# NAG Library Function Document nag_pde_parab_1d_keller_ode_remesh (d03prc) 

## 1 Purpose

nag_pde_parab_1d_keller_ode_remesh (d03prc) integrates a system of linear or nonlinear, first-order, time-dependent partial differential equations (PDEs) in one space variable, with scope for coupled ordinary differential equations (ODEs), and automatic adaptive spatial remeshing. The spatial discretization is performed using the Keller box scheme (see Keller (1970)) and the method of lines is employed to reduce the PDEs to a system of ODEs. The resulting system is solved using a Backward Differentiation Formula (BDF) method or a Theta method (switching between Newton's method and functional iteration).

## 2 Specification

```
#include <nag.h>
#include <nagd03.h>
void nag_pde_parab_ld_keller_ode_remesh (Integer npde, double *ts,
    double tout,
    void (*pdedef)(Integer npde, double t, double x, const double u[],
        const double udot[], const double ux[], Integer ncode,
        const double v[], const double vdot[], double res[], Integer *ires,
        Nag_Comm *comm),
    void (*bndary)(Integer npde, double t, Integer ibnd, Integer nobc,
        const double u[], const double udot[], Integer ncode,
        const double v[], const double vdot[], double res[], Integer *ires,
        Nag_Comm *comm),
        void (*uvinit)(Integer npde, Integer npts, Integer nxi,
            const double x[], const double xi[], double u[], Integer ncode,
            double v[], Nag_Comm *comm),
        double u[], Integer npts, double x[], Integer nleft, Integer ncode,
        void (*odedef)(Integer npde, double t, Integer ncode, const double v[],
            const double vdot[], Integer nxi, const double xi[],
            const double ucp[], const double ucpx[], const double ucpt[],
            double r[], Integer *ires, Nag_Comm *comm),
        Integer nxi, const double xi[], Integer neqn, const double rtol[],
        const double atol[], Integer itol, Nag_NormType norm,
        Nag_LinAlgOption laopt, const double algopt[], Nag_Boolean remesh,
        Integer nxfix, const double xfix[], Integer nrmesh, double dxmesh,
        double trmesh, Integer ipminf, double xratio, double con,
        void (*monitf)(double t, Integer npts, Integer npde, const double x[],
            const double u[], double fmon[], Nag_Comm *comm),
        double rsave[], Integer lrsave, Integer isave[], Integer lisave,
        Integer itask, Integer itrace, const char *outfile, Integer *ind,
        Nag_Comm *comm, Nag_DO3_Save *saved, NagError *fail)
```


## 3 Description

nag_pde_parab_1d_keller_ode_remesh (d03prc) integrates the system of first-order PDEs and coupled ODEs given by the master equations:

$$
\begin{gather*}
G_{i}\left(x, t, U, U_{x}, U_{t}, V, \dot{V}\right)=0, \quad i=1,2, \ldots, \text { npde }, \quad a \leq x \leq b, t \geq t_{0}  \tag{1}\\
R_{i}\left(t, V, \dot{V}, \xi, U^{*}, U_{x}^{*}, U_{t}^{*}\right)=0, \quad i=1,2, \ldots, \text { ncode } . \tag{2}
\end{gather*}
$$

In the PDE part of the problem given by (1), the functions $G_{i}$ must have the general form

$$
\begin{equation*}
G_{i}=\sum_{j=1}^{\text {npde }} P_{i, j} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {node }} Q_{i, j} \dot{V}_{j}+S_{i}=0, \quad i=1,2, \ldots, \text { npde }, \tag{3}
\end{equation*}
$$

where $P_{i, j}, Q_{i, j}$ and $S_{i}$ depend on $x, t, U, U_{x}$ and $V$.
The vector $U$ is the set of PDE solution values

$$
U(x, t)=\left[U_{1}(x, t), \ldots, U_{\text {npde }}(x, t)\right]^{\mathrm{T}},
$$

and the vector $U_{x}$ is the partial derivative with respect to $x$. The vector $V$ is the set of ODE solution values

$$
V(t)=\left[V_{1}(t), \ldots, V_{\text {ncode }}(t)\right]^{\mathrm{T}},
$$

and $\dot{V}$ denotes its derivative with respect to time.
In the ODE part given by (2), $\xi$ represents a vector of $n_{\xi}$ spatial coupling points at which the ODEs are coupled to the PDEs. These points may or may not be equal to some of the PDE spatial mesh points. $U^{*}, U_{x}^{*}$ and $U_{t}^{*}$ are the functions $U, U_{x}$ and $U_{t}$ evaluated at these coupling points. Each $R_{i}$ may only depend linearly on time derivatives. Hence equation (2) may be written more precisely as

$$
\begin{equation*}
R=A-B \dot{V}-C U_{t}^{*}, \tag{4}
\end{equation*}
$$

where $R=\left[R_{1}, \ldots, R_{\text {ncode }}\right]^{\mathrm{T}}, A$ is a vector of length ncode, $B$ is an ncode by ncode matrix, $C$ is an ncode by ( $n_{\xi} \times$ npde) matrix and the entries in $A, B$ and $C$ may depend on $t, \xi, U^{*}, U_{x}^{*}$ and $V$. In practice you only need to supply a vector of information to define the ODEs and not the matrices $B$ and $C$. (See Section 5 for the specification of odedef.)
The integration in time is from $t_{0}$ to $t_{\text {out }}$, over the space interval $a \leq x \leq b$, where $a=x_{1}$ and $b=x_{\text {npts }}$ are the leftmost and rightmost points of a mesh $x_{1}, x_{2}, \ldots, x_{\text {npts }}$ defined initially by you and (possibly) adapted automatically during the integration according to user-specified criteria.
The PDE system which is defined by the functions $G_{i}$ must be specified in pdedef.
The initial $\left(t=t_{0}\right)$ values of the functions $U(x, t)$ and $V(t)$ must be specified in uvinit. Note that uvinit will be called again following any remeshing, and so $U\left(x, t_{0}\right)$ should be specified for all values of $x$ in the interval $a \leq x \leq b$, and not just the initial mesh points.
For a first-order system of PDEs, only one boundary condition is required for each PDE component $U_{i}$. The npde boundary conditions are separated into $n_{a}$ at the left-hand boundary $x=a$, and $n_{b}$ at the right-hand boundary $x=b$, such that $n_{a}+n_{b}=\mathbf{n p d e}$. The position of the boundary condition for each component should be chosen with care; the general rule is that if the characteristic direction of $U_{i}$ at the left-hand boundary (say) points into the interior of the solution domain, then the boundary condition for $U_{i}$ should be specified at the left-hand boundary. Incorrect positioning of boundary conditions generally results in initialization or integration difficulties in the underlying time integration functions.
The boundary conditions have the master equation form:

$$
\begin{equation*}
G_{i}^{L}\left(x, t, U, U_{t}, V, \dot{V}\right)=0 \quad \text { at } x=a, \quad i=1,2, \ldots, n_{a}, \tag{5}
\end{equation*}
$$

at the left-hand boundary, and

$$
\begin{equation*}
G_{i}^{R}\left(x, t, U, U_{t}, V, \dot{V}\right)=0 \quad \text { at } x=b, \quad i=1,2, \ldots, n_{b}, \tag{6}
\end{equation*}
$$

at the right-hand boundary.
Note that the functions $G_{i}^{L}$ and $G_{i}^{R}$ must not depend on $U_{x}$, since spatial derivatives are not determined explicitly in the Keller box scheme functions. If the problem involves derivative (Neumann) boundary conditions then it is generally possible to restate such boundary conditions in terms of permissible variables. Also note that $G_{i}^{L}$ and $G_{i}^{R}$ must be linear with respect to time derivatives, so that the boundary conditions have the general form:

$$
\begin{equation*}
\sum_{j=1}^{\text {npde }} E_{i, j}^{L} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} H_{i, j}^{L} \dot{V}_{j}+K_{i}^{L}=0, \quad i=1,2, \ldots, n_{a} \tag{7}
\end{equation*}
$$

at the left-hand boundary, and

$$
\begin{equation*}
\sum_{j=1}^{\text {npde }} E_{i, j}^{R} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} H_{i, j}^{R} \dot{V}_{j}+K_{i}^{R}=0, \quad i=1,2, \ldots, n_{b} \tag{8}
\end{equation*}
$$

at the right-hand boundary, where $E_{i, j}^{L}, E_{i, j}^{R}, H_{i, j}^{L}, H_{i, j}^{R}, K_{i}^{L}$ and $K_{i}^{R}$ depend on $x, t, U$ and $V$ only.
The boundary conditions must be specified in bndary.
The problem is subject to the following restrictions:
(i) $P_{i, j}, Q_{i, j}$ and $S_{i}$ must not depend on any time derivatives;
(ii) $t_{0}<t_{\text {out }}$, so that integration is in the forward direction;
(iii) The evaluation of the function $G_{i}$ is done approximately at the mid-points of the mesh $\mathbf{x}[i-1]$, for $i=1,2, \ldots$, npts, by calling pdedef for each mid-point in turn. Any discontinuities in the function must therefore be at one or more of the fixed mesh points specified by xfix;
(iv) At least one of the functions $P_{i, j}$ must be nonzero so that there is a time derivative present in the PDE problem.

