

# NAG C Library Function Document

## nag\_1d\_quad\_inf\_wt\_trig\_1 (d01ssc)

### 1 Purpose

nag\_1d\_quad\_inf\_wt\_trig\_1 (d01ssc) calculates an approximation to the sine or the cosine transform of a function  $g$  over  $[a, \infty)$ :

$$I = \int_a^{\infty} g(x) \sin(\omega x) dx \quad \text{or} \quad I = \int_a^{\infty} g(x) \cos(\omega x) dx$$

(for a user-specified value of  $\omega$ ).

### 2 Specification

```
#include <nag.h>
#include <nagd01.h>

void nag_1d_quad_inf_wt_trig_1(double (*g)(double x, Nag_User *comm),
                                double a, double omega, Nag_TrigTransform wt_func,
                                Integer maxintervals, Integer maxsubints_per_interval,
                                double epsabs, double *result, double *abserr,
                                Nag_QuadSubProgress *qpsub, Nag_User *comm, NagError *fail)
```

### 3 Description

This function is based upon the QUADPACK routine QAWFE (Piessens *et al.* (1983)). It is an adaptive routine, designed to integrate a function of the form  $g(x)w(x)$  over a semi-infinite interval, where  $w(x)$  is either  $\sin(\omega x)$  or  $\cos(\omega x)$ . Over successive intervals

$$C_k = [a + (k - 1) \times c, a + k \times c], \quad k = 1, 2, \dots, \mathbf{qpsub.intervals}$$

integration is performed by the same algorithm as is used by nag\_1d\_quad\_wt\_trig\_1 (d01snc). The intervals  $C_k$  are of constant length

$$c = \{2[\omega]\} + 1\} \pi / |\omega|, \quad \omega \neq 0,$$

where  $[\omega]$  represents the largest integer less than or equal to  $|\omega|$ . Since  $c$  equals an odd number of half periods, the integral contributions over succeeding intervals will alternate in sign when the function  $g$  is positive and monotonically decreasing over  $[a, \infty)$ . The algorithm, described by Piessens *et al.* (1983), incorporates a global acceptance criterion (as defined by Malcolm and Simpson (1976)) together with the  $\epsilon$ -algorithm (Wynn (1956)) to perform extrapolation. The local error estimation is described by Piessens *et al.* (1983).

If  $\omega = 0$  and **wt\_func** = Nag\_Cosine, the routine uses the same algorithm as nag\_1d\_quad\_inf\_1 (d01smc) (with **epsrel** = 0.0).

In contrast to most other functions in Chapter d01, nag\_1d\_quad\_inf\_wt\_trig\_1 works only with a user-specified absolute error tolerance (**epsabs**). Over the interval  $C_k$  it attempts to satisfy the absolute accuracy requirement

$$EPSA_k = U_k \times \mathbf{epsabs},$$

where  $U_k = (1 - p)p^{k-1}$ , for  $k = 1, 2, \dots$  and  $p = 0.9$ .

However, when difficulties occur during the integration over the  $k$ th interval  $C_k$  such that the error flag **qpsub→interval\_flag** $[k - 1]$  is non-zero, the accuracy requirement over subsequent intervals is relaxed. See Piessens *et al.* (1983) for more details.

## 4 Parameters

1: **g** – function supplied by user *Function*

The function **g**, supplied by the user, must return the value of the function  $g$  at a given point.

The specification of **g** is:

```
double g(double x, Nag_User *comm)
```

1: **x** – double *Input*

*On entry:* the point at which the function  $g$  must be evaluated.

2: **comm** – Nag\_User \*

*On entry/on exit:* pointer to a structure of type **Nag\_User** with the following member:

**p** – Pointer *Input/Output*

*On entry/on exit:* the pointer **comm**–**p** should be cast to the required type, e.g., `struct user *s = (struct user *)comm->p;`, to obtain the original object's address with appropriate type. (See the argument **comm** below.)

2: **a** – double *Input*

*On entry:* the lower limit of integration,  $a$ .

3: **omega** – double *Input*

*On entry:* the parameter  $\omega$  in the weight function of the transform.

4: **wt\_func** – Nag\_TrigTransform *Input*

*On entry:* indicates which integral is to be computed:

if **wt\_func** = **Nag\_Cosine**,  $w(x) = \cos(\omega x)$ ;

if **wt\_func** = **Nag\_Sine**,  $w(x) = \sin(\omega x)$ .

