

Algorithms

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ALGORITHM 345 AN ALGOL CONVOLUTION PROCEDURE BASED ON THE FAST FOURIER TRANSFORM [C6]

RICHARD C. SINGLETON* (Recd. 30 Dec. 1966, 26 July
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Stanford Research Institute, Menlo Park, CA 94025

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Stockham [6] and Gentleman and Sande [3] have shown the practical advantages of computing the circular convolution

$$C_k = \sum_{j=0}^{n-1} A_j B_{(j+k) \bmod n}, \quad k = 0, 1, \dots, n-1,$$

of two real vectors A and B of period n by the fast Fourier transform [2, 3, 4]. The Fourier transforms

$$\alpha_j = \sum_{p=0}^{n-1} A_p \exp(i2\pi pj/n)$$

and

$$\beta_j = \sum_{q=0}^{n-1} B_q \exp(i2\pi qj/n)$$

are first computed, then the convolution

$$C_k = \frac{1}{n} \sum_{j=0}^{n-1} \alpha_j \beta_j^* \exp(i2\pi jk/n)$$

where β_j^* is the complex conjugate of β_j . By this method the number of arithmetic operations increases by a factor slightly more than 2 when n is doubled, as compared with a factor of 4 for the direct method. Tests show a 16 to 1 time advantage for the transform method at $n = 256$.

The operation of convolution is used in computing autocorrelation and cross-correlation functions, in digital filtering of time series, and many other applications.

Procedure *CONVOLUTION* computes the convolution of two real vectors of dimension $n = 2^m$. The special features of this procedure are: (1) the usual reordering of the fast Fourier transform results is avoided, and (2) the return from frequency to time is made with a transform of dimension $n/2$ instead of n . The two vectors A and B are first transformed with a single complex Fourier transform of dimension n . The complex product $\alpha\beta^*$ is then formed, leaving the result in reverse binary order. Since the

convolution is real-valued, the real part x of the complex product is an even function and the imaginary part y is an odd function; thus the Fourier transform of x is real and that of y is imaginary. These properties lead to the identity

$$\begin{aligned} T(x + iy) &= \text{Re}(Tx) - \text{Im}(Ty) \\ &= \text{Re}(T(x - y)) + \text{Im}(T(x - y)) \end{aligned}$$

where T represents the Fourier transform and $T(x + iy)$ is the desired convolution. We subtract y from x , yielding a real vector of dimension n , then transform using a complex transform of dimension $n/2$ and add the resulting cosine and sine coefficients to give the convolution. Thus with procedure *CONVOLUTION* we make maximum use of the complex Fourier transform in each direction and avoid any reverse binary to binary permutation. The Fourier transform

$$T(A + iB) = \alpha + i\beta$$

of the two original vectors is available in reverse binary order on exit from the procedure. We can permute this transform to normal order with procedure *REVERSEBINARY* and readily compute the power spectra and cross spectrum of the two data vectors.

Procedure *CONVOLUTION* uses procedure *REALTRAN*, given in Algorithm 338 [5], but repeated here with revisions to improve accuracy on computers using truncated floating-point arithmetic. Procedures *FFT4* and *REVFFT4* are also used and perform the same computation as procedures *FFT2* and *REVFFT2* given in Algorithm 338 for use on a system with virtual memory. The transform procedures given here are organized without regard to the problem of memory overlay. This change yields a 10 percent reduction in computing time on the Burroughs B5500 for transforms of dimension $n = 512$ or smaller. Procedure *FFT4* is based on an organization of the fast Fourier transform due to Sande [3], and procedure *REVFFT4* is similar to the method proposed by Cooley and Tukey [2], except that the data is in reverse binary order. In both cases, trigonometric functions are used in normal sequence, rather than reverse binary sequence, thus eliminating the need for a reverse binary counter. Another gain in efficiency comes from reducing the time for computing trigonometric function values. The following difference-equation method is used:

$$\cos((k+1)\theta) = \cos(k\theta) - (C \times \cos(k\theta) + S \times \sin(k\theta))$$

and

$$\sin((k+1)\theta) = \sin(k\theta) + (S \times \cos(k\theta) - C \times \sin(k\theta)),$$

where the constant multipliers are $C = 2 \sin^2(\theta/2)$ and $S = \sin(\theta)$, and the initial values are $\cos(0) = 1$ and $\sin(0) = 0$.

These initial values should be computed to full machine precision; if necessary, a stored table of $\sin(\theta)$ for $\theta = \pi/2, \pi/4, \pi/8, \dots, \pi/n$ can be added to procedures *FFT4* and *REVFFT4*. Using the standard sine function to compute initial values, the ratio of rms error to rms data is about 2×10^{-11} for the transform-inverse pair at $n = 512$ on the Burroughs B5500 computer; this error is about the same as that obtained when the sine and cosine functions

are used for all trigonometric function values. On a computer using truncated, rather than rounded, arithmetic operations, the sequence of values for $\cos(k\theta) + i \sin(k\theta)$ tends to spiral inward from the unit circle. Since the error is primarily one of magnitude, rather than angle, rescaling to the unit circle at each step gives a satisfactory correction. This correction is included in procedures *FFT4* and *REVFFT4* but may be removed to improve running speed if rounded arithmetic is used.

Procedures *FFT8* and *REVFFT8* are included as possible substitutes for *FFT4* and *REVFFT4*. These procedures use radix 8 arithmetic [1], rather than radix 4, and run about 20 percent faster on the Burroughs B5500 computer; however, the compiled code is twice as long. The code could be shortened by use of subscripted variables and FOR statements, but this change would probably eliminate most of the time-saving.

The permutation procedure *REVERSEBINARY* is based on a modified dual counter, one in normal sequence and the other in reverse binary sequence. In permuting a vector of dimension n , the normal sequence counter goes from 1 to $n/2 - 1$, and the elements indexed 1, 3, \dots , $n/2 - 1$ are exchanged with their reverse-binary counterparts (indexed greater than or equal to $n/2$) without need of a test. The reverse binary counter is incremented only $n/4$ times, and exchanges of pairs of elements below $n/2$ are done jointly with pair exchanges in the upper half of the array; i.e. if x_j and x_k are exchanged, where $j, k < n/2$, then x_{n-1-j} and x_{n-1-k} are also exchanged. This procedure is twice as fast on the Burroughs B5500 as *REORDER* given in Algorithm 338 [5] and is the better choice when the additional features of *REORDER* are not needed. For a single-variate, complex Fourier transform of dimension $n = 2^m$,

REVERSEBINARY(A, B, m);

REVFFT8($A, B, n, m, 1$)

was found to be the best combination for $n \leq 512$ on the B5500 computer, giving a time of 0.79 sec. for $n = 512$.

