_           /usr/local/lib/liblapack.a(dgetf2.o)
ld: fatal: Symbol referencing errors. No output written to tdgesv
make: [tdgesv] Error 1 (ignored)
#####
# OOPS ... those are references to BLAS routines! I haven't
# defined the environment variable LIBBLAS which is used in the
# Makefile. Best to add a line in ~/.cshrc.user. Arguably,
# '-lblas' should just be in the Makefile. However, on SOME
# systems, BLAS is effectively "built-in", and then explicit
# reference to it can cause problems at load time. With the
# current mechanism, we can easily deal with such a case
# simply by leaving the environment variable undefined on
# those systems.
#####
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physics% vi ~/.cshrc.user

INSERT ---->    setenv LIBBLAS '-lblas'

#####
# "Activate" the changed .cshrc_user; among other things this
# will set LIBBLAS properly.
#####
physics% source !$
source ~/.cshrc.user

[47]physics{phys410} set prompt="physics% "

physics% pwd; ls
/export/ugrad/phys410/linsys/ex1
Makefile  tdgesv.f  tdgesv.o

#####
# This time the build works ...
#####
physics% make
f77 -O -L/home5/choptuik/lib tdgesv.o -llapack -lblas -o tdgesv

#####
# ... and we get output consistent with the other systems.
#####
physics% tdgesv
5.4263644124316  -0.32577537681739  -0.40835080698946

```

```
c=====
c      Solves 1-d linear boundary value problem
c
c      u''(x) = f(x)   on   x = [0,1] ; u(0) = u0, u(1) = u1
c
c      using second-order finite difference technique and
c      LAPACK tridiagonal solver DGTSV.
c=====
```

program bvp1d

implicit none

integer i4arg

```

c-----
c      Extrema of problem domain; note that this approach
c      of defining extrema as parameters makes it easier
c      to generalize program to arbitrary domains.
c-----

      real*8           xmin,           xmax
      parameter        ( xmin = 0.0d0,  xmax = 1.0d0 )

c-----
c      Maximum problem size (2**20 + 1)
c-----

      integer          maxn
      parameter        ( maxn = 1 048 577 )

c-----
c      Storage for discrete x-values, unknowns, exact
c      solution and right hand side values
c-----

      real*8           x(maxn),       u(maxn),
      &                   ueexact(maxn),   f(maxn)

c-----
c      Storage for main, upper and lower diagonals of
c      tridiagonal system, and right-hand-side vector
c      for use with LAPACK routine DGTSV
c-----

      real*8           d(maxn),       du(maxn),
      &                   dl(maxn),       rhs(maxn)
      integer          nrhs,          info

c-----
c      Discretization level and size of system (# of discrete
c      unknowns)
c-----

      integer          level,          n,
      &                   option

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c-----
c      Mesh spacing and related constants (1/h**2, -2/h**2)
c-----
      real*8          h,                  hm2,                  m2hm2
      real*8          rmserr

c-----
c      Argument parsing.
c-----
      level = i4arg(1,-1)
      if( level .lt. 0 ) go to 900
      n = 2 ** level + 1
      if( n .gt. maxn ) then
          write(0,*) 'Insufficient internal storage'
          stop
      end if
      option = i4arg(2,0)
c-----
c      Set up finite-difference 'mesh' (discrete x-values)
c      and define some useful constants.
c-----
      h      = 1.0d0 / (n - 1)
      do j = 1 , n
          x(j) = xmin + (j - 1) * h
      end do
      hm2   = 1.0d0 / (h * h)
      m2hm2 = -2.0d0 / (h * h)

c-----
c      This only ensures that x(n) = xmax EXACTLY and is not
c      essential.
c-----
      x(n) = xmax

