NAG Fortran Library Routine Document

D02NCF

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

D02NCF is a forward communication routine for integrating stiff systems of explicit ordinary differential equations when the Jacobian is a banded matrix.

2 Specification

```
SUBROUTINE D02NCF(NEQ, NEQMAX, T, TOUT, Y, YDOT, RWORK, RTOL, ATOL,1ITOL, INFORM, FCN, YSAVE, NY2DIM, JAC, WKJAC, NWKJAC,2JACPVT, NJCPVT, MONITR, ITASK, ITRACE, IFAIL)INTEGERNEQ, NEQMAX, ITOL, INFORM(23), NY2DIM, NWKJAC,1JACPVT(NJCPVT), NJCPVT, ITASK, ITRACE, IFAILrealT, TOUT, Y(NEQMAX), YDOT(NEQMAX), RWORK(50+4*NEQMAX),1RTOL(*), ATOL(*), YSAVE(NEQMAX,NY2DIM), WKJAC(NWKJAC)EXTERNALFCN, JAC, MONITR
```

3 Description

C C

С

D02NCF is a general purpose routine for integrating the initial value problem for a stiff system of explicit ordinary differential equations,

y' = g(t, y).

It is designed specifically for the case where the Jacobian $\frac{\partial g}{\partial u}$ is a banded matrix.

Both interval and step oriented modes of operation are available and also modes designed to permit intermediate output within an interval oriented mode.

An outline of a typical calling program for D02NCF is given below. It calls the banded matrix linear algebra setup routine D02NTF, the Backward Differentiation Formula (BDF) integrator setup routine D02NVF, and its diagnostic counterpart D02NVF.

```
declarations
EXTERNAL FCN, JAC, MONITR
...
IFAIL = 0
CALL D02NVF(...,IFAIL)
CALL D02NTF(NEQ, NEQMAX, JCEVAL, ML, MU, NWKJAC, NJCPVT,
+ RWORK, IFAIL)
IFAIL = -1
CALL D02NCF(NEQ, NEQMAX, T, TOUT, Y, YDOT, RWORK, RTOL,
+ ATOL, ITOL, INFORM, FCN, YSAVE, NY2DIM, JAC, WKJAC, NWKJAC,
+ JACPVT, NJCPVT, MONITR, ITASK, ITRACE, IFAIL)
IF (IFAIL.EQ.1 .OR. IFAIL.GE.14) STOP
IFAIL = 0
CALL D02NYF(...)
...
STOP
END
```

The linear algebra setup routine D02NTF and one of the integrator setup routines, D02NVF or D02NWF, must be called prior to the call of D02NCF. The integrator diagnostic routine D02NYF may be called after the call to D02NCF. There is also a routine, D02NZF, designed to permit the user to change step size on a continuation call to D02NCF without restarting the integration process.

4 References

None.

5 Parameters

1: NEQ – INTEGER

On entry: the number of differential equations to be solved.

Constraint: NEQ \geq 1.

2: NEQMAX – INTEGER

On entry: a bound on the maximum number of differential equations to be solved during the integration.

Constraint: NEQMAX \geq NEQ.

3: T – *real*

On entry: the value of the independent variable, t. The input value of T is used only on the first call as the initial point of the integration.

On exit: the value at which the computed solution y is returned (usually at TOUT).

4: TOUT – *real*

On entry: the next value of t at which a computed solution is desired. For the initial t, the input value of TOUT is used to determine the direction of integration. Integration is permitted in either direction (see also ITASK).

Constraint: TOUT \neq T.

5: Y(NEQMAX) – *real* array

On entry: the values of the dependent variables (solution). On the first call the first NEQ elements of y must contain the vector of initial values.

On exit: the computed solution vector, evaluated at T (usually T = TOUT).

6: YDOT(NEQMAX) – *real* array

On exit: the time derivatives y' of the vector y at the last integration point.

