# Definition and Implementation of a SLICOT Interface and a MATLAB Gateway for the Solution of Nonlinear Equations Systems <sup>1</sup>

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# 1 Introduction

The solution of nonlinear equations systems is necessary in order to cope with nonlinear control problems (Task V in NICONET). Following an approach similar to the work done in NICONET for other problems in task V, such as the solution of ordinary differential equations (ODE) and differential algebraic equations (DAE) systems, SLICOT and MATLAB interfaces have been implemented on top of existing software packages for the solution of nonlinear equations systems. For the moment, only one such library, known as KINSOL, has been considered.

This paper presents SLICOT and MATLAB interfaces for the KINSOL package. The SLICOT interface enables the user to call the KINSOL package by means of a subroutine with a SLICOT-compliant calling sequence ([3]). By means of the MATLAB interface the user can call the package from MATLAB, defining the problem by means of MATLAB functions.

The interfaces could be extended in the future in order to consider other nonlinear equations

systems solvers, although some restructuring of the interfaces would be necessary.

# 2 The KINSOL Package

The KINSOL package ([1]) has been developed at the Center for Applied Scientific Computing, Lawrence Livermore National Laboratory and can be freely downloaded from the web (http://www.llnl.gov/CASC/PVODE). Strongly connected with KINSOL are the packages PVODE and IDA, developed by the same Center. The three packages are complementary, forming a software tool for the solution of ODE and DAE systems.

KINSOL (Krylov Inexact Newton SOLver) is a general purpose solver for nonlinear systems of equations. Its most notable feature is that it uses Krylov Inexact Newton techniques in the system's approximate solution. KINSOL is written in ANSI C, although it contains interface routines so that it can be called also from Fortran programs.

KINSOL is available both in serial and parallel forms. In fact, the package is arranged so that selecting one of two forms of a single module in the compilation process allows the entire package to be created in either serial or parallel form. This document is only concerned with the serial form of KINSOL.

The nonlinear system of equations

$$F(u) = 0$$
,

where F(u) is a nonlinear function from  $\mathbb{R}^n$  to  $\mathbb{R}^n$ , is solved by this package. The Newton method used results in the solution of linear systems of the form

$$J(u)x = b,$$

where J(u) is the Jacobian of F at u. The solution of these systems by a Krylov method requires products of the form J(u)v, which are approximated by a difference quotient of the form

$$J(u)v \approx \frac{F(u+\sigma v) - F(u)}{\sigma}.$$

Thus, the Jacobian need not be formed explicitly.

A Fortran program making use of the KINSOL package must follow the following sequence of steps:

- 1. Allocate space for KINSOL, by calling the function FSKINMALLOC.
- 2. Set up the linear solver. This is done by calling a routine with the name FKINSPGMR<A><B>, where <A> is a digit that can be 0, 1 or 2, and <B> is a digit with value 0 or 1. Thus, there are 6 different routines. Which one to choose depends on the user-defined routines that are provided, as we will see later.
- 3. Solve the nonlinear system F(u) = 0, by calling the function FKINSOL.
- 4. Free the memory, by calling the function FKINFREE.

In addition to this, there are some user-defined routines that have to be taken into account. One of them must be necessarily defined by the user, while the others are optional. These routines must have the following names and calling sequences:

- SUBROUTINE KFUN (NEQ, UU, FVAL) (Required). This function evaluates the nonlinear function F.
- SUBROUTINE KPSOL (NEQ, UU, USCALE, FVAL, FSCALE, VTEM, FTEM, UROUND, NFE, IER) (Optional). This is used to solve the linear system Px = b, where P is a preconditioner matrix. If this function is not provided, then no preconditioning is done.
- SUBROUTINE KPRECO (NEQ, UU, USCALE, FVAL, FSCALE, VTEMP1, VTEMP2, UROUND, NFE, IER)

  (Optional). This is used to evaluate and preprocess any Jacobian-related data needed by KPSOL.
- SUBROUTINE FATIMES (V, Z, NEWU, UU, IER) (Optional). This function performs the matrix-vector multiply J(u)v, where J(u) is an approximate Jacobian matrix for that iteration.

The subroutine FKINSPGRM<A><B> to be used in step 2 above depends on which optional user-defined routines are provided. <A>= 0 means that neither KPRECO nor KPSOL routines are provided; <A>= 1 means that KPSOL is provided, but KPRECO is not; <A>= 2 indicates that both KPRECO and KPSOL are provided. <B>= 0 indicates that the routine FATIMES is not provided, while <B>= 1 indicates that FATIMES is provided.

# 3 SLICOT Interface

The SLICOT interface contains a single user-callable routine that performs the sequence of 4 steps described above. The calling sequence of the routine presents the following form:

```
SUBROUTINE KINSOL( GSTRAT, LINSU, NEQ, OPTIN, MAXL, MAXLRST, MSBPRE, UU, USCALE, FSCALE, CONSTR, IOPT, ROPT, TOL1, TOL2, INFO)
```

NEQ corresponds to the number of equations (and unknowns) in the sistem. MAXL, MAXLRST and MSBPRE are arguments related to the Krylov method for the linear system. UU contains on input the initial point, and on exit the solution of the system. IOPT and ROPT are integer and real arrays, respectively, used to provide additional input/output options. The argument LINSU corresponds to the name of the FKINSPGMR<A><B> routine to be used to set up the linear solver (step 2). Arguments indicating tolerances (TOL1 and TOL2) and the error indicator INFO, are named according to the SLICOT standard ([3]). Note that no work arrays are used, since KINSOL allocates and deallocates its own memory space. For a more detailed description of the arguments, see Appendix A of this Working Note.

# SLICOT-User's code CALL KINSOL(...) KFUN KPRECO KPSOL FATIMES SUBROUTINE KINSOL (...) KINSOL Interface FKINSPGMR<A><B> FKINSPGMR<A><B> FKINFREE

Figure 1: Application using the SLICOT interface.

The user-defined subroutines (KFUN, KPRECO, KPSOL and FATIMES) are not altered by the SLI-COT interface. Figure 1 shows a diagram where we can see the different layers of an application that makes use of the SLICOT interface described here.

# 4 MATLAB Interface

The main user-callable routine in the MATLAB interface for KINSOL presents the following form

function [uu, kinout] = kinsol(kfun, uu0, opts)

where kinout and opts are optional arguments. As we can see, the number of arguments reduces considerably with respect to the SLICOT interface. This has been done by grouping the input arguments in the structure opts, and the output arguments in the structure kinout. This keeps the interface easy to use while at the same time powerful.

The argument kfun is the function that evaluates F. uu0 is the initial point. uu is the solution of the nonlinear system, if no error ocurred.

The argument opts is a structure containing input options. Each option corresponds either to an argument of the user-callable routine of the SLICOT interface, or to a parameter in the arrays IOPT or ROPT, which are arguments of the same routine. To create this structure, the user can call the function

function kopt = kinsoptions()

which creates a structure of input options with empty fields. The user can then assign a value to specific fields as needed, before passing the structure to the kinsol function.

The output argument kinout is a structrure containing additional information about the solution of the nonlinear system (e.g. number of nonlinear iterations or number of function evaluations). Each element of the structure corresponds to an output parameter in the arrays IOPT or ROPT.

The MATLAB user-defined functions present the following forms (see Appendix B for a more detailed description):

function fval = kfun(uu)

function [nfe, ierr] = kpreco(uu, uscale, fval, fscale, kfun, uround, nfe)

function [x, nfe, ierr] = kpsol(uu, uscale, fval, fscale, r, kfun, uround, nfe)

function [z, ierr] = fatimes(v, new\_uu, uu)

The MATLAB interface is implemented by means of a gateway written in Fortran, plus two m-files. The gateway has an entry point from MATLAB, which calls the KINSOL package, and one routine for each of the user-provided functions (kfun, kpreco, kpsol and fatimes). These gateway routines only make the call to the corresponding MATLAB user function, returning the results to the KINSOL package.

The two m-files of the interface correspond to the two user-callable functions of this MATLAB interface. These functions deal with the structures opts and kinout, constructing them, forming the calling sequence of the KINSOL gateway from the information contained in opts, and filling kinout with the information returned by the gateway. These functions can be found in Appendix B of this Working Note.

Figure 2 shows a diagram where we can see the different layers involved when the MATLAB interface for KINSOL is used.

# 5 Examples of use

## 5.1 A trivial diagonal example

We consider a trivial diagonal example, taken from the KINSOL package, only for illustrating the use of the interfaces described here. In particular, the nonlinear function F is

$$F(u)_i = x_i^2 - i^2, \quad i = 1, \dots, 128,$$

and we choose as starting point  $x^*$ , with  $x_i^* = 2i$ .

#### 5.1.1 SLICOT interface

A Fortran program using the SLICOT interface for solving the problem could be as follows.

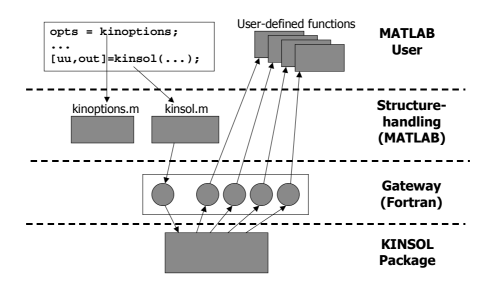


Figure 2: Diagram of the MATLAB interface.

```
PROGRAM DIAGSF
С
    ***********
С
С
     file: diagsf.f
С
       Simple diagonal test with SLICOT interface, using user-supplied
C
       preconditioner setup and solve routines (supplied in fortran, below).
       This example does a basic test of the code suite by solving the system
С
С
                            f(u) = 0. for
C
                            f(u) = u(i)^2 - i^2.
C
       no scaling is done.
C
       execute line: diagsf
С
С
     CONTRIBUTOR
С
С
     Fernando Alvarruiz, Vicente Hernandez
С
     Universidad Politecnica de Valencia, Spain
С
С
     PARAMETER (PROBSIZE=128)
     INTEGER NEQ, IER, GSTRAT
```