The algebraic-differential equation system which is defined by the functions $R_{i}$ must be specified in odedef. You must also specify the coupling points $\xi$ in the array $\mathbf{x i}$.
The first-order equations are approximated by a system of ODEs in time for the values of $U_{i}$ at mesh points. In this method of lines approach the Keller box scheme is applied to each PDE in the space variable only, resulting in a system of ODEs in time for the values of $U_{i}$ at each mesh point. In total there are npde $\times$ npts + ncode ODEs in time direction. This system is then integrated forwards in time using a Backward Differentiation Formula (BDF) or a Theta method.
The adaptive space remeshing can be used to generate meshes that automatically follow the changing time-dependent nature of the solution, generally resulting in a more efficient and accurate solution using fewer mesh points than may be necessary with a fixed uniform or non-uniform mesh. Problems with travelling wavefronts or variable-width boundary layers for example will benefit from using a moving adaptive mesh. The discrete time-step method used here (developed by Furzeland (1984)) automatically creates a new mesh based on the current solution profile at certain time-steps, and the solution is then interpolated onto the new mesh and the integration continues.
The method requires you to supply monitf which specifies in an analytic or numeric form the particular aspect of the solution behaviour you wish to track. This so-called monitor function is used to choose a mesh which equally distributes the integral of the monitor function over the domain. A typical choice of monitor function is the second space derivative of the solution value at each point (or some combination of the second space derivatives if more than one solution component), which results in refinement in regions where the solution gradient is changing most rapidly.

You must specify the frequency of mesh updates along with certain other criteria such as adjacent mesh ratios. Remeshing can be expensive and you are encouraged to experiment with the different options in order to achieve an efficient solution which adequately tracks the desired features of the solution.
Note that unless the monitor function for the initial solution values is zero at all user-specified initial mesh points, a new initial mesh is calculated and adopted according to the user-specified remeshing criteria. uvinit will then be called again to determine the initial solution values at the new mesh points (there is no interpolation at this stage) and the integration proceeds.

## 4 References

Berzins M (1990) Developments in the NAG Library software for parabolic equations Scientific Software Systems (eds J C Mason and M G Cox) 59-72 Chapman and Hall

Berzins M, Dew P M and Furzeland R M (1989) Developing software for time-dependent problems using the method of lines and differential-algebraic integrators Appl. Numer. Math. 5375-397
Berzins M and Furzeland R M (1992) An adaptive theta method for the solution of stiff and nonstiff differential equations Appl. Numer. Math. 9 1-19

Furzeland R M (1984) The construction of adaptive space meshes TNER.85.022 Thornton Research Centre, Chester
Keller H B (1970) A new difference scheme for parabolic problems Numerical Solutions of Partial Differential Equations (ed J Bramble) 2 327-350 Academic Press
Pennington S V and Berzins M (1994) New NAG Library software for first-order partial differential equations ACM Trans. Math. Softw. 20 63-99

## 5 Arguments

1: npde - Integer
On entry: the number of PDEs to be solved.
Constraint: npde $\geq 1$.
2: $\quad$ ts - double *
Input/Output
On entry: the initial value of the independent variable $t$.
Constraint: ts $<$ tout.
On exit: the value of $t$ corresponding to the solution values in $\mathbf{u}$. Normally $\mathbf{t s}=$ tout.
3: tout - double
Input
On entry: the final value of $t$ to which the integration is to be carried out.
4: pdedef - function, supplied by the user
External Function
pdedef must evaluate the functions $G_{i}$ which define the system of PDEs. pdedef is called approximately midway between each pair of mesh points in turn by nag_pde_parab_1d_keller_ ode_remesh (d03prc).

```
The specification of pdedef is:
void pdedef (Integer npde, double t, double x, const double u[],
        const double udot[], const double ux[], Integer ncode,
        const double v[], const double vdot[], double res[],
        Integer *ires, Nag_Comm *comm)
1: npde - Integer
    On entry: the number of PDEs in the system.
2: t - double
                            Input
    On entry: the current value of the independent variable t.
3: \mathbf{x - double Input}
    On entry: the current value of the space variable }x\mathrm{ .
```

4: $\quad \mathbf{u}[$ npde $]$ - const double
Input
On entry: $\mathbf{u}[i-1]$ contains the value of the component $U_{i}(x, t)$, for $i=1,2, \ldots$, npde.
5: udot[npde] - const double Input On entry: udot $[i-1]$ contains the value of the component $\frac{\partial U_{i}(x, t)}{\partial t}$, for $i=1,2, \ldots$, npde.

6: ux[npde] - const double
Input
On entry: $\mathbf{u x}[i-1]$ contains the value of the component $\frac{\partial U_{i}(x, t)}{\partial x}$, for $i=1,2, \ldots$, npde.
ncode - Integer
Input
On entry: the number of coupled ODEs in the system.
8: $\quad \mathbf{v}[$ ncode $]$ - const double
Input
On entry: if ncode $>0, \mathbf{v}[i-1]$ contains the value of the component $V_{i}(t)$, for $i=1,2, \ldots$, ncode.

9: $\quad \operatorname{vdot}[$ ncode $]$ - const double
Input
On entry: if ncode $>0, \operatorname{vdot}[i-1]$ contains the value of component $\dot{V}_{i}(t)$, for $i=1,2, \ldots$, ncode.

10: $\quad$ res[npde] - double
Output
On exit: $\mathbf{r e s}[i-1]$ must contain the $i$ th component of $G$, for $i=1,2, \ldots$, npde, where $G$ is defined as

$$
\begin{equation*}
G_{i}=\sum_{j=1}^{\text {npde }} P_{i, j} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} Q_{i, j} \dot{V}_{j} \tag{9}
\end{equation*}
$$

i.e., only terms depending explicitly on time derivatives, or

$$
\begin{equation*}
G_{i}=\sum_{j=1}^{\text {npde }} P_{i, j} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} Q_{i, j} \dot{V}_{j}+S_{i} \tag{10}
\end{equation*}
$$

i.e., all terms in equation (3).

The definition of $G$ is determined by the input value of ires.
11: $\quad$ ires - Integer *
Input/Output
On entry: the form of $G_{i}$ that must be returned in the array res.
ires $=-1$
Equation (9) must be used.
ires $=1$
Equation (10) must be used.
On exit: should usually remain unchanged. However, you may set ires to force the integration function to take certain actions, as described below:
ires $=2$
Indicates to the integrator that control should be passed back immediately to the calling function with the error indicator set to fail.code $=$ NE_USER_STOP.
ires $=3$
Indicates to the integrator that the current time step should be abandoned and a smaller time step used instead. You may wish to set ires $=3$ when a physically meaningless input or output value has been generated. If you consecutively set ires $=3$, then nag_pde_parab_1d_keller_ode_remesh (d03prc) returns to the calling function with the error $\overline{\text { indicator set to }} \overline{\text { fail.code }}=$ NE_FAILED_DERIV.

12: comm - Nag_Comm *
Pointer to structure of type Nag_Comm; the following members are relevant to pdedef.
user - double *
iuser - Integer *
p - Pointer
The type Pointer will be void *. Before calling nag_pde_parab_1d_keller_ ode_remesh (d03prc) you may allocate memory and initialize these pointers with various quantities for use by pdedef when called from nag_pde_parab_1d_kel ler_ode_remesh (d03prc) (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).

5: bndary - function, supplied by the user
External Function
bndary must evaluate the functions $G_{i}^{L}$ and $G_{i}^{R}$ which describe the boundary conditions, as given in (5) and (6).

The specification of bndary is:
void bndary (Integer npde, double $t$, Integer ibnd, Integer nobc, const double u[], const double udot[], Integer ncode, const double v[], const double vdot[], double res[], Integer *ires, Nag_Comm *comm)
1: npde - Integer Input
On entry: the number of PDEs in the system.
2: $\mathbf{t}$ - double $\quad$ Input
On entry: the current value of the independent variable $t$.
3: ibnd - Integer Input
On entry: specifies which boundary conditions are to be evaluated.
ibnd $=0$
bndary must compute the left-hand boundary condition at $x=a$.
ibnd $\neq 0$ bndary must compute of the right-hand boundary condition at $x=b$.

4: nobc - Integer Input
On entry: specifies the number $n_{a}$ of boundary conditions at the boundary specified by ibnd.

5: u[npde] - const double Input
On entry: $\mathbf{u}[i-1]$ contains the value of the component $U_{i}(x, t)$ at the boundary specified by ibnd, for $i=1,2, \ldots$, npde.

6: udot[npde] - const double
Input
On entry: udot $[i-1]$ contains the value of the component $\frac{\partial U_{i}(x, t)}{\partial t}$, for
$i=1,2, \ldots$, npde.
ncode - Integer
Input
On entry: the number of coupled ODEs in the system.
v[ncode] - const double
Input
On entry: if ncode $>0, \mathbf{v}[i-1]$ contains the value of the component $V_{i}(t)$, for $i=1,2, \ldots$, ncode.
vdot[ncode] - const double
Input
On entry: if ncode $>0, \operatorname{vdot}[i-1]$ contains the value of component $\dot{V}_{i}(t)$, for $i=1,2, \ldots$, ncode.

Note: $\operatorname{vdot}[i-1]$, for $i=1,2, \ldots$, ncode, may only appear linearly as in (11) and (12).
res[nobc] - double
Output
On exit: res $[i-1]$ must contain the $i$ th component of $G^{L}$ or $G^{R}$, depending on the value of ibnd, for $i=1,2, \ldots$, nobc, where $G^{L}$ is defined as

$$
\begin{equation*}
G_{i}^{L}=\sum_{j=1}^{\text {npde }} E_{i, j}^{L} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} H_{i, j}^{L} \dot{V}_{j}, \tag{11}
\end{equation*}
$$

i.e., only terms depending explicitly on time derivatives, or

$$
\begin{equation*}
G_{i}^{L}=\sum_{j=1}^{\text {npde }} E_{i, j}^{L} \frac{\partial U_{j}}{\partial t}+\sum_{j=1}^{\text {ncode }} H_{i, j}^{L} \dot{V}_{j}+K_{i}^{L}, \tag{12}
\end{equation*}
$$

i.e., all terms in equation (7), and similarly for $G_{i}^{R}$.

