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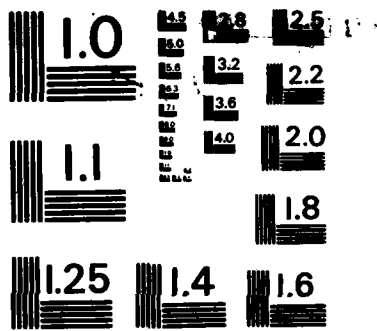
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COMPUTER-ASSISTED STATISTICAL ANALYSIS.

III. The *Drosophila melanogaster* Sex-Linked Recessive Lethal Assay

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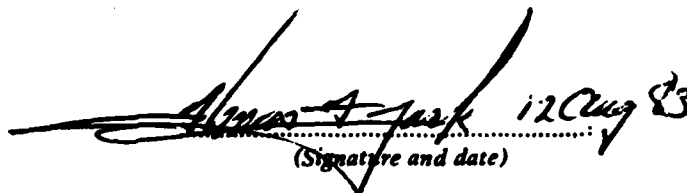
Computer-Assisted Statistical Analysis. III. The Drosophila melanogaster Sex-Linked Recessive Lethal Assay (Toxicology Series 61)--Powers and Jederberg

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PREFACE

This technical note is the third in a series on the utilization of the computer to assist mutagenicity testing at Letterman Army Institute of Research. In this report we detail three functions: searching of a specific file to summarize the data; calculating the mean mutation frequency with its variance expressed as percentages; and using a statistical program for analysis of the data. Using the program in the Drosophila melanogaster Sex-Linked Recessive Lethal Assay provides an easy manner to view the data and perform statistical analyses.

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Computer-Assisted Statistical Analysis III.
The Drosophila melanogaster Sex-Linked
Recessive Lethal Assay

The Drosophila melanogaster Sex-Linked Recessive Lethal Assay requires the testing of 8000 to 10,000 X-chromosomes for each compound and concurrent negative control in addition to 500 X-chromosomes for a positive control. A mass of data must be processed. Therefore, two computer programs were developed to summarize the results. By totalling the number of lethal mutations and non-lethal mutations for each brood of each replicate yielding the total number of lethal mutations and number of tests, this first program calculates the percentage mean mutation frequency and its variance. The second program performs the statistical analysis using the BMDP statistical softwareTM package by testing the null hypothesis if the number of lethal mutations resulting from the compound is non-significantly from different from the number resulting from the negative control.

DESCRIPTION OF THE PROGRAM

The program DRUNSUM.FR (Appendix A) searches the data file and summarizes the data accumulated in the program DROSTOXDATA.FR (1) for a specified subgroup of a given run number selected by the user. This program uses the subroutines RDDROSDATA.FR and STRING.FR (Appendix B and C) to read the data sequentially and accumulates the values necessary for the number of lethal, non-lethal, and total tests performed for each brood, and grand total for the specified run number. The program also calculates the mean mutation frequency and its variance and displays them as percentages. The detection of three error conditions is provided to see (a) if more than 25 records are present for the specified subgroup within the run; (b) if the file name under which the data from the DROSTOXDATA are stored is present; (c) if the specified subgroup is present.

The program DRUNSUM.FR (2) requests the following information from the user (Figure 1):

- The name of the file to be searched.
- The run number to be searched.
- The group designator of the male code number (this specifies if the male was from the control, positive, or test compound).

The second program is part of the BMDP statistical softwareTM (3). Instructions for its use are given in this statistical package. This package analyzes the data in a 2 x 2 table and two statistics are used, the Yates corrected chi-squares and Fisher's Exact test. The Yates correction is intended to improve the approximation to the chi-square distribution. The Fisher's Exact test is computed when the

minimum value is less than 20. Both 1-tail and 2-tail probabilities are tested. However the 2-tail probability for Fisher's Exact test should be used.

The program requests the following information from the user:

- Enter the number of lethal, lethal mutations and non-lethal mutations resulting from the negative control.

- Enter the number of lethal mutations and the number of non-lethal mutations resulting from the concurrent test compound.

The computer will then type the values and probabilities associated with the Yates corrected chi-square and the Fisher's Exact test.

COMMENT

By utilizing these programs and the subroutines presented in this report, the raw data may be summarized and statistical analysis performed.

CONCLUSION

None

RECOMMENDATION

None

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1. Jederberg WW, Powers NR. Computer-assisted data storage and retrieval in mutagenicity testing II. The Drosophila melanogaster sex-linked recessive lethal assay. Technical Note No. 82-42-TN. Presidio of San Francisco, CA: Letterman Army Institute of Research, 1982.
2. LAIR SOP-OP-STX-63, Execution of FORTRAN V Program DRUN: 14 June 1982.
3. Brown BM. Frequency tables. In: Dixon WJ, Brown MB, Engleman L, Frane JW, Hill MA, Jennrich RI, Toporek JD, eds. BMDP statistical software, Berkeley: University of California Press, 1981: 1943-161.