INTEGER IOPT(40) LOGICAL INOPT

DOUBLE PRECISION TOL1, TOL2

```
DOUBLE PRECISION ROPT(40)
     DOUBLE PRECISION UU(PROBSIZE), SCALE(PROBSIZE)
     DOUBLE PRECISION CONSTR(PROBSIZE)
     EXTERNAL FKINSPGMR20
С
        Preconditioner data
     DOUBLE PRECISION PP(PROBSIZE)
     COMMON /PCOM/ PP
     NEQ = PROBSIZE
     GSTRAT = 0
     TOL1 = 0.00001
     T0L2 = 0.0001
     INOPT = .FALSE.
     MAXL = 10
     MAXLRST = 2
     MSBPRE = 5
     MU = 0
     ML = 0
     DO 20 I = 1 , PROBSIZE
        UU(I) = 2.*I
        SCALE(I) = 1.
        CONSTR(I) = 0.
 20
    CONTINUE
WRITE(6,1240)
 1240 FORMAT('diagsf example case. this kinsol example code does a ')
      WRITE(6,1241)
 1241 FORMAT('128 eqn diagonal algebraic system. its purpose is to ')
      WRITE(6,1242)
 1242 FORMAT('demonstrate the use of the SLICOT interface.')
      WRITE(6,1244)
 1244 FORMAT('GSTRAT = INEXACT_NEWTON')
     CALL KINSOL (GSTRAT, FKINSPGMR2O, NEQ, INOPT, MAXL, MAXLRST,
                  MSBPRE, UU, SCALE, SCALE, CONSTR, IOPT, ROPT, TOL1,
    &
    &
                  TOL2, INFO)
      WRITE(6,1245)INFO
 1245 FORMAT(1X,' Return code is ',I5)
```

```
WRITE(6,1246)
 1246 FORMAT(1X,'The resultant values of UU are:',///)
      DO 30 I = 1, PROBSIZE, 4
         WRITE(6,1256)I, UU(I), UU(I+1), UU(I+2), UU(I+3)
 1256
         FORMAT(I4,4(1X,F10.6))
 30
    CONTINUE
       WRITE(6,1267) IOPT(4), IOPT(11), IOPT(5),
          IOPT(12), IOPT(13), IOPT(14)
 1267 FORMAT(1X,'NNI=',I4,' NLI=',I4,' NFE=',I4,' NPE=',I4,
            ' NPS=',I4,' NCFL=',I4)
      STOP
      END
   Following is the definition of the user-defined routines KFUN, KPRECO and KPSOL. It is not
necessary for this simple example to use preconditionning. However, it is used in order to
illustrate how to do it.
      The function defining the system f(u) = 0. Must be defined by a Fortran
С
      function of the following form.
      SUBROUTINE KFUN(NEQ, UU, FVAL)
      INTEGER NEQ
      DOUBLE PRECISION FVAL(*), UU(*)
      DO 10 I = 1 , NEQ
         FVAL(I) = UU(I)*UU(I) - I*I
 10
      CONTINUE
      RETURN
      END
    The routine KPRECO is the preconditioner setup routine. It is required that
    this specific name be used in order that the KINSOL code can find and
    link to it.
    The argument list must also be as illustrated below:
      SUBROUTINE KPRECO(NEQ, UDATA, USCALE, FDATA, FSCALE,
                         VTEMP1, VTEMP2, UROUND, NFE, IER)
     1
```

```
INTEGER NFE, IER
     DOUBLE PRECISION UDATA(*), USCALE(*), FDATA(*), FSCALE(*)
     DOUBLE PRECISION VTEMP1(*), VTEMP2(*)
     DOUBLE PRECISION UROUND
С
     COMMON PCOM * * * * * * * * *
     PARAMETER (PROBSIZE=128)
     DOUBLE PRECISION PP(PROBSIZE)
     COMMON /PCOM/ PP
C
     IER = 0
     DO 10 I = 1 , NEQ
       PP(I) = 0.5/(UDATA(I)+5.)
 10
     CONTINUE
     RETURN
     END
The routine KPSOL is the preconditioner solve routine. It is required that
   this specific name be used in order that the KINSOL code can find and
С
С
   link to it.
   The argument list must also be as illustrated below:
     SUBROUTINE KPSOL (NEQ, UDATA, USCALE, FDATA, FSCALE,
                   VTEM, FTEM, UROUND, NFE, IER)
    1
     INTEGER NFE, IER
     DOUBLE PRECISION UDATA(*), USCALE(*), FDATA(*), FSCALE(*)
     DOUBLE PRECISION VTEM(*), FTEM(*)
     DOUBLE PRECISION UROUND
С
     COMMON PCOM * * * * * * * * * *
     PARAMETER (PROBSIZE=128)
     DOUBLE PRECISION PP(PROBSIZE)
     COMMON /PCOM/ PP
С
```

```
IER = 0

DO 10 I = 1 , NEQ
    VTEM(I) = VTEM(I) * PP(I)

10    CONTINUE

RETURN
END
```

When running the program (in a Linux PC), the following output is obtained:

diagsf example case. This kinsol example code does a 128 eqn diagonal algebraic system. Its purpose is to demonstrate the use of the SLICOT interface.

GSTRAT = INEXACT\_NEWTON

return code is 0

The resultant values of UU are:

```
1.000000
                2.000000
                            3.000000
1
                                       4.000000
     5.000000
                6.000000
                            7.000000
                                       8.000000
5
9
     9.000000
               10.000000
                          11.000000
                                      12.000000
13
   13.000000
               14.000000
                          15.000000
                                      16.000000
17
   17.000000
               18.000000
                          19.000000
                                      20.000000
21
    21.000000
               22.000000
                          23.000000
                                      24.000000
25
   25.000000
               26.000000
                          27.000000
                                      28.000000
29
    29.000000
               30.000000
                          31.000000
                                      32.000000
33
   33.000000
               34.000000
                           35.000000
                                      36.000000
37
   37.000000
               38.000000
                           39.000000
                                      40.000000
41
   41.000000
               42.000000
                           43.000000
                                      44.000000
45
   45.000000
               46.000000
                          47.000000
                                      48.000000
               50.000000
49
   49.000000
                           51.000000
                                      52.000000
   53.000000
               54.000000
                           55.000000
                                      56.000000
57
   57.000000
               58.000000
                           59.000000
                                      60.000000
61
   61.000000
               62.000000
                           63.000000
                                      64.000000
   65.000000
               66.000000
                          67.000000
65
                                      68.000000
69
   69.000000
               70.000000
                          71.000000
                                      72.000000
73
   73.000000
               74.000000
                          75.000000
                                      76.000000
77
   77.000000
               78.000000
                          79.000000
                                      80.000000
81
   81.000000
               82.000000
                           83.000000
                                      84.000000
85
   85.000000
               86.000000
                           87.000000
                                      88.000000
89
   89.000000
               90.000000
                           91.000000
                                      92.000000
93
   93.000000
               94.000000
                          95.000000
                                      96.000000
```

```
97 97.000000 98.000000 99.000000 100.000000
101 101.000000 102.000000 103.000000 104.000000
105 105.000000 106.000000 107.000000 108.000000
109 109.000000 110.000000 111.000000 112.000000
113 113.000000 114.000000 115.000000 116.000000
117 117.000000 118.000000 119.000000 120.000000
121 121.000000 122.000000 123.000000 124.000000
125 125.000000 126.000000 127.000000 128.000000
NNI= 7 NLI= 21 NFE= 36 NPE= 2 NPS= 28 NCFL= 0
```

#### 5.1.2 MATLAB interface

To solve the problem from MATLAB, the following simple call is enough

```
[uu, kinout] = kinsol('func', 2*[1:128]');
```

and the function func is defined as follows (in this case we do not use preconditionning).

```
function fu = func (uu)
fu = uu.^2 - [1:length(uu)]'.^2;
```

The contents of uu after calling the kinsol function are as expected, while kinout contains the following infomation (results may vary slightly depending on the computer used):

#### kinout =

NonLinIters: 29
NumFuncEvals: 315
NumBetaCondFail: 0
NumBacktracks: 0
FNorm: 0

StepLength: 1.2645e-014

LinIters: 256
NumPrecEvals: 0
NumPSolve: 0
NumLinConvFails: 24

# 5.2 Predator-prey system

The following example, taken from the KINSOL package, is more illustrative of the power of KINSOL with real problems. The example problem is a Partial Differential Equation model of a multi-species food web [2], in which mutual competition and/or predator-prey relationships in a spatial domain are simulated. For this problem the dependent variable c (concentration of a species in a spatial point) replaces the generic dependent variable u used above. We consider a

model with s=2p species, where both species  $1, \ldots, p$  (the prey) and  $p+1, \ldots, s$  (the predators) have infinitely fast reaction rates:

$$0 = f_i(x, y, c) + d_i(c_{xx}^i + c_{yy}^i) \quad (i = 1, 2, \dots, s),$$

where (x, y) are the coordinates of a point in the plane;  $c_{xx}^i$  and  $c_{yy}^i$  denote the second derivatives with respect to x and y respectively, of the concentration of the i-th species, and

$$f_i(x, y, c) = c^i(b_i + \sum_{j=1}^s a_{ij}c^j).$$

The interaction and diffusion coefficients  $(a_{ij}, b_i, d_i)$  could be functions of (x, y) in general. The choices made for this test problem are for a simple model of p prey and p predator species, arranged in that order in the vector c. We take the various coefficients to be as follows:

$$\begin{cases} a_{ii} = -1 & (i = 1, 2, \dots, s) \\ a_{ij} = -0.5 \cdot 10^{-6} & (i \le p, j > p) \\ a_{ij} = 10^4 & (i > p, j \le p) \\ a_{ij} = 0 & \text{for all other } i, j \end{cases}$$

$$\begin{cases} b_i = b_i(x, y) = (1 + \alpha xy) & (i \le p) \\ b_i = b_i(x, y) = -(1 + \alpha xy) & (i > p) \end{cases}$$

and

$$\begin{cases} d_i = 1 & (i \le p) \\ d_i = 0.5 & (i > p). \end{cases}$$

The domain is the unit square  $0 \le x, y \le 1$ . The boundary conditions are of Neumann type (zero normal derivatives) everywhere. The coefficients are such that a unique stable equilibrium is guaranteed to exist when  $\alpha$  is zero [2]. Empirically, a stable equilibrium appears to exist when  $\alpha$  is positive, although it may not be unique. In this problem we take  $\alpha=1$ . The initial species concentrations used are constant functions by species type, in particular

$$\begin{cases} c_i = 1 & (i \le p) \\ c_i = 30000 & (i > p). \end{cases}$$

The PDE system (with boundary conditions) was discretized using central differencing on an  $MX \times MY$  mesh, with the resulting nonlinear system having size  $N = s \cdot MX \cdot MY$ .

#### 5.2.1 SLICOT Interface

The following Fortran program implements the solution of this problem using the SLICOT interface to KINSOL.

```
* File: predprey.f
     CONTRIBUTOR
     Fernando Alvarruiz, Vicente Hernandez
     Universidad Politecnica de Valencia, Spain
     Adapted from:
        kinxs.c, by Allan G. Taylor and Alan C. Hindmarsh @ LLNL
        Version of 16 January 2001
* Example problem for KINSol, serial machine version.
* This example solves a nonlinear system that arises from a system of
* partial differential equations. The PDE system is a food web
* population model, with predator-prey interaction and diffusion on the
* unit square in two dimensions. The dependent variable vector is
* c = (c, c, ..., c)
                                     (denoted by the variable cc)
* and the PDE's are as follows:
                         + c ) + f (x,y,c) (i=1,...,ns)
          0 = d(i)*(c
                          уу
   where
                                ns
   f(x,y,c) = c * (b(i) + sum a(i,j)*c)
                                j=1
* The number of species is ns = 2 * np, with the first np being prey and
* the last np being predators. The number np is both the number of prey and
* predator species. The coefficients a(i,j), b(i), d(i) are
   a(i,i) = -AA \quad (all \ i)
   a(i,j) = -GG \quad (i \le np, j > np)
   a(i,j) = EE (i > np, j \le np)
   b(i) = BB * (1 + alpha * x * y) (i \le np)
   b(i) = -BB * (1 + alpha * x * y) (i > np)
   d(i) = DPREY (i \le np)
```

```
d(i) = DPRED (i > np)
  The various scalar parameters are set using parameter statements
  The boundary conditions are: normal derivative = 0.
  The initial guess is constant in x and y, although the final
  solution is not.
  The PDEs are discretized by central differencing on a MX by MY mesh.
  The nonlinear system is solved by KINSOL using the method specified in
  local variable gstrat .
  The preconditioner matrix is a block-diagonal matrix based on the
  partial derivatives of the interaction terms f only.
* References:
* 1. Peter N. Brown and Youcef Saad,
    Hybrid Krylov Methods for Nonlinear Systems of Equations
    LLNL report UCRL-97645, November 1987.
* 2. Peter N. Brown and Alan C. Hindmarsh,
    Reduced Storage Matrix Methods in Stiff ODE systems,
    Lawrence Livermore National Laboratory Report UCRL-95088, Rev. 1,
    June 1987, and Journal of Applied Mathematics and Computation, Vol. 31
    (May 1989), pp. 40-91. (Presents a description of the time-dependent
    version of this test problem.)
implicit none
     ****** Common parameters ************************
     integer NS, NP, MX, MY, NEQ
     double precision AA, EE, GG, BB, DPREY, DPRED, ALPHA, AX, AY,
                     DX, DY, FTOL, STOL, PREYIN, PREDIN
     Number of species
     parameter(NS = 6)
     Number of prey/predator species
С
     parameter(NP = NS/2)
```

```
parameter(MX = 8)
     Number of y mesh points
     parameter(MY = 8)
     Total range of x variable
С
     parameter(AX
                 = 1.0)
     Total range of y variable
С
     parameter(AY = 1.0)
     Distance between x mesh points
С
     parameter(DX = AX/(MX-1))
     Distance between y mesh points
     parameter(DY
                 = AY/(MY-1)
С
     Number of nonlinear equations
     parameter(NEQ = NS * MX * MY)
     parameter(AA
                  = 1.0)
     parameter(EE = 10000.0)
     parameter(GG = 0.5e-6)
     parameter(BB = 1.0)
     parameter(DPREY= 1.0)
     parameter(DPRED= 0.5)
     parameter(ALPHA= 1.0)
     Tolerances for the solution of the system
С
     parameter(FTOL = 1.e-7)
     parameter(STOL = 1.e-13)
     Initial guess for prey concentrations
     parameter(PREYIN = 1.0)
С
     Initial guess for predator concentrations
     parameter(PREDIN = 30000.0)
С
     ****** End of Common parameters *****************
     С
     double precision Acoef(NS,NS), Bcoef(NS)
```

