NAG Fortran Library Routine Document

D01ANF

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

D01ANF calculates an approximation to the sine or the cosine transform of a function g over [a, b]:

$$I = \int_{a}^{b} g(x) \sin(\omega x) dx$$
 or $I = \int_{a}^{b} g(x) \cos(\omega x) dx$

(for a user-specified value of ω).

2 Specification

```
      SUBROUTINE DO1ANF(G, A, B, OMEGA, KEY, EPSABS, EPSREL, RESULT, ABSERR,

      1
      W, LW, IW, LIW, IFAIL)

      INTEGER
      KEY, LW, IW(LIW), LIW, IFAIL

      real
      G, A, B, OMEGA, EPSABS, EPSREL, RESULT, ABSERR, W(LW)

      EXTERNAL
      G
```

3 Description

D01ANF is based upon the QUADPACK routine QFOUR (Piessens *et al.* (1983)). It is an adaptive routine, designed to integrate a function of the form g(x)w(x), where w(x) is either $\sin(\omega x)$ or $\cos(\omega x)$. If a sub-interval has length

 $L = |b - a|2^{-l}$

then the integration over this sub-interval is performed by means of a modified Clenshaw–Curtis procedure (Piessens and Branders (1975)) if $L\omega > 4$ and $l \le 20$. In this case a Chebyshev-series approximation of degree 24 is used to approximate g(x), while an error estimate is computed from this approximation together with that obtained using Chebyshev-series of degree 12. If the above conditions do not hold then Gauss 7-point and Kronrod 15-point rules are used. The algorithm, described in Piessens *et al.* (1983), incorporates a global acceptance criterion (as defined in Malcolm and Simpson (1976)) together with the ϵ -algorithm (Wynn (1956)) to perform extrapolation. The local error estimation is described in Piessens *et al.* (1983).

4 **References**

Malcolm M A and Simpson R B (1976) Local versus global strategies for adaptive quadrature *ACM Trans. Math. Software* **1** 129–146

Piessens R and Branders M (1975) Algorithm 002. Computation of oscillating integrals J. Comput. Appl. Math. 1 153–164

Piessens R, de Doncker-Kapenga E, Überhuber C and Kahaner D (1983) QUADPACK, A Subroutine Package for Automatic Integration Springer-Verlag

Wynn P (1956) On a device for computing the $e_m(S_n)$ transformation Math. Tables Aids Comput. 10 91–96

5 Parameters

1: G - real FUNCTION, supplied by the user.

External Procedure

G must return the value of the function g at a given point.

Its specification is:

	real FUNCTION G(X)		
	real 2	X	
1:	X – real		Input
	On entry: the point	at which the function g must be evaluated.	

G must be declared as EXTERNAL in the (sub)program from which D01ANF is called. Parameters denoted as *Input* must **not** be changed by this procedure.

2:	A – real Input
	On entry: the lower limit of integration, a.
3:	B – real Input
	On entry: the upper limit of integration, b. It is not necessary that $a < b$.
4:	OMEGA – real Input
	On entry: the parameter ω in the weight function of the transform.
5:	KEY – INTEGER Input
	On entry: indicates which integral is to be computed:
	if KEY = 1, $w(x) = \cos(\omega x)$;
	if KEY = 2, $w(x) = \sin(\omega x)$.
	Constraint: $KEY = 1$ or 2.
6:	EPSABS – real Input
	<i>On entry</i> : the absolute accuracy required. If EPSABS is negative, the absolute value is used. See Section 7.
7:	EPSREL – real Input
	<i>On entry</i> : the relative accuracy required. If EPSREL is negative, the absolute value is used. See Section 7.
8:	RESULT – real Output
	On exit: the approximation to the integral I.
9:	ABSERR – <i>real</i> Output
	On exit: an estimate of the modulus of the absolute error, which should be an upper bound for $ I - \text{RESULT} $.
10:	W(LW) – <i>real</i> array Output
	On exit: details of the computation, as described in Section 8.
11:	LW – INTEGER Input
	<i>On entry</i> : the dimension of the array W as declared in the (sub)program from which D01ANF is called. The value of LW (together with that of LIW below) imposes a bound on the number of sub-

intervals into which the interval of integration may be divided by the routine. The number of sub-

intervals cannot exceed LW/4. The more difficult the integrand, the larger LW should be.

