

NAG Library Routine Document

D01RAF

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.

Note: *this routine uses optional parameters to define choices in the problem specification and in the details of the algorithm. If you wish to use default settings for all of the optional parameters, you need only read Sections 1 to 10 of this document. If, however, you wish to reset some or all of the settings please refer to Section 11 for a detailed description of the specification of the optional parameters.*

1 Purpose

D01RAF is a general purpose adaptive integrator which calculates an approximation to a vector of definite integrals \mathbf{F} over a finite range $[a, b]$, given the vector of integrands $\mathbf{f}(x)$.

$$\mathbf{F} = \int_a^b \mathbf{f}(x) dx$$

If the same subdivisions of the range are equally good for functions $f_1(x)$ and $f_2(x)$, because $f_1(x)$ and $f_2(x)$ have common areas of the range where they vary slowly and where they vary quickly, then we say that $f_1(x)$ and $f_2(x)$ are 'similar'. D01RAF is particularly effective for the integration of a vector of similar functions.

2 Specification

```

SUBROUTINE D01RAF (IREVCM, NI, A, B, SID, NEEDI, X, LENX, NX, FM, LDFM,      &
                  DINEST, ERREST, IOPTS, OPTS, ICOM, LICOM, COM, LCOM,      &
                  IFAIL)
INTEGER          IREVCM, NI, SID, NEEDI(NI), LENX, NX, LDFM,              &
                IOPTS(100), ICOM(LICOM), LICOM, LCOM, IFAIL
REAL (KIND=nag_wp) A, B, X(LENX), FM(LDFM,*), DINEST(NI), ERREST(NI),  &
                OPTS(100), COM(LCOM)

```

3 Description

D01RAF is an extension to various QUADPACK routines, including QAG, QAGS and QAGP. The extensions made allow multiple integrands to be evaluated simultaneously, using a vectorized interface and reverse communication.

The quadrature scheme employed by D01RAF can be chosen by you. Six Gauss–Kronrod schemes are available. The algorithm incorporates a global acceptance criterion (as defined by Malcolm and Simpson (1976)), optionally together with the ϵ -algorithm (see Wynn (1956)) to perform extrapolation. The local error estimation is described in Piessens *et al.* (1983).

D01RAF is the integration routine in the suite of routines D01RAF and D01RCF. It also uses optional parameters, which can be set and queried using the routines D01ZKF and D01ZLF respectively. The options available are described in Section 11.

First, the option arrays IOPTS and OPTS must be initialized using D01ZKF. Thereafter any required options must be set before calling D01RAF, or the routine D01RCF.

A typical usage of this suite of routines is (in pseudo-code for clarity),

Setup phase

```

liopts = 100
lopts = 100
allocate(iopts(liopts),opts(lopts))
call D01ZKF('initialize = d01raf',iopts,liopts,opts,lopts,ifail)
call D01ZKF('option = value',iopts,liopts,opts,lopts,ifail)

```

```

...
call D01RCF(ni,lenxrq,ldfmrq,sdfmrq,licmin,licmax,lcmn,lcmx, &
           iopts,opts,ifail)
lenx = lenxrq
ldfm = ldfmrq
sdfm = sdfmrq
licom = licmax
lcom = lcmx
allocate(icom(licom),com(lcom),x(lenx),fm(ldfm,sdfm),needi(ni), &
         dinest(ni),errest(ni))

```

Solve phase

```

irevcm = 1
while irevcm≠0
  call D01RAF(irevcm,ni,a,b,sid,needi,x,lenx,nx,fm,ldfm, &
             dinest,errest,iopts,opts,icom,licom,com, &
             lcom,ifail)
  select case(irevcm)
  case(11)
    Initial solve phase
    evaluate fm(1:ni,1:nx)
  case(12)
    Adaptive solve phase
    evaluate fm(needi(1:ni)=1,1:nx)
  case(0)
    investigate ifail
  end select
end while

```

Diagnostic phase

```

call D01ZLF('option',ivalue,rvalue,cvalue,optype,iopts,opts,ifail)
...

```

During the initial solve phase, the first estimation of the definite integral and error estimate is constructed over the interval $[a, b]$. This will have been divided into s_{pri} level 1 segments, where s_{pri} is the number of **Primary Divisions**, and will use any provided break-points if **Primary Division Mode** = MANUAL.

Once a complete integral estimate over $[a, b]$ is available, i.e., after all the estimates for the level 1 segments have been evaluated, the routine enters the adaptive phase. The estimated errors are tested against the requested tolerances ϵ_a and ϵ_r , corresponding to the **Absolute Tolerance** and **Relative Tolerance** respectively. Should this test fail, and additional subdivision be allowed, a segment is selected for subdivision, and is subsequently replaced by two new segments at the next level of refinement. How this segment is chosen may be altered by setting **Prioritize Error** to either favour the segment with the maximum error, or the segment with the lowest level supporting an unacceptable (although potentially non-maximal) error. Up to \max_{sdiv} subdivisions are allowed if sufficient memory is provided, where \max_{sdiv} is the value of **Maximum Subdivisions**.

Once a sufficient number of error estimates have been constructed for a particular integral, the routine may optionally use **Extrapolation** to attempt to accelerate convergence. This may significantly lower the amount of work required for a given integration. To minimize the risk of premature convergence from extrapolation, a safeguard ϵ_{safe} can be set using **Extrapolation Safeguard**, and the extrapolated solution will only be considered if $\epsilon_{safe}\epsilon_q \leq \epsilon_{ex}$, where ϵ_q and ϵ_{ex} are the estimated error directly from the quadrature and from the extrapolation respectively. If extrapolation is successful for the computation of integral j , the extrapolated solution will be returned in $DINEST(j)$ on completion of D01RAF. Otherwise the direct solution will be returned in $DINEST(j)$. This is indicated by the value of $NEEDI(j)$ on completion.

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 (1973) An algorithm for automatic integration *Angew. Inf.* **15** 399–401

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 Arguments

Note: this routine uses **reverse communication**. Its use involves an initial entry, intermediate exits and re-entries, and a final exit, as indicated by the argument **IREVCM**. Between intermediate exits and re-entries, **all arguments other than IREVCM, NEEDI and FM must remain unchanged**.

1: IREVCM – INTEGER *Input/Output*

On initial entry: IREVCM = 1.

IREVCM = 1

Sets up data structures in ICOM and COM and starts a new integration.

Constraint: IREVCM = 1 on initial entry.

On intermediate exit: IREVCM = 11 or 12.

IREVCM requests the integrands $f_j(x_i)$ be evaluated for all required $j \in 1, \dots, n_i$ as indicated by NEEDI, and at all the points x_i , for $i = 1, 2, \dots, n_x$. Abscissae x_i are provided in X(i) and $f_j(x_i)$ must be returned in FM(j, i).

