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Chapter 3. Interpolation and Extrapolation

3.0 Introduction

We sometimes know the value of a function f(x) at a set of points x_1, x_2, \ldots, x_N (say, with $x_1 < \ldots < x_N$), but we don't have an analytic expression for f(x) that lets us calculate its value at an arbitrary point. For example, the $f(x_i)$'s might result from some physical measurement or from long numerical calculation that cannot be cast into a simple functional form. Often the x_i 's are equally spaced, but not necessarily.

The task now is to estimate f(x) for arbitrary x by, in some sense, drawing a smooth curve through (and perhaps beyond) the x_i . If the desired x is in between the largest and smallest of the x_i 's, the problem is called *interpolation*; if x is outside that range, it is called *extrapolation*, which is considerably more hazardous (as many former stock-market analysts can attest).

Interpolation and extrapolation schemes must model the function, between or beyond the known points, by some plausible functional form. The form should be sufficiently general so as to be able to approximate large classes of functions which might arise in practice. By far most common among the functional forms used are polynomials (§3.1). Rational functions (quotients of polynomials) also turn out to be extremely useful (§3.2). Trigonometric functions, sines and cosines, give rise to *trigonometric interpolation* and related Fourier methods, which we defer to Chapters 12 and 13.

There is an extensive mathematical literature devoted to theorems about what sort of functions can be well approximated by which interpolating functions. These theorems are, alas, almost completely useless in day-to-day work: If we know enough about our function to apply a theorem of any power, we are usually not in the pitiful state of having to interpolate on a table of its values!

Interpolation is related to, but distinct from, *function approximation*. That task consists of finding an approximate (but easily computable) function to use in place of a more complicated one. In the case of interpolation, you are given the function f at points *not of your own choosing*. For the case of function approximation, you are allowed to compute the function f at f any desired points for the purpose of developing your approximation. We deal with function approximation in Chapter 5.

One can easily find pathological functions that make a mockery of any interpolation scheme. Consider, for example, the function

$$f(x) = 3x^2 + \frac{1}{\pi^4} \ln\left[(\pi - x)^2 \right] + 1 \tag{3.0.1}$$

which is well-behaved everywhere except at $x=\pi$, very mildly singular at $x=\pi$, and otherwise takes on all positive and negative values. Any interpolation based on the values x=3.13,3.14,3.15,3.16, will assuredly get a very wrong answer for the value x=3.1416, even though a graph plotting those five points looks really quite smooth! (Try it on your calculator.)

Because pathologies can lurk anywhere, it is highly desirable that an interpolation and extrapolation routine should return an estimate of its own error. Such an error estimate can never be foolproof, of course. We could have a function that, for reasons known only to its maker, takes off wildly and unexpectedly between two tabulated points. Interpolation always presumes some degree of smoothness for the function interpolated, but within this framework of presumption, deviations from smoothness can be detected.

Conceptually, the interpolation process has two stages: (1) Fit an interpolating function to the data points provided. (2) Evaluate that interpolating function at the target point x.

However, this two-stage method is generally not the best way to proceed in practice. Typically it is computationally less efficient, and more susceptible to roundoff error, than methods which construct a functional estimate f(x) directly from the N tabulated values every time one is desired. Most practical schemes start at a nearby point $f(x_i)$, then add a sequence of (hopefully) decreasing corrections, as information from other $f(x_i)$'s is incorporated. The procedure typically takes $O(N^2)$ operations. If everything is well behaved, the last correction will be the smallest, and it can be used as an informal (though not rigorous) bound on the error.

In the case of polynomial interpolation, it sometimes does happen that the coefficients of the interpolating polynomial are of interest, even though their use in *evaluating* the interpolating function should be frowned on. We deal with this eventuality in §3.5.

Local interpolation, using a finite number of "nearest-neighbor" points, gives interpolated values f(x) that do not, in general, have continuous first or higher derivatives. That happens because, as x crosses the tabulated values x_i , the interpolation scheme switches which tabulated points are the "local" ones. (If such a switch is allowed to occur anywhere else, then there will be a discontinuity in the interpolated function itself at that point. Bad idea!)

In situations where continuity of derivatives is a concern, one must use the "stiffer" interpolation provided by a so-called *spline* function. A spline is a polynomial between each pair of table points, but one whose coefficients are determined "slightly" nonlocally. The nonlocality is designed to guarantee global smoothness in the interpolated function up to some order of derivative. Cubic splines (§3.3) are the most popular. They produce an interpolated function that is continuous through the second derivative. Splines tend to be stabler than polynomials, with less possibility of wild oscillation between the tabulated points.

The number of points (minus one) used in an interpolation scheme is called the *order* of the interpolation. Increasing the order does not necessarily increase the accuracy, especially in polynomial interpolation. If the added points are distant from the point of interest x, the resulting higher-order polynomial, with its additional constrained points, tends to oscillate wildly between the tabulated values. This oscillation may have no relation at all to the behavior of the "true" function (see Figure 3.0.1). Of course, adding points *close* to the desired point usually does help,

3.0 Introduction 101

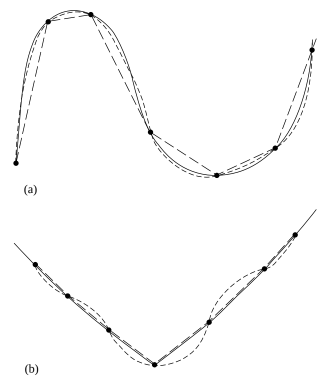


Figure 3.0.1. (a) A smooth function (solid line) is more accurately interpolated by a high-order polynomial (shown schematically as dotted line) than by a low-order polynomial (shown as a piecewise linear dashed line). (b) A function with sharp corners or rapidly changing higher derivatives is *less* accurately approximated by a high-order polynomial (dotted line), which is too "stiff," than by a low-order polynomial (dashed lines). Even some smooth functions, such as exponentials or rational functions, can be badly approximated by high-order polynomials.

but a finer mesh implies a larger table of values, not always available.

Unless there is solid evidence that the interpolating function is close in form to the true function f, it is a good idea to be cautious about high-order interpolation. We enthusiastically endorse interpolations with 3 or 4 points, we are perhaps tolerant of 5 or 6; but we rarely go higher than that unless there is quite rigorous monitoring of estimated errors.

When your table of values contains many more points than the desirable order of interpolation, you must begin each interpolation with a search for the right "local" place in the table. While not strictly a part of the subject of interpolation, this task is important enough (and often enough botched) that we devote §3.4 to its discussion.

The routines given for interpolation are also routines for extrapolation. An important application, in Chapter 16, is their use in the integration of ordinary differential equations. There, considerable care is taken with the monitoring of errors. Otherwise, the dangers of extrapolation cannot be overemphasized: An interpolating function, which is perforce an extrapolating function, will typically go berserk when the argument x is outside the range of tabulated values by more than the typical spacing of tabulated points.

Interpolation can be done in more than one dimension, e.g., for a function

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3.1 Polynomial Interpolation and Extrapolation

Through any two points there is a unique line. Through any three points, a unique quadratic. Et cetera. The interpolating polynomial of degree N-1 through the N points $y_1 = f(x_1), y_2 = f(x_2), \ldots, y_N = f(x_N)$ is given explicitly by Lagrange's classical formula,

$$P(x) = \frac{(x-x_2)(x-x_3)...(x-x_N)}{(x_1-x_2)(x_1-x_3)...(x_1-x_N)} y_1 + \frac{(x-x_1)(x-x_3)...(x-x_N)}{(x_2-x_1)(x_2-x_3)...(x_2-x_N)} y_2 + \dots + \frac{(x-x_1)(x-x_2)...(x-x_{N-1})}{(x_N-x_1)(x_N-x_2)...(x_N-x_{N-1})} y_N$$
(3.1.1)

There are N terms, each a polynomial of degree N-1 and each constructed to be zero at all of the x_i except one, at which it is constructed to be y_i .

