nag_opt_lsq_no_deriv (e04fcc)

1. Purpose

nag_opt_lsq_no_deriv (e04fcc) is a comprehensive algorithm for finding an unconstrained minimum of a sum of squares of m nonlinear functions in n variables $(m \ge n)$. No derivatives are required.

The function nag_opt_lsq_no_deriv is intended for objective functions which have continuous first and second derivatives (although it will usually work even if the derivatives have occasional discontinuities).

2. Specification

```
#include <nag.h>
#include <nage04.h>
```

3. Description

This function is applicable to problems of the form:

Minimize
$$F(x) = \sum_{i=1}^{m} [f_i(x)]^2$$

where $x = (x_1, x_2, ..., x_n)^T$ and $m \ge n$. (The functions $f_i(x)$ are often referred to as 'residuals'.) The user must supply a C function, **lsqfun**, to calculate the values of the $f_i(x)$ at any point x.

From a starting point $x^{(1)}$, supplied by the user, nag_opt_lsq_no_deriv generates a sequence of points $x^{(2)}, x^{(3)}, \ldots$, which is intended to converge to a local minimum of F(x). The sequence of points is given by

$$x^{(k+1)} = x^{(k)} + \alpha^{(k)} p^{(k)}$$

where the vector $p^{(k)}$ is a direction of search, and $\alpha^{(k)}$ is chosen such that $F(x^{(k)} + \alpha^{(k)}p^{(k)})$ is approximately a minimum with respect to $\alpha^{(k)}$.

The vector $p^{(k)}$ used depends upon the reduction in the sum of squares obtained during the last iteration. If the sum of squares was sufficiently reduced, then $p^{(k)}$ is an approximation to the Gauss-Newton direction; otherwise additional function evaluations are made so as to enable $p^{(k)}$ to be a more accurate approximation to the Newton direction.

The method is designed to ensure that steady progress is made whatever the starting point, and to have the rapid ultimate convergence of Newton's method.

4. Parameters

\mathbf{m}

Input: the number m of residuals, $f_i(x)$

n

Input: the number n of variables, x_j . Constraint: $1 \leq n \leq m$.

lsqfun

The function **lsqfun**, supplied by the user, must calculate the vector of values $f_i(x)$ at any point x. (However, if the user does not wish to calculate the residuals at a particular x, there is the option of setting a parameter to cause nag_opt_lsq_no_deriv to terminate immediately.)

The specification of **lsqfun** is:

void lsq	<pre>fun(Integer m, Integer n, double x[], double fvec[], Nag_Comm *comm)</pre>
m n	
	Input: the numbers m and n of residuals and variables, respectively.
$\mathbf{x}[\mathbf{n}]$	Input: the point x at which the values of the f_i are required.
fvec[[m] Output: unless comm->flag is reset to a negative number, on exit fvec $[i-1]$ must contain the value of f_i at the point x , for $i = 1, 2,, m$.
com	m Pointer to structure of type Nag_Comm; the following members are relevant to lsqfun.
	<pre>flag - Integer Input: comm->flag contains a non-negative number. Output: if lsqfun resets comm->flag to some negative number then nag_opt_lsq_no_deriv will terminate immediately with the error indicator NE_USER_STOP. If fail is supplied to nag_opt_lsq_no_deriv, fail.errnum will be set to the user's setting of comm->flag.</pre>
	<pre>first - Boolean Input: the value TRUE on the first call to lsqfun and FALSE for all subsequent calls.</pre>
	nf – Integer Input: the number of calls made to lsqfun including the current one.
	user - double * iuser - Integer * p - Pointer The tune Beinten will be weid a with a C compiler that defines weid a
	The type Pointer will be void * with a C compiler that defines void * and char * otherwise. Before calling nag_opt_lsq_no_deriv these pointers may be allocated memory by the user and initialised with various quantities for use by lsqfun when called from nag_opt_lsq_no_deriv.

Note: lsqfun should be tested separately before being used in conjunction with nag_opt_lsq_no_deriv. The array x must not be changed within lsqfun.

 $\mathbf{x}[\mathbf{n}]$

Input: $\mathbf{x}[j-1]$ must be set to a guess at the *j*th component of the position of the minimum, for j = 1, 2, ..., n.

Output: the final point x^* . On successful exit, $\mathbf{x}[j-1]$ is the *j*th component of the estimated position of the minimum.

fsumsq

Output: the value of F(x), the sum of squares of the residuals $f_i(x)$, at the final point given in **x**.

$\mathbf{fvec}[\mathbf{m}]$

Output: $\mathbf{fvec}[i-1]$ is the value of the residual $f_i(x)$ at the final point given in \mathbf{x} , for $i = 1, 2, \ldots, m$.

fjac[m][tdj]

Output: **fjac**[i-1][j-1] contains the estimate of the first derivative $\frac{\partial f_i}{\partial x_j}$ at the final point given in **x**, for i = 1, 2, ..., m; j = 1, 2, ..., n.

tdj

Input: the last dimension of the array ${\bf fjac}$ as declared in the function from which <code>nag_opt_lsq_no_deriv</code> is called.

