

Source file: bisect.f

```

c=====
c   bisect: Uses bisection to find approximate root
c   of f(x) on interval [xmin .. xmax].  Return value is      900
c   root located to (relative) tolerance 'xtol'.  Return code
c   'rc' is set to 0 on success, non-zero on failure
c   and routine succeeds (by definition) as long as initial
c   interval *does* bracket at least one root.  Routine
c   performs tracing of algorithm (on stderr) if input
c   argument 'trace' is .true.
c=====
real*8 function bisect(f,xmin,xmax,xtol,trace,rc)

    implicit none

    real*8 drelabs

    real*8 f
    external f

    real*8 xmin,      xmax,      xtol
    logical trace
    integer rc

c----- Other variables needed for search.
c-----
integer mxiter
parameter (mxiter = 50)

    real*8 xlo,      dx,      sgn
    integer iter

c----- Check that input interval is specified correctly
c   and that it manifestly brackets at least one root:
c   (i.e. the fcn changes sign).
c-----
& if( xmax .le. xmin .or.
&     f(xmin) * f(xmax) .gt. 0.0d0 ) then
&     write(0,*) 'bisect: Input interval is not //'
&                 'bracketing'
&     rc = 1

c----- Returned value is meaningless in this case,
c   but have to return *some* value.
c-----
    bisect = xmin
    return
end if

c----- Compute 'sgn' such that sgn * f(xmin) < 0, and
c   initialize bracketing interval.

c   Note that this could also be accomplished with
c   the 'sign' intrinsic (see tsign.f in the Misc. sec.
c   of the course Software page for instructive usage of
c   'sign')

c   sgn = sign(1.0d0,-f(xmin))

c----- if( f(xmin) .le. 0.0d0 ) then
c         sgn = 1.0d0
c     else
c         sgn = -1.0d0
c     end if

    xlo = xmin
    dx = xmax - xmin

c----- Bisection loop: continue until root found to
c   specified tolerance or until maximum number of
c   iterations taken
c-----
do iter = 1 , mxiter
    bisect = xlo + 0.5d0 * dx
    if( trace ) then
        write(0,*) xlo, xlo + dx, f(bisect)
    end if
    if( sgn * f(bisect) .lt. 0.0d0 ) then
        xlo = bisect
    end if

    if( drelabs(dx,bisect,1.0d-10) .le. xtol ) go to 900
    dx = 0.5d0 * dx
end do

    continue
    rc = 0
    if( trace ) write(0,*)
    return

end

c=====
c   drelabs: Function useful for 'relativizing' quantity
c   being monitored for detection of convergence.
c=====
real*8 function drelabs(dx,x,xfloor)
    implicit none

    real*8 dx,      x,      xfloor

    if( abs(x) .lt. abs(xfloor) ) then
        drelabs = abs(dx)
    else
        drelabs = abs(dx/x)
    end if

    return
end

```

Source file: tbisect.f

```

c=====
c   tbisect: Illustrates root finding using bisection
c   routine 'bisect'.
c
c   Initial bracketing interval must be specified via the
c   command-line, along with optional convergence criteria
c   and output option.
c
c   This program also illustrates the general Fortran
c   techniques (briefly discussed previously) for:
c
c   (1) Writing and using routines which take other routines
c       as arguments.
c   (2) Using a COMMON block to communicate information to
c       a routine in cases where the information cannot be
c       passed via the argument list.
c   (3) Using an "INCLUDE" file (in this case 'comf.inc')
c       to ensure that the same common block structure is defined
c       in all program units.
c
c   Currently set up for computing square roots i.e.
c   solves
c
c   f(x; a) = x**2 - a = 0
c
c   for 'a' specified on command-line
c
c   Outputs a, approximate root (x*) and f(x*; a) on stdout.
c=====
program tbisect
    implicit none

c----- Declaration of the bisection routine.
c-----
real*8 bisect
c----- Name of the specific function whose root we seek.
c   Note use of 'external' to let compiler know 'fsqr',
c   is the name of a function, not a variable.
c-----
real*8 fsqr
external fsqr

    integer i4arg,      iargc
    real*8 r8arg

c----- For use in detecting bad real*8 command-line value.
c-----
real*8 r8_never

```

```

parameter      ( r8_never = -1.0d-60 )
c-----
c   Use a common block to pass number whose square root
c   is sought to external function 'fsqr'.
c-----
include      'comf.inc'
c-----
c   Initial bracket, convergence tolerance and output
c   option from command-line; default value for conv.
c   tolerance.
c-----
real*8        xmin,           xmax,           xtol
logical       trace
c
real*8        default_xtol
parameter     ( default_xtol = 1.0d-8 )
c
c   Root and return code from bisection routine.
c-----
real*8        root
integer       rc
c
c   Argument parsing.
c-----
if( iargc() .lt. 3 ) go to 900
a    = r8arg(1,r8_never)
xmin = r8arg(2,r8_never)
xmax = r8arg(3,r8_never)
if( a .eq. r8_never .or. xmin .eq. r8_never .or.
&   xmax .eq. r8_never ) go to 900
c
xtol = r8arg(4,default_xtol)
trace = iargc() .gt. 4
c
c   Invoke root finder then write a, sqrt(a), and residual
c   to standard output.
c-----
root = bisect(fsqr,xmin,xmax,xtol,trace,rc)
if( rc .eq. 0 ) then
  write(*,*) a, root, fsqr(root)
else
  write(0,*) 'tbisect: Bisection failed.'
end if
c
c   Normal exit.
c-----
stop
c
c   Usage exit.
c-----
900 continue
  write(0,*) 'usage: tbisect <a> <xmin> <xmax> //'
  &           '[<xtol> <trace>]'
  stop
  end
c
c   Function whose root is sought. Again, note use of
c   COMMON block to pass additional information (in this
c   case 'a') to the routine.
c-----
real*8 function fsqr(x)
  implicit none
  real*8        x
  include      'comf.inc'
  fsqr = x**2 - a
  return
end

