

**APPENDIX C**  
**ORTEP-III FORTRAN SOURCE CODE LISTING**



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C *****
C ORTEP-III: Oak Ridge Thermal Ellipsoid Plot Program
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C Oak Ridge National Laboratory
C Version 1.0 April 1, 1996
C
C Send comments, questions, problems, suggestions, etc. to
C ortep@ornl.gov
C *****
C
PROGRAM ORTEP
REAL*8 TD
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
COMMON /QUEUE/ NED,NQUE,NEXT,NBACK,INQ,QUE(96),hque(96)
CHARACTER*73 INQ,QUE,hque
common /ns/ npf,ndraw,norient,nvar
logical tmpopn

C *** Drawing Output Options
C *** ndraw=0: no drawing output
C *** ndraw=1: screen output
C *** ndraw=2: Postscript file output
C *** ndraw=3: HPGl file output

C *** Logical Unit Numbers ***
C *** 15,16 are used in subroutine EDITR
C *** 18 (variable iu) is used in subroutine PRELIM
C *** NOUT is set in subroutine UINPUT
IN=3
NED=7
NSR=8
NPF=10

call uinput(in,nout)

2 CALL PRIME

C *** open ORTEP scratch file ***
C *** if already open, close it first ***
inquire(NSR,opened=tmpopn)
if (tmpopn) close(NSR)
open(NSR,status='scratch',form='unformatted')

C *** open a temporary file - needed by the editor ***
C *** if already open, close it first ***
inquire(NED,opened=tmpopn)

if (tmpopn) close(NED)
open(NED,status='scratch')

C *****
C
**** READ JOB TITLE CARD *****
READ (IN,4)(TITLE(I),I=1,18)
4 FORMAT(18A4)
write (NED,4)(TITLE(I),I=1,18)
IF (NOUT.GE.0)
&WRITE (NOUT,6)(TITLE(I),I=1,18)
5 FORMAT(1H0,10X,18A4)
6 FORMAT(1H1,10X,18A4)
CALL PRELIM
IF (NOUT.GE.0)
&WRITE (NOUT,6)(TITLE(I),I=1,18)
**** LOAD INSTRUCTION QUE *****
NQUE=0
2005 NQUE=NQUE+1
2010 READ (IN,2012,END=2015,ERR=3000) QUE(NQUE)
if (que(nque)(1:1).eq.'#') go to 2010
hque(nque)=que(nque)
if (que(nque)(4:9).eq.' -2') go to 2020
2012 FORMAT(A72)
IF (NQUE.LT.96) GO TO 2005
GO TO 2020
2015 NQUE=NQUE-1
**** REPOSITION TO POINT BEFORE EOF *****
BACKSPACE IN
2020 NBACK=NQUE
NEXT=1
ISAVE=0
GO TO 507
7 ISAVE=0
**** ZERO AIN ARRAY *****
8 DO 10 J=1,140
10 AIN(J)=0.
11 FORMAT(1H0,4X,17H(((( INSTRUCTION,I5,6H )))))
12 FORMAT(I3,I6,7F9.0)
13 FORMAT(I3,I6,7D15.8)
14 FORMAT(1H,9X,7D15.7)
**** READ NEW INSTRUCTION CARD *****
NCD=0
NI=-6
16 NI=NI+7
N2=NI+6
IF (ISAVE) 22,18,18
18 INQ=QUE(NEXT)
NEXT=NEXT+1
READ (INQ,12) IIC,NF,(AIN(I),I=NI,N2)
IF (ISAVE) 24,24,20
20 WRITE (NSR) IIC,NF,(AIN(I),I=NI,N2)
GO TO 24
22 READ (NSR) IIC,NF,(AIN(I),I=NI,N2)
IF (IIC) 7,24,24
24 IF (NI-1) 26,26,30
26 IF (NOUT.GE.0)
&WRITE (NOUT,11) NF
NPF=NPF
IF (NPF) 28,8,30
c *** run editor?
28 call go2edr
if (next.lt.nque) go to 8

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IF(NF1+2,2,2,3000
30 CONTINUE
CCC IF (NOUT.GE.0)
CCC &WRITE (NOUT,14)(AIN(I),I=NI,N2)
32 IIC=IIC+1
GO TO (90,16,38,50),IIC
33 FORMAT(I3,6X,5I3,8F6.0)
34 FORMAT(6I3,8E12.5)
35 FORMAT(1H,11X,5I3,8F11.5)
C ***** READ FORMAT 2 TRAILER CARDS *****
38 NCD=NCD+1
IF (ISAVE)44,40,40
40 INQ=QUE(NEXT)
NEXT=NEXT+1
READ (INQ,33)IIC,(KD(I,NCD),I=1,5),(CD(I,NCD),I=1,8)
IF(ISAVE)46,46,42
42 WRITE (NSR)IIC,(KD(I,NCD),I=1,5),(CD(I,NCD),I=1,8)
GO TO 46
44 READ (NSR)IIC,(KD(I,NCD),I=1,5),(CD(I,NCD),I=1,8)
46 IF (NOUT.GE.0)
&WRITE (NOUT,35)(KD(I,NCD),I=1,5),(CD(I,NCD),I=1,8)
GO TO 32
C ***** READ FORMAT 3 TRAILER CARD *****
50 IF (ISAVE)52,54,54
52 READ (NSR)(TITLE2(I),I=1,18)
GO TO 55
54 INQ=QUE(NEXT)
NEXT=NEXT+1
READ (INQ,4)(TITLE2(I),I=1,18)
55 IF (NOUT.GE.0)
&WRITE (NOUT,5)(TITLE2(I),I=1,18)
IF (ISAVE)90,90,56
56 WRITE (NSR)(TITLE2(I),I=1,18)
C ***** EXECUTE INSTRUCTION *****
90 NJ=NF1/100
NJ2=NF1-NJ*100
NJ3=MOD(NJ2,10)
IF (NJ-12)98,92,92
92 CALL SPARE(NF1)
IF (NG)94,8,94
94 CALL ERENT(0.DO,NF1)
GO TO 8
C *****BRANCH TABLE FOR FUNCTION TYPES*****
98 GO TO(100,200,300,400,500,600,700,800,900,1000,1100),NJ
C *****100 INSTRUCTIONS-STRUCTURE ANALYSIS FUNCTIONS*****
100 GO TO (101,101,104,104,101,101,94),NJ2
101 CALL SEARC
GO TO 8
C ***** ANISOTROPIC TEMP FACTOR OUTPUT *****
104 DO 164 I=1,NATOM
IF (MOD(I,14)-1)134,114,134
114 IF (NOUT.GE.0)
&WRITE (NOUT,6)(TITLE(J),J=1,18)
IF (NOUT.GE.0)
&WRITE (NOUT,129)
129 FORMAT(1H010X,4HATOM3X,16HRMS DISPLACEMENT3X,31HROW VECTORS, BASED
1 ON REFERENCE17X,29HPROBABILITY COVARIANCE MATRIX)
134 TD=55501.+FLOAT(1)*100000.
CALL PAXES(TD,-3)
IF (NG)144,154,144
144 CALL ERENT(TD,104)
149 FORMAT(1H0,10X,A6,F10.6,6X,3F12.7,10X,3F12.7)
154 IF (NOUT.GE.0)
&WRITE (NOUT,149)CHEM(I),RMS(1),(PAC(J,1),J=1,3),(Q(J,1)
1,J=1,3)
164 IF (NOUT.GE.0)
&WRITE (NOUT,159)(RMS(K),(PAC(J,K),J=1,3),(Q(J,K),J=1,3)
1,K=2,3)
159 FORMAT(1H,16X,F10.6,6X,3F12.7,10X,3F12.7)
GO TO 8
C *****200 INSTRUCTIONS-PLOTTER CONTROL FUNCTIONS*****
200 CALL F200
GO TO 8
C *****300 INSTRUCTIONS-DRAWING CONTROL FUNCTIONS*****
300 GO TO (301,302,303,304,94),NJ2
C *****PLOT DIMENSIONS*****
301 IF(AIN(1))321,321,311
311 XLNG(1)=AIN(1)
321 IF(AIN(2))341,341,331
331 XLNG(2)=AIN(2)
341 IF(AIN(3))361,351,351
351 VLEW=AIN(3)
361 IF(AIN(4))381,381,371
371 BRDR=AIN(4)
381 IF (NOUT.GE.0)
&WRITE (NOUT,389)XLNG(1),XLNG(2),BRDR
389 FORMAT(1H010X,11HPLOT LIMITSF6.2,3H BYF6.2,15H IN. INCLUDINGF6.2,
112H IN. MARGIN)
391 IF (NOUT.GE.0)
&WRITE (NOUT,399)VIEW
399 FORMAT(1H,10X,13HVIEW DISTANCE,F7.3,7H INCHES)
GO TO 8
C *****LEGEND ROTATION*****
302 THETA=AIN(1)
TI=THETA*.01745329252
COSTH=COS(TI)
SINTH=SIN(TI)
DO 312 J=1,9
312 SYMB(J,1)=0.
SYMB(1,1)=COSTH
SYMB(2,2)=COSTH
SYMB(3,3)=1.
SYMB(1,2)=-SINTH
SYMB(2,1)=SINTH
IF (NOUT.GE.0)
&WRITE (NOUT,319)THETA
319 FORMAT(1H010X,44HREGULAR TITLE AND SYMBOL ROTATION IN DEGREESF8.2)
GO TO 8
C ***** RETRACE DISPLACEMENT *****
303 DISP=AIN(1)
IF (NOUT.GE.0)
&WRITE (NOUT,313)DISP
313 FORMAT(1H0,10X,22HRETRACE DISPLACEMENT =,F7.4,5H INCH)
GO TO 8
C ***** change resolution (smoothness) of ellipses *****
304 res(1)=AIN(1)*.75
res(2)=.5*res(1)
res(3)=.25*res(2)
GO TO 8
C *****400 INSTRUCTIONS-ATOM LIST FUNCTIONS*****
400 GO TO (401,401,401,401,401,401,490,94,410,
1 401,401,401,401,401,94),NJ2

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401 CALL F400
GO TO 490
410 LATM=0
DO 420 I=1,500
  ATOMID(I)=0.
DO 420 J=1,73
  420 ATOMS(J,I)=0.
490 IF (LATM).8.491
491 IF (NOUT.GE.0)
&WRITE (NOUT,499)(ATOMID(I), I=1,LATM)
499 FORMAT(IH0,10X,23HCONTENTS OF ATOMS ARRAY/(15X,10F10.0))
GO TO 8
C *****500 INSTRUCTIONS-CARTESIAN COORDINATE SYSTEM FUNCTIONS*****
500 CALL F500
IF (NOUT.GE.0)
&WRITE (NOUT,503)(ORGN(J),J=1,3)
503 FORMAT(IH0,10X, 44HORIGIN FOR PROJECTION AXIS IN CRYSTAL COORD.
1,3F15.6)
IF (NJ3-3)507,539,504
504 IF (NJ3-6)601,507,601
507 IF (NOUT.GE.0)
&WRITE (NOUT,529)
IF (NOUT.GE.0)
&WRITE (NOUT,519)((REFV(J,I), I=1,3), (AAREV(J,I), I=1,3), J
1=1,3)
GO TO 8
509 FORMAT(IH010X, 49HORTHONORMAL WORKING VECTORS BASED ON CRYSTAL AXE
1S18X,33HPOST-FACTOR TRANSFORMATION MATRIX/16X,8HX VECTOR8X,8HY VEC
2TOR8X,8HZ VECTOR)
519 FORMAT(IH ,10X,3E16.7,8X,3E16.7)
529 FORMAT(IH010X,51HORTHONORMAL REFERENCE VECTORS BASED ON CRYSTAL AX
1ES16X,33HPOST-FACTOR TRANSFORMATION MATRIX/16X,8HX VECTOR8X,8HY VE
2CTOR8X,8HZ VECTOR)
539 IF (NOUT.GE.0)
&WRITE (NOUT,509)
IF (NOUT.GE.0)
&WRITE (NOUT,519)((WRKV(J,I), I=1,3), (AARWK(J,I), I=1,3), J
1=1,3)
GO TO 8
C *****600 INSTRUCTIONS-PLOT CENTERING FUNCTIONS*****
600 CALL F600
601 IF (NOUT.GE.0)
&WRITE (NOUT,609)XO(1),XO(2),SCAL1,SCAL2
609 FORMAT(IH010X,31HORIGIN POINT IN PLOTTER COORD.(F6.2,2H ,F6.2,8H )
1 IN. / 11X,15HOVERALL SCALE =F6.3,32H INCH/ANGSITROM ELLIPSOID SCA
2LE =F6.3)
GO TO 391
C *****700 INSTRUCTIONS-ELLIPSOID AND SYMBOL PLOT FUNCTIONS*****
C *****FILL OUT DETAILS FOR SPECIAL MODELS*****
700 GO TO (701,702,704,705,709,7006,94),NJ3
7006 AIN(3)=1
GO TO 703
701 AIN(3)=8.
GO TO 703
702 AIN(3)=0.
703 AIN(1)=4.
AIN(2)=0.
AIN(4)=0.
GO TO 709
704 AIN(1)=3.
AIN(2)=-5.
GO TO 706
705 AIN(1)=1.
AIN(2)=0.
706 AIN(3)=1.
AIN(4)=5.
GO TO 8
C *****800 INSTRUCTIONS-BOND FUNCTIONS*****
800 CALL F800
GO TO 8
C *****900 INSTRUCTIONS-TITLE FUNCTIONS*****
900 CALL F900
GO TO 8
C *****1000 INSTRUCTIONS-OVERLAP FUNCTIONS*****
1000 CALL F1000
GO TO 8
C *****1100 INSTRUCTIONS-SAVE SEQUENCE FUNCTIONS*****
1100 IF (NJ2-2)1101,1102,1103
1101 ISAVE=1
GO TO 1104
1102 ISAVE=0
J=-1
CCC END FILE NSR
GO TO 1104
1103 ISAVE=-1
1104 REWIND NSR
GO TO 8
3000 CALL EXITING(NG)
END
FUNCTION ARCCOS(X)
ARCCOS(X) IN DEGREES
IF(1.0-ABS(X))1,2,2
1 X=SIGN(1.0,X)
2 IF(X)3,4,5
3 ARCCOS=180.0+ATAN(SQRT(1.0-X*X)/X)*57.29577951
GO TO 6
4 ARCCOS=90.0
GO TO 6
5 ARCCOS=ATAN (SQRT(1.0-X*X)/X)*57.29577951
6 RETURN
END
SUBROUTINE ATOM(OA,Z)
ATOM COORDINATE SUBROUTINE
REAL*8 OA,TA,D100K
DIMENSION X(3),Z(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AAREV(3,3),AARWK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),VRKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
D100K=100000.0
K=QA/D100K

