NAG Fortran Library Routine Document

D02RAF

Note: before using this routine, please read the Users' Note for your implementation to check the interpretation of *bold italicised* terms and other implementation-dependent details.

1 Purpose

D02RAF solves the two-point boundary-value problem with general boundary conditions for a system of ordinary differential equations, using a deferred correction technique and Newton iteration.

2 Specification

SUBROUTINE DO2RAF	(N, MNP, NP, NUMBEG, NUMMIX, TOL, INIT, X, Y, IY, ABT,
1	FCN, G, IJAC, JACOBF, JACOBG, DELEPS, JACEPS, JACGEP,
2	WORK, LWORK, IWORK, LIWORK, IFAIL)
INTEGER	N, MNP, NP, NUMBEG, NUMMIX, INIT, IY, IJAC, LWORK,
1	IWORK(LIWORK), LIWORK, IFAIL
double precision	TOL, X(MNP), Y(IY,MNP), ABT(N), DELEPS, WORK(LWORK)
EXTERNAL	FCN, G, JACOBF, JACOBG, JACEPS, JACGEP

3 Description

D02RAF solves a two-point boundary-value problem for a system of *n* ordinary differential equations in the interval (a, b) with b > a. The system is written in the form

$$y'_{i} = f_{i}(x, y_{1}, y_{2}, \dots, y_{n}), \quad i = 1, 2, \dots, n$$
 (1)

and the derivatives f_i are evaluated by a (sub)program FCN supplied by you. With the differential equations (1) must be given a system of n (nonlinear) boundary conditions

$$g_i(y(a), y(b)) = 0, \quad i = 1, 2, \dots, n,$$

where

$$y(x) = [y_1(x), y_2(x), \dots, y_n(x)]^{\mathrm{T}}.$$
(2)

The functions g_i are evaluated by a (sub)program G supplied by you. The solution is computed using a finite-difference technique with deferred correction allied to a Newton iteration to solve the finite-difference equations. The technique used is described fully in Pereyra (1979).

You must supply an absolute error tolerance and may also supply an initial mesh for the finite-difference equations and an initial approximate solution (alternatively a default mesh and approximation are used). The approximate solution is corrected using Newton iteration and deferred correction. Then, additional points are added to the mesh and the solution is recomputed with the aim of making the error everywhere less than your tolerance and of approximately equidistributing the error on the final mesh. The solution is returned on this final mesh.

If the solution is required at a few specific points then these should be included in the initial mesh. If, on the other hand, the solution is required at several specific points then you should use the interpolation routines provided in Chapter E01 if these points do not themselves form a convenient mesh.

The Newton iteration requires Jacobian matrices

$$\left(\frac{\partial f_i}{\partial y_j}\right), \left(\frac{\partial g_i}{\partial y_j(a)}\right) \text{ and } \left(\frac{\partial g_i}{\partial y_j(b)}\right).$$

These may be supplied by you through (sub)programs JACOBF for $\left(\frac{\partial f_i}{\partial y_j}\right)$ and JACOBG for the others.

Alternatively the Jacobians may be calculated by numerical differentiation using the algorithm described in Curtis *et al.* (1974).

For problems of the type (1) and (2) for which it is difficult to determine an initial approximation from which the Newton iteration will converge, a continuation facility is provided. You must set up a family of problems

$$y' = f(x, y, \epsilon), \quad g(y(a), y(b), \epsilon) = 0, \tag{3}$$

where $f = [f_1, f_2, \dots, f_n]^T$ etc., and where ϵ is a continuation parameter. The choice $\epsilon = 0$ must give a problem (3) which is easy to solve and $\epsilon = 1$ must define the problem whose solution is actually required. The routine solves a sequence of problems with ϵ values

$$0 = \epsilon_1 < \epsilon_2 < \dots < \epsilon_p = 1. \tag{4}$$

The number p and the values ϵ_i are chosen by the routine so that each problem can be solved using the solution of its predecessor as a starting approximation. Jacobians $\frac{\partial f}{\partial \epsilon}$ and $\frac{\partial g}{\partial \epsilon}$ are required and they may be supplied by you via the user-supplied (sub)programs JACEPS and JACGEP respectively or may be computed by numerical differentiation.