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c-----
c      Set up exact solution and right hand side vector.
c-----
c      call exact(uexact,f,x,n)

c=====
c      Set up tridiagonal system. Note that indexing on
c      lower diagonal is always (j-1) when implementing the
c      j'th equation.
c=====

c-----
c      Left boundary: u(1) = u_0
c-----
d(1)      = 1.0d0
du(1)     = 0.0d0
rhs(1)    = uexact(1)
c-----
c      Interior: Second order FDA of ODE.
c-----
do j = 2 , n - 1
  dl(j-1) = hm2
  d(j)    = m2hm2
  du(j)   = hm2
  rhs(j)  = f(j)
end do
c-----
c      Right boundary: u(n) = u_1
c-----
dl(n-1)   = 0.0d0
d(n)      = 1.0d0
rhs(n)    = uexact(n)

```

```

c=====
c      Solve tridiagonal system.
c=====

nrhs = 1
call dgtsv( n, nrhs, dl, d, du, rhs, n, info )

if( info .eq. 0 ) then
c-----
c      Solver successful, output either (x_j, u_j) or
c      (x_j, error_j) to stdout. Also compute rms error
c      and output to standard error.
c-----

rmserr = 0.0d0
do j = 1 , n
    if( option .eq. 0 ) then
        write(*,*) x(j), rhs(j)
    else
        write(*,*) x(j), (uexact(j) - rhs(j))
    end if
    rmserr = rmserr + (uexact(j) - rhs(j)) ** 2
end do
rmserr = sqrt(rmserr / n)
write(0,*) 'rmserr = ', rmserr
else
c-----
c      Solver failed.
c-----

write(0,*) 'bvp1d: dgtsv() failed, info = ', info
end if

stop

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900 continue
    write(0,*) 'usage: bvp1d <level> [<option>]'
    write(0,*)
    write(0,*) '          Specify option .ne. 0 for output'
    write(0,*) '          of error instead of solution'
    stop

end

c=====
c      Computes exact values for u(x) (unknown function)
c      and f(x) (right hand side function).  x array must
c      have been previously defined.
c=====

subroutine exact(u,f,x,n)

implicit none
integer n
real*8 u(n), f(n), x(n)

real*8 pi2
integer j

pi2 = 8.0d0 * atan(1.0d0)
do j = 1 , n
    u(j) = sin(pi2 * x(j))
    f(j) = -pi2 * pi2 * u(j)
end do

return

end

```

```

        SUBROUTINE DGTSV( N, NRHS, DL, D, DU, B, LDB, INFO )
*
* -- LAPACK routine (version 2.0) --
* Univ. of Tennessee, Univ. of California Berkeley, NAG Ltd.,
* Courant Institute, Argonne National Lab, and Rice University
* September 30, 1994
*
* .. Scalar Arguments ..
INTEGER           INFO, LDB, N, NRHS
*
* ..
* .. Array Arguments ..
DOUBLE PRECISION  B( LDB, * ), D( * ), DL( * ), DU( * )
*
* ..
*
* Purpose
* ======
*
* DGTSV solves the equation
*
*      A*X = B,
*
* where A is an N-by-N tridiagonal matrix, by Gaussian elimination with
* partial pivoting.
*
* Note that the equation A'*X = B may be solved by interchanging the
* order of the arguments DU and DL.
*
* Arguments
* ======
*
* N          (input) INTEGER
*             The order of the matrix A.  N >= 0.
*
* NRHS       (input) INTEGER