The definitions of $G^{L}$ and $G^{R}$ are determined by the input value of ires.
11: ires - Integer *
Input/Output
On entry: the form of $G_{i}^{L}$ ( or $G_{i}^{R}$ ) that must be returned in the array res.
ires $=-1$
Equation (11) must be used.
ires $=1$
Equation (12) must be used.
On exit: should usually remain unchanged. However, you may set ires to force the integration function to take certain actions as described below:
$\boldsymbol{\operatorname { i r e s }}=2$
Indicates to the integrator that control should be passed back immediately to the calling function with the error indicator set to fail.code $=$ NE_USER_STOP.
ires $=3$
Indicates to the integrator that the current time step should be abandoned and a smaller time step used instead. You may wish to set ires $=3$ when a physically meaningless input or output value has been generated. If you consecutively set ires $=3$, then nag_pde_parab_1d_keller_ode_remesh (d03prc) returns to the calling function with the error indicator set to fail.code = NE_FAILED_DERIV.

12: comm - Nag_Comm *
Pointer to structure of type Nag_Comm; the following members are relevant to bndary.
user - double *
iuser - Integer *
p - Pointer
The type Pointer will be void *. Before calling nag_pde_parab_1d_keller_ ode_remesh (d03prc) you may allocate memory and initialize these pointers with various quantities for use by bndary when called from nag_pde_parab_1d_kel ler_ode_remesh (d03prc) (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).
uvinit - function, supplied by the user
External Function
uvinit must supply the initial $\left(t=t_{0}\right)$ values of $U(x, t)$ and $V(t)$ for all values of $x$ in the interval $[a, b]$.

The specification of uvinit is:
void uvinit (Integer npde, Integer npts, Integer nxi, const double x[], const double xi[], double u[], Integer ncode, double v[], Nag_Comm *comm)
npde - Integer Input

On entry: the number of PDEs in the system.
2: npts - Integer Input
On entry: the number of mesh points in the interval $[a, b]$.
3: nxi - Integer Input
On entry: the number of ODE/PDE coupling points.
4: $\mathbf{x}[\mathbf{n p t s}]$ - const double Input
On entry: the current mesh. $\mathbf{x}[i-1]$ contains the value of $x_{i}$, for $i=1,2, \ldots$, npts.
5: $\quad \mathbf{x i}[\mathbf{n x i}]$ - const double
Input
On entry: if $\mathbf{n x i}>0, \mathbf{x i}[i-1]$ contains the $\mathrm{ODE} / \mathrm{PDE}$ coupling point, $\xi_{i}$, for $i=1,2, \ldots, \mathbf{n x i}$.

6: $\quad \mathbf{u}[$ npde $\times \mathbf{n p t s}]-$ double
Output
On exit: if nxi $>0, \mathbf{u}[\mathbf{n p d e} \times(j-1)+i-1]$ contains the value of the component $U_{i}\left(x_{j}, t_{0}\right)$, for $i=1,2, \ldots$, npde and $j=1,2, \ldots$, npts.

7: ncode - Integer
Input
On entry: the number of coupled ODEs in the system.
8: $\quad \mathbf{v}[$ ncode $]$ - double
Output
On exit: if ncode $>0, \mathbf{v}[i-1]$ must contain the value of component $V_{i}\left(t_{0}\right)$, for $i=1,2, \ldots$, ncode.

9: $\quad$ comm - Nag_Comm *
Pointer to structure of type Nag_Comm; the following members are relevant to uvinit.

```
user - double *
iuser - Integer *
p - Pointer
```

The type Pointer will be void *. Before calling nag_pde_parab_1d_keller_ ode_remesh (d03prc) you may allocate memory and initialize these pointers with various quantities for use by uvinit when called from nag_pde_parab_1d_keller_ ode_remesh (d03prc) (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).

7: $\quad \mathbf{u}[$ neqn $]$ - double
Input/Output
On entry: if $\mathbf{i n d}=1$, the value of $\mathbf{u}$ must be unchanged from the previous call.
On exit: $\mathbf{u}[$ npde $\times(j-1)+i-1]$ contains the computed solution $U_{i}\left(x_{j}, t\right)$, for $i=1,2, \ldots$, npde and $j=1,2, \ldots$, npts, evaluated at $t=\mathbf{t s}$.
npts - Integer
Input
On entry: the number of mesh points in the interval $[a, b]$.
Constraint: npts $\geq 3$.
9: $\quad \mathbf{x}[\mathbf{n p t s}]-$ double
Input/Output
On entry: the initial mesh points in the space direction. $\mathbf{x}[0]$ must specify the left-hand boundary, $a$, and $\mathbf{x}[$ npts -1$]$ must specify the right-hand boundary, $b$.
Constraint: $\mathbf{x}[0]<\mathbf{x}[1]<\cdots<\mathbf{x}[$ npts -1$]$.
On exit: the final values of the mesh points.
nleft - Integer
Input
On entry: the number $n_{a}$ of boundary conditions at the left-hand mesh point $\mathbf{x}[0]$.
Constraint: $0 \leq$ nleft $\leq$ npde.
11: ncode - Integer Input
On entry: the number of coupled ODE components.
Constraint: ncode $\geq 0$.

12: odedef - function, supplied by the user
External Function
odedef must evaluate the functions $R$, which define the system of ODEs, as given in (4).
If ncode $=0$, odedef will never be called and the NAG defined null void function pointer, NULLFN, can be supplied in the call to nag_pde_parab_1d_keller_ode_remesh (d03prc).

```
The specification of odedef is:
void odedef (Integer npde, double t, Integer ncode, const double v[],
    const double vdot[], Integer nxi, const double xi[],
    const double ucp[], const double ucpx[], const double ucpt[],
    double r[], Integer *ires, Nag_Comm *comm)
1: npde - Integer
                                    Input
    On entry: the number of PDEs in the system.
2: t - double
Input
On entry: the current value of the independent variable \(t\).
```

3: ncode - Intege
On entry: the number of coupled ODEs in the system.
v[ncode] - const double Input

On entry: if ncode $>0, \mathbf{v}[i-1]$ contains the value of the component $V_{i}(t)$, for $i=1,2, \ldots$, ncode.
$\operatorname{vdot}[$ ncode $]$ - const double
Input
On entry: if ncode $>0, \operatorname{vdot}[i-1]$ contains the value of component $\dot{V}_{i}(t)$, for $i=1,2, \ldots$, ncode.
nxi - Integer
Input
On entry: the number of ODE/PDE coupling points.
7: $\quad \mathbf{x i}[\mathbf{n x i}]$ - const double
Input
On entry: if $\mathbf{n x i}>0, \mathbf{x i}[i-1]$ contains the $\mathrm{ODE} / \mathrm{PDE}$ coupling point, $\xi_{i}$, for $i=1,2, \ldots, \mathbf{n x i}$.
ucp $[$ npde $\times \mathbf{n x i}]$ - const double
Input
On entry: if nxi $>0$, ucp[npde $\times(j-1)+i-1]$ contains the value of $U_{i}(x, t)$ at the coupling point $x=\xi_{j}$, for $i=1,2, \ldots$, npde and $j=1,2, \ldots$, nxi.
$\mathbf{u c p x}[$ npde $\times \mathbf{n x i}]$ - const double
Input
On entry: if $\mathbf{n x i}>0, \mathbf{u c p x}[\mathbf{n p d e} \times(j-1)+i-1]$ contains the value of $\frac{\partial U_{i}(x, t)}{\partial x}$ at the coupling point $x=\xi_{j}$, for $i=1,2, \ldots$, npde and $j=1,2, \ldots$, nxi.
ucpt $[$ npde $\times \mathbf{n x i}]$ - const double
Input
On entry: if nxi $>0$, $\mathbf{u c p t}[\mathbf{n p d e} \times(j-1)+i-1]$ contains the value of $\frac{\partial U_{i}}{\partial t}$ at the coupling point $x=\xi_{j}$, for $i=1,2, \ldots$, npde and $j=1,2, \ldots$, nxi.
r[ncode] - double
Output
On exit: if ncode $>0, \mathbf{r}[i-1]$ must contain the $i$ th component of $R$, for $i=1,2, \ldots$, ncode, where $R$ is defined as

$$
\begin{equation*}
R=-B \dot{V}-C U_{t}^{*} \tag{13}
\end{equation*}
$$

i.e., only terms depending explicitly on time derivatives, or

$$
\begin{equation*}
R=A-B \dot{V}-C U_{t}^{*} \tag{14}
\end{equation*}
$$

i.e., all terms in equation (4). The definition of $R$ is determined by the input value of ires.

12: $\quad$ ires - Integer *
Input/Output
On entry: the form of $R$ that must be returned in the array $\mathbf{r}$.
ires $=-1$
Equation (13) must be used.
ires $=1$
Equation (14) must be used.

On exit: should usually remain unchanged. However, you may reset ires to force the integration function to take certain actions, as described below:
ires $=2$
Indicates to the integrator that control should be passed back immediately to the calling function with the error indicator set to fail.code $=$ NE_USER_STOP. ires $=3$

Indicates to the integrator that the current time step should be abandoned and a smaller time step used instead. You may wish to set ires $=3$ when a physically meaningless input or output value has been generated. If you consecutively set ires $=3$, then nag_pde_parab_1d_keller_ode_remesh (d03prc) returns to the calling function with the error indicator set to fail.code $=$ NE_FAILED_DERIV.