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Suggested value: a value in the range 800 to 2000 is adequate for most problems.

Constraint: $LW \ge 4$.

12: IW(LIW) - INTEGER array

On exit: IW(1) contains the actual number of sub-intervals used. The rest of the array is used as workspace.

13: LIW – INTEGER

On entry: the dimension of the array IW as declared in the (sub)program from which D01ANF is called. The number of sub-intervals into which the interval of integration may be divided cannot exceed LIW/2.

Suggested value: LIW = LW/2.

Constraint: $LIW \ge 2$.

14: IFAIL – INTEGER

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

The maximum number of subdivisions allowed with the given workspace has been reached without the accuracy requested being achieved. Look at the integrand in order to determine the integration difficulties. If the position of a local difficulty within the interval can be determined (e.g., a singularity of the integrand or its derivative, a peak, a discontinuity, etc.) you will probably gain from splitting up the interval at this point and calling the integrator on the subranges. If necessary, another integrator, which is designed for handling the type of difficulty involved, must be used. Alternatively consider relaxing the accuracy requirements specified by EPSABS and EPSREL, or increasing amount of workspace.

IFAIL = 2

Round-off error prevents the requested tolerance from being achieved. The error may be underestimated. Consider requesting less accuracy.

IFAIL = 3

Extremely bad local behaviour of g(x) causes a very strong subdivision around one (or more) points of the interval. The same advice applies as in the case of IFAIL = 1.

IFAIL = 4

The requested tolerance cannot be achieved because the extrapolation does not increase the accuracy satisfactorily; the returned result is the best which can be obtained. The same advice applies as in the case of IFAIL = 1.

Input/Output

Output

IFAIL = 5

The integral is probably divergent, or slowly convergent. It must be noted that divergence can occur with any non-zero value of IFAIL.

IFAIL = 6

On entry, KEY < 1, or KEY > 2.

IFAIL = 7

On entry, LW < 4, or LIW < 2.

7 Accuracy

The routine cannot guarantee, but in practice usually achieves, the following accuracy:

 $|I - \text{RESULT}| \le tol,$

where

 $tol = \max\{|\text{EPSABS}|, |\text{EPSREL}| \times |I|\},\$

and EPSABS and EPSREL are user-specified absolute and relative tolerances. Moreover, it returns the quantity ABSERR which in normal circumstances, satisfies

$$|I - \text{RESULT}| \leq \text{ABSERR} \leq tol.$$

8 Further Comments

The time taken by the routine depends on the integrand and on the accuracy required.

If IFAIL $\neq 0$ on exit, then the user may wish to examine the contents of the array W, which contains the end-points of the sub-intervals used by D01ANF along with the integral contributions and error estimates over these sub-intervals.

Specifically, for i = 1, 2, ..., n, let r_i denote the approximation to the value of the integral over the subinterval $[a_i, b_i]$ in the partition of [a, b] and e_i be the corresponding absolute error estimate. Then, $\int_{a_i}^{b_i} g(x)w(x) dx \simeq r_i$ and RESULT $= \sum_{i=1}^n r_i$ unless D01ANF terminates while testing for divergence of the integral (see Section 3.4.3 of Piessens *et al.* (1983)). In this case, RESULT (and ABSERR) are taken to be the values returned from the extrapolation process. The value of n is returned in IW(1), and the values a_i , b_i , e_i and r_i are stored consecutively in the array W, that is:

$$\begin{array}{rcl} a_i &=& {\rm W}(i), \\ b_i &=& {\rm W}(n+i), \\ e_i &=& {\rm W}(2n+i) \mbox{ and } \\ r_i &=& {\rm W}(3n+i). \end{array}$$

