



MICROCOPY RESOLUTION TEST CHART
NATIONAL BUREAU OF STANDARDS-1963-A

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K M SCIENCES

FINAL REPORT

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INTRODUCTION

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During a period 13 September 1985 to 31 March 1987, KM Sciences furnished support to the radiation effects programs of the Radiation-Matter Interactions Branch of the Naval Research Laboratory in the areas of collecting, manipulating, analyzing, and displaying experimental data, in writing, modifying, converting, and extending computer codes for modeling physical phenomena and in graphical presentation of experimental data and results of computations. Two major tasks were supported:

(1) computer simulation of the effects of particle beams on materials and (2) experimental measurements of the effects of particle beams on materials.

Keywords: MARPOP (Marlow Post Processor) computer program)

ACCOMPLISHMENTS

COMPUTER SIMULATION:

MARPOP, a computer program (written for the TI-ASC computer) that processes output from the MARLOW cascade simulation code, was converted to run on NRL's Cray computer. A new output routine for the Cray version of MARLOW was written to produce an output data file that serves as input to MARPOP. MARPOP performs calculations and manipulations on the data from MARLOW and produces tables and graphs summarizing and displaying the MARLOW results. A number of production runs were made with the Cray versions of MARLOW and MARPOP using input data designed to simulate experiments in which materials were irradiated with particle beams. In an evolutionary process, the experimental data and the results of the computer simulations led to frequent changes in the MARPOP processing and output specifications, requiring numerous changes in the MARPOP code. The results of these calculations were incorporated in a paper, "Cascade Simulation of the Crystal Orientation Dependence of Sputtering and Lattice Damage of Single Crystal Copper by Irradiation with 100 keV Copper Ions", by G.P. Mueller, M. Rosen, W.A. Fraser (KM Sciences), J.A. Sprague, P.R. Malmberg, J.M. Lambert, P.A. Treado, and G.W. Reynolds, published in Nuclear Instruments and Methods in Physics Research B18(1987)360-364. A source code listing of the most recent version of MARPOP is enclosed.

PARTICLE BEAM EXPERIMENTS:

Computer support was provided for five series of experiments using the NRL Linear Accelerator to study the effects of particle beams on materials. The support

consisted of computer data acquisition, data manipulation and calculations, and graphical and tabular display of results.

OTHER:

A hands-on guide with sample command sequences was developed to introduce users to graphics software installed on the Condensed Matter and Radiation Sciences Division VAX computer. This enabled members of the Radiation-Matter Interactions Branch to produce many types of data plots with less than an hour of self-training using the guide.

Hardware and software specifications were developed for a personal-computer-based pilot system for storage and retrieval of experimental data and results of theoretical calculations of effects of charged particle beams and deposition of energy in materials. After the personal computer systems were delivered, they were set up and software was installed.

PROGRAM MARPOP

VERSION 3.1

22 MAY 1986

Programmer: W. A. FRASER KM Sciences

MARPOP (MARlow P0st-Processor) reads Cray program MARLOW 'POPDAT' output file (data for each particle exiting from the target for zero surface binding energy) and recalculates the yield, energy and directional distributions for chosen surface binding energies.

THIS VERSION OF MARPOP PROCESSES OUTPUT FILES FROM MARLOW VERSION OF 18 (+ OR -) FEBRUARY 1982, ADDING TABLES OF MAXIMUM PARTICLE DEPTH, AND NUMBERS AND LENGTHS OF COLLISION SEQUENCES TO THE OUTPUT THAT WAS PRODUCED BY MARPOP VERSIONS 1.3 AND 1.4. THE "18" FEBRUARY MARLOW VERSION TRUNCATES MAXIMUM DEPTH VALUES TO THE NEXT LOWER 0.1 ALAT(1) UNIT. DATA FROM UP TO 10 MARLOW OUTPUT FILES CAN BE COMBINED. THE NUMBER OF PRIMARY PARTICLES FOR WHICH DATA WILL BE PROCESSED FROM A FILE CAN BE SET TO LESS THAN MAXRUN. SURFACE BINDING ENERGY FOR PRIMARY PARTICLES IS 0.0.

***** Input Parameter Records *****
***** read from logical unit LUSPEC *****

Up to 20 parameter sets may be input. Each set consists of the following:

Record		Format
1	Title - may contain any readable characters (maximum = 80 characters)	A80
2	NFILES Number of MARLOW 'POPDAT' files to be processed (maximum = 10).	10X, I2
3	LIMRUN(NFILES) NFILES values of LIMRUN, the number of primary particles for which data are to be processed from each 'POPDAT' file. The first value on record 3 will be used for file 'FT17', the next for 'FT18', etc. If LIMRUN(i)=0, the corresponding FTnn file will be processed to the end (MAXRUN primaries). If LIMRUN(i)>0, then data from the corresponding FTnn file will be processed until data produced by primary particle LIMRUN+1 are encountered.	10X, I015
4,5	SBND(1-10) Surface binding energies (e.v.) for up to 10 particle types. MARLOWE Version 12 provides for only 5 particle types. Ten types were permitted in Version 11. Both records 4 and 5 must be present (to process MARLOW Version 12 output there will be no values on record 5).	10X, 5E10.0
6,7	WIDTH(1-10) Widths (e.v.) of channels for binning the energies of up to 10 types of secondary particles. MARLOWE Version 12 provides for only 5 particle types. Ten types were permitted in Version 11. Both records	10X, 5E10.0

6 and 7 must be present (to process MARLOW Version 12 output there will be no values on record 7).

8 WIDTH(11) 10X, E10. 0
 Width (e.v.) of channels for binning the energies of primary particles.

9 DEPBIN, DEPMAX 10X, 2E10. 0
 DEPBIN = bin width (in units of ALAT(1)) for depth distribution tables.
 DEPMAX = bin width (in units of ALAT(1)) for maximum depth distribution tables.

10 NMUBIN, CHMU 10X, I2, 1X, A2
 NMUBIN = number of mu (cosine polar angle) to use for binning particles. Present maximum is 20, limited by processing and output routines, but input routine provides for 40 bins. Arrays in other routines and output formats must be modified before 40 bins can be used.
 CHMU = data type specifier. CHMU='MU' if bin boundary values on record(s) 11 are in units of mu (cosine polar angle). Leave CHMU blank if boundary values are in degrees.

11A, 11B, etc. GMU(0 to NMUBIN) or POLDEG(0 to NMUBIN) 10X, 7F10. 0
 As many records as are necessary for NMUBIN+1 values of bin boundaries for binning polar angle (theta) distribution of particles. If CHMU (record 10) is 'MU', bin boundaries must be cosines. If CHMU is anything else, bin boundaries must be in degrees.

12 NAZBIN 10X, I2
 Number of azimuthal angle (phi) bins to use for binning particles. Present maximum is 20, limited by processing and output routines, but input routine provides for 40 bins. Arrays in other routines and output formats must be modified before 40 bins can be used.

13A, 13B, etc. AZMDEG(0 to NAZBIN) 10X, 7F10. 0
 As many records as are necessary for NAZBIN+1 values of bin boundaries for binning azimuthal angle (phi) distribution of particles. Values must be in degrees.

14 XNORML 10X, F10. 0
 Factor (multiplier) to be used in normalizing values for ejected particle direction distribution table. See writeup describing normalization of array NANGLE in subroutine SUMRYZ.

15 OUTPUT CONTROL SWITCHES 10X, A70
 To eliminate undesired output, one or more of the following codes (separated by commas or spaces) may appear in columns 11-80, in any order.
 FPRIM ("Front PRIMaries") suppresses output for

reflected primaries.
RPRIM ("Rear PRIMaries") suppresses output for
transmitted primaries.
FTARG ("Front TARGet") suppresses output for
front sputtered target atoms.
RTARG ("Rear TARGet") suppresses output for
rear sputtered target atoms.
INFO causes file 'INFO' not to be included at
the end of the print file. The INFO file
may be saved or disposed of just as any
'other file.

The record must be present so if all output is
desired, columns 11-80 should be blank.

***** OTHER INPUT *****

Data files (named 'POPDAT') written by program MARLOW from routine
EXTRAI. From 1 to 10 POPDAT files may be processed in a single
MARPOP run. All POPDAT files are processed for each MARPOP input
parameter set described above. POPDAT files are read from logical
units 17, 18, . . . , 25, 26, and must be assigned the names FT17, FT18,
etc., up to FT26.
ETC.).

***** VARIABLE DECLARATIONS *****

Carriage control characters
CHARACTER*1 CHPLUS, CHWUN
Output switches
CHARACTER*5 FPRIM, RPRIM, FTARG, RTARG, INFO
CHARACTER*8 CHKODE
CHARACTER*80 CHKARD
CHARACTER*130 CHLINE
INTEGER I, ISET, KODE, LDATE, LIMRUN, LTIME, LUNIT,
1 MXFYLS, NAZBIN, NFILES
REAL DEPBIN, DEPMAX, SBND, WIDTH

PRIMARY PARTICLE PROCESSING LIMITS READ FROM CARD(S) 3
COMMON /LIMITS/ LIMRUN(10)

Output switches
COMMON /OUTSWT/ FPRIM, RPRIM, FTARG, RTARG, INFO
COMMON /READIN/ SBND(11), WIDTH(11), DEPBIN, DEPMAX
I/O logical units set in subprogram BLOKAA
COMMON /UNITS/ LUSPEC, LUPRT, LUIINFO

NEXT STATEMENT FORCES LINKING OF BLOCK DATA SUBPROGRAM THAT SETS
LOGICAL UNIT NUMBERS.