```

Source file: comf.inc

```

c-----
c   Common block for communicating value of 'a' from main
c   to 'fsqr'.
c-----
real*8          a
common / comf / a

```

Source file: Output on lnx1

```

#####
# Building 'tbisect' and sample output on lnx1
#
# 'tbisect' is set up to compute sqrt(a) via bisection.
#####
lnx1% pwd; ls
/home/phys410/nonlin/bisect
Makefile bisect.f comf.inc tbisect.f

lnx1% make
pgf77 -g -c tbisect.f
pgf77 -g -c bisect.f
pgf77 -g -L/usr/local/PGI/lib tbisect.o bisect.o -lp410f \
-o tbisect

#####
# Compute +sqrt(2) to default tolerance (1.0d-8)
#
# Note: Exact value to 16 digits is 1.414 2135 6237 3095
#####
lnx1% tbisect 2.0 1.0 2.0
  2.0000000000000000  1.414213564246893  5.2999007543741428E-009

#####
# Recompute with higher tolerance (1.0d-12)
#####
lnx1% tbisect 2.0 1.0 2.0 1.0e-12
  2.0000000000000000  1.414213562372879  -6.1080654423228964E-013

#####
# Enable tracing output by supplying 5th argument. Note
# supplying a '.' as an argument parsed by 'i4arg' or 'r8arg'
# is equivalent to specifying the default value.
#####
lnx1% tbisect 2.0 1.0 2.0 .1
  1.0000000000000000  2.0000000000000000  0.2500000000000000
  1.0000000000000000  1.5000000000000000  -0.4375000000000000
  1.2500000000000000  1.5000000000000000  -0.1093750000000000
  1.3750000000000000  1.5000000000000000  6.64062500000000E-002
  1.3750000000000000  1.4375000000000000  -2.24609375000000E-002
  1.4062500000000000  1.4375000000000000  2.17285156250000E-002
  1.4062500000000000  1.4218750000000000  -4.272460937500000E-004
  1.4140625000000000  1.4218750000000000  1.0635375976562500E-002
  1.4140625000000000  1.4179687500000000  5.1002502441406250E-003
  1.4140625000000000  1.4160156250000000  2.3355484008789063E-003
  1.4140625000000000  1.4150396250000000  9.5391273498535156E-004
  1.4140625000000000  1.414550781250000  2.6327371597290039E-004
  1.4140625000000000  1.414306640625000  -8.2001090049743652E-005
  1.414184570312500  1.414306640625000  9.0632587671279907E-005
  1.414184570312500  1.414245605468750  4.3148174881935120E-006
  1.414184570312500  1.414215087890625  -3.8843369111418724E-005
  1.414199829101563  1.414215087890625  -1.7264334019273520E-005
  1.414207458496094  1.414215087890625  -6.4747728174552321E-006
  1.414211273193359  1.414215087890625  -1.0799813026096672E-006
  1.414213180541992  1.414215087890625  1.6174171832972206E-006
  1.414213180541992  1.414214134216309  2.6871771297010127E-007
  1.414213180541992  1.414213657379150  -4.0563185166320181E-007
  1.414213418960571  1.414213657379150  -6.8457083557404985E-008
  1.414213538169861  1.414213657379150  1.0013031115363447E-007
  1.414213538169861  1.414213597774506  1.5836612909936321E-008
  1.414213538169861  1.414213567972183  -2.6310235545778937E-008
  1.414213553071022  1.414213567972183  -5.2368113734324595E-009
  1.414213560521603  1.414213567972183  5.2999007543741428E-009

  2.0000000000000000  1.414213564246893  5.2999007543741428E-009