9 Example

To compute

$$\int_0^1 \ln x \sin(10\pi x) \, dx.$$

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.

```
DO1ANF Example Program Text
      Mark 14 Revised. NAG Copyright 1989.
*
*
      .. Parameters ..
                        LW, LIW
      INTEGER
      PARAMETER
                        (LW=800,LIW=LW/2)
      INTEGER
                        NOUT
      PARAMETER
                        (NOUT=6)
      .. Scalars in Common ..
      INTEGER
                       KOUNT
      .. Local Scalars ..
                       A, ABSERR, B, EPSABS, EPSREL, OMEGA, PI, RESULT
      real
      INTEGER
                        IFAIL, KEY
      .. Local Arrays ..
*
      real
                        W(LW)
      INTEGER
                        IW(LIW)
      .. External Functions ..
                       FST, XO1AAF
FST, XO1AAF
      real
      EXTERNAL
      .. External Subroutines ..
      EXTERNAL
                       D01ANF
      .. Common blocks ..
                        /TELNUM/KOUNT
      COMMON
      .. Executable Statements ..
      WRITE (NOUT, *) 'DO1ANF Example Program Results'
      EPSREL = 1.0e - 04
      EPSABS = 0.0e+00
      A = 0.0e0
      B = 1.0e0
      OMEGA = 10.0e0 \times X01AAF(PI)
      KEY = 2
      KOUNT = 0
      IFAIL = -1
*
      CALL DO1ANF(FST,A,B,OMEGA,KEY,EPSABS,EPSREL,RESULT,ABSERR,W,LW,IW,
                   LIW, IFAIL)
*
      WRITE (NOUT, *)
      WRITE (NOUT, 99999) 'A
                                  - lower limit of integration = ', A
- upper limit of integration = ', B
      WRITE (NOUT, 99999) 'B
      WRITE (NOUT, 99998) 'EPSABS - absolute accuracy requested = ',
       EPSABS
      WRITE (NOUT, 99998) 'EPSREL - relative accuracy requested = ',
     + EPSREL
      WRITE (NOUT, *)
      IF (IFAIL.NE.O) WRITE (NOUT,99996) 'IFAIL = ', IFAIL
      IF (IFAIL.LE.5) THEN
         WRITE (NOUT, 99997) 'RESULT - approximation to the integral = ',
           RESULT
     +
         WRITE (NOUT, 99998) 'ABSERR - estimate of the absolute error = '
     +
           , ABSERR
         WRITE (NOUT,99996) 'KOUNT - number of function evaluations = '
     +
            , KOUNT
         WRITE (NOUT,99996) 'IW(1) - number of subintervals used = ',
           IW(1)
      END IF
      STOP
*
99999 FORMAT (1X,A,F10.4)
99998 FORMAT (1X,A,e9.2)
99997 FORMAT (1X,A,F9.5)
99996 FORMAT (1X,A,I4)
      END
      real FUNCTION FST(X)
*
      .. Scalar Arguments ..
```

```
real
                         Х
      .. Scalars in Common ..
*
      INTEGER KOUNT
      .. Intrinsic Functions ..
*
      INTRINSIC LOG
      INTRINSIC
.. Common blocks ..
/TELNUM/KOUNT
*
      COMMON
      .. Executable Statements ..
KOUNT = KOUNT + 1
*
      FST = 0.0e0
      IF (X.GT.0.0e^{0}) FST = LOG(X)
      RETURN
      END
```

9.2 Program Data

None.

9.3 Program Results

DO1ANF Example Program Results

```
A - lower limit of integration = 0.0000
B - upper limit of integration = 1.0000
EPSABS - absolute accuracy requested = 0.00E+00
EPSREL - relative accuracy requested = 0.10E-03
RESULT - approximation to the integral = -0.12814
ABSERR - estimate of the absolute error = 0.36E-05
KOUNT - number of function evaluations = 275
IW(1) - number of subintervals used = 8
```