Next statement forces linking of block data subprogram that sets
constant values for I/O logical units, etc.
EXTERNAL BLOKAA

DATA CHPLUS /'+'/, CHWUN /'1'/
Maximum number of POPDAT files that may be processed
DATA MXFYLS /10/

100 FORMAT(AB0)
101 FORMAT(10X, 5E10.0)
102 FORMAT(10X, 2E10.0, I5)
103 FORMAT(10X, I2)


```

104 FORMAT(10X,10I5)
105 FORMAT(A130)
200 FORMAT(A1/'0',7X,'PROGRAM MARPOP - MARLOW POST PROCESSOR',15X,
1 'VERSION 3.1',5X,'OF 22 MAY 1986')
201 FORMAT(' ',7X,'*** UNEXPECTED END-OF-FILE, INPUT DATA CARDS')
202 FORMAT(' ',7X,'*** READ ERROR, INPUT DATA CARDS')
203 FORMAT(' ',7X,'PROCESSING',I4,2X,'POPDAT FILES')
204 FORMAT(' ',7X,'PROGRAM MARPOP FINISHED')
205 FORMAT('/'0',27X,A80/)
206 FORMAT(' ',7X,'*** UNEXPECTED END-OF-FILE, UNIT ',I4)
207 FORMAT(' ',7X,'*** READ OR DATA ERROR, UNIT ',I4)
208 FORMAT(' ',7X,
1 '*** MAXIMUM NUMBER OF RECORDS PROCESSED FROM UNIT ',I4)
209 FORMAT(' ',7X,'*** END-OF-FILE READ, INPUT DATA CARDS')
210 FORMAT(' ',7X,'UNIT ',I4,5X,'MAXRUN PRIMARIES TO BE PROCESSED')
211 FORMAT(' ',7X,'UNIT ',I4,I10,2X,'PRIMARIES TO BE PROCESSED')
212 FORMAT('0',7X,'DATE',1X,A8,5X,'TIME',1X,A8)
213 FORMAT('1CONTENTS OF FILE INFO')
214 FORMAT(A130)
215 FORMAT(' LAST RECORD OF FILE INFO')

```

C

```

WRITE (LUPRT,200) CHPLUS
CALL DATE (LDATE)
CALL CLOCK (LTIME)
OPEN (UNIT=LUINFO,FILE='INFO',STATUS='NEW')
WRITE (LUPRT,212) LDATE, LTIME
WRITE (LUINFO,200) CHPLUS
WRITE (LUINFO,212) LDATE, LTIME

```

C

C

C

C

C

On each pass through DO 4000 ISET loop, one set of parameter records is read from logical unit LUSPEC and processed.

C

```
DO 4000 ISET=1,20
```

*** Record 1 ***

```

READ (LUSPEC,100,END=8000,ERR=10000) CHKARD
IF (ISET.GT.1) WRITE (LUPRT,200) CHWUN
WRITE (LUPRT,205) CHKARD
WRITE (LUINFO,205) CHKARD

```

C

*** Record 2 ***

```

READ (LUSPEC,103,END=9000,ERR=10000) NFILES
IF (NFILES.GT.MXFYLS) NFILES=MXFYLS
WRITE (LUPRT,203) NFILES

```

C

*** Record 3 ***

```
READ (LUSPEC,104,END=9000,ERR=10000) (LIMRUN(I),I=1,NFILES)
```

C

*** Records 4, 5 ***

```
READ (LUSPEC,101,END=9000,ERR=10000) (SBND(I),I=1,10)
```

C

*** Records 6, 7, 8 ***

```
READ (LUSPEC,101,END=9000,ERR=10000) WIDTH
```

C

*** Record 9 ***

```
READ (LUSPEC,102,END=9000,ERR=10000) DEPBIN, DEPMAX
```

C

C

C

C

Remaining parameter records are read from LUSPEC in subroutine INPTB.

```

CALL INPTB (CHKODE)
IF (CHKODE.EQ.'EOF      ') THEN
GO TO 9000
ELSE IF (CHKODE.EQ.'ERROR  ') THEN
GO TO 10000

```

END IF
CALL INITLZ

On each pass through DO 3000 JFYL loop one MARLOW
'POPDAT' file is read and particle data tabulated.

DO 3000 JFYL=1,NFILES
LUDAT = 16 + JFYL
IF (ISET.GT.1) REWIND LUDAT
IF (LIMRUN(JFYL).LE.0) THEN
WRITE (LUPRT,210) LUDAT
ELSE
WRITE (LUPRT,211) LUDAT, LIMRUN(JFYL)
END IF

Read first record from logical unit LUDAT

CALL INPTA (JFYL, LUDAT, KODE)

Values of KODE returned by INPTA:

- 1 = Data read OK
- 2 = End-of-file read, no FTxx file
- 3 = Read error, file FTxx

GO TO (1000, 5000, 7000) KODE
CONTINUE
CALL PRTCLS (ISET, JFYL, LUDAT, KODE)

1000

VALUES OF KODE RETURNED BY PRTCLS:

- 1 RECORDS READ AND PROCESSED OK
- 2 UNEXPECTED END-OF-FILE, NO DATA
- 3 UNEXPECTED END-OF-FILE, PARTIAL DATA
- 4 READ ERROR, NO DATA
- 5 READ ERROR, PARTIAL DATA
- 6 MAXIMUM NUMBER OF RECORDS READ AND PROCESSED

GO TO (3000, 5000, 6000, 7000, 10500, 2000) KODE
CONTINUE
WRITE (LUPRT,208) LUDAT

2000

CONTINUE
CALL SUMRYZ

3000

PRINT TABLES AND HISTOGRAMS

CALL DUPTAA

4000

CONTINUE
GO TO 12000

5000

CONTINUE
WRITE (LUPRT,206) LUDAT
IF (I.GT.1) GO TO 11000
GO TO 12000

6000

CONTINUE
WRITE (LUPRT,206) LUDAT
GO TO 11000

7000

CONTINUE
WRITE (LUPRT,207) LUDAT
IF (I.GT.1) GO TO 11000
GO TO 12000

8000

CONTINUE
IF (ISET.LE.1) GO TO 9000
WRITE (LUPRT,209)
GO TO 12000

```

9000 CONTINUE
WRITE (LUPRT,201)
GO TO 12000
10000 CONTINUE
WRITE (LUPRT,202)
GO TO 12000
10500 CONTINUE
WRITE (LUPRT,207) LUDAT
11000 CONTINUE
C
C PRINT TABLES AND HISTOGRAMS
CALL DUPTAA
12000 CONTINUE
WRITE (LUINFO,215)
ENDFILE LUINFO
WRITE (LUPRT,204)
IF (INFO.EQ.'INFO') THEN
REWIND LUINFO
WRITE (LUPRT,213)
DO 13000 I=1,600
READ (LUINFO,105,END=14000) CHLINE
WRITE (LUPRT,214) CHLINE
13000 CONTINUE
END IF
14000 CONTINUE

```

```

C
C Next statement is only normal program execution stop
C

```

```

STOP
END
SUBROUTINE ANGLPT (KTYPE, KSURF, AVYELD, ERROR)

```

```

C
C 13 MARCH 1986
C

```

```

C ANGLPT MAKES LINE PRINTER HISTOGRAM PLOT OF ANGULAR DISTRIBUTION
C

```

```

C ***** CALLING PARAMETERS *****
C

```

```

C KTYPE - (INTEGER, PASSED) PARTICLE TYPE NUMBER (1-10 FOR SECONDARY,
C 11 FOR PRIMARY PARTICLES)
C KSURF - (INTEGER, PASSED) TARGET SURFACE ID (1=FRONT, 2=BACK)
C AVYELD - (REAL, PASSED) MEAN SECONDARY PARTICLE YIELD PER PRIMARY
C ERROR - (REAL, PASSED) STANDARD DEVIATION OF MEAN YIELD (AVYELD)
C

```

```

C THE PLOT IS SET UP IN ARRAY LYNPLT, 81 COLUMNS (1=LEFTMOST) BY 42
C LINES (LINE 2 IS TOP OF PLOT FRAME, LINE 1 IS PRINTED ABOVE THE
C FRAME). IF COSINE BIN 20 (NPLOT(20,KTYPE,KSURF)) IS GREATER THAN
C ZERO, THE HEIGHT OF THE RIGHTMOST HISTOGRAM BAR = 1.0, AND THE OTHER
C BAR HEIGHTS ARE PROPORTIONED TO IT. IF NPLOT(20,KTYPE,KSURF) = 0,
C THE LARGEST BIN IS FOUND AND ASSIGNED A HEIGHT OF 1.0, AND THE OTHER
C BARS ARE PROPORTIONED TO THAT ONE. IF A BAR HAS A CALCULATED HEIGHT
C GREATER THAN 1.0, THE TOP OF THE BAR EXTENDS JUST ABOVE THE PLOT AND
C THE CALCULATED HEIGHT (ENCODED INTO ARRAY IXCEED) IS PRINTED ABOVE
C THE TOP OF THE BAR.
C