During the initial solve phase:

IREVCM = 11

Function values are required to construct the initial estimates of the definite integrals.

If NEEDI(j) = 1, $f_j(x_i)$ must be supplied in FM(j, i). This will be the case unless you have abandoned the evaluation of specific integrals on a previous call.

If NEEDI(j) = 0, you have previously abandoned the evaluation of integral j, and hence should not supply the value of $f_j(x_i)$.

DINEST and ERREST contain incomplete information during this phase. As such you should not abandon the evaluation of any integrals during this phase unless you do not require their estimate.

If IREVCM is set to a negative value during this phase, NEEDI(j), for $j = 1, 2, \dots, n_i$, will be set to this negative value and IFAIL = -1 will be returned.

During the adaptive solve phase:

IREVCM = 12

Function values are required to improve the estimates of the definite integrals.

If NEEDI(j) = 0, any evaluation of $f_j(x_i)$ will be discarded, so there is no need to provide them.

If NEEDI(j) = 1, $f_j(x_i)$ must be provided in FM(j, i).

If NEEDI(j) = 2, 3 or 4, the current error estimate of integral j does not require integrand j to be evaluated and provided in FM(j, i). Should you choose to, integrand j can be evaluated in which case NEEDI(j) must be set to 1.

DINEST and ERREST contain complete information during this phase.

If IREVCM is set to a negative value during this phase IFAIL = 1, 2 or 3 will be returned and the elements of NEEDI will reflect the current state of the adaptive process.

On intermediate re-entry: IREVCM should normally be left unchanged. However, if IREVCM is set to a negative value, D01RAF will terminate, (see IREVCM = 11 and IREVCM = 12 above).

On final exit: IREVCM = 0.

IREVCM = 0

Indicates the algorithm has completed.

2: NI – INTEGER *Input*

On entry: n_i , the number of integrands.

3: A – REAL (KIND=nag_wp) *Input*

On entry: a , the lower bound of the domain.

4: B – REAL (KIND=nag_wp) *Input*

On entry: b , the upper bound of the domain.

If $|b - a| < 10\epsilon$, where ϵ is the *machine precision* (see X02AJF), then D01RAF will return $\text{DINEST}(j) = \text{ERREST}(j) = 0.0$, for $j = 1, 2, \dots, n_i$.

5: SID – INTEGER *Output*

For advanced users.

On intermediate exit: SID identifies a specific set of abscissae, \mathbf{x} , returned during the integration process. When a new set of abscissae are generated the value of SID is incremented by 1. Advanced users may store calculations required for an identified set \mathbf{x} , and reuse them should D01RAF return the same value of SID, i.e., the same set of abscissae was used.

6: NEEDI(NI) – INTEGER array *Input/Output*

On initial entry: need not be set.

On intermediate exit: NEEDI(j) indicates what action must be taken for integral $j = 1, 2, \dots, n_i$ (see IREVCM).

NEEDI(j) = 0

Do not provide $f_j(x_i)$. Any provided values will be ignored.

NEEDI(j) = 1

The values $f_j(x_i)$ must be provided in FM(j, i), for $i = 1, 2, \dots, n_x$.

NEEDI(j) = 2

The values $f_j(x_i)$ are not required, however the error estimate for integral j is still above the requested tolerance. If you wish to provide values for the evaluation of integral j , set NEEDI(j) = 1, and supply $f_j(x_i)$ in FM(j, i), for $i = 1, 2, \dots, n_x$.

NEEDI(j) = 3

The error estimate for integral j cannot be improved to below the requested tolerance directly, either because no more new splits may be performed due to exhaustion, or due to the detection of extremely bad integrand behaviour. However, providing the values $f_j(x_i)$ may still lead to some improvement, and may lead to an acceptable error estimate indirectly using Wynn's epsilon algorithm. If you wish to provide values for the evaluation of integral j , set NEEDI(j) = 1, and supply $f_j(x_i)$ in FM(j, i), for $i = 1, 2, \dots, n_x$.

NEEDI(j) = 4

The error estimate of integral j is below the requested tolerance. If you believe this to be false, if for example the result in DINEST(j) is greatly different to what you may expect, you may force the algorithm to re-evaluate this conclusion by including the evaluations of integrand j at x_i , for $i = 1, 2, \dots, n_x$, and setting NEEDI(j) = 1. Integral and error estimation will be performed again during the next iteration.

On intermediate re-entry: NEEDI(j) may be used to indicate what action you have taken for integral j .

NEEDI(j) = 1

You have provided the values $f_j(x_i)$ in FM(j, i), for $i = 1, 2, \dots, n_x$.

NEEDI(j) < 0

You are abandoning the evaluation of integral j . The current values of DINEST(j) and ERREST(j) will be returned on final completion.

Otherwise you have not provided the value $f_j(x_i)$.

On final exit: NEEDI(j) indicates the final state of integral j .

NEEDI(j) = 0

The error estimate for F_j is below the requested tolerance.

NEEDI(j) = 1

The error estimate for F_j is below the requested tolerance after extrapolation.

NEEDI(j) = 2

The error estimate for F_j is above the requested tolerance.

NEEDI(j) = 3

The error estimate for F_j is above the requested tolerance, and extremely bad behaviour of integral j has been detected.

NEEDI(j) < 0

You prohibited further evaluation of integral j .

7: X(LENX) – REAL (KIND=nag_wp) array *Input/Output*

On initial entry: if **Primary Division Mode** = AUTOMATIC, X need not be set. This is the default behaviour.

If **Primary Division Mode** = MANUAL, X is used to supply a set of initial ‘break-points’ inside the domain of integration. Specifically, X(i) must contain a break-point x_i^0 , for $i = 1, 2, \dots, (s_{pri} - 1)$, where s_{pri} is the number of **Primary Divisions**.

Constraint: if break-points are supplied, $x_i^0 \in (a, b)$, $|x_i^0 - a| > 10.0\epsilon$, $|x_i^0 - b| > 10.0\epsilon$, for $i = 1, 2, \dots, (s_{pri} - 1)$.

On intermediate exit: X(i) is the abscissa x_i , for $i = 1, 2, \dots, n_x$, at which the appropriate integrals must be evaluated.

8: LENX – INTEGER *Input*

On entry: the dimension of the array X as declared in the (sub)program from which D01RAF is called. Currently $LENX = \max(122, s_{pri} - 1)$ will be sufficient for all cases.

Constraint: $LENX \geq lenxrq$, where $lenxrq$ is dependent upon the options currently set (see Section 11). $lenxrq$ is returned as LENXRQ from D01RCF.