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procedure *CONVOLUTION* ($A, B, C, D, m, scale$);
value $m, scale$; **integer** m ; **real** $scale$; **array** A, B, C, D ;
comment This procedure computes the circular convolution

$$C_k = scale \sum_{j=0}^{n-1} A_j B_{(j+k) \bmod n}, \quad k = 0, 1, \dots, n-1,$$

where $n = 2^m$ and $p \bmod n$ represents the remainder after division of p by n . (It is assumed that $m \geq 1$.) Arrays $A, B[0 : n-1]$ originally contain the two data vectors to be convoluted, and on exit, contain the Fourier transform of $A + iB$ arranged in reverse binary order. A and B must not be the same array. On exit, array $C[0 : n-1]$ contains the convolution multiplied by the factor $scale$. Array D is a scratch storage array with lower bound zero and upper bound at least $n \div 2$. If the Fourier

transform of the data is not needed, the procedure can be called with arrays A and B used for C and D in either order, for example, *CONVOLUTION* ($A, B, A, B, m, scale$). If the Fourier transform is used, it should first be permuted to normal order by the call *REVERSEBINARY*(A, B, m). After doing this, the Fourier cosine coefficients of the A vector are

$$(A_k + A_{n-k})/n, \quad k = 1, 2, \dots, n/2,$$

$$(2A_0)/n, \quad k = 0,$$

and the sine coefficients are

$$(B_k - B_{n-k})/n, \quad k = 1, 2, \dots, n/2 - 1.$$

The Fourier cosine coefficients of the B vector are

$$(B_k + B_{n-k})/n, \quad k = 1, 2, \dots, n/2,$$

$$(2B_0)/n, \quad k = 0,$$

and the sine coefficients are

$$(A_{n-k} - A_k)/n, \quad k = 1, 2, \dots, n/2 - 1.$$

The procedures *FFT4*, *REVFFT4*, and *REALTRAN* are used by this procedure and must also be declared. If convolutions of large dimension are to be computed on a system with virtual memory, procedures *FFT2* and *REVFFT2* (Algorithm 338) [5] should be substituted for procedures *FFT4* and *REVFFT4*;

begin integer j, kk, ks, n ; **real** aa, ab, ba, bb, im ;

$n := 2 \uparrow m$; $j := 1$;

FFT4(A, B, n, m, n);

$C[0] := 4 \times (A[0] \times B[0])$;

L: $kk := j$; $ks := j := j + j$;

L2: $ks := ks - 1$;

$aa := A[kk] + A[ks]$; $ab := A[kk] - A[ks]$;

$ba := B[kk] + B[ks]$; $bb := B[kk] - B[ks]$;

$im := ba \times bb + aa \times ab$; $aa := aa \times ba - ab \times bb$;

$C[kk] := aa - im$; $C[ks] := aa + im$;

$kk := kk + 1$; **if** $kk < ks$ **then go to** **L2**;

if $j < n$ **then go to** **L**;

$kk := n \div 2$; $ks := kk - 1$; $scale := scale / (8 \times n)$;

for $j := 0$ **step** 1 **until** ks **do** $D[j] := C[j + kk]$;

REVFFT4($C, D, kk, m-1, 1$);

REALTRAN(C, D, kk, false);

$C[0] := scale \times C[0]$; $C[kk] := scale \times C[kk]$;

for $j := 1$ **step** 1 **until** ks **do**

begin $C[n-j] := scale \times (C[j] - D[j])$;

end

$C[j] := scale \times (C[j] + D[j])$

end *CONVOLUTION*;

procedure *FFT4*(A, B, n, m, ks); **value** n, m, ks ;

integer n, m, ks ; **array** A, B ;

comment This procedure computes the fast Fourier transform for one variable of dimension 2^m in a multivariate transform. n is the number of data points, i.e. $n = n_1 \times n_2 \times \dots \times n_p$ for a p -variate transform, and $ks = n_k \times n_{k+1} \times \dots \times n_p$, where $n_k = 2^m$ is the dimension of the current variable. Arrays $A[0 : n-1]$ and $B[0 : n-1]$ originally contain the real and imaginary components of the data in normal order. Multivariate data is stored according to the usual convention, e.g. a_{jkl} is in $A[j \times n_2 \times n_3 + k \times n_3 + l]$ for $j = 0, 1, \dots, n_1 - 1$, $k = 0, 1, \dots, n_2 - 1$, and $l = 0, 1, \dots, n_3 - 1$. On exit, the Fourier coefficients for the current variable are in reverse binary order. Continuing the above example, if the "column" variable n_2 is the current one, column

$$k = k_{m-1}2^{m-1} + k_{m-2}2^{m-2} + \dots + k_12 + k_0$$

is permuted to position

$$k_02^{m-1} + k_12^{m-2} + \dots + k_{m-2}2 + k_{m-1}.$$

A separate procedure may be used to permute the results to normal order between transform steps or all at once at the end. If $n = ks = 2^m$, the single-variate transform

