Curve Fitting with a Digital Computer b_y C. W. Clenshaw

high-speed computers, uses orthogonal polynomials to overcome the problems of ill-conditioning which are usually associated with this approach. The present paper shows how this powerful method can be modified to save a substantial proportion of the machine storage. This is achieved by representing each polynomial within the machine by the coefficients in its Chebyshev series: The method, designed to exploit the advantages of Summary: Forsythe (1957) has described a method for fitting polynomials to a set of points, using the principle of least squares.

TRODUCTION

Curve fitting is essentially the process of finding a smooth curve which passes near to each of a number of prescribed points in a plane. In a numerical, as opposed to graphical, approach to this problem, it is customary to use a polynomial to provide the curve; polynomials are easy to evaluate, their unknown coefficients occur linearly, and their degree affords a convenient measure of smoothness. The nearness is then usually achieved by imposing the "least-squares" criterion. By this means the original vague requirement is converted into a definite and tractable problem.

The numerical solution of this problem involves a considerable amount of arithmetic in all but the most trivial cases. The classical method fits a number of function values by a polynomial in the form

$$k_0 + k_1 x + k_2 x^2 + \ldots + k_n x^n$$
,

using the least-squares criterion. It has the grave disadvantage of requiring the solution of a set of simultaneous equations for the coefficients *k* that may be ill-conditioned, often severely so when *n* is large. This defect can be removed by using orthogonal polynomials. A method, designed for desk machines, which makes some use of such polynomials has been described by Hayes and Vickers (1951). More recently, a method using essentially the same polynomials, but prepared for use with a digital computer, was discussed by Forsythe (1957) and again by Ascher and Forsythe (1958).

The main purpose of the present paper is to show how this powerful method can be modified to save a substantial proportion of the machine storage.

After a brief outline of Forsythe's proposals, we describe the modified method in detail. Next there is a discussion of the factors which help us decide upon the "best" degree for the approximating polynomial, followed by a description of a curve-fitting program, based on the modified method, which has been prepared for the DEUCE at the National Physical Laboratory. Finally, we show how the method is simplified in cases when the function value is available at *all* points of the range.

FORSYTHE'S METHOD

Let y_r ($r = 0, 1, 2, \ldots m$) be the observed or computed values of a dependent variable y at given values x_r

of the independent variable x. Then the polynomial $Y_i(x)$ of degree i which minimizes the residual sum of squares

$$\delta_I^2 = \sum_{r=0}^m \{Y_I(x_r) - y_r\}^2 \tag{1}$$

may be obtained by truncating the series

$$c_0p_0(x) + c_1p_1(x) + c_2p_2(x) + \cdots$$
 (2)

after (i + 1) terms. Here $p_i(x)$ is a polynomial of degree i satisfying the orthogonality condition

$$\sum_{r} p_{i}(x_{r}) p_{j}(x_{r}) = 0 \quad (i \neq j).$$
 (3)

The coefficients c_j in (2) are therefore given by

$$c_j = \sum_{r} y_r p_j(x_r) / \sum_{r} \{p_j(x_r)\}^2.$$
 (4)

The $p_i(x)$ may be computed successively from the following three-term recurrence formula, as suggested previously by Householder (1953), Stiefel (1955), and Lanczos (1957):

$$p_{i-1}(x) = \lambda_i(x - \alpha_{i-1})p_i(x) - \beta_i p_{i-1}(x).$$
 (5)

Here λ_i determines the normalization of the polynomials, $eta_0=0$ and

$$\alpha_{l+1} = \frac{\sum_{x} x_{t}^{l} \{p_{t}(x_{r})\}^{2}}{\sum_{x} \{p_{t}(x_{r})\}^{2}}, \quad \beta_{l} = \frac{\lambda_{l} \sum_{i} \{p_{t}(x_{r})\}^{2}}{\sum_{i} \{p_{l-1}(x_{r})\}^{2}}.$$
 (6)

Forsythe, following the previous writers, chooses $\lambda_i = 1$, so that the coefficient of x^i in the expression for $p_i(x)$ is independent of i, and in fact is unity.