13: comm - Nag_Comm *
Pointer to structure of type Nag_Comm; the following members are relevant to odedef.
user - double *
iuser - Integer *
p - Pointer
The type Pointer will be void *. Before calling nag_pde_parab_1d_keller_ ode_remesh (d03prc) you may allocate memory and initialize these pointers with various quantities for use by odedef when called from nag_pde_parab_1d_kel ler_ode_remesh (d03prc) (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).
nxi - Integer
Input
On entry: the number of $\mathrm{ODE} / \mathrm{PDE}$ coupling points.
Constraints:

> if ncode $=0, \mathbf{n x i}=0$
> if ncode $>0, \mathbf{n x i} \geq 0$

14: $\quad \mathbf{x i}[\mathrm{dim}]-$ const double
Input
Note: the dimension, dim, of the array $\mathbf{x i}$ must be at least $\max (1, \mathbf{n x i})$.
On entry: $\mathbf{x i}[i-1]$, for $i=1,2, \ldots, \mathbf{n x i}$, must be set to the ODE/PDE coupling points, $\xi_{i}$.
Constraint: $\mathbf{x}[0] \leq \mathbf{x i}[0]<\mathbf{x i}[1]<\cdots<\mathbf{x i}[\mathbf{n x i}-1] \leq \mathbf{x}[\mathbf{n p t s}-1]$.

15: neqn - Integer
Input
On entry: the number of ODEs in the time direction.
Constraint: neqn $=$ npde $\times$ npts + ncode.

16: $\quad \mathbf{r t o l}[d i m]-$ const double
Note: the dimension, dim, of the array rtol must be at least
1 when itol $=1$ or 2 ;
neqn when itol $=3$ or 4 .
On entry: the relative local error tolerance.
Constraint: $\mathbf{r t o l}[i-1] \geq 0.0$ for all relevant $i$.

17: $\quad \operatorname{atol}[\mathrm{dim}]$ - const double
Input
Note: the dimension, dim, of the array atol must be at least
1 when $\mathbf{i t o l}=1$ or 3 ;
neqn when $\mathbf{i t o l}=2$ or 4 .
On entry: the absolute local error tolerance.
Constraint: atol $[i-1] \geq 0.0$ for all relevant $i$.
Note: corresponding elements of rtol and atol cannot both be 0.0 .
18: itol - Integer
Input
A value to indicate the form of the local error test. itol indicates to nag_pde_parab_1d_keller_ ode_remesh (d03prc) whether to interpret either or both of rtol or atol as a vector or scalar. The error test to be satisfied is $\left\|e_{i} / w_{i}\right\|<1.0$, where $w_{i}$ is defined as follows:
On entry:

| itol | rtol | atol | $w_{i}$ |
| :---: | :---: | :---: | :---: |
| 1 | scalar | scalar | $\operatorname{rtol}[0] \times\|\mathbf{u}[i-1]\|+\mathbf{a t o l}[0]$ |
| 2 | scalar | vector | $\operatorname{rtol}[0] \times\|\mathbf{u}[i-1]\|+\mathbf{a t o l}[i-1]$ |
| 3 | vector | scalar | $\operatorname{rtol}[i-1] \times\|\mathbf{u}[i-1]\|+\mathbf{a t o l}[0]$ |
| 4 | vector | vector | $\operatorname{rtol}[i-1] \times\|\mathbf{u}[i-1]\|+\mathbf{a t o l}[i-1]$ |

In the above, $e_{i}$ denotes the estimated local error for the $i$ th component of the coupled PDE/ODE system in time, $\mathbf{u}[i-1]$, for $i=1,2, \ldots$, neqn.
The choice of norm used is defined by the argument norm.
Constraint: $\mathbf{i t o l}=1,2,3$ or 4.
19: norm - Nag_NormType
Input
On entry: the type of norm to be used.
norm = Nag_MaxNorm
Maximum norm.
norm $=$ Nag_TwoNorm $^{\text {_Th }}$
Averaged $L_{2}$ norm.
If $U_{\text {norm }}$ denotes the norm of the vector $\mathbf{u}$ of length neqn, then for the averaged $L_{2}$ norm

$$
U_{\text {norm }}=\sqrt{\frac{1}{\mathbf{n e q n}} \sum_{i=1}^{\text {neqn }}\left(U(i) / w_{i}\right)^{2}}
$$

while for the maximum norm

$$
U_{\text {norm }}=\max _{i}\left|\mathbf{u}[i-1] / w_{i}\right|
$$

See the description of itol for the formulation of the weight vector $w$.
Constraint: norm = Nag_MaxNorm or Nag_TwoNorm.
laopt - Nag_LinAlgOption
Input
On entry: the type of matrix algebra required.
laopt $=$ Nag_LinAlgFull
Full matrix methods to be used.
laopt $=$ Nag_LinAlgBand
Banded matrix methods to be used.

## laopt $=$ Nag_LinAlgSparse

Sparse matrix methods to be used.

## Constraint: laopt $=$ Nag_LinAlgFull, Nag_LinAlgBand or Nag_LinAlgSparse.

Note: you are recommended to use the banded option when no coupled ODEs are present (i.e., ncode $=0$ ).

21: algopt $[\mathbf{3 0}]$ - const double Input
On entry: may be set to control various options available in the integrator. If you wish to employ all the default options, then algopt $[0]$ should be set to 0.0 . Default values will also be used for any other elements of algopt set to zero. The permissible values, default values, and meanings are as follows:

## $\operatorname{algopt}[0]$

Selects the ODE integration method to be used. If algopt $[0]=1.0$, a BDF method is used and if algopt $[0]=2.0$, a Theta method is used. The default value is algopt $[0]=1.0$.
If algopt $[0]=2.0$, then $\operatorname{algopt}[i-1]$, for $i=2,3,4$, are not used.

```
algopt[1]
```

Specifies the maximum order of the BDF integration formula to be used. algopt[1] may be $1.0,2.0,3.0,4.0$ or 5.0 . The default value is algopt $[1]=5.0$.

## $\boldsymbol{\operatorname { a l g }} \boldsymbol{\operatorname { p o p t }}[2]$

Specifies what method is to be used to solve the system of nonlinear equations arising on each step of the BDF method. If algopt $[2]=1.0$ a modified Newton iteration is used and if algopt $[2]=2.0$ a functional iteration method is used. If functional iteration is selected and the integrator encounters difficulty, then there is an automatic switch to the modified Newton iteration. The default value is algopt $[2]=1.0$.

## $\operatorname{algopt}[3]$

Specifies whether or not the Petzold error test is to be employed. The Petzold error test results in extra overhead but is more suitable when algebraic equations are present, such as $P_{i, j}=0.0$, for $j=1,2, \ldots$, npde, for some $i$ or when there is no $\dot{V}_{i}(t)$ dependence in the coupled ODE system. If algopt $[3]=1.0$, then the Petzold test is used. If algopt $[3]=2.0$, then the Petzold test is not used. The default value is algopt $[3]=1.0$.
If algopt $[0]=1.0$, then $\operatorname{algopt}[i-1]$, for $i=5,6,7$, are not used.
$\operatorname{algopt}[4]$
Specifies the value of Theta to be used in the Theta integration method. $0.51 \leq \operatorname{algopt}[4] \leq 0.99$. The default value is algopt $[4]=0.55$.

## $\operatorname{algopt}[5]$

Specifies what method is to be used to solve the system of nonlinear equations arising on each step of the Theta method. If algopt $[5]=1.0$, a modified Newton iteration is used and if algopt $[5]=2.0$, a functional iteration method is used. The default value is $\operatorname{algopt}[5]=1.0$.

## $\operatorname{algopt}[6]$

Specifies whether or not the integrator is allowed to switch automatically between modified Newton and functional iteration methods in order to be more efficient. If $\operatorname{algopt}[6]=1.0$, then switching is allowed and if algopt $[6]=2.0$, then switching is not allowed. The default value is algopt $[6]=1.0$.

## $\operatorname{algopt}[10]$

Specifies a point in the time direction, $t_{\text {crit }}$, beyond which integration must not be attempted. The use of $t_{\text {crit }}$ is described under the argument itask. If algopt $[0] \neq 0.0$, a value of 0.0 , for algopt[10], say, should be specified even if itask subsequently specifies that $t_{\text {crit }}$ will not be used.

## $\operatorname{algopt}[11]$

Specifies the minimum absolute step size to be allowed in the time integration. If this option is not required, algopt[11] should be set to 0.0 .
$\boldsymbol{a l g o p t}[12]$
Specifies the maximum absolute step size to be allowed in the time integration. If this option is not required, algopt[12] should be set to 0.0 .

```
algopt[13]
```

Specifies the initial step size to be attempted by the integrator. If $\operatorname{algopt}[13]=0.0$, then the initial step size is calculated internally.

## $\operatorname{algopt}[14]$

Specifies the maximum number of steps to be attempted by the integrator in any one call. If algopt $[14]=0.0$, then no limit is imposed.

## $\operatorname{algopt}[22]$

Specifies what method is to be used to solve the nonlinear equations at the initial point to initialize the values of $U, U_{t}, V$ and $\dot{V}$. If $\operatorname{algopt}[22]=1.0$, a modified Newton iteration is used and if $\operatorname{algopt}[22]=2.0$, functional iteration is used. The default value is $\operatorname{algopt}[22]=1.0$.
$\operatorname{algopt}[28]$ and $\operatorname{algopt}[29]$ are used only for the sparse matrix algebra option, i.e., laopt $=$ Nag_LinAlgSparse.

## algopt[28]

Governs the choice of pivots during the decomposition of the first Jacobian matrix. It should lie in the range $0.0<\operatorname{algopt}[28]<1.0$, with smaller values biasing the algorithm towards maintaining sparsity at the expense of numerical stability. If algopt[28] lies outside this range then the default value is used. If the functions regard the Jacobian matrix as numerically singular then increasing algopt[28] towards 1.0 may help, but at the cost of increased fill-in. The default value is algopt $[28]=0.1$.

```
algopt[29]
```

Used as a relative pivot threshold during subsequent Jacobian decompositions (see algopt[28]) below which an internal error is invoked. algopt[29] must be greater than zero, otherwise the default value is used. If algopt[29] is greater than 1.0 no check is made on the pivot size, and this may be a necessary option if the Jacobian is found to be numerically singular (see algopt[28]). The default value is algopt $[29]=0.0001$.