It is not terribly wrong to implement the Lagrange formula straightforwardly, but it is not terribly right either. The resulting algorithm gives no error estimate, and it is also somewhat awkward to program. A much better algorithm (for constructing the same, unique, interpolating polynomial) is *Neville's algorithm*, closely related to and sometimes confused with *Aitken's algorithm*, the latter now considered obsolete.

Let P_1 be the value at x of the unique polynomial of degree zero (i.e., a constant) passing through the point (x_1, y_1) ; so $P_1 = y_1$. Likewise define P_2, P_3, \ldots, P_N . Now let P_{12} be the value at x of the unique polynomial of degree one passing through both (x_1, y_1) and (x_2, y_2) . Likewise $P_{23}, P_{34}, \ldots, P_{(N-1)N}$. Similarly, for higher-order polynomials, up to $P_{123...N}$, which is the value of the unique interpolating polynomial through all N points, i.e., the desired answer.

The various P's form a "tableau" with "ancestors" on the left leading to a single "descendant" at the extreme right. For example, with N=4,

$$x_{1}: y_{1} = P_{1}$$

$$p_{12}$$

$$x_{2}: y_{2} = P_{2} P_{123}$$

$$P_{23} P_{1234}$$

$$x_{3}: y_{3} = P_{3} P_{234}$$

$$P_{34}$$

$$x_{4}: y_{4} = P_{4}$$

$$(3.1.2)$$

Neville's algorithm is a recursive way of filling in the numbers in the tableau a column at a time, from left to right. It is based on the relationship between a "daughter" P and its two "parents,"

$$P_{i(i+1)\dots(i+m)} = \frac{(x - x_{i+m})P_{i(i+1)\dots(i+m-1)} + (x_i - x)P_{(i+1)(i+2)\dots(i+m)}}{x_i - x_{i+m}}$$
(3.1.3)

This recurrence works because the two parents already agree at points $x_{i+1} \dots x_{i+m-1}$.

An improvement on the recurrence (3.1.3) is to keep track of the small differences between parents and daughters, namely to define (for $m=1,2,\ldots,N-1$),

$$C_{m,i} \equiv P_{i...(i+m)} - P_{i...(i+m-1)}$$

$$D_{m,i} \equiv P_{i...(i+m)} - P_{(i+1)...(i+m)}.$$
(3.1.4)

Then one can easily derive from (3.1.3) the relations

$$D_{m+1,i} = \frac{(x_{i+m+1} - x)(C_{m,i+1} - D_{m,i})}{x_i - x_{i+m+1}}$$

$$C_{m+1,i} = \frac{(x_i - x)(C_{m,i+1} - D_{m,i})}{x_i - x_{i+m+1}}$$
(3.1.5)

At each level m, the C's and D's are the corrections that make the interpolation one order higher. The final answer $P_{1...N}$ is equal to the sum of $any\ y_i$ plus a set of C's and/or D's that form a path through the family tree to the rightmost daughter.

Here is a routine for polynomial interpolation or extrapolation:

```
SUBROUTINE polint(xa,ya,n,x,y,dy) INTEGER n,NMAX REAL dy,x,y,xa(n),ya(n) PARAMETER (NMAX=10) Largest anticipated value of n. Given arrays xa and ya, each of length n, and given a value x, this routine returns a value y, and an error estimate dy. If P(x) is the polynomial of degree N-1 such that P(xa_i) = ya_i, i = 1, \ldots, n, then the returned value y = P(x). INTEGER i,m,ns REAL den,dif,dift,ho,hp,w,c(NMAX),d(NMAX) ns=1 dif=abs(x-xa(1))
```

END

```
do 11 i=1,n
                                Here we find the index ns of the closest table entry,
    dift=abs(x-xa(i))
    if (dift.lt.dif) then
        ns=i
        dif=dift
    endif
    c(i)=ya(i)
                                and initialize the tableau of c's and d's.
    d(i)=ya(i)
                               This is the initial approximation to y.
y=ya(ns)
ns=ns-1
                                For each column of the tableau,
do 13 m=1,n-1
                                we loop over the current c's and d's and update them.
    do 12 i=1,n-m
        ho=xa(i)-x
        hp=xa(i+m)-x
        w=c(i+1)-d(i)
        den=ho-hp
        if(den.eq.0.)pause 'failure in polint'
          This error can occur only if two input xa's are (to within roundoff) identical.
        den=w/den
        d(i)=hp*den
                                Here the c's and d's are updated.
        c(i)=ho*den
    enddo 12
    if (2*ns.lt.n-m)then
                                After each column in the tableau is completed, we decide
        dv=c(ns+1)
                                   which correction, c or d, we want to add to our accu-
                                   mulating value of y, i.e., which path to take through
        dy=d(ns)
                                   the tableau—forking up or down. We do this in such a
        ns=ns-1
                                   way as to take the most "straight line" route through the
                                   tableau to its apex, updating ns accordingly to keep track
    y=y+dy
                                   of where we are. This route keeps the partial approxima-
enddo 13
                                   tions centered (insofar as possible) on the target x. The
                                   last dy added is thus the error indication.
return
```

Quite often you will want to call polint with the dummy arguments xa and ya replaced by actual arrays with offsets. For example, the construction call polint(xx(15),yy(15),4,x,y,dy) performs 4-point interpolation on the tabulated values xx(15:18), yy(15:18). For more on this, see the end of $\S 3.4$.

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3.2 Rational Function Interpolation and Extrapolation

Some functions are not well approximated by polynomials, but *are* well approximated by rational functions, that is quotients of polynomials. We denote by $R_{i(i+1)...(i+m)}$ a rational function passing through the m+1 points

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$$R_{i(i+1)\dots(i+m)} = \frac{P_{\mu}(x)}{Q_{\nu}(x)} = \frac{p_0 + p_1 x + \dots + p_{\mu} x^{\mu}}{q_0 + q_1 x + \dots + q_{\nu} x^{\nu}}$$
(3.2.1)

Since there are $\mu + \nu + 1$ unknown p's and q's (q_0 being arbitrary), we must have

$$m+1 = \mu + \nu + 1 \tag{3.2.2}$$

In specifying a rational function interpolating function, you must give the desired order of both the numerator and the denominator.

Rational functions are sometimes superior to polynomials, roughly speaking, because of their ability to model functions with poles, that is, zeros of the denominator of equation (3.2.1). These poles might occur for real values of x, if the function to be interpolated itself has poles. More often, the function f(x) is finite for all finite $real\ x$, but has an analytic continuation with poles in the complex x-plane. Such poles can themselves ruin a polynomial approximation, even one restricted to real values of x, just as they can ruin the convergence of an infinite power series in x. If you draw a circle in the complex plane around your m tabulated points, then you should not expect polynomial interpolation to be good unless the nearest pole is rather far outside the circle. A rational function approximation, by contrast, will stay "good" as long as it has enough powers of x in its denominator to account for (cancel) any nearby poles.