```

Source file: newtsqrt.f

```

c=====
c      newtsqrt: Uses Newton's method to find (positive)
c      square root of number supplied on command line, i.e.
c      solves
c
c      f(x) = x^2 - a = 0
c
c      for given 'a'. Optional second argument specifies
c      convergence criteria (relative dx).
c
c      Tracing output (written to standard error)
c      includes iteration number, estimated root (xn),
c      change in estimate (dxn), log10(dxn), residual and
c      log10(residual).
c=====
program      newtsqrt
implicit      none
integer        iargc
real*8         r8arg,      drelabs
real*8         r8_never
parameter      ( r8_never = -1.0d-60 )
c
c      Default convergence tolerance.
c-----
real*8         default_xtol
parameter      ( default_xtol = 1.0d-8 )
c
c      Maximum allowed number of Newton iterations.
c-----
integer        mxiter
parameter      ( mxiter = 50 )
c
c      Command-line arguments (see above).
c-----
real*8         a,           xtol
c
c      Locals used in Newton iteration.
c-----
integer        iter
real*8         xn,          resn,          dxn
c
c      Argument parsing.
c-----
if( iargc() .lt. 1 ) go to 900
a     = r8arg(1,r8_never)
if( a .eq. r8_never .or. a .lt. 0.0d0 ) go to 900
xtol = r8arg(2,1.0d-8)
if( xtol .le. 0.0d0 ) xtol = 1.0d-8
c
c      Un-inspired initial guess: x^(0) = a / 2.
c-----
xn = 0.5d0 * a
c
c      Newton loop.
c-----
write(0,*)
&      'Iter      xn      '//'
&      'dxn      log10(dxn)   rn      log10(rn)'
do iter = 1 , mxiter
    resn = xn**2 - a
    dxn = resn / (2.0d0 * xn)
    xn  = xn - dxn
    write(0,1000) iter, xn, dxn, log10(abs(dxn)),
&                  resn, log10(abs(resn))
1000  format(i2,1p,e26.16,e12.3,0p,f10.2,1p,e12.3,0p,f10.2)
c
c      Jump out of Newton loop if soln has converged.
c-----
if( drelabs(dxn,xn,1.0d-10) .le. xtol ) go to 100
end do
c
c      No-convergence exit.
c-----

```

```

write(0,*)
&      'No convergence after ', mxiter,
&      ' iterations'
stop

```

```

c-----
c      Normal exit, write input and estimated square root
c      to standard output.
c-----

```

```

100  continue
write(0,*)
write(*,*) a, xn
stop

```

```

c-----
c      Usage exit.
c-----

```

```

900  continue
write(0,*)
'usage: newtsqrt <a> [<xtol>]'
stop
end

```

```

c=====
c      drelabs: Function useful for 'relativizing' quantity
c      being monitored for detection of convergence.
c=====

```

```

real*8 function drelabs(dx,x,xfloor)

```

```

implicit      none
real*8         dx,          x,           xfloor
if( abs(x) .lt. abs(xfloor) ) then
    drelabs = abs(dx)
else
    drelabs = abs(dx/x)
end if
return
end

```

Source file: Makefile

```

.IGNORE:
F77_COMPILE  = $(F77) $(F77FLAGS) $(F77CFLAGS)
F77_LOAD     = $(F77) $(F77FLAGS) $(F77LFLAGS)

```

```

.f.o:
    $(F77_COMPILE) $*.f
EXECUTABLES = newtsqrt

```

```

all: $(EXECUTABLES)

```

```

newtsqrt: newtsqrt.o
    $(F77_LOAD) newtsqrt.o -lp410f -o newtsqrt

```

```

clean:
    rm *.o
    rm $(EXECUTABLES)

```

Source file: Output on lnx1

```

#####
# Building 'newtsqrt' and sample output on lnx1
#####
/home/phys410/nonlin/newtsqrt
Makefile newtsqrt.f

lnx1% make
pgf77 -g -c newtsqrt.f
pgf77 -g -L/usr/local/PGI/lib newtsqrt.o -lp410f -o newtsqrt

#####
# Compute +sqrt(10) to default tolerance (1.0d-8)
#
# Note: Exact value to 16 digits is 3.162 2776 6016 8379
#####
% newtsqrt 10

   Iter      xn          dxn      log10(dxn)      rn      log10(rn)
1   3.500000000000000E+00   1.500E+00       0.18   1.500E+01     1.18
2   3.1785714285714284E+00   3.214E-01      -0.49   2.250E+00     0.35
3   3.1623194221508828E+00   1.625E-02      -1.79   1.033E-01    -0.99
4   3.1622776604441363E+00   4.176E-05      -4.38   2.641E-04    -3.58
5   3.1622776601683795E+00   2.758E-10      -9.56   1.744E-09    -8.76

10.00000000000000          3.162277660168380

#####
# Recompute with higher tolerance---an extra Newton step
# is taken, but the solution was already accurate to
# roughly machine epsilon, so there is very little change
# in the output.
#####
% newtsqrt 10 1.0e-12

   Iter      xn          dxn      log10(dxn)      rn      log10(rn)
1   3.500000000000000E+00   1.500E+00       0.18   1.500E+01     1.18
2   3.1785714285714284E+00   3.214E-01      -0.49   2.250E+00     0.35
3   3.1623194221508828E+00   1.625E-02      -1.79   1.033E-01    -0.99
4   3.1622776604441363E+00   4.176E-05      -4.38   2.641E-04    -3.58
5   3.1622776601683795E+00   2.758E-10      -9.56   1.744E-09    -8.76
6   3.1622776601683795E+00   1.908E-16     -15.72   1.207E-15   -14.92

10.00000000000000          3.162277660168380

#####
# Compute +sqrt(1/2) to default tolerance (1.0d-8)
#
# Note: Exact value to 16 digits is 0.7071 0678 1186 5475
#####
lnx1% newtsqrt 0.5

   Iter      xn          dxn      log10(dxn)      rn      log10(rn)
1   1.125000000000000E+00   -8.750E-01      -0.06   -4.375E-01    -0.36
2   7.847222222222223E-01   3.403E-01      -0.47   7.656E-01    -0.12
3   7.109451819075130E-01   7.378E-02      -1.13   1.158E-01    -0.94
4   7.0711714297003674E-01   3.828E-03      -2.42   5.443E-03    -2.26
5   7.0710678126246602E-01   1.036E-05      -4.98   1.465E-05    -4.83
6   7.0710678118654755E-01   7.592E-11     -10.12   1.074E-10   -9.97

0.5000000000000000          0.7071067811865476