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*
*   The number of right hand sides, i.e., the number of columns
*   of the matrix B.  NRHS >= 0.
*
*   DL      (input/output) DOUBLE PRECISION array, dimension (N-1)
*   On entry, DL must contain the (n-1) subdiagonal elements of
*   A.
*   On exit, DL is overwritten by the (n-2) elements of the
*   second superdiagonal of the upper triangular matrix U from
*   the LU factorization of A, in DL(1), ..., DL(n-2).
*
*   D      (input/output) DOUBLE PRECISION array, dimension (N)
*   On entry, D must contain the diagonal elements of A.
*   On exit, D is overwritten by the n diagonal elements of U.
*
*   DU     (input/output) DOUBLE PRECISION array, dimension (N-1)
*   On entry, DU must contain the (n-1) superdiagonal elements
*   of A.
*   On exit, DU is overwritten by the (n-1) elements of the first
*   superdiagonal of U.
*
*   B      (input/output) DOUBLE PRECISION array, dimension (LDB,NRHS)
*   On entry, the N-by-NRHS right hand side matrix B.
*   On exit, if INFO = 0, the N-by-NRHS solution matrix X.
*
*   LDB    (input) INTEGER
*   The leading dimension of the array B.  LDB >= max(1,N).
*
*   INFO   (output) INTEGER
*   = 0:  successful exit
*   < 0:  if INFO = -i, the i-th argument had an illegal value
*   > 0:  if INFO = i, U(i,i) is exactly zero, and the solution
*         has not been computed.  The factorization has not been
*         completed unless i = N.
*

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* =====
*
* .. Parameters ..
    DOUBLE PRECISION    ZERO
    PARAMETER          ( ZERO = 0.0D+0 )
*
* ..
* .. Local Scalars ..
    INTEGER             J, K
    DOUBLE PRECISION    MULT, TEMP
*
* ..
* .. Intrinsic Functions ..
    INTRINSIC           ABS, MAX
*
* ..
* .. External Subroutines ..
    EXTERNAL            XERBLA
*
* ..
* .. Executable Statements ..
*
    INFO = 0
    IF( N.LT.0 ) THEN
        INFO = -1
    ELSE IF( NRHS.LT.0 ) THEN
        INFO = -2
    ELSE IF( LDB.LT.MAX( 1, N ) ) THEN
        INFO = -7
    END IF
    IF( INFO.NE.0 ) THEN
        CALL XERBLA( 'DGTSV ', -INFO )
        RETURN
    END IF
*
    IF( N.EQ.0 )
$      RETURN
*
```

```

DO 30 K = 1, N - 1
  IF( DL( K ).EQ.ZERO ) THEN
*
*      Subdiagonal is zero, no elimination is required.
*
*      IF( D( K ).EQ.ZERO ) THEN
*
*          Diagonal is zero: set INFO = K and return; a unique
*          solution can not be found.
*
*          INFO = K
*          RETURN
*          END IF
* ELSE IF( ABS( D( K ) ).GE.ABS( DL( K ) ) ) THEN
*
*      No row interchange required
*
*      MULT = DL( K ) / D( K )
*      D( K+1 ) = D( K+1 ) - MULT*DU( K )
*      DO 10 J = 1, NRHS
*          B( K+1, J ) = B( K+1, J ) - MULT*B( K, J )
10    CONTINUE
*      IF( K.LT.( N-1 ) )
*          DL( K ) = ZERO
*      ELSE
*
*          Interchange rows K and K+1
*
*          MULT = D( K ) / DL( K )
*          D( K ) = DL( K )
*          TEMP = D( K+1 )
*          D( K+1 ) = DU( K ) - MULT*TEMP
*          IF( K.LT.( N-1 ) ) THEN
*              DL( K ) = DU( K+1 )

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        DU( K+1 ) = -MULT*DL( K )
END IF
DU( K ) = TEMP
DO 20 J = 1, NRHS
    TEMP = B( K, J )
    B( K, J ) = B( K+1, J )
    B( K+1, J ) = TEMP - MULT*B( K+1, J )
20      CONTINUE
END IF
30 CONTINUE
IF( D( N ).EQ.ZERO ) THEN
    INFO = N
    RETURN
END IF
*
*     Back solve with the matrix U from the factorization.
*
DO 50 J = 1, NRHS
    B( N, J ) = B( N, J ) / D( N )
    IF( N.GT.1 )
        $     B( N-1, J ) = ( B( N-1, J )-DU( N-1 )*B( N, J ) ) / D( N-1 )
        DO 40 K = N - 2, 1, -1
            B( K, J ) = ( B( K, J )-DU( K )*B( K+1, J )-DL( K )*
                $                     B( K+2, J ) ) / D( K )
40      CONTINUE
50 CONTINUE
*
RETURN
*
*     End of DGTSV
*
END

```