## remesh - Nag_Boolean

Input
On entry: indicates whether or not spatial remeshing should be performed.

```
remesh = Nag_TRUE
```

Indicates that spatial remeshing should be performed as specified.
remesh $=$ Nag_FALSE
Indicates that spatial remeshing should be suppressed.
Note: remesh should not be changed between consecutive calls to nag_pde_parab_1d_keller_ ode_remesh (d03prc). Remeshing can be switched off or on at specified times by using appropriate values for the arguments nrmesh and trmesh at each call.
nxfix - Integer
Input
On entry: the number of fixed mesh points.
Constraint: $0 \leq \mathbf{n x f i x} \leq \mathbf{n p t s}-2$.
Note: the end points $\mathbf{x}[0]$ and $\mathbf{x}[\mathbf{n p t s}-1]$ are fixed automatically and hence should not be specified as fixed points.
$\mathbf{x f i x}[$ dim $]$ - const double Input
Note: the dimension, dim, of the array $\mathbf{x f i x}$ must be at least $\max (1, \mathbf{n x f i x})$.

On entry: $\mathbf{x f i x}[i-1]$, for $i=1,2, \ldots$, nxfix, must contain the value of the $x$ coordinate at the $i$ th fixed mesh point.
Constraint: $\mathbf{x f i x}[i-1]<\mathbf{x f i x}[i]$, for $i=1,2, \ldots, \mathbf{n x f i x}-1$, and each fixed mesh point must coincide with a user-supplied initial mesh point, that is $\mathbf{x f i x}[i-1]=\mathbf{x}[j-1]$ for some $j$, $2 \leq j \leq \mathbf{n p t s}-1$.
Note: the positions of the fixed mesh points in the array $\mathbf{x}$ remain fixed during remeshing, and so the number of mesh points between adjacent fixed points (or between fixed points and end points) does not change. You should take this into account when choosing the initial mesh distribution.
nrmesh - Integer
Input
On entry: indicates the form of meshing to be performed.
nrmesh $<0$
Indicates that a new mesh is adopted according to the argument dxmesh. The mesh is tested every $\mid$ nrmesh $\mid$ timesteps.
nrmesh $=0$
Indicates that remeshing should take place just once at the end of the first time step reached when $t>$ trmesh.
nrmesh $>0$
Indicates that remeshing will take place every nrmesh time steps, with no testing using dxmesh.

Note: nrmesh may be changed between consecutive calls to nag_pde_parab_1d_keller_ode_r emesh (d03prc) to give greater flexibility over the times of remeshing.
dxmesh - double
Input
On entry: determines whether a new mesh is adopted when nrmesh is set less than zero. A possible new mesh is calculated at the end of every $\mid$ nrmesh $\mid$ time steps, but is adopted only if

$$
x_{i}^{\text {new }}>x_{i}^{\text {old }}+\mathbf{d x m e s h} \times\left(x_{i+1}^{\text {old }}-x_{i}^{\text {old }}\right),
$$

or

$$
x_{i}^{\text {new }}<x_{i}^{\text {old }}-\mathbf{d x m e s h} \times\left(x_{i}^{\text {old }}-x_{i-1}^{\text {old }}\right) .
$$

dxmesh thus imposes a lower limit on the difference between one mesh and the next.
Constraint: dxmesh $\geq 0.0$.

27: trmesh - double
Input
On entry: specifies when remeshing will take place when nrmesh is set to zero. Remeshing will occur just once at the end of the first time step reached when $t$ is greater than trmesh.

Note: trmesh may be changed between consecutive calls to nag_pde_parab_1d_keller_ode_r emesh (d03prc) to force remeshing at several specified times.
ipminf - Integer
Input
On entry: the level of trace information regarding the adaptive remeshing.

## $\mathbf{i p m i n f}=0$

No trace information.
$\boldsymbol{\operatorname { i p m i n f }}=1$
Brief summary of mesh characteristics.
$\boldsymbol{i p m i n f}=2$
More detailed information, including old and new mesh points, mesh sizes and monitor function values.
Constraint: $\mathbf{i p m i n f}=0,1$ or 2.
xratio - double
Input
On entry: input bound on adjacent mesh ratio (greater than 1.0 and typically in the range 1.5 to 3.0). The remeshing functions will attempt to ensure that

$$
\left(x_{i}-x_{i-1}\right) / \mathbf{x r a t i o}<x_{i+1}-x_{i}<\mathbf{x r a t i o} \times\left(x_{i}-x_{i-1}\right)
$$

Suggested value: $\mathbf{x r a t i o}=1.5$.
Constraint: xratio $>1.0$.
con - double
Input
On entry: an input bound on the sub-integral of the monitor function $F^{\text {mon }}(x)$ over each space step. The remeshing functions will attempt to ensure that

$$
\int_{x_{1}}^{x_{i+1}} F^{\mathrm{mon}}(x) d x \leq \operatorname{con} \int_{x_{1}}^{x_{\mathrm{npts}}} F^{\mathrm{mon}}(x) d x
$$

(see Furzeland (1984)). con gives you more control over the mesh distribution e.g., decreasing con allows more clustering. A typical value is $2 /(\mathbf{n p t s}-1)$, but you are encouraged to experiment with different values. Its value is not critical and the mesh should be qualitatively correct for all values in the range given below.
Suggested value: con $=2.0 /(\mathbf{n p t s}-1)$.
Constraint: $0.1 /(\mathbf{n p t s}-1) \leq \mathbf{c o n} \leq 10.0 /(\mathbf{n p t s}-1)$.
31: monitf - function, supplied by the user

## External Function

monitf must supply and evaluate a remesh monitor function to indicate the solution behaviour of interest.
If ncode $=0$, monitf will never be called and the NAG defined null void function pointer, NULLFN, can be supplied in the call to nag_pde_parab_1d_keller_ode_remesh (d03prc).

```
The specification of monitf is:
void monitf (double t, Integer npts, Integer npde, const double x[],
    const double u[], double fmon[], Nag_Comm *comm)
1: t - double Input
    On entry: the current value of the independent variable t.
2: npts - Integer
    Input
    On entry: the number of mesh points in the interval [a,b].
3: npde - Integer Input
    On entry: the number of PDEs in the system.
4: }\quad\mathbf{x}[npts] - const double
    Input
    On entry: the current mesh. \mathbf{x}[i-1] contains the value of }\mp@subsup{x}{i}{}\mathrm{ , for }i=1,2,\ldots,npts
5: u[npde }\times\mathrm{ npts] - const double
                                    Input
        On entry: u[npde }\times(j-1)+i-1] contains the value of U U (x,t) at x=\mathbf{x}[j-1] and
        time t, for }i=1,2,\ldots,\mathrm{ npde and j=1,2,_., npts.
```

```
6: fmon[npts] - double
```

On exit: $\mathbf{f m o n}[i-1]$ must contain the value of the monitor function $F^{\mathrm{mon}}(x)$ at mesh point $x=\mathbf{x}[i-1]$.
Constraint: $\mathbf{f m o n}[i-1] \geq 0.0$.
comm - Nag_Comm *
Pointer to structure of type Nag_Comm; the following members are relevant to monitf.
user - double *
iuser - Integer *
p - Pointer
The type Pointer will be void *. Before calling nag_pde_parab_1d_keller_ ode_remesh (d03prc) you may allocate memory and initialize these pointers with various quantities for use by monitf when called from nag_pde_parab_1d_kel ler_ode_remesh (d03prc) (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).
rsave[Irsave] - double
Communication Array
If ind $=0$, rsave need not be set on entry.
If ind $=1$, rsave must be unchanged from the previous call to the function because it contains required information about the iteration.

Irsave - Integer
Input
On entry: the dimension of the array rsave. Its size depends on the type of matrix algebra selected.

If laopt $=$ Nag LinAlgFull, $\boldsymbol{l r s a v e} \geq$ neqn $\times$ neqn + neqn $+n w k r e s+$ lenode.
If laopt $=$ Nag_LinAlgBand, $\boldsymbol{l r s a v e} \geq(3 \times m l+m u+2) \times$ neqn $+n w k r e s+l e n o d e$.
If laopt $=$ Nag_LinAlgSparse, $\boldsymbol{I r s a v e} \geq 4 \times$ neqn $+11 \times$ neqn $/ 2+1+$ nwkres + lenode.
Where
$m l$ and $m u$ are the lower and upper half bandwidths given by $m l=\mathbf{n p d e}+$ nleft -1 such that $m u=2 \times$ npde - nleft -1 , for problems involving PDEs only; or
$m l=m u=$ neqn -1 , for coupled PDE/ODE problems.
$n w k r e s= \begin{cases}\text { npde } \times(3 \times \text { npde }+6 \times \mathbf{n x i}+\mathbf{n p t s}+15)+\mathbf{n x i}+\text { ncode }+7 \times \text { npts }+\mathbf{n x f i x}+1, & \text { when ncode }>0 \text { and } \mathbf{n x i}>0 ; \text { or } \\ \text { npde } \times(3 \times \text { npde }+\mathbf{n p t s}+21)+\text { ncode }+7 \times \text { npts }+\mathbf{n x f i x}+2, & \text { when ncode }>0 \text { and } \mathbf{n x i}=0 ; \text { or } \\ \text { npde } \times(3 \times \text { npde }+\mathbf{n p t s}+21)+7 \times \text { npts }+\mathbf{n x f i x}+3, & \text { when ncode }=0 .\end{cases}$
lenode $=\left\{\begin{array}{l}(6+\operatorname{int}(\operatorname{algopt}[1])) \times \operatorname{neqn}+50, \quad \text { when the BDF method is used; or } \\ 9\end{array}\right.$
Note: when using the sparse option, the value of Irsave may be too small when supplied to the integrator. An estimate of the minimum size of lrsave is printed on the current error message unit if itrace $>0$ and the function returns with fail.code $=\mathrm{NE}_{-}$INT_2.