The generation of the polynomials by equation (5), and the computation of the coefficients c_i in

$$Y_i(x) = c_0 p_0(x) + c_1 p_1(x) + \dots + c_i p_i(x),$$
 (7)

give the approximating polynomial of degree i, and we can allow i to take all values up to m, at which stage the approximating polynomial passes through all the (m+1) points (x_i, v_i) .

(m+1) points (x_r, y_r) . The polynomial of "best" fit may be conceived as that which most effectively compromises between smoothness, as represented by the degree of the polynomial, and

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closeness to the data, measured by δ_i^2 in equation (1). For practical purposes Forsythe regards the "best" value of i as that at which the mean square of residuals, defined as $\delta_i^2/(m-i)$, ceases to decrease significantly. This criterion is easy to apply: we have only to examine this quantity for each value of i as the computation proceeds; when its decrease is regarded as insignificant, the desired solution has been reached. Moreover, the quantity δ_i^2 can conveniently be produced as a byproduct of the main calculations. For, from (1), (3), (4), and (7), we find

$$\delta_{i}^{2} = \sum_{r} \{Y_{i-1}(x_{r}) + c_{i}p_{i}(x_{r}) - y_{r}\}^{2}$$

$$= \delta_{i-1}^{2} - 2c_{i} \sum_{r} p_{i}(x_{r})y_{r} + c_{i}^{2} \sum_{r} \{p_{i}(x_{r})\}^{2}$$

$$= \delta_{i-1}^{2} - c_{i}^{2} \sum_{r} \{p_{i}(x_{r})\}^{2}.$$
(6)

In considering the programming of this method, Forsythe suggests that the storage of the polynomials $p_i(x)$ should be effected by the retention of their numerical values at every x_r . Since the $p_i(x)$ are generated by a three-term recurrence relation, we need the values of two such polynomials at any given stage of the calculation. The data x_r and y_r are, of course, also needed throughout the calculation for all r points, so that a main store of 4(m+1) positions is required. We next consider how this requirement might be reduced.

THE MODIFIED METHOD

Essentially the modification consists of a more compact storage procedure. An obvious way to achieve economy of storage would be to store the coefficients of the powers of x in the explicit expression for $p_i(x)$, rather than the values of $p_i(x)$ at every x. Similarly, each $Y_i(x)$ could be stored, and punched out, in the same form. However, in arranging these polynomials as nower

However, in arranging these polynomials as power series, we run the risk of introducing large coefficients, with a possible loss of accuracy when the polynomial is evaluated. This risk becomes serious as *i* increases.

Similar economy can be gained, however, without incurring this risk, by representing each polynomial within the machine by the coefficients in its Chebyshev expansion. When we require, for a given x, the numerical value of a function f(x) which is represented by its Chebyshev series

$$f(x) = \frac{1}{2}a_0 + a_1T_1(x) + a_2T_2(x) + \ldots + a_nT_n(x), (9)$$

where $T_s(x) = \cos(s \cos^{-1}x)$ is the Chebyshev polynomial in x of degree s, we may use the method of recurrence given by Clenshaw (1955). Briefly, this entails evaluating successively the quantities b_n , b_{n-1} , ... b_0 , where

$$b_s = 2xb_{s+1} - b_{s+2} + a_s$$
, with $b_{n+1} = b_{n+2} = 0$. (10)

The required value is then given by

$$f(x) = \frac{1}{2}(b_0 - b_2)$$

as can be verified by substituting for the a_s in (9) their

expressions in terms of the b_s from (10), and using the recurrence relation

$$T_{s+1}(x) - 2xT_s(x) + T_{s-1}(x) = 0.$$

It is assumed here, without loss of generality, that the x_r all lie in the range (-1, 1).