- 7: RWORK(50+4*NEQMAX) *real* array
- 8: RTOL(*) *real* array

Note: the dimension of the array RTOL must be at least 1 or NEQ (see ITOL).

On entry: the relative local error tolerance.

Constraint: $RTOL(i) \ge 0.0$ for all relevant i (see ITOL).

9: ATOL(*) – *real* array

Note: the dimension of the array ATOL must be at least 1 or NEQ (see ITOL). On entry: the absolute local error tolerance. Constraint: $ATOL(i) \ge 0.0$ for all relevant *i* (see ITOL). Input

Input/Output

Input

Input

Output

Input/Output

Workspace

Input

Input

ITOL – INTEGER

10:

On entry: a value to indicate the form of the local error test. ITOL indicates to D02NCF whether to interpret either or both of RTOL or ATOL as a vector or a scalar. The error test to be satisfied is $||e_i/w_i|| < 1.0$, where w_i is defined as follows:

ITOL	RTOL	ATOL	w_i
1 2 3 4	scalar vector	vector scalar	$\begin{array}{l} \operatorname{RTOL}(1) \times y_i + \operatorname{ATOL}(1) \\ \operatorname{RTOL}(1) \times y_i + \operatorname{ATOL}(i) \\ \operatorname{RTOL}(i) \times y_i + \operatorname{ATOL}(1) \\ \operatorname{RTOL}(i) \times y_i + \operatorname{ATOL}(i) \end{array}$

 e_i is an estimate of the local error in y_i , computed internally, and the choice of norm to be used is defined by a previous call to an integrator setup routine.

Constraint: $1 \leq \text{ITOL} \leq 4$.

- INFORM(23) INTEGER array 11:
- FCN SUBROUTINE, supplied by the user. 12:

FCN must evaluate the derivative vector for the explicit ordinary differential equation system, defined by y' = g(t, y).

Its specification is:

	SUBROUTINE FCN(NEQ, T, Y, F, IRES)INTEGERNEQ, IRESrealT, Y(NEQ), F(NEQ)
1:	NEQ – INTEGER Input
	On entry: the number of differential equations being solved.
2:	T – real Input
	On entry: the current value of the independent variable, t.
3:	Y(NEQ) – <i>real</i> array Input
	On entry: the value of y_i for $i = 1, 2,, NEQ$.
4:	F(NEQ) – <i>real</i> array Output
	On exit: the value y'_i , given by $y'_i = g_i(t, y)$, for $i = 1, 2,, NEQ$.
5:	IRES – INTEGER Input/Output
	On entry: $IRES = 1$.
	<i>On exit</i> : the user may set IRES as follows to indicate certain conditions in FCN to the integrator:
	IRES = 1
	Indicates a normal return from FCN, that is IRES is not altered by the user and integration continues.
	IRES = 2
	Indicates to the integrator that control should be passed back immediately to the calling (sub)program with the error indicator set to $IFAIL = 11$.

Input

D02NCF.3

Workspace

External Procedure

Workspace

External Procedure

Input

IRES = 3

Indicates to the integrator that an error condition has occurred in the solution vector, its time derivative or in the value of t. The integrator will use a smaller time step to try to avoid this condition. If this is not possible the integrator returns to the calling (sub)program with the error indicator set to IFAIL = 7.

IRES = 4

Indicates to the integrator to stop its current operation and to enter the MONITR routine immediately with parameter IMON = -2.

FCN must be declared as EXTERNAL in the (sub)program from which D02NCF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

13: YSAVE(NEQMAX,NY2DIM) – *real* array

14: NY2DIM – INTEGER

On entry: the second dimension of the array YSAVE as declared in the (sub)program from which D02NCF is called. An appropriate value for NY2DIM is described in the specifications of the integrator setup routines D02NVF and D02NWF. This value must be the same as that supplied to the integrator setup routine.

15: JAC – SUBROUTINE, supplied by the user.