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IF(K)109,109,117
109 X(1)=0.0
X(2)=0.0
X(3)=0.0
GO TO 125
117 IF(K-NATOM)119,119,503
503 NG=5
GO TO 325
119 DO 123 J=1,3
123 X(J)=P(J,K)
125 TA=DABS(OA)
KSYM=DMOD(TA,D100K)
KT=KSYM/100
KS=KSYM-100*KT
IF(KS-NSYM)203,203,403
403 NG=4
GO TO 325
203 IF(KS)403,205,213
205 Z(1)=X(1)
Z(2)=X(2)
Z(3)=X(3)
GO TO 311
213 DO 223 K=1,3
Z(K)=TS(K,KS)
DO 223 J=1,3
223 Z(K)=Z(K)+FS(J,K,KS)*X(J)
311 IF(KT)403,325,313
313 IF(KT-555)317,315,317
315 KSYM=KS
GO TO 325
317 K1=KT/100
K=KT-100*K1
K3=K-10*K2
Z(1)=Z(1)+FLOAT(K1-5)
Z(2)=Z(2)+FLOAT(K2-5)
Z(3)=Z(3)+FLOAT(K3-5)
325 RETURN
END
SUBROUTINE AXQB(A1,X,B1,JJJ)
***** SOLUTION OF MATRIX EQUATION AX=B FOR X *****
C ***** USES METHOD OF TRIANGULAR ELIMINATION *****
C ***** B AND X HAVE DIMENSIONS (3,JJJ),A IS ALWAYS (3,3)
C ***** TO INVERT A MAKE B 3 BY 3 IDENTITY MATRIX *****
C DIMENSION A1(3,3),A(3,3),B1(3,3),X(3,3)
NV=JJJ
***** TRANSFER DATA *****
DO 2 I=1,3
DO 2 J=1,3
IF(NV-J)2,1,1
1 B(I,J)=B1(I,J)
2 A(I,J)=A1(I,J)
***** TRIANGULARIZE MATRIX A *****
DO 17 I=1,2
S=0.0
DO 4 J=I,3
R=ABS(A(J,I))
IF(R-S)/4,3,3
3 S=R
L=J
4 CONTINUE
IF(L-I)5,10,5
5 DO 6 J=I,3
S=A(I,J)
A(I,J)=A(L,J)
6 A(L,J)=S
DO 8 J=1,NV
S=B(I,J)
B(I,J)=B(L,J)
8 B(L,J)=S
10 TEM=A(I,I)
IF(TEM)11,17,11
11 IPO=I+1
DO 16 J=IPO,3
IF(A(J,I))12,16,12
12 S=A(J,I)/TEM
A(J,I)=0.0
DO 13 K=IPO,3
13 A(J,K)=A(J,K)-A(I,K)*S
DO 15 K=1,NV
15 B(J,K)=B(J,K)-B(I,K)*S
16 CONTINUE
17 CONTINUE
C ***** MODIFY SINGULAR MATRIX *****
DO 20 I=1,3
IF(A(I,I))20,19,20
19 A(I,I)=AMAX1(1.E-25,AMAX1(A(1,1),A(2,2),A(3,3)))*1.E-15
20 CONTINUE
DO 24 K=1,NV
DO 24 I=1,3
N=4-I
M=N+1
TEM=B(N,K)
IF(3-M)/23,21,21
21 DO 22 J=M,3
22 TEM=TEM-A(N,J)*B(J,K)
23 B(N,K)=TEM/A(N,N)
24 X(N,K)=B(N,K)
RETURN
END
SUBROUTINE AXES(U,V,X,ITYPE)
***** STORE THREE ORTHOGONAL VECTORS EACH 1 ANGSTROM LONG *****
C ***** ITYPE .GT.0 FOR CARTESIAN,.LE.0 FOR TRICLINIC *****
C ***** IABS(ITYPE)=1 W(1)=U,W(2)=(UXY),W(3)=UX(UXV) *****
C ***** IABS(ITYPE)=2 W(1)=U,W(2)=(UXY)XU,W(3)=(UXV) *****
C ***** ITYPE=0 W(1)=A,W(2)=(AXB)XA,W(3)=(AXB), ABC=CELL VECTORS ***
C DIMENSION U(3),V(3),W(3),X(3,3)
IT=ITYPE
IF(IT)115,105,115
105 U(1)=1.
U(2)=0.
U(3)=0.
V(1)=0.
V(2)=1.
V(3)=0.
115 DO 125 J=1,3
125 W(J,1)=U(J)
IF(IABS(IT)-1)145,135,145
135 CALL NORM(U,V,W(1,2),IT)
CALL NORM(U,W(1,2),W(1,3),IT)
GO TO 155
145 CALL NORM(U,V,W(1,3),IT)

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CALL NORM(W(1,3),U,W(1,2),IT)
155 DO 195 I=1,3
    IF(MOD(NJ2,2).EQ.0) GO TO 143
    IF(IT)165,165,175
165 IC=-1
    GO TO 195
175 IC=1
195 CALL UNITY(W(1,I),X(1,I),IC)
    RETURN
END
SUBROUTINE BOND(Z1,Z2,NB,NAL,NAZ)
REAL*8 Z1,Z2,WD(2),TD,D100,D1000,D100K
DIMENSION BI(3,3),E(3,3),S(3,3),U(3,3),VUE(3)
DIMENSION V7(3),W(13,2),Z(3),RESB(2)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,CONT(5),P(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCI,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
***** OBTAIN POSITIONAL PARAMETERS *****
DATA RESB/.2,.08/
D100=100.
D1000=1000.
D100K=100000.
NG1=0
DO 105 J=1,26
    W(J,1)=0.
    WD(1)=Z1
    WD(2)=Z2
    DO 135 I=1,2
        CALL XYZ(WD(I),W(4,I),2)
    IF(NG)125,110,125
110 DO 115 J=1,3
    115 W(J+6,I)=XT(J)
    K=WD(I)/D100K
    L=DMOD(WD(I)/D100,D1000)
    LI=DMOD(WD(I),D100)
    CALL PLXY(W(4,I),W(2,I))
    IF(EDGE=BRDR*.25)120,128,128
120 NG=10
125 NG1=1
    IF (NOUT.GE.0)
        &WRITE(NOUT,136)CHEM(K),K,L,LI,(W(J,I),J=2,9)
    CALL ERENT(WD(I),800)
GO TO 134
128 IF (NJ2=10)130,134,134
130 IF (NOUT.GE.0)
    &WRITE(NOUT,136)CHEM(K),K,L,LI,(W(J,I),J=2,9)
134 continue
135 CONTINUE
136 FORMAT(1H,10X,A6,3H (13,1H,I3,I2,4H) 2F8.2,5X,3F8.3,13X,3F8.4)
    IF (NG1)999,137,999
137 CALL DIFV(W(7,1),W(7,2),V7)
DIST=SQRT(WMV(V7,AA,V7))
IF(MOD(NJ2,2).EQ.0) GO TO 143
IF(MOD(NJ2,10).EQ.1) GO TO 143
***** LINE BONDS AND CENTERED SYMBOLS (803,813) *
HGT=SCL*.12
C *** ORTEP-II calls
C CALL SIMBOL(W(2,1),W(3,1),HGT,MOD(IDINT(WD(1)/D100K),10),0.,-
C 11)
C CALL SIMBOL(W(2,2),W(3,2),HGT,MOD(IDINT(WD(2)/D100K),10),0.,-
C 12)
C *** Only one centered symbol (*) is available in ORTEP-III.
C *** It is triggered by the negative value for argument 6.
C *** Argument 4 is ignored by SIMBOL.
CALL SIMBOL(W(2,1),W(3,1),HGT,' ',0.,-1)
CALL SIMBOL(W(2,2),W(3,2),HGT,' ',0.,-2)
GO TO 570
C ***** STICK BONDS FOR 801,802,811,812 *****
143 KODE=KD(5,NB)
    IF(KODE)145,144,146
144 NEND=0
    GO TO 148
145 KODE=-KODE
146 NEND=128/2*KODE
C ***** FIND UPPERMOST ATOM PUT IN POSITION ONE *****
148 IF(VIEW)152,150,152
150 W(12,1)=1.
    W(12,2)=1.
    IF(W(6,1)-W(6,2))165,175,175
C ***** VECTOR FROM ATOM TO VIEWPOINT *****
152 DO 160 I=1,2
    155 W(J,I)=-W(J-6,I)
    W(12,I)=W(12,I)+VIEW
C ***** DISTANCE SQUARED TO VIEWPOINT *****
160 W(13,I)=VV(W(10,I),W(10,I))
    IF(W(13,2)-W(13,1))165,175,175
C ***** SWITCH ATOMS *****
165 DO 170 J=1,13
    TI=W(J,1)
    W(J,1)=W(J,2)
    W(J,2)=TI
    TD=WD(1)
    WD(1)=WD(2)
    WD(2)=TD
C ***** FORM IDEMFACTOR MATRIX *****
175 DO 180 J=1,3
    E(J,J)=1.
    E(J+1,1)=0.
    E(J+5,1)=0.
C ***** FORM VECTOR SET RADIAL TO BOND *****
CALL DIFV(W(4,2),W(4,1),DA(1,3))
CALL UNITY(DA(1,3),V3,1)
C ***** UNIT VECTOR FROM BOND MIDPOINT TO REFERENCE VIEWPOINT *****
DO 183 I=1,3
    V2(I)=0.0
181 V2(I)=V2(I)+AREV(J,3)*WRKV(J,I)
    IF(VIEW)183,183,182
182 V2(I)=V2(I)*VIEW-0.5*(W(I+3,1)+W(I+3,2))
183 CONTINUE
    CALL UNITY(V2,V2,1)

```

```

T6=ABS(VV(V3,V2))
IF(.9994-T6)185,185,187
***** ALTERNATE CALC IF BOND IS ALONG REFERENCE VIEW DIRECTION ***
185 DO 186 J=1,3
186 V2(J)=W(J+9,1)+W(J+9,2)
CALL UNITY(V2,V2,1)
T6=ABS(VV(V3,V2))
IF(.9994-T6)390,390,187
187 CALL AXES(V3,V2,B1,1)
188 T1=CD(3,NB)/SCAL2
DO 190 J=1,3
DA(J,1)=-B1(J,2)*T1
DA(J,2)=-B1(J,3)*T1
190 IF(NEND)500,500,195
***** SFT PLOTTING RESOLUTION FOR BOND *****
195 T1=CD(3,NB)*SCL
NRESOL=4
NBIS=3
DO 200 J=1,2
IF(T1.GE.RESB(J)) GO TO 202
IF(NEND.LE.NRESOL) GO TO 202
NBIS=NBIS-1
200 NRESOL=NRESOL*2
202 CALL RADIAL(NBIS)
***** DERIVE QUADRICS FOR EACH ATOM *****
DO 380 II=1,2
CALL PAXES(WD(II),2)
IF(NG)205,210,205
205 GO TO 999
C ***** DOES BOND GO TO ELLIPSOID OR TO ENVELOPE *****
210 T1=3-II*2
DO 212 J=1,3
V3(J)=V3(J)*T1
212 VUE(J)=0.
IF(KD(5,NB))260,260,215
215 IF(VMV(V3,0,W(10,II)))220,260,260
220 IBND=0
IF(VIEW)240,240,225
C ***** DERIVE TANGENT CONE DIRECTLY WITHOUT ROTATING COORDINATES **
225 T2=- (SCAL2*RMS(1)*RMS(2)*RMS(3))**2
DO 230 J=1,3
V1(J)=-W(J+9,II)/SCAL1
VUE(J)=V1(J)/SCAL2
C ***** INVERT ELLIPSOID MATRIX *****
DO 230 K=J,3
T1=0.0
DO 228 I=1,3
T1=T1+PAC(J,I)*PAC(K,I)*RMS(I)**2
U(J,K)=T1
230 U(K,J)=T1
C ***** ADD POLARIZED COFACTOR MATRIX TO ELLIPSOID MATRIX *****
DO 235 J=1,3
J1=MOD(J,3)+1
VJ1=V1(J1)
J2=MOD(J+1,3)+1
VJ2=V1(J2)
DO 235 K=J,3
K1=MOD(K,3)+1
K2=MOD(K+1,3)+1
S(J,K)=T2*Q(J,K)+(U(J1,K1)*V1(K2)-U(J1,K2)*V1(K1))
+ VJ1*(U(J2,K2)*V1(K1)-U(J2,K1)*V1(K2))
1
235 S(K,J)=S(J,K)
T5=0.0
GO TO 300
C ***** DERIVE TANGENT CYLINDER WITH AXIS ALONG Z *****
240 T1=-1.0/Q(3,3)
DO 250 J=1,2
DO 245 K=1,2
245 S(K,J)=Q(K,J)+Q(K,3)*Q(J,3)*T1
250 S(J,3)=0.0
S(3,3)=0.0
GO TO 270
C ***** TRANSFER ELLIPSOID *****
260 DO 265 J=1,9
265 S(J,1)=Q(J,1)
IBND=II
270 T5=1.
C ***** CHECK FOR BOND TAPER *****
300 IF(II-2)305,310,310
305 RADIUS=1.+T6*TAPER
GO TO 320
310 RADIUS=1.-T6*TAPER
320 CALL MV(S,V3,V4)
T2=VV(V3,V4)
C ***** COMPUTE BOND INTERSECTION *****
KL=5-II-II
KSTP=NRESOL
IF(NJ2-21)324,322,322
322 KSTP=32
324 DO 335 K=1,65,KSTP
DO 325 J=1,3
V6(J)=D(J,K)*RADIUS
T3=VV(V5,V4)
T4=T3*T3-T2*(VMV(V5,S,V5))-T5
IF(T4)345,330,330
T4=SQRT(T4)
T1=(T4-T3)/T2
T3=(-T4-T3)/T2
L=K+KL-1
DO 335 J=1,3
D(J,L)=(V6(J)+T1*V3(J))*SCL
D(J,L+1)=(-V6(J)-T3*V3(J))*SCL
IF(IBND+21-NJ2)360,338,360
338 IF(KD(5,NB))360,360,340
C ***** FOR LOCAL OVERLAP, MAKE BOND QUADRANGLE TANGENT TO ENVELOPING CONE
340 T3=VV(VUE,V4)
T4=T3**2-T2*(VMV(VUE,S,VUE)-T5)
IF(T4)345,350,350
345 NG=13
CALL ERENT(WD(II),800)
GO TO 999
350 T1=(SQRT(T4)-T3)/T2
DO 355 J=1,3
T4=(T1*V3(J)*SCL-0.5*(D(J,KL)+D(J,KL+64)))*1.001
D(J,KL)=D(J,KL)+T4
355 D(J,KL+64)=D(J,KL+64)+T4
360 CALL PROJ(D(1,KL),DP(1,II),W(4,II),XO,VIEW,1,65,KSTP)
IF(IBND-1)370,365,370

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365 CALL PROJ(D(1,KL+KSTP+1),DP(1,II+64+KSTP),W(4,II),XO,VIEW,1,
& 65-KSTP,KSTP)
GO TO 380
C ***** RETRACE TOP HALF *****
370 KK=64-(II-1)*KSTP
DO 375 K=KSTP,KK,KSTP
L=K+II
M=L+64
N=66-L
DP(1,M)=DP(1,N)
375 DP(2,M)=DP(2,N)
380 CONTINUE
C ***** CHECK FOR LOCAL OVERLAP OR HIDDEN BOND *****
DO 395 K=1,65,32
T1=0.
T2=0.
DO 385 J=1,2
T1=T1+(DP(J,K)-W(J+1,1))**2
T2=T2+(DP(J,K+1)-W(J+1,1))**2
IF(T2-T1)390,390,395
395 CONTINUE
C ***** CALL GLOBAL OVERLAP ROUTINE *****
ICQ=0
CALL LAP800(NA1,NA2,ICQ)
IF(NJ2-21)400,999,999
400 IF(ICQ)390,405,405
C ***** DRAW BOND OUTLINE *****
405 CALL DRAW(DP(1,1),0.,0.,.3)
DO 415 K=NRESL,I29,NRESOL
415 CALL DRAW(DP(1,K),0.,0.,.2)
DO 420 K=2,66,NRESOL
420 CALL DRAW(DP(1,K),0.,0.,.2)
CALL DRAW(DP(1,65),0.,0.,.2)
C ***** DRAW BOND DETAIL *****
425 K=65
430 K=K-NBND
IF(K-1)500,500,435
435 CALL DRAW(DP(1,K),0.,0.,.3)
CALL DRAW(DP(1,K+1),0.,0.,.2)
K=K-NBND
IF(K-1)500,500,440
440 CALL DRAW(DP(1,K+1),0.,0.,.3)
CALL DRAW(DP(1,K),0.,0.,.2)
GO TO 430
500 HGT=CD(4,NB)
OFF=CD(5,NB)
IF(HGT)570,570,510
OFF=CD(5,NB)
C ***** PERSPECTIVE BOND LABEL ROUTINE *****
C ***** BASE DECISIONS ON REFERENCE SYSTEM *****
510 K=0
CALL DIFV(W(7,2),W(7,1),V7)
CALL VM(V7,AREV,V1)
CALL AXES(V1,E(1,3),U,1)
DO 535 I=1,3
T1=1.
IF(I-2)515,515,520
515 IF(VV(U(1,I),SYMB(1,I)))525,530,530
520 IF(MOD(K,2))530,525,530
525 T1=-1.
K=K+1
530 DO 535 J=1,3
U(J,I)=U(J,I)*T1
535 VT(J,I)=B1(J,I)*T1
DO 540 J=1,3
540 VT(J,4)=.5*(W(J+3,1)+W(J+3,2))
C ***** CHECK FOR EXCESS FORESHORTENING *****
IF(FORE-ABS(U(3,1)))545,550,550
545 CALL NORM(U(1,2),SYMB(1,3),VT(1,1),1)
VT(1,3)=SYMB(1,3)
VT(2,3)=SYMB(2,3)
VT(3,3)=SYMB(3,3)
HGT=CD(6,NB)
OFF=CD(7,NB)
DO 550 T1=CD(8,NB)
IF(HGT)550,999,550
Z(1)=VT(1,4)-HGT*(11.+3.*T1)/7.
Z(2)=VT(2,4)+OFF-HGT*.5
Z(3)=VT(3,4)
XO(3)=Z(3)
ITILT=1
I9=T1+2.
T9=10.**I9
DISTR=AINT((DIST*T9)+0.5)/T9+.0001
CALL NMBUR(Z(1),Z(2),HGT,DISTR,0.,I9)
570 ITILT=0
IF(NJ2-10)580,999,999
580 IF(NOUT.GE.0)
&WRITE(NOUT,571)DIST
571 FORMAT(1H,59X,10HDISTANCE =,F8.3/1H)
GO TO 999
390 NG=14
CALL ERENT(WD(2),800)
999 RETURN
END
SUBROUTINE DIFV(X,Y,Z)
VECTOR - VECTOR
Z(3)=X(3)-Y(3)
DIMENSION X(3),Y(3),Z(3)
Z(1)=X(1)-Y(1)
Z(2)=X(2)-Y(2)
Z(3)=X(3)-Y(3)
RETURN
END
SUBROUTINE DRAW(W,DX,DY,NPEN)
DIMENSION W(3),X(3),Y(3),Z(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AARV(3,3),AID(3,3)
1,AIN(1,40),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2,CONT(5),P(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4,ORGN(3),PAC(3,5),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5,SCAL2,SC3,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),W(3),W(3),W(3),W(3)
7,XLING(3),XO(3),XT(3)
Y(1)=W(1)+DX
Y(2)=W(2)+DY
IF(ITILT)115,140,115
C ***** ROTATE FOR PERSPECTIVE TITLE *****
115 Y(3)=XO(3)
DO 120 I=1,3
120 Z(I)=Y(I)-VT(I,4)

