

```

c
  IMPLICIT DOUBLE PRECISION (a-h,o-z)
  parameter (N=10,N1=N+1)
  DOUBLE PRECISION A(N1,N1),B(N1),Y(N1)
  INTEGER IPVT(N1)
  COMMON /CHEB/ z1,z2,PI,Alpha,Beta
  COMMON /TRANS/ C(N1),CBAR(N1),CT(N1,N1),CTINV(N1,N1),Z(N1)
  OPEN(11,FILE='chebyshive_tau.txt')
c
  PI    = 4.d0*ATAN(1.d0)
  z1    = 0.d0
  z2    = 1.d0
c
  call Chebyshev
c
  call MATRIX(A)
c
  call dgefa(A,N1,N1,ipvt,info)
c
  do i = 1, N1-2
    B(i) = 0.d0
  enddo
  B(N1-1) = 0.d0
  B(N1)    = 1.d0
c
  call dgesl(A,N1,N1,ipvt,B,job)
  call SPRE(B,Y)
c
  do i = 1, N1
    ye = 1.d0/(dexp(-1.d0)-dexp(-4.d0))*
!      (dexp(-Z(i))-dexp(-4.d0*Z(i)))
    write(11,'(3f16.6)') Z(i),Y(i),ye
    write(* , '(3f16.6)') Z(i),Y(i),ye
  enddo
c
1000 stop
end
c
c
c
  SUBROUTINE MATRIX(A)
  IMPLICIT DOUBLE PRECISION (a-h,o-z)
  parameter (N=10,N1=N+1)
  DOUBLE PRECISION A(N1,N1)
  COMMON /CHEB/ z1,z2,PI,Alpha,Beta
  COMMON /TRANS/ C(N1),CBAR(N1),CT(N1,N1),CTINV(N1,N1),Z(N1)
c
  do i = 1, N1
    do j = 1, N1
      A(i,j) = 0.d0
    enddo
  enddo
c
  Construct the residuals in the spectral domain
c
  do i = 1, N1-2
    ri = dble(i) - 1.d0
c
    A(i,i) = A(i,i) + 4.d0
c
    1st-order derivatives
    do ip = i + 1, N1, 2
      rp = dble(ip) - 1.d0
      A(i,ip) = A(i,ip) + 5.d0*(2.d0/C(i)*rp)/Beta
    enddo
c
    2nd-order derivatives
    do ip = i + 2, N1, 2
      rp = dble(ip) - 1.d0
      A(i,ip) = A(i,ip) + 1.d0/C(i)*rp*(rp*rp-ri*ri)/Beta/Beta
    enddo
  enddo
c
  Implement boundary conditions
c

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c
do j = 1, N1
  A(N1-1,j) = 1.d0
  A(N1 ,j) = (-1.d0)**(j-1)
enddo

c
return
end

c
c
c
SUBROUTINE Chebyshev
IMPLICIT DOUBLE PRECISION (a-h,o-z)
parameter (N=10,N1=N+1)
DOUBLE PRECISION XCHEB(N1)
COMMON /CHEB/ z1,z2,PI,Alpha,Beta
COMMON /TRANS/ C(N1),CBAR(N1),CT(N1,N1),CTINV(N1,N1),Z(N1)

c
c Vector XCHEB (1,-1) contains the Gauss-Lobatto grid.
