

J. G. HERRIOT, Editor

ALGORITHM 314

FINDING A SOLUTION OF N FUNCTIONAL EQUATIONS IN N UNKNOWNS [C5]

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procedure ndinvt (functions, initstep, error, cycles, x, f, accest, n);
value n; procedure functions; real initstep, error;
integer cycles, n; array x, f, accest;

comment This procedure performs inverse interpolation in n dimensions, i.e., it will find a set of values for n variables x, such that n functions f(x) are zero. A more sophisticated technique, suitable for large values of n, has been developed by S. M. Robinson (Interpolative Solution of Systems of Nonlinear Equations, SIAM Journal of Numerical Analysis, 3 (1966), 650-658). It can also be used to fit a curve with n arbitrary parameters to a set of points, the n functions being formed, in this case, by equating to zero the differential of the sum of the squares of the residues with respect to each parameter in turn.

The functions required are specified by a procedure of the form functions (f, x) where f and x are declared as arrays from 1 to n. This procedure should calculate the n functions from a set of values given in x, placing the results in f. The first step is made by forming partial derivatives over an interval initstep. $1_{10} - 6$ should be suitable for values of x of the order 1 to 10. Exit from the procedure will occur if:

- (i) the root sum square of the x increments is less than error. If error is negative, this condition must be satisfied for | error |, and in addition this process is continued until the root sum square of the incrementsfails to decrease
- or (ii) the number of iterations is greater than cycles, implying that too much accuracy has been requested
- or (iii) the specified equations are singular. In this case exit is by a jump to a label *fails*.

On entry, the array x should contain the starting values. On exit, the array x will contain the accurate root, f the residues and accest the last increments made to x as a measure of the accuracy.

This procedure calls on a global procedure equations in (A, b, n, label), which solves n linear simultaneous equations in n unknowns Ax = b, placing the result in b. If A is singular, it is assumed that an exit is made by a jump to label;

begin

```
real work, sumsgres, prevres;
integer i, j, count;
Boolean switch;
array prevf[1:n], copydelf[1:n, 1:n], delx, delf[1:n, 1:n+1];
functions(prevf, x);
for i := 1 step 1 until n do
begin
    x[i] := x[i] + initstep;
functions (f, x);
for j := 1 step 1 until n do
begin
    delf[i, j] := f[j] - prevf[j];
```

```
delx[i, j] := 0;
    end differencing initial point;
    delx[i, i] := initstep;
    x[i] := x[i] - initstep;
  end setting up the initial matrix of points;
  sumsqres := 1_{10}30;
  count := 0;
iterate:
  switch := true;
  prevres := sumsqres;
tryagain:
  for i := 1 step 1 until n do
  begin
    f[i] := prevf[i];
    for j := 1 step 1 until n do copydelf[i, j] := delf[i, j]
  end copying delf for destructive use in procedure eqnsolve;
  eqnsolve (copydelf, f, n, inline);
  sumsqres := 0;
  for := 1 step 1 until n do
  begin
    work := 0;
    for j := 1 step 1 until n do work := work - delx[i, j] \times f[j];
    accest[i] := work;
    x[i] := x[i] + work;
    sumsqres := sumsqres + work \times work
  end calculation of next point;
  count := count + 1;
  functions (f, x);
  if count > cycles \lor sumsqres < error \lor error \land
     (error > 0 \lor sumsqres > prevres) then go to exit;
  for i := 1 step 1 until n do
  begin
    work := f[i] - prevf[i];
     prevf[i] := f[i];
     for j := n \text{ step } -1 \text{ until } 1 \text{ do}
       delx[i, j+1] := delx[i, j] - accest[i];
       delf[i, j+1] := delf[i, j] - work
     end calculation of new differences;
     delx[i, 1] := -accest[i];
     delf[i, 1] := -work
  end moving points up one place in tables;
  go to iterate;
inline:
  for i := 1 step 1 until n do
  begin
     \operatorname{delx}[i,\,n]\,:=\,\operatorname{delx}\,[i,\,n{+}1];
     delf[i, n] := delf[i, n+1]
  end discarding alternative point;
  switch := \neg switch;
  if switch then go to fails else go to tryagain;
 exit:
 end ndinvt
```

ALGORITHM 315

THE DAMPED TAYLOR'S SERIES METHOD FOR MINIMIZING A SUM OF SQUARES AND FOR SOLVING SYSTEMS OF NONLINEAR EQUATIONS [E4, C5]