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DO 130 I=1,3
130 X(I)=VT(I,1)*Z(1)+VT(I,2)*Z(2)+VT(I,3)*Z(3)+VT(I,4)
CALL PLPXY(X,Y)
***** CHECK BOUNDARY *****
140 DO 160 J=1,2
IF(Y(J)-XING(J)+.1)150,150,145
145 Y(J)=XING(J)-.1
150 IF(Y(J)-.1)155,160,160
155 Y(J)=.1
160 CONTINUE
C ***** CHECK FOR OVERLAP *****
NCQ=0
CALL LAPDRW(Y,NPEN,NCQ)
IF(NCQ)165,165,170
***** CALL PLOTTING ROUTINE IF NO OVERLAPPING ELEMENTS ARE STORED
165 CALL SCRIBE(Y,NPEN)
170 RETURN
END
SUBROUTINE EDITR
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1,AIN(1,40),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2,CONT(5),P(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4,ORGN(3),PAC(3,3),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5,SCAL2,SCI,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7,XLING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1,IDENT(2,505),MAXATM
COMMON /QUEUE/ NED,NQUE,NEXT,NBACK,INQ,QUE(96),hque(96)
common /ns/ npl,ndraw,norient,nvar
CHARACTER*73 INQ,QUE,hque,tline
character*80 answer,card
CHARACTER*1 CH
10 FORMAT(6X,1111111111222222222233333333334444444444555',
*5555555666666666777',/,lx,LINE',lx,'1234567890123456789012345',
*67890123456789012345678901234567890123456789012')
NUM=1
C *** PRINT PART OR ALL OF COMMAND QUE ***
NUM1=MAXO(1,NUM-2)
100 WRITE(*,103)
103 FORMAT(1X)
111 WRITE(*,111)(J,QUE(J),J=NUM1,NQUE)
C *** DISPLAY PROMPT ***
115 WRITE(*,121)
121 FORMAT(/,' C=Change line #
&) # [#,/, I=Insert line before #
& #] [#,
&/, S=Save modified instruction set
&struction set',
&/, P=Save drawing as Postscript
&L',
&/, R=Redraw structure on screen
&Quit',/, >-> ', $)
C *** READ COMMAND CHARACTER AND LINE NUMBER(S) ***
read(*,131)answer
131 format(a)

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if (answer(1:1).eq.' ') go to 133
if (answer(1:1).ge.'1'.and.answer(1:1).le.'9') go to 133
last=iend(answer)
do 132 i=2,last
ch=answer(i:i)
if ((ch.ge.'a'.and.ch.le.'z') .or.
(ch.ge.'A'.and.ch.le.'Z')) go to 133
*
ich=ichar(ch)-48
if ((ch.ne.' ') .and. ((ich.lt.0) .or. (ich.gt.9))) go to 133
132 continue
go to 135
133 write(*,134)
134 format (/,'***INVALID INPUT! Enter 1 letter and 0, 1, or 2 integer
*s separated by spaces.***')
go to 115
135 answer=answer(1:last)//' 0 0'
read(answer,*)ch,num,num2
open(15,status='scratch')
write(15,136) answer(1:1),answer(2:75)
136 format(' ','al',' ',' ','lx,a)
rewind(15)
read(15,*)ch,num,num2
close(15)
numz=num
NUM=MAXO(1,NUM)
if (num2.gt.nque) then
write(*,137)
format(/,' **** value out of range ****')
137 go to 115
end if
IF(NUM.GT.NUM2) NUM2=NUM
write(6,*),
IF(CH.EQ.'T'.or.ch.eq.'t') GO TO 210
IF(CH.EQ.'D'.or.ch.eq.'d') GO TO 240
IF(CH.EQ.'C'.or.ch.eq.'c') GO TO 270
IF(CH.EQ.'I'.or.ch.eq.'i') GO TO 310
IF(CH.EQ.'O'.or.ch.eq.'o') GO TO 410
IF(CH.EQ.'S'.or.ch.eq.'s') GO TO 420
IF(CH.EQ.'R'.or.ch.eq.'r') GO TO 540
IF(CH.EQ.'Q'.or.ch.eq.'q') GO TO 590
IF(CH.EQ.'P'.or.ch.eq.'p') GO TO 510
IF(CH.EQ.'H'.or.ch.eq.'h') GO TO 520
GO TO 115
C *** TYPE LINES ***
210 if (numz.eq.0) num2=nque
WRITE(*,110)
GO TO 115
C *** DELETE LINES ***
240 if (numz.eq.0) then
write(6,*), *** Supply line number(s) with command ***
go to 115
end if
DO 260 I=NUM,NUM2
NQUE=NQUE-1
NEXT=NEXT-1
DO 250 J=NUM,NQUE
250 QUE(J)=QUE(J+1)
260 QUE(NQUE+1)=
GO TO 100
C *** CHANGE AN OLD LINE ***

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270 if (numz.eq.0) then
  write(6,*) ' *** Supply line number with command ***'
  go to 115
end if
NUM3=MAX0(1,NUM-3)
WRITE(*,111)(J,QUE(J),J=NUM3,NUM)
write(*,271)
271 format(' *** NOTE: Type @ to substitute a space in the original li
&ne. ***',/)
WRITE(*,10)
WRITE(*,111)NUM,QUE(NUM)
WRITE(*,276)NUM
276 FORMAT(1X,I3,' ',)$
read(*,281) tline
281 FORMAT(A72)
do 282 i=1,72
  if (tline(i:i).ne.' ') then
    if (tline(i:i).eq.'@') then
      que(num)(i:i)=' '
    else
      que(num)(i:i)=tline(i:i)
    end if
  end if
282 continue
write(*,283)
283 format(/,' Line now reads:')
284 format(6x,a)
write(*,285)
285 format(/,' Hit ENTER or RETURN key ',)$
read(*,131) CH
GO TO 100
C *** INSERT A NEW LINE ***
310 if (numz.eq.0) then
  write(6,*) ' *** Supply line number with command ***'
  go to 115
end if
IF(NQUE+1.GT.96) GO TO 115
NQUE=NQUE+1
NN=NQUE
N=NQUE-NUM
DO 320 J=1,N
  QUE(NN)=QUE(NN-1)
  NUM4=MAX0(1,NUM-4)
  NUM1=MAX0(1,NUM-1)
  write(*,10)
  WRITE(*,111)(J,QUE(J),J=NUM4,NUM1)
  WRITE(*,276)NUM
  READ(*,281) QUE(NUM)
  NEXT=NEXT+1
GO TO 100
C *** RETRIEVE OLD SET OF INSTRUCTIONS ***
410 DO 415 J=1,NBACK
415 que(j)=bque(j)
NUM1=1
NQUE=NBACK
NUM2=NQUE
GO TO 100
C *** SAVE CURRENT SET OF INSTRUCTIONS ***
420 CONTINUE
421 write (*,422)
422 format(' Enter file name: ',)$
read (*,131) answer
open (16,file=answer,status='new',err=460)
rewind(NED)
430 read (NED,131,end=440) card
write (16,131) card(1:iend(card))
go to 430
440 WRITE(16,450)(QUE(I),I=1,NQUE)
450 FORMAT(A73)
close(16)
GO TO 115
460 write(6,*) 'File already exists. Choose a different name.'
go to 421
C *** SAVE PICTURE AS POSTSCRIPT***
510 ndraw=2
go to 541
C *** SAVE PICTURE AS HPGL***
520 ndraw=3
C *** REDRAW PICTURE ***
540 ndraw=1
c *** default instruction save
541 open (16,file='TEP.NEW',status='unknown')
rewind(NED)
542 read (NED,131,end=543) card
write (16,131) card(1:iend(card))
go to 542
543 WRITE(16,450)(QUE(I),I=1,NQUE)
close(16)
call recycle
590 RETURN
END
SUBROUTINE EIGEN (W,VALU,VECT)
***** EIGENVALUES AND EIGENVECTORS OF 3X3 MATRIX *****
DIMENSION W(3,3),VALU(3),VECT(3,3),A(3,3),B(3,3),V(3),U(3)
COMMON NG
***** STATEMENT FUNCTION *****
PHIF(Z)=((B2-Z)*Z+B1)*Z+B0
***** START OF PROGRAM *****
ERRND=5.E-7
SIGMA=0.
DO 115 J=1,3
DO 115 I=1,3
TEM=W(I,J)
A(I,J)=TEM
115 SIGMA=SIGMA+TEM*TEM
***** CHECK FOR NULL MATRIX *****
IF(SIGMA)230,230,120
SIGMA=SQRT(SIGMA)
***** FORM CHARACTERISTIC EQUATION *****
B2=A(1,1)+A(2,2)+A(3,3)
B1=-A(1,1)*A(2,2)-A(1,1)*A(3,3)-A(2,2)*A(3,3)+A(1,3)*A(3,1)
1+A(2,3)*A(3,2)+A(1,2)*A(2,1)
B0=A(1,1)*A(2,2)*A(3,3)+A(1,2)*A(2,3)+A(1,3)*A(3,2)+A(2,1)*A(3,1)-
1A(1,3)*A(3,1)*A(2,2)-A(1,1)*A(2,3)-A(1,2)*A(3,2)-A(1,2)*A(3,3)
***** FIRST ROOT BY BISECTION *****
X=0.
Y=SIGMA
TEM=PHIF(SIGMA)

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VNEW=0.0
IF(B0)135,250,145
135 IF(TEM)140,140,165
140 Y=-Y
145 GO TO 165
145 Y=0.
X=SIGMA
IF(TEM)165,165,150
150 X=-X
***** NOW PHIF(X).LT.0.AND.PHIF(Y).GT.0. *****
165 VNEW=(X+Y)*.5
DO 225 I=1,40
175 IF(PHIF(VNEW))180,250,185
180 X=VNEW
GO TO 200
185 Y=VNEW
200 VOLD=VNEW
VNEW=(X+Y)*.5
TEM=ABS(VOLD-VNEW)
IF(TEM-ERRND)250,250,205
205 IF(VOLD)210,225,210
210 IF(ABS(TEM/VOLD)-ERRND)250,250,225
225 CONTINUE
C ***** DID NOT CONVERGE, SET ERROR INDICATOR *****
230 NG=6
GO TO 400
C ***** STORE FIRST ROOT *****
250 U(3)=VNEW
C ***** DEFLATE *****
CI=B2-VNEW
CO=B1+CI*VNEW
***** SOLVE QUADRATIC *****
TEM=C1*CI+4.*CO
IF(TEM)255,265,260
C ***** IGNORE IMAGINARY COMPONENT OF COMPLEX ROOT *****
255 TEM=0.
GO TO 265
260 TEM=SQRT(TEM)
265 U(1)=.5*(CI-TEM)
U(2)=.5*(CI+TEM)
***** SORT ROOTS *****
DO 275 J=1,2
270 IF(U(J)-U(3))275,275,270
TEM=U(J)
U(J)=U(3)
U(3)=TEM
275 CONTINUE
L=1
DO 375 I=1,2
C ***** CHECK FOR MULTIPLE ROOTS *****
TEM=ERRND*100.
NG=0
L=L-1
DO 305 I=1,2
IF(U(I+1)-U(I)-TEM)300,300,290
290 IF(U(I))295,305,295
295 IF(ABS((U(I+1)-U(I))/U(I))-TEM)300,300,305
300 L=L-1
305 CONTINUE
IF(L=1)308,400,400

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308 LLL=L
***** EIGENVECTOR ROUTINE *****
DO 375 I=1,3
TI=U(II)
IF(L)315,310,322
***** TWO VECTORS NULL FOR DOUBLE ROOT *****
310 IF(NG+5-II)315,322,315
***** ALL VECTORS NULL FOR TRIPLE ROOT *****
315 DO 320 J=1,3
320 VECT(J,II)=0.0
GO TO 375
322 DO 325 J=1,3
325 A(J,J)=W(J,J)-TI
SMAX=0.0
DO 355 I=1,3
II=I
IF(I-2)335,335,340
335 II=I+1
340 B(I,1)=A(I,2)*A(II,3)-A(I,3)*A(II,2)
B(I,2)=A(I,3)*A(II,1)-A(I,1)*A(II,3)
B(I,3)=A(I,1)*A(II,2)-A(I,2)*A(II,1)
TEM=B(I,1)**2+B(I,2)**2+B(I,3)**2
IF(TEM-SMAX)355,355,350
350 SMAX=TEM
IMAX=I
355 CONTINUE
IF(SMAX)353,353,360
353 NG=7
GO TO 375
360 SMAX=SQRT(SMAX)
DO 365 J=1,3
365 V(J)=B(IMAX,J)/SMAX
***** REFINE EIGENVECTOR *****
CALL AXEQB(A,V,V,1)
TEM=AMAX1(ABS(V(1)),ABS(V(2)),ABS(V(3)))
DO 370 J=1,3
370 V(J)=V(J)/TEM
CALL UNLTY(V,VECT(1,II),1)
***** REFINE EIGENVALUE *****
TI=VMV(VECT(1,II),W,VECT(1,II))
U(II)=TI
375 VALU(II)=TI
400 RETURN
END
SUBROUTINE ERENT(TD,N)
REAL*8 TD
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 IN,TITLT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCALI
5 SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),VRKV(3,3)
7 XLNG(3),XO(3),XT(3)
character*63 msg(18)
data msg /
1 'No sentinel found after reading 96 symmetry cards',
2 'No sentinel found after reading parameter cards for 100 atoms',
3 'Aniso temp factor coeffs form non-positive definite matrix',

