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VOLUME 2 - MAINTENANCE MANUAL

A COMPUTER PROGRAM FOR
POWER SPECTRAL ANALYSIS
OF UNEQUALLY SPACED POINTS

By

Matthew Lybanon

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ABSTRACT

The computer program whose mathematical method is derived in Volume 1 is discussed in some detail. The discussion and tables cover the contents of COMMON blocks, major calculations, subroutine argument lists, and program logic. Appendix 1 gives a listing of the program system, and Appendix 2 is an overall logic flowchart.

INTRODUCTION

This Maintenance Manual gives detailed information about the power spectral density computer program whose mathematical method and general form are discussed in Volume 1 of this report, the User's Manual. All equation numbers given in this volume refer to equations in the User's Manual.

This manual contains discussions of the functions of individual program elements, and information on the contents of COMMON blocks and argument lists. Appendix 1 is a listing of the entire program. Appendix 2, for convenience, shows the same overall logic flowchart contained in Appendix 1 of the User's Manual.

The entire program is written in FORTRAN V for the UNIVAC 1108. Each subprogram's source and relocatable elements, the MAP source element, and the absolute element, are in file LSQFIT*MLTSAN. in Bellcomm's computer system. The name of the absolute element is LSQFIT*MLTSAN.APSD.

In the subsequent discussion, sequence numbers refer to the program listings in Appendix 1. Equation numbers are those of equations in Volume 1, the User's Manual.

PROGRAM DESCRIPTION

Contents of COMMON Blocks

Blank COMMON:

PSPECT (1000) - array of spectral powers
Y (2000) - array of observations (d. p.)*
T (2000) - array of times (independent variable) (d. p.)
NAME (3) - identifying name of data
A (1000) - array of sine coefficients (d. p.)
B (1000) - array of cosine coefficients (d. p.)
PERCNT (1000) - array of percent powers

NOPEAK:

KNO (30) - array of subscripts of false peaks
NOPEAK - number of false peaks found since last removal
MOPEAK - number of false peaks from latest search

MORN:

NORM - norm of inverse times matrix, with 1 subtracted
from diagonal elements (d. p.)

CUCPSA

Important Quantities Calculated

<u>Quantity</u>	<u>Sequence Number</u>	<u>Equation</u>	<u>Comments</u>
OMEGAZ	135	56	$2\pi f$ (d. p.)
KMAX	140	57	Index of highest frequency term
A(KK)	185-186	47	Sine coefficients (d. p.)
B(KK)	187-188	46	Cosine coefficients (d. p.)
PSPECT(KK)	192-194	54	Spectral power
PERCNT(KK)	195		Percent power
Y(I)	408-409		Prewhitened signal (d. p.)
YBAR	418		Mean of data (or residuals) (d. p.)
YYBAR2	424-425		Total power (variance) (d. p.)

* "d. p." denotes a double precision variable.

Subroutine Calls

<u>Subroutine</u>	<u>Arguments</u>	<u>Comments</u>
PRNTCN		System routine - interface to print-page controls.
	2, 'M, 66, 5, 3.'	Changes margins to give 66-line page with 5-line top and 3-line bottom margins.
READ		Tape read subroutine for input data.
	N	Number of points.
	IY	Position of observation in record.
	IT	Position of time in record.
	TPC	Time scale factor.
RPLOT		Plots spectrum on one page.
	ITAPE	Scratch unit containing plot information.
GRAPH		Plots expanded spectrum.
	PERCNT	Array of percent powers.
	PSPECT	Array of spectral powers.
	KLIMIT	N/2 (N=number of points).
	NAME	Identifying name of data.
	HEAD3A	'P VS K'
	PRD	Fundamental period of data.
	RS	Product of range and spacing factors.
CAROL3		Solves for dominant terms.
	AREMOV	Array of sine coefficients (output) (d. p.)
	BREMOV	Array of cosine coefficients (output) (d. p.)
	PREMOV	Array of spectral powers (output)
	OREMOV	Array of angular frequencies (output) (d. p.)
	N	Number of points.
	KREMOV	Array of indices of peaks.
	OMEGAZ	Fundamental angular frequency ($2\pi f$) (d. p.)
	NTERMS	Number of peaks found in search.

CUCPSA is the main program in the system. Its main functions are to calculate the power spectrum, search the spectrum for peaks, call the subroutine which improves the estimates of the peaks, remove the peaks, and calculate the spectrum of the residuals (prewhitened signal). It also takes care of input and output.

Sequence numbers 144-197 are the portion of the program which calculates the coefficients, spectral powers, and percent powers. The search for peaks is handled by 322-384. The call to CAROL3 (which improves the estimates of coefficients and frequencies of peaks) is sequence number 394. The removal is done by 400-411. After each removal, the spectrum of the residuals is calculated by the same section of coding which calculates the original spectrum.

Sequence numbers 31-34, 88-105, 207-225, and 293-313 are the spectrum-smoothing option described in the User's Manual. These statements are currently in the program as comments. To activate this feature, the "C" in column 1 of each FORTRAN statement should be removed.

The spectrum search section searches the unsmoothed spectrum for peaks. However, there is also a provision for smoothing the spectrum as the search for peaks is made. To activate this feature the "C" in column 1 of each FORTRAN statement in sequence numbers 333-344 must be removed, and sequence numbers 345-351 should be deleted.

There are two flags (other than the print, plot, and removal flags) which control the flow of logic through the program. IFLAG is set equal to 0 at the beginning of each case. Its value is changed to 1 the first time CAROL3 successfully solves for peaks. There are a number of branch points at which the value of IFLAG determines what is done next. The logical variable LFLG is given the value .TRUE. at the beginning of the first search for peaks. When no more peaks can be found, TOL is cut in half and LFLG becomes .FALSE., which prevents TOL from being diminished again.

A modification which would permit a "magnified" view of one or more portions of the spectrum might be useful. This feature could be implemented in addition to or in place of the usual spectrum calculation; it could be triggered by input or by logic based on the result of some calculation. For example, it might be

helpful to see the fine structure of regions of the spectrum where the search-and-removal process has trouble.

The variables KTOTAL, FMESH, and FBEGIN (sequence numbers 144-146) lend themselves to this application. For example, setting KTOTAL=101, FMESH=.02, FBEGIN=39, would cause the spectrum calculation to begin at k=39 and proceed to k=41, bracketing k=40 in 100 steps. If this modification is not made, the program could be optimized slightly by eliminating these variables and rewriting the beginning of the spectrum calculation.