SAMPLE RUN OF DRUN

) DRUN

The computer will type:

NAME OF FILE TO BE SEARCHED: DROSDATA

The computer will type:

****DROSOPHILA RECORD SEARCH AND SUMMARY****
 PLEASE ENTER RUN NUMBER TO BE SUMMARIZED: 24

The computer will type:

ENTER GROUP TO BE SEARCHED FOR:
 (T1, T2, P1 or C1 ETC. ALLOWED): C1

The computer will type:

SUMMARY TOTALS
 RUN NUMBER: 24
 GROUP TYPE: C1

BROOD DATA

LETHALS:	0	1	0	1
NONLETHALS:	596	524	547	547

GRAND TOTAL LETHALS: 2

GRAND TOTAL NONLETHALS: 2214

TOTAL RECORDS PROCESSED: 25

TOTAL TESTS REPRESENTED: 2216

MEAN MUTATION RATE: .08000%, WITH VAR. .07667%

The computer will then display ")", the user may then return to
 the program for the next file to be searched or log off.

Figure 1. Sample Run of DRUN
 (Underlined entries are sample user inputs)

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APPENDIX A

```

C *** PROGRAM NAME:      DRUNSUM.FR
C *** WRITTEN BY:       WARREN W. JEDERBERG
C *** PURPOSE:         TO SUMMARIZE THE DATA FROM SPECIFIED
C ***                   RUNS IN DROSDATA.
C *** THE SUBROUTINE RDDROSDATA IS USED TO READ EACH RECORD
C *** VARIABLE DEFINITIONS
C   N   =NUMBER OF RECORDS
C   FBL =FIRST BROOD LETHALS
C   FBNL=FIRST BROOD NONLETHALS
C   STL =SECOND BROOD LETHALS
C   STNL=SECOND BROOD NONLETHALS
C   TTL =THIRD BROOD LETHALS
C   TTNL=THIRD BROOD NONLETHALS
C   FTL =FOURTH BROOD LETHALS
C   FTNL=FOURTH BROOD NONLETHALS
C
C   GTT = GRAND TOTAL TESTS
C   GTL =GRAND TOTAL LETHALS
C   GTNL=GRAND TOTAL NONLETHALS
C   TMR =TOTAL MUTATION RATE
C   V1MR=ACCUMULATED MUTATION RATES SQUARED
C   NNR =DESIGNATED RUN NUMBER (USER INPUT AT RUN TIME)
C   NS  =SUBGROUP DESIGNATOR   (USER INPUT AT RUN TIME)
C   RMR =MEAN MUTATION RATE
C
C   VARIABLES MENTIONED IN THE COMMON STATEMENT ARE DEFINED IN
C   THE FORTRAN V PROGRAM DROSTOXDATA.
C   INTEGER NR,MNUM,CNAM,BFF,BFL,BFNL,BSF,BSL,BSNL,BTF,BTL,BTNL,
1  BLF,BLL,BLNL,TF,SL,ML,TL,TNL,TT,SFNAM

C   DIMENSION NS(2),SFNAM(20)
C   REAL NR

C   COMMON/DREC/NR,MNUM(6),CNAM(6),BFF,BFL,BFNL,BSF,BSL,BSNL,BTF,BTL,
1  BTNL,BLF,BLL,BLNL,TF,SL,ML,TL,TNL,TT,MR

```

APPENDIX A
(continued)

LOGICAL EOF

```
C *** SET ACCUMULATORS TO 0.00
N=0
FBL=0
FBNL=0
STL=0
STNL=0
TTL=0
TTNL=0
FTL=0
FTNL=0
GTT=0
GTL=0
GTNL=0
TMR=0
VIMR=0

C *** INITIALIZE EOF
EOF=.FALSE.

C *** GET FILE NAME TO CHECK FOR THE DATA
WRITE (10,220)
220 FORMAT ('NAME OF FILE TO BE SEARCHED: ',2)
READ (11,210) SFNAM
210 FORMAT (20A2)
CALL STRING (SFNAM,20)
OPEN 1,SFNAM,ATT="SIB",ERR=100
READ (1,5)
5 FORMAT (/)
GO TO 120
100 WRITE (10,6) SFNAM
6 FORMAT (//,"**** ERROR **** ERROR **** ERROR ****",/,
14X,"FILE ",20A2," NOT FOUND . . . .")
STOP
120 CONTINUE
C *** PRINT HEADER AND GET RUN NUMBER
WRITE (10,1)
1 FORMAT (/,10X,"**** DROSOPHILA RECORD SEARCH AND SUMMARY ****",/,
110X,"PLEASE ENTER RUN NUMBER TO BE SUMMARIZED: ",2)

ACCEPT MNR
```