С

Number of x mesh points

```
Preconditioning data
С
      double precision P(NS,NS,MX,MY), ipiv(NS,MX,MY)
      Interaction rates, i.e. result of evaluating f (x,y,c)
С
С
      double precision rates(NS,MX,MY)
      double precision cox(NS), coy(NS)
     Machine unit roundoff and its square root
     double precision uround, squrnd
     common /pcom/ Acoef, Bcoef, P, ipiv, rates, cox, coy, uround,
                    squrnd
      ***** End of common variables *****************
С
     Local variables
С
      integer i, info, gstrat, maxl, maxlrst, msbpre
      integer iopt(40)
      logical inopt
      double precision ropt(40)
      double precision cc(NS,MX,MY), sc(NS,MX,MY)
      double precision constr(NEQ)
      external fkinspgmr20
      call InitUserData()
      call SetInitialProfiles(cc, sc)
      do i = 1, NEQ
         constr(i) = 0.0
      end do
      gstrat = 0
      inopt = .false.
     maxl = 15
     maxlrst = 2
     msbpre = 0
     print *, 'Predator-prey test problem -- KINSol (serial version)'
      print *, 'Mesh dimensions =', MX, ' X', MY
      print *, 'Number of species =', NS
```

```
print *, 'Total system size =', NEQ
     print *, 'Flag globalstrategy =', gstrat,
    & ' (0 = Inex. Newton, 1 = Linesearch)'
     print *, 'Linear solver is SPGMR with maxl =', maxl,
    & ' maxlrst =', maxlrst
     print *, 'Preconditioning uses interaction-only block-diagonal ',
    & 'matrix'
     print *, 'Tolerance parameters: fnormtol =', FTOL,
    & 'scsteptol =', STOL
     print *, 'Initial profile of concentration'
     print *, 'At all mesh points:', PREYIN, PREYIN,
    & PREDIN, PREDIN, PREDIN
     Call KINSol and print output concentration profile
     call kinsol (gstrat , fkinspgmr20, NEQ, inopt, maxl, maxlrst,
    &
                 msbpre, cc, sc, sc, constr, iopt, ropt, FTOL,
    &
                 STOL, info)
     if (info .ne. 0) then
        print *, 'KINSOL failed, returning ', info
        stop
     end if
     print *, ''
     print *, ''
     print *, 'Computed equilibrium species concentrations:'
     call PrintOutput(cc)
     Print final statistics
     print *, ',
     call PrintFinalStats(iopt)
     stop
     end
The function defining the system f(u) = 0 must be defined by a fortran
     function of the following form.
     subroutine kfun(neqn, cc, fval)
     implicit none
```

```
(... Deleted lines correponding to common parameters and variables ...)
      Function arguments
      integer neqn
      double precision fval(NS,MX,MY), cc(NS,MX,MY)
С
      Local variables
      double precision xx, yy, dcx, dcy
      integer jx, jy, is, up, down, left, right
      Loop over all mesh points, evaluating rate array at each point
      do jy=1,MY
          yy = (jy-1)*DY
          if (jy .ne. 1) then
              down = jy-1
          else
              down = 2
          end if
          if (jy .ne. MY) then
              up = jy+1
          else
              up = MY-1
          end if
          do jx=1,MX
              xx = (jx-1)*DX
              if (jx .ne. 1) then
                  left = jx-1
              else
                  left = 2
              end if
              if (jx .ne. MX) then
                  right = jx+1
              else
                  right = MX-1
              end if
              call WebRate(xx, yy, cc(1,jx,jy), rates(1,jx,jy))
              do is = 1,NS
                 dcy = coy(is) *
                       (cc(is,jx,down)+cc(is,jx,up)-2*cc(is,jx,jy))
     &
```

```
dcx = cox(is) *
                      (cc(is,left,jy)+cc(is,right,jy)-2*cc(is,jx,jy))
    &
                fval(is, jx, jy) = dcy + dcx + rates(is, jx, jy)
             end do
         end do
     end do
     return
     end
The routine kpreco is the preconditioner setup routine. It is required that
   this specific name be used in order that the KINSOL code can find and
С
   link to it.
   The argument list must also be as illustrated below:
     subroutine kpreco(neqn, cc, cscale, fval, fscale,
                      vtemp1, vtemp2, urnd, nfe, ier)
    1
     implicit none
(... Deleted lines correponding to common parameters and variables ...)
     Function arguments
С
     integer neqn, nfe, ier
     double precision cc(NS,MX,MY), cscale(NS,MX,MY), fval(NS,MX,MY),
                     fscale(NS,MX,MY)
     double precision vtemp1(*), vtemp2(*)
     double precision urnd
     Local variables
С
     integer jx, jy, j, i, info
     double precision csave, xx, yy, r, r0, fac
     double precision PertRates(NS)
     double precision WNorm
     ier = 0
     fac = WNorm(NEQ, fval, fscale)
     r0 = 1000.0 * uround * fac * NEQ
     if (r0 .eq. 0.0) r0 = 1
```

```
С
      Loop over all spatial points; get NS-sized Jacobian block at each
      do jy = 1, MY
          yy = (jy-1)*DY
          do jx = 1, MX
              xx = (jx-1)*DX
С
              Compute difference quotients of interaction rate function
              do j = 1,NS
                  Save the j,jx,jy element of cc
С
                  csave = cc(j,jx,jy)
                  r = max(squrnd*abs(csave), r0/cscale(j,jx,jy))
                  Perturb the j,jx,jy element of cc
С
                  cc(j,jx,jy) = cc(j,jx,jy) + r
                  fac = 1.0/r
                  call WebRate(xx, yy, cc(1,jx,jy), PertRates)
                  Restore the j,jx,jy element of cc
С
                  cc(j,jx,jy) = csave
                  Load the j-th column of difference quotients
С
                  do i = 1, NS
                      P(i,j,jx,jy) = (PertRates(i)-rates(i,jx,jy))*fac
                  end do
              end do
С
              Do LU decomposition of preconditioner block
              call dgetrf(NS, NS, P(1,1,jx,jy), NS, ipiv(1,jx,jy), info)
              if (info .ne. 0) then
                  ier = info
                  return
              end if
          end do
      end do
      return
      end
```

```
The routine kpsol is the preconditioner solve routine. It is required that
   this specific name be used in order that the KINSOL code can find and
С
   link to it.
С
   The argument list must also be as illustrated below:
     subroutine kpsol(neqn, cc, cscale, fval, fscale,
                     vv, ftem, urnd, nfe, ier)
     implicit none
(... Deleted lines correponding to common parameters and variables ...)
     Function arguments
     integer neqn, nfe, ier
     double precision cc(NS,MX,MY), cscale(NS,MX,MY), fval(NS,MX,MY),
                     fscale(NS,MX,MY)
     double precision vv(NS,MX,MY), ftem(*)
     double precision urnd
     Local variables
С
     integer jx, jy, info
     ier = 0
     do jy = 1,MY
         do jx = 1,MX
             For each (jx,jy), solve a linear system of size NS
С
             call dgetrs('N', NS, 1, P(1,1,jx,jy), NS, ipiv(1,jx,jy),
    &
                        vv(1,jx,jy), NS, info)
         end do
     end do
     return
     end
subroutine WebRate (xx, yy, cxy, rxy)
     implicit none
С
     This function computes f(x,y,c) at point (xx, yy)
(... Deleted lines correponding to common parameters and variables ...)
```

```
С
     Function arguments
     double precision xx, yy
     double precision cxy(NS), rxy(NS)
     Local variables
С
     double precision ONE
     parameter (ONE=1.0)
     integer is
     double precision fac
     (rxy = Bcoef)
     call dcopy(NS, Bcoef, 1, rxy, 1)
     fac = 1 + ALPHA*xx*yy
     ( rxy = fac*rxy + Acoef*cxy )
     call dgemv('N', NS, NS, ONE, Acoef, NS, cxy, 1, fac, rxy, 1)
     do is = 1,NS
         rxy(is) = cxy(is) * rxy(is)
     end do
     return
     end
subroutine InitUserData()
     implicit none
(... Deleted lines correponding to common parameters and variables ...)
     Local variables
     integer i, j
     double precision dx2, dy2
     double precision dlamch
     uround = dlamch('E')
     squrnd = sqrt(uround)
     dx2 = DX*DX
     dy2 = DY*DY
     do j=1,NP
         Fill in the portion of Acoef in the four quadrants,
С
```

```
column by column
С
         do i=1,NP
            Acoef(i,j) = 0.0
             Acoef(i, NP+j) = -GG
             Acoef(NP+i, j) = EE
             Acoef(NP+i, NP+j) = 0.0
         end do
         and then change the diagonal elements of Acoef to -AA
С
         Acoef(j,j) = -AA
         Acoef(NP+j, NP+j) = -AA
         Bcoef(j) = BB
         Bcoef(NP+j) = -BB
         cox(j) = DPREY/dx2
         cox(NP+j) = DPRED/dx2
         coy(j) = DPREY/dy2
         coy(NP+j) = DPRED/dy2
     end do
     return
     end
subroutine SetInitialProfiles(cc, sc)
     implicit none
(... Deleted lines correponding to common parameters and variables ...)
     Local variables
     double precision cc(NS,MX,MY), sc(NS,MX,MY)
     integer i, j, is
     do j=1,MY
         do i=1,MX
            do is = 1,NP
                cc(is,i,j) = PREYIN
                sc(is,i,j) = 1
             end do
             do is = NP+1, NS
                cc(is,i,j) = PREDIN
                sc(is,i,j) = 0.00001
```

```
end do
         end do
     end do
     return
     end
     subroutine PrintOutput(cc)
     implicit none
(... Deleted lines correponding to common parameters ...)
     double precision cc(NS,MX,MY)
     Local variables
     integer is
     print *, 'At bottom left:'
     do is = 1,NS
         print *, cc(is,1,1)
     end do
     print *, 'At top right:'
     do is = 1,NS
         print *, cc(is,MX,MY)
     end do
     return
     end
     subroutine PrintFinalStats(iopt)
     implicit none
     integer iopt(40)
     print *, 'Final Statistics:'
     print *, ' Non-linear iterations
                                       = ', iopt(4)
     print *, ' Linear iterations
                                         = ', iopt(11)
     print *, ' Num. function evals
                                         = ', iopt(5)
     print *, ' Num. precond evals
                                         = ', iopt(12)
     print *, ' Num. precond solves = ', iopt(13)
     print *, ' Num. linear conv. failures = ', iopt(14)
```

```
return
      end
      double precision function WNorm(n, x, w)
      Weighted 2-norm
С
      implicit none
      integer n
      double precision x(n), w(n)
      integer i
      double precision s, prodi
      s = 0.0
      do i = 1,n
          prodi = x(i) * w(i);
          s = s + prodi * prodi;
      end do
      WNorm = sqrt(s);
      return
      end
```

The program defines in the first place different constants and variables common to the main program and other subroutines. Other coefficients defining the problem, such as  $a_{ij}$  and machine unit roundoff, are initialized in subroutine InitUserData. Subroutine SetInitialProfiles provides the initial guess for the species concentrations, and also a vector that will be used for scaling both function values and concentrations within KINSOL. Note that the dependent variable c is implemented by means of a 3-dimensional array cc, although the KINSOL solver will see it as a 1-dimensional array, ordered in the following way

$$c = [c_1^{x_1 y_1} \dots c_s^{x_1 y_1} \dots c_1^{x_{MX}, y_1} \dots c_s^{x_{MX}, y_1} \ c_1^{x_1 y_2} \dots c_s^{x_1 y_2} \dots c_1^{x_{MX}, y_{MY}} \dots c_s^{x_{MX}, y_{MY}}]^T$$

where  $c_i^{x_j y_k}$  denotes the concentration of species i at the point  $(x_j, y_k)$ .

The implementation of the nonlinear function is contained in subroutine **kfun**, which in turn calls **WebRates** in order to compute the terms  $f_i(x, y, c)$ .

If we look at the call to the kinsol subroutine (the SLICOT interface) in the main program, we can see that the subroutine name fkinspgmr20 is used as the second argument, wich means that preconditioning will be used, and the user-defined subroutines kpreco and kpsol must be provided. The preconditioner matrix is a block-diagonal matrix based on the partial derivatives of the interaction terms f(x, y, c) only. This matrix is built and factorized in kpreco, using finite differences for computing the partial derivatives. LAPACK is used for LU factorization of the diagonal blocks. Subroutine kpsol solves a linear system with the factorized preconditioner as the coefficient matrix. Again LAPACK is used for solving the triangular systems.