```

.IGNORE:

F77_COMPILE = $(F77) $(F77FLAGS) $(F77CFLAGS)
F77_LOAD    = $(F77) $(F77FLAGS) $(F77LFLAGS)

.f.o:
    $(F77_COMPILE) $*.f

EXECUTABLES = bvp1d

all: $(EXECUTABLES)

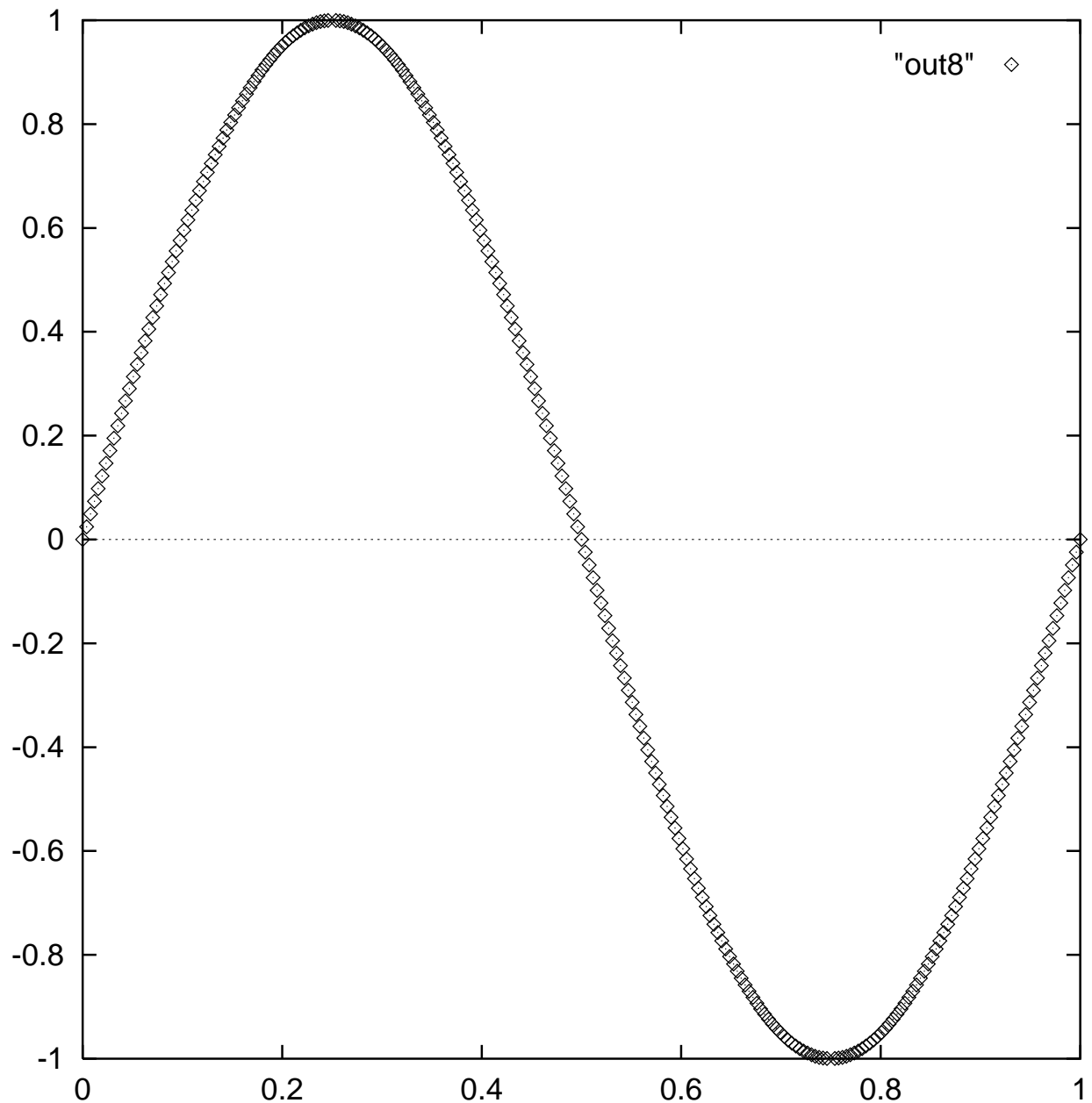
bvp1d: bvp1d.o
    $(F77_LOAD) bvp1d.o -lp410f -llapack $(LIBBLAS) -o bvp1d