For the interpolation problem, a rational function is constructed so as to go through a chosen set of tabulated functional values. However, we should also mention in passing that rational function approximations can be used in analytic work. One sometimes constructs a rational function approximation by the criterion that the rational function of equation (3.2.1) itself have a power series expansion that agrees with the first m+1 terms of the power series expansion of the desired function f(x). This is called $Pad\acute{e}$ approximation, and is discussed in §5.12.

Bulirsch and Stoer found an algorithm of the Neville type which performs rational function extrapolation on tabulated data. A tableau like that of equation (3.1.2) is constructed column by column, leading to a result and an error estimate. The Bulirsch-Stoer algorithm produces the so-called *diagonal* rational function, with the degrees of numerator and denominator equal (if m is even) or with the degree of the denominator larger by one (if m is odd, cf. equation 3.2.2 above). For the derivation of the algorithm, refer to [1]. The algorithm is summarized by a recurrence relation exactly analogous to equation (3.1.3) for polynomial approximation:

$$\begin{split} R_{i(i+1)\dots(i+m)} &= R_{(i+1)\dots(i+m)} \\ &+ \frac{R_{(i+1)\dots(i+m)} - R_{i\dots(i+m-1)}}{\left(\frac{x-x_i}{x-x_{i+m}}\right)\left(1 - \frac{R_{(i+1)\dots(i+m)} - R_{i\dots(i+m-1)}}{R_{(i+1)\dots(i+m)} - R_{(i+1)\dots(i+m-1)}}\right) - 1} \\ &\qquad \qquad (3.2.3) \end{split}$$

This recurrence generates the rational functions through m+1 points from the ones through m and (the term $R_{(i+1)...(i+m-1)}$ in equation 3.2.3) m-1 points. It is started with

$$R_i = y_i \tag{3.2.4}$$

and with

enddo 11

y=ya(ns) ns=ns-1 do 13 m=1,n-1 do 12 i=1,n-m

$$R \equiv [R_{i(i+1)...(i+m)} \quad \text{with} \quad m = -1] = 0$$
 (3.2.5)

Now, exactly as in equations (3.1.4) and (3.1.5) above, we can convert the recurrence (3.2.3) to one involving only the small differences

$$C_{m,i} \equiv R_{i...(i+m)} - R_{i...(i+m-1)}$$

$$D_{m,i} \equiv R_{i...(i+m)} - R_{(i+1)...(i+m)}$$
(3.2.6)

Note that these satisfy the relation

$$C_{m+1,i} - D_{m+1,i} = C_{m,i+1} - D_{m,i} (3.2.7)$$

which is useful in proving the recurrences

$$D_{m+1,i} = \frac{C_{m,i+1}(C_{m,i+1} - D_{m,i})}{\left(\frac{x - x_i}{x - x_{i+m+1}}\right) D_{m,i} - C_{m,i+1}}$$

$$C_{m+1,i} = \frac{\left(\frac{x - x_i}{x - x_{i+m+1}}\right) D_{m,i}(C_{m,i+1} - D_{m,i})}{\left(\frac{x - x_i}{x - x_{i+m+1}}\right) D_{m,i} - C_{m,i+1}}$$
(3.2.8)

This recurrence is implemented in the following subroutine, whose use is analogous in every way to polint in §3.1.

```
SUBROUTINE ratint(xa,ya,n,x,y,dy)
INTEGER n, NMAX
REAL dy,x,y,xa(n),ya(n),TINY
PARAMETER (NMAX=10,TINY=1.e-25)
                                      Largest expected value of n, and a small number.
   Given arrays xa and ya, each of length n, and given a value of x, this routine returns a
   value of y and an accuracy estimate dy. The value returned is that of the diagonal rational
   function, evaluated at x, which passes through the n points (xa_i, ya_i), i = 1...n.
INTEGER i,m,ns
REAL dd,h,hh,t,w,c(NMAX),d(NMAX)
hh=abs(x-xa(1))
do 11 i=1,n
    h=abs(x-xa(i))
    if (h.eq.0.)then
        y=ya(i)
        dy=0.0
        return
    else if (h.lt.hh) then
        ns=i
        hh=h
    endif
    c(i)=ya(i)
    d(i)=ya(i)+TINY
                                       The TINY part is needed to prevent a rare zero-over-
```

zero condition.

```
w=c(i+1)-d(i)
        h=xa(i+m)-x
                                       h will never be zero, since this was tested in the ini-
        t=(xa(i)-x)*d(i)/h
                                           tializing loop.
        dd=t-c(i+1)
        if(dd.eq.0.)pause 'failure in ratint'
          This error condition indicates that the interpolating function has a pole at the re-
          quested value of x.
        dd=w/dd
        d(i)=c(i+1)*dd
        c(i)=t*dd
    enddo 12
    if (2*ns.lt.n-m)then
        dy=c(ns+1)
        dy=d(ns)
        ns=ns-1
    endif
    y=y+dy
enddo 13
return
END
```

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3.3 Cubic Spline Interpolation

Given a tabulated function $y_i = y(x_i)$, i = 1...N, focus attention on one particular interval, between x_j and x_{j+1} . Linear interpolation in that interval gives the interpolation formula

$$y = Ay_i + By_{i+1} (3.3.1)$$

where

$$A \equiv \frac{x_{j+1} - x}{x_{j+1} - x_j} \qquad B \equiv 1 - A = \frac{x - x_j}{x_{j+1} - x_j}$$
(3.3.2)

Equations (3.3.1) and (3.3.2) are a special case of the general Lagrange interpolation formula (3.1.1).

Since it is (piecewise) linear, equation (3.3.1) has zero second derivative in the interior of each interval, and an undefined, or infinite, second derivative at the abscissas x_j . The goal of cubic spline interpolation is to get an interpolation formula that is smooth in the first derivative, and continuous in the second derivative, both within an interval and at its boundaries.

Suppose, contrary to fact, that in addition to the tabulated values of y_i , we also have tabulated values for the function's second derivatives, y'', that is, a set

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personal use. Further reproduction, or any copying of machine-To order Numerical Recipes books, diskettes, or CDROMS of numbers y_i'' . Then, within each interval, we can add to the right-hand side of equation (3.3.1) a cubic polynomial whose second derivative varies linearly from a value y_j'' on the left to a value y_{j+1}'' on the right. Doing so, we will have the desired continuous second derivative. If we also construct the cubic polynomial to have zero *values* at x_j and x_{j+1} , then adding it in will not spoil the agreement with the tabulated functional values y_j and y_{j+1} at the endpoints x_j and x_{j+1} .

A little side calculation shows that there is only one way to arrange this construction, namely replacing (3.3.1) by

$$y = Ay_j + By_{j+1} + Cy_j'' + Dy_{j+1}''$$
(3.3.3)

where A and B are defined in (3.3.2) and

$$C \equiv \frac{1}{6}(A^3 - A)(x_{j+1} - x_j)^2 \qquad D \equiv \frac{1}{6}(B^3 - B)(x_{j+1} - x_j)^2 \qquad (3.3.4)$$

Notice that the dependence on the independent variable x in equations (3.3.3) and (3.3.4) is entirely through the linear x-dependence of A and B, and (through A and B) the cubic x-dependence of C and D.