In order to be able to calculate $p_i(x)$ at any point, we accordingly store the coefficients $P_j^{(i)}$ in the expression

$$p_i(x) = \frac{1}{2}P_0^{(i)} + P_1^{(i)}T_1(x) + P_2^{(i)}T_2(x) + \dots + P_{i-1}^{(i)}T_{i-1}(x) + T_i(x), (i > 0).$$
 (11)

The polynomials $p_i(x)$ have been normalized by making the coefficient of $T_i(x)$ in (11), namely $P_i^{(i)}$, equal to unity. The definition of the polynomials is completed by putting $p_0(x) = \frac{1}{2}$, and we set $\lambda_i = 2$ in equations (5) and (6).

The number of storage positions required to represent $p_i(x)$ has thus been reduced from (m+1) to i, so that for $m \geqslant i$, the main storage requirement has been nearly halved.

Substituting (11) in the equation (5) with $\lambda_i = 2$ and comparing coefficients of $T_j(x)$, we obtain

$$P_j^{(i+1)} = P_{j+1}^{(i)} + P_{j-11}^{(i)} - 2\alpha_{i-1}P_j^{(i)} - \beta_i P_j^{(i-1)}, \quad (12)$$

where $P_i^{(i)}=1$ for $i\geqslant 0$. Equation (12) is valid for all i and j.

From these equations we calculate the coefficients in the Chebyshev series for $p_{i+1}(x)$ from those for $p_i(x)$ and $p_{i+1}(x)$.

Similarly, the approximating polynomial $Y_i(x)$ can be represented within the machine by the coefficients $A_j^{(i)}$, say, in its Chebyshev series

$$Y_i(x) = \frac{1}{2}A_0^{(i)} + A_1^{(i)}T_1(x)$$

$$+ A_i^{(i)}T_2(x) + \ldots + A_i^{(i)}T_i(x).$$
 (13)

As each coefficient c_i in the expression (7) is found from (4), the coefficients $A^{(i)}$ are obtained from their predecessors by means of the relation

$$A_j^{(i)} = A_j^{(i-1)} + c_i P_j^{(i)}, \tag{14}$$

which may be derived by comparing the coefficients of $T_{f}(x)$ in the equation

$$Y_i(x) = Y_{i-1}(x) + c_i p_i(x). \tag{15}$$

While the modification which we have introduced certainly saves much storage, a loss in speed may sometimes result, since the computation of $p_i(x)$ from its Chebyshev series for each x_r would take longer than the extraction of each $p_i(x)$ from its store. On the other hand, the modified method requires the application of the recurrence relation (12) only (i + 2) times to define $p_i(x)$, whereas in the original method of Forsythe, equation (5) is used (m + 1) times. The final decision as to which method is faster will depend upon the number of data points, the highest degree of polynomial required, and the characteristics of the computer; in particular the original method is faster whenever 4(m + 1) locations are available in the immediate-access store.

CHOICE OF "BEST" DEGREE

coefficients $A_i^{(0)}$, which can be punched out at the conclusion of each *i*-cycle. These coefficients provide another indication of the "best" value of i. If we were observational errors, which affect the behaviour of the $A_j^{(i)}$. For values of j exceeding a certain value k, say, the coefficients will fluctuate about zero in an apparently random manner, and as i increases this behaviour persists difference between $Y_k(x)$ and each of its immediate neighbours $Y_{k-1}(x)$ and $Y_{k-1}(x)$ is then small for all x in the range, and any one of these three polynomials with little change in k. The polynomial $Y_k(x)$ may then be accepted as the desired solution. Although it may sometimes be difficult to alight confidently on a definite k, the choice in such a case is not critical, since the output of the program consists of the function, we would expect the $A_i^{(i)}$ to decrease with dealing with exact values of a well-behaved mathematical increasing j, for fixed large values of i. In practice, however, the readings y, invariably contain rounding and The main

all be changed during every *i*-cycle. When the "best" degree *k* has been passed, however, the changes in the $A_j^{(i)}$ will merely be of the order of the unwanted "noise" present in the data, and the smallness of may be regarded as a satisfactory solution.