4 References

Curtis A R, Powell M J D and Reid J K (1974) On the estimation of sparse Jacobian matrices J. Inst. Maths. Applics. 13 117–119

Pereyra V (1979) PASVA3: An adaptive finite-difference Fortran program for first order nonlinear, ordinary boundary problems *Codes for Boundary Value Problems in Ordinary Differential Equations. Lecture Notes in Computer Science* (ed B Childs, M Scott, J W Daniel, E Denman and P Nelson) **76** Springer–Verlag

5 **Parameters**

1: N – INTEGER

On entry: n, the number of differential equations.

Constraint: N > 0.

2: MNP – INTEGER

On entry: MNP must be set to the maximum permitted number of points in the finite-difference mesh. If LWORK or LIWORK are too small then internally MNP will be replaced by the maximum permitted by these values. (A warning message will be output if on entry IFAIL is set to obtain monitoring information.)

Constraint: MNP \geq 32.

3: NP – INTEGER

On entry: must be set to the number of points to be used in the initial mesh.

Constraint: $4 \leq NP \leq MNP$.

On exit: the number of points in the final mesh.

4: NUMBEG – INTEGER

On entry: the number of left-hand boundary conditions (that is the number involving y(a) only). Constraint: $0 \le \text{NUMBEG} < \text{N}$.

Input

Input

Input

Input/Output

5: NUMMIX - INTEGER

On entry: the number of coupled boundary conditions (that is the number involving both y(a) and y(b)).

Constraint: $0 \leq \text{NUMMIX} \leq \text{N} - \text{NUMBEG}$.

TOL – double precision 6:

On entry: a positive absolute error tolerance. If

is the final mesh, $z_i(x_i)$ is the *j*th component of the approximate solution at x_i , and $y_i(x)$ is the *j*th component of the true solution of (1) and (2), then, except in extreme circumstances, it is expected that

 $a = x_1 < x_2 < \cdots < x_{\rm NP} = b$

$$|z_j(x_i) - y_j(x_i)| \le \text{TOL}, \quad i = 1, 2, \dots, \text{NP}; j = 1, 2, \dots, n.$$
 (5)

Constraint: TOL > 0.0.

INIT - INTEGER 7:

On entry: indicates whether you wish to supply an initial mesh and approximate solution $(INIT \neq 0)$ or whether default values are to be used, (INIT = 0).

X(MNP) - double precision array 8:

On entry: you must set X(1) = a and X(NP) = b. If INIT = 0 on entry a default equispaced mesh will be used, otherwise you must specify a mesh by setting $X(i) = x_i$, for i = 2, 3, ... NP - 1.

Constraints:

if INIT = 0, X(1) < X(NP); if $INIT \neq 0$, $X(1) < X(2) < \cdots < X(NP)$.

On exit: $X(1), X(2), \ldots, X(NP)$ define the final mesh (with the returned value of NP) and X(1) = aand X(NP) = b.

Y(IY,MNP) – *double precision* array 9:

On entry: if INIT = 0, then Y need not be set.

If INIT $\neq 0$, then the array Y must contain an initial approximation to the solution such that Y(j, i)contains an approximation to

$$y_i(x_i), \quad i = 1, 2, \dots, \text{NP}; j = 1, 2, \dots, n.$$

On exit: the approximate solution $z_i(x_i)$ satisfying (5) on the final mesh, that is

 $Y(j,i) = z_i(x_i), \quad i = 1, 2, ..., NP; j = 1, 2, ..., n,$

where NP is the number of points in the final mesh. If an error has occurred then Y contains the latest approximation to the solution. The remaining columns of Y are not used.

10: IY - INTEGER

> On entry: the first dimension of the array Y as declared in the (sub)program from which D02RAF is called.

Constraint: $IY \ge N$.

ABT(N) - double precision array 11:

On exit: ABT(i), for i = 1, 2, ..., n, holds the largest estimated error (in magnitude) of the *i*th component of the solution over all mesh points.

Input

Input

Input/Output

Output

Input

Input/Output

Input

12: FCN – SUBROUTINE, supplied by the user.

External Procedure

FCN must evaluate the functions f_i (i.e., the derivatives y'_i) at a general point x for a given value of ϵ , the continuation parameter (see Section 3).