We can readily check that y'' is in fact the second derivative of the new interpolating polynomial. We take derivatives of equation (3.3.3) with respect to x, using the definitions of A, B, C, D to compute dA/dx, dB/dx, dC/dx, and dD/dx. The result is

$$\frac{dy}{dx} = \frac{y_{j+1} - y_j}{x_{j+1} - x_j} - \frac{3A^2 - 1}{6}(x_{j+1} - x_j)y_j'' + \frac{3B^2 - 1}{6}(x_{j+1} - x_j)y_{j+1}''$$
(3.3.5)

for the first derivative, and

$$\frac{d^2y}{dx^2} = Ay_j'' + By_{j+1}'' \tag{3.3.6}$$

for the second derivative. Since A = 1 at x_j , A = 0 at x_{j+1} , while B is just the other way around, (3.3.6) shows that y'' is just the tabulated second derivative, and also that the second derivative will be continuous across (e.g.) the boundary between the two intervals (x_{j-1}, x_j) and (x_j, x_{j+1}) .

The only problem now is that we supposed the y_i'' 's to be known, when, actually, they are not. However, we have not yet required that the *first* derivative, computed from equation (3.3.5), be continuous across the boundary between two intervals. The key idea of a cubic spline is to require this continuity and to use it to get equations for the second derivatives y_i'' .

The required equations are obtained by setting equation (3.3.5) evaluated for $x = x_j$ in the interval (x_{j-1}, x_j) equal to the same equation evaluated for $x = x_j$ but in the interval (x_j, x_{j+1}) . With some rearrangement, this gives (for j = 2, ..., N-1)

$$\frac{x_{j} - x_{j-1}}{6}y_{j-1}'' + \frac{x_{j+1} - x_{j-1}}{3}y_{j}'' + \frac{x_{j+1} - x_{j}}{6}y_{j+1}'' = \frac{y_{j+1} - y_{j}}{x_{j+1} - x_{j}} - \frac{y_{j} - y_{j-1}}{x_{j} - x_{j-1}}$$
(3.3.7)

These are N-2 linear equations in the N unknowns $y_i'', i=1,\ldots,N$. Therefore there is a two-parameter family of possible solutions.

For a unique solution, we need to specify two further conditions, typically taken as boundary conditions at x_1 and x_N . The most common ways of doing this are either

- set one or both of y₁" and y_N" equal to zero, giving the so-called natural cubic spline, which has zero second derivative on one or both of its boundaries, or
- set either of y_1'' and y_N'' to values calculated from equation (3.3.5) so as to make the first derivative of the interpolating function have a specified value on either or both boundaries.

One reason that cubic splines are especially practical is that the set of equations (3.3.7), along with the two additional boundary conditions, are not only linear, but also tridiagonal. Each y_j'' is coupled only to its nearest neighbors at $j\pm 1$. Therefore, the equations can be solved in O(N) operations by the tridiagonal algorithm (§2.4). That algorithm is concise enough to build right into the spline calculational routine. This makes the routine not completely transparent as an implementation of (3.3.7), so we encourage you to study it carefully, comparing with tridag (§2.4).

```
SUBROUTINE spline(x,y,n,yp1,ypn,y2)
INTEGER n, NMAX
REAL yp1, ypn, x(n), y(n), y2(n)
PARAMETER (NMAX=500)
    Given arrays x(1:n) and y(1:n) containing a tabulated function, i.e., y_i = f(x_i), with
    x_1 < x_2 < \ldots < x_N, and given values yp1 and ypn for the first derivative of the inter-
    polating function at points 1 and n, respectively, this routine returns an array y2(1:n) of
   length n which contains the second derivatives of the interpolating function at the tabulated points x_i. If yp1 and/or ypn are equal to 1\times 10^{30} or larger, the routine is signaled to set
    the corresponding boundary condition for a natural spline, with zero second derivative on
    that boundary.
    Parameter: NMAX is the largest anticipated value of n.
INTEGER i,k
REAL p,qn,sig,un,u(NMAX)
                                              The lower boundary condition is set either to be
if (yp1.gt..99e30) then
    y2(1)=0.
                                                   "natural"
    u(1)=0.
                                              or else to have a specified first derivative.
else
    y2(1) = -0.5
    u(1)=(3./(x(2)-x(1)))*((y(2)-y(1))/(x(2)-x(1))-yp1)
endif
                                              This is the decomposition loop of the tridiagonal
do 11 i=2,n-1
    sig=(x(i)-x(i-1))/(x(i+1)-x(i-1))
                                                  algorithm. y2 and u are used for temporary
    p=sig*y2(i-1)+2.
                                                  storage of the decomposed factors.
    y2(i)=(sig-1.)/p
    u(i)=(6.*((y(i+1)-y(i))/(x(i+1)-x(i))-(y(i)-y(i-1))
          /(x(i)-x(i-1)))/(x(i+1)-x(i-1))-sig*u(i-1))/p
enddo 11
if (ypn.gt..99e30) then
                                              The upper boundary condition is set either to be
    qn=0.
                                                   "natural'
    un=0.
else
                                              or else to have a specified first derivative.
    un=(3./(x(n)-x(n-1)))*(ypn-(y(n)-y(n-1))/(x(n)-x(n-1)))
y2(n)=(un-qn*u(n-1))/(qn*y2(n-1)+1.)
                                              This is the backsubstitution loop of the tridiago-
do 12 k=n-1,1,-1
    y2(k)=y2(k)*y2(k+1)+u(k)
                                                  nal algorithm.
return
END
```

It is important to understand that the program spline is called only *once* to process an entire tabulated function in arrays x_i and y_i . Once this has been done,

values of the interpolated function for any value of x are obtained by calls (as many as desired) to a separate routine splint (for "spline interpolation"):

```
SUBROUTINE splint(xa,ya,y2a,n,x,y)
INTEGER n
REAL x,y,xa(n),y2a(n),ya(n)
   Given the arrays xa(1:n) and ya(1:n) of length n, which tabulate a function (with the
   xa_i's in order), and given the array y2a(1:n), which is the output from spline above,
   and given a value of x, this routine returns a cubic-spline interpolated value y.
INTEGER k,khi,klo
REAL a,b,h
klo=1
                               We will find the right place in the table by means of bisection.
                               This is optimal if sequential calls to this routine are at random
khi=n
if (khi-klo.gt.1) then
                                   values of x. If sequential calls are in order, and closely
    k=(khi+klo)/2
                                   spaced, one would do better to store previous values of
    if(xa(k).gt.x)then
                                   klo and khi and test if they remain appropriate on the
        khi=k
    else
        klo=k
    endif
goto 1
endif
                               klo and khi now bracket the input value of x.
h=xa(khi)-xa(klo)
if (h.eq.0.) pause 'bad xa input in splint'
                                                  The xa's must be distinct.
a=(xa(khi)-x)/h
                               Cubic spline polynomial is now evaluated.
b=(x-xa(klo))/h
y=a*ya(klo)+b*ya(khi)+
         ((a**3-a)*y2a(klo)+(b**3-b)*y2a(khi))*(h**2)/6.
return
END
```

CITED REFERENCES AND FURTHER READING:

De Boor, C. 1978, A Practical Guide to Splines (New York: Springer-Verlag).

Forsythe, G.E., Malcolm, M.A., and Moler, C.B. 1977, Computer Methods for Mathematical Computations (Englewood Cliffs, NJ: Prentice-Hall), §§4.4–4.5.

Stoer, J., and Bulirsch, R. 1980, *Introduction to Numerical Analysis* (New York: Springer-Verlag), §2.4.

Ralston, A., and Rabinowitz, P. 1978, *A First Course in Numerical Analysis*, 2nd ed. (New York: McGraw-Hill), §3.8.

3.4 How to Search an Ordered Table

Suppose that you have decided to use some particular interpolation scheme, such as fourth-order polynomial interpolation, to compute a function f(x) from a set of tabulated x_i 's and f_i 's. Then you will need a fast way of finding your place in the table of x_i 's, given some particular value x at which the function evaluation is desired. This problem is not properly one of numerical analysis, but it occurs so often in practice that it would be negligent of us to ignore it.