The following output is obtained when running the program on a Linux PC (again we must note that results may vary slightly depending on the computer/compiler used).

```
Predator-prey test problem -- KINSol (serial version)
Mesh dimensions = 8 X 8
Number of species = 6
Total system size = 384
Flag globalstrategy = 0 (0 = Inex. Newton, 1 = Linesearch)
Linear solver is SPGMR with maxl = 15 maxlrst = 2
Preconditioning uses interaction-only block-diagonal matrix
Tolerance parameters: fnormtol = 1.00000001E-07 scsteptol = 9.99999982E-14
Initial profile of concentration
At all mesh points: 1. 1. 1. 30000. 30000. 30000.
Computed equilibrium species concentrations:
At bottom left:
 1.16427931
 1.16427931
 1.16427931
 34927.4876
 34927.4876
 34927.4876
At top right:
 1.25796688
 1.25796688
 1.25796688
 37736.6641
 37736.6641
 37736.6641
Final Statistics:
 Non-linear iterations
 Linear iterations
                           = 352
 Num. function evals
                           = 371
 Num. precond evals
                           = 1
 Num. precond solves
                          = 361
 Num. linear conv. failures = 7
```

#### 5.2.2 MATLAB Interface

The file predprey.m, which is listed next, is a MATLAB script for the solution of the predator-prey problem using the MATLAB interface to KINSOL.

global data

```
% Initialization
data = initprpr(8,8,6);
[cc0, sc] = SetInitialProfiles(data);

% Set KINSOL Options
opts = kinsoptions;
opts.Uscale = sc;
opts.Fscale = sc;
opts.FNormTol = 1e-7;
opts.ScStepTol = 1e-13;
opts.MaxLinDim = 15;
opts.MaxLinRestarts = 2;
opts.PrecondSetFunc = 'kpreco';
opts.PrecondSolveFunc = 'kpsol';

% Solve nonlinear system of equations
[cc, kinout] = kinsol('funcprpr', cc0, opts);
```

The script calls auxiliary functions initprpr and SetInitialProfiles. The former initializes the problem coefficients and stores them in the structure global variable data, while the latter gives the inital guess for the concentrations and the scaling vector for KINSOL. These functions are listed next.

```
function data = initprpr(mx, my, ns)
ns2 = ns/2;
data.ax = 1;
data.ay = 1;
data.mx = mx;
data.my = my;
data.dx = data.ax/(data.mx-1);
data.dy = data.ay/(data.my-1);
data.ns = ns;
data.alpha = 1;
BB = 1;
data.bcoef = [ones(1,ns2)*BB -ones(1,ns2)*BB]';
AA = 1;
GG = 0.5e-6;
EE = 10000;
data.acoef = [...
   -diag(ones(ns2,1)*AA) -ones(ns2)*GG; ...
```

```
ones(ns2)*EE -diag(ones(ns2,1)*AA)];
DPREY = 1;
DPRED = 0.5;
a = DPREY/(data.dx*data.dx);
b = DPRED/(data.dx*data.dx);
data.cox = [ones(1,ns2)*a ones(1,ns2)*b]';
a = DPREY/(data.dy*data.dy);
b = DPRED/(data.dy*data.dy);
data.coy = [ones(1,ns2)*a ones(1,ns2)*b]';
% Initialization of Preconditioning data
data.sqruround = sqrt(eps);
data.L = zeros(ns,ns,mx,my);
data.U = zeros(ns,ns,mx,my);
function [cc, sc] = SetInitialProfiles(data)
mx = data.mx;
my = data.my;
ns = data.ns;
np = ns/2;
PREYIN=1;
PREDIN=30000;
cc = ones(ns, mx, my);
sc = ones(ns,mx,my);
cc(1:np,:,:) = cc(1:np,:,:) * PREYIN;
cc(np+1:ns,:,:) = cc(np+1:ns,:,:) * PREDIN;
sc(1:np,:,:) = sc(1:np,:,:)
                                * 1;
sc(np+1:ns,:,:) = sc(np+1:ns,:,:) * 0.00001;
cc = reshape(cc, ns*mx*my, 1);
sc = reshape(sc, ns*mx*my, 1);
   The evaluation of the function and the preconditioning related functions are presented next:
function fval=funcprpr(cc)
% System function for the predator-prey system
global data;
dx = data.dx;
```

```
dy = data.dy;
mx = data.mx;
my = data.my;
ns = data.ns;
cox = data.cox;
coy = data.coy;
rates = zeros(ns, mx, my);
cc = reshape(cc, ns, mx, my);
fval = zeros(ns, mx, my);
down = [2 [1:my-1]];
     = [[2:my] my-1]';
left = [2 [1:mx-1]]';
right = [[2:mx] mx-1]';
yvec = dy * [0:my-1]';
xvec = dx * [0:mx-1]';
% Loop over all mesh points, evaluating function at each point
for jy = 1:my
    % Get species interaction rate at (:, yy)
    rates(:,:,jy) = WebRateY(xvec, yvec(jy), cc(:,:,jy));
    fval(:,:,jy) = ...
        diag(coy) * (cc(:, :, up(jy)) + cc(:, :, down(jy)) - 2*cc(:, :, jy)) + ...
        diag(cox) * (cc(:, right, jy) + cc(:, left, jy) - 2*cc(:, :, jy)) + ...
        rates(:,:,jy);
end
data.rates = rates;
fval = reshape(fval, ns*mx*my, 1);
function rates = WebRateY(xvec, yy, cy)
% Computes rates for Interaction of species at points
% (xvec(i), yy) for all i
global data;
alpha = data.alpha;
```

```
fac = 1 + alpha * yy * xvec;
rates = data.bcoef * fac' + data.acoef * cy;
rates = cy .* rates;
function [nfe, ierr] = kpreco(cc, cscale, fval, fscale, kfun, uround, nfe)
% Preconditioner setup routine. Generates P
  global data;
  ierr = 0;
  mx = data.mx;
  my = data.my;
  dx = data.dx;
  dy = data.dy;
  ns = data.ns;
  sqruround = data.sqruround;
  cc = reshape(cc, ns, mx, my);
  cscale = reshape(cscale, ns, mx, my);
  r0 = 1000 * eps * norm(fval.*fscale) * length(fval);
  if (r0 == 0) r0=1; end
  yvec = dy * [0:my-1]';
  xvec = dx * [0:mx-1]';
  % Loop over spatial points; get NS-sized Jacobian block at each point
  P = zeros(ns);
  for jy = 1:my
      for jx = 1:mx
          for j = 1:ns
              % Save element of cc
              csave = cc(j,jx,jy);
              % Perturb element of cc
              r = max(sqruround*abs(csave), r0/cscale(j,jx,jy));
              cc(j,jx,jy) = cc(j,jx,jy) + r;
              perturb_rates = WebRate(xvec(jx), yvec(jy), cc(:,jx,jy));
              % Restore element of cc
```

```
cc(j,jx,jy) = csave;
              % Load j-th column of difference quotients
              P(:,j) = (perturb_rates - data.rates(:,jx,jy)) ./ r;
          end
          % Do LU decomposition of preconditioner block
          [data.L(:,:,jx,jy), data.U(:,:,jx,jy)] = lu(P);
      end
  end
function rates = WebRate(xx, yy, cxy)
\% Computes rates for Interaction of species at point (xx, yy)
global data;
alpha = data.alpha;
fac = 1 + alpha*xx*yy;
rates = fac * data.bcoef + data.acoef * cxy;
rates = cxy .* rates;
function [x, nfe, ierr] = kpsol(cc, cscale, fval, fscale, r, funcprpr, uround, nfe)
% Preconditioner setup routine. Generates P
  global data;
  ierr = 0;
  mxmy = data.mx * data.my;
  ns = data.ns;
  r = reshape(r, ns, mxmy);
  x = zeros(ns, mxmy);
  % For each spatial point (jx, jy), solve a NS-sized linear system
  for jxy = 1:mxmy
          x(:,jxy) = data.L(:,:,jxy) \setminus r(:,jxy);
          x(:,jxy) = data.U(:,:,jxy) \setminus x(:,jxy);
  end
  x = reshape(x, ns*mxmy, 1);
```

The evaluation of the nonlinear function is done in funcprpr, which, similarly to the Fortran implementation, calls the function WebRateY in order to evaluate the terms f(x, y, c). However, in this case funcprpr presents a single loop, which corresponds to the loop over the y coordinates,

i.e. the outermost loop in the corresponding Fortran subroutine kfun. The other two loops have been in this case *vectorized* (i.e. they are now implicit, forming part of matrix/vector operations). This vectorization of loops is important in order to obtain acceptable execution times in MATLAB code.

As a consequence of this loop vectorization, WebRateY must in this case compute the rates f(x, y, c) not for a single (x, y) point, but for all points with a given y coordinate.

The code for the preconditioning is very similar to the corresponding Fortran code. Note that in order to compute the preconditioner we use function WebRate instead of WebRateY, because in this case we need to evaluate f(x, y, c) only one (x, y) point at a time.

The final concentrations of the bottom left and top right points, and KINSOL statistical information is presented next (results obtained on a Windows PC with Matlab 6):

#### Final concentrations at bottom left point:

- 1.0e+004 \*
- 0.00011642793077
- 0.00011642793077
- 0.00011642793077
- 3.49274875697278
- 3.49274875697278
- 3.49274875697278

#### Final concentrations at top right point:

- 1.0e+004 \*
- 0.00012579668753
- 0.00012579668753
- 0.00012579668753
- 3.77366640744681
- 3.77366640744681
- 3.77366640744681

#### Statistical KINSOL information:

#### kinout =

NonLinIters: 8
NumFuncEvals: 325
NumBetaCondFail: 0
NumBacktracks: 0

FNorm: 5.367633801921303e-008 StepLength: 1.449283603070963e-007 LinIters: 308
NumPrecEvals: 1
NumPSolve: 316
NumLinConvFails: 6

# References

- [1] ALLAN G. TAYLOR AND ALAN C. HINDMARSH, User Documentation for KINSOL, a Non-linear Solver for Sequential and Parallel Computers, Center for Applied Scientific Computing, L-561, LLNL, Livermore, CA 94551.
- [2] P. N. Brown, Decay to Uniform States in Food Webs, SIAM J. Appl. Math., 46 (1986), 376-392.
- [3] WORKING GROUP ON SOFTWARE (WGS), SLICOT Implementation and Documentation Standards, WGS-Report 96-1, Eindhoven University of Technology, February 1998, ftp://wgs.esat.kuleuven.ac.be/pub/WGS/REPORTS/rep96-1.ps.Z.

# A SLICOT interface for KINSOL

We present here the source code of the subroutine KINSOL, which implements the SLICOT interface described in this Working Note.

```
SUBROUTINE KINSOL (GSTRAT, LINSU, NEQ, OPTIN, MAXL, MAXLRST,
     &
                         MSBPRE, UU, USCALE, FSCALE, CONSTR,
     &
                          IOPT, ROPT, TOL1, TOL2, INFO)
C
C
      WGS COPYRIGHT 2000.
C
C
      PURPOSE
C
C
      To solve a nonlinear system of equations F(u)=0, where F(u) is
C
C
      a nonlinear function from R to R , using Krylov Inexact Newton
C
      techniques.
C
C
      ARGUMENTS
C
C
      GSTRAT
             (input) INTEGER
C
              Indicates the global strategy to apply the computed
C
              increment delta in the solution UU. Choices are:
C
              0 - Inexact Newton.
              1 - Linesearch.
C
```

C C

C

C

C

C

C

C

C

C

C

C

C

C

C

C

С

#### LINSU SUBROUTINE

Linear Solver Set-up Routine. This is the KINSOL routine to be called to set-up the linear solver. The user should specify here one of the 6 different Fortran-callable routines provided by KINSOL for this purpose. The choice to be used depends on which of the optional user-defined routines are provided by the user (see User-defined routines below).

LINSU can be one of the following routines: FKINSPGMR00, FKINSPGMR01, FKINSPGMR10, FKINSPGMR11, FKINSPGMR20, and FKINSPGMR21, where the first digit in the name of the function is: 0 if neither KPSOL nor KPRECO routines are provided; 1 if only the preconditioner solve routine (KPSOL) is provided; and 2 if both the preconditioner solve (KPSOL) and setup (KPRECO) routines are provided. The second digit is: 0 if a function FATIMES is not provided; and 1 if a function FATIMES is provided.