do i = 1, N1
  XCHEB(i) = dcos((i-1)*pi/N)
  CBAR(i) = 1.d0
  C(i) = 1.d0
enddo

c
C(1) = 2.d0
CBAR(1) = 2.d0
CBAR(N1) = 2.d0

c
c Vector X maps XCHEB to the interval (z1,z2) ==> X(i) = Alpha + Beta*XCHEB(i)
Alpha = (z1+z2)/2.d0
Beta = (z1-z2)/2.d0

c
do i = 1, N1
  Z(i) = Alpha + Beta*XCHEB(i)
enddo

c
do i = 1, N1
do j = 1, N1
  CT(i,j) = 2.d0/N/CBAR(i)/CBAR(j)*dcos(pi*(i-1)*(j-1)/N)
  CTINV(i,j) = dcos(pi*(i-1)*(j-1)/N)
enddo
enddo

c
return
end
c
c
c
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
subroutine RESP(AR,AS)
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
implicit double precision (a-h,o-z)
parameter (N=10,N1=N+1)
DOUBLE PRECISION AR(N1),AS(N1)
COMMON /TRANS/ C(N1),CBAR(N1),CT(N1,N1),CTINV(N1,N1),Z(N1)

c
c invert chebychev
c
do i = 1, N1
  AS(i) = 0.d0
  do j = 1, N1
    AS(i) = AS(i) + CT(i,j) * AR(j)
  enddo
enddo

c
return
end
c
c
c
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
subroutine SPRE(AS,AR)
XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
implicit double precision (a-h,o-z)
parameter (N=10,N1=N+1)
DOUBLE PRECISION AR(N1),AS(N1)
COMMON /TRANS/ C(N1),CBAR(N1),CT(N1,N1),CTINV(N1,N1),Z(N1)

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c
c  invert chebychev
c
do i = 1, N1
  AR(i) = 0.d0
  do j = 1, N1
    AR(i) = AR(i) + CTINV(i,j) * AS(j)
  enddo
enddo
c
return
end

c
c
c
c
c
c
c
c
c
c
c  subroutine dgefa to LU decompose matrix A
c  subroutine dgefa(a,lda,n,ipvt,info)
c  integer lda,n,ipvt(1),info
c  double precision a(lda,1)
c
c  dgefa factors a double precision matrix by gaussian elimination.
c
c  dgefa is usually called by dgeco, but it can be called
c  directly with a saving in time if rcond is not needed.
c  (time for dgeco) = (1 + 9/n)*(time for dgefa) .
c
c  on entry
c
c    a      double precision(lda, n)
c           the matrix to be factored.
c
c    lda    integer
c           the leading dimension of the array a .
c
c    n      integer
c           the order of the matrix a .
c
c  on return
c
c    a      an upper triangular matrix and the multipliers
c           which were used to obtain it.
c           the factorization can be written a = l*u where
c           l is a product of permutation and unit lower
c           triangular matrices and u is upper triangular.
c
c    ipvt   integer(n)
c           an integer vector of pivot indices.
c
c    info   integer
c           = 0 normal value.
c           = k if u(k,k) .eq. 0.0 . this is not an error
c           condition for this subroutine, but it does
c           indicate that dgesl or dgedi will divide by zero
c           if called. use rcond in dgeco for a reliable
c           indication of singularity.
c
c  linpack. this version dated 08/14/78 .
c  cleve moler, university of new mexico, argonne national lab.
c
c  subroutines and functions
c
c  blas daxpy,dscal,idamax
c
c  internal variables
c
c  double precision t
c  integer idamax,j,k,kpl,l,nml
c

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c
c gaussian elimination with partial pivoting
c
info = 0
nml = n - 1
if (nml .lt. 1) go to 70
do 60 k = 1, nml
    kpl = k + 1
c
c find l = pivot index
c
l = idamax(n-k+1,a(k,k),1) + k - 1
ipvt(k) = l
c
c zero pivot implies this column already triangularized
c
if (a(l,k) .eq. 0.0d0) go to 40
c
c interchange if necessary
c
if (l .eq. k) go to 10
    t = a(l,k)
    a(l,k) = a(k,k)
    a(k,k) = t
10 continue
c
c compute multipliers
c
t = -1.0d0/a(k,k)
call dscal(n-k,t,a(k+1,k),1)
c
c row elimination with column indexing
c
do 30 j = kpl, n
    t = a(l,j)
    if (l .eq. k) go to 20
        a(l,j) = a(k,j)
        a(k,j) = t
20 continue
    call daxpy(n-k,t,a(k+1,k),1,a(k+1,j),1)
30 continue
go to 50
40 continue
    info = k
50 continue
60 continue
70 continue
ipvt(n) = n
if (a(n,n) .eq. 0.0d0) info = n
return
end
subroutine dgesl(a,lda,n,ipvt,b,job)
integer lda,n,ipvt(1),job
double precision a(lda,1),b(1)
c
c dgesl solves the double precision system
c a * x = b or trans(a) * x = b
c using the factors computed by dgeco or dgefa.
c
c on entry
c
c a double precision(lda, n)
c the output from dgeco or dgefa.
c
c lda integer
c the leading dimension of the array a .
c
c n integer
c the order of the matrix a .
c
c ipvt integer(n)
c the pivot vector from dgeco or dgefa.