9: NX – INTEGER *Input/Output*

On initial entry: need not be set.

On intermediate exit: n_x , the number of abscissae at which integrands are required.

On intermediate re-entry: must not be changed.

- 10: FM(LDFM,*) – REAL (KIND=nag_wp) array Input
Note: the second dimension of the array FM must be at least $sdfmrq$, where $sdfmrq$ is dependent upon n_i and the options currently set. $sdfmrq$ is returned as SDFMRQ from D01RCF. If default options are chosen, $sdfmrq = lenxrq$.
On initial entry: need not be set.
On intermediate re-entry: if indicated by NEEDI(j) you must supply the values $f_j(x_i)$ in FM(j, i), for $i = 1, 2, \dots, n_x$ and $j = 1, 2, \dots, n_i$.
- 11: LDFM – INTEGER Input
On entry: the first dimension of the array FM as declared in the (sub)program from which D01RAF is called.
Constraint: $LDFM \geq ldfmrq$, where $ldfmrq$ is dependent upon n_i and the options currently set. $ldfmrq$ is returned as LDFMRQ from D01RCF. If default options are chosen, $ldfmrq = n_i$, implying $LDFM \geq NI$.
- 12: DINEST(NI) – REAL (KIND=nag_wp) array Input/Output
DINEST(j) contains the current estimate of the definite integral F_j .
On initial entry: need not be set.
On intermediate re-entry: must not be altered.
On exit: contains the current estimates of the NI integrals. If IREVCM = 0, this will be the final solution.
- 13: ERREST(NI) – REAL (KIND=nag_wp) array Input/Output
ERREST(j) contains the current error estimate of the definite integral F_j .
On initial entry: need not be set.
On intermediate re-entry: must not be altered.
On exit: contains the current error estimates for the NI integrals. If IREVCM = 0, ERREST contains the final error estimates of the NI integrals.
- 14: IOPTS(100) – INTEGER array Communication Array
15: OPTS(100) – REAL (KIND=nag_wp) array Communication Array
The arrays IOPTS and OPTS **must not** be altered between calls to any of the routines D01RAF, D01RCF, D01ZKF and D01ZLF.
- 16: ICOM(LICOM) – INTEGER array Communication Array
ICOM contains details of the integration procedure, including information on the integration of the n_i integrals over individual segments. This data is stored sequentially in the order that segments are created. For further information see Section 9.1.
- 17: LICOM – INTEGER Input
On entry: the dimension of the array ICOM.
Constraint: $LICOM \geq licmin$, where $licmin$ is dependent upon NI and the current options set. $licmin$ is returned as LICMIN from D01RCF. If the default options are set, then $licmin = 55 + 6 \times NI$. Larger values than $licmin$ are recommended if you anticipate that any integrals will require the domain to be further subdivided.
The maximum value that may be required, $licmax$, is returned as LICMAX from D01RCF. If default options are chosen, except for possibly increasing the value of s_{pri} , then $licmax = 50 + 5 \times NI + (s_{pri} + 100) \times (5 + NI)$.

- 18: COM(LCOM) – REAL (KIND=nag_wp) array *Communication Array*
 COM contains details of the integration procedure, including information on the integration of the n_i integrals over individual segments. This data is stored sequentially in the order that segments are created. For further information see Section 9.1.
- 19: LCOM – INTEGER *Input*
On entry: the dimension of the array COM.
Constraint: $LCOM > l_{min}$, where l_{min} is dependent upon NI, s_{pri} and the current options set. l_{min} is returned as LCMIN from D01RCF. If default options are set, then $l_{min} = 96 + 12 \times NI$. Larger values are recommended if you anticipate that any integrals will require the domain to be further subdivided.
 Given the current options and arguments, the maximum value, l_{max} , of LCOM that may be required, is returned as LCMAX from D01RCF. If default options are chosen, $l_{max} = 94 + 9 \times NI + \lceil NI/2 \rceil + (s_{pri} + 100) \times (2 + \lceil NI/2 \rceil + 2 \times NI)$.
- 20: IFAIL – INTEGER *Input/Output*
On initial entry: IFAIL must be set to 0, -1 or 1. If you are unfamiliar with this argument you should refer to Section 3.4 in How to Use the NAG Library and its Documentation for details.
 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 arguments 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.**
On final exit: $IFAIL = 0$ unless the routine detects an error or a warning has been flagged (see Section 6).

6 Error Indicators and Warnings

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

Note: D01RAF may return useful information for one or more of the following detected errors or warnings.

Errors or warnings detected by the routine:

IFAIL = 1

At least one error estimate exceeded the requested tolerances.

IFAIL = 2

Extremely bad behaviour was detected for at least one integral.

IFAIL = 3

Extremely bad behaviour was detected for at least one integral. At least one other integral error estimate was above the requested tolerance.

IFAIL = 11

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

IFAIL = 21

On entry, $NI = \langle value \rangle$.
 Constraint: $NI \geq 1$.

IFAIL = 71

On entry, **Primary Division Mode** = MANUAL and at least one supplied break-point in X is outside of the domain of integration.

IFAIL = 81

LENX is insufficient for the chosen options.

On entry, LENX = $\langle value \rangle$.

Constraint: LENX $\geq \langle value \rangle$.

IFAIL = 111

LDFM $< ldfmrq$. If default options are chosen, this implies LDFM $< NI$.

On entry, LDFM = $\langle value \rangle$.

Constraint: LDFM $\geq \langle value \rangle$.

IFAIL = 171

LICOM is insufficient for additional subdivision.

On entry, LICOM = $\langle value \rangle$.

Constraint: LICOM $\geq \langle value \rangle$.

IFAIL = 191

LCOM is insufficient for additional subdivision.

On entry, LCOM = $\langle value \rangle$.

Constraint: LCOM $\geq \langle value \rangle$.

IFAIL = 1001

Either the option arrays IOPTS and OPTS have not been initialized for D01RAF, or they have become corrupted.

IFAIL = 1101

On entry, one of ICOM and COM has become corrupted.

IFAIL = -1

Evaluation of all integrals has been stopped during the initial phase.

IFAIL = -99

An unexpected error has been triggered by this routine. Please contact NAG.

See Section 3.9 in How to Use the NAG Library and its Documentation for further information.

IFAIL = -399

Your licence key may have expired or may not have been installed correctly.

See Section 3.8 in How to Use the NAG Library and its Documentation for further information.

IFAIL = -999

Dynamic memory allocation failed.

See Section 3.7 in How to Use the NAG Library and its Documentation for further information.