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```
procedure TAYLOR (n, m, x, h, f, itmax, eps1, eps2, der, S, KENN,
   EXIT);
value n, m, eps1, eps2; integer n, m, itmax, KENN;
   real eps1, eps2, S;
```

Boolean der; array x, h, f; label EXIT; comment

Let

$$S(x_1,\dots,x_n) = \sum_{i=1}^m f_i^2(x_1,\dots,x_n) \qquad (m \ge n)$$
 (1)

the function to be minimized. Such functions always appear if you apply the method of least squares to estimate nonlinear parameters. The following sequence

$$x^{(k+1)} = x^{(k)} - \beta \Delta x^{(k)} = x^{(k)} - \beta (F'_{x(k)}^T F'_{x(k)})^{-1} F'_{x(k)}^T F(x^{(k)})$$

$$F = (f_1, \dots, f_m), \quad F'_x = \left(\frac{\partial f_i}{\partial x_j}\right) i = 1, \dots, m, j = 1, \dots, n$$
(2)

where β , which is always possible, is chosen to be such that

$$S(x^{(k)} - \beta \Delta x^{(k)}) \leqslant (1 - \beta \lambda) S(x^{(k)}) \qquad (0 < \lambda < 1) \tag{3}$$

is known to converge [1] for any $x^{(0)}$ to a stationary point of $S(grad S=2F_x^TF(x)=0)$, if on the carrying out of the iteration the matrix $F_x^TF_x'$ does not become singular.

For m = n you have $\Delta x = F'_x^{-1}F(x)$ and (2) becomes a damped version of Newton's method for solving the system of nonlinear equations

$$F(x) = 0 (4)$$

All zeros of (4) are stationary points of (1). Thus we are able to generate a sequence which converges for any $x^{(0)}$ to a stationary point of (1) and the possible divergence of Newton's method $(\beta=1)$ is avoided. It is not assured, however, that the method will always converge to a solution of (4). Numerical experience has shown that though Newton's method $(\beta=1)$ diverges for a certain $x^{(0)}$ the damped sequence converges to a solution of (4) for the same $x^{(0)}$.

In the program we have chosen $\lambda = .2$. At each iteration we set first $\beta = 1$ and then, if (3) is not valid, $\beta = 2^{-j}$ $(j=1,2,\ldots,16)$. If j is greater than 16 then $\beta < .00002$ and we assume to have reached a stationary point of S.

Meaning of the formal parameters:

n the number of variables x_i

m the number of functions f_i

the array x[1:n] which must first contain a starting value $x^{(0)}$ and finally will contain a stationary point of S, if $F'_x {}^T F'_x$ or for $m = n F'_x$, respectively, has not become singular

h h[1:n] is a step size vector for the approximation of F'_x (see below)

f the array f[1:m] will contain the function values at the last x calculated in TAYLOR

itmax must initially contain the maximum number of iterations to be performed. Leaving TAYLOR regularly, itmax contains the actual number of performed iterations

eps1 the iteration is stopped when S < eps1

eps2 the iteration is discontinued when $\sum_{i=1}^{n} |\Delta x_i^{(k)}| < eps2 \times \sum_{i=1}^{n} |x_i^{(k+1)}|$

der if $der = \mathbf{true}$ the matrix F'_x must be produced by a global procedure named DERIVE(x, dfdx) which adjoins to the vector x[1:n] the array dfdx[1:m, 1:n]. In this case the array h can be loaded by an arbitrary vector, for instance x.

if der = false the matrix F'_x is approximated by

$$\frac{\partial f_i}{\partial x_j} = \frac{f_i(x_1, \dots, x_j + h_j, \dots, x_n) - f_i(x_1, \dots, x_j - h_j, \dots, x_n)}{2h_j}$$

where h is a given step size vector. With a suitable choice of the h_j the convergence behavior of the sequence (2) is not destroyed. DERIVE(x, dfdx) must be formally declared outside of TAYLOR in this case.

[In some cases, particularly when solving nonlinear equations, the extra accuracy achieved by using central differences to estimate the derivatives is not necessary. A considerable saving in execution time can be obtained by using one-sided differences which means only minor changes in the program below.

—Ref.]

should initially contain the greatest positive number that the employed computer can store. Finally S contains $S = S(x^{(itmax)})$, if TAYLOR is regularly left.

KENN if after having called TAYLOR

KENN = 0 then one of the above interruptions applies (eps1, eps2),

KENN = 1 then itmax iterations were carried out and TAYLOR is left,

KENN = -1 then $\beta = 2^{-17}$ and TAYLOR is left.

EXIT TAYLOR goes to this global label if i encounters a singular matrix.

Further two global procedures must be made available to TAYLOR:

 i) FUNCTION(x, f) which is able to calculate for a given vector x[1:n] the function values f[1:m]

ii) GAUSS(n, A, b, x, EXIT) which solves the linear system of n equations Ax = b for x. If A is singular then GAUSS returns to the global label EXIT. Any linear equation solver may be used for GAUSS;