*Constraint:* **wt\_func** = **Nag\_Cosine** or **Nag\_Sine**.

5: **maxintervals** – Integer *Input*

*On entry:* an upper bound on the number of intervals  $C_k$  needed for the integration.

*Suggested value:* **maxintervals** = 50 is adequate for most problems.

*Constraint:* **maxintervals**  $\geq 3$ .

6: **maxsubints\_per\_interval** – Integer *Input*

*On entry:* the upper bound on the number of sub-intervals into which the interval of integration may be divided by the function. The more difficult the integrand, the larger **max\_num\_subint** should be.

*Suggested values:* a value in the range 200 to 500 is adequate for most problems.

*Constraint:* **max\_num\_subint**  $\geq 1$ .

7: **epsabs** – double *Input*

*On entry:* the absolute accuracy required. If **epsabs** is negative, the absolute value is used. See Section 6.1.

8:	<b>result</b> – double *	<i>Output</i>
<i>On exit:</i> the approximation to the integral $I$ .		
9:	<b>abserr</b> – double *	<i>Output</i>
<i>On exit:</i> an estimate of the modulus of the absolute error, which should be an upper bound for $ I - \text{result} $ .		
10:	<b>qpsub</b> – Nag_QuadSubProgress *	<i>Output</i>
Pointer to structure of type <b>Nag_QuadSubProgress</b> with the following members:		
	<b>intervals</b> – Integer	<i>Output</i>
<i>On exit:</i> the number of intervals $C_k$ actually used for the integration.		
	<b>fun_count</b> – Integer	<i>Output</i>
<i>On exit:</i> the number of function evaluations performed by nag_1d_quad_inf_wt_trig_1.		
	<b>subints_per_interval</b> – Integer *	<i>Output</i>
<i>On exit:</i> the maximum number of sub-intervals actually used for integrating over any of the intervals $C_k$ .		
	<b>interval_error</b> – double *	<i>Output</i>
<i>On exit:</i> the error estimate corresponding to the integral contribution over the interval $C_k$ , for $k = 1, 2, \dots, \text{qpsub.intervals}$ .		
	<b>interval_result</b> – double *	<i>Output</i>
<i>On exit:</i> the corresponding integral contribution over the interval $C_k$ , for $k = 1, 2, \dots, \text{qpsub.intervals}$ .		
	<b>interval_flag</b> – Integer *	<i>Output</i>
<i>On exit:</i> the error flag corresponding to <b>qpsub.interval_result</b> , for $k = 1, 2, \dots, \text{qpsub.intervals}$ . See also Section 5.		
When the information available in the arrays <b>interval_error</b> , <b>interval_result</b> and <b>interval_flag</b> is no longer useful, or before a subsequent call to nag_1d_quad_inf_wt_trig_1 with the same parameter <b>qpsub</b> is made, the user should free the storage contained in this pointer using the NAG macro <b>NAG_FREE</b> . Note that these arrays do not need to be freed if one of the error exits <b>NE_INT_ARG_LT</b> , <b>NE_BAD_PARAM</b> or <b>NE_ALLOC_FAIL</b> occurred.		
11:	<b>comm</b> – Nag_User *	<i>Input/Output</i>
<i>On entry/on exit:</i> pointer to a structure of type <b>Nag_User</b> with the following member:		
	<b>p</b> – Pointer	<i>Input/Output</i>
<i>On entry/on exit:</i> the pointer <b>p</b> , of type <b>Pointer</b> , allows the user to communicate information to and from the user-defined function <b>g()</b> . An object of the required type should be declared by the user, e.g., a structure, and its address assigned to the pointer <b>p</b> by means of a cast to <b>Pointer</b> in the calling program, e.g., <b>comm.p = (Pointer)&amp;s</b> . The type <b>Pointer</b> is <b>void *</b> .		
12:	<b>fail</b> – NagError *	<i>Input/Output</i>
The NAG error parameter (see the Essential Introduction).		
Users are recommended to declare and initialise <b>fail</b> and set <b>fail.print = TRUE</b> for this function.		

## 5 Error Indicators and Warnings

### NE\_INT\_ARG\_LT

On entry, **maxsubints\_per\_interval** must not be less than 1: **maxsubints\_per\_interval** = <*value*>.

On entry, **maxintervals** must not be less than 3: **maxintervals** = <*value*>.