7 Accuracy

D01RAF cannot guarantee, but in practice usually achieves, the following accuracy for each integral F_j :

$$|F_j - \text{DINEST}(j)| \leq \text{tol}$$

where

$$\text{tol} = \max(\epsilon_a, \epsilon_r \times |F_j|)$$

ϵ_a and ϵ_r are the error tolerances **Absolute Tolerance** and **Relative Tolerance** respectively. Moreover, it returns ERREST, the entries of which in normal circumstances satisfy,

$$|F_j - \text{DINEST}(j)| \leq \text{ERREST}(j) \leq \text{tol}.$$

8 Parallelism and Performance

D01RAF is threaded by NAG for parallel execution in multithreaded implementations of the NAG Library.

D01RAF 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 routine. Please also consult the Users' Note for your implementation for any additional implementation-specific information.

9 Further Comments

The time required by D01RAF is usually dominated by the time required to evaluate the values of the integrands f_j .

D01RAF will be most efficient if any badly behaved integrands provided have irregularities over similar subsections of the domain. For example, evaluation of the integrals,

$$\int_0^1 \begin{pmatrix} \log(x) \\ x^{-\frac{1}{2}} \\ x^2 \end{pmatrix} dx$$

will be quite efficient, as the irregular behaviour of the first two integrands is at $x = 0$. On the contrary, the evaluation of the integrals,

$$\int_0^1 \begin{pmatrix} \log(x) \\ \log(1-x) \end{pmatrix} dx$$

will be less efficient, as the two integrands have singularities at opposite ends of the domain, which will result in subdivisions which are only of use to one integrand. In such cases, it will be more efficient to use two sets of calls to D01RAF.

D01RAF will flag extremely bad behaviour if a sub-interval \bar{k} with bounds $[a_{\bar{k}}, b_{\bar{k}}]$ satisfying $|b_{\bar{k}} - a_{\bar{k}}| < \max(\delta_a, \delta_r \times |b - a|)$ has a local error estimate greater than the requested tolerance for at least one integral. The values δ_a and δ_r can be set through the optional parameters **Absolute Interval Minimum** and **Relative Interval Minimum** respectively.

9.1 Details of the Computation

This section is recommended for expert users only. It describes the contents of the arrays COM and ICOM upon exit from D01RAF with IFAIL = 0, 1, 2 or 3, and provided at least one iteration completed, failure due to insufficient LICOM or LCOM.

The arrays ICOM and COM contain details of the integration, including various scalars, one-dimensional arrays, and (effectively) two-dimensional arrays. The dimensions of these arrays vary

depending on the arguments and options used and the progress of the algorithm. Here we describe some of these details, including how and where they are stored in ICOM and COM.

Scalar quantities:

The indices in ICOM including the following scalars are available via query only options, see Section 11.2. For example, I_{ldi} is the integer value returned by the option **Index LDI**.

- ldi* The leading dimension of the two-dimensional integer arrays stored in ICOM detailed below.
 $ldi = \text{ICOM}(I_{ldi})$.
- ldr* The leading dimension of the two-dimensional real arrays stored in COM detailed below.
 $ldr = \text{ICOM}(I_{ldr})$.
- nsdiv* The number of segments that have been subdivided during the adaptive process.
 $nsdiv = \text{ICOM}(I_{nsdiv})$.
- nseg* The total number of segments formed.
 $nseg = 2nsdiv + s_{pri}$.
 $nseg = \text{ICOM}(I_{nseg})$.
- dsp* The reference of the first element of the array *ds* stored in COM.
 $dsp = \text{ICOM}(I_{dsp})$.
- esp* The reference of the first element of the array *es* stored in COM.
 $esp = \text{ICOM}(I_{esp})$.
- evalsp* The reference of the first element of the array *evals* stored in ICOM.
 $evalsp = \text{ICOM}(I_{evalsp})$.
- fcp* The reference of the first element of the array *fcount* stored in ICOM.
 $fcp = \text{ICOM}(I_{fcp})$.
- sinforp* The reference of the first element of the array *sinfor* stored in COM.
 $sinforp = \text{ICOM}(I_{sinforp})$.
- sinfoip* The reference of the first element of the array *sinfoi* stored in ICOM.
 $sinfoip = \text{ICOM}(I_{sinfoip})$.

One-dimensional arrays:

$fcount(n_i)$
 $fcount(1) = \text{ICOM}(fcp)$.

$fcount(j)$ contains the number of different approximations of integral j calculated, for $j = 1, 2, \dots, n_i$.

Two-dimensional arrays:

$sinfoi(5, nseg)$
 $sinfoi(1, 1) = \text{ICOM}(sinfoip)$.
 $sinfoi$ contains information about the hierarchy of splitting.

$sinfoi(1, k)$ contains the split identifier for segment k , for $k = 1, 2, \dots, nseg$.

$sinfoi(2, k)$ contains the parent segment number of segment k (i.e., the segment was split to create segment k), for $k = 1, 2, \dots, nseg$.

$sinfoi(3, k)$ and $sinfoi(4, k)$ contain the segment numbers of the two child segments formed from segment k , if segment k has been split. If segment k has not been split, these will be negative.

$sinfoi(5, k)$ contains the level at which the segment exists, corresponding to $n_a + 1$, where n_a is the number of ancestor segments of segment k , for $k = 1, 2, \dots, nseg$. A negative level indicates that segment k will not be split further, the level is then given by the absolute value of $sinfoi(5, k)$.

$sinfor(2, nseg)$
 $sinfor(1, 1) = \text{COM}(sinforp)$.
 $sinfor$ contains the bounds of each segment.

$sinfor(1, k)$ contains the lower bound of segment k , for $k = 1, 2, \dots, nseg$.

$\text{sinfor}(2, k)$ contains the upper bound of segment k , for $k = 1, 2, \dots, n\text{seg}$.

$\text{evals}(n_i, n\text{seg})$
 $\text{evals}(1, 1) = \text{ICOM}(\text{evalsp})$.

evals contains information to indicate whether an estimate of the integral j has been obtained over segment k , and if so whether this evaluation still contributes to the direct estimate of F_j , for $j = 1, 2, \dots, n_i$ and $k = 1, 2, \dots, n\text{seg}$.

$\text{evals}(j, k) = 0$ indicates that integral j has not been evaluated over segment k .

$\text{evals}(j, k) = 1$ indicates that integral j has been evaluated over segment k , and that this evaluation contributes to the direct estimate of F_j .

$\text{evals}(j, k) = 2$ indicates that integral j has been evaluated over segment k , that this evaluation contributes to the direct estimate of F_j , and that you have requested no further evaluation of this integral at this segment by setting $\text{NEEDI}(j) < 0$.

$\text{evals}(j, k) = 3$ indicates that integral j has been evaluated over segment k , and this evaluation no longer contributes to the direct estimate of F_j .