CAROL3

Important Quantities Calculated

<u>Quantity</u>	<u>Sequence Number</u>	<u>Equation</u>	<u>Comments</u>
AREMOV(M)	130	55	a_k (d. p.)
BREMOV(M)	131		b_k (d. p.)
OREMOV(M)	132		$2\pi f_k$ (d. p.)
PREMOV(M)	142	54	Powers

Subroutine Calls

<u>Subroutine</u>	<u>Arguments</u>	<u>Comments</u>
BIORTH	AMTRX	Matrix inversion routine
	AI	Matrix to be inverted (d. p.)
	NT3	(Left) inverse of AMTRX (d. p.)
		Rank of AMTRX

CAROL3 is the subroutine which calculates the frequencies and coefficients of dominant terms in the spectrum. It does so by an iterative differential correction least squares technique. Such a method fits a model $\hat{\phi}(p_1, \dots, p_m)$

to an observed variable ϕ by adjusting the parameters $p_1, \dots, p_m$ to minimize the mean square error. The values of the parameters are adjusted iteratively according to:

$$p^{(k)} = p^{(k-1)} + \Delta p^{(k)}$$

$$\Delta p^{(k)} = (A^T W A)^{-1} A^T W \left[\phi - \hat{\phi}(p^{(k-1)}) \right]$$

where p denotes the vector whose components are $p_1, \dots, p_m$, A is the matrix with elements $\frac{\partial \hat{\phi}_i(p)}{\partial p_j}$, $i=1, \dots, N$ (the number of observations) and $j=1, \dots, m$ (the number of parameters), W is a (diagonal) weighting matrix, and superscripts in parentheses label the stage of iteration. (In calculating $\Delta p^{(k)}$, A must be evaluated using $p^{(k-1)}$.)

In CAROL3 the parameters which are adjusted are the amplitudes, phases, and frequencies of spectral peaks. (This was found to lead to faster convergence than the two coefficients and frequency formulation of equation (55).)

Frequency is the most critical parameter, so it is handled in perturbation form. The f_k of (55) is represented by $(k+\alpha)f$, where f is defined by (56), kf is the frequency found in the spectrum search, and α , $|\alpha| \leq .5$, is the quantity solved for.

The parameters p_j , $j=1, \dots, m$ (NT3 stands for m in the program) are: amplitude, phase, α , amplitude, phase, α , $\dots$, one group of three parameters for each spectral peak. The identity matrix is used as the weighting matrix W in CAROL3. The square matrix $A^T A$ is called AMTRX;

$A^T \left[\phi - \hat{\phi}(p^{(k-1)}) \right]$ is called BMTRX. They are set up by sequence

numbers 56-67. Sequence number 69 is the call to BIORTH, which computes the inverse of AMTRX. The vector $p^{(k-1)}$ is called AKOLD, $\Delta p^{(k)}$ is called DA, and $p^{(k)}$ is called AKNEW. The section of coding which does the differential corrections is sequence numbers 72-77.

It is assumed that if the frequency corrections are too large, something is wrong (e. g. perhaps an apparent peak is only a sidelobe of a large peak). So the values of AKNEW(3), AKNEW(6), ... - that is, α for each peak - are checked. If any of them exceed 0.5, the corresponding peaks are considered "bad" peaks. They are "tagged" and a return is made to the main program to search the spectrum for the MTERMS largest peaks, not including the tagged ones, which exceed TOL. This may happen several times before CAROL3 converges on a solution. After a successful solution (and removal) the tags are eliminated from the bad peaks - which may or may not be peaks in the spectrum of the residuals.

This process is done by keeping track of the k-values of the bad peaks. KNO is the array of bad k-values. NOPEAK is the total number of bad peaks found. MOPEAK counts the number found on the latest call to CAROL3. Before the return to the main program, MOPEAK is set equal to the subscript (of KNO) which indexes the first bad peak found in the current call to CAROL3. This portion of the program consists of sequence numbers 78-89.

Convergence is checked by observing the change in the sum of squares of the residuals (data-fit). The current value of the sum of squares is called DELTYN (calculated in sequence numbers 96-106); the value from the preceding iteration is called DELTYO. The quantity checked is $TESTIT = DELTYO / DELTYN - 1.0$. The differential correction process is permitted to go through three iterations before TESTIT is checked. Then it is checked after each iteration. $TESTIT < 0$ implies divergence. In this event there is a return to the main program with

the values of the parameters from the preceding iteration chosen as the "best" values. If the number of iterations exceeds NITER (set equal to 50 in a DATA statement) before convergence, an error message is printed and execution is terminated. $TESTIT < EPSLN$ (set equal to 10^{-8} in a DATA statement) is the condition for convergence. Sequence numbers 109-119 are the convergence check calculations.

The coefficients and frequencies found in the coarse search are converted to the fit parameters (amplitudes, phases, frequency correction factors) in sequence numbers 26-33. After convergence, the transformation back to coefficients and frequencies, along with the computation of spectral powers, are handled by 128-143.

Other Subroutines

The remaining subroutines in the system are straightforward. The following notes illuminate the principal features of the programs:

READ - The data tape is assumed to be a binary tape consisting of N (the number of points) records, each containing 7 (or more) single-precision words. One of the words is time, identified by IT; there is no day variable. TPC is the time scale factor. IY labels the data word. The tape is assumed to be on logical unit 3.

RPLOT - Automatic scale selection is done. The entire plot appears on one printer page, with each scale running from 0 to the maximum value of power or k. The scaling information is computed in CUCPSA and passed to RPLOT, along with the other plotting information, on scratch unit ITAPE. ITAPE (set equal to 19 in CUCPSA) is assigned by the operating system; no ASG statement is needed.

GRAPH - Automatic scale selection is done for the percent power scale. Three scales are available, with maxima of 20%, 40%, or 100%.

BIORTH - This subroutine is internally documented by means of embedded comments. It should be noted that BIORTH calls two internal subroutines, DOT and SCLR.

PROGRAM FILE

The entire program is contained on FASTRAND file LSQFIT*MLTSAN. The name of the MAP source element is LSQFIT*MLTSAN.PSAMAP. The absolute element is LSQFIT*MLTSAN.APSD.

The file has a write key, JULIE. Therefore, the control card:

@ ASG,A LSQFIT*MLTSAN//JULIE.

must be used if it is necessary to write on the file. Tape number 1015 (permanent) is a save tape of LSQFIT*MLTSAN.

APPENDIX I

Program Listing

LSOFT*MLTSAN.CHIPSA

POWER SPECTRUM ANALYSIS PROGRAM

DOUBLE PRECISION T,Y,VYBAR2,VBAR2,A,B,SIMYS,SIMYC,SIMSC2,

SIMC2,SIMSC,DELTA,XSIN,XCOS,DEMOV(10),DEMOV(10),

TWOPI,DUR,OMEGA7,OMEGA,STEP,THETA,DEMOV(10),

COMMON PSPECT(1000),Y(2000),T(2000),TIME(3),

A(1000),R(1000),PERCENT(1000)

COMMON /NDBEAK/ KNO(20),NDBEAK,NDBEAK

LOGICAL LFLG

DATA XEND/4HEMD/,VTITLE/6H POWER/,HTITLE/6H K /,

IVERT/17,THOR7777,TSNAME/27,HEAD3A/6HP VS K/

DATA IRLANK /6H /

EQUIVALENCE (FMT(1),IEMT1)

DIMENSION FMT(14)

DOUBLE PRECISION V(3)

COMMON KREMOV(10),DEMOV(10),DEMOV(10),DEMOV(10)

CALL PRNTCN (2,M,66,5,3.1)

CONTINUE

100 ICOUNT = 0

START A NEW CASE.