APPENDIX A
(continued)

```

C *** GET GROUP DESIGNATOR
  99 CONTINUE
     WRITE (10,3)
     3 FORMAT (/,10X,"ENTER GROUP DESIGNATOR TO BE SEARCHED FOR :",/,
110X,"(T1,T2,P1,OR C1 ETC. ALLOWED):",Z)
     READ (11,4)(NS(I),I=1,2)
     4 FORMAT (2A1)
     IF (NS(1).NE."C ".AND.NS(1).NE."P ".AND.NS(1).NE."T ") GO TO 99
C *** READ DATA RECORDS
  140 CALL RDDROSDATA (EOF)
     IF (EOF) GO TO 225
     IF (NR.NE.NNR) GO TO 140
     IF (NS(1).NE.MNUM(1)) GO TO 140
     IF (NS(2).NE.MNUM(2)) GO TO 140
     N=N+1
     IF (N.GT.25) GO TO 500
     FBL=FBL+BFL
     FBNL=FBNL+BFNL
     STL=STL+BSL
     STNL=STNL+BSNL
     TTL=TTL+BTL
     TTNL=TTNL+BTNL
     FTL=FTL+BLL
     FTNL=FTNL+BLNL
     GTL=GTL+TL
     GTNL=GTNL+TNL
     GTT=GTT+TT
     TMR=TMR+MR
     V1MR=V1MR+(MR**2)
     GO TO 140

C *** IF N>25 ERROR
  500 CONTINUE
     WRITE (10,7)
     7 FORMAT (/,"*** ERROR *** ERROR *** ERROR ***",/,
14X,"MORE THAN 25 RECORDS FOUND . . . ")
     STOP

```

APPENDIX A
(continued)

```
225 CONTINUE
C *** IF NO SPECIFIED RECORDS FOUND
  IF(N.NE.O.O) GO TO 300
  WRITE (10,9) NNR,NS,SFNAME
  9 FORMAT (/,"*** ERROR *** ERROR *** ERROR ***",/,
    14X,"RUN NUMBER:",2X,I4,1X,"GROUP:",1X,2A1,/,
    24X,"NOT FOUND IN ",1X,20A2," . . .")
  STOP

300 CONTINUE
C *** CALCULATE MUTATION RATE AND VARIANCE
  RMR=(TMR/N)
  VMR=(V1MR-((TMR**2.0)/N))/(N-1)

C *** WRITE OUT TOTALS .
  WRITE (10,8) NNR,NS,FBL,STL,TTL,FTL,FBNL,STNL,TTNL,FTNL,
  1GTL,GTNL,N,GTT,RMR,VMR

  8 FORMAT (20X," SUMMARY TOTALS:",/,
    110X,"RUN" NUMBER:",2X,I4,/,
    210X,"GROUP TYPE:",2X,2A1,/,
    320X,"BROOD DATA:",/,
    410X,"LETHALS:",4(5X,I3),/,
    57X,"NONLETHALS:",4(5X,I3),//,
    610X,"GRAND TOTAL LETHALS:",5X,I5,/,
    77X,"GRAND TOTAL NONLETHALS:",5X,I5,//,
    810X,"TOTAL RECORDS PROCESSED:",5X,I2,//,
    910X,"TOTAL TESTS REPRESENTED:",5X,I5,/,
    110X,"MEAN MUTATION RATE:",1X,F10.5,"% WITH VAR.",1X,F10.5,"%",/)
  CLOSE 1
  END
```

APPENDIX B

```
COMPILER NOSTACK
C *** PROGRAM NAME:          RDDROSDATA.FR
C *** WRITTEN BY:           WARREN JEDERBERG
C *** PURPOSE:              TO READ DATA FOR MAKING REPORTS
C ***                       FOR THE SLRL-DROSOPHILA ASSAY
```

```
SUBROUTINE RDDROSDATA (EOF)
COMMON/DREC/NR,MNUM(6),CNAM(6),BFF,BFL,BFNL,BSF,BSL,BSNL,
1BTF,BTL,BTNL,BLF,BLL,BLNL,TF,SL,ML,TL,TNL,TT,MR
```

```
INTEGER NR,MNUM,CNAM,BFF,BFL,BFNL,BSF,BSL,BSNL,BTF,BTL,BTNL,
1BLF,BLL,BLNL,TF,SL,ML,TL,TNL,TT
```

```
REAL MR
```

```
LOGICAL EOF
```

```
READ (1,100, END= 120) NR,MNUM,CNAM,BFF,BFL,BFNL,BSF,BSL,BSNL,
1BTF,BTL,BTNL,BLF,BLL,BLNL,TF,SL,ML,TL,TNL,TT,MR
```

```
100 FORMAT(I4,1X,6A1,1X,6A1,4(1X,3I2),1X,I2,1X,3I2,
12(1X,I3),1X,F6.2)
RETURN
```

```
120 EOF = .TRUE.
RETURN
END
```

APPENDIX C

```
C --- TO CONVERT FILE NAMES TO USE IN OPEN STATEMENTS
SUBROUTINE STRING(LINE,LLEN)
COMPILER STATIC
DIMENSION LINE (LLEN)
DO 100 I=1,LLEN
IF (LINE(I).EQ." ") LINE(I)=0
IF (FLD(LINE(I),9,16).EQ.FLD(" ",9,16))FLD(LINE(I),9,16)=0
100 CONTINUE
RETURN
END
```

END

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