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# NEQ (input) INTEGER

Number of equations (and unknowns) in the algebraic system.

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#### OPTIN (input) LOGICAL

Flag indicating whether optional inputs from the user in the arrays IOPT and ROPT are to be used.

Pass FALSE to ignore all optional inputs and TRUE to use all optional inputs that are present. Either choice does NOT affect outputs in other positions of IOPT or ROPT.

C C

#### MAXL (input) INTEGER

Maximum Krylov dimension for the Linear Solver. Pass 0 to use the default value MIN(Neq, 10).

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## MAXLRST (input) INTEGER

Maximum number of linear solver restarts allowed. Values outside the range 0 to 2\*NEQ/MAXL will be restricted to that range. 0, meaning no restarts, is a safe starting value.

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#### MSBPRE (input) INTEGER

Maximum number of steps calling the solver KPSOL without calling the preconditioner KPRECO. (The default is 10).

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C C C C	UU	<pre>(input/output) DOUBLE PRECISION array, dimension (NEQ) On entry, UU is the initial guess. On exit, if no errors ocurr, UU is the solution of the system KFUN(UU) = 0.</pre>	
C C C C C C	USCALE	(input) DOUBLE PRECISION array, dimension (NEQ) Array of diagonal elements of the scaling matrix for UU. The elements of USCALE must be positive values. The scaling matrix USCALE should be chosen so that USCALE * UU (as a matrix multiplication) should have all its components with roughly the same magnitude when UU is close to a root of KFUN.	
C C C C C C	FSCALE	(input) DOUBLE PRECISION array, dimension (NEQ) Array of diagonal elements of the scaling matrix for KFUN. The elements of FSCALE must be positive values. The scaling matrix FSCALE should be chosen so that FSCALE * KFUN(UU) (as a matrix multiplication) should have all its components with roughly the same magnitude when UU is NOT too near a root of KFUN.	
C C C C C C C	CONSTR	<pre>(input) DOUBLE PRECISION array, dimension (NEQ) Constraints on UU. A positive value in CONSTR(I) implies that the Ith component of UU is to be constrained &gt; 0. A negative value in CONSTR(I) implies that the Ith component of UU is to be constrained &lt; 0. A zero value in CONSTR(I) implies there is no constraint on UU(I).</pre>	
C C C C	IOPT	(input/output) INTEGER array, dimension (40) Array of optional integer inputs and outputs. If OPTIN is TRUE, the user should preset to 0 those locations for which default values are to be used. See Optional Inputs and Outputs, below.	
C C C C	ROPT	(input/output) DOUBLE PRECISION array, dimension (40) Array of optional double precision inputs and outputs. If OPTIN is TRUE, the user should preset to 0 those locations for which default values are to be used. See Optional Inputs and Outputs, below.	
C C	Tolerances		
C	TOL1	DOUBLE PRECISION	

	Stopping tolerance on maxnorm( $FSCALE * KFUN(UU)$ ).
	If TOL1 is input as 0., then a default value of
	(uround) to the $1/3$ power will be used. uround is the
	unit roundoff for the machine in use for the calculation
TOL2	DOUBLE PRECISION
	Stopping tolerance on the maximum scaled step
	UU(K) - UU(K-1).
	If TOL2 is input as 0., then a default value of (uround)
	to the 2/3 power will be used. uround is the unit
	roundoff for the machine in use for the calculation.
Error I	ndicator
INFO	(output) INTEGER
	See Termination Codes below.
Termina	tion Codes
(Note:	in this documentation we use named constants for
<u>.</u>	integer constant values. To see the values of these
	see Named constants below.)
~ j ~ ~ ~ ~ ~	
The term	mination values KINS_***** are now given. These are the
	of the INFO argument.
varaob .	or one into arbanone.
SUCCESS	: means maxnorm(FSCALE*KFUN(UU) <= TOL1, where
рооопрр	maxnorm() is the maximum norm function N_VMaxNorm.
	Therefore, UU is probably an approximate root of KFUN.
	ATON.
TMTTTAT	CHECC OV. moons the initial successful has been found
TNT11AL	_GUESS_OK: means the initial guess UU has been found
	to already satisfy the system to the desired
	accuracy. No calculation was performed other
	than testing UU.
STEP_LT	
	_STPTOL: means the scaled distance between the last
	_STPTOL: means the scaled distance between the last two steps is less than TOL2. UU may be an
	two steps is less than TOL2. UU may be an
	two steps is less than TOL2. UU may be an approximate root of KFUN, but it is also possible
	two steps is less than TOL2. UU may be an approximate root of KFUN, but it is also possible that the algorithm is making very slow progress
	two steps is less than TOL2. UU may be an approximate root of KFUN, but it is also possible that the algorithm is making very slow progress and is not near a root or that TOL2 is too

LNSRCH\_NONCONV: means the LineSearch module failed to reduce norm(KFUN) sufficiently on the last global step. Either UU is close to a root of F and no more accuracy is possible, or the finite-difference approximation to J\*v is inaccurate, or TOL2 is too large. Check the outputs NCFL and NNI: if NCFL is close to NNI, it may be the case that the Krylov iteration is converging very slowly. In this case, the user may want to use preconditioning and/or increase the MAXL argument (that is, increase the max dimension of the Krylov subspace) by setting MAXL to nonzero (thus not using the default value of KINSPGMR\_MAXL) or if MAXL is being

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MAXITER\_REACHED: means that the maximum allowable number of nonlinear iterations has been reached. This is by default 200, but may be changed through optional input IOPT(MXITER).

set, increase its value.

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MXNEWT\_5X\_EXCEEDED: means 5 consecutive steps of length mxnewt

(maximum Newton stepsize limit) have been taken.

Either norm(F) asymptotes from above to a finite
value in some direction, or mxnewt is too small.

Mxnewt is computed internally (by default) as
mxnewt = 1000\*max(norm(USCALE\*UUO),1), where

UUO is the initial guess for UU, and norm() is
the Euclidean norm. Mxnewt can be set by the
user through optional input ROPT(MXNEWTSTEP).

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LINESEARCH\_BCFAIL: means that more than the allowed maximum number of failures (MXNBCF) occurred when trying to satisfy the beta condition in the linesearch algorithm. It is likely that the iteration is making poor progress.

 ${\tt C}$ 

KRYLOV\_FAILURE: means there was a failure of the Krylov iteration process to converge.

 ${\tt C}$ 

PRECONDSET\_FAILURE: means there was a nonrecoverable error in PrecondSet causing the iteration to halt.

C C

PRECONDSOLVE\_FAILURE: means there was a nonrecoverable error in PrecondSolve causing the iteration to halt.

C

NO_MEM: the KINSol memory pointer received was NULL.
INPUT_ERROR: one or more input parameters or arrays was in
error. See the program output for further info.
orior, boo one program eaches for rationer into.
LSOLV_NO_MEM: The linear solver memory pointer (lmem) was
received as NULL. The return value from the linear
solver needs to be checked and the cause found.
borver needs to be checked and the cause round.
Optional inputs and outputs
opotonal inputs and outputs
(Note: in this documentation we use named constants for
certain integer constant values. To see the values of these
symbols see Named constants below.)
bymbolb boo named comboanes below.
The user should declare two arrays for optional input and
output, an IOPT array for optional integer input and output
and an ROPT array for optional real input and output. These
arrays should both be of size OPT_SIZE.
So the user's declaration should look like:
bo the user b decidration should look like.
INTEGER IOPT(OPT_SIZE)
DOUBLE PRECISION ROPT(OPT_SIZE)
DOODEE TRECIBION ROLL(OLI_SIZE)
The following definitions are indices into the IOPT and ROPT
arrays. A brief description of the contents of these positions
follows.
TOTTOWD.
IOPT(PRINTFL) (input) Allows user to select from 4 levels
of output.
=0 no statistics printed (DEFAULT)
=1 output the nonlinear iteration count, the
scaled norm of KFUN(UU), and number of
KFUN calls.
=2 same as 1 with the addition of global
strategy statistics:
f1 = 0.5*norm(FSCALE*KFUN(UU))**2 and $f1nov = 0.5*norm(FSCALE*KFUN(unov))**2$
f1new = 0.5*norm(FSCALE*KFUN(unew))**2.
=3 same as 2 with the addition of further
Krylov iteration statistics.
TODE/MYTEED) (' ') M ' 33 13 13 13 13 13 13 13 13 13 13 13 13
<pre>IOPT(MXITER) (input) Maximum allowable number of nonlinear</pre>