READ DATA SET AND INITIALIZE CONSTANTS

IRLAGE=0

FILTER(1) = 0.25

FILTER(2) = 0.50

FILTER(3) = 0.25

LMAX = 3

TWOPI = 6.283185307600

ITAPE = 10

NTAPE=0

FMT IS FORMAT FOR READING OBSERVATION CARDS

IT IS OBSERVATION FIELD NUMBER (ON OBSERVATION CARDS)

IT IS TIME FIELD NUMBER

IN IS DAY FIELD NUMBER

TDC IS TIME UNITS PER COMPUTATIONAL UNIT (SEE TIME CALC.)

CON IS COMPUTATIONAL UNITS PER DAY (SEE TIME CALC.)

READ (5:1000) FMT,IT,IN,TDC,CON

1005 FORMAT (13A6,A2/15,2E10.5)

IF (IT.EQ. -999) STOP

IF (IEMT1.EQ. IRLANK)

NTAPE = 1

NAME IDENTIFIES DATA

N IS NUMBER OF OBSERVATIONS (LIMITED TO 2000)

S IS SPACING CONSTANT FOR OMEGA7

R IS RANGE CONSTANT FOR KMAX

IDA = 0 PLOTS DOLPH (AND SMOOTH) POWER SPECTRUM

```

57 C = 1 PLOTS ROUGH POWER SPECTRUM ONLY
58 C = 2 (PLOTS SMOOTH POWER SPECTRUM ONLY)
59 C = 3 NO PLOT
60 C JF = 0 REMOVE FREQUENCIES WHOSE PERCENT POWERS EXCEED TOL
61 C = 1 DO NOT REMOVE DOMINANT FREQUENCIES
62 C TOL IS REMOVAL TOLERANCE DESCRIBED ABOVE
63 C JS = 0 PRINT SPECTRUM
64 C = 1 DO NOT PRINT SPECTRUM
65 C
66 C READ (5,1000) NAME,N,S,R,JPO,JE,TOL,JIC
67 C IF (NTAPE.EQ.1) CALL READ (N,IY,IT,IPC)
68 C IF (N.GT.2000) N = 2000
69 C ** TOL MAY BE EITHER AN INPUT OR A CALCULATED QUANTITY.
70 C IF (TOL.LF.0)
71 C . TOL = 10. / FLOAT(N)
72 C . TOL = 10. / FLOAT(N)
73 C . TOL = 10. / FLOAT(N)
74 C IF (N.GT.2000) N = 2000
75 C IF (N.GT.2000) NAME,N,S,R,JPO,JE,TOL,JS,MTERMS
76 C 1000 FORMAT (3A6,15,2F10.5,2I1,5F10.5,11)
77 C 1001 FORMAT (15)
78 C 2000 FORMAT (1H1/ 5X,13HSET NAME = 3A6
79 C /5X,13HNO. POINTS = 15
80 C /5X,13HSPACING = 513.0
81 C /5X,13HRANGE = 513.0
82 C /5X,13HPOINT FLAG = 15
83 C /5X,13HREMOVAL = 15
84 C /5X,13HTOLERANCE = 510.5
85 C /5X,13HPOINT FLAG = 15
86 C /5X,13HMTERMS = 15)
87 C IF (NTAPE.EQ.1) GO TO 4321
88 C
89 C IF (S.EQ.1.0) GO TO 110
90 C READ 1001,LEND,(FILTER(I),I=1,LEND)
91 C 1001 FORMAT (15,7F10.5)
92 C LMAX = 2 * LEND - 1
93 C LEND
94 C LMAX
95 C 105 FILTER(LM) = FILTER(LF)
96 C LF = LF-1
97 C LM = LM-1
98 C IF (LE.GT.0) GO TO 105
99 C LY = LY
100 C LY = LY + 1
101 C LM = LM-1
102 C IF (LM.GT.0) GO TO 107
103 C 110 PRINT 2001,FILTER(I),I=1,LMAX)
104 C 2001 FORMAT ( 5X,12HFILTER = 7F11.5/17X,5F11.5 )
105 C LCENTP = (LMAX+1)/2
106 C
107 C READ FIRST OBSERVATION CARD
108 C READ (5,FMT) V
109 C
110 C COMPUTE INITIAL VALUES
111 C T(1) = V(TT)/TDC
112 C Y(1) = V(TV)
113 C NAYO = V(TO)

```

```

114      DO 130 I=2,N
115      READ (5,FMT) V
116      Y(I) = V(I*V)
117      C
118      T(I) = V(I*V)/TDC + (V(I*V)-D*Y(I))*CDM
119      130 CONTINUE
120      4321 CONTINUE
121      C
122      NUP = T(N) - T(1)
123      WRITE (6,1004) FMT,IY,IT,IO,TDC,CDM,NUP
124      1004 FORMAT ( /5X,13HFORMATT = 13A6,13,
125      1 /5X,13HY FIELD = T5,
126      2 /5X,13HT FIELD = T5,
127      3 /5X,13HDAY FIELD = T5,
128      4 /5X,13HTIME SCALE = 614.0,14H TIME UNITS PER COMPUTATIONAL
129      4L UNIT ,
130      5 /5X,13HDAY SCALE = 614.0,14H COMPUTATIONAL UNITS PER DAY
131      6 / /5X,13HDURATION = F10.4,20H COMPUTATIONAL UNITS )
132      C
133      NUP = S * DUR * FLOAT(N) / FLOAT(N-1)
134      DQR=DUR
135      OMEGAZ = TWOPI/DUR
136      KLIMIT = N/2
137      RS = P*S
138      RSMAX = 2000./FLOAT(N)
139      IF (RS.GT.RSMAX) RS = RSMAX
140      KMAX = RS*FLOAT(N)/2.
141      C
142      GO TO 950
143      C
144      400 KTOTAL = KMAX
145      FMESH = 1.0
146      RBEGIN = 1.
147      C
148      C .....
149      C ..... BEGIN OMEGA LOOP .....
150      C .....
151      C
152      300 OMEGA = OMEGAZ *
153      (RBEGIN - FMESH)
154      STEP = OMEGAZ * FMESH
155      C
156      DO 600 KK=1,KTOTAL
157      C
158      OMEGA = OMEGA + STEP
159      C
160      C ..... SIMULATIONS .....
161      C
162      SIMS2 = 0.0
163      SIMC2 = 0.0
164      SIMYS = 0.0
165      SIMYC = 0.0
166      SIMSC = 0.0
167      C
168      DO 500 I=1,N
169      C
170      THETA = OMEGA*T(I)