The values of all the coefficients $A_j^{(l)}$ will, of course, these changes is a most valuable check on the arithmetic.

punched out, together with the corresponding values of Another criterion for the choice of k depends on the sively laborious; in practice it is often sufficient to inspect the extreme values. In the DEUCE program Physical Laboratory, the largest positive residual P_i and the numerically largest negative residual N_i are behaviour of the successive sets of residuals $Y_i(x_r) - y_r$. Examination of the complete set would usually be excescompiled in the Mathematics Division of the National

x, ξ_i and η_i , say. Like δ_i^2 , the numerically larger of P_i and N_i will decrease appreciably as i increases until k is reached, after which it will usually decrease only slowly.

Therefore if ξ_i (or η_i) remains unchanged for several values of *i*, it may be desirable to examine the reading at The quantities P_i and N_i serve another and more useful purpose, however. If one reading y, has an outstandingly large error, there will be a tendency for P_i (or N_i) to this point, or to repeat the calculation with this reading occur at the corresponding x_i , for different values i. omitted.

more rapidly if $y_r - Y_{i-1}(x_r)$ were stored in place. This replacement would not affect the calculation It may be observed that P_i and N_i would be deterof c_i , since $\sum Y_{i-1}(x_r) p_i(x_r) = 0$. mined more rapidly if $y_r - Y_t$

DETAILED DESCRIPTION OF PROGRAM

To clarify the procedure, we now consider the arith-Let us suppose that at the metic involved in one cycle.

beginning of the ith cycle the machine store holds the following quantities:

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- (A) x_r and y_r for $r = 0, 1, 2, \dots m$. (B) $P_j^{(i+1)}$ for $j = 0, 1, 2, \dots i$; $P_j^{(i)}$ for $j = 0, 1, 2, \dots (i-1)$. (C) $A_j^{(i)}$ for $j = 0, 1, 2, \dots i$. (D) P_{i-1} , N_{i-1} , ξ_i , η_i , and δ_i .
- (D) P'_{i-1} , N_{i-1} , ξ_{i-1} , η_{i-1} and δ_i^2 . (E) Other numbers, not required for output.

(2i + 1) and (i + 1) positions respectively, while (D) and (E) each need only a few positions. The contents of (C) and (D) comprise the output at the end of the Of these stores, (A), (B) and (C) must have 2(m+1), previous cycle.

The first step is the calculation of those quantities which depend directly on the initial data. Each pair (cf. (10)), using the coefficients stored in (B). Then $\{p_{i-1}(x_r)\}^2$ is formed and added to the partial sum $\sum \{p_{i-1}(x_r)\}^2$ stored in (E). Similarly the sums $\sum x_r\{p_{i-1}(x_r)\}^2$ and $\sum y_rp_{i-1}(x_r)$ are accumulated. In conjunction with this computation it is convenient to calfor each x_r . If this exceeds numerically the largest so far obtained in this cycle we send it to the P_i (or N_i) position, and store the corresponding value of ξ_i (or η_i). (x_r, y_r) is extracted in turn from store (A), and $p_{i+1}(x_r)$ culate the function $Y_i(x_r)$, whose Chebyshev coefficients are in store (C), and thence the residual $\{Y_i(x_r) - y_r\}$ calculated by summation of its Chebyshev

culate α_{l+2} and β_{l+1} from (6) with $\lambda_{l+1} = 2$, c_{l+1} from (4) and hence δ_{l+1}^2 from (8). When the complete sums have been formed, we cal-

which overwrite the contents of store (C). Likewise equation (12) enables us to compute the $P_j^{(i+2)}$, which can replace the $P_j^{(i+1)}$ after the latter have been transferred to the $P_j^{(i)}$ position. The cycle is now complete; we can punch out the contents of stores (C) and (D), Now we can evaluate from (14) the coefficients and start a new cycle.