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С
                      iterations. The default is MXITER_DEFAULT.
C
C
      IOPT(PRECOND_NO_INIT) (input) Set to 1 to prevent the initial
C
                       call to the routine KPRECO upon a given
C
                       call to KINSol. Set to 0 or leave unset to
C
                       force the initial call to KPRECO.
C
                       Use the choice of 1 only after beginning the
C
                       first of a series of calls with a 0 value.
C
                       If a value other than 0 or 1 is encountered,
                       the default, 0, is set in this element of
C
C
                       IOPT and thus the routine KPRECO will
C
                       be called upon every call to KINSol, unless
C
                       IOPT(PRECOND_NO_INIT) is changed by the user.
C
C
      IOPT(ETACHOICE) (input) A flag indicating which of three
C
                       methods to use for computing eta, the
С
                       coefficient in the linear solver
C
                       convergence tolerance eps, given by
C
                         eps = (eta+u_round)*norm(KFUN(UU)).
                       Here, all norms are the scaled L2 norm.
C
C
                       The linear solver attempts to produce a step
C
                       p such that norm(KFUN(UU)+J(UU)*p) \le eps.
C
                       Two of the methods for computing eta
                       calculate a value based on the convergence
C
C
                       process in the routine KINForcingTerm.
C
                       The third method does not require
C
                       calculation; a constant eta is selected.
C
C
                       The default if IOPT(ETACHOICE) is not
C
                       specified is ETACHOICE1, (see below).
C
C
                       The allowed values (methods) are:
C
                  ETACONSTANT constant eta, default of 0.1 or user
C
                     supplied choice, for which see ROPT(ETACONST),
C
                  ETACHOICE1 (default) which uses choice 1 of
C
C
                     Eisenstat and Walker's paper of SIAM J. Sci.
C
                     Comput.,17 (1996), pp 16-32 wherein eta is:
C
                              eta(k) =
C
      ABS( norm(KFUN(UU(k))) - norm(KFUN(UU(k-1))+J(UU(k-1))*p))
C
                          / norm(KFUN(UU(k-1))),
C
С
                  ETACHOICE2
                               which uses choice 2 of
С
                     Eisenstat and Walker wherein eta is:
C
                     eta(k) = egamma *
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C
                 (norm(KFUN(UU(k))) / norm(KFUN(u(k-1))))ealpha
C
C
                     egamma and ealpha for choice 2, both required,
С
                     are from either defaults (egamma = 0.9,
С
                     ealpha = 2) or from user input,
C
                     see ROPT(ETAALPHA) and ROPT(ETAGAMMA), below.
C
C
                     For eta(k) determined by either Choice 1 or
C
                     Choice 2, a value eta_safe is determined, and
C
                     the safeguard
                                      eta(k) <- max(eta_safe,eta(k))</pre>
С
                     is applied to prevent eta(k) from becoming too
С
                     small too quickly.
C
                      For Choice 1,
                        eta_safe = eta(k-1)^((1.+sqrt(5.))/2.)
C
C
                      for Choice 2,
               and
С
                        eta_safe = egamma*eta(k-1)^ealpha.
С
                      (These safeguards are turned off if they drop
C
                     below 0.1 . Also, eta is never allowed to be
C
                     less than eta_min = 1.e-4).
C
С
      IOPT(NO_MIN_EPS) (input) Set to 1 or greater to remove
С
                     protection agains eps becoming too small.
C
                     This option is useful for debugging linear
C
                     and nonlinear solver interactions. Set to 0
C
                     for standard eps minimum value testing.
С
С
      IOPT(NNI)
                      (output) Total number of nonlinear iterations.
С
С
      IOPT(NFE)
                      (output) Total number of calls to the user-
C
                      supplied system function KFUN.
C
C
      IOPT(NBCF)
                      (output) Total number of times the beta
C
                      condition could not be met in the linesearch
C
                      algorithm. The nonlinear iteration is halted
C
                      if this value ever exceeds MXNBCF (10).
C
C
      IOPT(NBKTRK)
                      (output) Total number of backtracks in the
С
                      linesearch algorithm.
C
С
      IOPT(SPGMR_NLI) (output) Number of linear iterations.
С
C
      IOPT(SPGMR_NPE) (output) Number of preconditioner evaluations.
С
С
      IOPT(SPGMR_NPS) (output) Number of calls made to user's psolve
С
                      function.
```

C C C C C C	<pre>IOPT(SPGMR_NCFL) (output) Number of linear convergence failures.</pre>					
	ROPT(MXNEWTSTEP) (input) Maximum allowable length of a Newton step. The default value is calculated from 1000*max(norm(USCALE*UU(0),norm(USCALE)).					
C C	ROPT(RELFUNC) (input) Relative error in computing KFUN(UU) if known. Default is the machine epsilon.					
C C C C	ROPT(RELU) (input) A scalar constraint which restricts the update of UU to del(UU)/UU < ROPT(RELU)  The default is no constraint on the relative step in UU.					
C C C	ROPT(ETAGAMMA) (input) The coefficient egamma in the eta computation. See routine KINForcingTerm (SEE IOPT(ETACHOICE) above for additional info).					
C C C C C C	ROPT(ETAALPHA) (input) The coefficient ealpha in the eta computation. See routine KINForcingTerm (SEE IOPT(ETACHOICE) above for additional info).					
	ROPT(ETACONST) (input) A user specified constant value for eta, used in lieu of that computed by routine KINForcingTerm  (SEE IOPT(ETACHOICE) above for additional info).					
C C C	ROPT(FNORM) (output) The scaled norm at a given iteration: norm(FSCALE(KFUN(UU)).					
C C C	ROPT(STEPL) (output) Last step length in the global strategy routine:  KINLineSearch or KINInexactNewton.					
C						
C C	User-defined routines					
C C	In order to use this routine, some user-defined routines have to be provided. One of them is required, while the others are optional. These routines are described next.					
C C	KFUN Required					

C C SUBROUTINE KFUN (NEQ, UU, FVAL) C INTEGER NEQ C DOUBLE PRECISION UU(NEQ), FVAL(NEQ) C C **PURPOSE** C C Evaluates the KFUN function which defines the system C to be solved: C KFUN(UU)=0C С ARGUMENTS C C NEO C (input) INTEGER C Number of equations (and unknowns) in the algebraic С system C UU C C (input) DOUBLE PRECISION array, dimension (NEQ) C independent variable vector C C FVAL C (output) DOUBLE PRECISION array, dimension (NEQ) C Result of KFUN(UU) C C KPRECO Optional C C SUBROUTINE KPRECO (NEQ, UU, USCALE, FVAL, FSCALE, C VTEMP1, VTEMP2, UROUND, NFE, IER) C INTEGER NEQ, NFE, IER C DOUBLE PRECISION UROUND C DOUBLE PRECISION UU(NEQ), USCALE(NEQ), FVAL(NEQ), C FSCALE(NEQ), VTEMP1(NEQ), VTEMP2(NEQ) C C **PURPOSE** C C The user-supplied preconditioner setup function KPRECO and C the user-supplied preconditioner solve function KPSOL C together must define the right preconditioner matrix P С chosen so as to provide an easier system for the Krylov C solver to solve. KPRECO is called to provide any matrix С data required by the subsequent call(s) to KPSOL. The C data is expected to be stored in variables within a C COMMON block and the definition of those variables is up

to the user. More specifically, the user-supplied preconditioner setup function KPRECO is to evaluate and preprocess any Jacobian-related data needed by the preconditioner solve function KPSOL. This might include forming a crude approximate Jacobian, and performing an LU factorization on the resulting approximation to J. This function will not be called in advance of every call to KPSOL, but instead will be called only as often as necessary to achieve convergence within the Newton iteration in KINSol. If the KPSOL function needs no preparation, the KPRECO function need not be provided.

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 ${\tt KPRECO}$  should not modify the contents of the arrays  ${\tt UU}$  or  ${\tt FVAL}$  as those arrays are used elsewhere in the iteration process.

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Each call to the KPRECO function is preceded by a call to the system function KFUN. Thus the KPRECO function can use any auxiliary data that is computed by the KFUN function and saved in a way accessible to KPRECO.

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The two scaling arrays, FSCALE and USCALE, and unit roundoff UROUND are provided to the KPRECO function for possible use in approximating Jacobian data, e.g. by difference quotients. These arrays should also not be altered

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## ARGUMENTS

C C

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### NEQ

(input) INTEGER

Number of equations (and unknowns) in the algebraic system.

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(input) DOUBLE PRECISION array, dimension (NEQ) Independent variable vector.

C C C

#### USCALE

(input) DOUBLE PRECISION array, dimension (NEQ) See USCALE above.

C

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FVAL

(input) DOUBLE PRECISION array, dimension (NEQ)
Current value of KFUN(UU).

С		
С		FSCALE
С		(input) DOUBLE PRECISION array, dimension (NEQ)
С		See FSCALE above.
С		
С		VTEMP1
C		DOUBLE PRECISION array, dimension (NEQ)
C		Temporary work array.
C		
C		VTEMP2
C		DOUBLE PRECISION array, dimension (NEQ)
C		Temporary work array.
C		remporary work array.
C		UROUND
C		(input) DOUBLE PRECISION
C		Machine unit roundoff.
		Machine unit foundoif.
C		NEE
C		NFE
C		(input/output) INTEGER
C		Number of calls to KFUN made by the package. The KPRECO
C		routine should update this counter by adding on the
C		number of KFUN calls made in order to approximate the
C		Jacobian, if any. For example, if the routine calls
C		KFUN a total of W times, then the update is
C		NFE = NFE + W.
C		
C		IER
C		(output) INTEGER
C		Error indicator.
C		0 if successful,
C		1 if failure, in which case KINSOL stops.
C		
C	KPSOL	Optional
C		
C		SUBROUTINE KPSOL (NEQ, UU, USCALE, FVAL, FSCALE, VTEM,
C		FTEM, UROUND, NFE, IER)
C		INTEGER NEQ, NFE, IER
C		DOUBLE PRECISION UU(NEQ), USCALE(NEQ), FVAL(NEQ),
C		<pre>FSCALE(NEQ), VTEM(NEQ), FTEM(NEQ)</pre>
C		
C		PURPOSE
C		
C		The user-supplied preconditioner solve function KPSOL
C		is to solve a linear system $P x = r$ in which the matrix
C		P is the (right) preconditioner matrix P.

a	
C	WDGOT 1 11
C	KPSOL should not modify the contents of the iterate
C	array UU or the current function value array FVAL as
С	those are used elsewhere in the iteration process.
С	
C	ARGUMENTS
C	
C	NEQ
С	(input) INTEGER
С	Number of equations (and unknowns) in the algebraic
С	system.
C	- J
C	UU
C	(input) DOUBLE PRECISION array, dimension (NEQ)
	- · ·
C	Independent variable vector.
C	
С	USCALE
С	(input) DOUBLE PRECISION array, dimension (NEQ)
С	See USCALE above.
С	
C	FVAL
C	(input) DOUBLE PRECISION array, dimension (NEQ)
C	Current value of KFUN(UU).
С	
С	FSCALE
С	(input) DOUBLE PRECISION array, dimension (NEQ)
С	See FSCALE above.
C	
C	VTEM
C	(input/output) DOUBLE PRECISION array, dimension (NEQ)
C	On entry, holds the RHS vector r.
	·
C	On exit, holds the result x.
C	
С	FTEM
С	DOUBLE PRECISION array, dimension (NEQ)
С	Temporary work array.
С	
С	UROUND
C	(input) DOUBLE PRECISION
C	Machine unit roundoff.
C	
С	NFE
С	(input/output) INTEGER
C	Number of calls to KFUN made by the package. The KPRECO
C	routine should update this counter by adding on the
v	10 acting should aparage only counter by adding on the

```
C
              number of KFUN calls made in order to carry out the
C
              solution, if any. For example, if the routine calls
C
              KFUN a total of W times, then the update is
C
              NFE = NFE + W.
C
C
              IER
C
              (output) INTEGER
C
              Error indicator.
C
              0 if successful,
C
              1 if failure, in which case KINSOL stops.
C
С
      FATIMES Optional
C
C
              SUBROUTINE FATIMES (V, Z, NEWU, UU, IER)
C
                                 NEWU, IER
              INTEGER
C
              DOUBLE PRECISION V(:), Z(:), UU(:)
С
C
              PURPOSE
C
C
              The user-supplied A times V routine (optional) where
C
              A is the Jacobian matrix dF/du, or an approximation to
C
              it, and V is a given vector. This routine computes the
C
              product Z = J V.
C
              ARGUMENTS
C
C
C
C
              (input) DOUBLE PRECISION array, dimension (NEQ)
С
              Vector to be multiplied by J
C
              (preconditioned and unscaled as received).
C
C
              (output) DOUBLE PRECISION array, dimension (NEQ)
C
C
              Vector resulting from the application of J to V.
C
C
              NEW_UU
C
              (input) INTEGER
C
              Flag indicating whether or not the UU vector has been
C
              changed since the last call to this function (0 means
C
              FALSE, 1 TRUE).
С
              If this function computes and saves Jacobian data, then
C
              this computation can be skipped if NEW_UU = FALSE.
С
C
C
              (input) DOUBLE PRECISION array, dimension (NEQ)
```

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C
              Current iterate u.
C
C
              IER
С
              (output) INTEGER
С
              Error indicator.
C
              0 if successful,
C
              1 if failure, in which case KINSOL stops.
C
С
C
С
      Named constants
С
С
      Here we specify the value of the named integer constants used
С
      in this documentation. We use Fortran code for the specification,
С
      so that the user can copy and paste these lines in order to
С
      use the named constants in his/her programs.
С
CC
       KINSOL return values
CC
       Note that the value of these constants differ from those of
       the KINSOL package. This is due to the adaptation to the
CC
CC
       SLICOT standards.
С
       INTEGER KINS_NO_MEM, KINS_INPUT_ERROR, KINS_LSOLV_NO_MEM,
С
               KINS_SUCCESS, KINS_INITIAL_GUESS_OK, KINS_STEP_LT_STPTOL,
С
               KINS_LNSRCH_NONCONV, KINS_MAXITER_REACHED,
C
               KINS_MXNEWT_5X_EXCEEDED, KINS_LINESEARCH_BCFAIL,
С
               KINS_KRYLOV_FAILURE, KINS_PRECONDSET_FAILURE,
      &
С
               KINS_PRECONDSOLVE_FAILURE}
С
С
       PARAMETER (KINS_NO_MEM=101)
С
       PARAMETER (KINS_INPUT_ERROR=102)
С
       PARAMETER (KINS_LSOLV_NO_MEM=103)
С
       PARAMETER (KINS_SUCCESS=0)
С
       PARAMETER (KINS_INITIAL_GUESS_OK=2)
С
       PARAMETER (KINS_STEP_LT_STPTOL=3)
С
       PARAMETER (KINS_LNSRCH_NONCONV=4)
С
       PARAMETER (KINS_MAXITER_REACHED=5)
C
       PARAMETER (KINS_MXNEWT_5X_EXCEEDED=6)
С
       PARAMETER (KINS_LINESEARCH_BCFAIL=7)
С
       PARAMETER(KINS_KRYLOV_FAILURE = 8)
С
       PARAMETER (KINS_PRECONDSET_FAILURE=9)
С
       PARAMETER (KINS_PRECONDSOLVE_FAILURE=10)
C
С
       Size of IOPT, ROPT
С
       INTEGER OPT_SIZE
С
       PARAMETER (OPT_SIZE=40)
```

```
С
CC
       IOPT indices
С
       INTEGER PRINTFL, MXITER, PRECOND_NO_INIT, NNI ,NFE ,NBCF, NBKTRK,
С
               ETACHOICE, NO MIN EPS
С
       INTEGER SPGMR_NLI, SPGMR_NPE, SPGMR_NPS, SPGMR_NCFL
С
С
       PARAMETER (PRINTFL=1)
С
       PARAMETER (MXITER=2)
С
       PARAMETER (PRECOND_NO_INIT=3)
С
       PARAMETER(NNI=4)
С
       PARAMETER (NFE=5)
С
       PARAMETER (NBCF=6)
С
       PARAMETER (NBKTRK=7)
С
       PARAMETER (ETACHOICE=8)
С
       PARAMETER (NO_MIN_EPS=9)
С
       PARAMETER (SPGMR_NLI=11)
С
       PARAMETER (SPGMR_NPE=12)
С
       PARAMETER (SPGMR_NPS=13)
С
       PARAMETER (SPGMR_NCFL=14)
С
CC
       ROPT indices
       INTEGER MXNEWTSTEP, RELFUNC, RELU, FNORM, STEPL,
С
С
               ETACONST, ETAGAMMA, ETAALPHA
С
С
       PARAMETER (MXNEWTSTEP=1)
С
       PARAMETER (RELFUNC=2)
С
       PARAMETER (RELU=3)
С
       PARAMETER (FNORM=4)
С
       PARAMETER (STEPL=5)
С
       PARAMETER (ETACONST=6)
С
       PARAMETER (ETAGAMMA=7)
С
       PARAMETER (ETAALPHA=8)
С
CC
       Values for IOPT(ETACHOICE)
С
       INTEGER ETACHOICE1, ETACHOICE2, ETACONSTANT
С
С
       PARAMETER (ETACHOICE1=0)
С
       PARAMETER (ETACHOICE2=1)
С
       PARAMETER (ETACONSTANT=2)
С
С
C
С
      METHOD
С
      KINSOL (Krylov Inexact Newton SOLver) is a general purpose
```

```
С
      solver for nonlinear systems of equations. Its most notable
С
      feature is that it uses Krylov Inexact Newton techniques in the
С
      system's approximate solution.
С
      The Newton method used results in the solution of linear systems
С
      of the form
С
                              J(u)*x = b
      where J(u) is the Jacobian of F at u. The solution of these
С
C
      systems by a Krylov method requires products of the form J(u)*v,
С
      which are approximated by a difference quotient of the form
C
                            F(u+sigma*v)-F(u)
С
С
                                sigma
C
      Thus, the Jacobian need not be formed explicitly.
С
С
      REFERENCES
С
С
      [1] Allan G. Taylor and Alan C. Hindmarsh, "User Documentation
C
          for KINSOL, a Nonlinear Solver for Sequential and Parallel
C
          Computers", Center for Applied Scientific Computing, L-561,
C
          LLNL, Livermore, CA 94551.
С
С
      NUMERICAL ASPECTS
С
С
      CONTRIBUTOR
C
С
      Fernando Alvarruiz, Vicente Hernandez
С
      Universidad Politecnica de Valencia, Spain
С
С
      REVISIONS
С
С
      KEYWORDS
С
С
      Newton Method, Krylov methods
C
      IMPLICIT NONE
      LOGICAL OPTIN
      INTEGER GSTRAT, NEQ, MAXL, MAXLRST, MSBPRE, INFO
      INTEGER IOPT(40)
      DOUBLE PRECISION TOL1, TOL2
      DOUBLE PRECISION UU(NEQ), USCALE(NEQ), FSCALE(NEQ),
              CONSTR(NEQ), ROPT(40)
      EXTERNAL LINSU
```