```

THIS SECTION CALCULATES THE
COEFFICIENTS, POWER, AND
POWER OF SPECTRAL TERMS.

```

171 XSTH = SIN(THETA)
172 XCOS = COS(THETA)
173 SIMYS = SIMYC + Y(I)*XSTH
174 SIMYC = SIMYC + Y(I)*XCOS
175 SIMS2 = SIMSC2 + XSTH*XSTH
176 SIMC2 = SIMC2 + XCOS*XCOS
177 SIMSC = SIMSC + XSTH*XCOS
178 C
179 C 500 CONTINUE
180 C
181 DELTA = SIMS2+SIMC2 - SIMSC*SIMSC
182 C
183 C FORM SOLUTION VECTOR
184 C
185 A(KK) = (SIMC2 * SIMYS - C STIF
186 * SIMYC * SIMSC) / DELTA C COEFFICIENT
187 B(KK) = (SIMS2 * SIMYC - C POSIVE
188 * SIMYS * SIMSC) / DELTA C COEFFICIENT
189 C
190 C FORM POWER SPECTRUM
191 C
192 PSPECT(KK) = 2. * (A(KK) * C POWER
193 * SIMYS + B(KK) * SIMYC) C SPECTRAL
194 / FLOAT(M) C DENSITY
195 PERCENT(KK) = PSPECT(KK)/YYBAR22
196 C
197 C 600 CONTINUE
198 C
199 C .....
200 C .....
201 C ..... FROM OMEGA LOOP .....
202 C .....
203 C .....
204 C .....
205 C COMPUTE SMOOTH POWER SPECTRUM
206 C
207 C 601 DO 700 K=1,KMAX
208 C
209 C PSMOOTH(K) = 0.0
210 C WEIGHT = 0.0
211 C
212 C DO 650 L=1,LMAX
213 C
214 C LK = K+L-IFENR
215 C IF (LK,LE,1) .OR. (LK,GE,KMAX) ) GO TO 620
216 C PSMOOTH(K) = PSMOOTH(K) + FILTER(L)*PSPECT(LK)
217 C GO TO 650
218 C 620 WEIGHT = WEIGHT + FILTER(L)
219 C IF (LK,GT,KMAX) LK = KMAX
220 C IF (LK,FO,1) .OR. (L,EO,1MAX) ) PSMOOTH(K) = PSMOOTH(K)
221 C + WEIGHT*PSPECT(LK)
222 C 1
223 C 650 CONTINUE
224 C
225 C 700 CONTINUE
226 C
227 C PRINT POWER SPECTRUM

```

```

228 C
229 IF (IFLAG.EQ. 1) C SEARCH FOR MORE PEAKS
230 GO TO 700 C UNLESS FIRST TIME THROUGH.
231 WRITE (6,3000) NAME,YYBAR2
232 3000 FORMAT (1H1,X,30HINITIAL DATA SPECTRUM FOR SET ,3A6,X,
233 1 15H(TOTAL POWER = ,E10.5,1H) /4X,21H-----
234 2 //)
235 IF (JS.EQ.1) GO TO 750
236 GO TO 730
237 C
238 720 IF (LELG) GO TO 800 C DENIVE TOL AND SEARCH AGAIN.
239 IF (IFLAG.NE. 1) C GO TO NEXT CASE IF NO
240 GO TO 100 C PEAKS WERE FOUND.
241 IF (JS.EQ. 1) GO TO 750
242 WRITE (6,3001) NAME,YYBAR2
243 3001 FORMAT (1H1,X,30H FINAL DATA SPECTRUM FOR SET ,3A6,X,
244 1 15H(TOTAL POWER = ,E10.5,1H) /4X,21H-----
245 2 //)
246 C
247 730 IFFD = 0
248 WRITE (6,3002) IFFD,YYBAR
249 IFFD = 1
250 C
251 DO 750 K=1,KMAX
252 OMEGA = OMEGA2 * FLOAT(K)
253 PERD = TWOPI/OMEGA
254 IF (MON(K,52).EQ. 51) WRITE (6,3002) IFFD,YYBAR
255 WRITE (6,3010) K,PERD,OMEGA,A(K),B(K),PSPECT(K),PERCENT(K)
256 750 CONTINUE
257 C
258 3002 FORMAT (11,5X,25H DEPRESSION PERCENT
259 1 5X,5HPERIOD,7X,5HOMEGA,8X,1HA,11X,1HP,11X,5HPOWER,
260 2 7X,5HPWPER,5XWHC = ,F10.5/)
261 3010 FORMAT (2X13,2XF10.4,2VF11.6,2(2XFC10.5),2XG11.6,2XF10.6)
262 C
263 PLOT POWER SPECTRUM
264 C
265 750 PNT = 5.*YYBAR2/FLOAT(KLIMIT)
266 IF ( (JPO.NE.0).AND. (JPO.NE.1) ) GO TO 770
267 VERMAX = 0.0
268 DO 760 K=1,KMAX
269 IF (PSPECT(K).GT.VERMAX) VERMAX=PSPECT(K)
270 760 CONTINUE
271 C
272 VERMAX = KMAX
273 VERHT = VERMAX/50.
274 VERINT = VERMAX/120.
275 BEWIND ITAE
276 WRITE (ITAE) KMAX, VTITLE,VTITLE,NAME,HEAD3A,IVE9T,4H02,
277 1 TSHADE,VERMAX,VERINT,H0PMAX,H0PINT,DNT
278 DO 765 K=1,KMAX
279 XK = K
280 WRITE (ITAE) PSPECT(K),XK
281 765 CONTINUE
282 WRITE (ITAE) VERIN
283 BEWIND ITAE
284 C