### NE\_BAD\_PARAM

On entry, parameter **wt\_func** had an illegal value.

### NE\_ALLOC\_FAIL

Memory allocation failed.

### NE\_QUAD\_MAX\_SUBDIV

The maximum number of subdivisions has been reached: **maxsubints\_per\_interval** = <*value*>. The maximum number of subdivisions within an interval has been reached without the accuracy requirements 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 this function on the infinite subrange and an appropriate integrator on the finite subrange. Alternatively, consider relaxing the accuracy requirements specified by **epsabs** or increasing the value of **maxsubints\_per\_interval**.

### NE\_QUAD\_ROUNDOFF\_ABS\_TOL

Round-off error prevents the requested tolerance from being achieved: **epsabs** = <*value*>.

The error may be underestimated. Consider relaxing the accuracy requirements specified by **epsabs**.

### NE\_QUAD\_BAD\_SUBDIV

Extremely bad integrand behaviour occurs around the sub-interval (<*value*>, <*value*>).

The same advice applies as in the case of **NE\_QUAD\_MAX\_SUBDIV**.

### NE\_QUAD\_ROUNDOFF\_EXTRAPL

Round-off error is detected during extrapolation.

The requested tolerance cannot be achieved, because the extrapolation does not increase the accuracy satisfactorily; the returned result is the best that can be obtained.

The same advice applies as in the case of **NE\_QUAD\_MAX\_SUBDIV**.

### NE\_QUAD\_BAD\_SUBDIV\_INT

Bad integration behaviour has occurred within one or more intervals.

### NE\_QUAD\_MAX\_INT

Maximum number of intervals allowed has been achieved. Increase the value of **maxintervals**.

### NE\_QUAD\_EXTRAPL\_INT

The extrapolation table constructed for convergence acceleration of the series formed by the integral contribution over the integral does not converge.

In the cases where **fail.code** = **NE\_QUAD\_BAD\_SUBDIV\_INT**, **NE\_QUAD\_MAX\_INT** or **NE\_QUAD\_EXTRAPL\_INT**, additional information about the cause of the error can be obtained from the array **qpsub→interval\_flag**, as follows:

**qpsub.interval\_flag**[*k* − 1] = 1

The maximum number of subdivisions (= **maxsubints\_per\_interval**) has been achieved on the *k*th interval.

**qpsub.interval\_flag**[ $k - 1$ ] = 2

Occurrence of round-off error is detected and prevents the tolerance imposed on the  $k$ th interval from being achieved.

**qpsub.interval\_flag**[ $k - 1$ ] = 3

Extremely bad integrand behaviour occurs at some points of the  $k$ th interval.

**qpsub.interval\_flag**[ $k - 1$ ] = 4

The integration procedure over the  $k$ th interval does not converge (to within the required accuracy) due to round-off in the extrapolation procedure invoked on this interval. It is assumed that the result on this interval is the best which can be obtained.

**qpsub.interval\_flag**[ $k - 1$ ] = 5

The integral over the  $k$ th interval is probably divergent or slowly convergent. It must be noted that divergence can occur with any other value of **qpsub.interval\_flag**[ $k - 1$ ].

If users declare and initialise **fail** and set **fail.print** = **TRUE** as recommended then

#### NE\_QUAD\_NO\_CONV

The integral is probably divergent or slowly convergent.

Please note that divergence can also occur with any error exit other than **NE\_INT\_ARG\_LT**, **NE\_BAD\_PARAM** or **NE\_ALLOC\_FAIL**.

may be produced supplemented by messages indicating more precisely where problems were encountered by the function. However, if the default error handling, **NAGERR\_DEFAULT**, is used then one of the following errors may occur. Please note the program will terminate when the first of such errors is detected.

#### NE\_QUAD\_MAX\_SUBDIV\_SPEC\_INT

The maximum number of subdivisions has been reached,

**maxsubints\_per\_interval** = <value> on the <value> interval.

**interval\_flag**[<value>] = <value> over sub-interval (<value>, <value>).

#### NE\_QUAD\_ROUNDOFF\_TOL\_SPEC\_INT

Round-off error prevents the requested tolerance from being achieved on the <value> interval.

**interval\_flag**[<value>] = <value> over sub-interval (<value>, <value>).

#### NE\_QUAD\_BAD\_SPEC\_INT

Bad integrand behaviour occurs at some points of the <value> interval.