34: isave[lisave] - Integer
Communication Array
If ind $=0$, isave need not be set.
If ind $=1$, isave must be unchanged from the previous call to the function because it contains required information about the iteration. In particular the following components of the array isave concern the efficiency of the integration:
isave[0]
Contains the number of steps taken in time.
isave[1]
Contains the number of residual evaluations of the resulting ODE system used. One such evaluation involves evaluating the PDE functions at all the mesh points, as well as one evaluation of the functions in the boundary conditions.
isave [2]
Contains the number of Jacobian evaluations performed by the time integrator.
isave [3]
Contains the order of the ODE method last used in the time integration.
isave[4]
Contains the number of Newton iterations performed by the time integrator. Each iteration involves residual evaluation of the resulting ODE system followed by a back-substitution using the $L U$ decomposition of the Jacobian matrix.

The rest of the array is used as workspace.
lisave - Integer
Input
On entry: the dimension of the array isave. Its size depends on the type of matrix algebra selected:

$$
\begin{aligned}
& \text { if laopt }=\text { Nag_LinAlgFull, lisave } \geq 25+\text { nxfix; } \\
& \text { if laopt }=\text { Nag_LinAlgBand, lisave } \geq \text { neqn }+25+\text { nxfix; } \\
& \text { if laopt }=\text { Nag_LinAlgSparse, lisave } \geq 25 \times \mathbf{n e q n}+25+\mathbf{n x f i x} .
\end{aligned}
$$

Note: when using the sparse option, the value of lisave may be too small when supplied to the integrator. An estimate of the minimum size of lisave is printed if itrace $>0$ and the function returns with fail.code $=$ NE_INT_2.
itask - Integer
Input
On entry: the task to be performed by the ODE integrator.
$\boldsymbol{i t a s k}=1$
Normal computation of output values $\mathbf{u}$ at $t=$ tout (by overshooting and interpolating).
$\boldsymbol{i t a s k}=2$
Take one step in the time direction and return.
itask $=3$
Stop at first internal integration point at or beyond $t=$ tout.

## $\boldsymbol{i t a s k}=4$

Normal computation of output values $\mathbf{u}$ at $t=$ tout but without overshooting $t=t_{\text {crit }}$ where $t_{\text {crit }}$ is described under the argument algopt.

## itask $=5$

Take one step in the time direction and return, without passing $t_{\text {crit }}$, where $t_{\text {crit }}$ is described under the argument algopt.

Constraint: $\mathbf{i t a s k}=1,2,3,4$ or 5.

37: itrace - Integer
Input
On entry: the level of trace information required from nag_pde_parab_1d_keller_ode_remesh (d03prc) and the underlying ODE solver as follows:
itrace $\leq-1$
No output is generated.
itrace $=0$
Only warning messages from the PDE solver are printed.
itrace $=1$
Output from the underlying ODE solver is printed. This output contains details of Jacobian entries, the nonlinear iteration and the time integration during the computation of the ODE system.

## itrace $=2$

Output from the underlying ODE solver is similar to that produced when itrace $=1$, except that the advisory messages are given in greater detail.
itrace $\geq 3$
The output from the underlying ODE solver is similar to that produced when itrace $=2$, except that the advisory messages are given in greater detail.

38: outfile - const char * Input
On entry: the name of a file to which diagnostic output will be directed. If outfile is NULL the diagnostic output will be directed to standard output.

39: ind - Integer *
Input/Output
On entry: indicates whether this is a continuation call or a new integration.
ind $=0$
Starts or restarts the integration in time.
ind $=1$
Continues the integration after an earlier exit from the function. In this case, only the arguments tout and fail and the remeshing arguments nrmesh, dxmesh, trmesh, xratio and con may be reset between calls to nag_pde_parab_1d_keller_ode_remesh (d03prc).
Constraint: ind $=0$ or 1 .
On exit: ind $=1$.

40: comm - Nag_Comm *
The NAG communication argument (see Section 2.3.1.1 in How to Use the NAG Library and its Documentation).

41: saved - Nag_D03_Save *
Communication Structure
saved must remain unchanged following a previous call to a Chapter d03 function and prior to any subsequent call to a Chapter d03 function.

42: fail - NagError * Input/Output
The NAG error argument (see Section 2.7 in How to Use the NAG Library and its Documentation).

## 6 Error Indicators and Warnings

## NE_ACC_IN_DOUBT

Integration completed, but small changes in atol or rtol are unlikely to result in a changed solution.

## NE_ALLOC_FAIL

Dynamic memory allocation failed.
See Section 2.3.1.2 in How to Use the NAG Library and its Documentation for further information.

## NE_BAD_PARAM

On entry, argument $\langle$ value $\rangle$ had an illegal value.

## NE_FAILED_DERIV

In setting up the ODE system an internal auxiliary was unable to initialize the derivative. This could be due to your setting ires $=3$ in pdedef or bndary.

## NE_FAILED_START

atol and rtol were too small to start integration.

## NE_FAILED_STEP

Error during Jacobian formulation for ODE system. Increase itrace for further details.
Repeated errors in an attempted step of underlying ODE solver. Integration was successful as far as $\mathbf{t s}$ : $\mathbf{t s}=\langle$ value $\rangle$.
Underlying ODE solver cannot make further progress from the point ts with the supplied values of atol and rtol. ts $=\langle$ value $\rangle$.

## NE_INCOMPAT_PARAM

On entry, con $=\langle$ value $\rangle$, npts $=\langle$ value $\rangle$.
Constraint: con $\leq 10.0 /($ npts -1$)$.
On entry, con $=\langle$ value $\rangle$, npts $=\langle$ value $\rangle$.
Constraint: con $\geq 0.1 /($ npts -1$)$.
On entry, the point $\mathbf{x f i x}[I-1]$ does not coincide with any $\mathbf{x}[J-1]: I=\langle$ value $\rangle$ and $\mathbf{x f i x}[I-1]=\langle$ value $\rangle$.

## NE_INT

ires set to an invalid value in call to pdedef, bndary, or odedef.
On entry, ind $=\langle$ value $\rangle$.
Constraint: ind $=0$ or 1 .
On entry, ipminf $=\langle$ value $\rangle$.
Constraint: $\operatorname{ipminf}=0,1$ or 2 .
On entry, itask $=\langle$ value $\rangle$.
Constraint: itask $=1,2,3,4$ or 5 .
On entry, itol $=\langle$ value $\rangle$.
Constraint: $\mathbf{i t o l}=1,2,3$ or 4.
On entry, ncode $=\langle$ value $\rangle$.
Constraint: ncode $\geq 0$.
On entry, nleft $=\langle$ value $\rangle$.
Constraint: nleft $\geq 0$.
On entry, npde $=\langle$ value $\rangle$.
Constraint: npde $\geq 1$.
On entry, npts $=\langle$ value $\rangle$.
Constraint: npts $\geq 3$.
On entry, nxfix $=\langle$ value $\rangle$.
Constraint: $n x f i x \geq 0$.

## NE_INT_2

On entry, corresponding elements atol $[I-1]$ and $\operatorname{rtol}[J-1]$ are both zero: $I=\langle$ value $\rangle$ and $J=\langle$ value $\rangle$.

On entry, lisave is too small: lisave $=\langle$ value $\rangle$. Minimum possible dimension: $\langle$ value $\rangle$.
On entry, Irsave is too small: Irsave $=\langle$ value $\rangle$. Minimum possible dimension: $\langle$ value $\rangle$.
On entry, ncode $=\langle$ value $\rangle$ and $\mathbf{n x i}=\langle$ value $\rangle$.
Constraint: nxi $=0$ when ncode $=0$.
On entry, ncode $=\langle$ value $\rangle$ and $\mathbf{n x i}=\langle$ value $\rangle$.
Constraint: nxi $\geq 0$ when ncode $>0$.
On entry, nleft $=\langle$ value $\rangle$, npde $=\langle$ value $\rangle$.
Constraint: nleft $\leq$ npde.
On entry, nxfix $=\langle$ value $\rangle, \mathbf{n p t s}=\langle$ value $\rangle$.
Constraint: nxfix $\leq$ npts -2 .
When using the sparse option lisave or Irsave is too small: lisave $=\langle$ value $\rangle$, Irsave $=\langle$ value $\rangle$.

## NE_INT_4

On entry, neqn $=\langle$ value $\rangle$, npde $=\langle$ value $\rangle$, npts $=\langle$ value $\rangle$ and ncode $=\langle$ value $\rangle$.
Constraint: neqn $=$ npde $\times$ npts + ncode.

## NE_INTERNAL_ERROR

An internal error has occurred in this function. Check the function call and any array sizes. If the call is correct then please contact NAG for assistance.

An unexpected error has been triggered by this function. Please contact NAG.
See Section 2.7.6 in How to Use the NAG Library and its Documentation for further information.
Serious error in internal call to an auxiliary. Increase itrace for further details.

## NE_ITER_FAIL

In solving ODE system, the maximum number of steps algopt[14] has been exceeded. $\operatorname{algopt}[14]=\langle$ value $\rangle$.

## NE_NO_LICENCE

Your licence key may have expired or may not have been installed correctly.
See Section 2.7.5 in How to Use the NAG Library and its Documentation for further information.

## NE_NOT_CLOSE_FILE

Cannot close file $\langle$ value $\rangle$.