```

Source file: newt2.f

```

c=====
c   newt2: Uses multi-dimensional Newton's method
c   to compute a root of simple non-linear system
c   discussed in class
c
c   sin(xy) - 1/2 = 0
c   y^2 - 6x - 2 = 0
c
c   Command line input is initial guess (two numbers)
c   for root, and optional convergence criteria.
c   Estimated root written to standard output.
c   Tracing output similar to that from 'newtsqrt'.
c=====
program      newt2
implicit      none
integer        iargc
real*8         r8arg,          drelabs,      dvl2norm
real*8         r8_never
parameter      ( r8_never = -1.0d-60 )
c
c   Size of system.
c
integer        neq
parameter      ( neq = 2 )
c
c   Command-line arguments: Initial guess will be
c   input directly into 'x' array.
c
real*8         tol
c
c   Variables used in Newton iteration and solution of
c   linear systems via LAPACK routine 'dgesv'.
c
real*8         J(neq,neq),    res(neq),
&             x(neq)
integer        ipiv(neq)
integer        ieq,           info
integer        mxiter,        nrhs
parameter      ( mxiter = 50,  nrhs = 1 )
integer        iter
real*8         nrm2res,       nrm2dx,       nrm2x
c
c   Default convergence tolerance.
c
real*8         default_tol
parameter      ( default_tol = 1.0d-8 )
c
c   Argument parsing.
c
if( iargc() .lt. neq ) go to 900
do ieq = 1 , neq
  x(ieq) = r8arg(ieq,r8_never)
  if( x(ieq) .eq. r8_never ) go to 900
end do
tol = r8arg(neq+1,default_tol)
if( tol .le. 0.0d0 ) tol = default_tol
c
c   Newton loop.
c
write(0,*)
  'Iter          x          y  //'
  &           log10(dx) log10(res),
write(0,*)
  do iter = 1 , mxiter
c
c   Evaluate residual vector.
c
  res(1) = sin(x(1)*x(2)) - 0.5d0
  res(2) = x(2)**2 - 6.0d0 * x(1) - 2.0d0
  nrm2res = dvl2norm(res,2)
c
c   Set up Jacobian.
c
  J(1,1) = x(2) * cos(x(1) * x(2))
  J(1,2) = x(1) * cos(x(1) * x(2))
  J(2,1) = -6.0d0
  J(2,2) = 2.0d0 * x(2)
c
c   Solve linear system (J dx = res) for update
c   dx. Update returned in 'res' vector.
c
call dgesv( neq, nrhs, J, neq, ipiv, res, neq, info )
if( info .eq. 0 ) then
c
c   Update solution.
c
  nrm2x = dvl2norm(x,neq)
  nrm2dx = dvl2norm(res,neq)
  do ieq = 1 , neq
    x(ieq) = x(ieq) - res(ieq)
  end do
c
c   Tracing output: note use of max to prevent
c   taking log10 of 0.
c
  write(0,1000)
    iter, x(1), x(2),
    log10(max(nrm2dx,1.0d-60)),
    log10(max(nrm2res,1.0d-60))
1000  format(i2,1p,2e24.16,0p,2f8.2)
c
c   Check for convergence.
c
  if( drelabs(nrm2dx,nrm2x,1.0d-6) .le. tol ) go to 100
else
  write(0,*) 'newt2: dgesv failed.'
  stop
end if
end do
c
c   No-convergence exit.
c
  write(0,*) 'No convergence after ', mxiter,
  &           ' iterations'
  stop
c
c   Normal exit, write input and estimated square root
c   to standard output.
c
100  continue
  write(0,*) x
  stop
c
c   Usage exit.
c
900  continue
  write(0,*) 'usage: newt2 <x0> <y0> [<tol>]'
  stop
end
c=====
c   dvl2norm: Returns l2-norm of double precision vector.
c=====
real*8 function dvl2norm(v,n)
implicit      none
integer        n
real*8         v(n)
integer        i
dvl2norm = 0.0d0
do i = 1 , n
  dvl2norm = dvl2norm + v(i) * v(i)
end do
if( n .gt. 0 ) then
  dvl2norm = sqrt(dvl2norm / n)
end if
return
end