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c
c      b      double precision(n)
c             the right hand side vector.
c
c      job      integer
c              = 0          to solve  a*x = b ,
c              = nonzero    to solve  trans(a)*x = b  where
c                             trans(a)  is the transpose.
c
c on return
c
c      b      the solution vector  x .
c
c error condition
c
c      a division by zero will occur if the input factor contains a
c      zero on the diagonal.  technically this indicates singularity
c      but it is often caused by improper arguments or improper
c      setting of lda .  it will not occur if the subroutines are
c      called correctly and if dgeco has set rcond .gt. 0.0
c      or dgefa has set info .eq. 0 .
c
c to compute inverse(a) * c  where  c  is a matrix
c with  p  columns
c      call dgeco(a,lda,n,ipvt,rcond,z)
c      if (rcond is too small) go to ...
c      do 10 j = 1, p
c          call dgesl(a,lda,n,ipvt,c(1,j),0)
c      10 continue
c
c linpack. this version dated 08/14/78 .
c cleve moler, university of new mexico, argonne national lab.
c
c subroutines and functions
c
c blas daxpy,ddot
c
c internal variables
c
c double precision ddot,t
c integer k,kb,l,nml
c
c nml = n - 1
c if (job .ne. 0) go to 50
c
c      job = 0 , solve  a * x = b
c      first solve  l*y = b
c
c      if (nml .lt. 1) go to 30
c      do 20 k = 1, nml
c          l = ipvt(k)
c          t = b(l)
c          if (l .eq. k) go to 10
c          b(l) = b(k)
c          b(k) = t
c 10      continue
c      call daxpy(n-k,t,a(k+1,k),1,b(k+1),1)
c 20      continue
c 30      continue
c
c      now solve  u*x = y
c
c      do 40 kb = 1, n
c          k = n + 1 - kb
c          b(k) = b(k)/a(k,k)
c          t = -b(k)
c          call daxpy(k-1,t,a(1,k),1,b(1),1)
c 40      continue
c      go to 100
c 50      continue
c
c      job = nonzero, solve  trans(a) * x = b

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c      first solve  trans(u)*y = b
c
c      do 60 k = 1, n
c          t = ddot(k-1,a(1,k),1,b(1),1)
c          b(k) = (b(k) - t)/a(k,k)
60      continue
c
c      now solve trans(l)*x = y
c
c      if (nm1 .lt. 1) go to 90
c      do 80 kb = 1, nm1
c          k = n - kb
c          b(k) = b(k) + ddot(n-k,a(k+1,k),1,b(k+1),1)
c          l = ipvt(k)
c          if (l .eq. k) go to 70
c              t = b(l)
c              b(l) = b(k)
c              b(k) = t
70      continue
80      continue
90      continue
100     continue
c      return
c      end
c      integer function idamax(n,dx,incx)
c
c      finds the index of element having max. absolute value.
c      jack dongarra, linpack, 3/11/78.
c      modified 3/93 to return if incx .le. 0.
c      modified 12/3/93, array(1) declarations changed to array(*)
c
c      double precision dx(*),dmax
c      integer i,incx,ix,n
c
c      idamax = 0
c      if( n.lt.1 .or. incx.le.0 ) return
c      idamax = 1
c      if(n.eq.1)return
c      if(incx.eq.1)go to 20
c
c      code for increment not equal to 1
c
c      ix = 1
c      dmax = dabs(dx(1))
c      ix = ix + incx
c      do 10 i = 2,n
c          if(dabs(dx(ix)).le.dmax) go to 5
c          idamax = i
c          dmax = dabs(dx(ix))
5      ix = ix + incx
10     continue
c      return
c
c      code for increment equal to 1
c
c      20 dmax = dabs(dx(1))
c      do 30 i = 2,n
c          if(dabs(dx(i)).le.dmax) go to 30
c          idamax = i
c          dmax = dabs(dx(i))
30     continue
c      return
c      end
c      subroutine daxpy(n,da,dx,incx,dy,incy)
c
c      constant times a vector plus a vector.
c      uses unrolled loops for increments equal to one.
c      jack dongarra, linpack, 3/11/78.