## NE_NOT_STRICTLY_INCREASING

On entry, $I=\langle$ value $\rangle, \mathbf{x f i x}[I]=\langle$ value $\rangle$ and $\mathbf{x f i x}[I-1]=\langle$ value $\rangle$.
Constraint: $\mathbf{x f i x}[I]>\mathbf{x f i x}[I-1]$.
On entry, $I=\langle$ value $\rangle, \mathbf{x i}[I]=\langle$ value $\rangle$ and $\mathbf{x i}[I-1]=\langle$ value $\rangle$.
Constraint: $\mathbf{x i}[I]>\mathbf{x i}[I-1]$.
On entry, mesh points $\mathbf{x}$ appear to be badly ordered: $I=\langle$ value $\rangle, \mathbf{x}[I-1]=\langle$ value $\rangle, J=\langle$ value $\rangle$ and $\mathbf{x}[J-1]=\langle$ value $\rangle$.

## NE_NOT_WRITE_FILE

Cannot open file $\langle v a l u e\rangle$ for writing.

## NE_REAL

On entry, dxmesh $=\langle$ value $\rangle$.
Constraint: dxmesh $\geq 0.0$.

On entry, xratio $=\langle$ value $\rangle$.
Constraint: xratio $>1.0$.

## NE_REAL_2

On entry, at least one point in $\mathbf{x i}$ lies outside $[\mathbf{x}[0], \mathbf{x}[\mathbf{n p t s}-1]]: \mathbf{x}[0]=\langle$ value $\rangle$ and $\mathbf{x}[\mathbf{n p t s}-1]=\langle$ value $\rangle$.

On entry, tout $=\langle$ value $\rangle$ and $\mathbf{t s}=\langle$ value $\rangle$.
Constraint: tout $>$ ts.
On entry, tout $-\mathbf{t s}$ is too small: tout $=\langle$ value $\rangle$ and $\mathbf{t s}=\langle$ value $\rangle$.

## NE_REAL_ARRAY

On entry, $I=\langle$ value $\rangle$ and atol $[I-1]=\langle$ value $\rangle$. Constraint: atol $[I-1] \geq 0.0$.
On entry, $I=\langle$ value $\rangle$ and $\mathbf{r t o l}[I-1]=\langle$ value $\rangle$.
Constraint: $\operatorname{rtol}[I-1] \geq 0.0$.

## NE_REMESH_CHANGED

remesh has been changed between calls to nag_pde_parab_1d_keller_ode_remesh (d03prc).

## NE_SING_JAC

Singular Jacobian of ODE system. Check problem formulation.

## NE_USER_STOP

In evaluating residual of ODE system, ires $=2$ has been set in pdedef, bndary, or odedef. Integration is successful as far as ts: $\mathbf{t s}=\langle$ value $\rangle$.

## NE_ZERO_WTS

Zero error weights encountered during time integration.

## 7 Accuracy

nag_pde_parab_1d_keller_ode_remesh (d03prc) controls the accuracy of the integration in the time direction but not the accuracy of the approximation in space. The spatial accuracy depends on both the number of mesh points and on their distribution in space. In the time integration only the local error over a single step is controlled and so the accuracy over a number of steps cannot be guaranteed. You should therefore test the effect of varying the accuracy arguments, atol and rtol.

## 8 Parallelism and Performance

nag_pde_parab_1d_keller_ode_remesh (d03prc) is threaded by NAG for parallel execution in multithreaded implementations of the NAG Library.
nag_pde_parab_1d_keller_ode_remesh (d03prc) makes calls to BLAS and/or LAPACK routines, which may be threaded within the vendor library used by this implementation. Consult the documentation for the vendor library for further information.

Please consult the x06 Chapter Introduction for information on how to control and interrogate the OpenMP environment used within this function. Please also consult the Users' Note for your implementation for any additional implementation-specific information.

## 9 Further Comments

The Keller box scheme can be used to solve higher-order problems which have been reduced to firstorder by the introduction of new variables (see the example in Section 10). In general, a second-order problem can be solved with slightly greater accuracy using the Keller box scheme instead of a finite difference scheme (nag_pde_parab_1d_fd_ode_remesh (d03ppc) for example), but at the expense of increased CPU time due to the larger number of function evaluations required.
It should be noted that the Keller box scheme, in common with other central-difference schemes, may be unsuitable for some hyperbolic first-order problems such as the apparently simple linear advection equation $U_{t}+a U_{x}=0$, where $a$ is a constant, resulting in spurious oscillations due to the lack of dissipation. This type of problem requires a discretization scheme with upwind weighting (nag_pde_parab_1d_cd_ode_remesh (d03psc) for example), or the addition of a second-order artificial dissipation term.
The time taken depends on the complexity of the system, the accuracy requested, and the frequency of the mesh updates. For a given system with fixed accuracy and mesh-update frequency it is approximately proportional to neqn.

## 10 Example

This example is the first-order system

$$
\begin{aligned}
& \frac{\partial U_{1}}{\partial t}+\frac{\partial U_{1}}{\partial x}+\frac{\partial U_{2}}{\partial x}=0 \\
& \frac{\partial U_{2}}{\partial t}+4 \frac{\partial U_{1}}{\partial x}+\frac{\partial U_{2}}{\partial x}=0
\end{aligned}
$$

for $x \in[0,1]$ and $t \geq 0$.
The initial conditions are

$$
\begin{aligned}
U_{1}(x, 0) & =e^{x} \\
U_{2}(x, 0) & =x^{2}+\sin \left(2 \pi x^{2}\right)
\end{aligned}
$$

and the Dirichlet boundary conditions for $U_{1}$ at $x=0$ and $U_{2}$ at $x=1$ are given by the exact solution:

$$
\begin{aligned}
& U_{1}(x, t)=\frac{1}{2}\left\{e^{x+t}+e^{x-3 t}\right\}+\frac{1}{4}\left\{\sin \left(2 \pi(x-3 t)^{2}\right)-\sin \left(2 \pi(x+t)^{2}\right)\right\}+2 t^{2}-2 x t \\
& U_{2}(x, t)=e^{x-3 t}-e^{x+t}+\frac{1}{2}\left\{\sin \left(2 \pi(x-3 t)^{2}\right)+\sin \left(2 \pi(x+t)^{2}\right)\right\}+x^{2}+5 t^{2}-2 x t
\end{aligned}
$$

### 10.1 Program Text

```
/* nag_pde_parab_1d_keller_ode_remesh (d03prc) Example Program.
    *
    * NAGPRODCODE Version.
    *
    * Copyright 2016 Numerical Algorithms Group.
    *
    * Mark 26, 2016.
    */
#include <stdio.h>
#include <math.h>
#include <nag.h>
#include <nag_stdlib.h>
#include <nagd03.h>
#include <nagx01.h>
#ifdef __cplusplus
extern "C"
{
#endif
    static void NAG_CALL pdedef(Integer, double, double, const double[],
```

```
const double[], const double[], Integer,
const double[], const double[], double[],
Integer *, Nag_Comm *);
static void NAG_CALL bndary(Integer, double, Integer, Integer,
const double[], const double[], Integer,
const double[], const double[], double[],
Integer *, Nag_Comm *);
static void NAG_CALL uvinit(Integer, Integer, Integer, const double[],
const double[], double[], Integer, double[],
Nag_Comm *);
static void NAG_CALL monitf(double, Integer, Integer, const double[],
const double[], double[], Nag_Comm *);
#ifdef
```

$\qquad$

``` cplusplus
}
#endif
static void exact(double, Integer, Integer, double *, double *);
#define UE(I, J) ue[npde*((J) -1)+(I) -1]
#define U(I, J) u[npde*((J) -1)+(I) -1]
#define UOUT(I, J, K) uout[npde*(intpts*((K) -1)+(J) -1)+(I) -1]
int main(void)
{
    const Integer npde = 2, npts = 61, ncode = 0, nxi = 0, nxfix = 0, nleft = 1;
    const Integer itype = 1, intpts = 5, neqn = npde * npts + ncode;
    const Integer lisave = 25 + nxfix;
    const Integer nwkres = npde * (npts + 3 * npde + 21) + 7 * npts + nxfix + 3;
    const Integer lenode = 11 * neqn + 50, lrsave =
    neqn * neqn + neqn + nwkres + lenode;
    static double ruser[4] = { -1.0, -1.0, -1.0, -1.0 };
    double con, dxmesh, tout, trmesh, ts, xratio;
    Integer exit_status = 0, i, ind, ipminf, it, itask, itol, itrace, nrmesh;
    Nag_Boolean remesh, theta;
    double *algopt = 0, *atol = 0, *rsave = 0, *rtol = 0, *u = 0, *ue = 0;
    double *uout = 0, *x = 0, *xfix = 0, *xi = 0, *xout = 0;
    Integer *isave = 0;
    NagError fail;
    Nag_Comm comm;
    Nag_DO3_Save saved;
    INIT_FAIL(fail);
    printf("nag_pde_parab_1d_keller_ode_remesh (dO3prc) Example Program"
        " Results\n\n");
    /* For communication with user-supplied functions: */
    comm.user = ruser;
    /* Allocate memory */
    if (!(algopt = NAG_ALLOC(30, double)) ||
        !(atol = NAG_ALLOC(1, double)) ||
        !(rsave = NAG_ALLOC(lrsave, double)) ||
        !(rtol = NAG_ALLOC(1, double)) ||
        !(u = NAG_ALLOC(npde * npts, double)) ||
        !(ue = NAG_ALLOC(npde * npts, double)) ||
        !(uout = NAG_ALLOC(npde * intpts * itype, double)) ||
        !(x = NAG_ALLOC(npts, double)) ||
        !(xfix = NAG_ALLOC(1, double)) ||
        !(xi = NAG_ALLOC(1, double)) ||
        !(xout = NAG_ALLOC(intpts, double)) ||
        !(isave = NAG_ALLOC(lisave, Integer)))
    {
        printf("Allocation failure\n");
        exit_status = 1;
        goto END;
}
```

```
itrace = 0;
itol = 1;
atol[0] = 5.0e-5;
rtol[O] = atol[O];
printf(" Accuracy requirement =%12.3e", atol[0]);
printf(" Number of points = %3" NAG_IFMT "\n\n", npts);
/* Set remesh parameters */
remesh = Nag_TRUE;
nrmesh = 3;
dxmesh = 0.0;
trmesh = 0.0;
con = 5.0 / (npts - 1.0);
xratio = 1.2;
ipminf = 0;
printf(" Remeshing every %3" NAG_IFMT " time steps\n\n", nrmesh);
/* Initialize mesh */
for (i = 0; i < npts; ++i)
    x[i] = i/ (npts - 1.0);
xout[0] = 0.0;
xout[1] = 0.25;
xout[2] = 0.5;
xout[3] = 0.75;
xout[4] = 1.0;
printf(" x ");
for (i = 0; i < intpts; ++i) {
    printf("%10.4f", xout[i]);
    printf((i + 1) % 5 == 0 || i == 4 ? "\n" : " ");
}
printf("\n\n");
xi[0] = 0.0;
ind = 0;
itask = 1;
/* Set theta to TRUE if the Theta integrator is required */
theta = Nag_FALSE;
for (i = 0; i < 30; ++i)
    algopt[i] = 0.0;
if (theta) {
    algopt[0] = 2.0;
    algopt[5] = 2.0;
    algopt[6] = 1.0;
}
/* Loop over output value of t */
ts = 0.0;
for (it = 0; it < 5; ++it) {
    tout = 0.05 * (it + 1);
    /* nag_pde_parab_1d_keller_ode_remesh (d03prc).
    * General system of first-order PDEs, coupled DAEs, method
    * of lines, Keller box discretization, remeshing, one space
        * variable
        */
    nag_pde_parab_1d_keller_ode_remesh(npde, &ts, tout, pdedef, bndary,
                                    uvinit, u, npts, x, nleft, ncode,
                                    NULLFN, nxi, xi, neqn, rtol, atol,
                                    itol, Nag_TwoNorm, Nag_LinAlgFull,
                                    algopt, remesh, nxfix, xfix, nrmesh,
                                    dxmesh, trmesh, ipminf, xratio, con,
                                    monitf, rsave, lrsave, isave, lisave,
                                    itask, itrace, 0, &ind, &comm, &saved,
```