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4 'Symmetry operation no. is higher than no. of input operations',
5 'Atom number is higher than the number of input atoms',
6 'Null temp factor matrix or failure in bisection routine',
7 'Eigenvector routine failure due to null vector',
8 'Error initializing screen driver',
9 'Unidentified instruction number',
a 'Atom out of bounds',
b 'No vector search codes',
c 'Insufficient number of atoms in ATOMS list',
d 'Imaginary bond intersection (i.e., bond larger than atom)',
e 'Hidden (end-on) bond',
f 'Null vector as base line',
g 'ATOMS array is full',
h 'Maximum number of overlapping atoms (20) exceeded',
i 'Maximum number of overlapping bonds (30) exceeded' /
IF (NOUT.GE.0) then
WRITE (NOUT,115)NG,TD,N
115 FORMAT(1H,10X,10HFAULT NG =,I3,F10.0,I6)
WRITE (NOUT,116) msg(ng)
116 format(1H,10X,a,/)
end if
NG=0
RETURN
END
SUBROUTINE EXITING(ING)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAWK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DR(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
character*12 routin(18)
data routin /
1 'PRELIM', 'PRELIM', 'ATOM, PAXES', 'ATOM, PAXES',
2 'EIGEN', 'EIGEN', 'INITSC', 'MAIN, SPARE', 'BOND, BOND, F700',
3 'F800', 'F600, SEARCH', 'BOND', 'BOND', 'F900', 'STORE',
4 'LAP700, LAPAB', 'LAP800, LAPAB' /
data msg /
1 'No sentinel found after reading 96 symmetry cards',
2 'No sentinel found after reading parameter cards for 100 atoms',
3 'Aniso temp factor coeffs form non-positive definite matrix',
4 'Symmetry operation no. is higher than no. of input operations',
5 'Atom number is higher than the number of input atoms',
6 'Null temp factor matrix or failure in bisection routine',
7 'Eigenvector routine failure due to null vector',
8 'Error initializing screen driver',
9 'Unidentified instruction number',
a 'Atom out of bounds',
b 'No vector search codes',
c 'Insufficient number of atoms in ATOMS list',
d 'Imaginary bond intersection (i.e., bond larger than atom)',
e 'Hidden (end-on) bond',
f 'Null vector as base line',
g 'ATOMS array is full',
h 'Maximum number of overlapping atoms (20) exceeded',
i 'Maximum number of overlapping bonds (30) exceeded' /
IF (NOUT.GT.0) then
if (nout.gt.0) then
write (*,101) ing
write (nout,101) ing
write (nout,102) routin(ing)
write (nout,103) msg(ing)
end if
write (*,101) ing
write (*,102) routin(ing)
write (*,103) msg(ing)
end if
IF(NOUT.GT.0) CLOSE(NOUT,STATUS='KEEP')
101 format('Fault Indicator: ',i2)
102 format('Subroutine(s) Involved: ',a)
103 format('Fault: ',a)
STOP
END
SUBROUTINE F200
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAWK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DR(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
common /ns/ npf,ndraw,norient,nvavr
common /trifac/ xtrans,ytrans
c *** NO drawing
if (ndraw.eq.0) return
c *** initialize plotting (201 or 203 inst) ***
90 to (201,202,201,204,205),nj2
201 xtrans=0.
ytrans=0.
if (ndraw.eq.1) call initsc
if (ndraw.eq.2) call initps
if (ndraw.eq.3) call initfp
if (ndraw.eq.9) then
open(unit=npf,file='TEP.EDT',status='unknown')
nvavr=1
end if
return
c *** change origin of plotting area or terminate (202 inst) ***
202 if (ain(1).eq.0 .and. ain(2).eq.0.) then
if (ndraw.eq.2) call endps
if (ndraw.eq.3) call endhp
if (ndraw.eq.1) call endsc
if (ndraw.eq.9) close(npf)
else
xtrans=ain(1)
ytrans=ain(2)
if (ndraw.eq.9) write (npf,203) xtrans,ytrans
203 format('TRN',2(Lx,f10.6))
end if
return
c *** change plot color (204 inst) ***
204 icolor=ain(1)
if (ndraw.eq.1) call colrsc(icolor)

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if (ndraw .eq. 2) call colrps(icolor)
if (ndraw .eq. 3) call colrhp(icolor)
if (ndraw .eq. 9) call colrsc(icolor)
return
c *** change pen width (205 inst) ***
c *** parameter units are thousandths of an inch (default=5)
205 penw=ain(1)
if (ndraw .eq. 1) call penwsc(penw)
if (ndraw .eq. 2) call penwps(penw)
if (ndraw .eq. 3) call penwhp(penw)
if (ndraw .eq. 9) call penwsc(penw)
return
c
SUBROUTINE F400
***** ATOM LIST FUNCTIONS *****
REAL*8 DL00,DL1000,DL100K,TD,TD1,TD2
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAWRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TITLE,KD(5,20),LATM,NATOM,NCD,NG,NJ2,INOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCH,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V1(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
DL100=100.
DL1000=1000.
DL100K=100000.
NG=0
IF(LATM)402,402,400
400 DO 401 I=1,LATM
401 CALL ATOM(ATOMID(I),ATOMS(1,I))
402 IF(MOD(NJ2,10)-1)499,404,403
403 IF(MOD(NJ2,10)-7)406,404,499
406 CALL SEARC
GO TO 499
c
***** STORES (401) OR REMOVES (411) RUNS OF ATOMS *****
c
***** RUN HIERARCHY = ATOM NO./SYM/ A/B/C TRANS. *****
404 II=1
c
***** FIND RUNS IN AIN AREAY *****
405 TD1=AIN(II)
IF(TD1)410,410,420
410 II=II+1
IF(140-II)499,405,405
420 JU=II
c
***** SET INITIAL RUN VALUES *****
M1=TD1/DL100K
M2=DMOD(TD1,DL100)
M5=DMOD(TD1/DL100,DL1000)
IF(M5)422,422,423
422 M5=555
423 M3=M5/100
M4=MOD(M5/10,10)
M5=MOD(M5,10)
425 IF(140-JJ)435,430,430

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430 TD2=-AIN(JJ)
IF(TD2)435,425,440
435 II=JJ-1
***** SET TERMINAL VALUES FOR DEGENERATE RUN *****
M1=M1
M2=M2
M3=M3
M4=M4
M5=M5
GO TO 450
440 II=JJ
***** SET TERMINAL RUN VALUES *****
M1=TD2/DL100K
M2=DMOD(TD2,DL100)
M5=DMOD(TD2/DL100,DL1000)
IF(M5)445,445,446
445 M5=555
446 M3=M5/100
M4=MOD(M5/10,10)
M5=MOD(M5,10)
***** LOOP THROUGH ALL RUNS *****
450 DO 490 L5=M5,M5
DO 490 L4=M4,N4
DO 490 L3=M3,N3
DO 490 L2=M2,N2
DO 490 L1=M1,N1
TD=DBLE(L1)*DL100K+DBLE(L3*100000+L4*10000+L5*100+L2)
CALL ATOM(TD,VI(1))
IF(NG)455,458,455
455 CALL ERENT(TD,401)
GO TO 490
c
***** CHECK IDENT CODE IF 407/417 INSTRUCTION *****
458 IF(MOD(NJ2,10)-7)475,460,490
460 ID1=IDENT(1,II)
ID2=IDENT(2,II)
IF(NCD)490,490,465
465 DO 470 J=1,NCD
if ((id1.ge.kd(1,j) .and. id1.le.kd(2,j)) .and.
& (id2.ge.kd(3,j) .and. id2.le.kd(4,j))) go to 475
else if (kd(1,j).gt.0.) then
if (id1.ge.kd(1,j) .and. id1.le.kd(2,j)) go to 475
else if (kd(3,j).gt.0.) then
if (id2.ge.kd(3,j) .and. id2.le.kd(4,j)) go to 475
end if
end if
c
IF(ID1-KD(1,J))470,467,467
c 467 IF(KD(2,J)-ID1)470,468,468
c 468 IF(ID2-KD(3,J))470,469,469
c 469 IF(KD(4,J)-ID2)470,475,475
470 CONTINUE
GO TO 490
475 CALL STOR(TD)
490 CONTINUE
GO TO 410
499 RETURN
END
SUBROUTINE F500
DIMENSION RM(3,3),V(3,4)
REAL*8 TD,DL100,DL1000,DL100K
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2

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COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCALI
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XLING(3),XO(3),XT(3)
INTEGER*2 IDENT
CHARACTER*8 CHEM
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
NG=0
D100=100.
D1000=1000.
D100K=100000.
IF(NJ2-11)500,700,710
500 IF(NJ2-1)710,501,510
501 TD=AIN(1)
CALL ATOM(TD,ORGN)
IF(NG)502,504,502
502 CALL ERENT(TD,501)
CALL EXLTING(NG)
504 DO 506 K=1,4
TD=AIN(K+1)
CALL ATOM(TD,V(1,K))
IF(NG)502,506,502
506 CONTINUE
DO 507 J=1,3
V1(J)=V(J,2)-V(J,1)
V2(J)=V(J,4)-V(J,3)
IND=-1
IF(AIN(7))509,509,508
508 IND=-2
509 CALL AXES(V1,V2,REFV,IND)
GO TO 670
510 IF(NJ2-4)515,511,599
***** SHIFT ORIGIN FOR PROJECTION AXIS (IN INCHES) *****
511 DO 513 J=1,3
DO 512 K=1,3
TI=AIN(K)
512 ORGN(J)=ORGN(J)+REFV(J,K)*TI/SCALI
513 XO(J)=XO(J)+T2
GO TO 675
C ***** FORM ROTATION MATRIX *****
515 DO 514 J=1,3
DO 514 K=1,3
514 V(J,K)=REFV(J,K)
DO 517 L=1,139,2
I=AIN(L)
IF(I) 532,519,516
516 X=AIN(L+1)*1.7453293D-2
TI=COS(X)
T2=SIN(X)
I3=MOD(I+2,3)+1
I1=MOD(I+3,3)+1
I2=MOD(I+1,3)+1
RM(I1,I1)=TI
RM(I1,I2)=T2
RM(I1,I3)=0.0

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RM(I2,I1)=-T2
RM(I2,I2)=T1
RM(I2,I3)=0.0
RM(I3,I1)=0.0
RM(I3,I2)=0.0
RM(I3,I3)=1.0
517 CALL MM(V,RM,V)
519 IF(NJ2-3)518,525,599
518 DO 522 J=1,3
DO 522 I=1,3
522 REFV(I,J)=V(I,J)
GO TO 552
525 DO 528 J=1,3
DO 528 I=1,3
528 WRKV(I,J)=V(I,J)
GO TO 552
532 IF(NJ2-3)535,552,599
535 I=MOD(-I,3)
DO 542 J=1,3
DO 542 K=1,3
TI=REFV(K,3)
REFV(K,3)=REFV(K,2)
REFV(K,2)=REFV(K,1)
542 REFV(K,1)=TI
552 CONTINUE
IF(NJ2-3)670,582,599
582 CALL MM(AA,WRKV,AAWRK)
GO TO 699
599 IF(NJ2-5)699,600,607
600 IF(LATM-1)605,610,610
605 NG=12
606 CALL ERENT(0,D0,506)
CALL EXLTING(NG)
607 IF(NJ2-6)699,608,710
608 IF(LATM-3)605,610,610
610 DO 612 J=1,3
V2(J)=0.0
DO 612 I=1,3
RM(I,J)=0.0
AWT=0.0
DO 620 K=1,LATM
CALL ATOM(ATOMID(K),ATOMS(1,K))
T2=1.0
613 IF(NGD)618,618,613
I1=ATOMID(K)/D100K
DO 616 J=1,NCD
IF(KD(5,J).EQ.1) I1=IDENT(1,K)
IF(KD(5,J).EQ.2) I1=IDENT(2,K)
T2=CD(1,J)
IF(I1-KD(1,J))616,614,614
614 IF(KD(2,J)-I1)616,618,618
616 CONTINUE
GO TO 620
618 AWT=AWT+T2
DO 619 J=1,3
619 V2(J)=V2(J)+ATOMS(J,K)*T2
620 CONTINUE
IF(AWT)605,605,621
***** PUT ORIGIN AT CENTER OF GRAVITY *****
C 621 DO 622 J=1,3
622 ORGN(J)=V2(J)/AWT

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IF(NI2-6)699,624,710
***** FORM PRODUCT-MOMENT MATRIX FOR ATOMS IN ATOM LIST *****
C 624 DO 630 K=1,LATM
T2=1.0
IF(NCD)628,628,625
DO 627 J=1,NCD
IF(ATOMID(K)/D100K)
IF(KD(5,J).EQ.1) I1=IDENT(1,K)
IF(KD(5,J).EQ.2) I1=IDENT(2,K)
T2=CD(1,J)
IF(I1-KD(1,J))627,626,626
626 IF(KD(2,J)-I1)627,628,628
627 CONTINUE
GO TO 630
628 DO 629 J=1,3
T1=(ATOMS(J,K)-ORGN(J))*T2
DO 629 I=1,3
629 RM(I,J)=T1*(ATOMS(I,K)-ORGN(I))+RM(I,J)
T1=0.037*(RM(1,1)+RM(2,2)+RM(3,3))
630 CONTINUE
DO 632 J=1,3
DO 632 I=1,3
632 RM(I,J)=RM(I,J)*T1
C ***** TRANSFORM TO INERTIAL AXIS SYSTEM *****
CALL MM(RM,AA,DA)
CALL EIGEN(DA,RMS,PAT)
IF(RMS(2))605,605,635
635 IF(NG)640,633,606
C ***** MAKE SURE VECTORS ARE ORTHOGONAL --> NEW REFERENCE VECTORS *
633 CALL AXES(PAT(1,3),PAT(1,1),REFV,-1)
GO TO 665
C ***** TWO EQUAL EIGENVECTORS SPECIAL CASE *****
640 IF(NG+6)665,665,645
645 N=NG+5
CALL UNITY(PAT(1,N),V1,-1)
DO 650 K=1,3
DO 660 J=1,3
IF(ABS(VM(V1,AA,REFV(1,K)))-.58)655,650,650
650 CONTINUE
655 CALL AXES(V1,REFV(1,K),DA,-1)
DO 660 K=1,3
L=MOD(N-K+2,3)+1
DO 660 J=1,3
REFV(J,L)=DA(J,K)
660 NG=0
670 CALL MM(AA,REFV,AAREV)
675 DO 680 J=1,3
DO 680 I=1,3
WKKV(I,J)=REFV(I,J)
680 AAWRK(I,J)=AAREV(I,J)
C ***** ELIMINATE ALL PREVIOUSLY STORED OVERLAP INFORMATION *****
C ***** ALL INSTRUCTIONS FROM 501 THROUGH 510 DO THIS *****
699 CALL LAP500(0)
GO TO 710
C ***** STORE NEW OVERLAP INFORMATION (INSTRUCTION 511) *****
700 CALL LAP500(1)
710 RETURN
END
SUBROUTINE F600
***** SCALING AND CENTERING FUNCTIONS *****
DIMENSION MAX(3),SCAL(4),X(3),XMAX(3),XMIN(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TITLIT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WKKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
dimension crtvai(99)
data crtvai /
1 0.3389, 0.4299, 0.4951, 0.5479, 0.5932, 0.6334, 0.6699, 0.7035,
2 0.7349, 0.7644, 0.7924, 0.8192, 0.8447, 0.8694, 0.8932, 0.9162,
3 0.9386, 0.9605, 0.9818, 1.0026, 1.0230, 1.0430, 1.0627, 1.0821,
4 1.1012, 1.1200, 1.1386, 1.1570, 1.1751, 1.1932, 1.2110, 1.2288,
5 1.2464, 1.2638, 1.2812, 1.2985, 1.3158, 1.3330, 1.3501, 1.3672,
6 1.3842, 1.4013, 1.4183, 1.4354, 1.4524, 1.4695, 1.4866, 1.5037,
7 1.5209, 1.5382, 1.5555, 1.5729, 1.5904, 1.6080, 1.6257, 1.6436,
8 1.6616, 1.6797, 1.6980, 1.7164, 1.7351, 1.7540, 1.7730, 1.7924,
9 1.8119, 1.8318, 1.8519, 1.8724, 1.8932, 1.9144, 1.9360, 1.9580,
a 1.9804, 2.0034, 2.0269, 2.0510, 2.0757, 2.1012, 2.1274, 2.1544,
b 2.1824, 2.2114, 2.2416, 2.2730, 2.3059, 2.3404, 2.3767, 2.4153,
c 2.4563, 2.5003, 2.5478, 2.5997, 2.6571, 2.7216, 2.7955, 2.8829,
d 2.9912, 3.1365, 3.3682 /
***** DEL = 1. FOR INCREMENTING FUNCTIONS *****
***** DEL = 0. FOR REGULAR FUNCTIONS *****
C DEL=FLOAT(MOD(NJ2/10,2))
NJ2=MOD(NJ2,10)
***** EXPLICIT ORIGIN AND SCALE *****
TI=AIN(1)
IF(T1)602,604,602
602 XO(1)=TI+XO(1)*DEL
604 T2=AIN(2)
IF(T2)606,608,606
606 XO(2)=T2+XO(2)*DEL
608 T3=AIN(3)
IF(T3)612,612,609
609 IF(DEL)611,611,610
610 SCAL1=SCAL1*T3
GO TO 612
611 SCAL1=T3
612 T4=AIN(4)
***** SET ELLIPSOID SCALE FACTOR *****
614 SCAL2=T4
90 TO 616
615 T4=-t4
IF (t4.gt.0. .and. t4.lt.1.) t4=100.*t4
it4=t4
scal2=crtvai(it4)
***** AUTOMATIC ORIGIN AND/OR SCALE *****
616 IF(NJ2-2)790,622,620
620 XO(1)=XLNG(1)*.5
XO(2)=XLNG(2)*.5
622 IF(NJ2-3)625,640,625
625 SCAL1=1.
630 IF(LATM-1)635,635,640
635 NG=12
CALL ERENT(0.D0,602)
CALL EXITING(NG)
640 DO 650 J=1,3