Constraint: $\mathbf{tdj} \geq \mathbf{n}$.

options

Input/Output: a pointer to a structure of type Nag_E04_Opt whose members are optional parameters for nag_opt_lsq_no_deriv. These structure members offer the means of adjusting some of the parameter values of the algorithm and on output will supply further details of the results. A description of the members of **options** is given below in Section 7.

If any of these optional parameters are required then the structure **options** should be declared and initialised by a call to nag_opt_init (e04xxc) and supplied as an argument to nag_opt_lsq_no_deriv. However, if the optional parameters are not required the NAG defined null pointer, E04_DEFAULT, can be used in the function call.

comm

Input/Output: structure containing pointers for communication to user-supplied functions; see the above description of lsqfun for details. If the user does not need to make use of this communication feature the null pointer NAGCOMM_NULL may be used in the call to nag_opt_lsq_no_deriv; comm will then be declared internally for use in calls to user-supplied functions.

fail

The NAG error parameter, see the Essential Introduction to the NAG C Library. Users are recommended to declare and initialise fail and set fail.print = \mathbf{TRUE} for this function.

4.1. Description of Printed Output

Intermediate and final results are printed out by default. The level of printed output can be controlled by the user with the option **print_level** (see Section 7.2.).The default print level of **Nag_Soln_Iter** provides a single line of output at each iteration and the final result. The line of results printed at each iteration gives:

Itn	the current iteration number k .
Nfun	the cumulative number of calls to lsqfun .
Objective	the current value of the objective function, $F(x^{(k)})$.
Norm g	the Euclidean norm of the gradient of $F(x^{(k)})$.
Norm x	the Euclidean norm of $x^{(k)}$.
Norm(x(k-1)-x(k))	the Euclidean norm of $x^{(k-1)} - x^{(k)}$.
Step	the step $\alpha^{(k)}$ taken along the computed search direction $p^{(k)}$.

The printout of the final result consists of:

х	the final point x^* .
g	the estimate of the gradient of F at the final point.
Residuals	the values of the residuals f_i at the final point.
Sum of squares	the value of $F(x^*)$, the sum of squares of the residuals at the final point.

5. Comments

A list of possible error exits and warnings from nag_opt_lsq_no_deriv is given in Section 8.

5.1. Preliminary comments on accuracy

If the problem is reasonably well scaled and a successful exit is made, then, for a computer with a mantissa of t decimals, one would expect to get about t/2 - 1 decimals accuracy in the components of x and between t - 1 (if F(x) is of order 1 at the minimum) and 2t - 2 (if F(x) is close to zero at the minimum) decimals accuracy in F(x).

Further details about accuracy are given in Section 9.

6. Example 1

To find least-squares estimates of x_1, x_2 and x_3 in the model

$$y = x_1 + \frac{t_1}{x_2 t_2 + x_3 t_3}$$

using the 15 sets of data given in the following table.

y	t_1	t_2	t_3
0.14	1.0	15.0	1.0
0.18	2.0	14.0	2.0
0.22	3.0	13.0	3.0
0.25	4.0	12.0	4.0
0.29	5.0	11.0	5.0
0.32	6.0	10.0	6.0
0.35	7.0	9.0	7.0
0.39	8.0	8.0	8.0
0.37	9.0	7.0	7.0
0.58	10.0	6.0	6.0
0.73	11.0	5.0	5.0
0.96	12.0	4.0	4.0
1.34	13.0	3.0	3.0
2.10	14.0	2.0	2.0
4.39	15.0	1.0	1.0

The program uses (0.5, 1.0, 1.5) as the initial guess at the position of the minimum.

This example shows the simple use of nag_opt_lsq_no_deriv where default values are used for all optional parameters. An example showing the use of optional parameters is given in Section 12. There is however only one example program file, the main program of which calls both examples. The main program and example 1 are given below.