$$(x_j + iy_j) = \sum_{k=0}^{n-1} (a_k + ib_k) \exp(i2\pi jk/n)$$

for $j = 0, 1, \dots, n-1$ is computed, where $(a+ib)$ represent the initial values and $(x+iy)$ represent the transformed values;

```

begin integer k0, k1, k2, k3, k, span;
real A0, A1, A2, A3, B0, B1, B2, B3, re, im;
real rad, dc, ds, c1, c2, c3, s1, s2, s3;
span := ks; ks := 2 ↑ m; rad := 4.0 × arctan(1.0)/ks;
ks := span ÷ ks; n := n - 1; k := m;
for m := m - 2 while m ≥ 0 do
begin
  c1 := 1.0; s1 := 0; k0 := 0; k := ks;
  dc := 2.0 × sin(rad) ↑ 2; rad := rad + rad;
  ds := sin(rad); rad := rad + rad;
  span := span ÷ 4;
La: k1 := k0 + span; k2 := k1 + span; k3 := k2 + span;
A0 := A[k0]; B0 := B[k0];
A1 := A[k1]; B1 := B[k1];
A2 := A[k2]; B2 := B[k2];
A3 := A[k3]; B3 := B[k3];
A[k0] := A0 + A2 + A1 + A3;
B[k0] := B0 + B2 + B1 + B3;
if s1 = 0 then
begin
  A[k1] := A0 + A2 - A1 - A3;
  B[k1] := B0 + B2 - B1 - B3;
  A[k2] := A0 - A2 - B1 + B3;
  B[k2] := B0 - B2 + A1 - A3;
  A[k3] := A0 - A2 + B1 - B3;
  B[k3] := B0 - B2 - A1 + A3
end
else
begin
  re := A0 + A2 - A1 - A3; im := B0 + B2 - B1 - B3;
  A[k1] := re × c2 - im × s2;
  B[k1] := re × s2 + im × c2;
  re := A0 - A2 - B1 + B3; im := B0 - B2 + A1 - A3;
  A[k2] := re × c1 - im × s1;
  B[k2] := re × s1 + im × c1;
  re := A0 - A2 + B1 - B3; im := B0 - B2 - A1 + A3;
  A[k3] := re × c3 - im × s3;
  B[k3] := re × s3 + im × c3
end;
k0 := k3 + span; if k0 < n then go to La;
k0 := k0 - n; if k0 ≠ k then go to La;
comment If computing for the current factor of 4 is not
  finished then increment the sine and cosine values;
if k0 ≠ span then
begin
  c2 := c1 - (dc×c1+ds×s1);
  s1 := (ds×c1-dc×s1) + s1;
comment The following three statements compensate
  for truncation error. If rounded arithmetic is used, sub-
  stitute c1 := c2;
  c1 := 1.5-0.5 × (c2 ↑ 2+s1 ↑ 2);
  s1 := c1 × s1; c1 := c1 × c2;
  c2 := c1 ↑ 2 - s1 ↑ 2; s2 := 2.0 × c1 × s1;
  c3 := c2 × c1 - s2 × s1; s3 := c2 × s1 + s2 × c1;
  k := k + ks; go to La
end;
k := m
end;
comment If m is odd then compute for one factor of 2;

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if k ≠ 0 **then**

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begin
  span := span ÷ 2; k0 := 0;
Lb: k2 := k0 + span; A0 := A[k2]; B0 := B[k2];
  A[k2] := A[k0] - A0; A[k0] := A[k0] + A0;
  B[k2] := B[k0] - B0; B[k0] := B[k0] + B0;
  k0 := k2 + span; if k0 < n then go to Lb;
  k0 := k0 - n; if k0 ≠ span then go to Lb
end
end FFT4;
procedure REVFFT4(A, B, n, m, ks); value n, m, ks;
integer n, m, ks; array A, B;
comment This procedure computes the fast Fourier transform
  for one variable of dimension 2m in a multivariate transform.
  n is the number of data points, i.e. n = n1 × n2 × ... × np
  for a p-variate transform, and ks = nk+1 × nk+2 × ... × np,
  where nk = 2m is the dimension of the current variable. Arrays
  A[0:n-1] and B[0:n-1] originally contain the real and imagi-
  nary components of the data with the indices of each variable in
  reverse binary order, e.g. ajkl is in A[j'×n2×n3+k'×n3+l']
  for j = 0, 1, ..., n1-1, k = 0, 1, ..., n2-1, and l = 0,
  1, ..., n3-1, where j', k', and l' are the bit-reversed values of
  j, k, and l. On completion of the multivariate transform, the
  real and imaginary components of the resulting Fourier coeffi-
  cients are in A and B in normal order. If n = 2m and ks = 1,
  a single-variate transform is computed;
begin integer k0, k1, k2, k3, k, span;
real A0, A1, A2, A3, B0, B1, B2, B3;
real rad, dc, ds, c1, c2, c3, s1, s2, s3;
rad := 4.0 × arctan(1.0); n := n - 1;
k0 := 0; span := ks;
comment If m is odd then compute for one factor of 2;
if (m ÷ 2) × 2 ≠ m then
begin
La: k2 := k0 + span; A0 := A[k2]; B0 := B[k2];
  A[k2] := A[k0] - A0; A[k0] := A[k0] + A0;
  B[k2] := B[k0] - B0; B[k0] := B[k0] + B0;
  k0 := k2 + span; if k0 < n then go to La;
  k0 := k0 - n; if k0 ≠ span then go to La;
  span := span + span; rad := 0.5 × rad
end;
for m := m - 2 while m ≥ 0 do
begin
  c1 := 1.0; s1 := 0; k0 := 0; rad := 0.25 × rad;
  dc := 2.0 × sin(rad) ↑ 2;
  ds := sin(rad+rad); k := ks;
Lb: k1 := k0 + span; k2 := k1 + span; k3 := k2 + span;
  A0 := A[k0]; B0 := B[k0];
  if s1 = 0 then
begin
    A2 := A[k1]; B2 := B[k1];
    A1 := A[k2]; B1 := B[k2];
    A3 := A[k3]; B3 := B[k3]
end
else
begin
    A2 := A[k1] × c2 - B[k1] × s2;
    B2 := A[k1] × s2 + B[k1] × c2;
    A1 := A[k2] × c1 - B[k2] × s1;
    B1 := A[k2] × s1 + B[k2] × c1;
    A3 := A[k3] × c3 - B[k3] × s3;
    B3 := A[k3] × s3 + B[k3] × c3
end;
  A[k0] := A0 + A2 + A1 + A3;
  B[k0] := B0 + B2 + B1 + B3;
  A[k1] := A0 - A2 - B1 + B3;
  B[k1] := B0 - B2 + A1 - A3;

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A[k2] := A0 + A2 - A1 - A3;
B[k2] := B0 + B2 - B1 - B3;
A[k3] := A0 - A2 + B1 - B3;
B[k3] := B0 - B2 - A1 + A3;
k0 := k3 + span; if k0 < n then go to Lb;
k0 := k0 - n; if k0 ≠ k then go to Lb;
comment If computing for the current factor of 4 is not
finished then increment the sine and cosine values;
if k0 ≠ span then
begin
c2 := c1 - (dc×c1+ds×s1);
s1 := (ds×c1-dc×s1) + s1;
comment The following three statements compensate
for truncation error. If rounded arithmetic is used, sub-
stitute c1 := c2;
c1 := 1.5-0.5 × (c2↑2+s1↑2);
s1 := c1 × s1; c1 := c1 × c2;
c2 := c1↑2 - s1↑2; s2 := 2.0 × c1 × s1;
c3 := c2 × c1 - s2 × s1; s3 := c2 × s1 + s2 × c1;
k := k + ks; go to Lb
end;
span := 4 × span
end
end REVFFFT4;
procedure REALTRAN(A, B, n, evaluate);
value n, evaluate; integer n;
Boolean evaluate; array A, B;
comment If evaluate is false, this procedure unscrambles the
single-variate complex transform of the n even-numbered and
n odd-numbered elements of a real sequence of length 2n, where
the even-numbered elements were originally in A and the odd-
numbered elements in B. Then it combines the two real trans-
forms to give the Fourier cosine coefficients A[0], A[1], ...,
A[n] and sine coefficients B[0], B[1], ..., B[n] for the full
sequence of 2n elements. If evaluate is true, the process is
reversed, and a set of Fourier cosine and sine coefficients is
made ready for evaluation of the corresponding Fourier series
by means of the inverse complex transform. Going in either
direction, REALTRAN scales by a factor of two, which should
be taken into account in determining the appropriate overall
scaling;
begin integer k, nk, nh;
real aa, ab, ba, bb, re, im, ck, sk, dc, ds;
nh := n ÷ 2; ds := 2.0 × arctan(1.0)/n;
dc := 2.0 × sin(ds)↑2; ds := sin(ds+ds);
sk := 0;
if evaluate then
begin ck := -1.0; ds := -ds end
else begin ck := 1.0; A[n] := A[0]; B[n] := B[0] end;
for k := 0 step 1 until nh do
begin
nk := n - k;
aa := A[k] + A[nk]; ab := A[k] - A[nk];
ba := B[k] + B[nk]; bb := B[k] - B[nk];
re := ck × ba + sk × ab; im := sk × ba - ck × ab;
B[nk] := im - bb; B[k] := im + bb;
A[nk] := aa - re; A[k] := aa + re;
aa := ck - (dc×ck+ds×sk);
sk := (ds×ck-dc×sk) + sk;
comment The following three statements compensate for
truncation error. If rounded arithmetic is used, substitute
ck := aa;
ck := 1.5-0.5 × (aa↑2+sk↑2);
sk := ck × sk; ck := ck × aa
end
end REALTRAN;
procedure REVERSEBINARY(A, B, m); value m;
integer m; array A, B;