Formally, the problem is this: Given an array of abscissas xx(j), j=1, 2, ..., n, with the elements either monotonically increasing or monotonically decreasing, and given a number x, find an integer j such that x lies between xx(j) and xx(j+1).

For this task, let us define fictitious array elements xx(0) and xx(n+1) equal to plus or minus infinity (in whichever order is consistent with the monotonicity of the table). Then j will always be between 0 and n, inclusive; a returned value of 0 indicates "off-scale" at one end of the table, n indicates off-scale at the other end.

In most cases, when all is said and done, it is hard to do better than *bisection*, which will find the right place in the table in about $\log_2 n$ tries. We already did use bisection in the spline evaluation routine splint of the preceding section, so you might glance back at that. Standing by itself, a bisection routine looks like this:

```
SUBROUTINE locate(xx,n,x,j)
INTEGER j,n
REAL x, xx(n)
    Given an array xx(1:n), and given a value x, returns a value j such that x is between
    xx(j) and xx(j+1). xx(1:n) must be monotonic, either increasing or decreasing. j=0
    or j=n is returned to indicate that x is out of range.
INTEGER jl,jm,ju
j1=0
                                Initialize lower
ju=n+1
                                and upper limits.
if(ju-jl.gt.1)then
                                If we are not yet done,
     jm=(ju+j1)/2
                                compute a midpoint,
     if((xx(n).ge.xx(1)).eqv.(x.ge.xx(jm)))then\\
                                and replace either the lower limit
         jl=jm
     else
                                or the upper limit, as appropriate.
         ju=jm
     endif
goto 10
                                Repeat until
                                the test condition 10 is satisfied.
endif
if(x.eq.xx(1))then
                                Then set the output
     j=1
 else if(x.eq.xx(n))then
     j=n-1
 else
     j=j1
endif
return
                                and return.
END
```

Note the use of the logical equality relation .eqv., which is true when its two logical operands are either both true or both false. This relation allows the routine to work for both monotonically increasing and monotonically decreasing orders of xx(1:n).

Search with Correlated Values

Sometimes you will be in the situation of searching a large table many times, and with nearly identical abscissas on consecutive searches. For example, you may be generating a function that is used on the right-hand side of a differential equation: Most differential-equation integrators, as we shall see in Chapter 16, call for right-hand side evaluations at points that hop back and forth a bit, but whose trend moves slowly in the direction of the integration.

In such cases it is wasteful to do a full bisection, *ab initio*, on each call. The following routine instead starts with a guessed position in the table. It first "hunts," either up or down, in increments of 1, then 2, then 4, etc., until the desired value is bracketed. Second, it then bisects in the bracketed interval. At worst, this routine is about a factor of 2 slower than locate above (if the hunt phase expands to include the whole table). At best, it can be a factor of $\log_2 n$ faster than locate, if the desired point is usually quite close to the input guess. Figure 3.4.1 compares the two routines.

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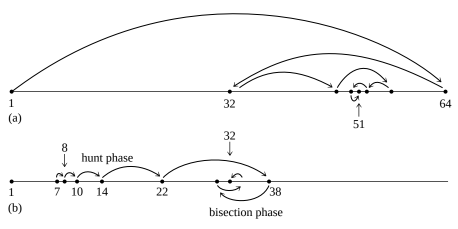


Figure 3.4.1. (a) The routine locate finds a table entry by bisection. Shown here is the sequence of steps that converge to element 51 in a table of length 64. (b) The routine hunt searches from a previous known position in the table by increasing steps, then converges by bisection. Shown here is a particularly unfavorable example, converging to element 32 from element 7. A favorable example would be convergence to an element near 7, such as 9, which would require just three "hops."

```
SUBROUTINE hunt(xx,n,x,jlo)
    INTEGER jlo,n
    REAL x, xx(n)
        Given an array xx(1:n), and given a value x, returns a value jlo such that x is between
        xx(jlo) and xx(jlo+1). xx(1:n) must be monotonic, either increasing or decreasing.
        jlo=0 or jlo=n is returned to indicate that x is out of range. jlo on input is taken as
       the initial guess for jlo on output.
    INTEGER inc, jhi, jm
    LOGICAL ascnd
                                           True if ascending order of table, false otherwise.
    ascnd=xx(n).ge.xx(1)
    if(jlo.le.0.or.jlo.gt.n)then
                                           Input guess not useful. Go immediately to bisection.
        jlo=0
        jhi=n+1
        goto 3
    endif
    inc=1
                                           Set the hunting increment.
    if(x.ge.xx(jlo).eqv.ascnd)then
                                           Hunt up:
        jhi=jlo+inc
        if(jhi.gt.n)then
                                           Done hunting, since off end of table.
            jhi=n+1
        else if(x.ge.xx(jhi).eqv.ascnd)then
                                                      Not done hunting,
            jlo=jhi
                                           so double the increment
            inc=inc+inc
            goto 1
                                           and try again.
        endif
                                           Done hunting, value bracketed.
    else
                                           Hunt down:
        jhi=jlo
2
        jlo=jhi-inc
        if(jlo.lt.1)then
                                           Done hunting, since off end of table.
            jlo=0
        else if(x.lt.xx(jlo).eqv.ascnd)then
                                                      Not done hunting,
             jhi=jlo
                                           so double the increment
            inc=inc+inc
            goto 2
                                           and try again.
        endif
                                           Done hunting, value bracketed.
    endif
                                           Hunt is done, so begin the final bisection phase:
3
    if(jhi-jlo.eq.1)then
        if(x.eq.xx(n))jlo=n-1
```

if(x.eq.xx(1))jlo=1

```
return
endif
jm=(jhi+jlo)/2
if(x.ge.xx(jm).eqv.ascnd)then
    jlo=jm
else
    jhi=jm
endif
goto 3
END
```

After the Hunt

The problem: Routines locate and hunt return an index j such that your desired value lies between table entries xx(j) and xx(j+1), where xx(1:n) is the full length of the table. But, to obtain an m-point interpolated value using a routine like polint (§3.1) or ratint (§3.2), you need to supply much shorter xx and yy arrays, of length m. How do you make the connection?

The solution: Calculate

```
k = min(max(j-(m-1)/2,1),n+1-m)
```

This expression produces the index of the leftmost member of an m-point set of points centered (insofar as possible) between j and j+1, but bounded by 1 at the left and n at the right. FORTRAN then lets you call the interpolation routine with array addresses offset by k, e.g.,

```
call polint(xx(k),yy(k),m,...)
```

CITED REFERENCES AND FURTHER READING:

Knuth, D.E. 1973, Sorting and Searching, vol. 3 of The Art of Computer Programming (Reading, MA: Addison-Wesley), §6.2.1.

3.5 Coefficients of the Interpolating Polynomial

Occasionally you may wish to know not the value of the interpolating polynomial that passes through a (small!) number of points, but the coefficients of that polynomial. A valid use of the coefficients might be, for example, to compute simultaneous interpolated values of the function and of several of its derivatives (see $\S 5.3$), or to convolve a segment of the tabulated function with some other function, where the moments of that other function (i.e., its convolution with powers of x) are known analytically.

However, please be certain that the coefficients are what you need. Generally the coefficients of the interpolating polynomial can be determined much less accurately than its value at a desired abscissa. Therefore it is not a good idea to determine the coefficients only for use in calculating interpolating values. Values thus calculated will not pass exactly through the tabulated points, for example, while values computed by the routines in $\S3.1-\S3.3$ will pass exactly through such points.