6.1. Program Text

```
/* nag_opt_lsq_no_deriv(e04fcc) Example Program
 * Copyright 1991 Numerical Algorithms Group.
 * Mark 2, 1991.
 */
#include <nag.h>
#include <stdio.h>
#include <nag_stdlib.h>
#include <math.h>
#include <nage04.h>
#include <nagx02.h>
#ifdef NAG_PROTO
static void lsqfun1(Integer m, Integer n, double x[], double fvec[],
                    Nag_Comm *comm);
static void lsqfun2(Integer m, Integer n, double x[], double fvec[],
                    Nag_Comm *comm);
static void ex1(void);
static void ex2(void);
#else
static void lsqfun1();
static void lsqfun2();
static void ex1();
static void ex2();
#endif
#define MMAX 15
#define NMAX 3
#define TMAX 3
```

```
/* Define a user structure template to store data in lsqfun. */
struct user
{
  double y[MMAX];
  double t[MMAX][TMAX];
};
main()
{
  /* Two examples are called, ex1() which uses the
   * default settings to solve the problem and
   * ex2() which solves the same problem with
   * some optional parameters set by the user.
   */
  Vprintf("e04fcc Example Program Results.\n");
  Vscanf(" %*[^\n]"); /* Skip heading in data file */
  ex1();
  ex2();
  exit(EXIT_SUCCESS);
}
static void ex1()
ł
  double fjac[MMAX][NMAX], fvec[MMAX], x[NMAX];
  Integer m, n, tdj;
  double fsumsq;
  static NagError fail;
  Vprintf("\ne04fcc example 1: no option setting.\n");
  Vscanf(" %*[^\n]"); /* Skip heading in data file */
  n = 3;
  m = 15;
  tdj = NMAX;
  /* Set up the starting point */
  x[0] = 0.5;
  x[1] = 1.0;
  x[2] = 1.5;
  /* Call the optimization routine */
  fail.print = TRUE;
e04fcc(m, n, lsqfun1, x, &fsumsq, fvec, (double *)fjac, tdj,
         E04_DEFAULT, NAGCOMM_NULL, &fail);
  if (fail.code != NE_NOERROR && fail.code != NW_COND_MIN) exit(EXIT_FAILURE);
}
                                  /* ex1 */
#ifdef NAG_PROTO
static void lsqfun1(Integer m, Integer n, double x[], double fvec[],
                    Nag_Comm *comm)
#else
     static void lsqfun1(m, n, x, fvec, comm)
     Integer m, n;
     double x[], fvec[];
     Nag_Comm *comm;
#endif
{
  /* Function to evaluate the residuals for example 1.
   * In this example a static variable is used to hold the
   * initial observations. The data is read into the structure
   * gs on the first call to lsqfun1(), it could alternatively
   * be read in from within main().
   */
  static struct user gs;
  Integer i, j, nt;
```

6.2. Program Data

3

eO4fcc Example Program Data

Example data for ex1: no option setting 0.14 1.0 15.0 1.0 0.18 2.0 14.0 2.0 0.22 3.0 13.0 3.0 0.25 4.0 12.0 4.0 0.29 5.0 11.0 5.0 0.32 6.0 10.0 6.0 0.35 7.0 9.0 7.0 0.39 8.0 8.0 8.0 0.37 9.0 7.0 7.0 0.58 10.0 6.0 6.0 0.73 11.0 5.0 5.0 0.96 12.0 4.0 4.0 1.34 13.0 3.0 3.0 2.10 14.0 2.0 2.0 4.39 15.0 1.0 1.0

6.3. Program Results

eO4fcc Example Program Results.

e04fcc example 1: no option setting.

Parameters to e04fcc

Number of residuals 15	Number of variables 3
optim_tol 1.05e-08 step_max 1.00e+05 print_levelNag_Soln_Iter outfile stdout	linesearch_tol 5.00e-01 max_iter 50 machine precision 1.11e-16
Memory allocation: sNag vNag Nag	tdv 3
Results from e04fcc:	

Iteration results:

Itn	Nfun	Objective	Norm g	Norm x	Norm $(x(k-1)-x(k))$	Step
0	4	1.0210e+01	3.2e+01	1.9e+00		
1	8	1.9873e-01	2.8e+00	2.4e+00	7.2e-01	1.0e+00
2	12	9.2324e-03	1.9e-01	2.6e+00	2.5e-01	1.0e+00
3	16	8.2149e-03	1.2e-03	2.6e+00	2.7e-02	1.0e+00
4	25	8.2149e-03	1.2e-07	2.6e+00	3.8e-04	1.0e+00
5	31	8.2149e-03	1.7e-10	2.6e+00	4.2e-06	1.0e+00

Final solution:

х	g	Residuals
8.24106e-02	-6.1762e-12	-5.8811e-03
1.13304e+00	1.4264e-10	-2.6535e-04
2.34370e+00	9.4150e-11	2.7469e-04
		6.5415e-03
		-8.2299e-04
		-1.2995e-03
		-4.4631e-03
		-1.9963e-02
		8.2216e-02
		-1.8212e-02
		-1.4811e-02
		-1.4710e-02
		-1.1208e-02
		-4.2040e-03
		6.8079e-03

The sum of squares is 8.2149e-03.

7. Optional Parameters

A number of optional input and output parameters to nag_opt_lsq_no_deriv are available through the structure argument **options**, type **Nag_E04_Opt**. A parameter may be selected by assigning an appropriate value to the relevant structure member; those parameters not selected will be assigned default values. If no use is to be made of any of the optional parameters the user should use the NAG defined null pointer, E04_DEFAULT, in place of **options** when calling nag_opt_lsq_no_deriv; the default settings will then be used for all parameters.

Before assigning values to **options** directly the structure **must** be initialised by a call to the function nag_opt_init (e04xxc). Values may then be assigned to the structure members in the normal C manner.

Option settings may also be read from a text file using the function nag_opt_read (e04xyc) in which case initialisation of the **options** structure will be performed automatically if not already done. Any subsequent direct assignment to the **options** structure must **not** be preceded by initialisation.

If assignment of functions and memory to pointers in the **options** structure is required, this must be done directly in the calling program, they cannot be assigned using nag_opt_read (e04xyc).

