NAG Library Function Document nag linf fit (e02gcc)

1 Purpose

nag linf fit (e02gcc) calculates an l_{∞} solution to an over-determined system of linear equations.

2 Specification

3 Description

Given a matrix A with m rows and n columns $(m \ge n)$ and a vector b with m elements, the function calculates an l_{∞} solution to the over-determined system of equations

$$Ax = b$$
.

That is to say, it calculates a vector x, with n elements, which minimizes the l_{∞} norm of the residuals (the absolutely largest residual)

$$r(x) = \max_{1 \le i \le m} |r_i|$$

where the residuals r_i are given by

$$r_i = b_i - \sum_{j=1}^n a_{ij} x_j, \quad i = 1, 2, \dots, m.$$

Here a_{ij} is the element in row i and column j of A, b_i is the ith element of b and x_j the jth element of x. The matrix A need not be of full rank. The solution is not unique in this case, and may not be unique even if A is of full rank.

Alternatively, in applications where a complete minimization of the l_{∞} norm is not necessary, you may obtain an approximate solution, usually in shorter time, by giving an appropriate value to the argument **relerr**.

Typically in applications to data fitting, data consisting of m points with coordinates (t_i, y_i) is to be approximated in the l_{∞} norm by a linear combination of known functions $\phi_j(t)$,

$$\alpha_1\phi_1(t) + \alpha_2\phi_2(t) + \cdots + \alpha_n\phi_n(t).$$

This is equivalent to finding an l_{∞} solution to the over-determined system of equations

$$\sum_{i=1}^{n} \phi_j(t_i)\alpha_j = y_i, \quad i = 1, 2, \dots, m.$$

Thus if, for each value of i and j the element a_{ij} of the matrix A above is set equal to the value of $\phi_j(t_i)$ and b_i is set equal to y_i , the solution vector x will contain the required values of the α_j . Note that the independent variable t above can, instead, be a vector of several independent variables (this includes the case where each ϕ_i is a function of a different variable, or set of variables).

The algorithm is a modification of the simplex method of linear programming applied to the dual formation of the l_{∞} problem (see Barrodale and Phillips (1974) and Barrodale and Phillips (1975)). The modifications are designed to improve the efficiency and stability of the simplex method for this particular application.

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4 References

Barrodale I and Phillips C (1974) An improved algorithm for discrete Chebyshev linear approximation *Proc. 4th Manitoba Conf. Numerical Mathematics* 177–190 University of Manitoba, Canada

Barrodale I and Phillips C (1975) Solution of an overdetermined system of linear equations in the Chebyshev norm [F4] (Algorithm 495) ACM Trans. Math. Software 1(3) 264–270

5 Arguments

1: **order** – Nag OrderType

Input

On entry: the **order** argument specifies the two-dimensional storage scheme being used, i.e., row-major ordering or column-major ordering. C language defined storage is specified by **order** = Nag_RowMajor. See Section 3.2.1.3 in the Essential Introduction for a more detailed explanation of the use of this argument.

Constraint: order = Nag_RowMajor or Nag_ColMajor.

2: \mathbf{m} - Integer Input

On entry: the number of equations, m (the number of rows of the matrix A).

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

3: \mathbf{n} - Integer Input

On entry: the number of unknowns, n (the number of columns of the matrix A).

Constraint: $\mathbf{n} \geq 1$.

4: $\mathbf{a}[dim]$ – double Input/Output

Note: the dimension, dim, of the array **a** must be at least $(\mathbf{n} + 3) \times (\mathbf{m} + 1)$.

Where A(j,i) appears in this document, it refers to the array element

$$\mathbf{a}[(i-1)\times(\mathbf{n}+3)+j-1]$$
 when $\mathbf{order}=\mathrm{Nag_ColMajor};$ $\mathbf{a}[(j-1)\times(\mathbf{m}+1)+i-1]$ when $\mathbf{order}=\mathrm{Nag_RowMajor}.$

On entry: A(j, i) must contain a_{ij} , the element in the *i*th row and *j*th column of the matrix A, for i = 1, 2, ..., m and j = 1, 2, ..., n, (that is, the **transpose** of the matrix). The remaining elements need not be set. Preferably, the columns of the matrix A (rows of the argument **a**) should be scaled before entry: see Section 7.