```

```

C ***** VARIABLE DECLARATIONS *****
C

```

```

C PLOT AXIS AND HEADING LABELS
C DIMENSION LABLAX(11), LABSRF(4,2)
C STORAGE FOR PLOT CHARACTERS
C

```

```

DIMENSION IXCEED(20), LYNPLT(81,42), LYNDOUT(324,10), LE01(24),
1      LE02(24), LE03(24), LE04(24)
C      TEMPORARY STORAGE FOR PLOT LEGEND LINES
C      DIMENSION LEGEND(3)

COMMON /BINS/ POLDEG(0:40), GMU(0:40), DMU(40), NMUBIN,
1      AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN
C      ADUMMY = ARRAY NOT USED IN THIS SUBPROGRAM
C      LTYPE = PARTICLE ID (LITERAL)
COMMON /FSTREC/ ADUMMY(48), LTYPE(10), DDUMMY(70)
C      SURFACE BINDING ENERGIES
COMMON /READIN/ SBND(11)
C      ENERGY (NENERG), YIELD (NYIELD), DEPTH (NDEPTH), AND
C      ANGULAR (NANGLE) DISTRIBUTION TABLES.
C      KSUM, KSUMSQ = SUM AND SUM OF SQUARES OF YIELD FOR
C      CALCULATING MEAN YIELD AND STANDARD DEVIATION.
C      NPRYMS = TOTAL NUMBER OF PRIMARY PARTICLES.
COMMON /TABLES/ NENERG(100,11,2), NYIELD(21,10,2),
1      NDEPTH(21,10,2), NRMLZD(21,21,11,2), FACNRM(11,2),
2      NANGLE(21,21,11,2), KSUM(10,2), KSUMSQ(10,2),
3      NPRYMS
C      COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

EQUIVALENCE (LE01(1),LYNPLT(4,4)), (LE02(1),LYNPLT(4,5)),
1      (LE03(1),LYNPLT(4,6)), (LE04(1),LYNPLT(4,7)),
2      (LYNDOUT(1,1),LYNPLT(1,2))
C      SAVE

DATA LABLAX /'0.0 ', '0.1 ', '0.2 ', '0.3 ', '0.4 ', '0.5 ',
1      '0.6 ', '0.7 ', '0.8 ', '0.9 ', '1.0 ' /
DATA LABSRF /' RE', 'FLEC', ' FR', 'ONT ', 'TRAN', 'SMIT',
1      ' R', 'EAR ' /
C      NUMBER OF CHARACTERS IN ARRAY LEGEND
DATA NLEG /28/
C      SYMBOLS FOR PLOT
DATA IBLANK /' ', IDASH /'-'/, IPLUS /'+'/
DATA MXAZM /21/

201 FORMAT(1H1////1H0,34X,'ANGULAR DISTRIBUTION',3X,'-',1X,2A4,
1      'SPUTTERED',2X,A2,I3,2X,'ATOMS')
202 FORMAT(1H1////1H0,33X,'ANGULAR DISTRIBUTION',3X,'-',3X,2A4,
1      'TED PRIMARY PARTICLES')
203 FORMAT(1H ,24X,20A4)
204 FORMAT((1H ,19X,A4,81A1,1X,A3/ 2(1H ,23X,81A1/), 1H ,23X,81A1))
205 FORMAT(20X,21(3X,'I') / 22X,21(F4.1) /
1      '0',49X,'Polar Angle (Degrees)')
206 FORMAT(1H0,7X,'*** NO DATA FOR ANGULAR DISTRIBUTION PLOT' / 1H1)
207 FORMAT('0',19X,'1.0=',18,2X,'Particles/dMu')
401 FORMAT(F4.1)
402 FORMAT('SURF. BINDG. EN. ',1PE9.2,1X)
403 FORMAT('PRIMARIES',5X,I7,3X)
404 FORMAT('MEAN YIELD',1X,F10.1,3X)
405 FORMAT('ERROR',6X,F10.1,3X)

C
C      PRINT HEADING
IF (KTYPE.LT.11) PRINT 201, (LABSRF(I,KSURF),I=3,4), LTYPE(KTYPE),
1      KTYPE
IF (KTYPE.EQ.11) PRINT 202, (LABSRF(I,KSURF),I=1,2)
C      BLANK PLOT ARRAY, PUT IN GRID LINES
DO 800 I=1,20

```

```

IXCEED(I) = IBLANK
800 CONTINUE
DO 900 I=1,42
DO 900 J=1,81
LYNPLT(J,I) = IBLANK
900 CONTINUE
DO 1000 I=2,42,8
DO 1000 J=1,81,2
LYNPLT(J,I) = IDASH
1000 CONTINUE
DO 2000 J=2,42
DO 2000 I=1,81,16
LYNPLT(I,J) = IDASH
2000 CONTINUE
C SET HISTOGRAM BAR HEIGHT REFERENCE
IYTOP = NRMLZD(1, MXAZM, KTYPE, KSURF)
DO 3000 I=NMUBIN, NMUBIN
IF (NRMLZD(I, MXAZM, KTYPE, KSURF).GT. IYTOP)
1 IYTOP=NRMLZD(I, MXAZM, KTYPE, KSURF)
3000 CONTINUE
IF (IYTOP.GT.0) GO TO 4000
PRINT 206
RETURN

C
4000 CONTINUE
YMAX = FLOAT(IYTOP)
C SET UP HISTOGRAM IN PLOT ARRAY
LYNPLT(1,42) = IPLUS
LAST = 42
C HISTOGRAM BAR FOR ONE OF THE 20
C COSINE BINS IN NRMLZD IS SET UP ON
C EACH PASS THROUGH DO 14000 I LOOP.
C I IS BIN SUBSCRIPT IN NRMLZD,
C I+1 IS CURRENT BAR IN ARRAY LYNPLT.
DO 14000 I=1, NMUBIN
NPLX40 = 40 * NRMLZD(I, MXAZM, KTYPE, KSURF)
LINTMP = NPLX40 / IYTOP
IF ((NPLX40-(LINTMP*IYTOP)).GT.0) LINTMP=LINTMP+1
LINE = 42 - LINTMP
IF (LINE.GT.1) GO TO 5000
RATIO = FLOAT(NRMLZD(I, MXAZM, KTYPE, KSURF)) / YMAX
ENCODE (4,401, IXCEED(I)) RATIO
LINE = 1
5000 CONTINUE
JSTART = 2 + (4*(I-1))
JSTOP = JSTART + 3
DO 6000 J=JSTART, JSTOP
LYNPLT(J,LINE) = IPLUS
6000 CONTINUE
IF (LINE.GE.41) GO TO 8000
KSTART = LINE + 1
DO 7000 K=KSTART, 41
DO 7000 J=JSTART, JSTOP
LYNPLT(J,K) = IBLANK
7000 CONTINUE
8000 CONTINUE
IF (LINE-LAST) 9000, 14000, 11000
9000 CONTINUE
ICOLUM = JSTART
DO 10000 J=LINE, LAST

```

```

      LYNPLT(ICOLUM,J) = IPLUS
10000  CONTINUE
      GO TO 13000
11000  CONTINUE
      ICOLUM = JSTART - 1
      DO 12000 J=LAST,LINE
          LYNPLT(ICOLUM,J) = IPLUS
12000  CONTINUE
13000  CONTINUE
      LAST = LINE
14000  CONTINUE
      DO 15000 J=LAST,42
          LYNPLT(81,J) = IPLUS
15000  CONTINUE
C
C          PUT LEGEND IN PLOT.  CIMOVE
C          EXPANDS LEGEND LINE FROM ARRAY
C          LEGEND INTO PLOT ARRAY.
      ENCODE (24,402,LEGEND(1)) SBND(KTYPE)
      CALL CIMOVE (LEGEND, LEG1)
      ENCODE (24,403,LEGEND(1)) NPRYMS
      CALL CIMOVE (LEGEND, LEG2)
C
C          OMIT OTHER LEGEND LINES FOR PRIMARY
C          PARTICLE
      IF (KTYPE.GT.10) GO TO 16000
      ENCODE (24,404,LEGEND(1)) AVYELD
      CALL CIMOVE (LEGEND, LEG3)
      ENCODE (24,405,LEGEND(1)) ERROR
      CALL CIMOVE (LEGEND, LEG4)
16000  CONTINUE
C
C          PRINT THE HISTOGRAM
      PRINT 204, IBLANK, (LYNPLT(I,1),I=1,81)
      PRINT 204, (LABLAX(12-I), (LYNOUT(J,I),J=1,81), LABLAX(12-I),
1      (LYNOUT(K,I),K=82,324), I=1,10)
      PRINT 204, LABLAX(1), (LYNPLT(I,42),I=1,81)
      WRITE (LUPRT,205) (POLDEG(I),I=0,NMUBIN)
      WRITE (LUPRT,207) IYTOP
      RETURN
      END
      BLOCK DATA BLOKAA
C
C 12 MARCH 1986
C
C BLOKAA sets values for some constants used in program MARPOP.
C
C   Array dimensions; values used mainly for loop indices.
C   MXSURF = Maximum number of target surfaces
C   MXTYPE = Maximum number of particle types.  Numbers 1-5
C           are used for secondary particles (10 were
C           permitted by MARLOWE Version 11).  Type 11 is
C           primaries.
C   MXPOLR = Maximum number of values for boundaries of polar
C           angle bins.
C   MXAZM  = Maximum number of values for boundaries of
C           azimuthal angle bins.
      COMMON /MXDMNS/ MXSURF, MXTYPE, MXPOLR, MXAZM
C
C   I/O Logical unit assignments
C   LUSPEC = Input parameter file
C   LUPRT  = Output print file
C   LUINFO = Output "information" file

```

COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

DATA MXSURF /2/, MXTYPE /11/, MXPOLR /21/, MXAZM /21/
DATA LUSPEC /5/, LUPRT /6/, LUINFO /9/

END
SUBROUTINE CIMOVE (ISORCE, ITARGET)

24 JANUARY 1986

CIMOVE EXPANDS EIGHT CHARACTERS FROM EACH WORD OF ISORCE INTO
ARRAY ITARGET, ONE CHARACTER PER WORD, IN LEFTMOST BYTE, WITH REST
OF WORD FILLED WITH ASCII SPACES (40 OCTAL, 20 HEXADECIMAL).

DIMENSION ISORCE(3), ITARGET(24)

DO 1000 I=1,3

DO 1000 J=1,8

ITARGET(8*(I-1)+J) = X'0020202020202020' +

1 AND(SHIFT(ISORCE(I),8*(J-1)),X'FF00000000000000')

1000 CONTINUE

RETURN

END

SUBROUTINE FRONT (KTYPE, DEEP, DEPMAX, LA, KARMA)

17 JANUARY 1986

FRONT extracts and tabulates the following data for particles
escaping from the front surface: (a) number of atomic collisions,
(b) numbers of replacements in replacement sequences, and (c) maximum
distance from surface. A second entry point, FRINIT, is used to
initialize the arrays in which data are tabulated.

***** CALLING PARAMETERS *****

KTYPE (integer,passed) is the particle type. MARLOWE Version 11
provided up to 10 types. Version 12 provides only 5 types.
Therefore, current permitted values of KTYPE are 1-5 and 11,
for up to 5 types of secondary particles (1-5) with a value
of 11 indicating a primary particle.

DEEP (real,passed) $d + (100 * (10 * z))$, where d is MARLOWE
variable DEEP, and z is the maximum depth (in MARLOWE
ALAT(1) units, truncated to the next lowest 0.1 unit)
reached by the particle.

DEPMAX (real,passed) Bin width (in MARLOWE ALAT(1) units) for
maximum depth distribution tables.

LA (integer,passed) $m + 256 * n$, where m is MARLOWE variable LA
(a particle type identifier) and n is the number of atomic
collisions of the particle between its deepest point in the
slab and the escape (front) surface.