Its specification is:

SUBROUTINE FCN (X, EPS, Y, F, N) INTEGER Ν X, EPS, Y(N), F(N)double precision X - double precision1: Input On entry: the value of the argument x. 2: **EPS** – *double precision* Input On entry: ϵ , the value of the continuation parameter. This is 1 if continuation is not being used. 3: Y(N) - double precision array Input On entry: the value of the argument y_i , for i = 1, 2, ..., n. F(N) - double precision array 4: Output On exit: the values of f_i , for i = 1, 2, ..., n. 5: N - INTEGER Input On entry: the number of equations.

FCN must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

13: G - SUBROUTINE, supplied by the user.

External Procedure

G must evaluate the boundary conditions in equation (3) and place them in the array BC. Its specification is:

SUBROUTINE G (EPS, YA, YB, BC, N) INTEGER double precision EPS, YA(N), YB(N), BC(N) EPS – double precision 1: Input On entry: ϵ , the value of the continuation parameter. This is 1 if continuation is not being used. 2: YA(N) - double precision array Input On entry: the value $y_i(a)$, for i = 1, 2, ..., n. YB(N) - double precision array 3: Input On entry: the value $y_i(b)$, for i = 1, 2, ..., n. BC(N) - double precision array Output 4: On exit: the values $g_i(y(a), y(b), \epsilon)$, for i = 1, 2, ..., n. These must be ordered as follows: (i) first, the conditions involving only y(a) (see NUMBEG description above);

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- (ii) next, the NUMMIX coupled conditions involving both y(a) and y(b) (see NUMMIX); and,
 (iii) finally, the conditions involving only y(b) (N NUMBEG NUMMIX).
- 5: N INTEGER
 - On entry: n, the number of equations.

G must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

14: IJAC – INTEGER

On entry: indicates whether or not you are supplying Jacobian evaluation routines.

IJAC $\neq 0$

You must supply (sub)programs JACOBF and JACOBG and also, when continuation is used, (sub)programs JACEPS and JACGEP.

IJAC = 0

Numerical differentiation is used to calculate the Jacobian and the routines D02GAZ, D02GAY, D02GAZ and D02GAX respectively may be used as the dummy parameters.

15: JACOBF - SUBROUTINE, supplied by the user.

JACOBF must evaluate the Jacobian $\left(\frac{\partial f_i}{\partial y_j}\right)$, for i, j = 1, 2, ..., n, given x and y_j , for j = 1, 2, ..., n.

Its specification is:

	SUBROUTINE JACOBF (X, EPS, Y, F, N)	
	INTEGER N double precision X, EPS, Y(N), F(N,N)	
1:	X – double precision Input	t
	On entry: the value of the argument x .	
2:	EPS – double precision Input	t
	On entry: the value of the continuation parameter ϵ . This is 1 if continuation is not being used.	g
3:	Y(N) – <i>double precision</i> array Input	t
	On entry: the value of the argument y_i , for $i = 1, 2,, n$.	
4:	F(N,N) – <i>double precision</i> array Output	t
	On exit: $F(i,j)$ must be set to the value of $\frac{\partial f_i}{\partial y_j}$, evaluated at the point (x,y) , for	
	$i,j=1,2,\ldots,n.$	
5:	N – INTEGER Input	t
	On entry: n, the number of equations.	

JACOBF must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

Input

Input

External Procedure

16: JACOBG – SUBROUTINE, supplied by the user.

External Procedure

JACOBG must evaluate the Jacobians $\left(\frac{\partial g_i}{\partial y_j(a)}\right)$ and $\left(\frac{\partial g_i}{\partial y_j(b)}\right)$. The ordering of the rows of AJ and BJ must correspond to the ordering of the boundary conditions described in the specification of (sub)program G.

Its specification is:

1		
	SUBROUTINE JACOBG (EPS, YA, YB, AJ, BJ, N)	
	INTEGER N double precision EPS, YA(N), YB(N), AJ(N,N), BJ(N,N)	
	<i>double precision</i> EPS, YA(N), YB(N), AJ(N,N), BJ(N,N)	
1:	EPS – double precision	Input
	On entry: ϵ , the value of the continuation parameter. This is 1 if continuation is no used.	ot being
2:	YA(N) - double precision array	Input
	On entry: the value $y_i(a)$, for $i = 1, 2,, n$.	-
3:	YB(N) - double precision array	Input
	On entry: the value $y_i(b)$, for $i = 1, 2,, n$.	
4:	AJ(N,N) – <i>double precision</i> array	Output
	On exit: AJ (i,j) must be set to the value $\frac{\partial g_i}{\partial y_j(a)}$, for $i,j = 1, 2,, n$.	
5:	BJ(N,N) – <i>double precision</i> array	Output
	On exit: BJ(<i>i</i> , <i>j</i>) must be set to the value $\frac{\partial g_i}{\partial y_j(b)}$, for $i, j = 1, 2,, n$.	
6:	N – INTEGER	Input
	On entry: n, the number of equations.	