```

```

=====
c     drelabs: Function useful for 'relativizing' quantity
c     being monitored for detection of convergence.
=====
      real*8 function drelabs(dx,x,xfloor)

      implicit none

      real*8      dx,      x,      xfloor

      if( abs(x) .lt. abs(xfloor) ) then
        drelabs = abs(dx)
      else
        drelabs = abs(dx/x)
      end if

      return

    end

Source file: Makefile

.IGNORE:

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

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

EXECUTABLES = newt2

all: $(EXECUTABLES)

newt2: newt2.o
$(F77_LOAD) newt2.o -lp410f -llapack $(LIBBLAS) -o newt2

clean:
rm *.o
rm $(EXECUTABLES)

Source file: Output on lnx1

#####
# Building 'newt2' and sample output on lnx1.
#
# Note how different roots are found depending on the initial
# guess and how, in each case, convergence of both dx and
# the residual is quadratic as the solution is approached.
#####
lnx1% pwd; ls
/home/phys410/nonlin/newt2
Makefile newt2.f

lnx1% make
pgf77 -g -c newt2.f
pgf77 -g -L/usr/local/PGI/lib newt2.o \
-lp410f -llapack -lblas -o newt2

lnx1% newt2
usage: newt2 <x0> <y0> [<tol>]

#####
# Start with initial guess (1.0,1.0) and use default tolerance
#####
lnx1% newt2 1.0 1.0
      Iter          x                  y      log10(dx) log10(res)
1   -3.2999966453609808E-02  1.4010001006391706E+00   -0.11   0.70
2    3.7660093320946681E-01  2.2207017966697333E+00   -0.19   -0.40
3    2.6508349149835868E-01  1.9187667230922997E+00   -0.64   -0.30
4    2.7416951525985471E-01  1.9092166705387069E+00   -2.03   -1.19
5    2.7423631305849172E-01  1.9092977465351673E+00   -4.13   -3.95
6    2.7423631371214592E-01  1.9092977458408303E+00   -9.17   -8.33
                                         0.2742363137121459           1.909297745840830

#####
# Start with initial guess (10.0,10.0)
#####
lnx1% newt2 10.0 10.0
      Iter          x                  y      log10(dx) log10(res)
1   1.1551311217431483E+01  8.5653933652294452E+00   0.17   1.43
2    5.2821340061726980E+00  6.2494950887340224E+00   0.67   0.26
3    7.9156169058357619E+00  7.0845635560826592E+00   0.29   0.58
4    8.0553488925966921E+00  7.0945184795080038E+00   -1.00   -0.08
5    8.047880096985382E+00  7.0913532277563132E+00   -2.24   -1.34
6    8.0480621354266226E+00  7.0914295327798467E+00   -3.86   -2.93
7    8.0480622340064549E+00  7.0914295740731097E+00   -7.12   -6.20
                                         8.048062234006455           7.091429574073110

#####
# Start with initial guess (100.0,100.0)
#####
lnx1% newt2 100.0 100.0
      Iter          x                  y      log10(dx) log10(res)
1   1.4561314470371519E+02  5.4378394341111459E+01   1.66   3.82
2    1.9021837653952545E+02  3.7701738714769562E+01   1.53   3.17
3    2.0349983567820647E+02  3.5070267397907138E+01   0.98   2.29
4    2.0392234856561166E+02  3.5007684984188501E+01   -0.52   0.70
5    2.0390326095147370E+02  3.5005993323580434E+01   -1.87   -0.53
6    2.0391023928640129E+02  3.5006591323292412E+01   -2.31   -0.59
7    2.0391061250942664E+02  3.5006623302706338E+01   -3.58   -1.92
8    2.0391061457091234E+02  3.5006623479357074E+01   -5.83   -4.18
9    2.0391061457097669E+02  3.5006623479362588E+01   -10.34   -8.68
                                         203.9106145709767           35.00662347936259

#####
# Start with initial guess (0.0,0.0), generates singular
# Jacobian
#####
lnx1% newt2 0.0 0.0
      Iter          x                  y      log10(dx) log10(res)
newt2: dgesv failed.

#####
# Start with initial guess (1.0,1.0) but use more stringent
# tolerance
#####
lnx1% newt2 1.0 1.0 1.0e-15
      Iter          x                  y      log10(dx) log10(res)
1   -3.2999966453609808E-02  1.4010001006391706E+00   -0.11   0.70
2    3.7660093320946681E-01  2.2207017966697333E+00   -0.19   -0.40
3    2.6508349149835868E-01  1.9187667230922997E+00   -0.64   -0.30
4    2.7416951525985471E-01  1.9092166705387069E+00   -2.03   -1.19
5    2.7423631305849172E-01  1.9092977465351673E+00   -4.13   -3.95
6    2.7423631371214592E-01  1.9092977458408303E+00   -9.17   -8.33
7    2.7423631371214592E-01  1.9092977458408303E+00   -16.28  -16.07
                                         0.2742363137121459           1.909297745840830