JAC must evaluate the Jacobian of the system. If this option is not required, the actual argument for JAC must be the dummy routine D02NCZ. (D02NCZ is included in the NAG Fortran Library and so need not be supplied by the user. Its name may be implementation dependent: see the Users' Note for your implementation for details.) The user indicates to the integrator whether this option is to be used by setting the parameter JCEVAL appropriately in a call to the linear algebra setup routine D02NTF.

First we must define the system of nonlinear equations which is solved internally by the integrator. The time derivative, y', generated internally has the form

$$y' = (y - z)/(hd),$$

where h is the current step size and d is a parameter that depends on the integration method in use. The vector y is the current solution and the vector z depends on information from previous time steps. This means that $\frac{d}{dy}() = \frac{1}{(hd)}\frac{d}{dy}()$. The system of nonlinear equations that is solved has the form

$$y' - q(t, y) = 0$$

but it is solved in the form

$$r(t,y) = 0,$$

where the function r is defined by

$$r(t, y) = hd((y - z)/(hd) - g(t, y)).$$

It is the Jacobian matrix $\frac{\partial r}{\partial u}$ that the user must supply in the routine JAC as follows:

$$\frac{\partial r_i}{\partial y_j} = 1 - (hd) \frac{\partial g_i}{\partial y_j}, \quad \text{if } i = j,$$

 $\frac{\partial r_i}{\partial y_j} = -(hd) \frac{\partial g_i}{\partial y_j}, \quad \text{otherwise.}$

Its specification is:

SUBROUTINE	JAC(NEQ,	т,	Υ,	Н,	D,	ML,	MU,	P)
INTEGER	NEQ,	ML,	, MU	J				
real	т, т	(NEÇ	<u>)</u> ,	Н,	D,	P(M)	L+MU-	+1,NEQ)

1		1
1:	NEQ – INTEGER	Input
	On entry: the number of differential equations being solved.	
2:	T – real	Input
	On entry: the current value of the independent variable, t.	
3:	Y(NEQ) – <i>real</i> array	Input
	On entry: the current solution component y_i , for $i = 1, 2,, NEQ$.	
4:	H – real	Input
	On entry: the current step size.	
5:	D – real	Input
	On entry: the parameter d which depends on the integration method.	
6: 7:	ML – INTEGER MU – INTEGER	Input Input
	On entry: the number of sub-diagonals and super-diagonals respectively in the b	and.
8:	P(ML+MU+1,NEQ) – <i>real</i> array	Output
	<i>On exit</i> : elements of the Jacobian matrix $\frac{\partial r}{\partial y}$ stored as specified by the following code:	pseudo-
	DO 20 I = 1, NEQ J1 = MAX(I-ML,1) J2 = MIN(I+MU,NEQ) DO 10 J = J1, J2 K = MIN(ML+1-I,0)+J $P(K,I) = \delta R/\delta Y(I,J)$ 10 CONTINUE 20 CONTINUE	
	See also F07BDF (SGBTRF/DGBTRF).	
	Only non-zero elements of this array need be set, since it is preset to zero before to JAC.	the call

JAC must be declared as EXTERNAL in the (sub)program from which D02NCF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

16: WKJAC(NWKJAC) – *real* array

Workspace Input

Workspace Input

17: NWKJAC – INTEGER

On entry: the dimension of the array WKJAC as declared in the (sub)program from which D02NCF is called. This value must be the same as that supplied to the linear algebra setup routine D02NTF.

Constraint: NWKJAC $\geq (2m_L + m_U + 1) \times \text{NEQMAX}$ where m_L and m_U are the number of subdiagonals and super-diagonals respectively in the band, defined by a call to D02NTF.

18: JACPVT(NJCPVT) - INTEGER array

On entry: the dimension of the array JACPVT as declared in the (sub)program from which D02NCF is called. This value must be the same as that supplied to the linear algebra setup routine D02NTF. *Constraint:* NICPVT > NEOMAX

Constraint: NJCPVT \geq NEQMAX.

20: MONITR - SUBROUTINE, supplied by the user.