```
begin integer i, j, k, z, l; real hf, hl, hs, hz;
  array fp, fm[1:m], b, dx[1:n], dfdx[1:m, 1:n], aa[1:n, 1:n];
  hs := S; KENN := z := 0;
ITERATION: z := z + 1;
  if z > itmax then begin KENN := 1; go to ENDE end;
    l := 0; hl := 1.0;
DAMP: l := l + 1;
  if l > 16 then begin KENN := -1; go to ENDE end;
  FUNCTION(x, f); hf := 0;
  for i := 1 step 1 until m do hf := hf + f[i] \times f[i];
  if hf > hs \times (1.0 - .2 \times hl) then
  begin hl := hl \times .5;
    for k := 1 step 1 until n do x[k] := x[k] + hl \times dx[k];
    go to DAMP
  end;
  hs := hf; if hs < eps 1 then go to ENDE;
  if der then DERIVE(x, dfdx) else
    for i := 1 step 1 until n do
    begin hf := h[i]; hz := 2.0 \times hf;
      x[i] := x[i] + hf; FUNCTION(x, fp);
      x[i] := x[i] - hz; FUNCTION(x, fm);

x[i] := x[i] + hf; hz := 1.0/hz;
      for k := 1 step 1 until m do
      dfdx[k, i] := hz \times (fp[k] - fm[k])
    end
  end;
if m = n then GAUSS(n, dfdx, f, dx, EXIT) else
  for i := 1 step 1 until n do
  begin hf := 0;
   for k := 1 step 1 until m do
      hf := hf + dfdx[k, i] \times f[k]; \quad b[i] := hf;
    for k := i step 1 until n do
   begin hf := 0;
      for j := 1 step 1 until m do
        hf := hf + dfdx[j, i] \times dfdx[j, k];
      aa[i, k] := aa[k, i] := hf
   end
 end:
 GAUSS(n, aa, b, dx, EXIT)
```

```
\begin{array}{l} hz := hf := 0; \\ \textbf{for } i := 1 \ \textbf{step 1 until } n \ \textbf{do} \\ \textbf{begin} \\ x[i] := x[i] - dx[i]; \quad hz := hz + abs(x[i]); \\ hf := hf + abs(dx[i]) \\ \textbf{end}; \\ \textbf{if } hf \geq eps2 \times hz \ \textbf{then go to } ITERATION; \\ ENDE: FUNCTION(x, f); \quad S := 0; \quad itmax := z; \\ \textbf{for } i := 1 \ \textbf{step 1 until } m \ \textbf{do } S := S + f[i] \times f[i] \\ \textbf{end } TAYLOR \end{array}
```

Reference:

[1] Braess, D. Über Dämpfung bei Minimalisierungsverfahren. Computing 1 (1966), 264-272.

ALGORITHM 316

SOLUTION OF SIMULTANEOUS NON-LINEAR EQUATIONS [C5]

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procedure nonlinear system (n, maxit, numsig, singular, x); value n, numsig; integer n, maxit, numsig, singular; array x; **comment** This procedure solves a system of *n* simultaneous nonlinear equations. The method is roughly quadratically convergent and requires only $((n^2/2)+(3n/2))$ function evaluations per iterative step as compared with (n^2+n) evaluations for Newton's Method. This results in a savings of computational effort for sufficiently complicated functions. A detailed description of the general method and proof of convergence are included in [1]. Basically the technique consists in expanding the first equation in a Taylor series about the starting guess, retaining only linear terms, equating to zero and solving for one variable, say x_k , as a linear combination of the remaining n-1 variables. In the second equation, x_k is eliminated by replacing it with its linear representation found above, and again the process of expanding through linear terms, equating to zero and solving for one variable in terms of the now remaining n - 2 variables is performed. One continues in this fashion,eliminating one variable per equation, until for the nth equation, we are left with one equation in one unknown. A single Newton step is now performed, followed by back-substitution in the triangularized linear system generated for the x_i 's. A pivoting effect is achieved by choosing for elimination at any step that variable having a partial derivative of largest absolute value. The pivoting is done without physical interchange of rows or columns.

The vector of initial guesses x, the number of significant digits desired numsig, the maximum number of iterations to be used, maxit, and the number of equations n, should be set up prior to the procedure call which activates nonlinear system. After execution of the procedure, the vector x is the solution of the system (or best approximation thereto), maxit is now the number of iterations used and singular = 0 is an indication that a Jacobian-related matrix was singular—indicative of the process "blowing-up," whereas singular = 1 is an indication that no such difficulty occurred. Storage space may be saved by implementing the algorithm in a way which takes advantage of the fact that the strict lower triangle of the array pointer and the same number of positions in the array coe are not used;