JACOBG must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

17: DELEPS – *double precision*

Input/Output

On entry: must be given a value which specifies whether continuation is required. If DELEPS ≤ 0.0 or DELEPS ≥ 1.0 then it is assumed that continuation is not required. If 0.0 < DELEPS < 1.0 then it is assumed that continuation is required unless DELEPS $< \sqrt{machine precision}$ when an error exit is taken. DELEPS is used as the increment $\epsilon_2 - \epsilon_1$ (see (4)) and the choice DELEPS = 0.1 is recommended.

On exit: an overestimate of the increment $\epsilon_p - \epsilon_{p-1}$ (in fact the value of the increment which would have been tried if the restriction $\epsilon_p = 1$ had not been imposed). If continuation was not requested then DELEPS = 0.0.

If continuation is not requested then the parameters JACEPS and JACGEP may be replaced by dummy actual parameters in the call to D02RAF. (D02GAZ and D02GAX respectively may be used as the dummy parameters.)

18: JACEPS – SUBROUTINE, supplied by the user.

External Procedure

JACEPS must evaluate the derivative $\frac{\partial f_i}{\partial \epsilon}$ given x and y if continuation is being used.

Its specification is:

	SUBROUTINE JACEPS (X, EPS, Y, F, N)	
	INTEGERNdouble precisionX, EPS, Y(N), F(N)	
1:	X – double precision	Input
	On entry: the value of the argument x.	
2:	EPS – double precision	Input
	On entry: ϵ , the value of the continuation parameter.	
3:	Y(N) – <i>double precision</i> array	Input
	On entry: the solution values y_i at the point x, for $i = 1, 2,, n$.	
4:	$F(N) - double \ precision$ array	Output
	On exit: $F(i)$ must contain the value $\frac{\partial f_i}{\partial \epsilon}$ at the point (x, y) , for $i = 1, 2,, n$.	
5:	N – INTEGER	Input
	On entry: n, the number of equations.	

JACEPS must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

19:JACGEP – SUBROUTINE, supplied by the user.External ProcedureJACGEP must evaluate the derivatives $\frac{\partial g_i}{\partial \epsilon}$ if continuation is being used.

Its specification is:

	SUBROUTINE JACGEP (EPS, YA, YB, BCEP, N)	
	INTEGER N double precision EPS, YA(N), YB(N), BCEP(N)	
1:	EPS – double precision	Input
	On entry: ϵ , the value of the continuation parameter.	
2:	$YA(N) - double \ precision$ array	Input
	On entry: the value of $y_i(a)$, for $i = 1, 2,, n$.	
3:	YB(N) - double precision array	Input
	On entry: the value of $y_i(b)$, for $i = 1, 2,, n$.	
4:	BCEP(N) – <i>double precision</i> array	Output
	On exit: BCEP(i) must contain the value of $\frac{\partial g_i}{\partial \epsilon}$, for $i = 1, 2,, n$.	
5:	N – INTEGER	Input
	On entry: n, the number of equations.	

JACGEP must be declared as EXTERNAL in the (sub)program from which D02RAF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

20: WORK(LWORK) – *double precision* array

21: LWORK – INTEGER

On entry: the dimension of the array WORK as declared in the (sub)program from which D02RAF is called.

Constraint: LWORK \geq MNP \times (3N² + 6N + 2) + 4N² + 3N.

22: IWORK(LIWORK) – INTEGER array

23: LIWORK – INTEGER

On entry: the dimension of the array IWORK as declared in the (sub)program from which D02RAF is called.