On exit: contains the last simplex tableau.

5: $\mathbf{b}[\mathbf{m}]$ - double Input/Output

On entry: $\mathbf{b}[i-1]$ must contain b_i , the *i*th element of the vector b, for $i=1,2,\ldots,m$.

On exit: the *i*th residual r_i corresponding to the solution vector x, for i = 1, 2, ..., m. Note however that these residuals may contain few significant figures, especially when **resmax** is within one or two orders of magnitude of **tol**. Indeed if **resmax** \leq **tol**, the elements $\mathbf{b}[i-1]$ may all be set to zero. It is therefore often advisable to compute the residuals directly.

6: **tol** – double *Input*

On entry: a threshold below which numbers are regarded as zero. The recommended threshold value is $10.0 \times \epsilon$, where ϵ is the **machine precision**. If **tol** ≤ 0.0 on entry, the recommended value is used within the function. If premature termination occurs, a larger value for **tol** may result in a valid solution.

Suggested value: 0.0.

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7: **relerr** – double *

Input/Output

On entry: must be set to a bound on the relative error acceptable in the maximum residual at the solution

If relerr ≤ 0.0 , then the l_{∞} solution is computed, and relerr is set to 0.0 on exit.

If **relerr** > 0.0, then the function obtains instead an approximate solution for which the largest residual is less than 1.0 +**relerr** times that of the l_{∞} solution; on exit, **relerr** contains a smaller value such that the above bound still applies. (The usual result of this option, say with **relerr** = 0.1, is a saving in the number of simplex iterations).

On exit: is altered as described above.

8: $\mathbf{x}[\mathbf{n}]$ – double

Output

On exit: if an optimal but not necessarily unique solution is found, $\mathbf{x}[j-1]$ contains the jth element of the solution vector x, for $j=1,2,\ldots,n$. Whether this is an l_{∞} solution or an approximation to one, depends on the value of **relerr** on entry.

9: **resmax** – double *

Output

On exit: if an optimal but not necessarily unique solution is found, **resmax** contains the absolute value of the largest residual(s) for the solution vector x. (See **b**.)

10: rank – Integer *

Output

On exit: if an optimal but not necessarily unique solution is found, \mathbf{rank} contains the computed rank of the matrix A.

11: iter – Integer *

Output

On exit: if an optimal but not necessarily unique solution is found, iter contains the number of iterations taken by the simplex method.

12: **fail** – NagError *

Input/Output

The NAG error argument (see Section 3.6 in the Essential Introduction).

6 Error Indicators and Warnings

NE ALLOC FAIL

Dynamic memory allocation failed.

NE BAD PARAM

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

NE_INT

```
On entry, \mathbf{n} = \langle value \rangle.
Constraint: \mathbf{n} \geq 1.
```

NE INT 2

```
On entry, \mathbf{m} = \langle value \rangle and \mathbf{n} = \langle value \rangle.
Constraint: \mathbf{m} \geq \mathbf{n}.
```

NE INTERNAL ERROR

An internal error has occurred in this function. Check the function call and any array sizes. If the call is correct then please contact NAG for assistance.

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NE NON UNIQUE

An optimal solution has been obtained, but may not be unique.

NE_TERMINATION_FAILURE

Premature termination due to rounding errors. Try using larger value of **tol**: **tol** = $\langle value \rangle$.

7 Accuracy

Experience suggests that the computational accuracy of the solution x is comparable with the accuracy that could be obtained by applying Gaussian elimination with partial pivoting to the n+1 equations which have residuals of largest absolute value. The accuracy therefore varies with the conditioning of the problem, but has been found generally very satisfactory in practice.

8 Parallelism and Performance

Not applicable.

9 Further Comments

The effects of m and n on the time and on the number of iterations in the simplex method vary from problem to problem, but typically the number of iterations is a small multiple of n and the total time is approximately proportional to mn^2 .