KARMA (integer,passed)
 $k + 65536 * (a + 16 * b + (16 ** 2) * c + (16 ** 3) * d + (16 ** 4) * e)$
where k is MARLOWE variable KARMA and a,b,c,d,e are the
numbers of replacements in each of five replacement
sequences.

C This processing was grouped together and put in a subroutine to
C minimize the modifications to subroutine CALCNS in MARPOP Version 2.0
C for TI-ASC.

C Data are tabulated in the arrays MXDEP, LENSEG, and NCOLIS. In
C those arrays the dimension 11 represents type of atom, passed to this
C routine in calling parameter KTYPE. Only values 1-5 and 11 are used
C by data from MARLOWE Version 12, so nearly half of each array is not
C used (see description of KTYPE above.)

C MXDEP Maximum depth distribution table. Depths are measured in
C units of MARLOWE variable ALAT(1). 21 bins are provided for
C each particle type. MXDEP(1,i) is count of particles
C reaching maximum depth less than 1 unit; MXDEP(2,i) counts
C particles reaching depth of 1 unit, but less than 2 units;
C ; MXDEP(20,i) counts particles reaching 19 units but
C less than 20 units; MXDEP(21,i) is count of particles
C reaching 20 units or more.

C LENSEG Length of replacement sequences table. 15 collisions in
C sequence X 5 sequences X 11 particle types.

C NCOLIS Number of collisions from deepest point table. First index
C runs 1-21 for 1-20 collisions and .GT.20 in 21st bin.
C Second index is for 11 particle types.

C
C INTEGER I, KARMA, KARTMP, KCOLIS, KMXDEP, KOUNT, KTYPE, LA,
1 LENSEG, MXDEP, NCOLIS, NSEQ
C REAL DEEP, DEPMAX
C DIMENSION NSEQ(5)

C
C COMMON /FRSURF/ MXDEP(21,11), LENSEG(15,5,11), NCOLIS(21,11)

C
C Next statement causes values of all variables to be saved
C on exit from this subroutine.

C
C SAVE

C
C ***** Extract and bin maximum depth data *****

C
C KMXDEP is first calculated as number of DEPMAX units,
C then converted to bin index.
C 1.0E-8 in next statements prevents roundoff error

C
C KMXDEP = IFIX(((DEEP-AMOD(DEEP,100.0))/(1000.0*DEPMAX))+1.0E-8)
C IF (KMXDEP.GT.20) KMXDEP=20
C MXDEP(KMXDEP+1,KTYPE) = 1 + MXDEP(KMXDEP+1,KTYPE)

C
C ***** Extract and bin collision data *****

C
C KCOLIS = LA / 65536
C IF (KCOLIS.GT.20) KCOLIS=20
C NCOLIS(KCOLIS+1,KTYPE) = 1 + NCOLIS(KCOLIS+1,KTYPE)

C
C ***** Extract and bin replacement sequence data *****

C
C KARTMP = KARMA / 256
C KOUNT = 0
C DO 1000 I=1,5
C NSEQ(I) = MOD(KARTMP,16)
C IF (NSEQ(I).GT.0) KOUNT=KOUNT+1


```

KARTMP = KARTMP / 16
1000 CONTINUE
DO 2000 I=1,5
  IF (NSEG(I).GT.0) LENSEG(NSEG(I),KOUNT,KTYPE) =
1    1 + LENSEG(NSEG(I),KOUNT,KTYPE)
2000 CONTINUE
RETURN

```

```

C
C *****
C *****
C ***** ENTRY POINT TO INITIALIZE ARRAYS *****
C *****
C *****
C *****

```

```

C ENTRY FRINIT
C
DO 3000 I=1,11
  DO 3000 J=1,21
    MXDEP(J,I) = 0
3000 CONTINUE
DO 4000 I=1,11
  DO 4000 J=1,21
    NCOLIS(J,I) = 0
4000 CONTINUE
DO 5000 I=1,11
  DO 5000 J=1,5
    DO 5000 K=1,15
      LENSEG(K,J,I) = 0
5000 CONTINUE
RETURN
END
SUBROUTINE INITLZ

```

```

C
C 12 MARCH 1986
C
LOGICAL LOJDEG
COMMON /BINS/ POLDEG(0:40), GMU(0:40), DMU(40), NMUBIN,
1 AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN
COMMON /MORBNS/ XMUMID(20), PHIMID(40)

```

```

C
C SAVE values.
C SAVE
C
DATA MXBINS /40/, PI /3.141592653590/
C
RADFAC = PI / 180.0

```

```

C *****
C *
C * THETA (POLAR ANGLE) BINS *
C *
C *****

```

```

C Initialize polar angle bin boundaries: Non-zero POLDEG value
C indicates input boundaries were in degrees and boundaries in
C cosine (GMU) units are calculated in DO 3000 I loop. If
C POLDEG values are all zero, input boundaries were in mu units
C and boundaries in degrees are calculated in DO 4000 I loop.
C

```

```

LOJDEG = .FALSE.

```

```

DO 1000 I=0, MXBINS
  IF (POLDEG(I).NE.0.0) THEN
    LOJDEG = .TRUE.
    GO TO 2000
  END IF
1000 CONTINUE
2000 CONTINUE
  IF (LOJDEG) THEN
    DO 3000 I=0, NMUBIN
      GMU(I) = COS(RADFAC*POLDEG(I))
    CONTINUE
  ELSE
    DO 4000 I=0, NMUBIN
      POLDEG(I) = ACOS(GMU(I)) / RADFAC
    CONTINUE
  END IF
4000 CONTINUE
  END IF

C
C      Calculate width (DMU) and midpoint (XMUMID) of each theta bin
C
DO 5000 I=1, NMUBIN
  DMU(I) = GMU(I) - GMU(I-1)
  XMUMID(I) = 0.5 * (GMU(I) + GMU(I-1))
5000 CONTINUE

C
C      *****
C      *
C      *   PHI (AZIMUTHAL ANGLE) BINS   *
C      *
C      *****
C
C      Calculate phi boundaries (AZMRAD) in radian, and width
C      (DAZBIN) and midpoint (PHIMID) of each phi bin.
C
DO 6000 I=0, NAZBIN
  AZMRAD(I) = RADFAC * AZMDEG(I)
6000 CONTINUE
DO 7000 I=1, NAZBIN
  DAZBIN(I) = AZMRAD(I) - AZMRAD(I-1)
  PHIMID(I) = 0.5 * (AZMRAD(I) + AZMRAD(I-1))
7000 CONTINUE
RETURN
END
SUBROUTINE INPTA (ITER, LUDAT, KODE)

C
C 24 JANUARY 1986
C
C INPTA READS FIRST RECORD WRITTEN BY PROGRAM MARLOW AND PRINTS MARLOW
C RUN DATE AND TIME IN HEADING
C
C      ***** CALLING PARAMETERS *****
C
C ITER (INTEGER, PASSED) FILE IDENTIFIER
C LUDAT (INTEGER, PASSED) LOGICAL UNIT FROM WHICH CURRENT INPUT FILE
C IS READ.
C KODE (INTEGER, RETURNED) VALUES OF KODE RETURNED BY INPTA:
C 1 DATA READ OK
C 2 End-of-file read, no file FTnn (nn=LUDAT)
C 3 Read error, file FTnn (nn=LUDAT)
C

```

CHARACTER*8 LDATE, LTIME, JOBID(5), IDENT(15), OLDTYP, NEWTYP

```

C      INTEGER I, INREC, INTEMP, ITER, KODE, LTYPE, NTYPE
C      ARRAY INTO WHICH RECORD IS READ
C      DIMENSION INTEMP(128)
C      DIMENSION LTYPE(10)
C      COMMON /FSTREC/ INREC(128)
C      EQUIVALENCE (NTYPE, INREC(7)), (LTYPE(1), INREC(49))
C      EQUIVALENCE (MAXRUN, INTEMP(6))
C      SAVE
C      201 FORMAT(' ', 7X,
C      1      'PROCESSING OUTPUT FROM PROGRAM MARLOW EXECUTION OF', 2(1X, A8)/
C      2      '0', 5A8, 5X, 'MAXRUN=', I10/' ', 15A8)
C      202 FORMAT('0', 7X, '*** UNIT', 2X, I4, 5X,
C      1      'NUMBER OF PARTICLE TYPES IS', I5, 4X,
C      2      'DOES NOT MATCH PREVIOUS:', I5)
C      203 FORMAT('0', 7X, '*** UNIT', 2X, I4, 5X, 'PARTICLE SYMBOL NUMBER',
C      1      I4, 2X, 'IS', 2X, A8, 4X, 'DOES NOT MATCH PREVIOUS:', 2X, A8)
C
C      READ (LUDAT, END=4000, ERR=5000) INTEMP
C      CALL MKCHAR (INTEMP(1), LDATE)
C      CALL MKCHAR (INTEMP(3), LTIME)
C      DO 800 I=1, 5
C          CALL MKCHAR (INTEMP(64+I), JOBID(I))
C      800 CONTINUE
C      DO 900 I=1, 15
C          CALL MKCHAR (INTEMP(69+I), IDENT(I))
C      900 CONTINUE
C      WRITE (6, 201) LDATE, LTIME, JOBID, MAXRUN, IDENT
C      IF (ITER.LE.1) GO TO 3000
C      IF (INTEMP(7).EQ.NTYPE) GO TO 1000
C      WRITE (6, 202) LUDAT, INTEMP(7), NTYPE
C      GO TO 5000
C      1000 CONTINUE
C      DO 2000 I=1, NTYPE
C          IF (INTEMP(48+I).EQ.LTYPE(I)) GO TO 2000
C          CALL MKCHAR (INTEMP(48+I), NEWTYP)
C          CALL MKCHAR (INREC(48+I), OLDTYP)
C          WRITE (6, 203) LUDAT, I, NEWTYP, OLDTYP
C          GO TO 5000
C      2000 CONTINUE
C      3000 CONTINUE
C      DO 3500 I=1, 128
C          INREC(I) = INTEMP(I)
C      3500 CONTINUE
C      KODE = 1
C      RETURN
C
C      4000 CONTINUE
C      KODE = 2
C      RETURN
C
C      5000 CONTINUE
C      KODE = 3
C      RETURN
C      END
C      SUBROUTINE INPTB (CHKODE)

```

C 12 MARCH 1986

C INPTB reads parameter records 10-

C Values of CHKODE returned by INPTB:

C 'OK ' if data records were read sucessfully.
C 'EOF ' if end-of-file read (all records did not exist).
C 'ERROR ' if error on attempt to read record.
C

CHARACTER*5 FPRIM, RPRIM, FTARG, RTARG, INFO
CHARACTER CHKODE*8, CHLINE*81, CHMU*2

C Following arrays provide for 40 bins, but only 20 are
C allowed in some other subprograms.