Constraints:

 $\begin{array}{l} \text{if IJAC} \neq 0, \ \text{LIWORK} \geq \text{MNP} \times (2 \times \text{N} + 1) + \text{N}; \\ \text{if IJAC} = 0, \ \text{LIWORK} \geq \text{MNP} \times (2 \times \text{N} + 1) + \text{N}^2 + 4 \times \text{N} + 2. \end{array}$

24: IFAIL – INTEGER

Input/Output

Workspace

Workspace

Input

Input

For this routine, the normal use of IFAIL is extended to control the printing of error and warning messages as well as specifying hard or soft failure (see Chapter P01).

On entry: IFAIL must be set to a value with the decimal expansion cba, where each of the decimal digits c, b and a must have a value of 0 or 1.

a = 0 specifies hard failure, otherwise soft failure;

b = 0 suppresses error messages, otherwise error messages will be printed (see Section 6);

c = 0 suppresses warning messages, otherwise warning messages will be printed (see Section 6).

The recommended value for inexperienced users is 110 (i.e., hard failure with all messages printed).

On exit: IFAIL = 0 unless the routine detects an error (see Section 6).

6 Error Indicators and Warnings

If on entry IFAIL = 0 or -1, explanatory error messages are output on the current error message unit (as defined by X04AAF).

Errors or warnings detected by the routine:

IFAIL = 1

One or more of the parameters N, MNP, NP, NUMBEG, NUMMIX, TOL, DELEPS, LWORK or LIWORK has been incorrectly set, or $X(1) \ge X(NP)$ or the mesh points X(i) are not in strictly ascending order.

IFAIL = 2

A finer mesh is required for the accuracy requested; that is MNP is not large enough. This error exit normally occurs when the problem being solved is difficult (for example, there is a boundary layer) and high accuracy is requested. A poor initial choice of mesh points will make this error exit more likely.

IFAIL = 3

The Newton iteration has failed to converge. There are several possible causes for this error:

- (i) faulty coding in one of the Jacobian calculation routines;
- (ii) if IJAC = 0 then inaccurate Jacobians may have been calculated numerically (this is a very unlikely cause); or,

(iii) a poor initial mesh or initial approximate solution has been selected either by you or by default or there are not enough points in the initial mesh. Possibly, you should try the continuation facility.

IFAIL = 4

The Newton iteration has reached round-off error level. It could be however that the answer returned is satisfactory. The error is likely to occur if too high an accuracy is requested.

IFAIL = 5

The Jacobian calculated by JACOBG (or the equivalent matrix calculated by numerical differentiation) is singular. This may occur due to faulty coding of JACOBG or, in some circumstances, to a zero initial choice of approximate solution (such as is chosen when INIT = 0).

IFAIL = 6

There is no dependence on ϵ when continuation is being used. This can be due to faulty coding of JACEPS or JACGEP or, in some circumstances, to a zero initial choice of approximate solution (such as is chosen when INIT = 0).

IFAIL = 7

DELEPS is required to be less than *machine precision* for continuation to proceed. It is likely that either the problem (3) has no solution for some value near the current value of ϵ (see the advisory print out from D02RAF) or that the problem is so difficult that even with continuation it is unlikely to be solved using this routine. If the latter cause is suspected then using more mesh points initially may help.

IFAIL = 8 (D02RAF) IFAIL = 9 (D02RAR)

A serious error has occurred in a call in the specified routine. Check all array subscripts and (sub)program parameter lists in calls to D02RAF. Seek expert help.

7 Accuracy

The solution returned by the routine will be accurate to your tolerance as defined by the relation (5) except in extreme circumstances. The final error estimate over the whole mesh for each component is given in the array ABT. If too many points are specified in the initial mesh, the solution may be more accurate than requested and the error may not be approximately equidistributed.

8 **Further Comments**

There are too many factors present to quantify the timing. The time taken by D02RAF is negligible only on very simple problems.

You are strongly recommended to set IFAIL to obtain self-explanatory error messages, and also monitoring information about the course of the computation.

In the case where you wish to solve a sequence of similar problems, the use of the final mesh and solution from one case as the initial mesh is strongly recommended for the next.

9 Example

We solve the differential equation

$$y''' = -yy'' - 2\epsilon (1 - {y'}^2)$$

with $\epsilon = 1$ and boundary conditions

$$y(0) = y'(0) = 0, \quad y'(10) = 1$$

to an accuracy specified by TOL = 1.0D - 4. The continuation facility is used with the continuation parameter ϵ introduced as in the differential equation above and with DELEPS = 0.1 initially. (The continuation facility is not needed for this problem and is used here for illustration.)