7.1. Optional Parameter Checklist and Default Values

For easy reference, the following list shows the members of **options** which are valid for nag_opt_lsq_no_deriv together with their default values where relevant. The number ϵ is a generic notation for **machine precision** (see nag_machine_precision (X02AJC)).

Boolean list	TRUE
Nag_PrintType print_level	Nag_Soln_Iter
char outfile[80]	stdout
<pre>void (*print_fun)()</pre>	NULL
Integer max_iter	$\max(50, 5\mathbf{n})$
double optim_tol	$\sqrt{\epsilon}$
double linesearch_tol	0.5 (0.0 if n = 1)
double step_max	100000.0
double *s	size n
double *v	size n*n
Integer tdv	n
Integer grade	
Integer iter	
Integer nf	

7.2. Description of Optional Parameters

list – Boolean

Default = TRUE

Input: if **options.list** = **TRUE** the parameter settings in the call to nag_opt_lsq_no_deriv will be printed.

Default = Nag_Soln_Iter

$print_level-Nag_PrintType$

Input: the level of results printout produced by nag_opt_lsq_no_deriv. The following values are available.

Nag_NoPrint	No output.
, Nag_Soln	The final solution.
Nag_Iter	One line of output for each iteration.
Nag_Soln_Iter	The final solution and one line of output for each iteration.
Nag_Soln_Iter_Full	The final solution and detailed printout at each iteration.

Details of each level of results printout are described in Section 7.3.

 $Constraint: options.print_level = Nag_NoPrint \text{ or } Nag_Soln \text{ or } Nag_Soln_Iter \text{ or } Nag_Soln_Iter_Full.$

outfile – char[80]

Input: the name of the file to which results should be printed. If **options.outfile** $[0] = ' \setminus 0'$ then the stdout stream is used.

print_fun – pointer to function

Input: printing function defined by the user; the prototype of **print_fun** is

void (*print_fun)(const Nag_Search_State *st, Nag_Comm *comm);

See Section 7.3.1. below for further details.

max_iter - Integer

Input: the limit on the number of iterations allowed before termination.

Constraint: **options.max_iter** ≥ 0 .

optim_tol – double

Input: the accuracy in x to which the solution is required.

If x_{true} is the true value of x at the minimum, then x_{sol} , the estimated position prior to a normal exit, is such that

$$||x_{sol} - x_{true}|| < \textbf{optim_tol} \times (1.0 + ||x_{true}||),$$

where $||y|| = \sqrt{\sum_{j=1}^{n} y_j^2}$. For example, if the elements of x_{sol} are not much larger than 1.0

in modulus and if **optim_tol** = 1.0×10^{-5} , then x_{sol} is usually accurate to about 5 decimal places. (For further details see Section 9.)

If F(x) and the variables are scaled roughly as described in Section 9)and ϵ is the **machine precision**, then a setting of order **optim_tol** = $\sqrt{\epsilon}$ will usually be appropriate. Constraint: $10\epsilon \leq \text{options.optim_tol} < 1.0$.

linesearch_tol - double

Default = 0.5. (If n = 1, default = 0.0)

Input: Every iteration of nag_opt_lsq_no_deriv involves a linear minimization i.e., minimization of $F(x^{(k)} + \alpha^{(k)}p^{(k)})$ with respect to $\alpha^{(k)}$.

linesearch_tol specifies how accurately the linear minimizations are to be performed. The minimum with respect to $\alpha^{(k)}$ will be located more accurately for small values of **linesearch_tol** (say 0.01) than for large values (say 0.9). Although accurate linear minimizations will generally reduce the number of iterations performed by nag_opt_lsq_no_deriv, they will increase the number of calls of **lsqfun** made each iteration. On balance it is usually more efficient to perform a low accuracy minimization.

Constraint: $0.0 \leq \text{options.linesearch_tol} < 1.0$.

step_max – double

1

Default = 100000.0

Input: an estimate of the Euclidean distance between the solution and the starting point supplied by the user. (For maximum efficiency, a slight overestimate is preferable.) nag_opt_lsq_no_deriv will ensure that, for each iteration,

$$\sum_{j=1}^{n} (x_{j}^{(k)} - x_{j}^{(k-1)})^{2} \leq (\text{step_max})^{2}$$

Default = max(50, 5n)

Default = stdout

Default = NULL

Default = $\sqrt{\epsilon}$

where k is the iteration number. Thus, if the problem has more than one solution, nag_opt_lsq_no_deriv is most likely to find the one nearest to the starting point. On difficult problems, a realistic choice can prevent the sequence $x^{(k)}$ entering a region where the problem is ill-behaved and can help avoid overflow in the evaluation of F(x). However, an underestimate of **options.step_max** can lead to inefficiency. Constraint: **options.step_max** \geq **options.optim_tol**.

 \mathbf{s} – double *

Default memory $= \mathbf{n}$

Input: **n** values of memory will be automatically allocated by nag_opt_lsq_no_deriv and this is the recommended method of use of **options.s**. However a user may supply memory from the calling program.