```

Source file: Maple verification of computations

```
#####
# Checking 'newt2' using numerical root finding capablities
# of Maple.
#####
lnx1% maple
    \^/|      Maple 6 (IBM INTEL LINUX)
._\|_ \|/_|. Copyright (c) 2000 by Waterloo Maple Inc.
\ MAPLE / All rights reserved. Maple is a registered trademark of
<---- ----> Waterloo Maple Inc.
    |      Type ? for help.
> Digits := 20;
Digits := 20

> f1 := sin(x*y) - 1/2;
f1 := sin(x y) - 1/2

> f2 := y^2 - 6*x - 2;
          2
f2 := y - 6 x - 2

#####
# Locates root found by 'newt2 1.0 1.0'
#####
> ans := fsolve( {f1,f2}, {x,y}, {x=0.25..0.30, y=1.8..2.0});
ans := {x = .27423631371214588082, y = 1.9092977458408301606}

#####
# Compute residuals of root
#####
> r1 := evalf(subs(ans,f1)); r2 := evalf(subs(ans,f2));
           -19
r1 := -.1 10
           -18
r2 := -.1 10

#####
# Locates root found by 'newt2 10.0 10.0'
#####
> ans := fsolve( {f1,f2}, {x,y}, {x=7..9, y=6..8});
ans := {x = 8.0480622340064835835, y = 7.0914295740731220704}

> r1 := evalf(subs(ans,f1)); r2 := evalf(subs(ans,f2));
           -18
r1 := -.35 10
r2 := 0

#####
# Locates root found by 'newt2 100.0 100.0'
#####
> ans := fsolve( {f1,f2}, {x,y}, {x=203.9..203.95, y=35.0..35.01});
ans := {x = 203.91061457097670060, y = 35.006623479362590528}

> r1 := evalf(subs(ans,f1)); r2 := evalf(subs(ans,f2));
           -16
r1 := -.5214 10
r2 := 0

#####
# Another nearby, but distinct, root
#####
> ans := fsolve( {f1,f2}, {x,y}, {x=203..204, y=35.0..35.1});
ans := {x = 203.95052002180667001, y = 35.010043132376172782}

> r1 := evalf(subs(ans,f1)); r2 := evalf(subs(ans,f2));
           -16
r1 := .4548 10
r2 := 0

> quit;
```

Source file: nlbvp1d.f

```

c=====
c      Solves 1-d non-linear boundary value problem
c
c      u''(x) + (u u')^2 + sin(u) = f(x)
c
c      on  x = [0,1]; u(0) = 0, u(1) = 0
c
c      using second-order finite difference technique,
c      Newton's method and LAPACK tridiagonal solver DGTSV.
c
c usage: nlbvp1d <level> <guess_factor> [<option> <tol>]
c
c      level:      Discretization level;
c                  FD mesh has 2**level + 1 pts.
c      guess_factor: Controls initial estimate of soln;
c                  u^(0) = guess_factor * u_exact
c      option:       Output option, zero for solution,
c                  non-zero for error.
c      tol:         Convergence criterion for Newton
c                  iteration.
c
c      Currently set up for solution
c
c      u(x) = sin(4 Pi x)
c=====
program nlbvp1d
implicit none
integer i4arg
real*8 r8arg, drelabs, dvl2norm
real*8 r8_never
parameter (r8_never = -1.0d-60)
c
c Extrema of problem domain.
c
real*8 xmin, xmax
parameter (xmin = 0.0d0, xmax = 1.0d0)
c
integer maxn
parameter (maxn = 32769)
c
c Storage for discrete x-values, unknowns, coefficient
c exact solution and right hand side values.
c
real*8 x(maxn), u(maxn),
& uexact(maxn), f(maxn)
c
c Storage for main, upper and lower diagonals of
c tridiagonal system (Jacobian matrix) and
c right-hand-side vector (residual vector) for use with
c LAPACK routine DGTSV. Other parameters needed for
c call to DGTSV.
c
real*8 d(maxn), du(maxn),
& dl(maxn), rhs(maxn)
integer nrhs
parameter (nrhs = 1)
integer info
c
c Discretization level and size of system (# of discrete
c unknowns)
c
integer level, n, i,
& option
c
c Variables used in Newton iteration.
c
integer mxiter
parameter (mxiter = 50)
integer iter
real*8 guess_factor, tol,
& nrm2res, nrm2du, nrm2u
c
c      Enable following parameter for full tracing of
c      Newton iteration.
c
c----- logical newton_trace
c----- parameter (newton_trace = .false.)
c
c----- Mesh spacing and related constants
c
real*8 h, hm1by2, hhm2, m2hm2,
& 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
guess_factor = r8arg(2,r8_never)
if( guess_factor .eq. r8_never ) go to 900
option = i4arg(3,0)
tol = r8arg(4,1.0d-8)
c
c----- Set up finite-difference 'mesh' (discrete x-values)
c and define some useful constants.
c
h = 1.0d0 / (n - 1)
do i = 1 , n
  x(i) = xmin + (i - 1) * h
end do
hm2 = 1.0d0 / (h * h)
m2hm2 = -2.0d0 / (h * h)
hm1by2 = 0.50d0 / h
hhm2 = 0.50d0 * hm2
qhm2 = 0.25d0 * hm2
c
c----- This only ensures that x(n) = xmax EXACTLY and is not
c essential.
c
x(n) = xmax
c
c----- Set up exact solution, coefficient functions and right
c hand side vector.
c
call exact(uexact,f,x,n)
c
c----- Initialize unknown (u) to constant (guess_factor)
c times exact solution.
c
do i = 1 , n
  u(i) = guess_factor * uexact(i)
end do
c
c----- N E W T O N   L O O P
c
do iter = 1 , mxiter
c
c----- Set up tridiagonal Jacobian matrix and evaluate
c right-hand-side (residuals)
c
c----- Left boundary: Dirichlet boundary condition has
c 0 residual.
c
d(1) = 1.0d0
du(1) = 0.0d0
rhs(1) = 0.0d0
c
c----- Interior: J[i,j] = d(F_i)/d_(u_j) and has non-zero
c elements only for j = i-1, i and i+1.
c