External Procedure

MONITR performs tasks requested by the user. If this option is not required, then the actual argument for MONITR must be the dummy routine D02NBY. (D02NBY is included in the NAG Fortran Library and so need not be supplied by the user. Its name may be implementation dependent: see the Users' Note for your implementation for details.)

Its specification is:

SUBROUTINE MONITR(NEQ, NEQMAX, T, HLAST, HNEXT, Y, YDOT, YSAVE, R, 1 ACOR, IMON, INLN, HMIN, HMAX, NOU) NEQ, NEQMAX, IMON, INLN, NQU INTEGER T, HLAST, HNEXT, Y(NEQMAX), YDOT(NEQMAX), real 1 YSAVE(NEQMAX,*), R(NEQMAX), ACOR(NEQMAX,2), HMIN, 2 HMAX NEQ - INTEGER Input 1: On entry: the number of differential equations being solved. 2: NEQMAX - INTEGER Input On entry: an upper bound on the number of differential equations to be solved. 3: T – real Input On entry: the current value of the independent variable. 4: HLAST - real Input On entry: the last step size successfully used by the integrator. 5: HNEXT - real Input/Output On entry: the step size that the integrator proposes to take on the next step. On exit: the next step size to be used. If this is different from the input value, then IMON must be set to 4. Y(NEQMAX) - real array 6: Input/Output On entry: the values of the dependent variables, y, evaluated at t. On exit: these values must not be changed unless IMON is set to 2. 7: YDOT(NEQMAX) - real array Input On entry: the time derivatives y' of the vector y. YSAVE(NEQMAX,*) - real array 8: Input On entry: workspace to enable the user to carry out interpolation using either of the routines D02XJF or D02XKF. 9. R(NEQMAX) - real array Input On entry: if IMON = 0 and INLN = 3, the first NEQ elements contain the residual vector y' - g(t, y).ACOR(NEQMAX,2) - real array 10: Input On entry: with IMON = 1, ACOR(i, 1) contains the weight used for the *i*th equation when the norm is evaluated, and ACOR(i, 2) contains the estimated local error for the *i*th equation. The scaled local error at the end of a timestep may be obtained by calling the

real function D02ZAF as follows:

IFAIL = 1ERRLOC = D02ZAF(NEQ, ACOR(1,2), ACOR(1,1), IFAIL)CHECK IFAIL BEFORE PROCEEDING С IMON - INTEGER 11: Input/Output On entry: a flag indicating under what circumstances MONITR was called: IMON = -2Entry from the integrator after IRES = 4 (set in FCN) caused an early termination (this facility could be used to locate discontinuities). IMON = -1The current step failed repeatedly. IMON = 0Entry after a call to the internal nonlinear equation solver (see below). IMON = 1The current step was successful. On exit: IMON may be reset to determine subsequent action in D02NCF: IMON = -2Integration is to be halted. A return will be made from the integrator to the calling (sub)program with IFAIL = 12. IMON = -1Allow the integrator to continue with its own internal strategy. The integrator will try up to 3 restarts unless IMON is set $\neq -1$ on exit. IMON = 0Return to the internal nonlinear equation solver, where the action taken is determined by the value of INLN (see below). IMON = 1Normal exit to the integrator to continue integration. IMON = 2Restart the integration at the current time point. The integrator will restart from order 1 when this option is used. The MONITR provided solution Y will be used for the initial conditions. IMON = 3Try to continue with the same step size and order as was to be used before the call to MONITR. HMIN and HMAX may be altered if desired. IMON = 4Continue the integration but using a new value HNEXT and possibly new values of HMIN and HMAX. INLN – INTEGER 12: Output On exit: the action to be taken by the internal nonlinear equation solver when MONITR is exited with IMON = 0. By setting INLN = 3 and returning to the integrator, the residual vector is evaluated and placed in the array R, and then MONITR is called again. At present this is the only option available: INLN must not be set to any other value.