COMMON /BINS/ POLDEG(0:40), GMU(0:40), DMU(40), NMUBIN,
1 AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN

COMMON /OUTSWT/ FPRIM, RPRIM, FTARG, RTARG, INFO

COMMON /SPECS/ XNORML

COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

C SAVE values.

C SAVE

C DATA MXBINS /40/

C 100 FORMAT(A80)

101 FORMAT(10X, I2, 1X, A2)

102 FORMAT(10X, 7F10.0)

103 FORMAT(10X, F10.0)

C CHKODE = 'OK '

C *****
C *
C * THETA (POLAR ANGLE) BINS *
C *
C *****

C Read number of mu (cosine theta) bins to use and data type
C specifier (CHMU). If CHMU='MU', read bin boundaries in
C units of mu and set array POLDEG to zero. If CHMU is not
C 'MU', bin boundaries are in degrees, read into array POLDEG.
C

*** Record 10 ***

C READ (LUSPEC, 101, END=3000, ERR=4000) NMUBIN, CHMU
C IF (CHMU.EQ. 'MU') THEN

*** Record(s) 11 ***

C READ (LUSPEC, 102, END=3000, ERR=4000) (GMU(I), I=0, NMUBIN)

DO 1000 I=0, MXBINS

POLDEG(I) = 0.0

1000 CONTINUE

C ELSE

READ (LUSPEC, 102, END=3000, ERR=4000) (POLDEG(I), I=0, NMUBIN)

C END IF

C *****
C *
C * PHI (AZIMUTHAL ANGLE) BINS *
C *
C *****

Read number of azimuthal angle bins and bin boundaries.

*** Record 12 ***

READ (LUSPEC, 101, END=3000, ERR=4000) NAZBIN

*** Record(s) 13 ***

READ (LUSPEC, 102, END=3000, ERR=4000) (AZMDEG(I), I=0, NAZBIN)

```
*****  
*  
*   ROUNDING FACTOR   *  
*  
*****
```

*** Record 14 ***

READ (LUSPEC, 103, END=3000, ERR=4000) XNORML

```
*****  
*  
*   OUTPUT ELIMINATION SWITCHES   *  
*  
*****
```

Set defaults, specifying all output is desired.

```
FPRIM = 'FPRIM'  
RPRIM = 'RPRIM'  
FTARG = 'FTARG'  
RTARG = 'RTARG'  
INFO  = 'INFO '
```

Read record from specification file and decode data in columns 11-80.

*** Record 15 ***

READ (LUSPEC, 100, END=3000, ERR=4000) CHLINE(1:80)

CHLINE(81:81) = ' '

DO 2000 I=11,77

IF (CHLINE(I:(I+4)).EQ. 'FPRIM') THEN

FPRIM = ' '

ELSE IF (CHLINE(I:(I+4)).EQ. 'RPRIM') THEN

RPRIM = ' '

ELSE IF (CHLINE(I:(I+4)).EQ. 'FTARG') THEN

FTARG = ' '

ELSE IF (CHLINE(I:(I+4)).EQ. 'RTARG') THEN

RTARG = ' '

ELSE IF (CHLINE(I:(I+3)).EQ. 'INFO') THEN

INFO = ' '

END IF

2000 CONTINUE

GO TO 5000

3000 CONTINUE

CHKODE = 'EOF'

GO TO 5000

4000 CONTINUE

CHKODE = 'ERROR'

5000 CONTINUE

RETURN

END

SUBROUTINE MKCHAR (INWORD, CHOUT)

C
C 23 JANUARY 1986 11:09
C
C MKCHAR CONVERTS THE VALUE IN INWORD, BYTE-BY-BYTE, INTO THE CHARACTER
C VARIABLE CHOUT.

C
C CHARACTER*8 CHOUT
C INTEGER INWORD

C
C DO 1000 I=1,7
C CHOUT(I:I) = CHAR(AND(X'00000000000000FF', SHIFT(INWORD, 8*I)))
1000 CONTINUE
C CHOUT(8:8) = CHAR(AND(X'00000000000000FF', INWORD))
C RETURN
C END
C SUBROUTINE CUPTAA

C
C 22 MAY 1986

C
C CUPTAA PRINTS DISTRIBUTION TABLES

C
C CHARACTER*5 FPRIM, RPRIM, FTARG, RTARG

C
C DIMENSION LABSRF(3,2)

C
C COMMON /BINS/ POLDEG(0:40), GMU(0:40), DMU(40), NMUBIN,
1 AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN
C COMMON /FORFAC/ FACDIV(11,2)
C COMMON /FSTREC/ ADUMMY(5), MAXRUN, NTYPE, BDUMMY(41), LTYPE(10),
1 DDUMMY(70)
C COMMON /MXDMNS/ MXSURF, MXTYPE, MXPOLR, MXAZM
C COMMON /MORBNS/ XMUMID(20), PHIMID(40)
C COMMON /OUTSWT/ FPRIM, RPRIM, FTARG, RTARG
C COMMON /READIN/ SBND(11), WIDTH(11), DEPBIN, DEPMAX
C COMMON /SPECS/ XNORML
C COMMON /TABLES/ NENERG(100,11,2), NYIELD(21,10,2),
1 NDEPTH(21,10,2), NRMLZD(21,21,11,2), FACNRM(11,2),
2 NANGLE(21,21,11,2), KSUM(10,2), KSUMSQ(10,2),
3 NPRYMS
C COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

C
C SAVE

C
C Factor for converting degrees to radians
C DATA DEGRAD /0.0174532925/
C DATA LABSRF /' REF', 'LECT', 'ION ', 'TRAN', 'SMIS', 'SION'/
C MAXIMUM NUMBER OF BINS IN 'TABLES' COMMON BLOCK ARRAY
C NENERG
C DATA MXMU /21/, MXPHI /21/, NBINLM /100/
C

C
C 201 FORMAT('1',43X,'PRIMARY PARTICLE ',3A4,' COEFFICIENTS')
C 202 FORMAT('1',45X,3A4,' SPUTTERING OF ',A2,I3,' ATOMS')/
C 1 1H,43X,'SURFACE BINDING ENERGY (E.V.)',G14.6)
C 203 FORMAT(1H0,52X,'DISTRIBUTION OF YIELD VALUES')/
C 1 1H,8X,'YIELD',8X,'0',20I5,'+')
C 204 FORMAT(1H,8X,'FREQUENCY',21I5)
C 205 FORMAT(1H0,25X,'DISTRIBUTION OF ORIGINAL DEPTHS OF SPUTTERED',
C 1 ' ATOMS (CHANNEL WIDTH',G13.6,'')/
C 2 1H,8X,'DEPTH',4X,21I5,'+')
C 206 FORMAT(1H0,43X,'DISTRIBUTION OF EJECTED PARTICLE DIRECTIONS')/
C

```

1 ' ', 39X, 'POLAR ANGLE (ACROSS)'/
2 1H, 39X, 'CHANNEL WIDTH (DOWN), AZIMUTHAL ANGLE (DEGREES)',
3 013.6/
4 ' ', 10X, 'CHAN', 2015, 2X, 'TOTALS' / (' ', 10X, I3, 1X, 2015, I7))
207 FORMAT(1H0, 13X, 'PRIMARY PARTICLES', 7X, 'MEAN YIELD', 14X, 'ERROR' /
1 1H, 18X, I5, 13X, F10.2, 10X, F10.2)
208 FORMAT('0', 'Totals (*dPhi)', 2015, I7)
209 FORMAT('1', 43X, 'DISTRIBUTION OF EJECTED PARTICLE DIRECTIONS' /
1 'OParticle Count/(dMu*dPhi), normalized by factor (' ,G11.4,
2 ' * ', G11.4, '/', G11.4, ') = ', G11.4/
3 5X, 'Channel', 1X, 2015/
4 9X, 'dMu', 3X, 20(1X, F4.2) /
5 2X, 'Phi', 5X, 'Mu', 1X, 21(1X, F4.2) /
6 ' Chan Rad Deg I', 20(4X, 'I'), 1X, 'Totals' /
7 4X, F5.1, F6.1, 102X, '(*dMu)'/
8 (' ', I3, F5.1, F6.1, 2015, I7))
210 FORMAT(' ', 14X, 21('I ') / ' POL. ANG. DEG', 21F5.1 /
1 ' POL. ANG. RAD', 21F5.1)
211 FORMAT('1'/'0', 43X, 'DISTRIBUTION OF EJECTED PARTICLE DIRECTIONS')
212 FORMAT('0', 57X, 'PARTICLE COUNTS')
213 FORMAT('0', 14X, 'Bin', 2X, 2015 / 15X, 'dMu', 3X, 20F5.2 /
1 16X, 'Mu', 3X, 20F5.2 /
2 9X, 'dPhi', 3X, 'Phi', 1X, 20(4X, 'I'), 4X, 'Row' /
3 3X, 'Bin', 1X, 2(1X, '(Deg)'), 1X, 20(4X, 'I'), 4X, 'Sums' //
4 (4X, I2, 1X, F5.1, 3X, I3, 3X, 2015, I7 /
5 4(4X, I2, 1X, F5.1, 3X, I3, 3X, 2015, I7)))
214 FORMAT(8X, 'Column Sums', 2X, 2015, I7 /
1 '0', 7X, 'Theta (Deg)', 2X, 2015 /
2 7X, 'dTheta (Deg)', 2X, 20F5.1)

```

C

DELAZM = 360.0 / FLOAT(NAZBIN)