```

```
295      CALL PLOT (ITAPE)      C      PLOT SPECTRUM ON ONE PAGE.
296      CALL GRAPH (PERCENT,PERCENT, C      PLOT SPECTRUM ON EXPANDED
297      . KLIMIT,NAME(1),NAME(2), C      GRAPH, ONE SPECTRAL
298      . NAME(3),HEAD3A,PRD,RS) C      TERM PER LINE.
299
300      PLOT SMOOTH POWER SPECTRUM
301
302      CONTINUE
303      C 770 TE ( (JPD,NE,0).AND. (JPD,NE,2) ) GO TO 700
304      C REVIND ITAPE
305      C VERMAX = 0.0
306      C DO 775 K=1,KMAX
307      C TE (PSMOTH(K),GT,VERMAX) VERMAX=PSMOTH(K)
308      C 775 CONTINUE
309      C HOPMAX = KMAX
310      C VERINT = VERMAX/50.
311      C HORINT = HOPMAX/120.
312
313      C ** HEAD3P MUST BE ADDED TO DATA STATEMENT
314      C WRITE (ITAPE) KMAX, VTITLE,UTITLE,NAME,HEAD3P,IVERT,YHODZ,
315      C 1 TSHADF,VERMAX,VERINT,HOPMAX,HORINT,DNT
316      C DO 780 K=1,KMAX
317      C XK = K
318      C WRITE (ITAPE) PSMOTH(K),XK
319      C 780 CONTINUE
320      C WRITE (ITAPE) YEND
321      C REVIND ITAPE
322
323      C CALL PLOT (ITAPE)
324
325      CONTINUE
326
327      C 790 CONTINUE
328
329      C IF (IFLAG.EQ. 1) C      FINAL SPECTRUM HAS BEEN
330      . GO TO 100 C      PLOTTED AND PRINTED.
331
332      C TEST FOR FREQUENCY REMOVAL
333
334      C NOPEAK = 0 C      FALSE PEAK COUNTER
335      C LFLG = .TRUE. C      TOL HAS ORIGINAL VALUE.
336      C IF (JF,NE. 0) GO TO 100
337      C IF (IFLAG.EQ. 0) WRITE (A,3030) NAME,VYGRAB
338      C 709 IF (ICOUNT.GT. KMAX) GO TO 725
339      C ** THIS SECTION FINDS PEAKS IN DESCENDING ORDER OF SIZE.
340      C NITERMS=0
341      C PEAK = VYGRAB
342      C 7095 CONTINUE
343      C KK = 1
344      C OR = TOL
345      C DT2 = .75 * PERCENT(1) + .25 * PERCENT(2)
346      C DT1 = DT2
347      C DO 810 K=1,KMAX C      BEGIN SEARCH FOR PEAK.
348      C ** SMOOTH SPECTRUM AS SEARCH FOR PEAK IS MADE.
349      C K1 = K + 1
350      C K2 = K + 2
351      C IF (K1 - KMAX) 813,812,811 C      COLLAPSE FILTER AT END OF SPECTRUM.
352      C 811 K1 = K
353      C 812 K2 = K1
```



```

399 C
400 DO 840 M=1,NTERMS
401   APEMV(M)=APEMV(M)/TWOPI
402   PDPEMV(M)=1./APEMV(M)
403 C
404 C REMOVE FREQUENCY
405 C
406 TCOUNT = TCOUNT + 1
407 DO 850 I=1,N
408   Y(I) = Y(I) - ( APEMV(M)*SIN(APEMV(M)*T(I))
409     + PDPEMV(M)*COS(APEMV(M)*T(I)) )
410 R50 CONTINUE
411 R40 CONTINUE
412 C
413 C COMPUTE AVERAGE Y
414 C
415 YBAR = 0.0
416 DO 200 I=1,N
417   YBAR = YBAR + Y(I)
418   YBAR = YBAR / FLOAT(N) C MEAN OF DATA
419 YBAR2 = 0.0
420 DO 250 I=1,N
421   Y(I) = Y(I) - YBAR
422   YBAR2 = Y(I) * Y(I) + YBAR2
423 R250 CONTINUE
424 YBAR2 = YBAR2 * 2. / C TOTAL
425   FLOAT(N) C A.C. POWER
426 C ** TOL IS USED AS A FRACTION OF INITIAL TOTAL POWER.
427 IF (IFLAG.FO. 0) TOL=TOL*YBAR2
428 C
429 IF (IFLAG.NE. 1) GO TO 400
430 C
431 C PRINT REMOVAL DATA
432 C
433 NTERMS=NTERMS-1
434 IF (NTERMS.EQ. 0) GO TO 970
435 DO 960 M=1,NTERMS
436   WRITE (6,3031) PDPEMV(M),APEMV(M),APEMV(M),APEMV(M),
437     YBAR,PDPEMV(M)
438 R960 CONTINUE
439 FORMAT (2X,F11.5,2X,F10.6,2X,F10.5,3(2X,F11.6,2X,F11.6))
440 CONTINUE
441 NTERMS=N+1
442 WRITE (6,3031) PDPEMV(M),APEMV(M),APEMV(M),APEMV(M),
443   YBAR,PDPEMV(M),YBAR2
444 C
445 GO TO 800
446 C
447 C ** PUT TOL IN HALF, THEN SEARCH FOR MORE PEAKS.
448 CONTINUE
449 LEIG=.FALSE.
450 TOL = TOL / 2.
451 WRITE (6,3000)
452 FORMAT (4X,SEARCH WILL NOW BE MADE WITH SMALLER TOL.)
453 GO TO 790
454 C
455 END

```

LISTING OF SPECTRUM PROGRAM FOR INEQUALLY SPACED DATA

```

LSQFIT*MLTSAN.READ
1 SUBROUTINE READ (N,TY,TT,TPC)
2 DOUBLE PRECISION T,Y
3 COMMON PSPECT(1000),Y(2000),T(2000)
4 DIMENSION V(7)
5 DO 10 I=1,N
6 READ (3) V
7 T(I) = V(1)/TPC
8 Y(I) = V(2)
9 10 CONTINUE
10 RETURN
11 ENB

```

Q PRT MLTSAN.RPLOT

LISTING OF SPECTRUM PROGRAM FOR LINEARLY SPACED DATA

LSOFIT*ULTSAN.RPLOT

```

1  SUBROUTINE RPLOT (ITAPE)
2  C
3  DIMENSION ALPHA(121),DATA(2,2000),HMAX(120)
4  DATA IEND,ITIMEP,7,ITIMEP,ITIMEP,ASTER,1144/
5  DATA BLANK/14 /,DASH/14-/,DOT/14./
6  C
7  READ PLOT DATA
8  C
9  7 READ (ITAPE) NBRCONS,VTTITLE,HTITLE,HEAD1,HEAD2,HEAD3,HEAD4,IVERT,
10  THORZ,TSIAPDE,VERMAX,VERTINT,HORMAX,HORINT,PNT
11  C
12  DO 11 I=1,NBRCONS
13  11 READ (ITAPE) (DATA(J,I),J=1,2)
14  C
15  DO 20 I=1,120
16  20 ALPHA(I) = BLANK
17  C
18  WRITE (6,055) HEAD1,HEAD2,HEAD3,HEAD4,VTTITLE
19  055 FORMAT (1H1,4X3A6,1X14/1X14)
20  C
21  PMARK = PNT + VERTINT/2.
22  VERMAX = VERMAX + VERTINT
23  DO 150 NM = 1,51
24  VERMAX = VERMAX - VERTINT
25  C
26  DO 100 NI = 1,NBRCONS
27  HMAX = HORMAX - (121. * HORINT)
28  IENDATA(IVERT,NI),DE,VERMAX,AND,DATA(IVERT,NI),LT,VERMAX+VERTINT)
29  1 GO TO 50
30  GO TO 100
31  C
32  50 DO 75 NJ = 1,120
33  HMAX = HMAX + HORINT
34  IENDATA(HORZ,NI),LT,HMAX,DP,DATA(THORZ,NI),DE,HMAX+HORINT)
35  1 GO TO 75
36  ALPHA(NJ) = ASTER
37  GO TO 100
38  C
39  75 CONTINUE
40  100 CONTINUE
41  C
42  80 WRITE (6,060) VERMAX,(ALPHA(I),I=1,120)
43  060 FORMAT (1X67.3,2H I,120A1)
44  C
45  110 IF (TSIAPDE.EQ.1) GO TO 150
46  C
47  STILL = BLANK
48  IF (VERMAX.GE.PMARK .AND. VERMAX.LT.(PMARK+VERTINT) ) STILL = DOT
49  DO 125 I=1,120
50  125 ALPHA(I) = STILL
51  150 CONTINUE
52  C
53  WRITE (6,065) (DASH,I=1,121)
54  065 FORMAT(8X,2H I,121A1 )
55  C
56  C