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XMAX(J)=-1.E5
XMIN(J)=1.E5
650 ***** FIT BOX AROUND SET OF ATOMS *****
C DO 670 I=1,LATM
CALL XYZ(ATOMID(I),ATOMS(1,I),3)
IF(NG)652,653,652
652 CALL EREPNT(ATOMID(I),600)
GO TO 670
653 DO 668 J=1,3
TI=ATOMS(J,I)
IF(XMAX(J)-TI)655,660,660
655 XMAX(J)=TI
MAX(J)=I
660 IF(TI-XMIN(J))665,668,668
665 XMIN(J)=TI
668 CONTINUE
670 CONTINUE
C ***** KW=TOP ATOM *****
KW=MAX(3)
SMULT=1.
DO 780 M=1,5
IF(M-2)740,675,678
C ***** CHECK VIEW DISTANCE *****
675 IF(VIEW)785,785,680
678 IF(NJ2-3)680,785,680
680 TI=ATOMS(3,KM)*SMULT
IF(VIEW*5-TI)685,690,690
C ***** INCREASE VIEW DISTANCE *****
685 VIEW=2.*TI
C ***** FIND PERSPECTIVE PROJECTION LIMITS *****
690 DO 700 J=1,2
XMAX(J)=-1.E5
700 XMIN(J)=1.E5
DO 725 I=1,LATM
DO 705 J=1,3
X(J)=ATOMS(J,I)*SMULT
T2=VIEW/(VIEW-X(3))
TI=X(J)*T2
IF(XMAX(J)-TI)710,715,715
710 XMAX(J)=TI
715 IF(TI-XMIN(J))720,725,725
720 XMIN(J)=TI
725 CONTINUE
C ***** REFINER PARAMETERS *****
740 IF(NJ2-3)745,742,755
742 SMUL2=1.
GO TO 765
C ***** AUTOMATIC SCALE ONLY *****
745 DO 750 J=1,2
T2=XO(J)
SCAL(J)=(BRDR-T2)/XMIN(J)
750 SCAL(J+2)=(XLING(J)-BRDR-T2)/XMAX(J)
SMUL2=AMINI(SCAL(1),SCAL(2),SCAL(3),SCAL(4))
GO TO 780
C ***** AUTOMATIC SCALE AND POSITION *****
755 DO 760 J=1,2
760 SCAL(J)=(XLING(J)-BRDR*2.)/(XMAX(J)-XMIN(J))
SMUL2=AMINI(SCAL(1),SCAL(2))
C ***** AUTOMATIC POSITION *****
765 DO 770 J=1,2
XMAX(J)=5*(XLING(J)-SMUL2*(XMAX(J)+XMIN(J)))
780 SMULF=SMULT*SMUL2
VIEW=VIEW*SMUL2
785 SCALI=SCALI*SMULT
790 SCLE=SCALI*SCALI
C ***** ELIMINATE ALL PREVIOUSLY STORED OVERLAP INFORMATION *****
CALL LAP500(0)
RETURN
END
SUBROUTINE F700
C ***** SUBROUTINE TO DRAW ELLIPSOIDS *****
DIMENSION EYE(3),VIEWV(3),X(3),Z(3)
REAL*8 TD,TD2,D100,D1000,D100K
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2,CONT(5),P(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4,ORGN(3),PAC(3,5),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7,XLING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1,IDENT(2,505),MAXATM
common /ns/ npf,ndraw,norient,nvar
C ***** SET ELLIPSOID GRAPHIC DETAILS *****
DI00=100.
DI100=1000.
DI100K=100000.
ITILT=0
NG=0
NFIRST=1
NPLANE=AIN(1)
IF(NPLANE-1)720,715,720
715 NFIRST=4
NPLANE=4
NDOT=64/2** (IABS(NSOLID))
LINES=AIN(3)
NDASH=AIN(4)
CHSYM=AIN(5)
T6=AIN(6)
DH=T6-CHSYM*17./7.
T7=AIN(7)
DV=T7-CHSYM*.5
C ***** ESTABLISH REFERENCE POINT OF VIEW *****
TI=1.E6
IF(VIEW)740,740,735
735 TI=VIEW/SCALI
740 DO 741 J=1,3
741 EYE(J)=REFV(J,3)*TI+ORGN(J)
LNS=-1
C ***** LOOP THROUGH ATOM LIST *****
DO 1105 IATOM=1,LATM
TD=ATOMID(ITOM)
K=TD/D100K
IF(AIN(10))744,744,7412

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7412 IF(AIN(1,2))-1.0D0)742,7414,7416
7414 TD2=IDENT(1,K)
GO TO 7422
7416 TD2=IDENT(2,K)
GO TO 7422
742 TD2=DINT(TD/D100K)
7422 IF(TD2-AIN(10))1105,743,743
743 IF(AIN(11)-TD2)1105,744,744
744 CALL XYZ(TD,X,2)
IF(NG)758,746,758
746 CALL PLTXY(X,Z)
L=DMOD(TD/D100,D1000)
L1=DMOD(TD,D100)
if (ndraw.eq.1) WRITE (npf,750) CHEM(K),K,L,L1,
Z(1)+xtrans,Z(2)+ytrans
747 LNS=MOD(LNS+1,18)
IF(LNS)749,746,749
748 IF (NOUT.GE.0)
&WRITE (NOUT,751)(TITLE(I),I=1,18)
IF (NOUT.GE.0)
&WRITE (NOUT,752)
749 IF (NOUT.GE.0)
&WRITE (NOUT,750) CHEM(K),K,L,L1,Z(1),Z(2),
1(X(I),I=1,3),(XT(I),I=1,3)
750 FORMAT(1H 10X,A6,3H (I3,1H,I3,I2,4H) 2F8.2,3X,3F8.3,11X,3F8.4)
751 FORMAT(1H1,10X,18A4)
752 FORMAT(1H10X,18SYMBOLE, ATOM CODE7X,16HPLOTTER X,Y(IN.) 3X,21HCA
IRTESIAN X,Y,Z (IN.)15X,20HCRYSTAL SYSTEM X,Y,Z/1H 19X,45H(DIRECTIO
3D ON WORKING SYSTEM/1H )
754 IF(EDGE-BRDR*.75)755,760,760
755 NG=10
758 CALL ERENT(TD,700)
GO TO 1105
***** CALL OVERLAP ROUTINE *****
C ICQ=0
CALL LAP700(ITOM,ICQ)
IF(ICQ)762,764,764
***** OMIT HIDDEN ATOM *****
C NG=14
GO TO 758
764 IF(CHSYM)775,775,765
***** PLOT CHEMICAL SYMBOLS *****
C T4=1.
IF(VIEW)767,767,766
766 T4=VIEW/(VLEW-X(3))
767 T3=CHSYM*T4
T4=DISP*T4*.5
V1(1)=X(1)+DH*SYMB(1,1)+DV*SYMB(1,2)
V1(2)=X(2)+DH*SYMB(2,1)+DV*SYMB(2,2)
V1(3)=X(3)
CALL PLTXY(V1,V3)
IF(EDGE-CHSYM)775,768,768
V2(3)=0.
DO 770 I=1,3,2
V2(1)=V3(1)+FLOAT(I-2)*T4
DO 770 J=1,3,2
V2(2)=V3(2)+FLOAT(J-2)*T4
CALL SIMBOLE(V2(1),V2(2),T3,CHEM(K),THETA,6)
IF(T4)775,775,770

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770 CONTINUE
775 IF(NPLANE)1105,1105,780
***** ELLIPSOID PRINC VECTORS TOWARD VIEWER *****
C 780 CALL FAXES(TD,2)
IF(NG)758,783,758
783 CALL DIFV(EYE,XT,VIEWV)
CALL UNITY(VIEWV,VIEWV,-1)
CALL VM(VIEWV,AA,V2)
DO 795 I=1,3
IF(VV(V2,PAT(1,I)))785,795,795
785 DO 790 J=1,3
PAC(J,I)=-PAC(J,I)
790 PAT(J,I)=-PAT(J,I)
795 CONTINUE
DO 800 J=1,3
PAC(J,4)=PAC(J,1)
800 PAC(J,5)=PAC(J,2)
IF(NJ2-10)802,803,803
801 FORMAT(1H ,13X,3(3X,3F8.4,F8.5)/1H )
802 IF (NOUT.GE.0)
&WRITE (NOUT,801) ((PAC(J,K),J=1,3),RMS(K),K=1,3)
***** V4 = VECTOR NORMAL TO POLAR PLANE *****
C 803 continue
CALL VM(VIEWV,AWRK,V6)
CALL UNITY(V6,V6,1)
CALL MV(O,V6,V4)
CALL UNITY(V4,V4,1)
***** SET PLOTTING RESOLUTION FOR ELLIPSOID *****
t3a=sqrt(rms(3)*rms(2))
t3b=sqrt(rms(2)*rms(1))
t3c=sqrt(rms(3)*rms(1))
T3=((t3a+t3b+t3c)/3.0)*SCL
NRESOL=1
NBIS=5
DO 805 J=1,3
IF(T3-RES(J))804,810,810
804 NBIS=NBIS-1
805 NRESOL=NRESOL*2
810 NRESI=NRESOL+1
***** LOOP THROUGH PRINC AND POLAR PLANES *****
C DO 1100 II=NFIRST,NPLANE
II0=MOD(II+2,3)+1
II1=MOD(II,3)+1
II2=MOD(II+1,3)+1
***** GENERATE CONJUGATE DIAMETERS *****
C IF(.99938-ABS(VV(V4,PAC(1,II2))))820,820,830
820 T1=RMS(II0)*SCL
T2=RMS(II1)*SCL
DO 825 J=1,3
DA(J,1)=PAC(J,II0)*T1
825 DA(J,2)=PAC(J,II1)*T2
GO TO 850
830 CALL NORM(PAC(1,II0),PAC(1,II1),V1,1)
CALL NORM(V1,V4,V2,1)
CALL UNITY(V2,V2,1)
CALL MV(O,V2,V3)
IF(II-4)835,840,840
835 CALL NORM(V3,V1,V5,1)
GO TO 843
840 CALL NORM(V3,V4,V5,1)
843 CALL UNITY(V5,V5,1)

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T1=SCL/SORT(VMV(V2,Q,V2))
T2=SCL/SORT(VMV(V5,Q,V5))
DO 845 J=1,3
DA(J,1)=V2(J)*T1
DA(J,2)=V5(J)*T2
C
845 ***** GENERATE ELLIPSE *****
C
850 CALL RADIAL(NBIS)
IF(II-4)900,851,851
851 IF(NSOLID)859,859,859
***** PLOT DOTTED BOUNDARY ELLIPSE *****
C
852 IF(NDOT-NRESOL)853,855,855
853 CALL RADIAL(NSOLID-1)
855 CALL PROJ(D,DP,X,XO,VIEW,1,129,NDOT)
DO 857 J=1,129,NDOT
CALL DRAW(DP(1,J),DISP,DISP,3)
DO 856 I=1,3,2
T1=FLOAT(I-2)*DISP
DO 856 K=1,3,2
T2=FLOAT(K-2)*DISP
CALL DRAW(DP(1,J),T1,T2,2)
IF(DISP)857,857,856
856 CONTINUE
857 CONTINUE
C
859 ***** PLOT SOLID BOUNDARY ELLIPSE *****
CALL PROJ(D,DP,X,XO,VIEW,1,129,NRESOL)
CALL DRAW(DP,0,0,3)
DO 860 J=NRESL,129,NRESOL
CALL DRAW(DP(1,J),0,0,0,2)
IF(DISP)1100,1100,865
C
865 ***** BOUNDARY ANNULUS AS A LINEAR FUNCTION OF HEIGHT *****
T5=VV(VI,AAREV(1,3))*SCAL1
T8=AIN(8)
T9=AIN(9)
NCYCLE=.5+(T8+T5*T9)/DISP
IF(NCYCLE)1100,1100,870
870 T3=(2.*DISP)/(T1+T2)
***** INCREASE ANNULAR THICKNESS *****
C
DO 875 I=1,NCYCLE
T4=T3*FLOAT(I)
DO 875 J=1,129,NRESOL
CALL DRAW(DP(1,J),D(1,J)*T4,D(2,J)*T4,2)
GO TO 1100
C
***** PLOT HALF AN ELLIPSE *****
CALL DRAW(DP,0,0,3)
DO 905 J=NRESL,65,NRESOL
CALL DRAW(DP(1,J),0,0,0,2)
IF(DISP)930,930,910
C
***** ACCENTUATE FRONT HALF *****
DO 910 DO 925 I=1,3,2
T2=FLOAT(I-2)*DISP
DO 915 J=1,65,NRESOL
K=66-J
CALL DRAW(DP(1,K),DISP,T2,2)
DO 925 K=1,65,NRESOL
CALL DRAW(DP(1,K),-DISP,-T2,2)
930 IF(NSOLID)940,967,935
935 L=NDOT
IF(NDOT-NRESOL)938,945,940

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938 CALL RADIAL(NSOLID-1)
GO TO 945
940 L=NRESOL
945 CALL PROJ(D(1,65),DE(1,65),X,XO,VIEW,1,65,L)
***** DOTTED LINE ON REVERSE SIDE *****
C
950 DO 958 J=65,129,NDOT
CALL DRAW(DP(1,J),DISP,DISP,3)
DO 955 I=1,3,2
T1=FLOAT(I-2)*DISP
DO 955 K=1,3,2
T2=FLOAT(K-2)*DISP
CALL DRAW(DP(1,J),T1,T2,2)
IF(DISP)958,958,955
955 CONTINUE
958 CONTINUE
GO TO 967
C
***** SINGLE LINE ON REVERSE SIDE *****
960 DO 965 J=65,129,NRESOL
CALL DRAW(DP(1,J),0,0,0,2)
***** DETAIL INTERIOR FEATURES *****
C
967 T2=NDASH*2
DO 975 J=1,3
T1=PAC(J,II0)*RMS(II0)*SCL
DA(J,1)=T1
DA(J,2)=PAC(J,III)*RMS(III)*SCL
DA(J,3)=0
IF(NDASH)975,975,970
970 V1(J)=-T1
V2(J)=T1/T2
975 CONTINUE
IF(NDASH)987,987,980
***** DASHED LINE FOR REVERSE AXIS *****
C
DO 985 J=1,NDASH
DO 985 K=1,2
L=4-K
CALL PROJ(V1,DP,X,XO,VIEW,1,1,1)
CALL DRAW(DP,0,0,L)
DO 985 I=1,3
V1(I)=V1(I)+V2(I)
C
***** SOLID LINE FOR FORWARD AXIS *****
987 IF(LINES)1100,1100,988
988 CALL PROJ(DA,DP,X,XO,VIEW,1,3,1)
T1=DISP*.5
DO 990 I=1,3,2
T2=FLOAT(2-I)*T1
CALL DRAW(DP,T1,T2,3)
CALL DRAW(DP(1,3),T1,T2,2)
IF(DISP)1000,1000,989
989 CALL DRAW(DP(1,3),-T1,T2,2)
990 CALL DRAW(DP,-T1,T2,2)
C
***** SHADE QUADRANT BETWEEN TWO PRINCIPAL AXES *****
1000 L=LINES-1
IF(L)1100,1100,1005
1005 T2=LINES
DO 1025 I=1,L
T1=FLOAT(I)/T2
T3=SQRT(1.-T1*T1)
IF(MOD(I,2))1010,1015,1010
1010 M=I*2
N=M-1