Output: the singular values of the estimated Jacobian matrix at the final point. Thus **options.s** may be useful as information about the structure of the user's problem.

v - double *

Default memory $= n^*n$

Input: $\mathbf{n}^*\mathbf{n}$ values of memory will be automatically allocated by nag_opt_lsq_no_deriv and this is the recommended method of use of **options.v**. However a user may supply memory from the calling program.

Output: the matrix V associated with the singular value decomposition

 $J = USV^T$

of the estimated Jacobian matrix at the final point, stored by rows. This matrix may be useful for statistical purposes, since it is the matrix of orthonormalised eigenvectors of $J^T J$.

tdv - Integer

 $\mathrm{Default}=\mathbf{n}$

Input: if memory is supplied by the user then **options.tdv** must contain the last dimension of the array assigned to **options.tdv** as declared in the function from which nag_opt_lsq_no_deriv is called.

Output: the trailing dimension used by **options.v**. If the Nag default memory allocation has been used this value will be \mathbf{n} .

Constraint: $options.tdv \ge n$.

grade – Integer

Output: the grade of the Jacobian at the final point. nag_opt_lsq_no_deriv estimates the dimension of the subspace for which the Jacobian matrix can be used as a valid approximation to the curvature (see Gill and Murray, 1978); this estimate is called the grade.

iter – Integer

Output: the number of iterations which have been performed in nag_opt_lsq_no_deriv.

 $\mathbf{nf} - \mathrm{Integer}$

Output: the number of times the residuals have been evaluated (i.e., number of calls of lsqfun).

7.3. Description of Printed Output

The level of printed output can be controlled with the structure members **options.list** and **options.print_level** (see Section 7.2.). If **list** = **TRUE** then the parameter values to nag_opt_lsq_no_deriv are listed, whereas the printout of results is governed by the value of **print_level**. The default of **print_level** = **Nag_Soln_Iter** provides a single line of output at each iteration and the final result. This section describes all of the possible levels of results printout available from nag_opt_lsq_no_deriv.

When **options.print_level** = **Nag_Iter** or **Nag_Soln_Iter** a single line of output is produced on completion of each iteration, this gives the following values:

Itn	the current iteration number k .
Nfun	the cumulative number of calls to lsqfun .
Objective	the value of the objective function, $F(x^{(k)})$.
Norm g	the Euclidean norm of the gradient of $F(x^{(k)})$.
Norm x	the Euclidean norm of $x^{(k)}$.
Norm(x(k-1)-x(k))	the Euclidean norm of $x^{(k-1)} - x^{(k)}$.

Step the step $\alpha^{(k)}$ taken along the computed search direction $p^{(k)}$.

When $options.print_level = Nag_Soln_Iter_Full$ more detailed results are given at each iteration. Additional values output are

Grade	the grade of the Jacobian matrix. (See description of grade , Section 7.2)
x	the current point $x^{(k)}$.
g	the current estimate of the gradient of $F(x^{(k)})$.
Singular values	the singular values of the current approximation to the Jacobian matrix.

If $options.print_level = Nag_Soln$ or Nag_Soln_Iter or Nag_Soln_Iter_Full the final result is printed out. This consists of:

x	the final point x^* .
g	the estimate of the gradient of F at the final point.
Residuals	the values of the residuals f_i at the final point.
Sum of squares	the value of $F(x^*)$, the sum of squares of the residuals at the final point.

If **options.print_level** = $Nag_NoPrint$ then printout will be suppressed; the user can print the final solution when $nag_opt_lsq_no_deriv$ returns to the calling program.

7.3.1. Output of results via a user defined printing function

Users may also specify their own print function for output of iteration results and the final solution by use of the **options.print_fun** function pointer, which has prototype

void (*print_fun)(const Nag_Search_State *st, Nag_Comm *comm);

The rest of this section can be skipped if the default printing facilities provide the required functionality.

When a user defined function is assigned to **options.print_fun** this will be called in preference to the internal print function of nag_opt_lsq_no_deriv. Calls to the user defined function are again controlled by means of the **options.print_level** member. Information is provided through **st** and **comm**, the two structure arguments to **print_fun**. If **comm->it_prt = TRUE** then the results from the last iteration of nag_opt_lsq_no_deriv are in the following members of **st**:

\mathbf{m} – Integer

the number of residuals.

 \mathbf{n} – Integer

the number of variables.

 \mathbf{x} – double *

points to the **n** memory locations holding the current point $x^{(k)}$.

fvec – double *

points to the ${\bf m}$ memory locations holding the values of the residuals f_i at the current point $x^{(k)}.$

fjac – double *

points to **m*st->tdj** memory locations. **fjac**[(i-1)***st->tdj** + (j-1)] contains the value of $\frac{\partial f_i}{\partial x_i}$, for i = 1, 2, ..., m; j = 1, 2, ..., n at the current point $x^{(k)}$.

tdj – Integer

the trailing dimension for **st->fjac**[].

step – double

the step $\alpha^{(k)}$ taken along the search direction $p^{(k)}$.

 $xk_norm - double$

the Euclidean norm of $x^{(k-1)} - x^{(k)}$.