```

LISTING OF SPECTRUM PROGRAM FOR IMMEDIATELY SAVED DATA

DATE 121560

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```

57      XY7 = 120.
58      GO 210 IF = 1,12
59      XY7 = XY7 - 10.
60      210 ARRAY(I) = HORDMAX - (XY7 * HORDINT)
61      C
62      C
63      WRITE (6,075) (ARRAY(I),I=1,12),HTITLE
64      975 FORMAT (11X,12(9X,14V),11X,12(4X,5A,1)/60Y,A6)
65      C
66      C
67      READ (11,085) KCODE
68      C
69      IF (KCODE .EQ. 1) THEN RETURN
70      IF (KCODE .EQ. 1) THEN GO TO 7
71      END

```

Q PRT MLTSM.GRAPH

LSOFT*WLTSA*.GRAPH

```

1  SUBROUTINE GRAPH (PCNT,PWP,KMAX,TITLE1,TITLE2,TITLE3,TITLE4,POW,R)
2  C
3  DIMENSION IARY(102),PCNT(1),PWP(1)
4  DATA IBLANK/1H 7,IASTER/1H 7,INOT/1H 7/
5  C
6  WRITE (6,1000) TITLE1,TITLE2,TITLE3,TITLE4
7  1000 FORMAT (1H1,1X,3A6,1X,A6,30X,13HPERCENT POWER)
8  C
9  PMAX = 0.
10 KPLOT = FLOAT(KMAX)*R
11 DO 5 I=1,KPLOT
12   IF(PCNT(I).GT.PMAX) PMAX = PCNT(I)
13   5 CONTINUE
14 C
15   ISCALE = 2
16   IF (PMAX.LE. 0.2) ISCALE = 1
17   IF (PMAX.GT.4) ISCALE = 5
18   PMAX = 500./FLOAT(ISCALE)
19   OFFSET = 1.0 + .005*PMAX
20   MARK = ( 5./FLOAT(KMAX)) * PMAX + OFFSET
21   IF (MARK.GT.101) MARK = 102
22 C
23   DO 10 I=1,21
24   10 IARY(I) = (I-1)*ISCALE
25 C
26   WRITE (6,1010) (IARY(I),I=1,21)
27   1010 FORMAT (/26X,21(2X,I3))
28 C
29   200 WRITE (6,2020)
30   2020 FORMAT (5X,5HPPOWER,5X,5HPFREQUENCY,1X,1H4,2X,1H4V,20(4X,1H4V))
31 C
32   1030 WRITE (6,1030)
33   1030 FORMAT (20X,2HI-,20(5H-----) )
34 C
35   DO 20 I=1,102
36   20 IARY(I) = IBLANK
37 C
38   DO 100 I=1,KPLOT
39   C
40   IARY(MARK) = INOT
41   JPOINT = PCNT(I)*PMAX + OFFSET
42   IF (JPOINT.GT.101) JPOINT = 101
43   IARY(JPOINT) = IASTER
44   PLENG=FLOAT(I)/PRN
45 C
46   2000 WRITE (6,2000) PWP(I),PLENG,I,IARY
47   2000 FORMAT (1XG11.7,2XG10.6,14,2H 7,102A1)
48 C
49   100 IARY(JPOINT) = IBLANK
50 C
51   RETURN
52 C
53   END

```

PDP10

PDP10

LSOFIT*MLTSAN.CAROL3

```
1 SUBROUTINE CAROL3 (AREMOV,PREMOV,OREMOV,N,KREMOV,  
2 OMEGAZ,NTERMS)  
3 C ** CALCULATES FREQUENCIES AND COEFFICIENTS OF DOMINANT TERMS IN SPECTRUM  
4 DOUBLE PRECISION AREMOV(NTERMS),PREMOV(NTERMS),OREMOV(NTERMS)  
5 DIMENSION KREMOV(NTERMS),PREMOV(NTERMS),FK(10),XK(10)  
6 DOUBLE PRECISION T,Y,A,R  
7 DOUBLE PRECISION F,DF(30),DY,DELTA,DELTA2,DELTA3,DELTA4  
8 DOUBLE PRECISION THETA,XSIN,XCOS,SUMYS,SUMYC,OMEGAZ  
9 DOUBLE PRECISION AMTRY(30,30),AMTRY(30),AKOLD(30),AKNEW(30),  
10 ATT(30,30)  
11 DOUBLE PRECISION DSIN,DCOS,DSORT,DATAN2,WOT  
12 DOUBLE PRECISION NORM  
13 COMMON /MORN/ NORM  
14 COMMON PSPECT(1000),Y(2000),T(2000),NAME(3),  
15 A(1000),R(1000),PERCENT(1000)  
16 COMMON /NOPEAK/ KNO(30),NOPEAK,NOPEAK  
17 DIMENSION FNO(30)  
18 DATA EPSLN /1.E-04/  
19 DATA NITER /50/  
20 1000 FORMAT (30H1 MAXIMUM NUMBER OF ITERATIONS PERFORMED//  
21 . 16X3 HOLD,15X3HNEW/14X6HVALUES,12X6HVALUES/  
22 . 3H AK,4X2HIR,10X7X2HNR,10H)  
23 1002 FORMAT (7H DELTY ,2019.10//12H RUN ABORTED)  
24 NOPEAK = 0  
25 NT3=3*NTERMS  
26 DO 5 ME1,NTERMS  
27 M3=3*M  
28 KRE=KREMOV(M)  
29 XK(M)=KRM  
30 AKNEW(M3-2)=DSORT(A(KRM)*A(KRM)+R(KRM)*R(KRM))  
31 AKNEW(M3-1)=DATAN2(R(KRM),A(KRM))  
32 AKNEW(M3)=0.00  
33 5 CONTINUE  
34 ITERS=0  
35 GO TO 400  
36 20 CONTINUE  
37 DO 10 J=1,NT3  
38 AMTRY(J)=0.00  
39 DA(J)=0.00  
40 DO 10 I=1,NT3  
41 AMTRY(I,J)=0.00  
42 10 CONTINUE  
43 DO 100 T=1,N  
44 F=0.00  
45 WOT=OMEGAZ*T(I)  
46 DO 50 ME1,NTERMS  
47 M3=3*M  
48 THETA=OREMOV(M)*T(I)+AKOLD(M3-1)  
49 XSIN=DSIN(THETA)  
50 XCOS=DCOS(THETA)  
51 F=F+AKOLD(M3-2)*XSIN  
52 DF(M3-2)=XSIN  
53 DF(M3-1)=XCOS*AKOLD(M3-2)  
54 DF(M3)=DF(M3-1)*WOT  
55 50 CONTINUE  
56 NY=Y(I)-F  
6 NY = RESIDUAL AT T(I)
```