$\text{evals}(j, k) = 4$ indicates that integral j has been evaluated over segment k , that this evaluation contributes to the direct estimate of F_j , and that this segment is too small for any further splitting to be performed. Integral j also has a local error estimate over this segment above the requested tolerance. Such segments cause D01RAF to return $\text{IFAIL} = 2$ or 3 , indicating extremely bad behaviour.

$\text{evals}(j, k) = 5$ indicates that integral j has been evaluated over segment k , that this evaluation contributes to the direct estimate of F_j , and that this segment is too small for any further splitting to be performed. The local error estimate is however below the requested tolerance.

$\text{ds}(n_i, n\text{seg})$
 $\text{ds}(1, 1) = \text{COM}(\text{dsp})$.

$\text{ds}(j, k)$ contains the definite integral estimate of the j th integral over the k th segment, $\text{ds}_{j,k}$, provided it has been evaluated, for $j = 1, 2, \dots, n_i$ and $k = 1, 2, \dots, n\text{seg}$.

$\text{es}(n_i, n\text{seg})$
 $\text{es}(1, 1) = \text{COM}(\text{esp})$.

$\text{es}(j, k)$ contains the definite integral error estimate of the j th integral over the k th segment, $\text{es}_{j,k}$, provided it has been evaluated, for $j = 1, 2, \dots, n_i$ and $k = 1, 2, \dots, n\text{seg}$.

For each integral j , the direct approximation D_j of F_j , and its error estimate E_j , may be constructed as,

$$F_j \approx D_j = \sum_{K_j} \text{ds}_{j,k},$$

$$|F_j - D_j| \approx E_j = \sum_{K_j} \text{es}_{j,k},$$

where K_j is the set of all contributing segments, $K_j = \{k \mid \text{evals}(j, k) = 1, 2, 4 \text{ or } 5, 1 \leq k \leq n\text{seg}\}$. D_j will have been returned in $\text{DINEST}(j)$, unless extrapolation was successful, as indicated by $\text{NEEDI}(j)$.

Similarly, E_j will have been returned in $\text{ERREST}(j)$ unless extrapolation was successful, in which case the error estimate from the extrapolation will have been returned. If for a given integral j one or more contributing segments have unacceptable error estimates, it may be possible to improve the direct approximation by replacing the contributions from these segments with more accurate estimates should these be calculable by some means. Indeed for any segment $\bar{k} \in k$, with lower bound $a_{\bar{k}} = \text{sinfor}(1, \bar{k})$ and upper bound $b_{\bar{k}} = \text{sinfor}(2, \bar{k})$, one may alter the direct approximation D_j by the following,

$$D_j = \sum_{K_j} \text{ds}_{j,k} - \text{ds}_{j,\bar{k}} + \text{ds}_{j,\bar{k}}^{\text{new}}$$

$$\text{ds}_{j,\bar{k}}^{\text{new}} \approx \int_{a_{\bar{k}}}^{b_{\bar{k}}} f_j(x) dx$$

The error estimate E_j may be altered similarly.

10 Example

This example integrates

$$\mathbf{F} = \int_0^{\pi} \begin{pmatrix} x \sin(2x) \cos(15x) \\ x^2 \sin(2x) \cos(50x) \end{pmatrix} dx.$$

10.1 Program Text

```

!   D01RAF Example Program Text
!   Mark 26 Release. NAG Copyright 2016.
!   Module d01rafe_mod

!
!   .. Use Statements ..
!   Use nag_library, Only: nag_wp
!   .. Implicit None Statement ..
!   Implicit None
!   .. Accessibility Statements ..
!   Private
!   Public
!
!   .. Parameters ..
!   Integer, Parameter, Public      :: nout = 6
!   Logical, Parameter, Public      :: disp_integration_info = .True.
Contains
!   Subroutine display_integration_details(ni,iopts,opts,icom,licom,com,
!   lcom)
!
!   .. Use Statements ..
!   Use nag_library, Only: d01zlf
!   .. Implicit None Statement ..
!   Implicit None
!   .. Scalar Arguments ..
!   Integer, Intent (In)            :: lcom, licom, ni
!   .. Array Arguments ..
!   Real (Kind=nag_wp), Intent (In) :: com(lcom), opts(100)
!   Integer, Intent (In)            :: icom(licom), iopts(100)
!   .. Local Scalars ..
!   Real (Kind=nag_wp)              :: rvalue
!   Integer                          :: dsp, esp, evalsp, fcp, ifail, index, &
!   ldi, ldr, nsdiv, nseg, optype, &
!   sinfoip, sinforp
!   Character (16)                  :: cvalue
!
!   .. Executable Statements ..

!
!   Request communication array indices.
!   ifail = 0
!   Call d01zlf('Index nseg',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   nseg = icom(index)
!   Call d01zlf('Index nsdiv',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   nsdiv = icom(index)
!   Call d01zlf('Index ldi',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   ldi = icom(index)
!   Call d01zlf('Index ldr',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   ldr = icom(index)
!   Call d01zlf('Index fcp',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   fcp = icom(index)
!   Call d01zlf('Index evalsp',index,rvalue,cvalue,optype,iopts,opts,
!   ifail)
!   evalsp = icom(index)
!   Call d01zlf('Index sinfoip',index,rvalue,cvalue,optype,iopts,opts,
!   ifail)
!   sinfoip = icom(index)
!   Call d01zlf('Index dsp',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   dsp = icom(index)
!   Call d01zlf('Index esp',index,rvalue,cvalue,optype,iopts,opts,ifail)
!   esp = icom(index)
!   Call d01zlf('Index sinforp',index,rvalue,cvalue,optype,iopts,opts,
!   ifail)

