

D03PWF – NAG Fortran Library Routine Document

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

D03PWF calculates a numerical flux function using a modified HLL (Harten-Lax-van Leer) Approximate Riemann Solver for the Euler equations in conservative form. The routine is designed primarily for use with the upwind discretisation routines D03PFF, D03PLF or D03PSF, but may also be applicable to other conservative upwind schemes requiring numerical flux functions.

2 Specification

```
SUBROUTINE D03PWF(ULEFT, URIGHT, GAMMA, FLUX, IFAIL)
INTEGER           IFAIL
real              ULEFT(3), URIGHT(3), GAMMA, FLUX(3)
```

3 Description

D03PWF calculates a numerical flux function at a single spatial point using a modified HLL (Harten-Lax-van Leer) Approximate Riemann Solver (see [1] [2] [3]) for the Euler equations (for a perfect gas) in conservative form. The user must supply the *left* and *right* solution values at the point where the numerical flux is required, i.e., the initial left and right states of the Riemann problem defined below. In the routines D03PFF, D03PLF and D03PSF, the left and right solution values are derived automatically from the solution values at adjacent spatial points and supplied to the subroutine argument NUMFLX from which the user may call D03PWF. The Euler equations for a perfect gas in conservative form are:

$$\frac{\partial U}{\partial t} + \frac{\partial F}{\partial x} = 0, \quad (1)$$

with

$$U = \begin{bmatrix} \rho \\ m \\ e \end{bmatrix} \quad \text{and} \quad F = \begin{bmatrix} m \\ \frac{m^2}{\rho} + (\gamma - 1) \left(e - \frac{m^2}{2\rho} \right) \\ \frac{me}{\rho} + \frac{m}{\rho} (\gamma - 1) \left(e - \frac{m^2}{2\rho} \right) \end{bmatrix}, \quad (2)$$

where ρ is the density, m is the momentum, e is the specific total energy and γ is the (constant) ratio of specific heats. The pressure p is given by

$$p = (\gamma - 1) \left(e - \frac{\rho u^2}{2} \right), \quad (3)$$

where $u = m/\rho$ is the velocity.

The routine calculates an approximation to the numerical flux function $F(U_L, U_R) = F(U^*(U_L, U_R))$, where $U = U_L$ and $U = U_R$ are the left and right solution values, and $U^*(U_L, U_R)$ is the intermediate state $\omega(0)$ arising from the similarity solution $U(y, t) = \omega(y/t)$ of the Riemann problem defined by

$$\frac{\partial U}{\partial t} + \frac{\partial F}{\partial y} = 0, \quad (4)$$

with U and F as in (2), and initial piecewise constant values $U = U_L$ for $y < 0$ and $U = U_R$ for $y > 0$. The spatial domain is $-\infty < y < \infty$, where $y = 0$ is the point at which the numerical flux is required.

4 References

- [1] Toro E F (1996) *Riemann Solvers and Upwind Methods for Fluid Dynamics* Springer-Verlag

- [2] Toro E F (1992) The weighted average flux method applied to the Euler equations *Phil. Trans. R. Soc. Lond.* **A341** 499–530
- [3] Toro E F, Spruce M and Spears W (1994) Restoration of the contact surface in the HLL Riemann solver *J. Shock Waves* **4** 25–34

5 Parameters

- 1: ULEFT(3) — **real** array *Input*
On entry: ULEFT(i) must contain the left value of the component U_i for $i = 1,2,3$. That is, ULEFT(1) must contain the left value of ρ , ULEFT(2) must contain the left value of m and ULEFT(3) must contain the left value of e .
- 2: URIGHT(3) — **real** array *Input*
On entry: URIGHT(i) must contain the right value of the component U_i for $i = 1,2,3$. That is, URIGHT(1) must contain the right value of ρ , URIGHT(2) must contain the right value of m and URIGHT(3) must contain the right value of e .
- 3: GAMMA — **real** *Input*
On entry: the ratio of specific heats γ .
Constraint: GAMMA > 0.0.
- 4: FLUX(3) — **real** array *Output*
On exit: FLUX(i) contains the numerical flux component \hat{F}_i for $i = 1,2,3$.
- 5: IFAIL — INTEGER *Input/Output*
On entry: IFAIL must be set to 0, -1 or 1. For users not familiar with this parameter (described in Chapter P01) the recommended value is 0.