c      modified 12/3/93, array(1) declarations changed to array(*)
c
c      double precision dx(*),dy(*),da
c      integer i,incx,incy,ix,iy,m,mp1,n

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c
  if(n.le.0)return
  if (da .eq. 0.0d0) return
  if(incx.eq.1.and.incy.eq.1)go to 20
c
c      code for unequal increments or equal increments
c      not equal to 1
c
  ix = 1
  iy = 1
  if(incx.lt.0)ix = (-n+1)*incx + 1
  if(incy.lt.0)iy = (-n+1)*incy + 1
  do 10 i = 1,n
    dy(iy) = dy(iy) + da*dx(ix)
    ix = ix + incx
    iy = iy + incy
10 continue
  return
c
c      code for both increments equal to 1
c
c      clean-up loop
c
20 m = mod(n,4)
  if( m .eq. 0 ) go to 40
  do 30 i = 1,m
    dy(i) = dy(i) + da*dx(i)
30 continue
  if( n .lt. 4 ) return
40 mpl = m + 1
  do 50 i = mpl,n,4
    dy(i) = dy(i) + da*dx(i)
    dy(i + 1) = dy(i + 1) + da*dx(i + 1)
    dy(i + 2) = dy(i + 2) + da*dx(i + 2)
    dy(i + 3) = dy(i + 3) + da*dx(i + 3)
50 continue
  return
  end
  double precision function ddot(n,dx,incx,dy,incy)
c
c  forms the dot product of two vectors.
c  uses unrolled loops for increments equal to one.
c  jack dongarra, linpack, 3/11/78.
c  modified 12/3/93, array(1) declarations changed to array(*)
c
  double precision dx(*),dy(*),dtemp
  integer i,incx,incy,ix,iy,m,mpl,n
c
  ddot = 0.0d0
  dtemp = 0.0d0
  if(n.le.0)return
  if(incx.eq.1.and.incy.eq.1)go to 20
c
c      code for unequal increments or equal increments
c      not equal to 1
c
  ix = 1
  iy = 1
  if(incx.lt.0)ix = (-n+1)*incx + 1
  if(incy.lt.0)iy = (-n+1)*incy + 1
  do 10 i = 1,n
    dtemp = dtemp + dx(ix)*dy(iy)
    ix = ix + incx
    iy = iy + incy
10 continue
  ddot = dtemp
  return
c
c      code for both increments equal to 1
c
c
c

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c      clean-up loop
c
20 m = mod(n,5)
   if( m .eq. 0 ) go to 40
   do 30 i = 1,m
     dtemp = dtemp + dx(i)*dy(i)
30 continue
   if( n .lt. 5 ) go to 60
40 m+1 = m + 1
   do 50 i = m+1,n,5
     dtemp = dtemp + dx(i)*dy(i) + dx(i + 1)*dy(i + 1) +
*     dx(i + 2)*dy(i + 2) + dx(i + 3)*dy(i + 3) + dx(i + 4)*dy(i + 4)
50 continue
60 ddot = dtemp
   return
   end
   subroutine dscal(n,da,dx,incx)

c
c   scales a vector by a constant.
c   uses unrolled loops for increment equal to one.
c   jack dongarra, linpack, 3/11/78.
c   modified 3/93 to return if incx .le. 0.
c   modified 12/3/93, array(1) declarations changed to array(*)
c
   double precision da,dx(*)
   integer i,incx,m,m+1,n,nincx

c
   if( n.le.0 .or. incx.le.0 )return
   if(incx.eq.1)go to 20

c
   code for increment not equal to 1
c
   nincx = n*incx
   do 10 i = 1,nincx,incx
     dx(i) = da*dx(i)
10 continue
   return

c
   code for increment equal to 1
c
c
c   clean-up loop
c
20 m = mod(n,5)
   if( m .eq. 0 ) go to 40
   do 30 i = 1,m
     dx(i) = da*dx(i)
30 continue
   if( n .lt. 5 ) return
40 m+1 = m + 1
   do 50 i = m+1,n,5
     dx(i) = da*dx(i)
     dx(i + 1) = da*dx(i + 1)
     dx(i + 2) = da*dx(i + 2)
     dx(i + 3) = da*dx(i + 3)
     dx(i + 4) = da*dx(i + 4)
50 continue
   return
   end
c

```