Input/Output

Input/Output

13: HMIN – *real*

On entry: the minimum step size to be taken on the next step.

On exit: the minimum step size to be used. If this is different from the input value, then IMON must be set to 3 or 4.

14: HMAX – *real*

On entry: the maximum step size to be taken on the next step.

On exit: the maximum step size to be used. If this is different from the input value, then IMON must be set to 3 or 4. If HMAX is set to zero, no limit is assumed.

15: NQU – INTEGER

On entry: the order of the integrator used on the last step. This is supplied to enable the user to carry out interpolation using either of the routines D02XJF or D02XKF.

MONITR must be declared as EXTERNAL in the (sub)program from which D02NCF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

21: ITASK – INTEGER

On entry: the task to be performed by the integrator. The permitted values for ITASK and their meanings are detailed below:

ITASK = 1

Normal computation of output values of y(t) at t = TOUT (by overshooting and interpolating).

ITASK = 2

Take one step only and return.

ITASK = 3

Stop at the first internal integration point at or beyond t = TOUT and return.

ITASK = 4

Normal computation of output values of y(t) at t = TOUT but without overshooting t = TCRIT. TCRIT must be specified as an option in one of the integrator setup routines prior to the first call to the integrator, or specified in the optional input routine prior to a continuation call. TCRIT may be equal to or beyond TOUT, but not before it, in the direction of integration.

ITASK = 5

Take one step only and return, without passing TCRIT. TCRIT must be specified as under ITASK = 4.

Constraint: $1 \leq \text{ITASK} \leq 5$.

22: ITRACE – INTEGER

On entry: the level of output that is printed by the integrator. ITRACE may take the value -1, 0, 1, 2 or 3. If ITRACE < -1, then -1 is assumed and similarly if ITRACE > 3, then 3 is assumed. If ITRACE = -1, no output is generated. If ITRACE = 0, only warning messages are printed on the current error message unit (see X04AAF). If ITRACE > 0, then warning messages are printed as above, and on the current advisory message unit (see X04ABF) output is generated which details Jacobian entries, the nonlinear iteration and the time integration. The advisory messages are given in greater detail the larger the value of ITRACE.

Input

Input

Input

Input/Output

On entry: IFAIL must be set to 0, -1 or 1. Users who are unfamiliar with this parameter should refer to Chapter P01 for details.

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

For environments where it might be inappropriate to halt program execution when an error is detected, the value -1 or 1 is recommended. If the output of error messages is undesirable, then the value 1 is recommended. Otherwise, because for this routine the values of the output parameters may be useful even if IFAIL $\neq 0$ on exit, the recommended value is -1. When the value -1 or 1 is used it is essential to test the value of IFAIL on exit.

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

An illegal input was detected on entry, or after an internal call to MONITR. If ITRACE > -1, then the form of the error will be detailed on the current error message unit (see X04AAF).

IFAIL = 2

The maximum number of steps specified has been taken (see the description of optional inputs in the integrator setup routines and the optional input continuation routine, D02NZF).

IFAIL = 3

With the given values of RTOL and ATOL no further progress can be made across the integration range from the current point T. The components $Y(1), Y(2), \ldots, Y(NEQ)$ contain the computed values of the solution at the current point T.

IFAIL = 4

There were repeated error test failures on an attempted step, before completing the requested task, but the integration was successful as far as T. The problem may have a singularity, or the local error requirements may be inappropriate.

IFAIL = 5

There were repeated convergence test failures on an attempted step, before completing the requested task, but the integration was successful as far as T. This may be caused by an inaccurate Jacobian matrix or one which is incorrectly computed.

IFAIL = 6

Some error weight w_i became zero during the integration (see description of ITOL). Pure relative error control (ATOL(i) = 0.0) was requested on a variable (the *i*th) which has now vanished. The integration was successful as far as T.

IFAIL = 7

The user-supplied subroutine FCN set its error flag (IRES = 3) continually despite repeated attempts by the integrator to avoid this.