С

Local variables

```
Executable statements
      IF (OPTIN) THEN
         IOPTIN = 1
      ELSE
         IOPTIN = 0
      ENDIF
      CALL FSKINMALLOC (NEQ, INFO)
      IF (INFO .NE. O) THEN
         INFO = 101
         RETURN
      ENDIF
С
      Linear solver setup
      CALL LINSU (MAXL, MAXLRST, MSBPRE)
      CALL FKINSOL (NEQ, UU, GSTRAT, USCALE, FSCALE, TOL1,
                    TOL2, CONSTR, OPTIN, IOPT, ROPT, INFO)
      CALL FKINFREE
С
      Transform the return code to SLICOT standards
      IF (INFO .EQ. 1) THEN
         INFO = O
      ENDIF
      IF (INFO .LT. O) THEN
         INFO=100-INFO
      ENDIF
      RETURN
C *** Last line of KINSOL ***
      END
```

INTEGER IOPTIN

# B Documentation of the MATLAB interface for KINSOL

We include here the source code of the two m-files for the KINSOL interface (kinsol.m and kinsoptions.m). This is included because the source code contains detailed comments describing the interface.

### B.1 The function kinsol

```
% kinsol - Krylov Inexact Newton Solver
% function [uu, kinout] = kinsol (kfun, uu0, opts)
%
%
      PURPOSE
%
%
      To solve a nonlinear system of equations F(u)=0, where F(u) is
%
%
      a nonlinear function from {\tt R} to {\tt R} , using Krylov Inexact Newton
%
      techniques.
%
%
      INPUT ARGUMENTS
%
%
      kfun
              String
%
              Name of a matlab .m file that contains a function with
%
              the following form:
%
%
              function fval = kfun(uu)
%
%
              This function evaluates the function F at uu, i.e. F(uu).
%
%
              uu
%
              Real vector.
%
              Independent variable vector.
%
%
              fval
%
              Real vector.
%
              Result of F(uu).
%
%
              Real vector
      uu0
%
              Initial guess.
%
%
      opts
              Structure of options (type help kinoptions for a description
%
              of the options).
%
%
      OUTPUT ARGUMENTS
%
%
      kinout Structure of output information about the solution
%
              of the problem. The fields of this structure are
%
              presented next.
%
%
         NonLinIters (output) Total number of nonlinear iterations.
%
%
         NumFuncEvals (output) Total number of calls to the user-
```

% supplied system function kfun. % % NumBetaCondFail (output) Total number of times the beta % condition could not be met in the linesearch % algorithm. The nonlinear iteration is halted % if this value ever exceeds MXNBCF (10). % % NumBacktracks (output) Total number of backtracks in the % linesearch algorithm. % % (output) The scaled norm at a given iteration: Fnorm % norm(fscale(kfun(uu)). % % StepLength (output) Last step length in the global % strategy routine: % KINLineSearch or KINInexactNewton. % % LinIters (output) Number of linear iterations. % % NumPrecEvals (output) Number of preconditioner evaluations. % % NumPSolve (output) Number of calls made to user's % PrecondSolveFunc function. % % NumLinConvFails (output) Number of linear convergence failures. % % METHOD % % KINSOL (Krylov Inexact Newton SOLver) is a general purpose % solver for nonlinear systems of equations. Its most notable % feature is that it uses Krylov Inexact Newton techniques in the % system's approximate solution. % The Newton method used results in the solution of linear systems % of the form % J(u)\*x = b% where J(u) is the Jacobian of F at u. The solution of these % systems by a Krylov method requires products of the form J(u)\*v, % which are approximated by a difference quotient of the form % F(u+sigma\*v)-F(u)% % sigma % Thus, the Jacobian need not be formed explicitly. % % REFERENCES %

```
%
      [1] Allan G. Taylor and Alan C. Hindmarsh, "User Documentation
          for KINSOL, a Nonlinear Solver for Sequential and Parallel
%
%
          Computers", Center for Applied Scientific Computing, L-561,
%
          LLNL, Livermore, CA 94551.
%
%
      NUMERICAL ASPECTS
%
%
      CONTRIBUTOR
%
%
      Fernando Alvarruiz, Vicente Hernandez
%
      Universidad Politecnica de Valencia, Spain
%
%
      REVISIONS
%
%
      KEYWORDS
%
%
      Newton Method, Krylov methods
%
function [uu, kinout] = kinsol (kfun, uu0, opts)
n = length(uu0);
iopt = zeros(1,40);
ropt = zeros(1,40);
% Set default values for empty fields of opt
if isempty(opts.GlobalStrategy)
    opts.GlobalStrategy = 'in';
end
if isempty(opts.Constr)
    opts.Constr = zeros(1,n);
end
if isempty(opts.Uscale)
    opts.Uscale = ones(1,n);
end
if isempty(opts.Fscale)
    opts.Fscale = ones(1,n);
if isempty(opts.FNormTol)
    opts.FNormTol = 0;
end
if isempty(opts.ScStepTol)
    opts.ScStepTol = 0;
end
```

```
if isempty(opts.MaxNewtStep)
    opts.MaxNewtStep = 0;
end
ropt(1) = opts.MaxNewtStep;
if isempty(opts.Verbosity)
    opts.Verbosity = 0;
end
iopt(1) = opts. Verbosity;
if isempty(opts.MaxIter)
    opts.MaxIter = 0;
end
iopt(2) = opts.MaxIter;
if isempty(opts.RelFunc)
    opts.RelFunc = 0;
end
ropt(2) = opts.RelFunc;
if isempty(opts.RelU)
    opts.RelU = 0;
end
ropt(3) = opts.RelU;
if isempty(opts.PrecondSetFunc)
    opts.PrecondSetFunc = '';
end
if isempty(opts.PreconsSolveFunc)
    opts.PreconsSolveFunc = '';
if isempty(opts.UserATimesFunc)
    opts.UserATimesFunc = '';
end
if isempty(opts.MaxLinDim)
    opts.MaxLinDim = 0;
end
if isempty(opts.MaxLinRestarts)
    opts.MaxLinRestarts = 0;
end
if isempty(opts.MaxBeforePrecond)
    opts.MaxBeforePrecond = 0;
end
if isempty(opts.PrecondNoInit)
    opts.PrecondNoInit = 0;
end
iopt(3) = opts.PrecondNoInit;
if isempty(opts.EtaChoice)
    opts.EtaChoice = 0;
end
```

```
iopt(8)=opts.EtaChoice;
if isempty(opts.EtaGamma)
    opts.EtaGamma = 0;
end
ropt(7) = opts.EtaGamma;
if isempty(opts.EtaAlpha)
    opts.EtaAlpha = 0;
end
ropt(8) = opts.EtaAlpha;
if isempty(opts.EtaConst)
    opts.EtaConst = 0;
end
ropt(6) = opts.EtaConst;
% Call the MEX-File
[uu, iopt, ropt, info] = kinsol_gtw (opts.GlobalStrategy, kfun,...
    opts.PrecondSetFunc, opts.PreconsSolveFunc, opts.UserATimesFunc,...
    opts.MaxLinDim, opts.MaxLinRestarts, opts.MaxBeforePrecond,...
    uuO, opts.Uscale, opts.Fscale, opts.Constr, iopt, ropt,...
    opts.FNormTol, opts.ScStepTol);
% Form kinout structure
kinout.NonLinIters = iopt(4);
kinout.NumFuncEvals = iopt(5);
kinout.NumBetaCondFail = iopt(6);
kinout.NumBacktracks = iopt(7);
kinout.FNorm = ropt(4);
kinout.StepLength = ropt(5);
kinout.LinIters = iopt(10);
kinout.NumPrecEvals = iopt(11);
kinout.NumPSolve = iopt(12);
kinout.NumLinConvFails = iopt(13);
      The function kinsoptions
% KINSOL options
% function kopt = kinsoptions()
\% This function creates a default structure of options to be passed to the
% kinsol function.
% In these comments, the following notation is used:
% UU - independent variable for the nonlinear function.
% KFUN - nonlinear function.
```

```
% NEQ - number of equations/unknowns.
%
% The members of the options structure are:
% GlobalStrategy: String.
%
      Indicates the global strategy to apply the computed
%
      increment delta in the solution UU. Choices are:
      'in' - Inexact Newton (default).
%
%
      'ls' - Linesearch.
% Constr: Vector, dimension (NEQ).
%
      {\tt Constraints} \ {\tt on} \ {\tt UU}\,.
%
      A positive value in Constr(i) implies that the ith
%
      component of UU is to be constrained > 0.
%
      A negative value in Constr(i) implies that the ith
%
      component of UU is to be constrained < 0.
%
      A zero value in Constr(i) implies there is no constraint
%
      on UU(i).
%
% Uscale: Vector, dimension (NEQ).
%
      Array of diagonal elements of the scaling matrix for UU.
%
      The elements of Uscale must be positive values. The
%
      scaling matrix Uscale should be chosen so that
%
      Uscale * UU (as a matrix multiplication) should have all
%
      its components with roughly the same magnitude when UU is
%
      close to a root of KFUN.
%
% Fscale: Vector, dimension (NEQ).
%
      Array of diagonal elements of the scaling matrix for
%
      KFUN. The elements of Fscale must be positive values.
%
      The scaling matrix Fscale should be chosen so that
%
      Fscale * KFUN(UU) (as a matrix multiplication) should
%
      have all its components with roughly the same magnitude
%
      when UU is NOT too near a root of KFUN.
%
% FNormTol: Real.
%
      Stopping tolerance on maxnorm( Fscale * KFUN(UU) ).
%
      If FNormTol is input as O., then a default value of
%
      (uround) to the 1/3 power will be used. uround is the
%
      unit roundoff for the machine in use for the calculation.
%
% ScStepTol: Real.
%
      Stopping tolerance on the maximum scaled step
%
      UU(K) - UU(K-1).
%
      If ScStepTol is input as O., then a default value of (uround)
```

```
%
      to the 2/3 power will be used. uround is the unit
%
      roundoff for the machine in use for the calculation.
%
% MaxNewtStep: Real.
%
      Maximum allowable length of a Newton step. The default value
%
      is calculated from 1000*max(norm(uscale*UU(0),norm(uscale)).
%
% Verbosity: Integer.
%
      Allows user to select from 4 levels of output.
%
      =0 no statistics printed (DEFAULT).
%
      =1 output the nonlinear iteration count, the scaled norm of
%
         KFUN(UU), and number of KFUN calls.
%
      =2 same as 1 with the addition of global strategy statistics:
%
         f1 = 0.5*norm(Fscale*KFUN(UU))**2
%
         f1new = 0.5*norm(Fscale*KFUN(unew))**2 .
%
      =3 same as 2 with the addition of further Krylov iteration
%
         statistics.
%
% MaxIter: Integer.
%
      Maximum allowable number of nonlinear iterations. The default
%
      is 200.
%
% RelFunc: Real.
%
      Relative error in computing KFUN(UU) if known. Default is the
%
      machine epsilon.
%
% RelU: Real.
%
      A scalar constraint which restricts
%
      the update of UU to del(UU)/UU < RelU.
%
      The default is no constraint on the relative
%
      step in UU.
%
% PrecondSetFunc: String.
%
      Name of an m-file with the definition of the following function
%
%
      [NFE, IERR] = KPRECO(UU, USCALE, FVAL, FSCALE, KFUN, UROUND, NFE)
%
%
      where the name of the function, here KPRECO, must be the same as the
%
      name of the m-file. This function is described next.
%
%
      PURPOSE
%
%
      The user-supplied preconditioner setup function KPRECO and
%
      the user-supplied preconditioner solve function KPSOL (see below
%
      together must define the right preconditoner matrix P
```