 \mathbf{g} – double *

points to the **n** memory locations holding the estimated gradient of F at the current point $x^{(k)}$.

 $\mathbf{grade} - \mathrm{Integer}$

the grade of the Jacobian matrix.

${\bf s}$ – double *

points to the ${\bf n}$ memory locations holding the singular values of the current approximation to the Jacobian.

iter – Integer

the number of iterations, k, performed by nag_opt_lsq_no_deriv.

nf – Integer

the cumulative number of calls made to lsqfun.

The relevant members of the structure **comm** are:

it_prt – Boolean

will be **TRUE** when the print function is called with the result of the current iteration.

sol_prt – Boolean

will be **TRUE** when the print function is called with the final result.

```
user – double *
```

iuser – Integer *

\mathbf{p} – Pointer

pointers for communication of user information. If used they must be allocated memory by the user either before entry to nag_opt_lsq_no_deriv or during a call to **lsqfun** or **print_fun**. The type Pointer will be **void** * with a C compiler that defines **void** * and **char** * otherwise.

8. Error Indications and Warnings

NE_USER_STOP

User requested termination, user flag value = $\langle value \rangle$.

This exit occurs if the user sets **comm->flag** to a negative value in **lsqfun**. If **fail** is supplied the value of **fail.errnum** will be the same as the user's setting of **comm->flag**.

NE_INT_ARG_LT

On entry, **n** must not be less than 1: $\mathbf{n} = \langle value \rangle$.

NE_2_INT_ARG_LT

On entry, $\mathbf{m} = \langle value \rangle$ while $\mathbf{n} = \langle value \rangle$. These parameters must satisfy $\mathbf{m} \ge \mathbf{n}$. On entry, $\mathbf{tdj} = \langle value \rangle$ while $\mathbf{n} = \langle value \rangle$. These parameters must satisfy $\mathbf{tdj} \ge \mathbf{n}$. On entry, **options.tdv** = $\langle value \rangle$ while $\mathbf{n} = \langle value \rangle$. These parameters must satisfy $\mathbf{tdv} \ge \mathbf{n}$.

NE_OPT_NOT_INIT

Options structure not initialised.

NE_BAD_PARAM

On entry parameter **options.print_level** had an illegal value.

NE_2_REAL_ARG_LT

On entry, **options.step_max** = $\langle value \rangle$ while **options.optim_tol** = $\langle value \rangle$. These parameters must satisfy **step_max** \geq **optim_tol**.

NE_INVALID_INT_RANGE_1

Value $\langle value \rangle$ given to **options.max_iter** not valid. Correct range is **max_iter** ≥ 0 .

NE_INVALID_REAL_RANGE_EF

Value $\langle value \rangle$ given to **options.optim_tol** not valid. Correct range is $\langle value \rangle \leq$ **optim_tol** < 1.0.

NE_INVALID_REAL_RANGE_FF

Value $\langle value \rangle$ given to **options.linesearch_tol** not valid. Correct range is $0.0 \leq \text{linesearch_tol} < 1.0$.

NE_ALLOC_FAIL

Memory allocation failed.

If one of the above exits occurs, no values will have been assigned to **fsumsq**, or to the elements of **fvec**, **fjac**, **options.s** or **options.v**.

NW_TOO_MANY_ITER

The maximum number of iterations, $\langle value \rangle$, have been performed.

If steady reductions in the sum of squares, F(x), were monitored up to the point where this exit occurred, then the exit probably occurred simply because **options.max_iter** was set too small, so the calculations should be restarted from the final point held in **x**. This exit may also indicate that F(x) has no minimum.

NW_COND_MIN

The conditions for a minimum have not all been satisfied, but a lower point could not be found.

This could be because **options.optim_tol** has been set so small that rounding errors in the evaluation of the residuals make attainment of the convergence conditions impossible.

NE_SVD_FAIL

The computation of the singular value decomposition of the Jacobian matrix has failed to converge in a reasonable number of sub-iterations.

It may be worth applying nag_opt_lsq_no_deriv again starting with an initial approximation which is not too close to the point at which the failure occurred.

The exits **NW_TOO_MANY_ITER**, **NW_COND_MIN**, and **NE_SVD_FAIL** may also be caused by mistakes in **lsqfun**, by the formulation of the problem or by an awkward function. If there are no such mistakes it is worth restarting the calculations from a different starting point (not the point at which the failure occurred) in order to avoid the region which caused the failure.

NE_NOT_APPEND_FILE

Cannot open file $\langle string \rangle$ for appending.

NE_WRITE_ERROR

Error occurred when writing to file $\langle string \rangle$.

NE_NOT_CLOSE_FILE

Cannot close file $\langle string \rangle$.

9. Further Comments

The number of iterations required depends on the number of variables, the number of residuals, the behaviour of F(x), the accuracy demanded and the distance of the starting point from the solution. The number of multiplications performed per iteration of nag_opt_lsq_no_deriv varies, but for m >> n is approximately $n \times m^2 + O(n^3)$. In addition, each iteration makes at least n + 1 calls of **lsqfun**. So, unless the residuals can be evaluated very quickly, the run time will be dominated by the time spent in **lsqfun**.