```

comment This procedure permutes the elements A[j] and B[j] of arrays A and B, for j = 0, 1, ..., 2↑m - 1, according to the reverse binary transformation. Element

$$k = k_{m-1}2^{m-1} + k_{m-2}2^{m-2} + \dots + k_12 + k_0$$

is moved to location

$$k_02^{m-1} + k_12^{m-2} + \dots + k_{m-2}2 + k_{m-1}.$$

Two successive calls of this procedure give an identity transformation;

begin integer j, jj, k, lim, jk, n2, n4, n8, nn;

real t;

integer array C[0:m];

C[0] := nn := 1; jj := 0;

for j := 1 **step** 1 **until** m **do** C[j] := nn := nn + nn;

if m > 1 **then** n4 := C[m-2]; **if** m > 2 **then** n8 := C[m-3];

n2 := C[m-1]; lim := n2 - 1; nn := nn - 1; m := m - 4;

for j := 1 **step** 1 **until** lim **do**

begin

jk := jj + n2;

t := A[j]; A[j] := A[jk]; A[jk] := t;

t := B[j]; B[j] := B[jk]; B[jk] := t;

j := j + 1;

if jj ≥ n4 **then**

begin

jj := jj - n4;

if jj ≥ n8 **then**

begin

jj := jj - n8; k := m;

L: **if** C[k] ≤ jj **then**

begin jj := jj - C[k]; k := k - 1; **go to** L **end**;

jj := C[k] + jj

end

else jj := jj + n8

end

else jj := jj + n4;

if jj > j **then**

begin

k := nn - j; jk := nn - jj;

t := A[j]; A[j] := A[jj]; A[jj] := t;

t := B[j]; B[j] := B[jj]; B[jj] := t;

t := A[k]; A[k] := A[jk]; A[jk] := t;

t := B[k]; B[k] := B[jk]; B[jk] := t

end

end

end REVERSEBINARY;

procedure FFT8(A, B, n, m, ks); value n, m, ks;

integer n, m, ks; **array** A, B;

comment This procedure computes the fast Fourier transform for one variable of dimension 2^m in a multivariate transform. n is the number of data points, i.e. n = n₁ × n₂ × ... × n_p for a p-variate transform, ks = n_k × n_{k+1} × ... × n_p, where n_k = 2^m is the dimension of the current variable. Arrays A[0:n-1] and B[0:n-1] originally contain the real and imaginary components of the data in normal order. Multivariate data is stored according to the usual convention, e.g. a_{jkl} is in A[j×n₂×n₃+k×n₃+l] for j = 0, 1, ..., n₁ - 1, k = 0, 1, ..., n₂ - 1, and l = 0, 1, ..., n₃ - 1. On exit, the Fourier coefficients for the current variable are in reverse binary order. Continuing the above example, if the "column" variable n₂ is the current one, column

$$k = k_{m-1}2^{m-1} + k_{m-2}2^{m-2} + \dots + k_12 + k_0$$

is permuted to position

$$k_02^{m-1} + k_12^{m-2} + \dots + k_{m-2}2 + k_{m-1}.$$

A separate procedure may be used to permute the results to

normal order between transform steps or all at once at the end.
If $n = ks = 2^m$, the single variate transform

$$(x_j + iy_j) = \sum_{k=0}^{n-1} (a_k + ib_k) \exp(i2\pi jk/n)$$

for $j = 0, 1, \dots, n-1$ is computed, where $(a+ib)$ represent the initial values and $(x+iy)$ represent the transformed values;
begin integer $k0, k1, k2, k3, k4, k5, k6, k7, k, span$;
real $A0, A1, A2, A3, A4, A5, A6, A7, B0, B1, B2, B3, B4, B5, B6, B7, x0, x1, x2, x3, x4, x5, x6, x7, y0, y1, y2, y3, y4, y5, y6, y7, c1, c2, c3, c4, c5, c6, c7, s1, s2, s3, s4, s5, s6, s7, c45, dc, ds, rad$;
 $span := ks$; $ks := 2 \uparrow m$; $rad := 4.0 \times \arctan(1.0)/ks$;
 $ks := span \div ks$; $n := n - 1$; $c45 := \text{sqr}(0.5)$; $k := m$;
comment Radix 8 transform;
for $m := m - 3$ **while** $m \geq 0$ **do**
begin
 $c1 := 1.0$; $s1 := 0$; $k0 := 0$; $k := ks$;
 $dc := 2.0 \times \sin(rad) \uparrow 2$; $rad := rad + rad$;
 $ds := \sin(rad)$; $rad := 4 \times rad$;
 $span := span \div 8$;
La: $k1 := k0 + span$; $k2 := k1 + span$; $k3 := k2 + span$;
 $k4 := k3 + span$; $k5 := k4 + span$; $k6 := k5 + span$;
 $k7 := k6 + span$; $A0 := A[k0]$; $B0 := B[k0]$;
 $A1 := A[k1]$; $B1 := B[k1]$;
 $A2 := A[k2]$; $B2 := B[k2]$;
 $A3 := A[k3]$; $B3 := B[k3]$;
 $A4 := A[k4]$; $B4 := B[k4]$;
 $A5 := A[k5]$; $B5 := B[k5]$;
 $A6 := A[k6]$; $B6 := B[k6]$;
 $A7 := A[k7]$; $B7 := B[k7]$;
 $x0 := A0 + A4$; $y0 := B0 + B4$;
 $x4 := A0 - A4$; $y4 := B0 - B4$;
 $x1 := A1 + A5$; $y1 := B1 + B5$;
 $x5 := (A1 - A5 - B1 + B5) \times c45$;
 $y5 := (A1 - A5 + B1 - B5) \times c45$;
 $x2 := A2 + A6$; $y2 := B2 + B6$;
 $x6 := B6 - B2$; $y6 := A2 - A6$;
 $x3 := A3 + A7$; $y3 := B3 + B7$;
 $x7 := (A7 - A3 - B3 + B7) \times c45$;
 $y7 := (A3 - A7 - B3 + B7) \times c45$;
 $A1 := x0 + x2 - x1 - x3$; $B1 := y0 + y2 - y1 - y3$;
 $A2 := x0 - x2 - y1 + y3$; $B2 := y0 - y2 + x1 - x3$;
 $A3 := x0 - x2 + y1 - y3$; $B3 := y0 - y2 - x1 + x3$;
 $A4 := x4 + x6 + x5 + x7$; $B4 := y4 + y6 + y5 + y7$;
 $A5 := x4 + x6 - x5 - x7$; $B5 := y4 + y6 - y5 - y7$;
 $A6 := x4 - x6 - y5 + y7$; $B6 := y4 - y6 + x5 - x7$;
 $A7 := x4 - x6 + y5 - y7$; $B7 := y4 - y6 - x5 + x7$;
 $A[k0] := x0 + x2 + x1 + x3$; $B[k0] := y0 + y2 + y1 + y3$;
if $s1 = 0$ **then**

begin
 $A[k1] := A1$; $B[k1] := B1$;
 $A[k2] := A2$; $B[k2] := B2$;
 $A[k3] := A3$; $B[k3] := B3$;
 $A[k4] := A4$; $B[k4] := B4$;
 $A[k5] := A5$; $B[k5] := B5$;
 $A[k6] := A6$; $B[k6] := B6$;
 $A[k7] := A7$; $B[k7] := B7$

end

else

begin

$A[k1] := c4 \times A1 - s4 \times B1$;
 $B[k1] := s4 \times A1 + c4 \times B1$;
 $A[k2] := c2 \times A2 - s2 \times B2$;
 $B[k2] := s2 \times A2 + c2 \times B2$;
 $A[k3] := c6 \times A3 - s6 \times B3$;
 $B[k3] := s6 \times A3 + c6 \times B3$;
 $A[k4] := c1 \times A4 - s1 \times B4$;
 $B[k4] := s1 \times A4 + c1 \times B4$;