**interval\_flag**[<value>] = <value> over sub-interval (<value>, <value>).

#### NE\_QUAD\_NO\_CONV\_SPEC\_INT

The integral has failed to converge on the <value> interval.

**interval\_flag**[<value>] = <value> over sub-interval (<value>, <value>).

#### NE\_QUAD\_BAD\_DIVERGENCE\_SPEC\_INT

The integral is probably divergent on the <value> interval.

**interval\_flag**[<value>] = <value> over sub-interval (<value>, <value>).

## 6 Further Comments

The time taken by **nag\_1d\_quad\_inf\_wt\_trig\_1** depends on the integrand and on the accuracy required.

## 6.1 Accuracy

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

$$|I - \text{result}| \leq |\text{epsabs}|$$

where **epsabs** is the user-specified absolute error tolerance. Moreover it returns the quantity **abserr** which, in normal circumstances, satisfies

$$|I - \text{result}| \leq \text{abserr} \leq |\text{epsabs}|.$$

## 6.2 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, 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

## 7 See Also

`nag_1d_quad_inf_1` (d01smc)  
`nag_1d_quad_wt_trig_1` (d01snc)

## 8 Example

To compute

$$\int_0^\infty \frac{1}{\sqrt{x}} \cos(\pi x/2) dx.$$

### 8.1 Program Text

```
/* nag_1d_quad_inf_wt_trig_1(d01ssc) Example Program
 *
 * Copyright 1998 Numerical Algorithms Group.
 *
 * Mark 5, 1998.
 *
 * Mark 6 revised, 2000.
 */

#include <nag.h>
#include <stdio.h>
#include <nag_stdl�.h>
#include <math.h>
#include <nagd01.h>
#include <nagx01.h>

static double g(double x, Nag_User *comm);

main()
{
    double a;
    double omega;
    double epsabs, abserr;
    Nag_TrigTransform wt_func;
    double result;
```

```

Nag_QuadSubProgress qpsub;
Integer maxintervals, maxsubint_per_int;
static NagError fail;
Nag_User comm;

Vprintf("d01ssc Example Program Results\n");
epsabs = 0.001;
a = 0.0;
omega = X01AAC * 0.5;
wt_func = Nag_Cosine;
maxintervals = 50;
maxsubint_per_int = 500;

d01ssc(g, a, omega, wt_func, maxintervals, maxsubint_per_int,
       epsabs, &result, &abserr, &qpsub, &comm, &fail);

Vprintf("a      - lower limit of integration = %10.4f\n", a);
Vprintf("b      - upper limit of integration = infinity\n");
Vprintf("epsabs - absolute accuracy requested = %9.2e\n", epsabs);
if (fail.code != NE_NOERROR)
    Vprintf("%s\n", fail.message);
if (fail.code != NE_INT_ARG_LT && fail.code != NE_BAD_PARAM &&
    fail.code != NE_ALLOC_FAIL)
{
    Vprintf("result - approximation to the integral = %9.5f\n", result);
    Vprintf("abserr - estimate of the absolute error = %9.2e\n", abserr);
    Vprintf("qpsub.fun_count - number of function evaluations = %4ld\n",
           qpsub.fun_count);
    Vprintf("qpsub.intervals - number of intervals used = %4ld\n",
           qpsub.intervals);
    Vprintf("qpsub.subints_per_interval - \n\
maximum number of subintervals used in any one interval = %4ld\n",
           qpsub.subints_per_interval);
    /* Free memory used by qpsub */
    NAG_FREE(qpsub.interval_error);
    NAG_FREE(qpsub.interval_result);
    NAG_FREE(qpsub.interval_flag);
    exit(EXIT_SUCCESS);
}
else
    exit(EXIT_FAILURE);
}

static double g(double x, Nag_User *comm)
{
    return (x > 0.0) ? 1.0/sqrt(x) : 0.0;
}

```

## 8.2 Program Data

None.

### 8.3 Program Results

```
d01ssc Example Program Results
a      - lower limit of integration =      0.0000
b      - upper limit of integration = infinity
epsabs - absolute accuracy requested = 1.00e-03

result - approximation to the integral = 1.00000
abserr - estimate of the absolute error = 5.92e-04
qpsub.fun_count - number of function evaluations = 380
qpsub.intervals - number of intervals used =      6
qpsub.subints_per_interval -
maximum number of subintervals used in any one interval =      8
```

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