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1015 GO TO 1020
N=I*2
M=N-1
1020 DO 1025 J=1,3
T4=DA(J,1)*T1
D(J,M)=T4
1025 D(J,N)=DA(J,2)*T3+T4
I=L*2
CALL PROJ(D,DP,X,XO,VIEW,1,L,1)
DO 1030 I=2,L,2
CALL DRAW(DP(I,I-1),0.,0.,3)
1030 CALL DRAW(DP(I,I),0.,0.,2)
1100 CONTINUE
1105 ***** ELIMINATE LOCAL OVERLAP INFORMATION BEFORE RETURNING *****
RETURN
END
SUBROUTINE F800
***** SUBROUTINE FINDS ATOM PAIRS FOR BONDS *****
DIMENSION IA(3),M1(6)
REAL*8 TD1,TD2,TD3,D100K
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AARX(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NU,NJ2,INOUT,NSR,NSIM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCH,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATOM
COMMON/OLAF/CONIC(7,500),COVER(6,20),KC(20),KQ(30),NCONIC,NCOVER,
1 ,NOOVER,NOVAD,OVNRGN,QCOVER(3,4,30),QUAD(9,600),SEGM(50,2)
D100K=100000.
C *** old
C *** new
IAN=AIN(2)
NJ4=(MOD(NJ2,10)-4)+(IAN*2)
if (nj.eq.10) nj4=(ian*2)-2
INS=-4
IF(MOD(NJ2,10)-2)805,848,848
***** EXPLICIT DESCRIPTION *****
805 II=0
IF(NCD)810,810,815
810 NG=11
CALL ERENT(0,D0,NJ*100+NJ2)
GO TO 980
815 II=II+1
IF(140-II)980,980,820
820 TD1=AIN(II)
IF(TD1)815,815,825
825 II=II+1
TD2=AIN(II)
IF(TD2)815,815,830
830 IF(NJ2-10)832,838,838

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832 LNS=MOD(LNS+4,56)
IF(LNS)838,834,838
834 IF (NOUT.GE.0)
&WRITE (NOUT,835)(TITLE(I),I=1,18)
835 FORMAT(1H1,10X,18A4)
IF (NOUT.GE.0)
&WRITE (NOUT,837)
837 FORMAT(1H010X,18HSYMBOL ATOM CODE6X,16HPLOTTER X,Y(IN.) 6X,21HCA
1RTEJIAN X,Y,Z (IN.,17X,20HCRYSTAL SYSTEM X,Y,Z/1H )
***** CHECK IF BOND ATOMS ARE IN ATOMS LIST (FOR OVERLAP CALC) ***
838 NA1=0
NA2=0
IF(LATM-2)845,839,839
839 N2=2
DO 844 K=1,LATM
TD3=ATOMID(K)
IF(TD3-TD1)841,840,841
840 NA1=K
GO TO 843
841 IF(TD3-TD2)844,842,844
842 NA2=K
843 N2=N2-1
IF(N2)845,845,844
844 CONTINUE
845 IF(NA2-NA1)846,847,847
846 NA3=NA1
NA1=NA2
NA2=NA3
TD3=TD1
TD1=TD2
TD2=TD3
847 CALL BOND(TD1,TD2,1,NA1,NA2)
GO TO 815
***** IMPLICIT DESCRIPTION *****
848 IF(LATM-2)810,850,850
850 SCAL3=SCAL1
SCAL1=1.
DO 855 I=1,LATM
CALL XYZ(ATOMID(I),ATOMS(1,I),2)
SCAL1=SCAL3
IF(NCD)810,810,860
860 IF (NOUT.GE.0)
&WRITE (NOUT,861)
861 FORMAT(1H010X,20HBOND SELECTION CODES//11X,94H(SEQUENCE(A)) (SEQUEN
1CE(B)) (BOND) (DISTANCES) ( BOND ) (PERSP.--LABELS) (NORMAL--LABELS) (
2DIGITS) /11X,93H( MIN MAX ) ( TYPE) ( MIN MAX) (RADI
3US) (HEIGHT OFFSET) (HEIGHT OFFSET) (NUMBER)
DMAX=0.
DO 870 I=1,NCD
IF(DMAX-CD(2,I))865,866,866
865 DMAX=CD(2,I)
866 IF (NOUT.GE.0)
&WRITE (NOUT,871)(KD(J,I),J=1,5),(CD(J,I),J=1,8)
870 CONTINUE
871 FORMAT(1H ,10X,I6,I5,I8,I5,I8,2F6.2,5F8.3,F7.0)
***** LOOP THROUGH ATOMS ARRAY *****
DO 977 N=1,LATM
NA1=N
TD1=ATOMID(M)
MI=TD1/D100K

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8718 IF(NJ4-2)8724,8724,8718
8720 IF(NJ4-4)8726,8726,8720
8722 IA(1)=TD1/D100K
GO TO 8728
8724 IA(1)=IDENT(1,MI)
GO TO 8728
8726 IA(1)=IDENT(2,MI)
8728 IA(3)=IA(1)
WI(1)=ATOMS(1,M)
WI(2)=ATOMS(2,M)
WI(3)=ATOMS(3,M)
L=M+1
IF(LATM=L)977,872,872
DO 975 N=L,LATM
NR2=N
DIST=(ATOMS(1,N)-WI(1))*2
IF(DMAX-DIST)975,873,873
873 DIST=DIST+(ATOMS(2,N)-WI(2))*2
IF(DMAX-DIST)975,874,874
874 DIST=DIST+(ATOMS(3,N)-WI(3))*2
IF(DMAX-DIST)975,875,875
875 DIST=SQRT(DIST)
TD2=ATOMID(N)
NI=TD2/D100K
IF(NJ4)876,877,878
C 876 IA(2)=TD2/D100K
C 877 IA(2)=IDENT(1,NI)
C 878 IA(2)=IDENT(2,NI)
IF(NJ4.LT.0) IA(2)=TD2/D100K
IF(NJ4.EQ.0.OR.NJ4.EQ.1) IA(2)=IDENT(1,NI)
IF(NJ4.GT.1) IA(2)=IDENT(2,NI)
C ***** SELECT BONDS ACCORDING TO CODES *****
C 879 DO 950 J=1,NCDD
JB=J
IF(DIST-CD(1,J)) 950,880,880
880 IF(CD(2,J)-DIST) 950,881,881
881 DO 885 K=1,2
IF(IA(K)-KD(1,J)) 885,882,882
882 IF(KD(2,J)-IA(K)) 885,883,883
883 IF(IA(K+1)-KD(3,J))885,884,884
884 IF(KD(4,J)-IA(K+1))885,890,890
885 CONTINUE
GO TO 950
C ***** CHECK FOR POLYHEDRA CODE *****
890 IF(CD(4,J)1900,955,955
900 W1(4)=ATOMS(1,N)
W1(5)=ATOMS(2,N)
W1(6)=ATOMS(3,N)
KML=ABS(CD(4,J))
KMZ=ABS(CD(5,J))
DSQ1=(CD(6,J))*2
DSQ2=(CD(7,J))*2
C ***** SEARCH FOR POLYHEDRA CENTER *****
DO 935 IM=1,LATM
K3=ATOMID(IM)/D100K
IF(IAN.EQ.1) K3=IDENT(1,IM)
IF(IAN.EQ.2) K3=IDENT(2,IM)
IF(K3-KM1)935,905,905

905 IF(KM2-K3)935,910,910
***** CHECK POLYHEDRA DISTANCE RANGE *****
910 DO 930 J1=1,4,3
DSQ=(ATOMS(1,IM)-WI(J1))*2
IF(DSQ2-DSQ)935,915,915
915 DSQ=DSQ+(ATOMS(2,IM)-WI(J1+1))*2
IF(DSQ2-DSQ)935,920,920
920 DSQ=DSQ+(ATOMS(3,IM)-WI(J1+2))*2
IF(DSQ2-DSQ)935,925,925
925 IF(DSQ-DSQ1)935,930,930
930 CONTINUE
GO TO 955
935 CONTINUE
***** END OF POLYHEDRA CHECK *****
C
GO TO 975
***** PREPARE TO DRAW BOND *****
955 IF(NJ2-10)960,970,970
960 LNS=MOD(LNS+4,56)
IF(LNS)965,965,970
965 IF(NOUT.GE.0)
&WRITE(NOUT,835)(TITLE(I),I=1,18)
IF(NOUT.GE.0)
&WRITE(NOUT,837)
970 CALL BOND(TD1,TD2,JB,NA1,NA2)
975 CONTINUE
977 CONTINUE
C ***** ELIMINATE LOCAL OVERLAP INFORMATION BEFORE RETURNING *****
980 IF(NJ2-21)985,990,990
985 CALL LAP500(-1)
990 IF(NJ2.EQ.22) THEN
IF(NQUAD)993,993,991
***** PRINT OUT NUMBER OF BOND QUADRANGLES STORED *****
C ***** PRINT OUT QUADRANGLE IDENTIFICATION ARRAY *****
991 IF(NOUT.GE.0)
&WRITE(NOUT,992)NQUAD,(QUAD(9,J),J=1,NQUAD)
992 FORMAT(1H0,10X,27HBOND OVERLAP ARRAY CONTAINS,14,23H BONDS (MAXIMU
IM IS 599)/ 11X, 66HATOM-PAIR NUMBERS IN ARRAY REFER TO SEQUENCE
2IN SORTED ATOMS ARRAY/(15X,10F10.0))
END IF
993 RETURN
END
SUBROUTINE F900
DIMENSION X(3),XW(3,5),Y(3),Z(3)
REAL*8 D100K
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 IN,ITILT,KD(5,20),IATM,NATOM,NCDD,NJ,NJ2,NOUT,NSR,NSYM
4 ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),VRKV(3,3)
7 XLNG(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
character*72 tmp1, tmp2
***** LABELING FUNCTION SUBROUTINE *****
C

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      if (tmpti(i:i) .ne. ' ') then
        klast = i
        go to 104
      end if
      continue
      ioffset = (72 - klast) / 2
      do 105 i=1,klast
        tmpti2(i+ioffset:i+ioffset) = tmpti(i:i)
      continue
      do 106 i=1,18
        titlez(i) = tmpti2(i*4-3:i*4)
      continue
      end if
      DO 1065 I=1, ILAST
      DO 1075 J=1, 3, 2
      Z(1)=Y(1)+FLOAT(J-2)*DISP*.5
      DO 1075 K=1, 3, 2
      Z(2)=Y(2)+FLOAT(K-2)*DISP*.5
      IF(NJ3-2)1065,1068,1068
      ***** PLOT CHEMICAL SYMBOL *****
      C 1065 CALL SYMBOL(Z(1),Z(2),HGT,CHEM(L),TH,6)
      GO TO 1070
      C ***** PLOT TITLES *****
      1068 CALL SYMBOL(Z(1),Z(2),HGT,TITLE2(I),TH,4)
      1070 IF(DISP)1080,1080,1075
      1075 CONTINUE
      1080 Y(1)=Y(1)+DXW
      1085 Y(2)=Y(2)+DYW
      GO TO 1199
      C ***** PLOT BOND DISTANCE LABELS *****
      I9=NJ3-3
      T9=10.**I9
      DISTP=AJNT((DIST*T9)+0.5)/T9 + 0001
      CALL NUMBER(Y(1),Y(2),HGT,DISTR,TH,I9)
      GO TO 1199
      C ***** PLOT CENTERED SYMBOLS *****
      I105 TT8=AIN(8)
      C *** ORTEP-II call
      CALL SYMBOL(Y(1),Y(2),HGT,ITX(TT8),TH,7-NJ3)
      C *** Only one centered symbol (*) is available in ORTEP-III.
      C *** It is triggered by the negative value for argument 6.
      C *** Argument 4 is ignored by SYMBOL.
      CALL SYMBOL(Y(1),Y(2),HGT,' ',TH,7-NJ3)
      GO TO 1199
      912 NG=15
      915 CALL ERENT(AIN(II),NU*100+NUJ2)
      1199 ITILT=0
      RETURN
      END
      SUBROUTINE F1000
      C *** 1001 identical to 511
      CALL LAP500(1)
      RETURN
      END
      function iend(string)
      C *** returns position of last non-space character in string
      C *** character string*(*)
      do 800 i=len(string),1,-1
      if (string(i:i) .ne. ' ') then
        iend = i
      return
    end if
    continue
    iend = 1
    return
  end
end
SUBROUTINE LAP500 (NTYPE)
***** STORE PROJECTED ATOM CONICS AND BOND QUADRANGLES *****
DIMENSION OC(3,3),OD(3,3),VDI(3),VD2(3)
REAL*8 OC,OD,VD1,VD2,TD1,TD2,TD3,TD
COMMON/OLAP/CONIC(7,500),COVER(6,20),KC(20),KQ(30),NCONIC,NCOVER,
1 NCOVER,NQUAD,OVMRGN,QCOVER(3,4,30),QUAD(9,600),SEGM(50,2)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AARWK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NU,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WERKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
***** ELIMINATE ALL PREVIOUSLY STORED LOCAL OVERLAP INFORMATION **
C NCOVER=0
NCOVER=0
C ***** CONSTANT FOR OVERLAP MARGIN (WHITE MARGIN AT OVERLAP) *****
IF(NTYPE)420,195,195
C ***** ELIMINATE ALL PREVIOUSLY STORED GLOBAL OVERLAP INFORMATION *
***** NCONIC=0
NQUAD=0
C IF(NTYPE)420,420,200
***** CONSTANT FOR OVERLAP MARGIN (WHITE MARGIN AT OVERLAP) *****
200 IF(AIN(1))205,215,210
***** NEGATIVE NUMBER OR POSITIVE INTEGER GIVES OVMRGN=0.0 *****
205 OVMRGN=0.0
GO TO 220
C ***** SET OVERLAP MARGIN WIDTH DIRECTLY IN INCHES *****
210 OVMRGN=AIN(1)-DINT(AIN(1))
GO TO 220
C ***** DEFAULT OPTION, OVERLAP MARGIN WIDTH AS A FUNCTION OF SCAL1
215 if (scal1.lt..25) then
else
OVMRGN=AMAX1(SQRT(SCAL1)*0.050,0.010)
end if
220 IF (NOUT.GE.0)
&WRITE (NOUT,2) OVMRGN
2 FORMAT(1H0,10X,17HOVERLAP MARGIN IS, F6.3,5H INCH)
225 IF(LATM)230,230,235
230 NG=12
CALL ERENT(0.D0,510+NUJ2)
GO TO 420
C ***** SORT ATOMS LIST BY -VIEWDISTANCE OR BY Z PARAMETER
235 IF(VIEW)250,250,240
***** CALCULATE VIEWDISTANCES**2*(-1) IF VIEW.GT.ZERO *****
240 DO 245 I=1,LATM
CALL XYZ(ATOMID(I),V3,2)
V3(3)=V3(3)-VIEW
245 ATOMS(3,I)=-VV(V3,V3)
GO TO 260
***** STORE CARTESIAN COORDINATES IF VIEW.EQ.ZERO *****
250 DO 255 I=1,LATM
255 CALL XYZ(ATOMID(I),ATOMS(1,I),2)