$A[k5] := c5 \times A5 - s5 \times B5$;
 $B[k5] := s5 \times A5 + c5 \times B5$;
 $A[k6] := c3 \times A6 - s3 \times B6$;
 $B[k6] := s3 \times A6 + c3 \times B6$;
 $A[k7] := c7 \times A7 - s7 \times B7$;
 $B[k7] := s7 \times A7 + c7 \times B7$

end;

$k0 := k7 + span$; **if** $k0 < n$ **then go to** *La*;

$k0 := k0 - n$; **if** $k0 \neq k$ **then go to** *La*;

comment Increment sine and cosine values;

if $k0 \neq span$ **then**

begin

$c2 := c1 - (dc \times c1 + ds \times s1)$;

$s1 := (ds \times c1 - dc \times s1) + s1$;

comment The following three statements compensate for truncation error. If rounded arithmetic is used, substitute $c1 := c2$;

$c1 := 1.5 - 0.5 \times (c2 \uparrow 2 + s1 \uparrow 2)$;

$s1 := c1 \times s1$; $c1 := c1 \times c2$;

$c2 := c1 \uparrow 2 - s1 \uparrow 2$; $s2 := 2.0 \times c1 \times s1$;

$c3 := c2 \times c1 - s2 \times s1$; $s3 := c2 \times s1 + s2 \times c1$;

$c4 := c2 \uparrow 2 - s2 \uparrow 2$; $s4 := 2.0 \times c2 \times s2$;

$c5 := c1 \times c4 - s1 \times s4$; $s5 := s1 \times c4 + c1 \times s4$;

$c6 := c3 \uparrow 2 - s3 \uparrow 2$; $s6 := 2.0 \times c3 \times s3$;

$c7 := c1 \times c6 - s1 \times s6$; $s7 := s1 \times c6 + c1 \times s6$;

$k := k + ks$; **go to** *La*

end;

$k3 := m$

end;

comment If m is not a multiple of 3, then complete the transform with radix 2 steps;

for $k3 := k3 - 1$ **while** $k3 \geq 0$ **do**

begin

$k0 := 0$; $span := span \div 2$;

Lb: $k2 := k0 + span$;

$A2 := A[k2]$; $B2 := B[k2]$;

$A[k2] := A[k0] - A2$; $B[k2] := B[k0] - B2$;

$A[k0] := A[k0] + A2$; $B[k0] := B[k0] + B2$;

$k0 := k2 + span$; **if** $k0 < n$ **then go to** *Lb*;

$k0 := k0 - n$; **if** $k0 < ks$ **then go to** *Lb*;

if $ks = span$ **then go to** *Ld*;

Lc: $k2 := k0 + span$;

$A2 := A[k0] - A[k2]$; $B2 := B[k0] - B[k2]$;

$A[k0] := A[k0] + A[k2]$; $B[k0] := B[k0] + B[k2]$;

$A[k2] := -B2$; $B[k2] := A2$;

$k0 := k2 + span$; **if** $k0 < n$ **then go to** *Lc*;

$k0 := k0 - n$; **if** $k0 < span$ **then go to** *Lc*;

Ld: **end**

end FFT8;

procedure *REVFFT8*(A, B, n, m, ks); **value** n, m, ks ;

integer n, m, ks ; **array** A, B ;

comment This procedure computes the fast Fourier transform for one variable of dimension 2^m in a multivariate transform. n is the number of data points, i.e., $n = n_1 \times n_2 \times \dots \times n_p$ for a p -variate transform, and $ks = n_{k+1} \times n_{k+2} \times \dots \times n_p$, where $n_k = 2^m$ is the dimension of the current variable. Arrays $A[0:n-1]$ and $B[0:n-1]$ originally contain the real and imaginary components of the data with the indices of each variable in reverse binary order, e.g. a_{jkl} is in $A[j' \times n_2 \times n_3 + k' \times n_3 + l']$ for $j = 0, 1, \dots, n_1 - 1$, $k = 0, 1, \dots, n_2 - 1$, and $l = 0, 1, \dots, n_3 - 1$, where $j', k',$ and l' are the bit-reversed values of $j, k,$ and l . On completion of the multivariate transform, the real and imaginary components of the resulting Fourier coefficients are in A and B in normal order. If $n = 2^m$ and $ks = 1$, a single-variate transform is computed;

begin integer $k0, k1, k2, k3, k4, k5, k6, k7, k, span$;

real $A0, A1, A2, A3, A4, A5, A6, A7, B0, B1, B2, B3, B4, B5, B6, B7, x0, x1, x2, x3, x4, x5, x6, x7, y0, y1, y2, y3, y4, y5, y6, y7,$

```

c1, c2, c3, c4, c5, c6, c7, s1, s2, s3, s4, s5, s6, s7, c45, dc, ds, rad;
rad := 4.0 × arctan(1.0); n := n - 1;
c45 := sqrt(0.5); span := ks;
comment Compute radix 2 steps if m is not a multiple of 3;
k3 := (m ÷ 3) × 3;
for k3 := k3 + 1 while k3 ≤ m do
begin
k0 := 0;
La: k2 := k0 + span;
A2 := A[k2]; B2 := B[k2];
A[k2] := A[k0] - A2; B[k2] := B[k0] - B2;
A[k0] := A[k0] + A2; B[k0] := B[k0] + B2;
k0 := k2 + span; if k0 < n then go to La;
k0 := k0 - n; if k0 < ks then go to La;
if ks = span then go to Lc;
Lb: k2 := k0 + span;
A2 := A[k2]; B2 := B[k2];
A[k2] := A[k0] + B2; B[k2] := B[k0] - A2;
A[k0] := A[k0] - B2; B[k0] := B[k0] + A2;
k0 := k2 + span; if k0 < n then go to Lb;
k0 := k0 - n; if k0 < span then go to Lb;
Lc: span := span + span; rad := 0.5 × rad
end;
comment Radix 8 transform;
for m := m - 3 while m ≥ 0 do
begin
c1 := 1.0; s1 := 0; k0 := 0; k := ks;
rad := 0.125 × rad; dc := 2.0 × sin(rad) ↑ 2;
ds := sin(rad+rad);
Ld: k1 := k0 + span; k2 := k1 + span; k3 := k2 + span;
k4 := k3 + span; k5 := k4 + span; k6 := k5 + span;
k7 := k6 + span; A0 := A[k0]; B0 := B[k0];
if s1 = 0 then
begin
A1 := A[k1]; B1 := B[k1];
A2 := A[k2]; B2 := B[k2];
A3 := A[k3]; B3 := B[k3];
A4 := A[k4]; B4 := B[k4];
A5 := A[k5]; B5 := B[k5];
A6 := A[k6]; B6 := B[k6];
A7 := A[k7]; B7 := B[k7];
end
else
begin
A1 := A[k1] × c4 - B[k1] × s4;
B1 := A[k1] × s4 + B[k1] × c4;
A2 := A[k2] × c2 - B[k2] × s2;
B2 := A[k2] × s2 + B[k2] × c2;
A3 := A[k3] × c6 - B[k3] × s6;
B3 := A[k3] × s6 + B[k3] × c6;
A4 := A[k4] × c1 - B[k4] × s1;
B4 := A[k4] × s1 + B[k4] × c1;
A5 := A[k5] × c5 - B[k5] × s5;
B5 := A[k5] × s5 + B[k5] × c5;
A6 := A[k6] × c3 - B[k6] × s3;
B6 := A[k6] × s3 + B[k6] × c3;
A7 := A[k7] × c7 - B[k7] × s7;
B7 := A[k7] × s7 + B[k7] × c7;
end;
x0 := A0 + A1 + A2 + A3; y0 := B0 + B1 + B2 + B3;
x1 := A0 - A1 - B2 + B3; y1 := B0 - B1 + A2 - A3;
x2 := A0 + A1 - A2 - A3; y2 := B0 + B1 - B2 - B3;
x3 := A0 - A1 + B2 - B3; y3 := B0 - B1 - A2 + A3;
x4 := A4 + A5 + A6 + A7; y4 := B4 + B5 + B6 + B7;
x5 := (A4 - A5 - B6 + B7) × c45;
y5 := (B4 - B5 + A6 - A7) × c45;
x6 := A4 + A5 - A6 - A7; y6 := B4 + B5 - B6 - B7;
x7 := (A4 - A5 + B6 - B7) × c45;