Note. If the left and/or right values of ρ or p (from (3)) are found to be negative, then the routine will terminate with an error exit (IFAIL = 2). If the routine is being called from the user-supplied subroutine NUMFLX in D03PFF etc., then a **soft fail** option (IFAIL = 1 or -1) is recommended so that a recalculation of the current time step can be forced using the IRES parameter.

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

6 Error Indicators and Warnings

Errors detected by the routine:

IFAIL = 1

On entry, GAMMA ≤ 0.0 .

IFAIL = 2

On entry, the left and/or right density or derived pressure value is less than 0.0.

7 Accuracy

The routine performs an exact calculation of the HLL numerical flux function, and so the result will be accurate to machine precision.

8 Further Comments

The routine must only be used to calculate the numerical flux for the Euler equations in exactly the form given by (2), with ULEFT(i) and URIGHT(i) containing the left and right values of ρ, m and e for $i = 1,2,3$ respectively. The time taken by the routine is independent of the input parameters.

9 Example

This example uses D03PLF and D03PWF to solve the Euler equations in the domain $0 \leq x \leq 1$ for $0 < t \leq 0.035$ with initial conditions for the primitive variables $\rho(x, t)$, $u(x, t)$ and $p(x, t)$ given by

$$\begin{aligned}\rho(x, 0) &= 5.99924, & u(x, 0) &= 19.5975, & p(x, 0) &= 460.894, & \text{for } x < 0.5, \\ \rho(x, 0) &= 5.99242, & u(x, 0) &= -6.19633, & p(x, 0) &= 46.095, & \text{for } x > 0.5.\end{aligned}$$

This test problem is taken from [1] and its solution represents the collision of two strong shocks travelling in opposite directions, consisting of a left facing shock (travelling slowly to the right), a right travelling contact discontinuity and a right travelling shock wave. There is an exact solution to this problem (see [1]) but the calculation is lengthy and has therefore been omitted.