Ideally, the problem should be scaled so that, at the solution, F(x) and the corresponding values of the x_j are each in the range (-1, +1), and so that at points one unit away from the solution, F(x) differs from its value at the solution by approximately one unit. This will usually imply that the Hessian matrix of F(x) at the solution is well-conditioned. It is unlikely that the user will be able to follow these recommendations very closely, but it is worth trying (by guesswork), as sensible scaling will reduce the difficulty of the minimization problem, so that nag_opt_lsq_no_deriv will take less computer time.

When the sum of squares represents the goodness of fit of a nonlinear model to observed data, elements of the variance-covariance matrix of the estimated regression coefficients can be computed by a subsequent call to nag_opt_lsq_covariance (e04ycc), using information returned in the arrays **options.s** and **options.v**. See nag_opt_lsq_covariance (e04ycc) for further details.

9.1. Accuracy

A successful exit (fail.code = NE_NOERROR) is made from nag_opt_lsq_no_deriv when (B1, B2 and B3) or B4 or B5 hold, where

$$\begin{split} B1 &\equiv \alpha^{(k)} \times \parallel p^{(k)} \parallel < (\textbf{optim_tol} + \epsilon) \times (1.0 + \parallel x^{(k)} \parallel) \\ B2 &\equiv |F^{(k)} - F^{(k-1)}| < (\textbf{optim_tol} + \epsilon)^2 \times (1.0 + F^{(k)}) \end{split}$$

and where $||.||, \epsilon$ and the optional parameter **optim_tol** are as defined in Section 7.2, while $F^{(k)}$ and $g^{(k)}$ are the values of F(x) and its vector of estimated first derivatives at $x^{(k)}$.

If **fail.code** = **NE_NOERROR** then the vector in **x** on exit, x_{sol} , is almost certainly an estimate of x_{true} , the position of the minimum to the accuracy specified by **options.optim_tol**.

If fail.code = NW_COND_MIN, then x_{sol} may still be a good estimate of x_{true} , but to verify this the user should make the following checks. If

(a) the sequence $\{F(x^{(k)})\}$ converges to $F(x_{sol})$ at a superlinear or a fast linear rate, and (b) $g(x_{sol})^T g(x_{sol}) < 10\epsilon$,

where T denotes transpose, then it is almost certain that x_{sol} is a close approximation to the minimum. When (b) is true, then usually $F(x_{sol})$ is a close approximation to $F(x_{true})$.

Further suggestions about confirmation of a computed solution are given in the Chapter Introduction.

10. References

Gill P E and Murray W (1978) Algorithms for the Solution of the Nonlinear Least-squares Problem SIAM J. Numer. Anal. 15 977–992.

11. See Also

nag_opt_lsq_covariance (e04ycc) nag_opt_init (e04xxc) nag_opt_read (e04xyc) nag_opt_free (e04xzc)

12. Example 2

Example 2 solves the same problem as Example 1 but shows the use of certain optional parameters. This example shows option values being assigned directly within the program text and by reading values from a data file. The **options** structure is declared and initialised by nag_opt_init (e04xxc), a value is then assigned directly to option **optim_tol** and two further options are read from the data file by use of nag_opt_read (e04xyc). The memory freeing function nag_opt_free (e04xzc) is used to free the memory assigned to the pointers in the option structure. Users should **not** use the standard C function **free(**) for this purpose.

12.1. Program Text

```
static void ex2()
{
   double fjac[MMAX][NMAX], fvec[MMAX], x[NMAX];
   Integer i, j, m, n, nt, tdj;
   double fsumsq;
   Boolean print;
   Nag_E04_Opt options;
   Nag_Comm comm;
   static NagError fail, fail2;
   struct user s;

   Vprintf("\n\ne04fcc example 2: using option setting.\n");
   Vscanf(" %*[^\n]"); /* Skip heading in data file */
   n = 3;
   m = 15;
   tdj = NMAX;
   nt = 3;
```

```
/* Read data into structure.
   * Observations t (j = 0, 1, 2) are held in s \rightarrow t[i][j]
* (i = 0, 1, 2, . . . , 14)
   */
  nt = 3;
  for (i = 0; i < m; ++i)
    {
      Vscanf("%lf", &s.y[i]);
      for (j = 0; j < nt; ++j) Vscanf("%lf", &s.t[i][j]);</pre>
    }
  /* Set up the starting point */
  x[0] = 0.5;
  x[1] = 1.0;
  x[2] = 1.5;
  e04xxc(&options); /* Initialise options structure */
  /* Set one option directly. */
  options.optim_tol = 10.0*sqrt(X02AJC);
  /* Read remaining option values from file */
  fail.print = TRUE;
  print = FALSE;
  e04xyc("e04fcc", "stdin", &options, print, "", &fail);
  /* Assign address of user defined structure to
   * comm.p for communication to lsqfun2().
   */
  comm.p = (Pointer)&s;
  if (fail.code == NE_NOERROR)
    eO4fcc(m, n, lsqfun2, x, &fsumsq, fvec, (double *)fjac, tdj,
           &options, &comm, &fail);
  /* Free memory allocated to pointers s and v */
fail2.print = TRUE;
  e04xzc(&options, "all", &fail2);
  if ((fail.code != NE_NOERROR && fail.code != NW_COND_MIN)
      || fail2.code != NE_NOERROR) exit(EXIT_FAILURE);
}
                                  /* ex2 */
#ifdef NAG_PROTO
static void lsqfun2(Integer m, Integer n, double x[], double fvec[],
                     Nag_Comm *comm)
#else
     static void lsqfun2(m, n, x, fvec, comm)
     Integer m, n;
     double x[], fvec[];
     Nag_Comm *comm;
#endif
ł
  /* Function to evaluate the residuals for example 2.
   *
   * To avoid the use of a global varibale this example assigns the address
   * of a user defined structure to comm.p in the main program (where the
   * data was also read in).
   * The address of this structure is recovered in each call to lsqfun2()
   * from comm->p and the structure used in the calculation of the residuals.
   */
  Integer i;
  struct user *s = (struct user *)comm->p;
 for (i = 0; i < m; ++i)
    fvec[i] = x[0] + s->t[i][0] / (x[1]*s->t[i][1] + x[2]*s->t[i][2]) - s->y[i];
}
                                   /* lsqfun2 */
```