```

```

y7 := (B4 - B5 - A6 + A7) × c45;
A[k0] := x0 + x4; B[k0] := y0 + y4;
A[k1] := x1 + x5 - y5; B[k1] := y1 + x5 + y5;
A[k2] := x2 - y6; B[k2] := y2 + x6;
A[k3] := x3 - x7 - y7; B[k3] := y3 + x7 - y7;
A[k4] := x0 - x4; B[k4] := y0 - y4;
A[k5] := x1 - x5 + y5; B[k5] := y1 - x5 - y5;
A[k6] := x2 + y6; B[k6] := y2 - x6;
A[k7] := x3 + x7 + y7; B[k7] := y3 - x7 + y7;
k0 := k7 + span; if k0 < n then go to Ld;
k0 := k0 - n; if k0 < k then go to Ld;
comment Increment the sine and cosine values;
if k0 ≠ span then
begin
c2 := c1 - (dc × c1 + ds × s1);
s1 := (ds × c1 - dc × s1) + s1;
comment The following three statements compensate
for truncation error. If rounded arithmetic is used,
substitute c1 := c2;
c1 := 1.5 - 0.5 × (c2 ↑ 2 + s1 ↑ 2);
s1 := c1 × s1; c1 := c1 × c2;
c2 := c1 ↑ 2 - s1 ↑ 2; s2 := 2.0 × c1 × s1;
c3 := c1 × c2 - s1 × s2; s3 := s1 × c2 + c1 × s2;
c4 := c2 ↑ 2 - s2 ↑ 2; s4 := 2.0 × c2 × s2;
c5 := c1 × c4 - s1 × s4; s5 := s1 × c4 + c1 × s4;
c6 := c3 ↑ 2 - s3 ↑ 2; s6 := 2.0 × c3 × s3;
c7 := c1 × c6 - s1 × s6; s7 := s1 × c6 + c1 × s6;
k := k + ks; go to Ld
end;
span := 8 × span
end
end REVFFTS

```

ALGORITHM 346

F-TEST PROBABILITIES [S14]

JOHN MORRIS (Recd. 10 Apr. 1968, 12 Sept. 1968, and 6 Nov. 1968)

Computer Institute for Social Science Research, Michigan State University, East Lansing, MI 48823

KEY WORDS AND PHRASES: F-test, Snedecor F-statistic, Fisher test, distribution function

CR CATEGORIES: 5.5

procedure *Ftest* (*f*, *df1*, *df2*, *maxn*, *prob*, *gauss*, *error*);

value *f*, *df1*, *df2*, *maxn*; **real** *f*, *prob*; **integer** *df1*, *df2*, *maxn*;
real procedure *gauss*; **label** *error*;

comment This procedure gives the probability that *F* will be greater than the value of *f* where

$$f = \sigma_1^2 / \sigma_2^2,$$

σ_1^2 is the variance of the sample with size N_1 , σ_2^2 is the variance of the sample with size N_2 , $df1 = N_1 - 1$, $df2 = N_2 - 1$, and *F* is the Snedecor-Fisher statistic as defined and tabled by Snedecor [4].

The present algorithm computes a value which is directly related to that of Algorithm 322, such that *prob* = 1 - *Fisher*. A number of test runs on various computers suggest that *Ftest* may be considerably faster than *Fisher*.

An approximation is included to limit execution time when sample size is large. It should be used when register overflow would otherwise result, and the appropriate value for *maxn* will therefore depend upon the specific implementation. When *maxn* = 500 the approximation appears to give three-digit

accuracy. The **real procedure gauss** computes the area under the left-hand portion of the normal curve. Algorithm 209 [3] may be used for this purpose. If $f < 0$ or if $df1 < 1$ or if $df2 < 1$ then exit to the label *error* occurs.

National Bureau of Standards formulas 26.6.4, 26.6.5, and 26.6.8 are used for computation of the statistic, and 26.6.15 is used for the approximation [2].

Thanks to Mary E. Rafter for extensive testing of this procedure and to the referee for a number of suggestions.

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```
begin
  if df1 < 1 ∨ df2 < 1 ∨ f < 0.0 then go to error;
  if f = 0.0 then prob := 1.0
  else
    begin
```

```
      real f1, f2, x, ft, vp;
      f1 := df1; f2 := df2; ft := 0.0;
      x := f2/(f2+f1×f); vp := f1 + f2 - 2.0;
      if 2 × (df1÷2) = df1 ∧ df1 ≤ maxn then
        begin
          real xx; xx := 1.0 - x;
          for f1 := f1 - 2.0 step - 2.0 until 1.0 do
            begin
              vp := vp - 2.0;
              ft := xx × vp/f1 × (1.0+ft)
            end;
          ft := x ↑ (0.5×f2) × (1.0+ft)
        end
```

```
      else if 2 × (df2 ÷ 2) = df2 ∧ df2 ≤ maxn then
        begin
          for f2 := f2 - 2.0 step - 2.0 until 1.0 do
            begin
              vp := vp - 2.0;
              ft := x × vp/f2 × (1.0+ft)
            end;
          ft := 1.0 - (1.0-x) ↑ (0.5×f1) × (1.0+ft)
        end
```