```

```

    sinforp = icom(index)

    Write (nout,*)
    Write (nout,99999)
    Write (nout,99998) ni, nseg, nsdiv
    Call display_subdivision_strategy(ni,nseg,icom(fcp),icom(sinfoip),      &
        icom(evalsp),ldi,com(sinforp),com(dsp),com(esp),ldr)

    Return
99999  Format (' Information on integration: ')
99998  Format (' NI = ',I3,', nseg = ',I3,', nsdiv = ',I3,','.')
End Subroutine display_integration_details
Subroutine display_subdivision_strategy(ni,nseg,fcount,sinfoi,evals,ldi, &
    sinfor,ds,es,ldr)

!      .. Scalar Arguments ..
Integer, Intent (In)          :: ldi, ldr, ni, nseg
!      .. Array Arguments ..
Real (Kind=nag_wp), Intent (In) :: ds(ldr,*), es(ldr,*), sinfor(ldr,*)
Integer, Intent (In)          :: evals(ldi,*), fcount(ni),      &
    sinfoi(ldi,*)

!      .. Local Scalars ..
Real (Kind=nag_wp)           :: lbnd, ubnd
Integer                       :: child1, child2, j, k, level, parent, &
    sid

!      .. Executable Statements ..

!      Display information on individual segments.
Do j = 1, ni
    Write (nout,99991) j, fcount(j)
End Do
Write (nout,*)
Write (nout,99992)
Do k = 1, nseg
    Write (nout,*)
    sid = sinfoi(1,k)
    parent = sinfoi(2,k)
    child1 = sinfoi(3,k)
    child2 = sinfoi(4,k)
    level = sinfoi(5,k)
    lbnd = sinfor(1,k)
    ubnd = sinfor(2,k)
    Write (nout,99999) k
    Write (nout,99998) sid, parent, level
    If (child1>0) Then
        Write (nout,99997) child1, child2
    End If
    Write (nout,99996) lbnd, ubnd
    Do j = 1, ni
        If (evals(j,k)/=0) Then
            Write (nout,99995) j, ds(j,k)
            Write (nout,99994) j, es(j,k)
            If (evals(j,k)==3) Then
                Write (nout,99993) j
            End If
        End If
    End Do
End Do
Return
99999  Format (' Segment ',I3,','.')
99998  Format (' Sid = ',I3,', Parent = ',I3,', Level = ',I3,','.')
99997  Format (' Children = (',I3,',',I3,','.')
99996  Format (' Bounds (',Es11.4,',',Es11.4,','.')
99995  Format (' Integral ',I2,' approximation :',lX,Es11.4,','.')
99994  Format (' Integral ',I2,' error estimate:',lX,Es11.4,','.')
99993  Format (' Integral ',I2,
    ' evaluation has been superseded by descendants.')      &
99992  Format (' Information on subdivision and evaluations over segments.')
99991  Format (' Integral ',I2,' total approximations: ',I3,','.')
End Subroutine display_subdivision_strategy
Subroutine display_option(optstr,optype,ivalue,rvalue,cvalue)

```

```

!      Subroutine to query optype and print the appropriate option values

!      .. Scalar Arguments ..
Real (Kind=nag_wp), Intent (In) :: rvalue
Integer, Intent (In)          :: ivalue, optype
Character (*), Intent (In)    :: cvalue, optstr
!      .. Executable Statements ..

      Select Case (optype)
      Case (1)
        Write (nout,99999) optstr, ivalue
      Case (2)
        Write (nout,99998) optstr, rvalue
      Case (3)
        Write (nout,99997) optstr, cvalue
      Case (4)
        Write (nout,99996) optstr, ivalue, cvalue
      Case (5)
        Write (nout,99995) optstr, rvalue, cvalue
      Case Default
      End Select

      Flush (nout)

      Return
99999  Format (3X,A30,' : ',I16)
99998  Format (3X,A30,' : ',Es16.4)
99997  Format (3X,A30,' : ',12X,A16)
99996  Format (3X,A30,' : ',I16,3X,A16)
99995  Format (3X,A30,' : ',Es16.4,3X,A16)
      End Subroutine display_option
End Module d01rafe_mod

Program d01rafe

!      .. Use Statements ..
Use nag_library, Only: d01raf, d01rcf, d01z kf, d01zlf, nag_wp, x0laaf
Use d01rafe_mod, Only: display_integration_details, display_option,      &
                        disp_integration_info, nout
!      .. Implicit None Statement ..
Implicit None
!      .. Local Scalars ..
Real (Kind=nag_wp)          :: a, b, pi, rvalue
Integer                     :: ifail, irevcm, ivalue, j, lcomax,      &
                                lcomin, lcom, ldfm, ldfmrq, lenx,      &
                                lenxrq, licmax, licmin, licom, ni,      &
                                nx, optype, sdfm, sdfmrq, sid
Character (16)              :: cvalue
!      .. Local Arrays ..
Real (Kind=nag_wp), Allocatable :: com(:), dinest(:), errest(:),      &
                                fm(:, :), opts(:), x(:)
Integer, Allocatable         :: icom(:), iopts(:), needi(:)
!      .. Intrinsic Procedures ..
Intrinsic                   :: cos, sin
!      .. Executable Statements ..
Continue
Write (nout,*) 'D01RAF Example Program Results'
Write (nout,*)

pi = x0laaf(pi)

!      Setup phase.

!      Set problem parameters
ni = 2
!      Lower (a) and upper (b) bounds
a = 0.0E0_nag_wp
b = pi
Allocate (opts(100),iopts(100))

!      Initialize option arrays

```

```

ifail = 0
Call d01zkg('Initialize = d01raf',iopts,100,opts,100,ifail)
! Set any non-default options required
Call d01zkg('Quadrature Rule = gk41',iopts,100,opts,100,ifail)
Call d01zkg('Absolute Tolerance = 1.0e-7',iopts,100,opts,100,ifail)
Call d01zkg('Relative Tolerance = 1.0e-7',iopts,100,opts,100,ifail)

! Determine maximum required array lengths
ifail = -1
Call d01rcf(ni,lenxrq,ldfmrq,sdfmrq,licmin,licmax,lcmn,lcmx,iopts,      &
  opts,ifail)
ldfm = ldfmrq
sdfm = sdfmrq
lenx = lenxrq
licom = licmax
lcom = lcmx

! Allocate remaining arrays
Allocate (icom(licom),needi(ni),com(lcom),fm(ldfm,sdfm),dinst(ni),      &
  errest(ni),x(lenx))

! Solve phase.
! Use D01RAF to evaluate the definite integrals of:
! f_1 = (x*sin(2*x))*cos(15*x)
! f_2 = (x*sin(2*x))*(x*cos(50*x))

! Set initial irevcm
irevcm = 1
ifail = -1

Do While (irevcm/=0)

  Call d01raf(irevcm,ni,a,b,sid,needi,x,lenx,nx,fm,ldfm,dinst,errest,      &
    iopts,opts,icom,licom,com,lcom,ifail)

  Select Case (irevcm)
  Case (11)
! Initial returns.
! These will occur during the non-adaptive phase.
! All values must be supplied.
! DINEST and ERREST do not contain approximations
! over the complete interval at this stage.

! Calculate x*sin(2*x), storing the result in fm(2,1:nx) for re-use.
fm(2,1:nx) = x(1:nx)*sin(2.0E0_nag_wp*x(1:nx))

! Calculate f1
fm(1,1:nx) = fm(2,1:nx)*cos(15.0E0_nag_wp*x(1:nx))

! Complete f2 calculation.
fm(2,1:nx) = fm(2,1:nx)*x(1:nx)*cos(50.0E0_nag_wp*x(1:nx))

  Case (12)
! Intermediate returns.
! These will occur during the adaptive phase.
! All requested values must be supplied.
! DINEST and ERREST do not contain approximations
! over the complete interval at this stage.

! Calculate x*sin(2*x).
fm(2,1:nx) = x(1:nx)*sin(2.0E0_nag_wp*x(1:nx))

! Calculate f1 if required.
If (needi(1)==1) Then
  fm(1,1:nx) = fm(2,1:nx)*cos(15.0E0_nag_wp*x(1:nx))
End If

! Complete f2 calculation if required.
If (needi(2)==1) Then
  fm(2,1:nx) = fm(2,1:nx)*x(1:nx)*cos(50.0E0_nag_wp*x(1:nx))
End If