```

```

do i = 2 , n - 1
  dl(i-1) = hm2 - hhm2 * u(i)**2 * (u(i+1) - u(i-1))
  &           + hhm2 * u(i) * (u(i+1) - u(i-1))**2
  &           + cos(u(i))
  du(i)   = hm2 + hhm2 * u(i)**2 * (u(i+1) - u(i-1))
  rhs(i)  = hm2 * (u(i+1) - 2.0d0 * u(i) + u(i-1))
  &           + qhm2 * u(i)**2 * (u(i+1) - u(i-1))**2
  &           + sin(u(i)) - f(i)
end do
c-----
c Right boundary: Dirichlet boundary condition has
c 0 residual.
c-----
dl(n-1) = 0.0d0
d(n)   = 1.0d0
rhs(n) = 0.0d0
c-----
c Compute l-2 norm of residuals.
c-----
nrm2res = dvl2norm(rhs,n)
if( newton_trace ) then
  write(0,*) 'iter = ', iter
  write(0,*) 'res = ', nrm2res
end if
c-----
c Solve tridiagonal system for Newton update, delu,
c which satisfies
c
c J delu = residuals
c-----

call dgtsv( n, nrhs, dl, d, du, rhs, n, info )
if( info .eq. 0 ) then
c----- 
c Solver successful: compute norms of u and delu,
c update solution and check for convergence.
c----- 
nrm2u = dvl2norm(u,n)
nrm2du = dvl2norm(rhs,n)
if( newton_trace ) then
  write(0,*) 'du = ', nrm2du
  write(0,*) 'u = ', nrm2u
end if
do i = 1 , n
  u(i) = u(i) - rhs(i)
end do
if( drelabs(nrm2du,nrm2u,1.0d-6) .le. tol )
& go to 500
else
c----- 
c Solver failed, write error message and exit.
c----- 
write(0,*) 'nlbvp1d: dgtsv() failed, info = ',
&           info
end if
end do
c----- 
c Newton iteration failed to converge: write error
c message and exit.
c----- 
write(0,*) 'nlbvp1d: No convergence after ', mxiter,
&           ' iterations'
stop
c----- 
c Newton iteration converged: output solution or error
c to stdout, depending on output option. Also compute
c rms error and output to stderr.
c----- 
500 continue

rmserr = 0.0d0
do i = 1 , n
  if( option .eq. 0 ) then
    write(*,*) x(i), u(i)
  else
    write(*,*) x(i), (uexact(i) - u(i))
  end if
  rmserr = rmserr + (uexact(i) - u(i)) ** 2
end do
c-----
c----- 
rmserr = sqrt(rmserr / n)
write(0,*) 'rmserr = ', rmserr
stop
900 continue
write(0,*) 'usage: nlbvp1d <level> <guess_factor> //'
&           '[<option> <tol>]'
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 pi4
integer i

pi4 = 16.0d0 * atan(1.0d0)
do i = 1 , n
  u(i) = sin(pi4 * x(i))
  f(i) = -pi4**2 * sin(pi4 * x(i)) +
&           pi4**2 * (sin(pi4 * x(i)) *
&           cos(pi4 * x(i)))**2 +
&           sin(sin(pi4 * x(i)))
end do

return
end

c=====
c dvl2norm: Returns l2-norm of double precision vector.
c=====

real*8 function dvl2norm(v,n)

implicit none
integer n
real*8 v(n)
integer i

dvl2norm = 0.0d0
do i = 1 , n
  dvl2norm = dvl2norm + v(i) * v(i)
end do
if( n .gt. 0 ) then
  dvl2norm = sqrt(dvl2norm / n)
end if

return
end

c=====
c drelabs: Function useful for 'relativizing' quantity
c being monitored for detection of convergence.
c=====

real*8 function drelabs(dx,x,xfloor)

implicit none
real*8 dx, x, xfloor

if( abs(x) .lt. abs(xfloor) ) then
  drelabs = abs(dx)
else
  drelabs = abs(dx/x)
end if