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Also, you should not mistake the interpolating polynomial (and its coefficients) for its cousin, the *best fit* polynomial through a data set. Fitting is a *smoothing* process, since the number of fitted coefficients is typically much less than the number of data points. Therefore, fitted coefficients can be accurately and stably determined even in the presence of statistical errors in the tabulated values. (See §14.8.) Interpolation, where the number of coefficients and number of tabulated points are equal, takes the tabulated values as perfect. If they in fact contain statistical errors, these can be magnified into oscillations of the interpolating polynomial in between the tabulated points.

As before, we take the tabulated points to be $y_i \equiv y(x_i)$. If the interpolating polynomial is written as

$$y = c_1 + c_2 x + c_3 x^2 + \dots + c_N x^{N-1}$$
(3.5.1)

then the c_i 's are required to satisfy the linear equation

$$\begin{bmatrix} 1 & x_1 & x_1^2 & \cdots & x_1^{N-1} \\ 1 & x_2 & x_2^2 & \cdots & x_2^{N-1} \\ \vdots & \vdots & \vdots & & \vdots \\ 1 & x_N & x_N^2 & \cdots & x_N^{N-1} \end{bmatrix} \cdot \begin{bmatrix} c_1 \\ c_2 \\ \vdots \\ c_N \end{bmatrix} = \begin{bmatrix} y_1 \\ y_2 \\ \vdots \\ y_N \end{bmatrix}$$
(3.5.2)

This is a *Vandermonde matrix*, as described in §2.8. One could in principle solve equation (3.5.2) by standard techniques for linear equations generally (§2.3); however the special method that was derived in §2.8 is more efficient by a large factor, of order N, so it is much better.

Remember that Vandermonde systems can be quite ill-conditioned. In such a case, *no* numerical method is going to give a very accurate answer. Such cases do not, please note, imply any difficulty in finding interpolated *values* by the methods of §3.1, but only difficulty in finding *coefficients*.

Like the routine in §2.8, the following is due to G.B. Rybicki.

```
SUBROUTINE polcoe(x,y,n,cof)
INTEGER n, NMAX
REAL cof(n), x(n), y(n)
                                      Largest anticipated value of n.
PARAMETER (NMAX=15)
   Given arrays x(1:n) and y(1:n) containing a tabulated function y_i = f(x_i), this routine
   returns an array of coefficients cof(1:n), such that y_i = \sum_j \text{cof}_j x_i^{j-1}.
INTEGER i, j, k
REAL b,ff,phi,s(NMAX)
do 11 i=1,n
    s(i)=0.
    cof(i)=0.
enddo 11
s(n) = -x(1)
do 13 i=2,n
                                      Coefficients s_i of the master polynomial P(x) are found
    do 12 j=n+1-i,n-1
                                          by recurrence.
         s(j)=s(j)-x(i)*s(j+1)
    s(n)=s(n)-x(i)
enddo 13
do 16 j=1,n
    phi=n
                                      The quantity \mathtt{phi} = \prod_{j \neq k} (x_j - x_k) is found as a deriva-
    do 14 k=n-1,1,-1
                                          tive of P(x_j).
```

```
\begin{array}{c} \text{phi=k*s(k+1)+x(j)*phi}\\ \text{enddo }_{14}\\ \text{ff=y(j)/phi}\\ \text{b=1.}\\ \text{do }_{15}\text{ k=n,1,-1}\\ \text{cof(k)=cof(k)+b*ff}\\ \text{b=s(k)+x(j)*b}\\ \text{enddo }_{15}\\ \text{enddo }_{15}\\ \text{enddo }_{16}\\ \text{return}\\ \text{END} \end{array} \\ \begin{array}{c} \text{Coefficients of polynomials in each term of the Lagrange}\\ \text{formula are found by synthetic division of } P(x) \text{ by}\\ (x-x_j). \text{ The solution } c_k \text{ is accumulated.}\\ \text{expression}\\ \text{the solution } c_k \text{ is accumulated.}\\ \text{the
```

Another Method

END

Another technique is to make use of the function value interpolation routine already given (polint $\S 3.1$). If we interpolate (or extrapolate) to find the value of the interpolating polynomial at x=0, then this value will evidently be c_1 . Now we can subtract c_1 from the y_i 's and divide each by its corresponding x_i . Throwing out one point (the one with smallest x_i is a good candidate), we can repeat the procedure to find c_2 , and so on.

It is not instantly obvious that this procedure is stable, but we have generally found it to be somewhat *more* stable than the routine immediately preceding. This method is of order N^3 , while the preceding one was of order N^2 . You will find, however, that neither works very well for large N, because of the intrinsic ill-condition of the Vandermonde problem. In single precision, N up to 8 or 10 is satisfactory; about double this in double precision.

```
SUBROUTINE polcof(xa,ya,n,cof)
INTEGER n, NMAX
REAL cof(n), xa(n), ya(n)
PARAMETER (NMAX=15)
                                                     Largest anticipated value of n.
USES polint
    Given arrays xa(1:n) and ya(1:n) of length n containing a tabulated function ya_i =
    f(xa_i), this routine returns an array of coefficients cof (1:n), also of length n, such that
\mathtt{ya}_i = \sum_j \mathtt{cof}_j \mathtt{xa}_i^{j-1} INTEGER i,j,k
REAL dy, xmin, x(NMAX), y(NMAX)
do 11 j=1,n
    x(j)=xa(j)
    y(j)=ya(j)
enddo 11
do 14 i=1.n
    call polint(x,y,n+1-j,0.,cof(j),dy)
                                                     This is the polynomial interpolation rou-
    xmin=1.e38
                                                         tine of §3.1. We extrapolate to x =
    k=0
    do 12 i=1,n+1-j
                                                     Find the remaining x_i of smallest abso-
         if (abs(x(i)).lt.xmin)then
                                                         lute value.
             xmin=abs(x(i))
         endif
         if(x(i).ne.0.)y(i)=(y(i)-cof(j))/x(i)
                                                         (meanwhile reducing all the terms)
                                                         and eliminate it.
    do 13 i=k+1,n+1-j
         y(i-1)=y(i)
         x(i-1)=x(i)
    enddo 13
enddo 14
return
```

If the point x=0 is not in (or at least close to) the range of the tabulated x_i 's, then the coefficients of the interpolating polynomial will in general become very large. However, the real "information content" of the coefficients is in small differences from the "translation-induced" large values. This is one cause of ill-conditioning, resulting in loss of significance and poorly determined coefficients. You should consider redefining the origin of the problem, to put x=0 in a sensible place.

Another pathology is that, if too high a degree of interpolation is attempted on a smooth function, the interpolating polynomial will attempt to use its high-degree coefficients, in combinations with large and almost precisely canceling combinations, to match the tabulated values down to the last possible epsilon of accuracy. This effect is the same as the intrinsic tendency of the interpolating polynomial values to oscillate (wildly) between its constrained points, and would be present even if the machine's floating precision were infinitely good. The above routines polcoe and polcof have slightly different sensitivities to the pathologies that can occur.

Are you still quite certain that using the *coefficients* is a good idea?

CITED REFERENCES AND FURTHER READING:

Isaacson, E., and Keller, H.B. 1966, Analysis of Numerical Methods (New York: Wiley), §5.2.