AVYELD = 0.0

ERROR = 0.0

DO 7000 KSURF=1,2

```

1 IF (((KSURF.EQ.1).AND.(FPRIM.EQ.'FPRIM')).OR.
((KSURF.EQ.2).AND.(RPRIM.EQ.'RPRIM')))) THEN
DO 1000 I=1,NBINLM
IF (NENERG(I,11,KSURF).NE.0) GO TO 2000

```

1000 CONTINUE

END IF

GO TO 3000

2000 CONTINUE

C

PRINT PRIMARY PARTICLE TABLES

PRINT 201, (LABSRF(I,KSURF), I=1,3)

KTYPE = 11

CALL DUPTBB (KTYPE, KSURF)

PRINT 206, DELAZM, (I, I=1,20),

```

1 ((J, (NANGLE(K, J, 11, KSURF), K=1, 21)), J=1, NAZBIN)

```

IF (NAZBIN.GT.1) WRITE (6,208) (NANGLE(J,21,11,KSURF), J=1,20)

IF (KSURF.EQ.1) CALL DUPTCC (KTYPE, DEPMAX)

CALL ANOPLT(KTYPE, KSURF, AVYELD, ERROR)

3000 CONTINUE

IF (((KSURF.EQ.1).AND.(FTARG.EQ.'FTARG')).OR.

```

1 ((KSURF.EQ.2).AND.(RTARG.EQ.'RTARG')))) THEN

```

DO 6000 KTYPE=1, NTYPE

DO 4000 I=1, NBINLM

IF (NENERG(I, KTYPE, KSURF).GT.0) GO TO 5000

4000 CONTINUE

GO TO 6000

5000 CONTINUE

C

PRINT SECONDARY PARTICLE TABLES

```

1 PRINT 202, (LABSRF(I,KSURF), I=1,3), LTYPE(KTYPE), KTYPE,
  SBND(KTYPE)
  SUM = FLOAT(KSUM(KTYPE,KSURF))
  PRIMES = FLOAT(NPRYMS)
  AVYELD = SUM / PRIMES
  IF (NPRYMS.LE.1) ERROR = 0.0
  IF (NPRYMS.GT.1) ERROR = SQRT((FLOAT(KSUMSQ(KTYPE,KSURF))-
1 ((SUM*SUM)/PRIMES))/FLOAT(NPRYMS*(NPRYMS-1)))
  PRINT 207, NPRYMS, AVYELD, ERROR
  CALL OUP TBB (KTYPE, KSURF)
  PRINT 203, (I, I=1,20)
  PRINT 204, (NYIELD(I,KTYPE,KSURF), I=1,21)
  PRINT 205, DEPBIN, (I, I=1,21)
  PRINT 204, (NDEPTH(I,KTYPE,KSURF), I=1,21)
  IF (KSURF.EQ.1) CALL OUP TCC (KTYPE,DEPMAX)
  IF (NANGLE(MXPOLR, MXAZM, KTYPE, KSURF).GT.0) THEN
    WRITE (LUPRT, 211)
    WRITE (LUPRT, 212)
    WRITE (LUPRT, 213) (I, I=1, NMUBIN), (DMU(J), J=1, NMUBIN),
1 ((0.5*(GMU(K)+GMU(K-1))), K=1, NMUBIN),
2 ((L, (AZMDEG(L)-AZMDEG(L-1))),
3 (NINT(0.5*(AZMDEG(L)+AZMDEG(L-1)))),
4 (NANGLE(M, L, KTYPE, KSURF), M=1, NMUBIN),
5 (NANGLE(MXPOLR, L, KTYPE, KSURF))), L=1, NAZBIN)
    WRITE (LUPRT, 214)
1 ((NANGLE(I, MXAZM, KTYPE, KSURF)), I=1, NMUBIN),
2 NANGLE(MXPOLR, MXAZM, KTYPE, KSURF),
3 ((NINT(0.5*(POLDEG(J)+POLDEG(J-1)))), J=1, NMUBIN),
4 ((POLDEG(K-1)-POLDEG(K)), K=1, NMUBIN)
  END IF
  WRITE (LUPRT, 209) AVYELD, XNORML, FACDIV(KTYPE,KSURF),
1 FACNRM(KTYPE,KSURF), (I, I=1, NMUBIN),
2 (DMU(J), J=1, NMUBIN), (GMU(K), K=0, NMUBIN),
2 AZMRAD(0), AZMDEG(0), ((L, AZMRAD(L), AZMDEG(L),
3 (NRMLZD(M, L, KTYPE, KSURF), M=1, NMUBIN),
4 NRMLZD(MXMU, L, KTYPE, KSURF)), L=1, NAZBIN)
  IF (NAZBIN.GT.1) WRITE (6, 208) (NRMLZD(J, MXPHI, KTYPE, KSURF),
1 J=1, NMUBIN)
  WRITE (6, 210) (POLDEG(I), I=0, NMUBIN),
1 ((DEGRAD*POLDEG(I)), I=0, NMUBIN)
  CALL ANOPLT(KTYPE, KSURF, AVYELD, ERROR)
6000 CONTINUE
  END IF
7000 CONTINUE
  RETURN
  END
  SUBROUTINE OUP TBB (KTYPE, KSURF)

```

C
C
C
C
C
C
C
C
C
C
C

4 MARCH 1986

OUP TBB PRINTS ENERGY DISTRIBUTION TABLE

***** CALLING PARAMETERS *****

KTYPE (INTEGER, PASSED) PARTICLE TYPE (1-10 SECONDARY, 11 PRIMARY)
KSURF (INTEGER, PASSED) 1 = TARGET FRONT SURFACE, 2 = BACK SURFACE

***** VARIABLE DECLARATIONS *****

INTEGER I, J, KOUNT, KSURF, KTYPE, NBIN, NBINLM, NENERG, NUM
REAL ADUMMY, WIDTH

DIMENSION KOUNT(20), NBIN(20)

ADUMMY IS NOT USED IN THIS SUBPROGRAM

COMMON /READIN/ ADUMMY(11), WIDTH(11)

ENERGY (NENERG), YIELD (NYIELD), DEPTH (NDEPTH), AND
ANGULAR (NANGLE) DISTRIBUTION TABLES.

KSUM, KSUMSQ = SUM AND SUM OF SQUARES OF YIELD FOR
CALCULATING MEAN YIELD AND STANDARD DEVIATION.

NPRYMS = TOTAL NUMBER OF PRIMARY PARTICLES.

COMMON /TABLES/ NENERG(100,11,2), NYIELD(21,10,2),

1 NDEPTH(21,10,2), XANGLE(21,21,11,2), FACNRM(11,2),

2 NANGLE(21,21,11,2), KSUM(10,2), KSUMSQ(10,2),

3 NPRYMS

COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

SAVE

ENERGY BIN DIMENSION IN ARRAY NENERG

DATA NBINLM /100/

200 FORMAT(1H0,33X,'EJECTED PARTICLE ENERGY SPECTRUM ('.G13.6,

1 ' E.V. PER CHANNEL)'/ 1H0,8X,'CHANNEL ',2X,20I5)

201 FORMAT(1H0,8X,'CHANNEL ',2X,20(1X,A4))

202 FORMAT(1H ,8X,'FREQUENCY',20I5)

401 FORMAT(I4)

402 FORMAT(I3,'+')

Print counts for first 20 channels

PRINT 200, WIDTH(KTYPE), (I,I=1,20)

PRINT 202, (NENERG(I,KTYPE,KSURF),I=1,20)

Beyond channel 20 print only non-zero channels

Last channel is count of "all greater than next-to-last"

NUM = 0

DO 2000 I=21,NBINLM

IF (NENERG(I,KTYPE,KSURF).LE.0) GO TO 1000

NUM = NUM + 1

KOUNT(NUM) = NENERG(I,KTYPE,KSURF)

IF (I.LT.NBINLM) ENCODE(4,401,NBIN(NUM))I

IF (I.GE.NBINLM) ENCODE(4,402,NBIN(NUM))I

1000 CONTINUE

IF ((NUM.LE.0).OR.((NUM.LT.20).AND.(I.LT.NBINLM))) GO TO 2000

PRINT 201, (NBIN(J),J=1,NUM)

PRINT 202, (KOUNT(J),J=1,NUM)

NUM = 0

2000 CONTINUE

RETURN

END

SUBROUTINE OOPTCC (KTYPE, DEPMAX)