9.1 Program Text

```
DO2RAF Example Program Text
*
      Mark 14 Revised. NAG Copyright 1989.
*
      .. Parameters ..
*
                       N, MNP, IY, LWORK, LIWORK
      INTEGER
                       (N=3,MNP=40,IY=N,LWORK=MNP*(3*N*N+6*N+2)
      PARAMETER
     +
                       +4*N*N+3*N,LIWORK=MNP*(2*N+1)+N)
      INTEGER
                       NOUT
      PARAMETER
                       (NOUT=6)
      .. Local Scalars ..
      DOUBLE PRECISION DELEPS, TOL
                       I, IFAIL, IJAC, INIT, J, NP, NUMBEG, NUMMIX
      INTEGER
      .. Local Arrays ..
      DOUBLE PRECISION ABT(N), WORK(LWORK), X(MNP), Y(IY,MNP)
      TNTEGER
                       IWORK(LIWORK)
      .. External Subroutines ...
                       D02RAF, FCN, G, JACEPS, JACGEP, JACOBF, JACOBG,
      EXTERNAL
     +
                       X04ABF
      .. Executable Statements ..
      WRITE (NOUT,*) 'DO2RAF Example Program Results'
      WRITE (NOUT, *)
      WRITE (NOUT, *) 'Calculation using analytic Jacobians'
      CALL X04ABF(1,NOUT)
      TOL = 1.0D-4
      NP = 17
      NUMBEG = 2
      NUMMIX = 0
      X(1) = 0.0D0
      X(NP) = 10.0D0
      INIT = 0
      DELEPS = 0.1D0
      IJAC = 1
*
      * Set IFAIL to 111 to obtain monitoring information *
      IFAIL = 11
*
      CALL D02RAF(N,MNP,NP,NUMBEG,NUMMIX,TOL,INIT,X,Y,N,ABT,FCN,G,IJAC,
     +
                  JACOBF, JACOBG, DELEPS, JACEPS, JACGEP, WORK, LWORK, IWORK,
     +
                  LIWORK, IFAIL)
*
      IF (IFAIL.EQ.0 .OR. IFAIL.EQ.4) THEN
         IF (IFAIL.EQ.4) WRITE (NOUT,99996)
             'On exit from DO2RAF IFAIL = ', IFAIL
     +
         WRITE (NOUT, *)
         WRITE (NOUT,99999) 'Solution on final mesh of ', NP, ' points'
         WRITE (NOUT, *)
     +
                  X(I)
                               Y1(I)
                                            Y2(T)
                                                          Y3(I)'
         WRITE (NOUT, 99998) (X(J), (Y(I,J), I=1,N), J=1,NP)
         WRITE (NOUT, *)
         WRITE (NOUT, *) 'Maximum estimated error by components'
         WRITE (NOUT, 99997) (ABT(I), I=1, N)
      ELSE
         WRITE (NOUT, 99996) 'On exit from DO2RAF IFAIL = ', IFAIL
      END IF
      STOP
99999 FORMAT (1X,A,I2,A)
99998 FORMAT (1X,F10.3,3F13.4)
99997 FORMAT (11X,1P,3E13.2)
99996 FORMAT (1X,A,I3)
     END
*
      SUBROUTINE FCN(X,EPS,Y,F,M)
      .. Scalar Arguments .
      DOUBLE PRECISION EPS, X
```

```
INTEGER
                     М
      .. Array Arguments ..
*
     DOUBLE PRECISION F(M), Y(M)
      .. Executable Statements ..
*
      F(1) = Y(2)
     F(2) = Y(3)
     F(3) = -Y(1)*Y(3) - 2.0D0*(1.0D0-Y(2)*Y(2))*EPS
     RETURN
     END
*
     SUBROUTINE G(EPS,Y,Z,AL,M)
*
      .. Scalar Arguments ..
      DOUBLE PRECISION EPS
     INTEGER
                  М
      .. Array Arguments ..
*
     DOUBLE PRECISION AL(M), Y(M), Z(M)
      .. Executable Statements ..
     AL(1) = Y(1)
     AL(2) = Y(2)
     AL(3) = Z(2) - 1.0D0
     RETURN
     END
*
     SUBROUTINE JACEPS(X, EPS, Y, F, M)
*
     .. Scalar Arguments ..
     DOUBLE PRECISION EPS, X
     INTEGER
                        М
      .. Array Arguments ..
*
     DOUBLE PRECISION F(M), Y(M)
      .. Executable Statements ..
*
      F(1) = 0.0D0
     F(2) = 0.0D0
      F(3) = -2.0D0 * (1.0D0 - Y(2) * Y(2))
     RETURN
     END
*
      SUBROUTINE JACGEP(EPS,Y,Z,AL,M)
*
      .. Scalar Arguments ..
     DOUBLE PRECISION EPS
      INTEGER
                        М
      .. Array Arguments .
*
     DOUBLE PRECISION AL(M), Y(M), Z(M)
      .. Local Scalars ..
*
      INTEGER
                        Ι
      .. Executable Statements ..
      DO 20 I = 1, M
        AL(I) = 0.0D0
  20 CONTINUE
     RETURN
     END
*
     SUBROUTINE JACOBF(X,EPS,Y,F,M)
      .. Scalar Arguments ..
     DOUBLE PRECISION EPS, X
     INTEGER
                        М
      .. Array Arguments ..
*
     DOUBLE PRECISION F(M,M), Y(M)
      .. Local Scalars ..
*
     INTEGER
                        I. J
      .. Executable Statements ..
*
     DO 40 I = 1, M
         DO 20 J = 1, M
            F(I,J) = 0.0D0
  20
        CONTINUE
  40 CONTINUE
     F(1,2) = 1.0D0
     F(2,3) = 1.0D0
     F(3,1) = -Y(3)
     F(3,2) = 4.0D0 * Y(2) * EPS
     F(3,3) = -Y(1)
     RETURN
```

```
END
*
      SUBROUTINE JACOBG(EPS,Y,Z,A,B,M)
*
      .. Scalar Arguments ..
      DOUBLE PRECISION EPS
     INTEGER
                        М
      .. Array Arguments ..
*
     DOUBLE PRECISION A(M,M), B(M,M), Y(M), Z(M)
     . Local Scalars ..
I, J
*
      .. Executable Statements ..
*
     DO 40 I = 1, M
DO 20 J = 1, M
           A(I,J) = 0.0D0
            B(I,J) = 0.0D0
      CONTINUE
   20
   40 CONTINUE
     A(1,1) = 1.0D0
      A(2,2) = 1.0D0
      B(3,2) = 1.0D0
      RETURN
      END
```