```
same number of positions in the array coe are not used;

begin integer converge, m, j, k, i, jsub, itemp, kmax, kplus, tally;

real f, hold, h, fplus, dermax, test, factor, relconvg;

integer array pointer[1:n, 1:n], isub[1:n-1];

array temp, part[1:n], coe[1:n, 1:n+1];

procedure backsubstitution (k, n, x, isub, coe, pointer);

value k, n;

integer k, n; integer array isub, pointer; array x, coe;
```

```
comment This procedure back-solves a triangular linear
    system for improved x[i] values in terms of old ones;
  begin integer km, kmax, jsub;
    for km := k \text{ step } -1 \text{ until } 2 \text{ do}
    begin kmax := isub[km-1]; x[kmax] := 0;
      for j := km step 1 until n do
      begin jsub := pointer[km, j];
        x[kmax] := x[kmax] + coe[km-1, jsub] \times x[jsub]
      x[kmax] := x[kmax] + coe[km-1, n+1]
    end;
  end backsubstitution;
  procedure evaluatekthfunction (x, y, k);
    integer k; real y; array x;
 begin comment the body of this procedure must be provided
    by the user. One call of the procedure should cause the value
    of the kth function at the current value of the vector x to be
    placed in y;
  end evaluatekthfunction;
  converge := 1; \quad singular := 1; \quad relconvg := 10 \uparrow (-numsig);
 for m := 1 step 1 until maxit do
 begin
   comment An intermediate output statement may be in-
      serted at this point in the procedure to print the successive
      approximation vectors x generated by each complete itera-
      tive step;
    for j := 1 step 1 until n do pointer [1, j] := j;
    for k := 1 step 1 until n do
    begin if k > 1 then backsubstitution (k, n, x, isub, coe, pointer);
      evaluatekthfunction (x, f, k); factor := .001;
AAA:
            tally := 0; for i := k step 1 until n do
      begin itemp := pointer[k, i]; hold := x[itemp];
        h := factor \times hold; if h = 0 then h := .001;
        x[itemp] := hold + h;
        if k > 1 then backsubstitution (k, n, x, isub, coe, pointer);
        evaluatekth function (x, fplus, k);
        part[itemp] := (fplus-f)/h;
        x[itemp] := hold; if (abs(part[itemp]) = 0) \lor
        (abs(f/part[itemp]) > 1.0_{10}20) then tally := tally + 1;
      end;
      if tally \leq n - k then go to AA; factor := factor \times 10.0;
      if factor > .5 then go to SING; go to AAA;
AA: if k < n then go to A; if abs (part[itemp]) = 0
      then go to SING;
      coe[k, n+1] := 0; kmax := itemp; go to ENDK;
      kmax := pointer[k, k]; dermax := abs(part[kmax]);
      kplus := k + 1;
      for i := kplus step 1 until n do
      \mathbf{begin}\ jsub\ :=\ pointer[k,\ i];\ \ test\ :=\ abs(part[jsub]);
        if test < dermax then go to B; dermax := test;
        pointer[kplus, i] := kmax; kmax := jsub;
        go to ENDI;
B:
        pointer [kplus, i] := jsub;
ENDI:
      if abs(part[kmax]) = 0 then go to SING; isub[k] := kmax;
      coe[k, n+1] := 0;
      for j := kplus step 1 until n do
      begin jsub := pointer[kplus, j];
        coe[k, jsub] := -part[jsub]/part[kmax];
        coe[k, n+1] := coe[k, n+1] + part[jsub] \times x[jsub]
      end:
ENDK:
      coe[k, n+1] := (coe[k, n+1]-f)/part[kmax] + x[kmax]
    end k:
    x[kmax] := coe[n, n+1];
    if n > 1 then backsubstitution (n, n, x, isub, coe, pointer);
    if m = 1 then go to D;
    for i := 1 step 1 until n do
      if abs((temp[i]-x[i])/x[i]) > relconvg then go to C;
```