chosen so as to provide an easier system for the Krylov solver to solve. KPRECO is called to provide any matrix data required by the subsequent call(s) to KPSOL. The data is expected to be stored in variables within a COMMON block and the definition of those variables is up to the user. More specifically, the user-supplied preconditioner setup function KPRECO is to evaluate and preprocess any Jacobian-related data needed by the preconditioner solve function KPSOL. This might include forming a crude approximate Jacobian, and performing an LU factorization on the resulting approximation to J. This function will not be called in advance of every call to KPSOL, but instead will be called only as often as necessary to achieve convergence within the Newton iteration in KINSol. If the KPSOL function needs no preparation, the KPRECO function need not be provided.

% % %

%

%

%

%

%

%

%

%

%

%

%

%

%

%

%

%

%

Each call to the KPRECO function is preceded by a call to the system function KFUN. Thus the KPRECO function can use any auxiliary data that is computed by the KFUN function and saved in a way accessible to KPRECO.

% % %

%

%

The two scaling arrays, FSCALE and USCALE, and unit roundoff UROUND are provided to the KPRECO function for possible use in approximating Jacobian data, e.g. by difference quotients.

% % %

### INPUT ARGUMENTS

% %

UU: Real array, dimension (NEQ). Independent variable vector.

% % %

USCALE: Real array, dimension(NEQ). See Uscale above.

% % %

FVAL: Real array, dimension (NEQ). Current value of KFUN(UU).

% % %

FSCALE: Real array, dimension(NEQ). See FSCALE above.

% %

KFUN: String.

% %

Name of an m-file defining the KFUN function.

% %

UROUND: Real.

```
%
      Machine unit roundoff.
%
%
      NFE: Integer.
%
      Number of calls to KFUN made by the package so far.
%
%
      OUTPUT ARGUMENTS
%
%
      NFE: Integer.
%
      NFE should be set to the result of updating NFE by adding to it the
%
      number of KFUN calls made in order to approximate the
%
      Jacobian, if any. For example, if the routine calls
%
      KFUN a total of W times, then the update is
%
      NFE = NFE + W.
%
%
      IER: Integer.
%
      Error indicator.
%
      0 if successful,
%
      1 if failure, in which case KINSOL stops.
%
% PrecondSolveFunc: String.
%
      Name of an m-file with the definition of the following function
%
%
      [X, NFE, IERR] = KPSOL(UU, USCALE, FVAL, FSCALE, R, KFUN, UROUND, NFE)
%
%
      where the name of the function, here KPSOL, must be the same as the
%
      name of the m-file. This function is described next.
%
%
      PURPOSE
%
%
      The user-supplied preconditioner solve function KPSOL
      is to solve a linear system P x = r in which the matrix
%
%
      P is the (right) preconditioner matrix P.
%
%
      ARGUMENTS
%
%
      UU: Real array, dimension (NEQ)
%
      Independent variable vector.
%
%
      USCALE: Real array, dimension(NEQ).
%
      See Uscale above.
%
%
      FVAL: Real array, dimension (NEQ).
%
      Current value of KFUN(UU).
%
%
      FSCALE: Real array, dimension(NEQ).
```

```
%
      See Fscale above.
%
%
      R: Real array, dimension (NEQ).
%
      RHS vector r.
%
      On exit, holds the result x.
%
%
      KFUN: String.
%
      Name of an m-file defining the KFUN function.
%
%
      UROUND: Real.
%
      Machine unit roundoff.
%
%
      NFE: Integer.
%
      Number of calls to KFUN made by the package so far.
%
%
      OUTPUT ARGUMENTS
%
%
      X: Real array, dimension (NEQ).
%
      The result x.
%
%
      NFE: Integer.
%
      NFE should be set to the result of updating NFE by adding to it the
%
      number of KFUN calls made in order to carry out the solution,
%
      if any. For example, if the routine calls KFUN a total of W times,
%
      then the update is
%
      NFE = NFE + W.
%
%
      IER: Integer.
%
      Error indicator.
%
      0 if successful,
%
      1 if failure, in which case KINSOL stops.
%
% UserATimesFunc: String.
%
      Name of an m-file with the definition of the following function
%
%
      [Z, IERR] = FATIMES(V, NEW_UU, UU)
%
%
      where the name of the function, here FATIMES, must be the same as the
%
      name of the m-file. This function is described next.
%
%
      PURPOSE
%
%
      The user-supplied A times v routine (optional) where
%
      A is the Jacobian matrix dF/du, or an approximation to
%
      it, and v is a given vector. This routine computes the
```

```
%
      product z = J*v.
%
%
      INPUT ARGUMENTS
%
%
      V: Real array, dimension (NEQ).
%
      Vector to be multiplied by J
%
      (preconditioned and unscaled as received).
%
%
      NEW_UU: Integer.
%
      Flag indicating whether or not the UU vector has been
%
      changed since the last call to this function (0 means
%
      FALSE, 1 TRUE).
%
      If this function computes and saves Jacobian data, then
%
      this computation can be skipped if NEW_UU = FALSE.
%
%
      UU: Real array, dimension (NEQ).
%
      Current iterate u.
%
%
      OUTPUT ARGUMENTS
%
%
      Z: Real array, dimension (NEQ).
%
      Vector resulting from the application of J to v.
%
%
      IER: Integer.
%
      Error indicator.
%
      0 if successful,
%
      1 if failure, in which case KINSOL stops.
%
% MaxLinDim: Integer.
%
      Maximum Krylov dimension. This is an
%
      optional input to the KINSpgmr solver. Pass 0 to
%
      use the default value MIN(NEQ, KINSPGMR_MAXL=10).
% MaxLinRestarts: Integer.
%
      Is the maximum number of linear solver restarts
%
      allowed. Values outside the range O to 2*NEQ/MaxLinDim
%
      will be restricted to that range. O, meaning no
%
      restarts, is a safe starting value.
% MaxBeforePrecond: Integer.
%
      Is the maximum number of steps calling the solver
%
      PrecondSolveFunc without calling the preconditioner
%
      PrecondSetFunc (the default is KINSPGMR_MSBPRE = 10).
% PrecondNoInit: Integer.
```

```
%
      Set to 1 to prevent the initial
%
      call to the routine PrecondSetFunc upon a given
%
      call to KINSol. Set to 0 or leave unset to
%
      force the initial call to PrecondSetFunc.
%
      Use the choice of 1 only after beginning the
%
      first of a series of calls with a 0 value.
%
      If a value other than 0 or 1 is encountered,
%
      the default, 0, is set in this option
%
      and thus the routine PrecondSetFunc will
%
      be called upon every call to KINSol, unless
%
      PrecondNoInit is changed by the user.
%
% EtaChoice: Integer.
%
      Flag indicating which of three
%
      methods to use for computing eta, the
%
      coefficient in the linear solver
%
      convergence tolerance eps
%
      eps = (eta+u_round)*norm(KFUN(UU)).
%
      Here, all norms are the scaled L2 norm.
%
      The linear solver attempts to produce a step
%
      p such that norm(KFUN(UU)+J(UU)*p) \le eps.
%
      Two of the methods for computing eta
%
      calculate a value based on the convergence
      process in the routine KINForcingTerm.
%
%
      The third method does not require
%
      calculation; a constant eta is selected.
%
%
      The default if EtaChoice is not
%
      specified is ETACHOICE1 (0), (see below).
%
%
      The allowed values (methods) are:
%
      ETACONSTANT (2) constant eta, default of 0.1 or user
%
         supplied choice, for which see EtaConst,
%
%
      ETACHOICE1 (0) [default] which uses choice 1 of
%
         Eisenstat and Walker's paper of SIAM J. Sci.
%
         Comput.,17 (1996), pp 16-32 wherein eta is:
%
                           eta(k) =
%
    ABS( norm(KFUN(UU(k))) - norm(KFUN(UU(k-1))+J(UU(k-1))*p) )
%
                        / norm(KFUN(UU(k-1))),
%
%
      ETACHOICE2 (1) which uses choice 2 of
         Eisenstat and Walker wherein eta is:
%
%
                      eta(k) = egamma *
%
          ( norm(KFUN(UU(k))) / norm(KFUN(u(k-1))) )^ealpha
```

```
%
%
         egamma and ealpha for choice 2, both required,
%
         are from either defaults (egamma = 0.9,
%
         ealpha = 2) or from user input,
%
         see EtaAlpha and EtaGamma, below.
%
%
      For eta(k) determined by either Choice 1 or
%
      Choice 2, a value eta_safe is determined, and
%
      the safeguard eta(k) <- max(eta_safe,eta(k))
%
      is applied to prevent eta(k) from becoming too
%
      small to quickly.
%
      For Choice 1,
%
             eta_safe = eta(k-1)^((1.+sqrt(5.))/2.)
%
      and for Choice 2,
%
               eta_safe = egamma*eta(k-1)^ealpha.
%
      (These safeguards are turned off if they drop
%
      below 0.1 . Also, eta is never allowed to be
%
      less than eta_min = 1.e-4).
%
% EtaGamma: Real.
%
      The coefficient egamma in the eta
%
      computation. See routine KINForcingTerm
%
      (see EtaChoice above for additional info).
%
% EtaAlpha: Real.
%
      The coefficient ealpha in the eta
%
      computation. See routine KINForcingTerm
%
      (see EtaChoice above for additional info).
% EtaConst: Real.
%
      A user specified constant value for
%
      eta, used in lieu of that computed by
%
      routine KINForcingTerm
%
      (see EtaChoice above for additional info).
%
function kopt = kinsoptions()
kopt = struct('GlobalStrategy',[],...
    'Constr',[],...
    'Uscale',[],...
    'Fscale',[],...
    'FNormTol',[],...
    'ScStepTol',[],...
    'MaxNewtStep',[],...
    'Verbosity',[],...
```

```
'MaxIter',[],...
'RelFunc',[],...
'RelU',[],...
'PrecondSetFunc',[],...
'PreconsSolveFunc',[],...
'UserATimesFunc',[],...
'MaxLinDim',[],...
'MaxLinRestarts',[],...
'MaxBeforePrecond',[],...
'PrecondNoInit',[],...
'EtaChoice',[],...
'EtaGamma',[],...
'EtaAlpha',[],...
'EtaConst',[]);
```