```

```

      Case (0)
!      Final return. Test IFAIL.
      Select Case (ifail)
      Case (0:3)
!      Useful information has been returned.
      Case Default
!      An unrecoverable error has been detected.
      Go To 100
      End Select
End Select

End Do

!      Query some currently set options and statistics.
      ifail = 0
      Call d01zlf('Quadrature rule',ivalue,rvalue,cvalue,optype,iopts,opts, &
        ifail)
      Call display_option('Quadrature rule',optype,ivalue,rvalue,cvalue)

      Call d01zlf('Maximum Subdivisions',ivalue,rvalue,cvalue,optype,iopts, &
        opts,ifail)
      Call display_option('Maximum Subdivisions',optype,ivalue,rvalue,cvalue)

      Call d01zlf('Extrapolation',ivalue,rvalue,cvalue,optype,iopts,opts, &
        ifail)
      Call display_option('Extrapolation',optype,ivalue,rvalue,cvalue)

      Call d01zlf('Extrapolation Safeguard',ivalue,rvalue,cvalue,optype,iopts, &
        opts,ifail)
      Call display_option('Extrapolation safeguard',optype,ivalue,rvalue, &
        cvalue)

!      Print solution
      Write (nout,*)
      Write (nout,99999)
      Do j = 1, ni
        Write (nout,99998) j, needi(j), dinest(j), errest(j)
      End Do

!      Investigate integration strategy
      If (disp_integration_info) Then
        Call display_integration_details(ni,iopts,opts,icom,licom,com,lcom)
      End If

100      Continue

99999 Format (' Integral | NEEDI |   DINEST   |   ERREST   ')
99998 Format (2(1X,I9),2(1X,Es12.4))
      End Program d01rafe

```

10.2 Program Data

None.

10.3 Program Results

D01RAF Example Program Results

```

      Quadrature rule :           GK41
      Maximum Subdivisions :       50
      Extrapolation :             ON
      Extrapolation safeguard :    1.0000E-12

```

```

Integral |  NEEDI |   DINEST   |   ERREST
      1   |    0   | -2.8431E-02 | 1.1234E-14
      2   |    0   | 7.9083E-03  | 2.6600E-09

```

```

Information on integration:
NI = 2, nseg = 7, nsdiv = 3.
Integral 1 total approximations: 2.

```


Integral 2 total approximations: 4.

Information on subdivision and evaluations over segments.

Segment 1.
 Sid = 1, Parent = 0, Level = 1.
 Children = (2, 3).
 Bounds (0.0000E+00, 3.1416E+00).
 Integral 1 approximation : -2.8431E-02.
 Integral 1 error estimate: 8.0372E-04.
 Integral 1 evaluation has been superseded by descendants.
 Integral 2 approximation : -3.6050E-01.
 Integral 2 error estimate: 4.2596E+00.
 Integral 2 evaluation has been superseded by descendants.

Segment 2.
 Sid = 2, Parent = 1, Level = 2.
 Children = (6, 7).
 Bounds (0.0000E+00, 1.5708E+00).
 Integral 1 approximation : -1.2285E-03.
 Integral 1 error estimate: 2.8161E-15.
 Integral 2 approximation : 1.9771E-03.
 Integral 2 error estimate: 4.0437E-01.
 Integral 2 evaluation has been superseded by descendants.

Segment 3.
 Sid = 2, Parent = 1, Level = 2.
 Children = (4, 5).
 Bounds (1.5708E+00, 3.1416E+00).
 Integral 1 approximation : -2.7202E-02.
 Integral 1 error estimate: 8.4182E-15.
 Integral 2 approximation : 5.9313E-03.
 Integral 2 error estimate: 3.0259E+00.
 Integral 2 evaluation has been superseded by descendants.

Segment 4.
 Sid = 3, Parent = 3, Level = 3.
 Bounds (1.5708E+00, 2.3562E+00).
 Integral 2 approximation : 1.0922E-01.
 Integral 2 error estimate: 7.9151E-10.

Segment 5.
 Sid = 3, Parent = 3, Level = 3.
 Bounds (2.3562E+00, 3.1416E+00).
 Integral 2 approximation : -1.0329E-01.
 Integral 2 error estimate: 1.6413E-09.

Segment 6.
 Sid = 4, Parent = 2, Level = 3.
 Bounds (0.0000E+00, 7.8540E-01).
 Integral 2 approximation : 1.2343E-02.
 Integral 2 error estimate: 5.2456E-11.

Segment 7.
 Sid = 4, Parent = 2, Level = 3.
 Bounds (7.8540E-01, 1.5708E+00).
 Integral 2 approximation : -1.0365E-02.
 Integral 2 error estimate: 1.7467E-10.

11 Optional Parameters

This section can be skipped if you wish to use the default values for all optional parameters, otherwise, the following is a list of the optional parameters available. A full description of each optional parameter is provided in Section 11.1.

Absolute Interval Minimum

Absolute Tolerance

Extrapolation

Extrapolation Safeguard**Maximum Subdivisions****Primary Division Mode****Primary Divisions****Prioritize Error****Quadrature Rule****Relative Interval Minimum****Relative Tolerance**

The following optional parameters, see Section 11.2, may be utilized by expert users in conjunction with the information provided in Section 9.1.

Index LDI**Index LDR****Index NSDIV****Index NSEG****Index FCP****Index EVALSP****Index DSP****Index ESP****Index SINFOIP****Index SINFORP****11.1 Description of the Optional Parameters**

For each option, we give a summary line, a description of the optional parameter and details of constraints.

The summary line contains:

the keywords, where the minimum abbreviation of each keyword is underlined;

a parameter value, where the letters a , i and r denote options that take character, integer and real values respectively;

the default value.

The following symbol represents various machine constants:

ϵ represents the *machine precision* (see X02AJF).

All options accept the value 'DEFAULT' in order to return single options to their default states.

Keywords and character values are case insensitive, however they must be separated by at least one space.

Unsettable options will return the appropriate value when calling D01ZLF. They will have no effect if passed to D01ZKF.

For D01RAF the maximum length of the argument CVALUE used by D01ZLF is 15.