```

57 DO 30 J=1,NT3
58 AMTRX(J)=AMTRX(J)+DE(J)*DY
59 DO 30 K=J,NT3
60 AMTRX(K,J)=AMTRX(K,J)+DE(K)*DE(J)
61 30 CONTINUE
62 100 CONTINUE
63 DO 40 I=2,NT3
64 IJ=I-1
65 DO 40 J=1,IJ
66 AMTRX(J,I)=AMTRX(I,J)
67 40 CONTINUE
68 C
69 CALL RTORTH (AMTRX,AI,NT3) C PUTS LEFT INVERSE OF AMTRX IN AI
70 C
71 C ** CORRECT THE AK'S
72 DO 250 I=1,NT3
73 DO 200 J=1,NT3
74 DA(I)=DA(I)+AI(I,J)*AMTRX(J)
75 200 CONTINUE
76 AKNEW(I)=AKOLD(I)+DA(I)
77 250 CONTINUE
78 DO 325 I=1,NT3,3
79 IF (ABS(AKNEW(I)) .LE. 0.5) C CHECK FREQUENCY CORRECTIONS.
80 . GO TO 325 C CONTINUE PROCESSING TO FREQUENCY
81 MOPEAK = MOPEAK + 1 C CORRECTIONS ARE IN ROUNDS.
82 I3 = I / 3 C COUNT FALSE PEAKS.
83 KNOWNOPEAK = KPEMOV(I3) C SUBSCRIPTS OF FALSE PEAKS
84 FNO(MOPEAK) = FLOAT(KNO(MOPEAK)) * OMEGA2 / 6.2831853076
85 MOPEAK = MOPEAK + 1 C COUNT BAD PEAKS THIS TIME THROUGH.
86 325 CONTINUE
87 IF (MOPEAK .LE. 0) GO TO 400
88 MOPEAK = MOPEAK - MOPEAK + 1
89 RETURN
90 400 CONTINUE
91 DO 25 M=1,NTPEAKS
92 OREMOV(M)=(XK(M)+AKNEW(3*M))*OMEGA2
93 IF (ITER .GT. 0) GO TO 25
94 FK(M)=OREMOV(M)/6.2831853076
95 25 CONTINUE
96 DELTYN=0.00
97 DO 300 I=1,N
98 F=0.00 C CALCULATE DELTY USING CURRENT
99 DO 350 M=1,NTPEAKS C VALUES OF PARAMETERS.
100 M3=3*M
101 THETA=OREMOV(M)*T(I)+AKNEW(M3-1)
102 F=F+AKNEW(M3-2)*DSIN(THETA)
103 350 CONTINUE
104 DY=Y(I)-F
105 DELTYN=DELTYN+DY*DY
106 300 CONTINUE
107 IF (ITER .LT. 3) GO TO 375
108 C ** TEST CONVERGENCE.
109 TESTIT = DELTYN / DELTYN - 1.0
110 IF (TESTIT .LT. 0.1) C CUT OFF IN CASE
111 . GO TO 500 C OF DIVERGENCE.
112 IF (ITER .EQ. NITER) GO TO 475
113 375 CONTINUE

```

LISTING OF SPECTRUM PROGRAM FOR LINEARLY SPACED DATA

```

114 DO 450 I=1,N13
115 AKOLD(I)=AKNEW(I)
116 450 CONTINUE
117 IF (ITER .LT. 3) GO TO 465
118 IF (TESTIT .LT. FPELN) C PUT OFF IF CONVERGENCE
119 . GO TO 500 C IS SUFFICIENTLY SLOW.
120 465 CONTINUE
121 DELTYO=DELTYN
122 ITER=ITER+1
123 GO TO 20
124 475 WRITE (6,1000) (AKOLD(I),AKNEW(I),I=1,N13)
125 WRITE (6,1002) DELTYO,DELTYN
126 STOP
127 500 CONTINUE
128 DO 550 M=1,NTERMS
129 M3=3*M
130 AREMOV(M)=AKOLD(M3-2)*ACOS(AKOLD(M3-1))
131 AREMOV(M)=AKOLD(M3-2)*OSIN(AKOLD(M3-1))
132 OREMOV(M)=(XK(M)+AKOLD(M3))*OMEGA2
133 C
134 C ** COMPUTE POWER.
135 SIMYS=0.00
136 SIMYC=0.00
137 DO 600 I=1,N
138 THETA=OREMOV(M)*I(I)
139 SIMYS=SIMYS+Y(I)*OSIN(THETA)
140 SIMYC=SIMYC+Y(I)*OCOS(THETA)
141 600 CONTINUE
142 AREMOV(M)=2.*(AREMOV(M)*SIMYS+OREMOV(M)*SIMYC)/FLOAT(N)
143 550 CONTINUE
144 RETURN
145 END

```

Q PRT MLTSAN.PIORTH

LSQFIT*WLTSA*RIORTH