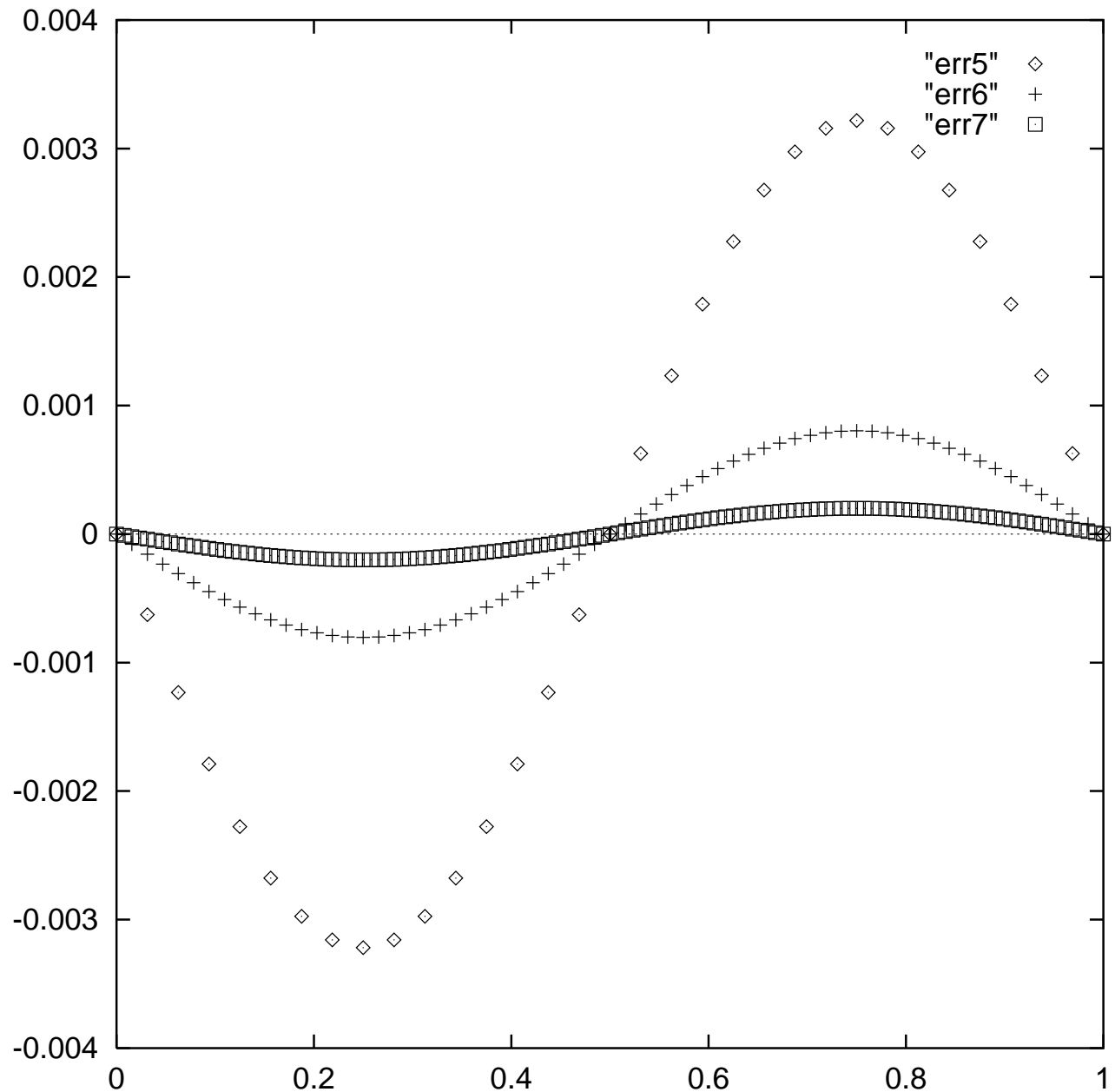
clean:
    rm *.o
    rm $(EXECUTABLES)