9.2 Program Data

None.

9.3 Program Results

D02RAF Example Program Results

Calculation using analytic Jacobians

	final mesh of	33 points	
X(I)	Y1(I)	Y2(I)	Y3(I)
0.000	0.0000	0.0000	1.6872
0.062	0.0032	0.1016	1.5626
0.125	0.0125	0.1954	1.4398
0.188	0.0275	0.2816	1.3203
0.250	0.0476	0.3605	1.2054
0.375	0.1015	0.4976	0.9924
0.500	0.1709	0.6097	0.8048
0.625	0.2530	0.6999	0.6438
0.703	0.3095	0.7467	0.5563
0.781	0.3695	0.7871	0.4784
0.938	0.4978	0.8513	0.3490
1.094	0.6346	0.8977	0.2502
1.250	0.7776	0.9308	0.1763
1.458	0.9748	0.9598	0.1077
1.667	1.1768	0.9773	0.0639
1.875	1.3815	0.9876	0.0367
2.031	1.5362	0.9922	0.0238
2.188	1.6915	0.9952	0.0151
2.500	2.0031	0.9983	0.0058
2.656	2.1591	0.9990	0.0035
2.812	2.3153	0.9994	0.0021
3.125	2.6277	0.9998	0.0007
3.750	3.2526	1.0000	0.0001
4.375	3.8776	1.0000	0.0000
5.000	4.5026	1.0000	0.0000
5.625	5.1276	1.0000	-0.0000
6.250	5.7526	1.0000	0.0000
6.875	6.3776	1.0000	-0.0000
7.500	7.0026	1.0000	0.0000
8.125	7.6276	1.0000	-0.0000
8.750	8.2526	1.0000	0.0000
9.375	8.8776	1.0000	-0.0000
10.000	9.5026	1.0000	0.0000

Maximum estimated error by components

6.92E-05 1.81E-05 6.42E-05