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C ***** SORTING PROCEDURE BY SHELL, COMM ACM 2,30 (1959) *****
260 M=LATM
265 M=M/2
270 IF(M)300,300,270
    J=1
275 I=J
280 IM=I+M
285 TD=ATOMID(I)
    ATOMID(I)=ATOMID(IM)
    ATOMID(IM)=TD
    TL=ATOMS(3,I)
    ATOMS(3,I)=ATOMS(3,IM)
    ATOMS(3,IM)=TL
    I=I-M
    IF(I)295,295,280
    J=J+1
295 IF(J-K)275,275,265
C ***** LOOP THROUGH ALL ATOMS IN SORTED ATOMS LIST *****
300 DO 405 IA=1, LATM
    CALL XYZ(ATOMID(IA),ATOMS(1,IA),2)
    CALL FAXES(ATOMID(IA),2)
305 DO 305 J=1,3
    QD(J,K)=Q(J,K)
C ***** CALCULATE ENVELOPING CONE WITH ORIGIN AT VIEWPOINT *****
310 V1(3)=V1(3)-V1EW
    VD1(3)=V1(3)
C ***** FORM COFACTOR MATRIX *****
315 DO 315 J=1,3
    J1=MOD(J,3)+1
    J2=MOD(J+1,3)+1
    DO 315 K=J,3
    K1=MOD(K,3)+1
    K2=MOD(K+1,3)+1
    QC(J,K)=QD(J1,K1)*QD(J2,K2)-QD(J1,K2)*QD(J2,K1)
315 QC(K,J)=QC(J,K)
C ***** FORM POLARIZED COFACTOR MATRIX AND ADD TO ELLIPSOID MATRIX *
    TD2=-SCL**2
C ***** TD1 IS AN ARBITRARY SCALING FACTOR *****
    TD1=VMV(V1,0,V1)
    DO 325 J=1,3
    J1=MOD(J,3)+1
    J2=MOD(J+1,3)+1
    DO 320 K=J,3
    K1=MOD(K,3)+1
    K2=MOD(K+1,3)+1
    QD(J,K)=(VD1(J2)*QC(J1,K1)*VD1(K2)-QC(J1,K2)*VD1(K1))
    1 +VD1(J1)*VD1(K1)*QC(J2,K2)-VD1(K2)*QC(J2,K1))+TD2*QD(J,K))/TD1
320 QD(K,J)=QD(J,K)
C ***** PROJECTED ELLIPSE IN HOMOGENEOUS COORD OF WORKING SYSTEM ***
    QD(J,3)=-QD(J,3)*VIEW
325 QD(3,J)=-QD(3,J)*VIEW
C ***** PROJECT CENTER OF ATOM ONTO PROJECTION PLANE *****
    TD1=-VIEW/VD1(3)
    VD2(1)=VD1(1)*TD1
    VD2(2)=VD1(2)*TD1
C ***** TRANSFORM TO NEW ORIGIN TO IMPROVE CONDITION OF MATRIX Q ***
330 J=1,3
    DO 330 K=1,2
    330 QD(J,3)=QD(J,3)+QD(J,K)*VD2(K)
    DO 335 J=1,3
    DO 335 K=1,2
    335 QD(3,J)=QD(3,J)+VD2(K)*QD(K,J)
    V6(1)=XO(1)+VD2(1)
    V6(2)=XO(2)+VD2(2)
    GO TO 355
C ***** CALCULATE ENVELOPING CYLINDER ALONG Z OF WORKING SYSTEM *****
340 DO 345 J=1,2
    DO 345 K=1,2
    345 QD(J,K)=QD(J,K)-QD(J,3)*QD(K,3)/QD(3,3)
    DO 350 J=1,2
    QD(J,3)=0.0
    QD(3,J)=0.0
350 V6(J)=XO(J)+ATOMS(J,IA)
C ***** PROJECTED ELLIPSE IN HOMOGENEOUS COORD ABOUT CENTER OF ATOM
    QD(3,3)=-SCL**2
C ***** FIT RECTANGLE AROUND ELLIPSE ALLOWING OVERLAP MARGIN *****
C ***** FORM MATRIX OF COFACTORS *****
355 DO 360 J=1,3
    J1=MOD(J,3)+1
    J2=MOD(J+1,3)+1
    DO 360 K=J,3
    K1=MOD(K,3)+1
    K2=MOD(K+1,3)+1
    360 QC(J,K)=QD(J1,K1)*QD(J2,K2)-QD(J1,K2)*QD(J2,K1)
C ***** RESCALE MATRIX OF COFACTORS SO THAT QC(3,3)=1.0 *****
    DO 365 J=1,3
    DO 365 K=J,3
    QC(J,K)=QC(J,K)/QC(3,3)
    365 QC(K,J)=QC(J,K)
    TD2=QD(3,3)
    NDG=0
C ***** SOLVE QUADRATIC EQUATION *****
    DO 385 J=1,2
    TL=QC(3,J)**2-QC(J,J)
    IF(TL)370,370,375
C ***** ROUND OFF PROBLEMS, RESET LIMITS IN X OR Y *****
    370 NDG=1
    V5(J)=0.001+OVMRGN
    GO TO 380
375 V5(J)=SORT(TL)+OVMRGN
    V6(J)=V6(J)+QC(3,J)
    TD2=TD2+QD(3,J)*QC(3,J)
380 CONIC(2*J-1,IA)=V6(J)-V5(J)
385 CONIC(2*J,IA)=V6(J)+V5(J)
    IF(NDG)390,390,395
390 IF(TD2)400,395,395
C ***** ELLIPSE IMAGINARY DUE TO ROUND OFF, RESET TO REAL VALUE *****
    395 CONIC(5,IA)=1.0/((CONIC(2,IA)-CONIC(1,IA))*0.5)**2
    CONIC(6,IA)=0.0
    CONIC(7,IA)=1.0/((CONIC(4,IA)-CONIC(3,IA))*0.5)**2
    GO TO 405
C ***** STORE NORMALIZED QUADRATIC COEFFICIENTS FOR ELLIPSE *****
C ***** SCALED BY OVERLAP MARGIN PARAMETER *****
    400 TD3=-((1.0-2.0*OVMRGN/(V5(1)+V5(2)))**2)/TD2
    CONIC(5,IA)=QD(1,1)*TD3
    CONIC(6,IA)=QD(1,2)*TD3

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405 CONIC(7,IA)=QB(2,2)*TD3
CONTINUE
NCONIC=LATM
***** PRINT OUT SORTED ATOMS ARRAY *****
C IF (NOUT.GE.0)
&WRITE (NOUT,4) (ATOMID(J),J=1,LATM)
4 FORMAT(1H0,10X,30HCNENTS OF SORTED ATOMS ARRAY/(15X,10F10.0))
***** STORE BOND QUADRANGLES IF SEARCH CODES ARE GIVEN *****
C IF(NCD)420,420,410
***** GENERATE PSEUDO-INSTRUCTION 822 TO CALCULATE BONDS *****
410 NJ2=22
CALL F800
C *** the lines below have been moved to the end of F800
C IF(NQUAD)420,420,415
C ***** PRINT OUT NUMBER OF BOND QUADRANGLES STORED *****
C ***** PRINT OUT QUADRANGLE IDENTIFICATION ARRAY *****
C 415 IF (NOUT.GE.0)
&WRITE (NOUT,6)NQUAD,(QUAD(9,J),J=1,NQUAD)
6 FORMAT(1H0,10X,27HBOND OVERLAP ARRAY CONTAINS,I4,23H BONDS (MAXIMU
IM IS 599)/ 11X, 66HATOM-PAIR NUMBERS IN ARRAY REFER TO SEQUENCE
2IN SORTED ATOMS ARRAY/(15X,10F10.0))
C *** the lines above have been moved to the end of F800
420 RETURN
END
SUBROUTINE LAP700(NA,ICQ)
DIMENSION DETER(2),QA(3,3,2),QC(3,3,2),VL2(3,2),YMIN(2),YMAX(2)
DIMENSION OVMR(2)
REAL*8 AOV3,AOV3SQ,BOV3,DETER,PI,PHI,POV3,POV3CU,QA,OC,QOV2,QOV2SQ
REAL*8 ROOF,TD,TIDD
REAL*8 AIN,ATOMID
1 NCOVER,NQUAD,OVMRGN,QCOVER(3,4,30),QUAD(9,600),SEGM(50,2)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AARWK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FSR,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XLING(3),XO(3),XT(3)
PI=3.1415926535897932
ICQ=0
NCOVER=0
NCOVER=0
OVMR(1)=OVMRGN
OVMR(2)=0.0
IF(NCONIC=NA)200,200,205
200 RETURN
C ***** ROUGH CHECK FOR OVERLAPPING ATOMS *****
205 DO 210 J=1,2
YMIN(J)=CONIC(2*J-1,NA)
210 YMAX(J)=CONIC(2*J,NA)
L=0
DO 420 IA=NA,NCONIC
IF(IA=NA)230,230,215
215 DO 225 J=1,2
IF(YMAX(J)-CONIC(2*J-1,IA))420,420,220
220 IF(YMIN(J)-CONIC(2*J,IA))225,420,420
225 CONTINUE
C ***** EXACT CHECK FOR OVERLAPPING ATOMS *****

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230 IF(L-1)235,235,240
235 L=L+1
240 CALL LAPCON(ONIC(1,IA),DA,VL2(1,L),OVMR(L))
DO 245 J=1,3
DO 245 K=1,3
245 QA(J,K,L)=DA(J,K)
***** CALCULATE COFACTORS AND DETERMINANTS *****
DETER(L)=0.0
DO 250 J=1,3
J1=MOD(J+3,3)+1
J2=MOD(J+1,3)+1
DO 250 K=1,3
K1=MOD(K+3,3)+1
K2=MOD(K+1,3)+1
TD=QA(J1,K1,L)*QA(J2,K2,L)-QA(J1,K2,L)*QA(J2,K1,L)
DETER(L)=DETER(L)+TD*QA(J,K,L)
250 QC(J,K,L)=TD
C ***** DETER(L) IS THE DETERMINANT TIMES 3 *****
IF(L-1)420,420,255
***** FORM CHARACTERISTIC EQUATION AND EXAMINE ITS ROOTS *****
255 AOV3=0.0
BOV3=0.0
DO 260 J=1,3
DO 260 K=1,3
AOV3=AOV3+QC(J,K,2)*QA(J,K,1)
BOV3=BOV3+QC(J,K,1)*QA(J,K,2)
AOV3SQ=AOV3/DETER(2)
BOV3=BOV3/DETER(2)
POV3=BOV3-AOV3SQ
QOV2=AOV3*(AOV3SQ-BOV3*1.5D0)+DETER(1)/(DETER(2)*2.0D0)
***** CHECK DISCRIMINANT OF CHARACTERISTIC CUBIC EQUATION *****
ITYPE=0
POV3CU=POV3**3
QOV2SQ=QOV2**2
IF(POV3CU+QOV2SQ)270,310,310,400
265 IF(POV3CU*1.00001+QOV2SQ)310,310,400
270 IF(POV3CU+1.00001*QOV2SQ)275,310,310
***** THREE REAL ROOTS, ALL DIFFERENT *****
C 275 ITYPE=1
C ***** NO INTERSECTION IF A/3 AND B/3 INVARIANTS ARE NEGATIVE *****
IF(AOV3)280,285,285
280 IF(BOV3)420,285,285
***** CALCULATE ONE ROOT OF CHARACTERISTIC CUBIC EQUATION *****
285 IF(QOV2)295,290,295
290 PHI=PI/2.0D0
GO TO 305
295 PHI=DATAN(-DSQRT(-POV3CU-QOV2SQ)/QOV2)
300 PHI=PHI+PI
305 ROOT=-2.0D0*DSQRT(-POV3)*DCOS(PHI/3.0D0)-AOV3
GO TO 325
C ***** THREE REAL ROOTS, AT LEAST TWO ARE EQUAL *****
310 ITYPE=2
C ***** CHECK SIGNS OF INVARIANTS A/3 AND B/3 *****
IF(AOV3)315,320,320
315 IF(BOV3)420,320,320
***** CALCULATE REPEATED ROOT OF CUBIC EQUATION *****
320 ROOT=DSIGN(DSQRT(-POV3),QOV2)-AOV3
C ***** FORM DEGENERATE CONIC (LINE PAIR WHICH MAY BE COINCIDENT) **
325 DO 330 J=1,3

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330 DA(J,K)=QA(J,K,1)+ROOT*QA(J,K,2)
***** EXAMINE INVARIANTS OF THE DEGENERATE CONIC *****
C
T6=DA(1,1)*DA(2,2)
T7=DA(1,2)**2
C
***** NEGATIVE DENOTES REAL INTERSECTING LINE PAIR *****
C
***** POSITIVE DENOTES IMAGINARY LINES INTERSECTING AT REAL POINT
C
335 IF(T6-T7)335,345,340
340 IF(T6*1.0001-T7)400,345,345
345 T8=DA(3,3)*(DA(1,1)+DA(2,2))
T9=DA(1,3)**2+DA(2,3)**2
C
***** NEGATIVE DENOTES REAL PARALLEL LINE PAIR *****
C
***** POSITIVE DENOTES IMAGINARY PARALLELS *****
C
***** ZERO DENOTES ONE REAL LINE (COINCIDENT PARALLELS) *****
C
IF(T8-T9)350,360,355
350 IF(T8*1.0001-T9)400,360,360
355 IF(T8-1.0001*T9)360,360,365
C
***** COINCIDENT LINE PAIR FOUND FOR THE REPEATED ROOT *****
C
360 ITYPE=3
***** COMPARE AREAS OF CONICS *****
C
365 KA=1
KB=2
IF(QC(3,3,KA)-QC(3,3,KB))370,375,375
370 KA=2
KB=1
C
***** SEE IF ONE CONIC IS INSIDE THE OTHER CONIC *****
C
375 T1=0.0
DO 385 J=1,3
T2=QA(J,3,KB)
DO 380 K=1,2
380 T2=T2+QA(J,K,KB)*V12(K,KA)
385 T1=T1+V12(J,KA)*T2
C
***** DISCARD IF KA IS OUTSIDE KB *****
C
IF(T1)390,390,420
390 IF(KA-1)395,395,400
C
***** THE OVERLAPPING ATOM HIDES THE ORIGINAL ATOM *****
C
395 ICQ=-1
RETURN
C
***** STORE OVERLAPPING ATOM *****
C
400 ICQ=ICQ+1
IF(NCOVER=20)410,405,405
405 NG=17
CALL ERENT(ATOMID(IA),700)
NCOVER=NCOVER-1
410 NCOVER=NCOVER-1
IU=1
DO 415 I=1,3
DO 415 J=I,3
COVER(IJ,NCOVER)=QA(I,J,2)
415 IJ=IJ+1
KC(NCOVER)=IA
420 CONTINUE
C
***** SECOND PART OF SUBROUTINE CHECKS FOR BONDS OVER THE ATOM ***
C
425 IF(NQUAD)470,470,430
430 ITY=0
***** ROUGH CHECK FOR OVERLAPPING BONDS *****
C
DO 465 IQ=1,NQUAD
TIDD=TID
NA1=TTID/1000.