23 JANUARY 1986

OPTCC PRINTS TABLES OF MAXIMUM PARTICLE DEPTHS, NUMBER OF
COLLISIONS, AND NUMBERS AND LENGTHS OF COLLISION SEQUENCES ADDED TO
OUTPUT FOR PROGRAM MARLOW VERSION OF 6 FEBRUARY 1982

```

      INTEGER I, J, K, KTYPE, L, LENSEQ, MXDEP, NCOL, NCOLIS, NROW
      REAL    DEPMAX
C          COLUMN AND ROW TOTALS FOR COLLISION SEQUENCE TABLE
      DIMENSION NCOL(15), NROW(5)
C          FRONT-SURFACE-SPUTTERED PARTICLE DATA
C          MXDEP = MAXIMUM DEPTH DISTRIBUTION TABLE
C          LENSEQ = NUMBER AND LENGTH OF REPLACEMENT SEQUENCES TABLE
C          NCOLIS = NUMBER OF COLLISIONS FROM DEEPEST POINT TABLE.
C          FIRST INDEX RUNS 1-21 FOR BIN VALUES 0-20.
      COMMON /FRSURF/ MXDEP(21,11), LENSEQ(15,5,11), NCOLIS(21,11)
      SAVE
C
201  FORMAT(1H0,26X,'DISTRIBUTION OF MAXIMUM DEPTHS OF SPUTTERED ',
1     'ATOMS (CHANNEL WIDTH',G13.4,')')
2     1H ,8X,'MAX. DEPTH',1X,21I5,'+')
202  FORMAT(1H ,8X,'FREQUENCY ',21I5)
203  FORMAT(/1H0,9X,'NO. OF ',25X,'NUMBER OF REPLACEMENTS IN SEQUENCE',
1     22X,'NUMBER OF '/
2     1H ,8X,'SEQUENCES',1X,15I5,3X,'PARTICLES'//
3     5(1H ,12X,I1,5X,15I5,5X,I5/),
4     1H0,10X,'TOTALS',2X,15I5)
204  FORMAT(1H0,39X,
1     'NUMBER OF COLLISIONS FROM MAXIMUM DEPTH TO SURFACE'//
2     1H ,8X,'COLLISIONS',4X,'0',20I5,'+')
C
      PRINT 201, DEPMAX, (I, I=1, 21)
      PRINT 202, (MXDEP(I, KTYPE), I=1, 21)
      DO 900 I=1, 5
          NROW(I) = 0
900  CONTINUE
      DO 1000 I=1, 5
          DO 1000 J=1, 15
              NROW(I) = NROW(I) + LENSEQ(J, I, KTYPE)
1000 CONTINUE
C          TO GET NUMBER OF PARTICLES PER ROW, MUST
C          DIVIDE NROW(I) BY NUMBER OF SEQUENCES PER
C          PARTICLE IN ITH ROW.
      DO 2000 I=2, 5
          NROW(I) = NROW(I)/I
2000 CONTINUE
      DO 2500 I=1, 15
          NCOL(I) = 0
2500 CONTINUE
      DO 3000 I=1, 15
          DO 3000 J=1, 5
              NCOL(I) = NCOL(I) + LENSEQ(I, J, KTYPE)
3000 CONTINUE
      PRINT 203, (I, I=1, 15), (J, (LENSEQ(K, J, KTYPE), K=1, 15),
1     NROW(J), J=1, 5), (NCOL(L), L=1, 15)
      PRINT 204, (I, I=1, 20)
      PRINT 202, (NCOLIS(J, KTYPE), J=1, 21)
      RETURN
      END
      SUBROUTINE PRTCLS (ISET, JFYL, LUDAT, KODE)
C
C 12 MARCH 1986
C
C One MARLOW 'POPDAT' file (except file's first record) is processed
C on each call to PRTCLS.
C PRTCLS CALCULATES NEW VALUES FOR EK (PARTICLE ENERGY) AND RCOS3

```

C ("DIRECTION" COSINE FROM NORMAL TO TARGET SURFACE) AND COUNTS
C PARTICLES FOR ENERGY (NENERG), YIELD (NYIELD), DEPTH (NDEPTH),
C AND POLAR AND AZIMUTHAL ANGLE (XANGLE) DISTRIBUTIONS.
C THIS VERSION OF PRTCLS (FOR PROGRAM MARPOP VERSION 2.0) READS AND
C PROCESSES RECORDS FROM FILE FT17F001 WRITTEN BY PROGRAM MARLOW
C VERSION OF 6 FEBRUARY 1982 AND CALLS SUBROUTINE FRONT TO TABULATE
C MAXIMUM DEPTH DISTRIBUTION AND NUMBERS AND LENGTHS OF DISPLACEMENT
C SEQUENCES.

C C C ***** CALLING PARAMETERS *****

C ISET (integer, passed)
C JFYL (integer, passed) Sequence number of current POPDAT data file.
C LUDAT (integer, passed) Logical unit from which POPDAT file is to be
C read.

C KODE (INTEGER, RETURNED) VALUES OF KODE RETURNED BY PRTCLS:
C 1 RECORDS READ AND DATA PROCESSED OK.
C 2 UNEXPECTED END-OF-FILE, NO DATA.
C 3 UNEXPECTED END-OF-FILE, PARTIAL DATA READ AND PROCESSED
C 4 READ ERROR, NO DATA PROCESSED
C 5 READ ERROR, PARTIAL DATA READ AND PROCESSED.
C 6 MAXIMUM NUMBER OF RECORDS READ AND PROCESSED. NO EXIT FLAG.

C TEMPORARY STORAGE FOR PARTICLE YIELD COUNTS

C DIMENSION KYELD(10000,10,2)

C COMMON /BINS/ POLDEG(0:40), GMU(0:40), DMU(40), NMUBIN,

1 AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN

C STORAGE FOR RECORD READ FROM FILE FT17F001

C DIMENSION INREC(8,16), XINREC(8,16)

C ADUMMY = ARRAY NOT USED IN THIS ROUTINE

C MAXRUN = NUMBER OF PRIMARY PARTICLES IN MARLOW RUN

C NTYPE = NUMBER OF TARGET PARTICLE TYPES

C COMMON /FSTREC/ ADUMMY(5), MAXRUN, NTYPE, BDUMMY(121)

C LIMITS ON NRUN - NUMBER OF PRIMARY PARTICLES TO PROCESS
C FROM EACH FT17FXXX FILE

C COMMON /LIMITS/ LIMRUN(10)

C SBND = SURFACE BINDING ENERGIES

C WIDTH = WIDTH OF ENERGY DISTRIBUTION BINS FOR (1-10)
C SECONDARY AND (11) PRIMARY PARTICLES

C DEPBIN = WIDTH OF BIN FOR DEPTH DISTRIBUTION

C DEPMAX = BIN WIDTH FOR MAXIMUM DEPTH DISTRIBUTION

C COMMON /READIN/ SBND(11), WIDTH(11), DEPBIN, DEPMAX

C ENERGY (NENERG), YIELD (NYIELD), DEPTH (NDEPTH), AND
C ANGULAR (NANGLE) DISTRIBUTION TABLES.

C KSUM, KSUMSQ = SUM AND SUM OF SQUARES OF YIELD FOR
C CALCULATING MEAN YIELD AND STANDARD DEVIATION.

C NPRYMS = TOTAL NUMBER OF PRIMARY PARTICLES.

C COMMON /TABLES/ NENERG(100,11,2), NYIELD(21,10,2),

1 NDEPTH(21,10,2), XANGLE(21,21,11,2), FACNRM(11,2),

2 NANGLE(21,21,11,2), KSUM(10,2), KSUMSQ(10,2),

3 NPRYMS

C COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

C EQUIVALENCE (INREC(1,1),XINREC(1,1))

C Save all values on RETURN from this subroutine.

C SAVE

C MAXIMUM NUMBER OF PRIMARY PARTICLES PERMITTED BY ARRAY
C KYELD

C DATA MXPTCL /10000/

```
C          MAXIMUM NUMBER OF BINS IN 'TABLES' COMMON BLOCK ARRAY
C          NENERG
C          DATA NBINLM /100/
C          TWOPI = 2*PI
C          DATA PI /3.141592653590/, TWOPI /6.283185307179/
```

```
C
C
201 FORMAT('0',7X,'***',I5,' PRIMARY PARTICLES EXCEEDS LIMIT OF',I5,
1      5X,' YIELD FROM EXCESS WILL NOT BE COUNTED')
202 FORMAT('0',7X,'*** NUMBER OF AZIMUTHAL ANGLE BINS,',I5,2X,
1      ' IS TOO LARGE. CHANGING TO MAXIMUM =',I5)
203 FORMAT(' BAD COSINE IN ROUTINE PRCLs=',G20.10,3X,' OFF BY',G20.10,
1      ', RESET TO 1 OR -1')
2      5X,' FILE',I3,', RECORD',I7,', PARTICLE',I3,' DIRECTION COSINES',
3      2(G20.10,', '),G20.10)
204 FORMAT(' MU OUT-OF-BOUNDS, SPECIFICATION SET',I3,
1      ', LOGICAL UNIT',I3,', RECORD',I8,', PARTICLE',I3,', OFF BY',
2      G12.4,', CHANGED TO',F3.0)
```

```
C
C          Zero arrays before processing data from first POPDAT file.
C
```

```
IF (JFYL.LE.1) THEN
DO 910 I=1,2
DO 910 J=1,11
DO 910 K=1,100
NENERG(K,J,I) = 0
910 CONTINUE
DO 920 I=1,2
DO 920 J=1,10
DO 920 K=1,21
NYIELD(K,J,I) = 0
920 CONTINUE
DO 930 I=1,2
DO 930 J=1,10
DO 930 K=1,21
NDEPTH(K,J,I) = 0
930 CONTINUE
DO 960 I=1,2
DO 960 J=1,10
KSUM(J,I) = 0
960 CONTINUE
DO 970 I=1,2
DO 970 J=1,10
KSUMSG(J,I) = 0
970 CONTINUE
DO 985 I=1,2
DO 985 J=1,11
DO 985 K=1,21
DO 985 L=1,21
NANGLE(L,K,J,I) = 0
985 CONTINUE
NPRYMS = 0
```

```
C
C          Initialize subroutine FRONT.
C
```

```
CALL FRINIT
END IF
```

```
C
C          Set binding energy for primary particle and zero array
C          KYELD before processing data from each POPDAT file.
C
```

```

C
SBND(11) = 0.0
DO 990 I=1,2
  DO 990 J=1,10
    DO 990 K=1,10000
      KYELD(K,J,I) = 0
990 CONTINUE
C
C
C      Set MAXRUN, the number of particles to be processed from
C      current POPDAT file.
C
IF ((LIMRUN(JFYI). GT. 0). AND. (LIMRUN(JFYI). LT. MAXRUN))
1  MAXRUN = LIMRUN(JFYI)
IF (MAXRUN. LE. MXPTCL) GO TO 2000
PRINT 201, MAXRUN, MXPTCL
MAXRUN = MXPTCL
2000 CONTINUE
AZMBIN = TWOPI / FLOAT(NAZBIN)
C
C
C      Each pass through DO 11000 I loop processes one record
C      from POPDAT file.
C
DO 11000 I=1,1000000
READ (LUDAT,END=12000,ERR=14000) INREC
C
C
C      Each pass through DO 10000 J loop processes data for one
C      particle.
C
DO 10000 J=1,16
C
C
C      INREC(B,J) (Marlowe variable NRUN) out-of-bounds is flag
C      that last particle has been processed.
C
IF ((INREC(B,J). LE. 0). OR. (INREC(B,J). GT. MAXRUN)) GO TO 16000
CALCULATE EK CORRECTED FOR SURFACE
BINDING ENERGY AND NEW Z-DIRECTION
COSINE. INCREMENT NENERG BIN IF
PARTICLE ESCAPES.
C
C      Extract particle type (11=primary)
C
KTYPE = MOD(INREC(7,J),256)
IF (MOD(INREC(6,J),2). EQ. 1) KTYPE=11
C
C
C      Recalculate particle energy as (EK*RCOS3**2) and process
C      if particle escapes (new energy > SBND)
C
TEMP = XINREC(1,J)*XINREC(4,J)*XINREC(4,J) - SBND(KTYPE)
IF (TEMP. LE. 0.0) GO TO 10000
C
C
C      Calculate new direction cosine (ZMU) of escaped particle.
C      If calculated ZMU is invalid cosine, adjust it and write
C      message on file INFO.
C
ZMU = SQRT(TEMP/(XINREC(1,J)-SBND(KTYPE)))
IF ((ZMU. GT. 1.0). OR. (ZMU. LT. 0.0)) THEN
  IF (ZMU. GT. 1.0) THEN
    ERRSIZ = ZMU - 1.0
    ZMU = 1.0
  ELSE