#####
# Note the 'vclean' target: 'make vclean' results in
# 'make clean' followed by removal of input and output
# data files and postscript files.
#####

vclean: clean
    rm err[0-9]*
    rm out[0-9]*
    rm *.ps

```





```

c=====
c      Solves 1-d linear boundary value problem
c
c      u''(x) = f(x)   on   x = [0,1] ; u(0) = u0, u(1) = u1
c
c      using mixed fourth-order and second order finite
c      difference technique and LAPACK banded solver DGSBV.
c=====

      program          bvp1d4

      implicit          none

      integer           i4arg

c-----
c      Domain extrema and maximum system size.
c-----
      real*8            xmin,           xmax
      parameter         ( xmin = 0.0d0,   xmax = 1.0d0 )

      integer           maxn
      parameter         ( maxn = 2**19 + 1 )

c-----
c      Storage for discrete x-values, unknowns, exact
c      solution and right hand side values.
c-----
      real*8            x(maxn),        u(maxn),
      &                  ueexact(maxn),  f(maxn)

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```

c-----
c      Number of lower and upper bands.
c-----
c      integer          kl,                 ku
c      parameter        ( kl = 2,           ku = 2  )
c-----
c      Storage for LAPACK-banded-form of linear system,
c      right-hand-side of system and pivot vector,
c      for use with DGBSV.
c
c      Note that for pivoting purposes (row interchanges)
c      DGBSV requires an additional 'kl' rows of workspace.
c      Leading dimension of 'ab' is thus
c
c      ku + kl + kl + 1 = 7
c-----
c      integer          ldab
c      parameter        ( ldab = 7  )
c      real*8          ab(ldab,maxn), rhs(maxn)
c      integer          ipiv(maxn)
c-----
c      Other standard LAPACK parameters.
c-----
c      integer          nrhs,             info
c-----
c      Discretization level, size of system (# of discrete
c      unknowns) and output option.
c-----
c      integer          level,            n,           option
c-----
c      Storage for difference coefficients. Note: these
c      arrays have elements -2, -1, 0, 1 and 2.
c-----
c      real*8          cdd2(-2:2),       cdd4(-2:2),       c0(-2:2)

```

```
c-----
c      Mesh spacing and related constants.
c-----
      real*8           h,           hm2,           hm2by12
c-----
c      Other locals.
c-----
      integer           i,           j,           k
      real*8           rmserr
c-----
c      Argument parsing.
c-----
      level = i4arg(1,-1)
      if( level .lt. 0 ) go to 900
      n = 2 ** level + 1
      if( n .gt. maxn ) then
          write(0,*) 'Insufficient internal storage'
          stop
      end if
      option = i4arg(2,0)
```