```

```

        return
    end

    quit
END

# Generate 3 distinct solutions and plot the results

nlbvp1d 6 0.7 > out6-0.7
nlbvp1d 6 1.0 > out6-1.0
nlbvp1d 6 1.1 > out6-1.1

gnuplot<<END
set terminal postscript portrait
set output "allsolns.ps"
set size square
set title "Three Distinct Solutions \
of Nonlinear BVP"
set xlabel "x"
set ylabel "u(x)"
plot \
    "out6-0.7" title "guess_factor = 0.7" with lines, \
    "out6-1.0" title "guess_factor = 1.0" with lines, \
    "out6-1.1" title "guess_factor = 1.1" with lines
quit
END

gnuplot<<END
set terminal postscript portrait
set output "allsolnsz.ps"
set size square
set title "Three Distinct Solutions \
of Nonlinear BVP - Detail"
set xlabel "x"
set ylabel "u(x)"
plot [0.025:0.25] [0.5:1.1] \
    "out6-0.7" title "guess_factor = 0.7" with lines, \
    "out6-1.0" title "guess_factor = 1.0" with lines, \
    "out6-1.1" title "guess_factor = 1.1" with lines
quit
END

ls -lt *.ps

Source file: Nlbvp1d

#!/bin/sh -x
P='basename $0'

#-----
# Nlbvp1d: script which runs 'nlbvp1d' and plots results.
#-----

# Test for executable and make it if it doesn't exist.
test -f nlbvp1d || make

# Generate level-6 solution with guess_factor = 1.0

nlbvp1d 6 1.0 > out6

gnuplot<<END
set terminal postscript portrait
set output "soln6.ps"
set size square
set title "Solution of Nonlinear BVP\nguess_factor = 1.0"
set xlabel "x"
set ylabel "u(x)"
plot [0:1] [-1:1] \
    sin(12.56637061435917*x) title "exact solution", \
    "out6" notitle
quit
END

# Perform convergence test for guess_factor = 1.0

nlbvp1d 5 1.0 1 > err5
nlbvp1d 6 1.0 1 | nf _1 '4 * _2' > err6
nlbvp1d 7 1.0 1 | nf _1 '16 * _2' > err7

gnuplot<<END
set terminal postscript portrait
set output "err567.ps"
set size square
set title "Convergence Test of Solution \
of Nonlinear BVP\nguess_factor = 1.0"
set xlabel "x"
set ylabel ""
set key 0.725,0.014
plot \
    "err5" title "Level-5 error", \
    "err6" title "4 * Level-6 error", \
    "err7" title "16 * Level-7 error"

```

Figure file: soln6.ps

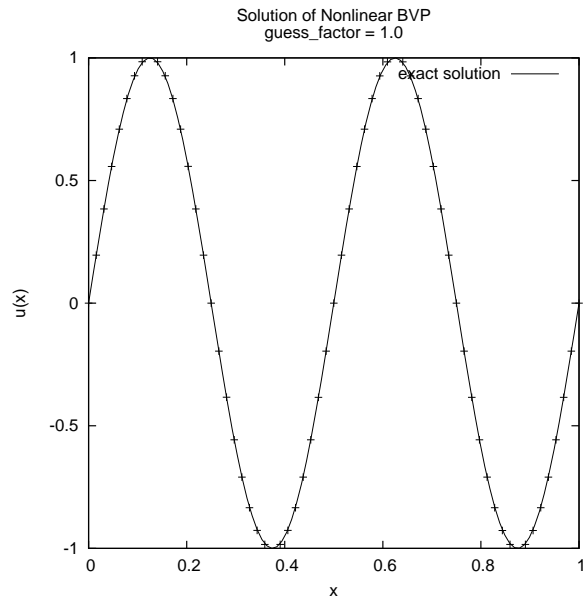


Figure file: allsolns.ps

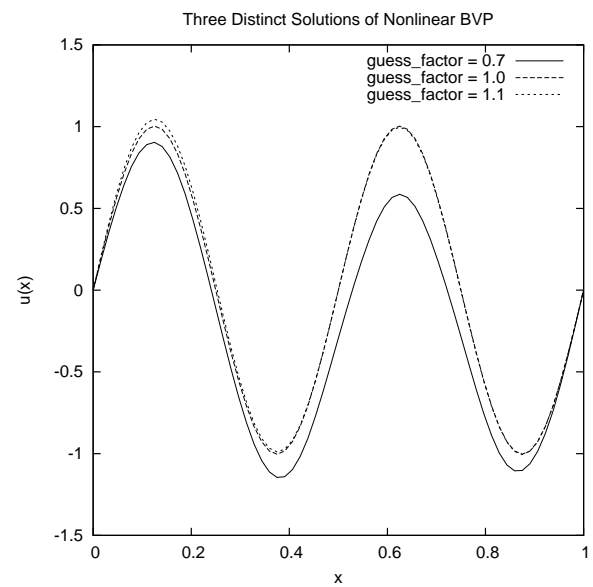


Figure file: err567.ps

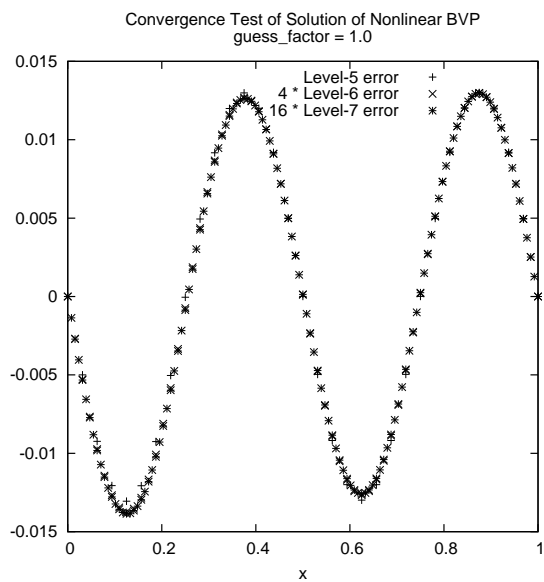


Figure file: allsolnsz.ps