```
converge := converge + 1;
    if converge ≥ 3 then go to TERMINATE else go to D;
C: converge := 1;
D: for i := 1 step 1 until n do temp[i] := x[i]
    end m;
    go to THROUGH;
TERMINATE:
    maxit := m; go to THROUGH;
SING:
    singular := 0;
THROUGH:
    end nonlinear system
```

APPENDIX

We include a sample procedure evaluatekth function for the 2×2 system:

$$\left(1 - \frac{1}{4\pi}\right)(e^{2x_1} - e) + \frac{e}{\pi}x_2 - 2ex_1 = 0$$

$$\frac{1}{2}\sin(x_1x_2) - \frac{x_2}{4\pi} - \frac{x_1}{2} = 0,$$

one solution of which is $(.5, \pi)$ see [2]

procedure evaluate kthfunction(x, y, k);

integer k; real y; array x; begin switch functionnumber := F1, F2;

go to function number [k];

F1: $y := 2.71828183 \times (.920422528 \times (exp(2 \times x[1]-1)-1) + x[2]/3.14159265-2 \times x[1]);$

go to RETURN;

F2: $y := .5 \times sin(x[1] \times x[2]) - x[2]/12.5663706 - x[1]/2;$ RETURN:

end evaluatekthfunction;

References:

- Brown, K. M. A quadratically convergent method for solving simultaneous non-linear equations. Doctoral Thesis, Dept. Computer Sciences, Purdue U., Lafayette, Ind., Aug., 1966.
- Brown, K. M., and Conte, S. D. The solution of simultaneous nonlinear equations. Proc. ACM 22nd Nat. Conf., pp 111-114.

ALGORITHM 317* PERMUTATION [G6]

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procedure permute(n, k, v); value n, k; integer array v; integer n, k;

comment This procedure produces in the vector v the kth permutation on n variables. When k=0, v takes on the value $1, 2, 3, 4, \cdots, n$. This algorithm is not as efficient as previously published algorithms [1], [2], [3] for generating a complete set of permutations but it is significantly better for generating a random permutation, a property useful in certain simulation applications. Any non-negative value of k will produce a valid permutation. To generate a random permutation, k should be chosen from the uniform distribution over the integers from 0 to n! - 1 inclusive;

```
begin integer i, q, r, x, j;

for i := 1 step 1 until n do v[i] := 0;

for i := n step -1 until 1 do

begin

q := k \div i; r := k - q \times i; x := 0; j := n;
```

```
no: 	ext{ if } v[j] = 0 	ext{ then } 
	ext{begin}
	ext{ if } x = r 	ext{ then go to } it 	ext{ else } x := x + 1 
	ext{ end;}
	ext{ } j := j - 1; 	ext{ go to } no;
it: 	ext{ } v[j] := i; 	ext{ } k := q;
	ext{ end}
	ext{ end}
```

References:

- COVEYOU, R. R., AND SULLIVAN, J. G. Algorithm 71, Permutation. Comm. ACM 4 (Nov. 1961), 497.
- PECK, J. E. L., AND SCHRACK, G. F. Algorithm 86, Permute. Comm. ACM 5 (Apr. 1962), 208.
- TROTTER, H. F. Algorithm 115, Perm. Comm. ACM 5 (Aug. 1962), 434.

Algorithms Policy • Revised August, 1966

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For ALGOL 60 programs, input and output should be achieved by procedure statements, using any of the following eleven procedures (whose body is not specified in ALGOL) [See "Report on Input-Output Procedures for ALGOL 60," Comm. ACM 7 (Oct. 1964), 628-629]:

insymbol inreal outarray ininteger outsymbol outreal outboolean outinteger length inarray outstring

If only one channel is used by the program for output, it should be designated by 1 and similarly a single input channel should be designated by 2. Examples:

```
outstring (1, 'x='); outreal (1,x); for i := 1 step 1 until n do outreal (1,A[i]); ininteger (2, digit [17]):
```

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