IFAIL = 8

Not used for this integrator.

IFAIL = 9

A singular Jacobian $\frac{\partial r}{\partial y}$ has been encountered. This error exit is unlikely to be taken when solving explicit ordinary differential equations. The user should check the problem formulation and Jacobian calculation.

IFAIL = 10

An error occurred during Jacobian formulation or back-substitution (a more detailed error description may be directed to the current error message unit, see X04AAF).

IFAIL = 11

The user-supplied subroutine FCN signalled the integrator to halt the integration and return (IRES = 2). Integration was successful as far as T.

IFAIL = 12

The user-supplied subroutine MONITR set IMON = -2 and so forced a return but the integration was successful as far as T.

IFAIL = 13

The requested task has been completed, but it is estimated that a small change in RTOL and ATOL is unlikely to produce any change in the computed solution. (Only applies when the user is not operating in one step mode, that is when ITASK $\neq 2$ or 5.)

IFAIL = 14

The values of RTOL and ATOL are so small that the routine is unable to start the integration.

IFAIL = 15

The linear algebra setup routine D02NTF was not called prior to calling D02NCF.

7 Accuracy

The accuracy of the numerical solution may be controlled by a careful choice of the parameters RTOL and ATOL, and to a much lesser extent by the choice of norm. Users are advised to use scalar error control unless the components of the solution are expected to be poorly scaled. For the type of decaying solution typical of many stiff problems, relative error control with a small absolute error threshold will be most appropriate (that is the user is advised to choose ITOL = 1 with ATOL(1) small but positive).

8 Further Comments

The cost of computing a solution depends critically on the size of the differential system and to a lesser extent on the degree of stiffness of the problem. For D02NCF the cost is proportional to NEQ × $(ML + MU + 1)^2$ though for problems which are only mildly nonlinear the cost may be dominated by factors proportional to NEQ × (ML + MU + 1) except for very large problems.

In general the user is advised to choose the Backward Differentiation Formula option (setup routine D02NVF) but if efficiency is of great importance and especially if it is suspected that $\frac{\partial g}{\partial y}$ has complex eigenvalues near the imaginary axis for some part of the integration, the user should try the BLEND option (setup routine D02NWF).

9 Example

We solve the well-known stiff Robertson problem

$$a' = -0.04a + 1.0E4bc$$

 $b' = 0.04a - 1.0E4bc - 3.0E7b^2$
 $c' = 3.0E7b^2$

over the range [0,10] with initial conditions a = 1.0 and b = c = 0.0 using scalar relative error control and vector absolute error control (ITOL = 2). We obtain the solution at TOUT = 5.0 and TOUT = 10.0 by overshooting and internal C^0 interpolation (ITASK = 1). D02NBY is used for MONITR, we use the BLEND integrator (setup routine D02NWF) and we choose the option of an analytical Jacobian.