The main program is entered at the beginning of the above cycle, with i = -1. The stores (C), (D) and (E) above cycle, with i = -1. The stores (C), (D) and (E) then contain zeros, while (B) holds only $P_0^{(0)} = 1$. For this first cycle only, we can arrange to omit that part of the program which evaluates the residuals $\{Y_i(x_r) - y_r\}$,

and to insert the value $\beta_0=0$. If m is less than 30, the program may be allowed to find the best least-squares polynomial of degree m. The corresponding P_m and N_m would be zero if there were no rounding errors, so their actual size affords a valuable check on the build-up of such errors.

n general, the results required from a specific problem quently with an auxiliary program. A few such programs will cover most of the common requirements, and might at equal intervals of x, and one which rearranges $Y_k(x)$ will not be simply the Chebyshev coefficients $A_j^{(i)}$, and residuals δ_i for a given k, one to form a table of $Y_k(x)$ into a power series (where such rearrangement is possible it will usually be necessary to operate on these subsefor example, one which calculates all without loss of accuracy).

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SPEED AND STORAGE

punch all the least-squares polynomials $Y_0(x)$, $Y_1(x)$, $Y_1(x)$, using (m+1) points, for various values of mately. The table shows the number of minutes taken to read in the program and data, and calculate and out, we present in Table 1 examples of the time taken and a multiplication and division time of 2 msec approxiin order to give an indication of the rate of operation of the program, which uses fixed-point arithmetic throughby DEUCE, which has an addition time of 0.062 msec,

TABLE 1

| 300 | 31 5 7 10 10 14 25 39 | 28 |
|-------|--|----|
| 100 | 11 2 3 4 4 5 9 15 15 | 22 |
| 20 | 2 | |
| _ m/: | 20 10 10 10 20 | 25 |

application has so far been restricted to data with m not exceeding 100; the greatest value of i required has been 12. The numbers in Table 1 which refer to larger The DEUCE has a high-speed store of 400 words and a magnetic-drum store of 8,192 words, each word consisting of 32 binary digits. The program described above allows for values of (m+1) up to 3,840, and for obtained from data specially Its practical polynomials of degree not exceeding 29. prepared to test the present program. problems have been

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are given at the points $x_r = \cos \pi r/m$, as has been shown by Lanczos (N.B.S., 1952). We can make direct use of The general procedure may be simplified when the y, this solution in problems where we are given a formula or graph from which the value of y, corresponding to any chosen x, may be readily obtained.

We first suppose y to be a polynomial of degree less than or equal to m, given by

$$y = \frac{1}{2}a_0 + a_1T_1(x) + a_2T_2(x) + \dots$$

$$+ a_{m-1}T_{m-1}(x) + \frac{1}{2}a_mT_m(x)$$
 (16)

$$=\sum_{i=0}^{m} a_i T_i(x)$$
, say,

Then, as Lanczos shows, the where Σ'' denotes a summation in which the first and coefficients in (16) are given by last terms are halved.

$$a_i = \frac{2}{m} \sum_{r=0}^{m} y_r \cos \frac{\pi i r}{m}.$$
 (17)

nomial; but if it is continuous and of bounded variation, its expansion in Chebyshev polynomials converges, often we can approximate the function to any desired degree of accuracy by a polynomial of the form (16), and find very rapidly. By choosing a sufficiently large value of m, of course, the function y is not a polyits coefficients from (17). In general,

The solution of this problem is a special case of that $x_r = \cos \frac{m}{m}$, the polynomials $p_l(x)$ become the Chebyshev of the general problem considered earlier, since polynomials $T_i(x)$.

CONCLUSION

Considerable economy of storage space and a more concise primary output have been achieved by using Charachaev expansions in programming Forsythe's having ample high-speed storage, when polynomials of particularly on machines whose high-speed store cannot accommodate all the data. It is possible, however, that Forsythe's original method may be faster on machines method of curve fitting. Some time may also be saved, in programming high degree are required. Chebyshev expansions

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