```
      end
      else if df1 + df2 ≤ maxn then
        begin
          real theta, sth, cth, sts, cts, a, b, xi, gamma;
          theta := arctan(sqrt(f1×f2));
          sth := sin(theta); cth := cos(theta);
          sts := sth ↑ 2; cts := cth ↑ 2;
          a := b := 0.0;
```

```
          if df2 > 1 then
            begin
              for f2 := f2 - 2.0 step - 2.0 until 2.0 do
                a := cts × (f2-1.0)/f2 × (1.0+a);
                a := sth × cth × (1.0+a)
              end;
          a := theta + a;
          if df1 > 1 then
            begin
              for f1 := f1 - 2.0 step - 2.0 until 2.0 do
                begin
                  vp := vp - 2.0;
                  b := sts × vp/f1 × (1.0+b)
                end;
          gamma := 1.0; f2 := 0.5 × df2;
```

```
          for xi := 1.0 step 1.0 until f2 do
            gamma := xi × gamma/(xi-0.5);
            b := gamma × sth × cth ↑ df2 × (1.0+b)
          end;
          ft := 1.0 + 0.636619772368 × (b-a);
          comment 0.6366197723675813430755351 ... = 2.0/π;
        end
      else
        begin
          real cbrf;
          f1 := 2.0/(9.0 × f1); f2 := 2.0/(9.0×f2);
          cbrf := f ↑ 0.333333333333;
          ft := gauss(-(1.0-f2)×cbrf+f1-1.0)/
            sqrt(f2×cbrf ↑ 2+f1)
        end;
      prob := if ft < 0.0 then 0.0 else ft
    end
  end Ftest
```

ALGORITHM 347

AN EFFICIENT ALGORITHM FOR SORTING WITH MINIMAL STORAGE [M1]

RICHARD C. SINGLETON* (Recd. 17 Sept. 1968)

Mathematical Statistics and Operations Research Department, Stanford Research Institute, Menlo Park, CA 94025

* This work was supported by Stanford Research Institute with Research and Development funds.

KEY WORDS AND PHRASES: sorting, minimal storage sorting, digital computer sorting

CR CATEGORIES: 5.31

```
procedure SORT(A, i, j);
  value i, j; integer i, j;
  array A;
```

comment This procedure sorts the elements of array *A* into ascending order, so that

$$A[k] \leq A[k+1], \quad k = i, i+1, \dots, j-1.$$

The method used is similar to *QUICKERSORT* by R. S. Scowen [5], which in turn is similar to an algorithm given by Hibbard [2, 3] and to Hoare's *QUICKSORT* [4]. *QUICKERSORT* is used as a standard, as it was shown in a recent comparison to be the fastest among four ACM algorithms tested [1]. On the Burroughs B5500 computer, the present algorithm is about 25 percent faster than *QUICKERSORT* when tested on random uniform numbers (see Table I) and about 40 percent faster on numbers in natural order (1, 2, ..., *n*), in reverse order (*n*, *n*-1, ..., 1), and sorted by halves (2, 4, ..., *n*, 1, 3, ..., *n*-1). *QUICKERSORT* is slow in sorting data with numerous "tied" observations, a problem that can be corrected by changing the code to exchange elements $a[k] \geq t$ in the lower segment with elements $a[q] \leq t$ in the upper segment. This change gives a better split of the original segment, which more than compensates for the additional interchanges.

In the earlier algorithms, an element with value *t* was selected from the array. Then the array was split into a lower segment with all values less than or equal to *t* and an upper segment with all values greater than or equal to *t*, separated by a third segment of length one and value *t*. The method was then applied

TABLE I. SORTING TIMES IN SECONDS FOR SORT AND QUICKERSORT, ON THE BURROUGHS B5500 COMPUTER—AVERAGE OF FIVE TRIALS

Original order and number of items	Algorithm	
	SORT	QUICKERSORT
Random uniform:		
500	0.48	0.63
1000	1.02	1.40
Natural order:		
500	0.29	0.48
1000	0.62	1.00
Reverse order:		
500	0.30	0.51
1000	0.63	1.08
Sorted by halves:		
500	0.73	1.15
1000	1.72	2.89
Constant value:		
500	0.43	10.60
1000	0.97	41.65

recursively to the lower and upper segments, continuing until all segments were of length one and the data were sorted. The present method differs slightly—the middle segment is usually missing—since the comparison element with value t is not removed from the array while splitting. A more important difference is that the median of the values of $A[i]$, $A[(i+j)\div 2]$, and $A[j]$ is used for t , yielding a better estimate of the median value for the segment than the single element used in the earlier algorithms. Then while searching for a pair of elements to exchange, the previously sorted data (initially, $A[i] \leq t \leq A[j]$) are used to bound the search, and the index values are compared only when an exchange is about to be made. This leads to a small amount of overshoot in the search, adding to the fixed cost of splitting a segment but lowering the variable cost. The longest segment remaining after splitting a segment of n has length less than or equal to $n - 2$, rather than $n - 1$ as in QUICKERSORT.

For efficiency, the upper and lower segments after splitting should be of nearly equal length. Thus t should be close to the median of the data in the segment to be split. For good statistical properties, the median estimate should be based on an odd number of observations. Three gives an improvement over one and the extra effort involved in using five or more observations may be worthwhile on long segments, particularly in the early stages of a sort.

Hibbard [3] suggests using an alternative method, such as Shell's [6], to complete the sort on short sequences. An experimental investigation of this idea using the splitting algorithm adopted here showed no improvement in going beyond the final stage of Shell's algorithm, i.e. the familiar "sinking" method of sorting by interchange of adjacent pairs. The minimum time was obtained by sorting sequences of 11 or fewer items by this method. Again the number of comparisons is reduced by using the data themselves to bound the downward search. This requires

$$A[i-1] \leq A[k], \quad i \leq k \leq j.$$

Thus the initial segment cannot be sorted in this way. The initial segment is treated as a special case and sorted by the splitting algorithm. Because of this feature, the present algorithm lacks the pure recursive structure of the earlier algorithms.

For n elements to be sorted, where $2^k \leq n < 2^{k+1}$, a maximum of k elements each are needed in arrays IL and IU . On the B5500 computer, single-dimensional arrays have a maximum length of 1023. Thus the array bounds [0:8] suffice.

This algorithm was developed as a FORTRAN subroutine, then translated to ALGOL. The original FORTRAN version follows:

```

SUBROUTINE SORT(A,II,JJ)
C SORTS ARRAY A INTO INCREASING ORDER, FROM A(II) TO A(JJ)
C ORDERING IS BY INTEGER SUBTRACTION, THUS FLOATING POINT
C NUMBERS MUST BE IN NORMALIZED FORM.
C ARRAYS IU(K) AND IL(K) PERMIT SORTING UP TO 2**(K+1)-1 ELEMENTS
DIMENSION A(1),IU(16),IL(16)
INTEGER A,T,TT
M=1
I=II
J=JJ
5 IF(I .GE. J) GO TO 70
10 K=I
IJ=(J+I)/2
T=A(IJ)
IF(A(I) .LE. T) GO TO 20
A(IJ)=A(I)
A(I)=T
T=A(IJ)
20 L=J
IF(A(J) .GE. T) GO TO 40
A(IJ)=A(J)
A(J)=T
T=A(IJ)
IF(A(I) .LE. T) GO TO 40
A(IJ)=A(I)
A(I)=T
T=A(IJ)
GO TO 40
30 A(L)=A(K)
A(K)=TT
40 L=L-1
IF(A(L) .GT. T) GO TO 40
TT=A(L)
50 K=K+1
IF(A(K) .LT. T) GO TO 50
IF(K .LE. L) GO TO 30
IF(L-I .LE. J-K) GO TO 60
IL(M)=I
IU(M)=L
I=K
M=M+1
GO TO 80
60 IL(M)=K
IU(M)=J
J=L
M=M+1
GO TO 80
70 M=M-1
IF(M .EQ. 0) RETURN
I=IL(M)
J=IU(M)
80 IF(J-I .GE. 11) GO TO 10
IF(I .EQ. II) GO TO 5
I=I-1
90 I=I+1
IF(I .EQ. J) GO TO 70
T=A(I+1)
IF(A(I) .LE. T) GO TO 90
K=I
100 A(K+1)=A(K)
K=K-1
IF(T .LT. A(K)) GO TO 100
A(K+1)=T
GO TO 90
END

```

This FORTRAN subroutine was tested on a CDC 6400 computer. For random uniform numbers, sorting times divided by $n \log_2 n$ were nearly constant at 20.2×10^{-6} for $100 \leq n \leq 10,000$, with a time of 0.202 seconds for 1000 items. This subroutine was also hand-compiled for the same computer to produce a more efficient machine code. In this version the constant of proportionality was 5.2×10^{-6} , with a time of 0.052 seconds for 1000 items. In both cases, integer comparisons were used to order normalized floating-point numbers.

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```

begin
  real t, tt;
  integer ii, ij, k, L, m;
  integer array IL, IU[0:8];
  m := 0; ii := i; go to L4;
L1: ij := (i+j) ÷ 2; t := A[ij]; k := i; L := j;
  if A[i] > t then
    begin A[ij] := A[i]; A[i] := t; t := A[ij] end;
  if A[j] < t then
    begin
      A[ij] := A[j]; A[j] := t; t := A[ij];
      if A[i] > t then
        begin A[ij] := A[i]; A[i] := t; t := A[ij] end
    end;
L2: L := L - 1;
  if A[L] > t then go to L2;
  tt := A[L];
L3: k := k + 1;
  if A[k] < t then go to L3;
  if k ≤ L then
    begin A[L] := A[k]; A[k] := tt; go to L2 end;
  if L - i > j - k then
    begin IL[m] := i; IU[m] := L; i := k end
  else
    begin IL[m] := k; IU[m] := j; j := L end;
  m := m + 1;
L4: if j - i > 10 then go to L1;
  if i = ii then
    begin if i < j then go to L1 end;
    for i := i + 1 step 1 until j do
      begin
        t := A[i]; k := i - 1;
        if A[k] > t then
          begin
L5: A[k+1] := A[k]; k := k - 1;
            if A[k] > t then go to L5;
            A[k+1] := t
          end
        end;
      m := m - 1; if m ≥ 0 then
        begin i := IL[m]; j := IU[m]; go to L4 end
    end SORT

```

REMARK ON ALGORITHM 329 [G6]
 DISTRIBUTION OF INDISTINGUISHABLE OBJECTS INTO DISTINGUISHABLE SLOTS [Robert R. Fenichel, *Comm. ACM* 11 (June 1968), 430]
 M. GRAY (Recd. 20 Sept. 1968)
 Computing Science Department, University of Adelaide,
 South Australia

As the procedure stands it is incorrect. Preceding
 end skip 99,189,198, etc.
 the following statement should be inserted:
 if $q[k] \neq 0$ then $LeftmostZero := k + 1$
 Thus the compound statement becomes:

```

begin
  LeftmostZero := LeftmostZero - 1;
  q[k] := q[LeftmostZero] - 1;
  q[LeftmostZero] := 0;
  q[LeftmostZero-1] := q[LeftmostZero-1] + 1;
  if  $q[k] \neq 0$  then  $LeftmostZero := k + 1$ 
end skip 99, 189, 198, etc.

```

REMARK ON ALGORITHM 339 [C6]
 AN ALGOL PROCEDURE FOR THE FAST FOURIER
 TRANSFORM WITH ARBITRARY FACTORS
 [Richard C. Singleton, *Comm. ACM* 11 (Nov. 1968),
 776]

RICHARD C. SINGLETON (Recd. 27 Nov. 1968)
 Stanford Research Institute, Menlo Park, CA 94025

KEY WORDS AND PHRASES: fast Fourier transform, complex
 Fourier transform, multivariate Fourier transform, Fourier
 series, harmonic analysis, spectral analysis, orthogonal poly-
 nomials, orthogonal transformation, virtual core memory,
 permutation
 CR CATEGORIES. 3.15, 3.83, 5.12, 5.14

On page 778, column 2, the 7th and 6th lines from the bottom
 should be corrected to read:

```

LJ: jj := C[i-2] + jj; if jj ≥ C[i-1] then
  begin i := i - 1; jj := jj - C[i]; go to LJ end;

```

On page 779, column 1, the 9th and 8th lines from the bottom
 should be corrected to read:

```

LX: jj := D[k+1] + jj; if jj ≥ D[k] then
  begin jj := jj - D[k]; k := k + 1; go to LX end;

```

In both cases jj was originally used as the controlled variable of
 a for clause and thus was undefined after exit; the corrections
 preserve the value of jj for later use.

If the user prefers to compute constants with library functions,
 line 5 in column 2 on page 777 may be replaced by:

```

rad := 8.0 × arctan(1.0); c30 := sqrt(0.75);

```

Algorithms 338 [*Comm. ACM* 11 (Nov. 1968), 773] and 339 were
 punched from the printed page and tested on the CDC 6400
 ALGOL compiler. After changing a colon to a semicolon at the end
 of line 37 in column 2 on page 775, the test results agreed with
 those obtained earlier with this compiler.

When computing a single-variate Fourier transform of real
 data, procedure *REALTRAN* may be used with procedure *FFT*
 (Algorithm 339) to reduce computing time. Two versions of
REALTRAN have been given (Algorithms 338 and 345 [*Comm.*
ACM 12 (Mar. 1969), 179-184]); the first version is the faster of
 the two, but the second should be used if arithmetic results for
 real quantities are truncated rather than rounded.

In describing the evaluation of a real Fourier series, in the
 middle of column 2 on page 776, the necessary steps of reversing
 the signs of the B array values both before and after calling *FFT*
 were omitted. The correct steps, including scaling, are as follows:

```

REALTRAN(A, B, n, true);
for j := n - 1 step -1 until 0 do B[j] := -B[j];
FFT(A, B, n, n, n);
for j := n - 1 step -1 until 0 do
  begin A[j] := 0.5 × A[j]; B[j] := -0.5 × B[j] end;

```

The policy concerning the contributions of algorithms to
Communications of the ACM appears, most recently, in the
 January 1969 issue, page 39. A contribution should be in the
 form of an algorithm, a certification, or a remark. An al-
 gorithm must normally be written in the ALGOL 60 Refer-
 ence Language or in USASI Standard FORTRAN or Basic
 FORTRAN.