9.1 Program Text

Note: the listing of the example program presented below uses *bold italicised* terms to denote precision-dependent details. Please read the Users' Note for your implementation to check the interpretation of these terms. As explained in the Essential Introduction to this manual, the results produced may not be identical for all implementations.

```
DO2NCF Example Program Text
*
      Mark 14 Revised. NAG Copyright 1989.
*
+
      .. Parameters ..
      TNTEGER
                       NOUT
      PARAMETER
                        (NOUT=6)
                       NEQ, NEQMAX, NRW, NINF, ML, MU, NJCPVT, NWKJAC,
      INTEGER
                       MAXORD, NY2DIM, MAXSTP, MXHNIL
     +
      PARAMETER
                       (NEQ=3,NEQMAX=NEQ,NRW=50+4*NEQMAX,NINF=23,ML=1,
     +
                       MU=2,NJCPVT=NEQMAX,NWKJAC=NEQMAX*(2*ML+MU+1),
     +
                       MAXORD=11,NY2DIM=MAXORD+3,MAXSTP=200,MXHNIL=5)
                       HO, HMAX, HMIN, TCRIT
      real
      PARAMETER
                        (H0=0.0e0, HMAX=10.0e0, HMIN=1.0e-10, TCRIT=0.0e0)
      .. Local Scalars ..
*
      real
                        H, HU, T, TCUR, TOLSF, TOUT
                        I, IFAIL, IMXER, ITASK, ITOL, ITRACE, NITER, NJE,
      INTEGER
                       NO, NOU, NRE, NST
      .. Local Arrays ..
*
                       ATOL(NEQMAX), CONST(6), RTOL(NEQMAX), RWORK(NRW),
      real
                        WKJAC(NWKJAC), Y(NEQMAX), YDOT(NEQMAX),
     +
     +
                       YSAVE (NEQMAX, NY2DIM)
      INTEGER
                       INFORM(NINF), JACPVT(NJCPVT)
                       ALGEQU(NEQMAX)
      LOGICAL
      .. External Subroutines ..
EXTERNAL DO2NBY, DO2NCF, DO2NTF, DO2NWF, DO2NYF, DO2NZF,
*
      EXTERNAL
                       FCN, JAC, XO4ABF
      .. Executable Statements .
*
      WRITE (NOUT, *) 'DO2NCF Example Program Results'
      CALL X04ABF(1,NOUT)
*
*
      Integrate to TOUT (ITASK=1 i.e. overshooting and internal
*
      interpolation) using the blend method. Default values for the
*
      array CONST are used. Employ scalar relative tolerance and vector
      absolute tolerance. The Jacobian is evaluated by JAC.
*
*
      MONITR subroutine replaced by NAG dummy routine DO2NBY.
      T = 0.0e0
      TOUT = 5.0e0
      ITASK = 1
      Y(1) = 1.0e0
      Y(2) = 0.0e0
      Y(3) = 0.0e0
      ITOL = 2
      RTOL(1) = 1.0e-4
      ATOL(1) = 1.0e-7
      ATOL(2) = 1.0e - 8
      ATOL(3) = 1.0e-7
      DO 20 I = 1, 6
        CONST(I) = 0.0e0
   20 CONTINUE
      IFAIL = 0
```

```
*
      CALL D02NWF(NEQMAX,NY2DIM,MAXORD,CONST,TCRIT,HMIN,HMAX,H0,MAXSTP,
     +
                  MXHNIL, 'Average-L2', RWORK, IFAIL)
      CALL D02NTF(NEQ,NEQMAX,'Analytical',ML,MU,NWKJAC,NJCPVT,RWORK,
     +
                  IFAIL)
      WRITE (NOUT,*)
      WRITE (NOUT, *) '
                          Х
                                      Y(1)
                                                      Y(2)
                                                                     Y(3)′
      WRITE (NOUT, 99999) T, (Y(I), I=1, NEQ)
      Soft fail and error messages only
      ITRACE = 0
      IFAIL = 1