```

1  C
2  SUBROUTINE RIORTH(A,R,N)  C
3  C
4  C WHERE: (1) A = AN NBY-N DIMENSIONAL MATRIX
5  C (2) R = THE LEFT INVERSE OF THE MATRIX A
6  C (3) N = ROW- AND COLUMN-DIMENSION OF THE MATRICES
7  C
8  C DOUBLE PRECISION A(30,30),R(30,30),DOT,NORM,NSORT,EPS,CJK
9  C DOUBLE PRECISION NSA
10  C
11  C DATA EPS/1D-9/  C
12  C
13  C COMMON/WORN/NORM  C
14  C
15  C
16  C NITE=0
17  C DO 100 I=1,N
18  C DO 100 J=1,N
19  C R(I,J)=A(I,J)
20  C CONTINUE
21  C
22  C CONTINUE
23  C NITE=NITE+1
24  C DO 300 K=1,N
25  C CALL SCLP(1./DOT(R(1,K),
26  C . A(1,K)),R(I,K),
27  C . R(1,K))
28  C DO 300 J=1,N
29  C IF(XOR(J,K).EQ.0)GO TO 300
30  C CJK=DOT(R(1,J),A(I,K))
31  C DO 301 I=1,N
32  C R(I,J)=R(I,J)-CJK*A(I,K)
33  C
34  C CONTINUE
35  C NORM=0.0
36  C DO 400 I=1,N
37  C DO 400 J=1,N
38  C NORM=NORM+(DOT(R(1,I),
39  C . A(1,J))-DOT(R(1,J),
40  C . J/I))**2
41  C
42  C CONTINUE
43  C NORM=NSORT(NORM)
44  C IF(NORM.GT.((2**NITE)*EPS))
45  C GO TO 200
46  C
47  C DO 500 I=1,N
48  C DO 500 J=1,N
49  C NSA=R(I,J)
50  C R(I,J)=R(I,J)+I
51  C R(J,I)=NSA
52  C CONTINUE
53  C
54  C RETURN
55  C
56  C FUNCTION DOT(Y,Y)

```

C INITIALIZE ITERATION COUNTER
 C AS A 1-ST GUESS, SET THE IN-
 C VERSE MATRIX, *R*, EQUAL TO THE
 C INPUT MATRIX, *A* (THE R-MATRIX IS
 C BEING COMPUTED AS ITS TRANSPOSE)
 C INCREMENT ITERATION COUNTER
 C THERE ARE *N* CYCLES
 C COMPUTE THE K-TH ROW (STORED AS A
 C COLUMN) OF THE INVERSE MATRIX
 C DO FOR THE REMAINING ROWS
 C BUT ONLY IF *J* NOT EQUALS *K*
 C COMPUTE THE SCALAR, C(J,K)
 C DO FOR EACH OF THE *N* COMPONENTS
 C COMPUTE THE (I,J)-TH ELEMENT OF THE
 C INVERSE FOR THE K-TH CYCLE
 C INITIALIZE THE NORM OF THE MATRIX
 C THE *NORM*, OF THE PRODUCT OF THE
 C LEFT-INVERSE TIMES THE MATRIX IS
 C DEFINED AS THE SQUARE-ROOT OF THE
 C SUM OF THE TERMS, WHERE *I* IS SUB-
 C TRACTED FROM THE DIAGONAL TERMS
 C FINALLY, TAKE THE SQUARE-ROOT
 C GO TO AN ADDITIONAL ITERATION IF
 C THE CONVERGENCE CRITERIA WAS NOT
 C MET
 C TRANSPOSE THE R-MATRIX, SINCE IT
 C WAS BEING COMPUTED AS COLUMN VEC-
 C OF ROW VECTORS
 C DOT-PRODUCT FUNCTION

```

57 DOUBLE PRECISION X,Y
58 DIMENSION X(1),Y(1)
59 N=0.0
60 DO 600 K=1,N
61   N=N+X(K)*Y(K)
62 CONTINUE
63 RETURN
64 C
65 SUBROUTINE SCALR(Z,X,Y)
66   DOUBLE PRECISION X,Y,Z
67   DIMENSION X(1),Y(1)
68   DO 700 K=1,N
69     Y(K)=Z*X(K)
70 CONTINUE
71 RETURN
72 END

```

Q FIN

RUNIN: MXL ACCOUNT: 002 PROJECT: LSQFIT

TIME: 00:00:01.381 IN: 10 OUT: 0 PAGES: 20

INITIATION TIME: 18:43:07-DEC 15,1969

TERMINATION TIME: 18:43:20-DEC 15,1969

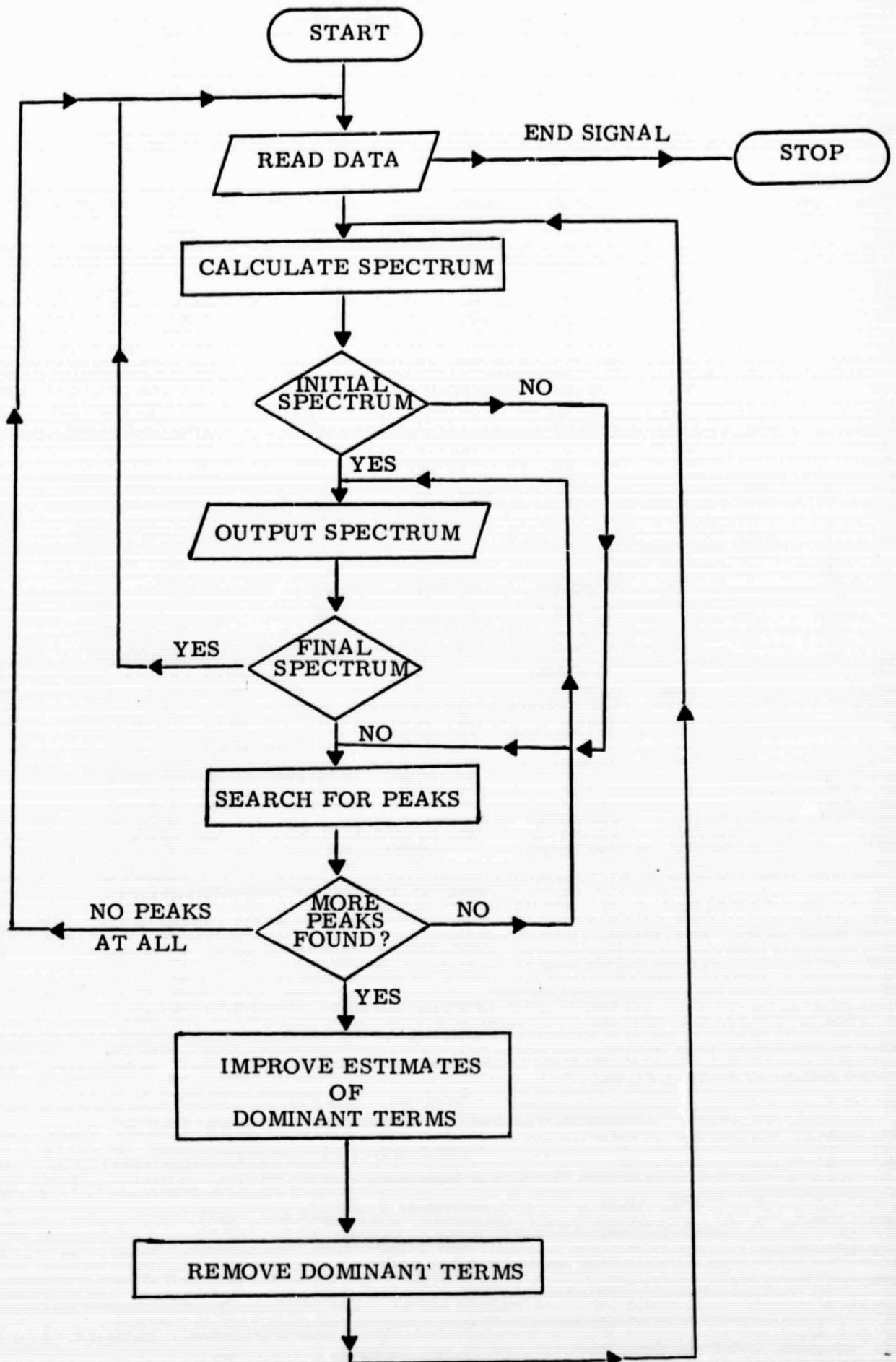
CORE-SECONDS: 9

IC COUNT: 41

CHARGE: 0.331

APPENDIX II

Overall Logic Flowchart



END

DATE

FILMED

JUL 16 1970