It is recommended that, before the function is entered, the columns of the matrix A are scaled so that the largest element in each column is of the order of unity. This should improve the conditioning of the matrix, and also enable the argument **tol** to perform its correct function. The solution x obtained will then, of course, relate to the scaled form of the matrix. Thus if the scaling is such that, for each $j=1,2,\ldots,n$, the elements of the jth column are multiplied by the constant k_j , the element x_j of the solution vector x must be multiplied by k_j if it is desired to recover the solution corresponding to the original matrix A.

10 Example

This example approximates a set of data by a curve of the form

$$y = Ke^t + Le^{-t} + M$$

where K, L and M are unknown. Given values y_i at 5 points t_i we may form the over-determined set of equations for K, L and M

$$e^{t_i}K + e^{-t_i}L + M = y_i, \quad i = 1, 2, \dots, 5.$$

nag linf fit (e02gcc) is used to solve these in the l_{∞} sense.

10.1 Program Text

```
/* nag_linf_fit (e02gcc) Example Program.
    * Copyright 2001 Numerical Algorithms Group.
    * Mark 7, 2001.
    */
#include <stdio.h>
#include <math.h>
#include <nag.h>
#include <naged2.h>
int main(void)
{
```

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```
/* Scalars */
 double
                relerr, resmax, t, tol;
 Integer
                exit_status, i, irank, iter, m, n, pda;
 NagError
                fail;
 Nag_OrderType order;
  /* Arrays */
                *a = 0, *b = 0, *x = 0;
 double
#ifdef NAG COLUMN MAJOR
#define A(I, J) a[(J-1)*pda + I - 1]
 order = Nag_ColMajor;
#else
#define A(I, J) a[(I-1)*pda + J - 1]
 order = Nag_RowMajor;
#endif
 INIT_FAIL(fail);
 exit_status = 0;
 printf("nag linf fit (e02qcc) Example Program Results\n");
  /* Skip heading in data file */
 scanf("%*[^\n] ");
 n = 3;
 scanf("%ld%*[^\n] ", &m);
 if (m > 0)
      /* Allocate memory */
      if (!(a = NAG\_ALLOC((n+3)*(m+1), double)))
          !(b = NAG_ALLOC(m, double)) ||
          !(x = NAG\_ALLOC(n, double)))
          printf("Allocation failure\n");
          exit_status = -1;
          goto END;
      if (order == Nag_ColMajor)
       pda = n + 3;
      else
       pda = m + 1;
      for (i = 1; i \leq m; ++i)
          scanf("%lf%lf%*[^\n] ", &t, &b[i-1]);
          A(1, i) = exp(t);

A(2, i) = exp(-t);
         A(3, i) = 1.0;
      tol = 0.0;
      relerr = 0.0;
      /* nag_linf_fit (e02gcc).
      * L_infinity-approximation by general linear function
      nag_linf_fit(order, m, n, a, b, tol, &relerr, x, &resmax, &irank, &iter,
                   &fail);
      if (fail.code != NE_NOERROR)
        {
          printf("Error from nag_linf_fit (e02gcc).\n%s\n",
                  fail.message);
          exit_status = 1;
          goto END;
        }
      else
        {
          printf("\n");
          printf("resmax = %11.2e Rank = %5ld Iterations ="
                  " %5ld\n", resmax, irank, iter);
```

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```
printf("\n");
    printf("Solution\n");

for (i = 1; i <= n; ++i)
        printf("%10.4f", x[i-1]);
    printf("\n");
}
END:
    NAG_FREE(a);
    NAG_FREE(b);
    NAG_FREE(x);
return exit_status;
}</pre>
```

10.2 Program Data

```
nag_linf_fit (e02gcc) Example Program Data
5
    0.0 4.501
    0.2 4.360
    0.4 4.333
    0.6 4.418
    0.8 4.625
```

10.3 Program Results

```
nag_linf_fit (e02gcc) Example Program Results
resmax = 1.03e-03 Rank = 3 Iterations = 4
Solution
    1.0049    2.0149    1.4822
```

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