9.1 Program Text

```
*      D03PWF Example Program Text
*      Mark 18 Release. NAG Copyright 1997.
*      .. Parameters ..
  INTEGER           NIN, NOUT
  PARAMETER        (NIN=5,NOUT=6)
  INTEGER           NPDE, NPTS, NCODE, NXI, NEQN, NIW, NWKRES,
+                  LENODE, MLU, NW
  PARAMETER        (NPDE=3,NPTS=141,NCODE=0,NXI=0,
+                  NEQN=NPDE*NPTS+NCODE,NIW=NEQN+24,
+                  NWKRES=NPDE*(2*NPTS+3*NPDE+32)+7*NPTS+4,
+                  LENODE=9*NEQN+50,MLU=3*NPDE-1,NW=(3*MLU+1)
+                  *NEQN+NWKRES+LENODE)
*      .. Scalars in Common ..
  real              ELO, ERO, GAMMA, RLO, RR0, ULO, UR0
*      .. Local Scalars ..
  real              D, P, TOUT, TS, V
  INTEGER          I, IFAIL, IND, ITASK, ITOL, ITRACE, K
  CHARACTER         LAOPT, NORM
*      .. Local Arrays ..
  real              ALGOPT(30), ATOL(1), RTOL(1), U(NPDE,NPTS),
+                  UE(3,9), W(NW), X(NPTS), XI(1)
  INTEGER          IW(NIW)
*      .. External Subroutines ..
  EXTERNAL          BNDARY, D03PEK, D03PLF, D03PLP, NUMFLX
*      .. Common blocks ..
  COMMON            /INIT/ELO, ERO, RLO, RR0, ULO, UR0
  COMMON            /PARAMS/GAMMA
*      .. Executable Statements ..
  WRITE (NOUT,*) 'D03PWF Example Program Results'
*      Skip heading in data file
  READ (NIN,*)
*
*      Problem parameters
*
  GAMMA = 1.4e0
  RLO = 5.99924e0
  RRO = 5.99242e0
  ULO = 5.99924e0*19.5975e0
  UR0 = -5.99242e0*6.19633e0
  ELO = 460.894e0/(GAMMA-1.0e0) + 0.5e0*RL0*19.5975e0**2
  ERO = 46.095e0/(GAMMA-1.0e0) + 0.5e0*RR0*6.19633e0**2
*
```

```

*      Initialise mesh
*
      DO 20 I = 1, NPTS
         X(I) = 1.0e0*(I-1.0e0)/(NPTS-1.0e0)
20 CONTINUE
*
*      Initial values
*
      DO 40 I = 1, NPTS
         IF (X(I).LT.0.5e0) THEN
            U(1,I) = RL0
            U(2,I) = UL0
            U(3,I) = EL0
         ELSE IF (X(I).EQ.0.5e0) THEN
            U(1,I) = 0.5e0*(RL0+RR0)
            U(2,I) = 0.5e0*(UL0+UR0)
            U(3,I) = 0.5e0*(EL0+ER0)
         ELSE
            U(1,I) = RR0
            U(2,I) = UR0
            U(3,I) = ER0
         END IF
40 CONTINUE
*
      ITRACE = 0
      ITOL = 1
      NORM = '2'
      ATOL(1) = 0.5e-2
      RTOL(1) = 0.5e-3
      XI(1) = 0.0e0
      LAOPT = 'B'
      IND = 0
      ITASK = 1
      DO 60 I = 1, 30
         ALGOPT(I) = 0.0e0
60 CONTINUE
*
*      Theta integration
*
      ALGOPT(1) = 2.0e0
      ALGOPT(6) = 2.0e0
      ALGOPT(7) = 2.0e0
*
*      Max. time step
*
      ALGOPT(13) = 0.5e-2
*
      TS = 0.0e0
      TOUT = 0.035e0
      IFAIL = 0
*
      CALL D03PLF(NPDE,TS,TOUT,D03PLP,NUMFLX,BNDARY,U,NPTS,X,NCODE,
      +           D03PEK,NXI,XI,NEQN,RTOL,ATOL,ITOL,NORM,LAOPT,ALGOPT,W,
      +           NW,IW,NIW,ITASK,ITRACE,IND,IFAIL)
*
      WRITE (NOUT,99998) TS
      WRITE (NOUT,99999)

```

```

*
*      Read exact data at output points
*
      DO 80 I = 1, 9
          READ (NIN,*) UE(1,I), UE(2,I), UE(3,I)
80 CONTINUE
*
*      Calculate density, velocity and pressure
*
      K = 0
      DO 100 I = 15, NPTS - 14, 14
          D = U(1,I)
          V = U(2,I)/D
          P = D*(GAMMA-1.0e0)*(U(3,I)/D-0.5e0*V**2)
          K = K + 1
          WRITE (NOUT,99996) X(I), D, UE(1,K), V, UE(2,K), P, UE(3,K)
100 CONTINUE
*
      WRITE (NOUT,99997) IW(1), IW(2), IW(3), IW(5)
      STOP
*
99999 FORMAT (4X,'X',6X,'APPROX D',3X,'EXACT D',4X,'APPROX V',3X,'EXAC',
+           'T V',4X,'APPROX P',3X,'EXACT P')
99998 FORMAT ('/ T = ',F6.3,/)
99997 FORMAT ('/ Number of integration steps in time = ',I6,' Number ',
+           'of function evaluations = ',I6,' Number of Jacobian ',
+           'evaluations = ',I6,' Number of iterations = ',I6,/)

99996 FORMAT (1X,e8.2,6(1X,e10.4))
      END
*
      SUBROUTINE BNDARY(NPDE,NPTS,T,X,U,NCODE,V,VDOT,IBND,G,IRES)
* .. Scalar Arguments ..
      real             T
      INTEGER          IBND, IRES, NCODE, NPDE, NPTS
* .. Array Arguments ..
      real             G(NPDE), U(NPDE,NPTS), V(*), VDOT(*), X(NPTS)
* .. Scalars in Common ..
      real             ELO, ERO, RLO, RRO, ULO, URO
* .. Common blocks ..
      COMMON            /INIT/ELO, ERO, RLO, RRO, ULO, URO
* .. Executable Statements ..
      IF (IBND.EQ.0) THEN
          G(1) = U(1,1) - RLO
          G(2) = U(2,1) - ULO
          G(3) = U(3,1) - ELO
      ELSE
          G(1) = U(1,NPTS) - RRO
          G(2) = U(2,NPTS) - URO
          G(3) = U(3,NPTS) - ERO
      END IF
      RETURN
      END
*
      SUBROUTINE NUMFLX(NPDE,T,X,NCODE,V,ULEFT,URIGHT,FLUX,IRES)
* .. Scalar Arguments ..
      real             T, X
      INTEGER          IRES, NCODE, NPDE