3.6 Interpolation in Two or More Dimensions

In multidimensional interpolation, we seek an estimate of $y(x_1, x_2, \ldots, x_n)$ from an n-dimensional grid of tabulated values y and n one-dimensional vectors giving the tabulated values of each of the independent variables x_1, x_2, \ldots, x_n . We will not here consider the problem of interpolating on a mesh that is not Cartesian, i.e., has tabulated function values at "random" points in n-dimensional space rather than at the vertices of a rectangular array. For clarity, we will consider explicitly only the case of two dimensions, the cases of three or more dimensions being analogous in every way.

In two dimensions, we imagine that we are given a matrix of functional values ya(j,k), where j varies from 1 to m, and k varies from 1 to n. We are also given an array x1a of length m, and an array x2a of length n. The relation of these input quantities to an underlying function $y(x_1,x_2)$ is

$$ya(j,k) = y(x1a(j), x2a(k))$$
 (3.6.1)

We want to estimate, by interpolation, the function y at some untabulated point (x_1, x_2) .

An important concept is that of the *grid square* in which the point (x_1, x_2) falls, that is, the four tabulated points that surround the desired interior point. For convenience, we will number these points from 1 to 4, counterclockwise starting from the lower left (see Figure 3.6.1). More precisely, if

$$x1a(j) \le x_1 \le x1a(j+1)$$

 $x2a(k) \le x_2 \le x2a(k+1)$ (3.6.2)

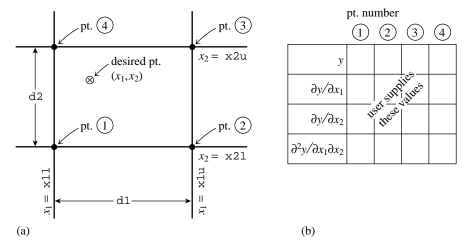


Figure 3.6.1. (a) Labeling of points used in the two-dimensional interpolation routines bcuint and bcucof. (b) For each of the four points in (a), the user supplies one function value, two first derivatives, and one cross-derivative, a total of 16 numbers.

defines j and k, then

$$y_1 \equiv ya(j,k)$$

 $y_2 \equiv ya(j+1,k)$
 $y_3 \equiv ya(j+1,k+1)$
 $y_4 \equiv ya(j,k+1)$ (3.6.3)

The simplest interpolation in two dimensions is *bilinear interpolation* on the grid square. Its formulas are:

$$t \equiv (x_1 - x1a(j))/(x1a(j+1) - x1a(j))$$

$$u \equiv (x_2 - x2a(k))/(x2a(k+1) - x2a(k))$$
(3.6.4)

(so that t and u each lie between 0 and 1), and

$$y(x_1, x_2) = (1 - t)(1 - u)y_1 + t(1 - u)y_2 + tuy_3 + (1 - t)uy_4$$
 (3.6.5)

Bilinear interpolation is frequently "close enough for government work." As the interpolating point wanders from grid square to grid square, the interpolated function value changes continuously. However, the gradient of the interpolated function changes discontinuously at the boundaries of each grid square.

There are two distinctly different directions that one can take in going beyond bilinear interpolation to higher-order methods: One can use higher order to obtain increased accuracy for the interpolated function (for sufficiently smooth functions!), without necessarily trying to fix up the continuity of the gradient and higher derivatives. Or, one can make use of higher order to enforce smoothness of some of these derivatives as the interpolating point crosses grid-square boundaries. We will now consider each of these two directions in turn.

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Higher Order for Accuracy

The basic idea is to break up the problem into a succession of one-dimensional interpolations. If we want to do m-1 order interpolation in the x_1 direction, and n-1 order in the x_2 direction, we first locate an m \times n sub-block of the tabulated function matrix that contains our desired point (x_1, x_2) . We then do m one-dimensional interpolations in the x_2 direction, i.e., on the rows of the sub-block, to get function values at the points $(x1a(j), x_2)$, $j = 1, \ldots, m$. Finally, we do a last interpolation in the x_1 direction to get the answer. If we use the polynomial interpolation routine polint of $\S 3.1$, and a sub-block which is presumed to be already located (and copied into an m by n array ya), the procedure looks like this:

```
SUBROUTINE polin2(x1a,x2a,ya,m,n,x1,x2,y,dy)
INTEGER m,n,NMAX,MMAX
REAL dy,x1,x2,y,x1a(m),x2a(n),ya(m,n)
PARAMETER (NMAX=20,MMAX=20)
                                                  Maximum expected values of n and m.
USES polint
   Given arrays x1a(1:m) and x2a(1:n) of independent variables, and an m by n array of
   function values ya(1:m,1:n), tabulated at the grid points defined by x1a and x2a; and
   given values x1 and x2 of the independent variables; this routine returns an interpolated
   function value y, and an accuracy indication dy (based only on the interpolation in the x1
   direction, however).
INTEGER j,k
REAL ymtmp(MMAX), yntmp(NMAX)
do 12 j=1,m
                                                  Loop over rows.
    do 11 k=1,n
                                                  Copy the row into temporary storage.
        yntmp(k)=ya(j,k)
    enddo 11
    call polint(x2a,yntmp,n,x2,ymtmp(j),dy)
                                                  Interpolate answer into temporary stor-
                                                     age.
                                                  Do the final interpolation.
call polint(x1a,ymtmp,m,x1,y,dy)
return
END
```

Higher Order for Smoothness: Bicubic Interpolation

We will give two methods that are in common use, and which are themselves not unrelated. The first is usually called *bicubic interpolation*.

Bicubic interpolation requires the user to specify at each grid point not just the function $y(x_1,x_2)$, but also the gradients $\partial y/\partial x_1 \equiv y_{,1}$, $\partial y/\partial x_2 \equiv y_{,2}$ and the cross derivative $\partial^2 y/\partial x_1\partial x_2 \equiv y_{,12}$. Then an interpolating function that is *cubic* in the scaled coordinates t and u (equation 3.6.4) can be found, with the following properties: (i) The values of the function and the specified derivatives are reproduced exactly on the grid points, and (ii) the values of the function and the specified derivatives change continuously as the interpolating point crosses from one grid square to another.

It is important to understand that nothing in the equations of bicubic interpolation requires you to specify the extra derivatives *correctly!* The smoothness properties are tautologically "forced," and have nothing to do with the "accuracy" of the specified derivatives. It is a separate problem for you to decide how to obtain the values that are specified. The better you do, the more *accurate* the interpolation will be. But it will be *smooth* no matter what you do.