## \&fail);

```
    if (fail.code != NE_NOERROR) {
        printf("Error from "
                "nag_pde_parab_1d_keller_ode_remesh (d03prc).\n%s\n",
                fail.message);
    exit_status = 1;
    goto END;
}
/* Interpolate at output points */
/* nag_pde_interp_1d_fd (d03pzc). PDEs, spatial interpolation with
    * nag_pde_parab_1d_keller_ode_remesh (d03prc).
    */
nag_pde_interp_1d_fd(npde, 0, u, npts, x, xout, intpts, itype, uout,
                                    &fail);
    if (fail.code != NE_NOERROR) {
    printf("Error from nag_pde_interp_1d_fd (d03pzc).\n%s\n", fail.message);
    exit_status = 1;
    goto END;
}
/* Check against exact solution */
exact(ts, npde, intpts, xout, ue);
printf(" t = %6.3f\n", ts);
printf(" Approx u1");
for (i = 1; i <= intpts; ++i) {
        printf("%10.4f", UOUT(1, i, 1));
        printf(i % 5 == 0 || i == 5 ? "\n" : "");
    }
printf(" Exact u1");
for (i = 1; i <= 5; ++i) {
        printf("%10.4f", UE(1, i));
        printf(i % 5 == 0 || i == 5 ? "\n" : "");
    }
    printf(" Approx u2");
    for (i = 1; i <= 5; ++i) {
        printf("%10.4f", UOUT(2, i, 1));
        printf(i % 5 == 0 || i == 5 ? "\n" : "");
    }
    printf(" Exact u2");
    for (i = 1; i <= 5; ++i) {
        printf("%10.4f", UE(2, i));
        printf(i % 5 == 0 || i == 5 ? "\n" : "");
    }
    printf("\n");
}
printf(" Number of integration steps in time = %6" NAG_IFMT "\n", isave[0]);
printf(" Number of function evaluations = %6" NAG_IFMT "\n", isave[1]);
printf(" Number of Jacobian evaluations =%6" NAG_IFMT "\n", isave[2]);
printf(" Number of iterations = %6" NAG_IFMT "\n\n", isave[4]);
NAG_FREE(algopt);
NAG_FREE(atol);
NAG_FREE(rsave);
NAG_FREE(rtol);
NAG_FREE(u);
```

END:

```
    NAG_FREE(ue);
    NAG_FREE(uout);
    NAG_FREE(x);
    NAG_FREE(xfix);
    NAG_FREE(xi);
    NAG_FREE(xout);
    NAG_FREE(isave);
    return exit_status;
}
static void NAG_CALL uvinit(Integer npde, Integer npts, Integer nxi,
                                    const double x[], const double xi[], double u[],
                                    Integer ncode, double v[], Nag_Comm *comm)
{
    Integer i;
    if (comm->user[0] == -1.0) {
        printf("(User-supplied callback uvinit, first invocation.)\n");
        comm->user[0] = 0.0;
    }
    for (i = 1; i <= npts; ++i) {
        U(1, i) = exp(x[i - 1]);
        U(2, i) = x[i - 1] * x[i - 1] + sin(2.0 * nag_pi * (x[i - 1] * x[i - 1]));
    }
    return;
}
static void NAG_CALL pdedef(Integer npde, double t, double x,
                                    const double u[], const double udot[],
                                    const double ux[], Integer ncode,
                                    const double v[], const double vdot[],
                                    double res[], Integer *ires, Nag_Comm *comm)
{
    if (comm->user[1] == -1.0) {
        printf("(User-supplied callback pdedef, first invocation.)\n");
        comm->user[1] = 0.0;
    }
    if (*ires == -1) {
        res[0] = udot[0];
        res[1] = udot[1];
    }
    else {
        res[0] = udot[0] + ux[0] + ux[1];
        res[1] = udot[1] + 4.0 * ux[0] + ux[1];
    }
    return;
}
static void NAG_CALL bndary(Integer npde, double t, Integer ibnd,
                    Integer nobc, const double u[],
                    const double udot[], Integer ncode,
                    const double v[], const double vdot[],
                    double res[], Integer *ires, Nag_Comm *comm)
{
    double pp;
    if (comm->user[2] == -1.0) {
        printf("(User-supplied callback bndary, first invocation.)\n");
        comm->user[2] = 0.0;
}
    pp = 2.0 * nag_pi;
    if (ibnd == 0) {
        if (*ires == -1) {
            res[0] = 0.0;
        }
        else {
            res[0] = u[0] - 0.5 * (exp(t) + exp(-3.0 * t))
                    -0.25 * ( sin(9.0 * pp * t * t) - sin(pp * t * t)) - 2.0 * t * t;
```

```
        }
    }
    else {
        if (*ires == -1) {
            res[0] = 0.0;
        }
        else {
            res[0] = u[1] - (exp(1.0 - 3.0 * t) - exp(1.0 + t) +
                        0.5 * sin(pp * (1.0 - 3.0 * t) * (1.0 - 3.0 * t)) +
                        0.5 * sin(pp * (1.0 + t) * (1.0 + t))
                        + 1.0 + 5.0 * t * t - 2.0 * t);
        }
    }
    return;
}
static void NAG_CALL monitf(double t, Integer npts, Integer npde,
                    const double x[], const double u[], double fmon[],
                    Nag_Comm *comm)
{
    double d2x1, d2x2, h1, h2, h3;
    Integer i;
    if (comm->user[3] == -1.0) {
        printf("(User-supplied callback monitf, first invocation.)\n");
        comm->user[3] = 0.0;
    }
    for (i = 2; i <= npts - 1; ++i) {
        h1 = x[i - 1] - x[i - 2];
        h2 = x[i] - x[i - 1];
        h3 = 0.5 * (x[i] - x[i - 2]);
        /* Second derivatives */
        d2x1 = fabs(((U(1, i + 1) - U(1, i)) / h2 -
                (U(1, i) - U(1, i - 1)) / h1) / h3);
        d2x2 = fabs(((U(2, i + 1) - U(2, i)) / h2 -
                (U(2, i) - U(2, i - 1)) / h1) / h3);
            fmon[i - 1] = d2x1;
        if (d2x2 > d2x1)
            fmon[i - 1] = d2x2;
    }
    fmon[0] = fmon[1];
    fmon[npts - 1] = fmon[npts - 2];
    return;
}
static void exact(double t, Integer npde, Integer npts, double *x, double *u)
{
    /* Exact solution (for comparison purposes) */
    double pp;
    Integer i;
    pp = 2.0 * nag_pi;
    for (i = 1; i <= npts; ++i) {
        U(1, i) = 0.5 * (exp(x[i - 1] + t) + exp(x[i - 1] - 3.0 * t)) +
                        0.25 * (sin(pp * (x[i - 1] - 3.0 * t) * (x[i - 1] - 3.0 * t)) -
                                    sin(pp * (x[i - 1] + t) * (x[i - 1] + t)) ) +
                2.0 * t * t - 2.0 * x[i - 1] * t;
        U(2, i) = exp(x[i - 1] - 3.0* t) - exp(x[i - 1] + t) +
                        0.5* ( sin(pp* *(x[i - 1] - 3.0* t)* (x[i - 1] - 3.0*t))) +
                                    sin(pp * ((x[i - 1] + t) * (x[i - 1] + t)))) +
            x[i - 1] * x[i - 1] + 5.0 * t * t - 2.0* *[i - 1] * t;
    }
    return;
}
```


### 10.2 Program Data

None.

### 10.3 Program Results



## Example Program

Solution of First-order System using Moving Mesh
$U(1, x, t)$


Solution of First-order System using Moving Mesh
$U(2, x, t)$