```

```

*      .. Array Arguments ..
*      real           FLUX(NPDE), ULEFT(NPDE), URIGHT(NPDE), V(*)
*      .. Scalars in Common ..
*      real           GAMMA
*      .. Local Scalars ..
*      INTEGER         IFAIL
*      .. External Subroutines ..
*      EXTERNAL        D03PWF
*      .. Common blocks ..
*      COMMON          /PARAMS/GAMMA
*      .. Save statement ..
*      SAVE            /PARAMS/
*      .. Executable Statements ..
*
IFAIL = 0
CALL D03PWF(ULEFT,URIGHT,GAMMA,FLUX,IFAIL)
RETURN
END

```

9.2 Program Data

D03PWF Example Program Data

| | | |
|------------|------------|------------|
| 0.5999E+01 | 0.1960E+02 | 0.4609E+03 |
| 0.1428E+02 | 0.8690E+01 | 0.1692E+04 |
| 0.1428E+02 | 0.8690E+01 | 0.1692E+04 |
| 0.1428E+02 | 0.8690E+01 | 0.1692E+04 |
| 0.3104E+02 | 0.8690E+01 | 0.1692E+04 |

9.3 Program Results

D03PWF Example Program Results

T = 0.035

| X | APPROX D | EXACT D | APPROX V | EXACT V | APPROX P | EXACT P |
|----------|------------|------------|------------|------------|------------|------------|
| 0.10E+00 | 0.5999E+01 | 0.5999E+01 | 0.1960E+02 | 0.1960E+02 | 0.4609E+03 | 0.4609E+03 |
| 0.20E+00 | 0.5999E+01 | 0.5999E+01 | 0.1960E+02 | 0.1960E+02 | 0.4609E+03 | 0.4609E+03 |
| 0.30E+00 | 0.5999E+01 | 0.5999E+01 | 0.1960E+02 | 0.1960E+02 | 0.4609E+03 | 0.4609E+03 |
| 0.40E+00 | 0.5999E+01 | 0.5999E+01 | 0.1960E+02 | 0.1960E+02 | 0.4609E+03 | 0.4609E+03 |
| 0.50E+00 | 0.5999E+01 | 0.5999E+01 | 0.1960E+02 | 0.1960E+02 | 0.4609E+03 | 0.4609E+03 |
| 0.60E+00 | 0.1422E+02 | 0.1428E+02 | 0.8658E+01 | 0.8690E+01 | 0.1687E+04 | 0.1692E+04 |
| 0.70E+00 | 0.1426E+02 | 0.1428E+02 | 0.8670E+01 | 0.8690E+01 | 0.1688E+04 | 0.1692E+04 |
| 0.80E+00 | 0.1944E+02 | 0.1428E+02 | 0.8678E+01 | 0.8690E+01 | 0.1691E+04 | 0.1692E+04 |
| 0.90E+00 | 0.3100E+02 | 0.3104E+02 | 0.8676E+01 | 0.8690E+01 | 0.1687E+04 | 0.1692E+04 |

Number of integration steps in time = 699
 Number of function evaluations = 1714
 Number of Jacobian evaluations = 1
 Number of iterations = 2