      CALL D02NCF(NEQ,NEQMAX,T,TOUT,Y,YDOT,RWORK,RTOL,ATOL,ITOL,INFORM,
     +
                  FCN, YSAVE, NY2DIM, JAC, WKJAC, NWKJAC, JACPVT, NJCPVT,
     +
                  D02NBY,ITASK,ITRACE,IFAIL)
*
      IF (IFAIL.EO.O) THEN
         WRITE (NOUT, 99999) T, (Y(I), I=1, NEQ)
      ELSE
         WRITE (NOUT, *)
         WRITE (NOUT, 99998) 'Exit DO2NCF with IFAIL = ', IFAIL,
           ' and T = ', T
         STOP
      END IF
      Reset TOUT and call DO2NZF to override internal choice for
      stepsize. No changes to other parameters.
      H = 0.7e0
      IFAIL = 0
*
      CALL D02NZF(NEQMAX,TCRIT,H,HMIN,HMAX,MAXSTP,MXHNIL,RWORK,IFAIL)
*
      TOUT = 10.0e^{0}
      IFAIL = 1
*
      CALL D02NCF(NEQ,NEQMAX,T,TOUT,Y,YDOT,RWORK,RTOL,ATOL,ITOL,INFORM,
     +
                  FCN, YSAVE, NY2DIM, JAC, WKJAC, NWKJAC, JACPVT, NJCPVT,
     +
                  DO2NBY, ITASK, ITRACE, IFAIL)
*
      IF (IFAIL.EO.O) THEN
         WRITE (NOUT, 99999) T, (Y(I), I=1, NEQ)
         CALL D02NYF(NEQ,NEQMAX,HU,H,TCUR,TOLSF,RWORK,NST,NRE,NJE,NQU,
     +
                     NQ,NITER,IMXER,ALGEQU,INFORM,IFAIL)
         WRITE (NOUT, *)
         WRITE (NOUT,99997) ' HUSED = ', HU, ' HNEXT = ', H,
            TCUR = ', TCUR
     +
         WRITE (NOUT,99996) ' NST = ', NST, '
                                                 NRE = ', NRE,
                NJE = ', NJE
     +
         WRITE (NOUT, 99996) ' NQU = ', NQU, '
                                                 NQ = ', NQ,
           ' NITER = ', NITER
     +
         WRITE (NOUT, 99995) ' Max Err Comp = ', IMXER
         WRITE (NOUT, *)
      ELSE
         WRITE (NOUT, *)
         WRITE (NOUT, 99998) 'Exit DO2NCF with IFAIL = ', IFAIL,
     +
           ' and T = ', T
      END IF
      STOP
99999 FORMAT (1X,F8.3,3(F13.5,2X))
99998 FORMAT (1X,A,I2,A,e12.5)
99997 FORMAT (1X,A,e12.5,A,e12.5,A,e12.5)
99996 FORMAT (1X,A,I6,A,I6,A,I6)
99995 FORMAT (1X,A,I4)
      END
*
      SUBROUTINE FCN(NEQ,T,Y,R,IRES)
      .. Scalar Arguments ..
```

```
real
                   Т
     INTEGER
                   IRES, NEQ
     .. Array Arguments ..
*
     real
                  R(NEQ), Y(NEQ)
     .. Executable Statements ..
*
     R(3) = 3.0e7 * Y(2) * Y(2)
     RETURN
     END
*
     SUBROUTINE JAC(NEQ,T,Y,H,D,ML,MU,P)
*
     .. Scalar Arguments ..
                  D, H, Т
     real
     INTEGER
                  ML, MU, NEQ
     .. Array Arguments ..
*
              P(ML+MU+1,NEQ), Y(NEQ)
     real
     .. Local Scalars ..
*
     real
           HXD
*
     .. Executable Statements ..
     HXD = H*D
*
     P(1,1) = 1.0e0 - HXD * (-0.04e0)
     P(2,1) = -HXD*(1.0e4*Y(3))
     P(3,1) = -HXD * (1.0e4 * Y(2))
*
     P(1,2) = -HXD * (0.04e0)
     P(2,2) = 1.0e0 - HXD*(-1.0e4*Y(3)-6.0e7*Y(2))
     P(3,2) = -HXD*(-1.0e4*Y(2))
*
     P(1,3) = -HXD * (6.0e7 * Y(2))
     P(2,3) = 1.0e0 - HXD * (0.0e0)
     RETURN
     END
```

9.2 Program Data

None.

9.3 Program Results

DO2NCF Example Program Results Y(1) Y(2) Y(3) Х 0.00000 0.000 1.00000 0.00000 5.000 0.00002 0.10846 0.89152 10.000 0.84137 0.00002 0.15861 HUSED = 0.11280E+01 HNEXT = 0.11280E+01 TCUR = 0.10034E+02 NST = 63 NRE = 274 NJE = 14 NQU = 4 NQ = 4 NITER = 272 NQ = Max Err Comp = 3