```

```
ERRSIZ = ZMU
ZMU = 0.0
END IF
WRITE (LUINFO,204) ISET, LUDAT, I, J, ERRSIZ, ZMU
END IF
```

```
Reset XINREC(1,J) to energy of escaped particle.
```

```
XINREC(1,J) = XINREC(1,J) - SBND(KTYPE)
```

```
Extract KARMA from INREC(6,j) and divide by 10 to
determine surface from which particle escapes (KSURF=1
for front, 2 for rear). KARMA is expected to be in
range 10-29. Increment energy distribution count.
```

```
KSURF = MOD(INREC(6,J),65536)/10
NENBIN = 1 + IFIX(XINREC(1,J)/WIDTH(KTYPE))
IF (NENBIN.GT.NBINLM) NENBIN=NBINLM
NENERG(NENBIN,KTYPE,KSURF) = NENERG(NENBIN,KTYPE,KSURF) + 1
```

```
If secondary particle, increment yield count, extract
original depth of particle, and increment depth
distribution table.
```

```
IF (KTYPE.EQ.11) GO TO 5000
KYELD(INREC(8,J),KTYPE,KSURF) =
KYELD(INREC(8,J),KTYPE,KSURF) + 1
KDEP = 1 + IFIX(AMOD(XINREC(5,J),100.0)/DEPBIN)
IF (KDEP.GT.21) KDEP=21
NDEPTH(KDEP,KTYPE,KSURF) = NDEPTH(KDEP,KTYPE,KSURF) + 1
CONTINUE
```

```
Calculate indices of mu and phi bins and increment
angular distribution table
```

```
DO 6000 KMU=0,NMUBIN
IF (ZMU.GT.GMU(KMU)) GO TO 6000
GO TO 7000
CONTINUE
KMU = NMUBIN + 1
CONTINUE
```

```
FIND QUADRANT OF AZIMUTHAL ANGLE AND
CALCULATE ANGLE IN RADIANS
```

```
COSPHI = XINREC(2,J)/SQRT((XINREC(2,J)*XINREC(2,J))+
(XINREC(3,J)*XINREC(3,J)))
IF ((COSPHI.GT.1.0).OR.(COSPHI.LT.(-1.0))) THEN
DELTA = ABS(COSPHI) - 1.0
WRITE (LUINFO,203) COSPHI, DELTA, LUDAT, I, J,
XINREC(2,J), XINREC(3,J), XINREC(4,J)
IF (COSPHI.GT.1.0) THEN
COSPHI = 1.0
ELSE
COSPHI = -1.0
END IF
END IF
AZIMTH = ACOS(COSPHI)
IF (XINREC(3,J).LT.0.0) AZIMTH = TWOPI - AZIMTH
IF (AZIMTH.LE.AZMRAD(NAZBIN)) THEN
PHI = AZIMTH
ELSE
```

```

      PHI = AZIMTH - TWOPI
      END IF
      DO 8000 KPHI=0,NAZBIN
        IF (PHI.GT.AZMRAD(KPHI))GO TO 8000
        GO TO 9000
8000    CONTINUE
        KPHI = NAZBIN + 1
9000    CONTINUE
        IF ((KMU.GT.0).AND.(KMU.LE.NMUBIN).AND.(KPHI.GT.0).AND.
1         (KPHI.LE.NAZBIN)) NANGLE(KMU,KPHI,KTYPE,KSURF) = 1 +
2         NANGLE(KMU,KPHI,KTYPE,KSURF)
C         TABULATE MAXIMUM DEPTH, NUMBER AND LENGTH OF
C         DISPLACEMENT SEQUENCES AND NUMBER OF
C         COLLISIONS DATA FOR PARTICLES LEAVING FRONT
C         SURFACE
        IF (KSURF.EQ.1) CALL FRONT (KTYPE, XINREC(5,J), DEPMAX,
1         INREC(7,J), INREC(6,J))
10000   CONTINUE
11000   CONTINUE
        KODE = 6
        GO TO 17000
12000   CONTINUE
        IF (I.GT.1) GO TO 13000
        KODE = 2
        RETURN
C
13000   CONTINUE
        KODE = 3
        GO TO 17000
14000   CONTINUE
        IF (I.GT.1) GO TO 15000
        KODE = 4
        RETURN
C
15000   CONTINUE
        KODE = 5
        GO TO 17000
16000   CONTINUE
        KODE = 1
17000   CONTINUE
C
C         ADD YIELD DATA FROM THIS FILE TO
C         NYIELD, KSUM, AND KSUMSQ
        DO 18000 KSURF=1,2
          DO 18000 KTYPE=1,NTYPE
            DO 18000 I=1,MAXRUN
              KSUM(KTYPE,KSURF) = KSUM(KTYPE,KSURF) +
1              KYELD(I,KTYPE,KSURF)
              KSUMSQ(KTYPE,KSURF) = KSUMSQ(KTYPE,KSURF) +
1              (KYELD(I,KTYPE,KSURF) * KYELD(I,KTYPE,KSURF))
              IBIN = KYELD(I,KTYPE,KSURF) + 1
              IF (IBIN.GT.21) IBIN=21
              NYIELD(IBIN,KTYPE,KSURF) = NYIELD(IBIN,KTYPE,KSURF) + 1
18000   CONTINUE
          NPRYMS = NPRYMS + MAXRUN
          RETURN
        END
        SUBROUTINE SUMRYZ
C
C 22 MAY 1986
C

```

```

COMMON /BINS/ POLDEG(0:40), QMU(0:40), DMU(40), NMUBIN,
1      AZMDEG(0:40), AZMRAD(0:40), DAZBIN(40), NAZBIN
COMMON /FORFAC/ FACDIV(11,2)
COMMON /FSTREC/ ADUMMY(5), MAXRUN, NTYPE, BDUMMY(121)
COMMON /MXDMNS/ MXSURF, MXTYPE, MXPOLR, MXAZM
COMMON /SPECS/ XNORML
COMMON /TABLES/ NENERG(100,11,2), NYIELD(21,10,2),
1      NDEPTH(21,10,2), NRMLZD(21,21,11,2), FACNRM(11,2),
2      NANGLE(21,21,11,2), KSUM(10,2), KSUMSQ(10,2),
3      NPRYMS
COMMON /UNITS/ LUSPEC, LUPRT, LUINFO

```

```

C
SAVE

```

```

C
C
C      CALCULATE ROW AND COLUMN SUMS FOR
      PARTICLE DIRECTION TABLE

```

```

DO 7000 KSURF=1, MXSURF
  DO 7000 KTYPE=1, MXTYPE
    IF ((KTYPE. LE. NTYPE). OR. (KTYPE. EQ. MXTYPE)) THEN
      DO 2000 I=1, NMUBIN
        NANGSM = 0
        DO 1500 J=1, NAZBIN
          NANGSM = NANGSM + NANGLE(I, J, KTYPE, KSURF)
1500      CONTINUE
          NANGLE(I, MXAZM, KTYPE, KSURF) = NANGSM
2000      CONTINUE
        DO 4000 I=1, NAZBIN
          NANGSM = 0
          DO 3500 J=1, NMUBIN
            NANGSM = NANGSM + NANGLE(J, I, KTYPE, KSURF)
3500      CONTINUE
            NANGLE(MXPOLR, I, KTYPE, KSURF) = NANGSM
4000      CONTINUE
          NCROSM = 0
          DO 6200 I=1, NMUBIN
            NCROSM = NCROSM + NANGLE(I, MXAZM, KTYPE, KSURF)
6200      CONTINUE
            NANGLE(MXPOLR, MXAZM, KTYPE, KSURF) = NCROSM
          END IF
        7000 CONTINUE

```

```

C
C      Normalize target values in array NRMLZD by factor
C      (average yield*10000) / total of column with largest Mu
C

```

```

DO 7500 I=1, MXSURF
  DO 7500 J=1, MXTYPE
    DO 7500 K=1, MXAZM
      DO 7500 L=1, MXPOLR
        NRMLZD(L, K, J, I) = 0
7500 CONTINUE
  DO 9000 KSURF=1, MXSURF
    DO 9000 KTYPE=1, NTYPE
      IF (NANGLE(NMUBIN, MXAZM, KTYPE, KSURF). GT. 0) THEN
        FACDIV(KTYPE, KSURF) = FLOAT(
1      NANGLE(NMUBIN, MXAZM, KTYPE, KSURF)) / DMU(NMUBIN)
        FACTOR = (FLOAT(KSUM(KTYPE, KSURF)) / FLOAT(NPRYMS)) *
1      XNORML / FACDIV(KTYPE, KSURF)
        FACNRM(KTYPE, KSURF) = FACTOR
        DO 8000 I=1, NAZBIN
          DO 8000 J=1, NMUBIN

```



```
NRMLZD(J, I, KTYPE, KSURF) = NINT(FACTOR * FLOAT(
1
8000 CONTINUE
DO 8200 I=1, NAZBIN
1
8200 NRMLZD(MXPOLR, I, KTYPE, KSURF) = NINT(FACTOR * (
CONTINUE
DO 8400 I=1, NMUBIN
1
8400 NRMLZD(I, MXAZM, KTYPE, KSURF) = NINT(FACTOR * (
CONTINUE
END IF
9000 CONTINUE
RETURN
END
```

END

7-89

Dotic