```

c-----
c      Set up finite-difference 'mesh' (discrete x-values)
c      and difference coefficient arrays.
c-----

      h      =  1.0d0 / (n - 1)
      do j = 1 , n
         x(j) = xmin + (j - 1) * h
      end do
      x(n) = xmax

      hm2      =  1.0d0 / (h * h)
      hm2by12 = hm2 / 12.0d0

      c0(-2)   =  0.0d0
      c0(-1)   =  0.0d0
      c0( 0)   =  1.0d0
      c0( 1)   =  0.0d0
      c0( 2)   =  0.0d0

      cdd2(-2) =  0.0d0
      cdd2(-1) =  hm2
      cdd2( 0) = -2.0d0 * hm2
      cdd2( 1) =  hm2
      cdd2( 2) =  0.0d0

      cdd4(-2) = -hm2by12
      cdd4(-1) =  16.0d0 * hm2by12
      cdd4( 0) = -30.0d0 * hm2by12
      cdd4( 1) =  16.0d0 * hm2by12
      cdd4( 2) = -hm2by12

```

```

c-----
c      Set up exact solution and right hand side vector.
c-----
call exact(uexact,f,x,n)

c=====
c      Set up banded system. Recall that for LAPACK
c      banded storage for LU decomposition
c
c      a( i , j ) -> ab( kl + ku + 1 + i - j , j )
c=====

c-----
c      i = 1: (Left boundary) u(1) = u_0
c-----
i = 1

do k = 0 , 2
    j = i + k
    ab(kl + ku + 1 + i - j,j) = c0(k)
end do
rhs(i) = uexact(i)
c-----
c      i = 2: O(h^2) approximation of u''(x) = f(x)
c-----
i = 2

do k = -1 , 2
    j = i + k
    ab(kl + ku + 1 + i - j,j) = cdd2(k)
end do
rhs(i) = f(i)

```

```

c-----
c      i = 3, . . . , n-2: O(h^4) approximation of u''(x) = f(x)
c-----
      do i = 3 , n - 2
        do k = -2 , 2
          j = i + k
          ab(kl + ku + 1 + i - j,j) = cdd4(k)
        end do
        rhs(i) = f(i)
      end do
c-----
c      i = n-1:   O(h^2) approximation of u''(x) = f(x)
c-----
      i = n - 1

      do k = -2 , 1
        j = i + k
        ab(kl + ku + 1 + i - j,j) = cdd2(k)
      end do
      rhs(i) = f(i)
c-----
c      i = n:   (Right boundary) u(n) = u_1
c-----
      i = n

      do k = -2 , 0
        j = i + k
        ab(kl + ku + 1 + i - j,j) = c0(k)
      end do
      rhs(i) = uexact(i)

```

```

c=====
c      Solve banded system.
c=====

      nrhs = 1
      call dgbsv( n, kl, ku, nrhs, ab, ldab, ipiv, rhs, n,
&                  info )

      if( info .eq. 0 ) then
c-----
c      Solver successful, output either (x_j, u_j) or
c      (x_j, error_j) to stdout. Also compute rms error
c      and output to standard error.
c-----

      rmserr = 0.0d0
      do j = 1 , n
          if( option .eq. 0 ) then
              write(*,*) x(j), rhs(j)
          else
              write(*,*) x(j), (uexact(j) - rhs(j))
          end if
          rmserr = rmserr + (uexact(j) - rhs(j)) ** 2
      end do
      rmserr = sqrt(rmserr / n)
      write(0,*) 'rmserr = ', rmserr
      else
c-----
c      Solver failed.
c-----

      write(0,*) 'bvp1d4: dgbsv() failed, info = ', info
      end if

      stop

```

```

900 continue
    write(0,*) 'usage: bvp1d4 <level> [<option>]'
    write(0,*)
    write(0,*) '           Specify option .ne. 0 for output'
    write(0,*) '           of error instead of solution'
    stop

end

c=====
c      Computes exact values for u(x) (unknown function)
c      and f(x) (right hand side function).  x array must
c      have been previously defined.
c=====

subroutine exact(u,f,x,n)

implicit none
integer n
real*8 u(n), f(n), x(n)

real*8 pi2
integer j

pi2 = 8.0d0 * atan(1.0d0)
do j = 1 , n
    u(j) = sin(pi2 * x(j))
    f(j) = -pi2 * pi2 * u(j)
end do

return

end

```