12.2. Program Data

Example data for ex2: using option setting

0.14	1.0	15.0	1.0
0.18	2.0	14.0	2.0
0.22	3.0	13.0	3.0
0.25	4.0	12.0	4.0
0.29	5.0	11.0	5.0
0.32	6.0	10.0	6.0
0.35	7.0	9.0	7.0
0.39	8.0	8.0	8.0
0.37	9.0	7.0	7.0
0.58	10.0	6.0	6.0
0.73	11.0	5.0	5.0
0.96	12.0	4.0	4.0
1.34	13.0	3.0	3.0
2.10	14.0	2.0	2.0
4.39	15.0	1.0	1.0

Following optional parameter settings are read by e04xyc

begin e04fcc

/* Results printout set to fullest detail */
print_level = Nag_Soln_Iter_Full

```
/* Estimate minimum will be within 10 units of the
 * starting point.
 */
step_max = 10.0
```

 ${\tt end}$

12.3. Program Results

eO4fcc example 2: using option setting.

Parameters to e04fcc

Number of res	iduals 15	Number of variables	. 3
step_max print_level	1.05e-07 1.00e+01 Nag_Soln_Iter_Full stdout	<pre>max_iter machine precision 1.11e</pre>	50
Memory alloca sv	Nag	tdv	3
Results from	e04fcc:		
Iteration res	ults:		
Itn Nfun 0 4	Objective Norm g 1.0210e+01 3.2e+01		cep Grade 3
x 5.00000e-01 1.00000e+00 1.50000e+00	-1.6838e+01		
Itn Nfun 1 8	Objective Norm g 1.9873e-01 2.8e+00		cep Grade De+00 3
x 8.24763e-02 1.13575e+00		Singular values 4.1973e+00 1.8396e+00	

2.06664e+00 -1.5073e+00 6.6356e-02 ItnNfunObjectiveNorm gNorm xNorm (x(k-1)-x(k))Step2129.2324e-031.9e-012.6e+002.5e-011.0e+00 Grade 1.0e+00 3 xgSingular valu8.24402e-021.3523e-014.1026e+001.13805e+00-9.4890e-021.6131e+002.31707e+00-9.4630e-026.1372e-02 Singular values ItnNfunObjectiveNormNormNorm(x(k-1)-x(k))Step3168.2149e-031.2e-032.6e+002.7e-021.0e+00 Grade 3 xgSingular values8.24150e-028.1961e-044.0965e+001.13323e+00-5.7539e-041.5951e+002.34337e+00-5.7660e-046.1250e-02 ItnNfunObjectiveNormNormNorm(x(k-1)-x(k))Step4258.2149e-031.2e-072.6e+003.8e-041.0e+00 Grade 2 g -7.4836e-08 8.5498e-08 4.1969e-08 Singular values 8.24107e-02 1.13304e+00 2.34369e+00 4.0965e+00 1.5950e+00 6.1258e-02 ItnNfunObjectiveNorm gNorm xNorm (x(k-1)-x(k))Step5318.2149e-031.7e-102.6e+004.2e-061.0e+00 Grade 2 xgSingular values8.24106e-02-6.1762e-124.0965e+001.13304e+001.4264e-101.5950e+002.34370e+009.4150e-116.1258e-02 Final solution: g -6.1762e-12 1.4264e-10 Residuals x 8.24106e-02 -5.8811e-03 1.13304e+00 -2.6535e-04 1.420-10 9.4150e-11 2.7469e-04 2.34370e+00 6.5415e-03 -8.2299e-04 -1.2995e-03 -4.4631e-03 -1.9963e-02 8.2216e-02 -1.8212e-02 -1.4811e-02 -1.4710e-02 -1.1208e-02 -4.2040e-03 6.8079e-03 The sum of squares is 8.2149e-03.