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Best of all is to know the derivatives analytically, or to be able to compute them accurately by numerical means, at the grid points. Next best is to determine them by numerical differencing from the functional values already tabulated on the grid. The relevant code would be something like this (using centered differencing):

```
\begin{array}{l} y1a(j,k) = & (ya(j+1,k) - ya(j-1,k)) / (x1a(j+1) - x1a(j-1)) \\ y2a(j,k) = & (ya(j,k+1) - ya(j,k-1)) / (x2a(k+1) - x2a(k-1)) \\ y12a(j,k) = & (ya(j+1,k+1) - ya(j+1,k-1) - ya(j-1,k+1) + ya(j-1,k-1)) \\ & / ((x1a(j+1) - x1a(j-1)) * (x2a(k+1) - x2a(k-1))) \end{array}
```

To do a bicubic interpolation within a grid square, given the function y and the derivatives y1, y2, y12 at each of the four corners of the square, there are two steps: First obtain the sixteen quantities c_{ij} , $i,j=1,\ldots,4$ using the routine bcucof below. (The formulas that obtain the c's from the function and derivative values are just a complicated linear transformation, with coefficients which, having been determined once in the mists of numerical history, can be tabulated and forgotten.) Next, substitute the c's into any or all of the following bicubic formulas for function and derivatives, as desired:

$$y(x_1, x_2) = \sum_{i=1}^{4} \sum_{j=1}^{4} c_{ij} t^{i-1} u^{j-1}$$

$$y_{,1}(x_1, x_2) = \sum_{i=1}^{4} \sum_{j=1}^{4} (i-1)c_{ij} t^{i-2} u^{j-1} (dt/dx_1)$$

$$y_{,2}(x_1, x_2) = \sum_{i=1}^{4} \sum_{j=1}^{4} (j-1)c_{ij} t^{i-1} u^{j-2} (du/dx_2)$$

$$y_{,12}(x_1, x_2) = \sum_{i=1}^{4} \sum_{j=1}^{4} (i-1)(j-1)c_{ij} t^{i-2} u^{j-2} (dt/dx_1) (du/dx_2)$$
(3.6.6)

where t and u are again given by equation (3.6.4).

```
SUBROUTINE bcucof(y,y1,y2,y12,d1,d2,c) REAL d1,d2,c(4,4),y(4),y1(4),y12(4),y2(4)
```

Given arrays y, y1, y2, and y12, each of length 4, containing the function, gradients, and cross derivative at the four grid points of a rectangular grid cell (numbered counterclockwise from the lower left), and given d1 and d2, the length of the grid cell in the 1- and 2-directions, this routine returns the table c(1:4,1:4) that is used by routine bcuint for bicubic interpolation.

C

END

```
x(i+4)=y1(i)*d1
    x(i+8)=y2(i)*d2
    x(i+12)=y12(i)*d1d2
do 13 i=1.16
                               Matrix multiply by the stored table.
    xx=0.
    do 12 k=1.16
        xx=xx+wt(i,k)*x(k)
    enddo 12
    cl(i)=xx
enddo 13
1=0
                               Unpack the result into the output table.
do 15 i=1,4
    do_{14} j=1,4
        1=1+1
        c(i,j)=cl(1)
    enddo 14
enddo 15
return
END
```

The implementation of equation (3.6.6), which performs a bicubic interpolation, returns the interpolated function value and the two gradient values, and uses the above routine bcucof, is simply:

```
SUBROUTINE bcuint(y,y1,y2,y12,x11,x1u,x21,x2u,x1,x2,ansy,
     ansy1, ansy2)
REAL ansy, ansy1, ansy2, x1, x11, x1u, x2, x21, x2u, y(4), y1(4),
    y12(4),y2(4)
USES bcucof
   Bicubic interpolation within a grid square. Input quantities are y, y1, y2, y12 (as described
   in bcucof); x11 and x1u, the lower and upper coordinates of the grid square in the 1-
   direction; x21 and x2u likewise for the 2-direction; and x1,x2, the coordinates of the
   desired point for the interpolation. The interpolated function value is returned as ansy,
   and the interpolated gradient values as ansy1 and ansy2. This routine calls bcucof.
INTEGER i
REAL t,u,c(4,4)
call bcucof(y,y1,y2,y12,x1u-x11,x2u-x21,c)
                                                     Get the c's.
if(x1u.eq.x11.or.x2u.eq.x21)pause 'bad input in bcuint'
                                                     Equation (3.6.4).
t=(x1-x11)/(x1u-x11)
u=(x2-x21)/(x2u-x21)
ansy=0.
ansy2=0
ansv1=0.
                                                     Equation (3.6.6).
do = i=4,1,-1
    ansy=t*ansy+((c(i,4)*u+c(i,3))*u+c(i,2))*u+c(i,1)
    ansy2=t*ansy2+(3.*c(i,4)*u+2.*c(i,3))*u+c(i,2)
    ansy1=u*ansy1+(3.*c(4,i)*t+2.*c(3,i))*t+c(2,i)
ansy1=ansy1/(x1u-x11)
ansy2=ansy2/(x2u-x21)
return
```

Higher Order for Smoothness: Bicubic Spline

The other common technique for obtaining smoothness in two-dimensional interpolation is the *bicubic spline*. Actually, this is equivalent to a special case

of bicubic interpolation: The interpolating function is of the same functional form as equation (3.6.6); the values of the derivatives at the grid points are, however, determined "globally" by one-dimensional splines. However, bicubic splines are usually implemented in a form that looks rather different from the above bicubic interpolation routines, instead looking much closer in form to the routine polin2 above: To interpolate one functional value, one performs m onedimensional splines across the rows of the table, followed by one additional one-dimensional spline down the newly created column. It is a matter of taste (and trade-off between time and memory) as to how much of this process one wants to precompute and store. Instead of precomputing and storing all the derivative information (as in bicubic interpolation), spline users typically precompute and store only one auxiliary table, of second derivatives in one direction only. Then one need only do spline evaluations (not constructions) for the m row splines; one must still do a construction and an evaluation for the final column spline. (Recall that a spline construction is a process of order N, while a spline evaluation is only of order $\log N$ — and that is just to find the place in the table!)

Here is a routine to precompute the auxiliary second-derivative table:

```
SUBROUTINE splie2(x1a,x2a,ya,m,n,y2a)
INTEGER m,n,NN
REAL x1a(m), x2a(n), y2a(m,n), ya(m,n)
PARAMETER (NN=100)
                                                   Maximum expected value of n and m.
USES spline
   Given an m by n tabulated function ya(1:m,1:n), and tabulated independent variables
   x2a(1:n), this routine constructs one-dimensional natural cubic splines of the rows of ya
    and returns the second-derivatives in the array y2a(1:m,1:n). (The array x1a is included
   in the argument list merely for consistency with routine splin2.)
INTEGER j,k
REAL y2tmp(NN),ytmp(NN)
do 13 j=1,m
    do 11 k=1,n
        ytmp(k)=ya(j,k)
    enddo 11
                                                      Values 1 \times 10^{30} signal a natural spline.
    call spline(x2a,ytmp,n,1.e30,1.e30,y2tmp)
    do 12 k=1.n
        y2a(j,k)=y2tmp(k)
    enddo 12
enddo 13
return
END
```

After the above routine has been executed once, any number of bicubic spline interpolations can be performed by successive calls of the following routine:

```
SUBROUTINE splin2(x1a,x2a,ya,y2a,m,n,x1,x2,y)
INTEGER m,n,NN
REAL x1,x2,y,x1a(m),x2a(n),y2a(m,n),ya(m,n)
PARAMETER (NN=100) Maximum expected value of n and m.

C USES spline,splint
Given x1a, x2a, ya, m, n as described in splie2 and y2a as produced by that routine; and given a desired interpolating point x1,x2; this routine returns an interpolated function value y by bicubic spline interpolation.

INTEGER j,k
REAL y2tmp(NN),ytmp(NN),yytmp(NN)
```

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```
Perform m evaluations of the row splines
do 12 j=1,m
                                                      constructed by splie2, using the one-
    do 11 k=1,n
        ytmp(k)=ya(j,k)
                                                      dimensional spline evaluator splint.
        y2tmp(k)=y2a(j,k)
    enddo 11
    call splint(x2a,ytmp,y2tmp,n,x2,yytmp(j))
enddo 12
call spline(x1a,yytmp,m,1.e30,1.e30,y2tmp)
                                                  Construct the one-dimensional column spline
call splint(x1a,yytmp,y2tmp,m,x1,y)
                                                      and evaluate it.
return
END
```

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