Absolute Interval Minimum r Default = 128.0 ϵ

$r = \delta_a$, the absolute lower limit for a segment to be considered for subdivision. See also **Relative Interval Minimum** and Section 9.

Constraint: $r \geq 128\epsilon$.

Absolute Tolerance r Default = 1024 ϵ

$r = \epsilon_a$, the absolute tolerance required. See also **Relative Tolerance** and Section 3.

Constraint: $r \geq 0.0$.

Extrapolation *a* Default = ON

Activate or deactivate the use of the ϵ algorithm (Wynn (1956)). **Extrapolation** often reduces the number of iterations required to achieve the desired solution, but it can occasionally lead to premature convergence towards an incorrect answer.

ON

Use extrapolation.

OFF

Disable extrapolation.

Extrapolation Safeguard *r* Default = 1.0E–12

$r = \epsilon_{safe}$. If ϵ_q is the estimated error from the quadrature evaluation alone, and ϵ_{ex} is the error estimate determined using extrapolation, then the extrapolated solution will only be accepted if $\epsilon_{safe}\epsilon_q \leq \epsilon_{ex}$.

Maximum Subdivisions *i* Default = 50

$i = \max_{div}$, the maximum number of subdivisions the algorithm may use in the adaptive phase, forming at most an additional $(2 \times \max_{div})$ segments.

Primary Divisions *i* Default = 1

$i = s_{pri}$, the number of initial segments of the domain $[a, b]$. By default the initial segment is the entire domain.

Constraint: $0 < i < 1000000$.

Primary Division Mode *a* Default = AUTOMATIC

Determines how the initial set of s_{pri} segments will be generated.

AUTOMATIC

D01RAF will automatically generate s_{pri} segments of equal size covering the interval $[a, b]$.

MANUAL

D01RAF will use the break-points x_i^0 , for $i = 1, 2, \dots, s_{pri} - 1$, supplied in X on initial entry to generate the initial segments covering $[a, b]$. These may be supplied in any order, however it will be more efficient to supply them in ascending (or descending if $a > b$) order. Repeated break-points are allowed, although this will generate fewer initial segments.

Note: an absolute bound on the size of an initial segment of 10.0ϵ is automatically applied in all cases, and will result in fewer initial subdivisions being generated if automatically generated or supplied break-points result in segments smaller than this.

Prioritize Error *a* Default = LEVEL

Indicates how new subdivisions of segments sustaining unacceptable local errors for integrals should be prioritized.

LEVEL

Segments with lower level with unsatisfactory error estimates will be chosen over segments with greater error on higher levels. This will probably lead to more integrals being improved in earlier iterations of the algorithm, and hence will probably lead to fewer repeated returns (see argument SID), and to more integrals being satisfactorily estimated if computational exhaustion occurs.

MAXERR

The segment with the worst overall error will be split, regardless of level. This will more rapidly improve the worst integral estimates, although it will probably result in the fewest integrals being improved in earlier iterations, and may hence lead to more repeated returns (see argument SID), and potentially fewer integrals satisfying the requested tolerances if computational exhaustion occurs.

Quadrature Rule a Default = GK15

The basic quadrature rule to be used during the integration. Currently 6 Gauss–Kronrod rules are available, all identifiable by the letters GK followed by the number of points required by the Kronrod rule. Higher order rules generally provide higher accuracy with fewer subdivisions. However, for integrands with sharp singularities, lower order rules may be more efficient, particularly if the integrand away from the singularity is well behaved. With higher order rules, you may need to increase the **Absolute Interval Minimum** and the **Relative Interval Minimum** to maintain numerical difference between the abscissae and the segment bounds.

GK15

The Gauss–Kronrod rule based on 7 Gauss points and 15 Kronrod points.

GK21

The Gauss–Kronrod rule based on 10 Gauss points and 21 Kronrod points. This is the rule used by D01ATF

GK31

The Gauss–Kronrod rule based on 15 Gauss points and 31 Kronrod points.

GK41

The Gauss–Kronrod rule based on 20 Gauss points and 41 Kronrod points.

GK51

The Gauss–Kronrod rule based on 25 Gauss points and 51 Kronrod points.

GK61

The Gauss–Kronrod rule based on 30 Gauss points and 61 Kronrod points. This is the highest order rule, most suitable for highly oscillatory integrals.

Relative Interval Minimum r Default = 1.0E–6

$r = \delta_r$, the relative factor in the lower limit, $\delta_r|b - a|$, for a segment to be considered for subdivision. See also **Absolute Interval Minimum** and Section 9.

Constraint: $r \geq 0.0$.

Relative Tolerance r Default = $\sqrt{\epsilon}$

$r = \epsilon_r$, the required relative tolerance. See also **Absolute Tolerance** and Section 3.

Constraint: $r \geq 0.0$.

Note: setting both $\epsilon_r = \epsilon_a = 0.0$ is possible, although it will most likely result in an excessive amount of computational effort.

11.2 Diagnostic Options

These options are provided for expert users who wish to examine and modify the precise details of the computation. They should only be used **after** D01RAF returns, as opposed to the options listed in Section 11.1 which must be used **before** the first call to D01RAF.

Index LDI i query only

I_{ldi} , the index of ICOM required for obtaining ldi . See Section 9.1.

Index LDR i query only

I_{ldr} , the index of ICOM required for obtaining ldr . See Section 9.1.

Index NSDIV i query only

I_{nsdiv} , the index of ICOM required for obtaining $nsdiv$. See Section 9.1.

Index NSEG i query only

I_{nseg} , the index of ICOM required for obtaining $nseg$. See Section 9.1.

<u>Index FCP</u>	<i>i</i>	query only
<i>I_{fcp}</i> , the index of ICOM required for obtaining <i>fcp</i> . See Section 9.1.		
<u>Index EVALSP</u>	<i>i</i>	query only
<i>I_{evalsp}</i> , the index of ICOM required for obtaining <i>evalsp</i> . See Section 9.1.		
<u>Index DSP</u>	<i>i</i>	query only
<i>I_{dsp}</i> , the index of ICOM required for obtaining <i>dsp</i> . See Section 9.1.		
<u>Index ESP</u>	<i>i</i>	query only
<i>I_{esp}</i> , the index of ICOM required for obtaining <i>esp</i> . See Section 9.1.		
<u>Index SINFOIP</u>	<i>i</i>	query only
<i>I_{sinfoip}</i> , the index of ICOM required for obtaining <i>sinfoip</i> . See Section 9.1.		
<u>Index SINFORP</u>	<i>i</i>	query only
<i>I_{sinforp}</i> , the index of ICOM required for obtaining <i>sinforp</i> . See Section 9.1.		