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NA2=AMOD(TID,1000.)
IF(NA-NA2)435,435,465
DO 445 J=1,2
IF(YMAX(J)-AMIN1(QUAD(J,IQ),QUAD(J+2,IQ),QUAD(J+4,IQ),QUAD(J+6,IQ),
1)465,465,440
1)465,465,440
440 IF(YMIN(J)-AMAX1(QUAD(J,IQ),QUAD(J+2,IQ),QUAD(J+4,IQ),QUAD(J+6,IQ),
1)445,445,465
445 CONTINUE
***** EXACT CHECK FOR OVERLAPPING BONDS *****
C
450 ITY=ITY-1
IQQ=0
IQR=IQ
CALL LAPAB(IQR,NA,IQQ,ITY)
IF(IQQ)455,460,460
455 ICQ=-1
RETURN
460 ICQ=ICQ+1
IF(NCOVER=30)465,470,470
465 CONTINUE
470 RETURN
END
SUBROUTINE LAP800(NA1,NA2,ICQ)
***** SUBROUTINE CHECKS FOR ATOMS AND BONDS OVERLAPPING A BOND ***
DIMENSION FL(4,4),Y1(2),Y2(2),YMAX(2),YMIN(2),QUA(3,4)
DIMENSION VUE(3)
REAL*8 TIDD
COMMON/OLAP/CONIC(7,500),COVER(6,20),KC(20),KQ(30),NCONIC,NCOVER,
1 NCOVER,NQUAD,OVNRGN,QOVER(3,4,30),QUAD(9,600),SEGM(50,2)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 CONI(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCALI
5 ,SCAL2,SCI,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XLANG(3),XO(3),XT(3)
IQ=0
ICQ=0
IF(NA1*NA2)245,245,195
TID1=FLOAT(NA1)*1000.+FLOAT(NA2)
IF(NCONIC)245,245,200
200 IF(NJ2-21)250,205,205
***** PART 1, CALLED FROM BOND, STORES BOND OUTLINE INFORMATION **
C
205 IF(NQUAD=599)215,210,210
210 NG=16
CALL ERENT(ATOMID(NA1),822)
GO TO 245
215 NQUAD=NQUAD+1
***** CALCULATE OVERLAP MARGIN FOR BOND QUADRANGLE *****
C
T1=0.0
T2=0.0
DO 220 J=1,2
Y1(J)=DP(J,1)-DP(J,65)
Y2(J)=DP(J,2)-DP(J,66)
T1=T1+Y1(J)**2
220 T2=T2+Y2(J)**2
IF(T1*T2)225,225,230
225 T1=0.0
T2=0.0

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GO TO 235
230 T1=OVMRGN/SORT(T1)
    T2=OVMRGN/SORT(T2)
    ***** STORE BOND QUADRANGLE *****
C
235 DO 240 J=1,2
    Y1(J)=Y1(J)*T1
    Y2(J)=Y2(J)*T2
    QUAD(J,NQUAD)=DP(J,1)+Y1(J)
    QUAD(J+2,NQUAD)=DP(J,2)+Y2(J)
    QUAD(J+4,NQUAD)=DP(J,66)-Y2(J)
    QUAD(J+6,NQUAD)=DP(J,65)-Y1(J)
240 QUAD(J+6,NQUAD)=DP(J,65)-Y1(J)
245 RETURN
C
250 ***** PART 2, CALLED FROM BOND, OVERLAP CHECK FOR BOND NAL1-NA2 ***
    NCOVER=0
    NCOVER=0
    TOL=1.E-5
    IF(NCONIC-NAL1)245,245,255
    ***** SAVE QUADRANGLE TEMPORARILY *****
C
255 IQ=NQUAD+1
    DO 260 J=1,2
    QUAD(J,IQ)=DP(J,1)
    QUAD(J+2,IQ)=DP(J,2)
    QUAD(J+4,IQ)=DP(J,66)
    QUAD(J+6,IQ)=DP(J,65)
260 QUAD(J+6,IQ)=DP(J,65)
C
265 DO 270 J=1,2
    YMIN(J)=AMINI(DP(J,1),DP(J,2),DP(J,66),DP(J,65))
    YMAX(J)=AMAX1(DP(J,1),DP(J,2),DP(J,66),DP(J,65))
    ***** ROUGH CHECK FOR ATOM-OVER-BOND OVERLAP *****
C
    NALP1=NA1+1
    ITY=0
    DO 305 IA=NA1P1,NCONIC
    DO 285 J=1,2
    IF(IA-NA2)275,305,275
    IF(YMAX(J)-CONIC(2*J-1,IA))305,305,280
    275 IF(YMIN(J)-CONIC(2*J,IA))285,305,305
    280 IF(YMIN(J)-CONIC(2*J,IA))285,305,305
    285 CONTINUE
    ***** CHECK FOR TRUE ATOM-OVER-BOND OVERLAP *****
C
    ITY=ITY+1
    IAQ=IA
    CALL LAPAB(IQ,IAQ,IQO,ITY)
    IF(IQO)290,305,300
    300 ICQ=ICQ+1
    IF(NCOVER-20)305,310,310
    305 CONTINUE
    310 IF(NQUAD)295,295,315
    ***** HIDDEN BOND *****
C
    290 ICQ=-1
    295 RETURN
    ***** ROUGH CHECK FOR BOND-OVER-BOND OVERLAP *****
C
    315 CALL DIFV(ATOMS(1,NA2),ATOMS(1,NA1),V1)
    CALL UNITY(V1,V1,1)
    VUE(1)=ATOMS(1,NA1)
    VUE(2)=ATOMS(2,NA1)
    VUE(3)=ATOMS(3,NA1)-VIEW
    DO 495 IB=1,NQUAD
    TID2=QUAD(9,IB)
    IF(TID1-TID2)320,495,320
320 NB2=AMOD(TID2,1000.)
    NB1=TID2/1000.
    IF(NAL-NB2)325,495,495
    DO 335 J=1,2
    IF(YMAX(J)-AMINI(QUAD(J,IB),QUAD(J+2,IB),QUAD(J+4,IB),QUAD(J+6,IB)
    1)495,495,330
    330 IF(YMIN(J)-AMAX1(QUAD(J,IB),QUAD(J+2,IB),QUAD(J+4,IB),QUAD(J+6,IB)
    1)335,495,495
    335 CONTINUE
    ***** SET UP LINEAR FORMS FOR EDGES OF QUADRANGLE *****
    DO 345 L=1,4
    K=2*L
    KL=MOD(K,8)+2
    QUA(1,L)=QUAD(K,IB)-QUAD(KL,IB)
    QUA(2,L)=QUAD(KL-1,IB)-QUAD(K-1,IB)
    QUA(3,L)=QUAD(K-1,IB)*QUAD(KL,IB)-QUAD(K,IB)*QUAD(KL-1,IB)
    ***** NORMALIZE LINE EQUATION COEFFICIENTS *****
    T1=SQRT(QUA(1,L)**2+QUA(2,L)**2)
    IF(T1)495,495,340
    340 DO 345 J=1,3
    345 QUA(J,L)=QUA(J,L)/T1
    ***** EVALUATE LINEAR FORMS AND SIGNATURES FOR QUADRANGLE *****
    T3=3.0
    DO 365 K=1,4
    T2=3.0
    J=K*2
    DO 355 L=1,4
    T1=QUAD(J-1,IQ)*QUA(1,L)+QUAD(J,IQ)*QUA(2,L)+QUA(3,L)
    IF(T1)350,355,355
    T2=T2-1.0
    350 T2=T2-1.0
    355 FL(L,K)=T1
    IF(T2)360,365,365
    360 T3=T3-1.0
    365 CONTINUE
    ***** CHECK FOR 4 POINTS INSIDE QUADRANGLE *****
    IF(T3)370,375,375
    370 ITYPE=-1
    GO TO 415
    ***** CHECK FOR 1 TO 3 POINTS INSIDE QUADRANGLE *****
    375 IF(T3-3.0)380,385,385
    380 ITYPE=0
    GO TO 415
C
    ***** DETERMINE WHICH EDGES ARE CROSSED BY THE 4 LINE SEGMENTS ***
    DO 405 L=1,4
    LI=MOD(L,4)+1
    ***** LINE SEGMENT L FROM POINT Y1 TO POINT Y2 *****
    Y1(L)=QUAD(L*2-1,IQ)
    Y1(2)=QUAD(L*2,IQ)
    Y2(L)=QUAD(L1*2-1,IQ)
    Y2(2)=QUAD(L1*2,IQ)
    DO 405 K=1,4
    T1=FL(K,L)
    T2=FL(K,L1)
    T3=T1-T2
C
    ***** T1 AND T2 MUST HAVE OPPOSITE SIGNS FOR INTERSECTION TO OCCUR
    IF(T1*T2)390,390,405
C
    ***** COMPONENT OF SEGMENT L PERPENDICULAR TO EDGE K OF IB IS T3
    390 IF(ABS(T3)-1.E-5)405,405,395
C
    ***** CALCULATE COORDINATES OF INTERSECTION *****
    395 T4=(T1*Y2(1)-T2*Y1(1))/T3
    T5=(T1*Y2(2)-T2*Y1(2))/T3
    KO=2*K

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C      K1=2*(MOD(K,4)+1)
C      ***** IS INTERSECTION WITHIN QUADRANGLE IQ *****
C      T6=(T4-QUAD(K0-1,IB))*(QUAD(K1-1,IB)-T4)+(T5-QUAD(K0,IB))*
1 QUAD(K1,IB)-T5)
C      IF(ABS(T6)-1.E-4)410,410,400
400 IF(T6)405,410,410
405 CONTINUE
C      GO TO 495
410 ITYPE=1
C      ***** CHECK OVER/UNDER AMBIGUITY *****
415 IF((NA1-NB1)*(NA2-NB2)*(NA2-NB1))425,420,425
C      ***** BONDS SHARE AN ATOM *****
420 IF(NA1+NA2-NB1-NB2)465,495,495
425 CALL DIFV(ATOMS(1,NB2),ATOMS(1,NB1),V2)
CALL DIFV(ATOMS(1,NB1),ATOMS(1,NA1),V4)
CALL UNITY(V2,V2,1)
CALL UNITY(V4,V4,1)
CALL NORM(V1,V2,V3,1)
IF(VV(V3,V3)-TOL)430,430,435
C      ***** PARALLEL BONDS, RECALCULATE V3 *****
430 CALL NORM(V1,V4,V5,1)
CALL NORM(V5,V1,V3,1)
C      ***** CHECK FOR COLLINEAR BONDS *****
435 IF(VV(V3,V3)-TOL)465,465,450
440 DO 445 J=1,3
445 V3(J)=-V3(J)
C      ***** V3 IS NORMAL TO BONDS IQ AND IB GOING FROM IQ TOWARD IB ***
450 IF(VIEW)455,455,460
455 IF(V3(3))495,495,465
460 IF(VV(VUE,V3))465,495,495
C      ***** OVERLAPPING BOND FOUND *****
465 ICQ=ICQ+1
IF(ITYPE)470,475,475
C      ***** HIDDEN BOND *****
470 ICQ=-1
RETURN
C      ***** STORE INTERFERING QUADRANGLE *****
475 IF(NCOVER=30)485,480,480
480 NG=18
TIDD=TIDD2
CALL ERENT(TIDD,800)
RETURN
485 NCOVER=NCOVER+1
DO 490 K=1,4
DO 490 J=1,3
490 COVER(J,K,NCOVER)=QUA(J,K)
495 CONTINUE
500 RETURN
END
C      SUBROUTINE LAPAB(IQ,IA,ICQ,ITY)
C      ***** SUBROUTINE CHECKS FOR OVERLAP BETWEEN ATOMS AND BONDS *****
C      ***** CALLED BY SUBROUTINES LAP700 AND LAP800 *****
C      DIMENSION BF(4),CON(3,3),QF(5),QUA(3,4)
COMMON/OLAP/CONIC(7,500),COVER(6,20),KC(20),KQ(30),NCONIC,NCOVER,
1 NCOVER,NOUAD,OVVRGN,COVER(3,4,30),QUAD(9,600),SEGM(50,2)
REAL*8 TIDD
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BE(3,3),BRDR,CD(8,20)
2 CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 IN_ITILT,KD(5,20),LATM,NATOM,PCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCALI
5 ,SCAL2,SC3,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),V6K(3,3)
7 ,XLNG(3),XO(3),XT(3)
TIDD=QUAD(9,IQ)
NA1=TID/1000
NA2=AMOD(TIDD,1000)
C      ***** ITY.GT.0, CHECK FOR ATOMS OVER A BOND *****
C      ***** ITY.LT.0, CHECK FOR BONDS OVER AN ATOM *****
ICQ=0
IF(ITY)210,200,205
200 RETURN
205 CALL LAPCON(CONIC(1,IA),CON,V1,0.0)
IF(ITY-2)220,240,240
210 IF(ITY+2)220,220,215
215 CALL LAPCON(CONIC(1,IA),CON,V1,OVVRGN)
C      ***** SET UP LINEAR FORMS FOR EDGES OF QUADRANGLE *****
220 DO 235 L=1,4
K=2*L
K1=MOD(K,8)+2
QUA(1,L)=QUAD(K,IQ)-QUAD(K1,IQ)
QUA(2,L)=QUAD(K-1,IQ)-QUAD(K-1,IQ)
QUA(3,L)=QUAD(K-1,IQ)*QUAD(K1,IQ)-QUAD(K,IQ)*QUAD(K1-1,IQ)
T1=SORT(QUA(1,L)**2+QUA(2,L)**2)
IF(T1)225,225,230
225 ITY=0
ICQ=0
GO TO 430
C      ***** TRANSFORM COEFFICIENTS FOR EDGES TO NORMAL FORM *****
230 DO 235 J=1,3
235 QUA(J,L)=QUA(J,L)/T1
C      ***** EVALUATE 4 QUADRATIC AND 4 BILINEAR FORMS *****
240 V2(3)=1.0
V3(3)=1.0
T2=3.0
DO 265 L=1,4
L1=(MOD(L,4)+1)*2
V2(1)=QUAD(2*L-1,IQ)
V2(2)=QUAD(2*L,IQ)
V3(1)=QUAD(L1-1,IQ)
V3(2)=QUAD(L1,IQ)
QF(L)=0.0
BE(L)=0.0
DO 250 K=1,3
T1=CON(3,K)
DO 245 J=1,2
245 T1=TI+V2(J)*CON(J,K)
250 BE(L)=BE(L)+TI*V2(K)
255 IF(QF(L))260,255,265
260 T2=T2-0.8
GO TO 265
265 T2=T2-1.0
CONTINUE
265 QF(5)=QF(1)
C      ***** CHECK FOR 4 POINTS OF QUADRANGLE INSIDE OR ON ELLIPSE *****
IF(T2)270,275,275
270 ITYPE=-1

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GO TO 330
C ***** CHECK FOR 1 TO 3 POINTS OF QUADRANGLE INSIDE THE ELLIPSE ***
275 IF(T2-2.2)280,285,285
280 ITYPE=0
C ***** CHECK FOR QUADRANGLE-ELLIPSE INTERSECTION *****
285 DO 305 K=1,4
C ***** EVALUATE DISCRIMINANT *****
TI=BF(K)**2-QF(K)*QF(K+1)
IF(TI)305,305,290
290 TI=SQRT(TI)
C ***** IS INTERSECTION WITHIN BOUNDS OF QUADRANGLE *****
T3=QF(K)-BF(K)
T4=T3+QF(K+1)-BF(K)
IF(ABS(T4)-1.E-5)305,305,295
295 T5=(T3-TI)/T4
IF(T5)305,280,300
300 IF(1.0-T5)305,305,280
305 CONTINUE
C ***** NO VALID INTERSECTION FOUND *****
C ***** CHECK FOR CENTER OF ELLIPSE WITHIN THE QUADRANGLE *****
T3=3.0
DO 320 K=1,4
TI=QUA(3,K)
DO 310 J=1,2
310 TI=TI+V1(J)*QUA(J,K)
315 T3=T3-1.0
320 CONTINUE
325 IF(T3)325,370,370
ITYPE=1
C ***** CHECK OVER/UNDER AMBIGUITY *****
330 IF(NA2-IA)375,375,335
335 IF(IA-NA1)375,375,340
340 CALL DIFV(ATOMS(1,NA2),ATOMS(1,NA1),V2)
CALL UNIFY(V2,V2,1)
CALL UNIFY(V3,V3,1)
CALL NORM(V2,V3,V4,1)
IF(VV(V4,V4)-1.E-5)345,345,350
345 IF(ITV)370,370,385
C ***** CENTER OF ATOM IQ IS ON THE BOND LINE *****
C ***** CENTER OF ATOM IQ IS NOT ON THE BOND LINE *****
350 CALL NORM(V4,V2,V5,1)
TI=-V5(3)
IF(VIEW)365,365,355
355 TI=V5(3)*(ATOMS(3,IA)-VIEW)
DO 360 J=1,2
360 TI=TI+V5(J)*ATOMS(J,IA)
365 IF(TI*FLOAT(ITV))375,375,370
C ***** NO INTERFERENCE FOUND *****
370 ICQ=0
GO TO 430
C ***** ITYPE=1 ENCLOSED ELLIPSE / ITYPE=-1 ENCLOSED QUADRANGLE *****
375 IF(ITYP*ITY)380,385,385
C ***** HIDDEN ATOM OR HIDDEN BOND *****
380 ICQ=-1
GO TO 430
385 ICQ=1
IF(ITV)410,390,390
C ***** STORE INTERFERING ELLIPSE *****

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390 IF(NCOVER-20)400,395,395
395 NG=17
CALL ERENT(ATOMID(IA),800)
NCOVER=NCOVER-1
400 NCOVER=NCOVER+1
IJ=1
DO 405 I=1,3
DO 405 J=1,3
COVER(IJ,NCOVER)=CON(I,J)
405 IJ=IJ+1
KC(NCOVER)=IA
GO TO 430
C ***** STORE INTERFERING QUADRANGLE *****
410 IF(NCOVER-30)420,415,415
415 NG=18
TIDD=TID
CALL ERENT(TIDD,700)
NCOVER=NCOVER-1
420 NCOVER=NCOVER+1
DO 425 K=1,4
DO 425 J=1,3
425 QOVER(J,K,NCOVER)=QUA(J,K)
KQ(NCOVER)=IQ
430 RETURN
END
SUBROUTINE LAPCON(CON1,CON,Y,OVMR)
C ***** TRANSFORM CONIC TO PLOTTER HOMOGENEOUS COORDINATE SYSTEM ***
C ***** CALLED BY SUBROUTINES LAP700 AND LAPAB *****
DIMENSION CON1(7),CON(3,3),Y(3)
Y(1)=(CON1(1)+CON1(2))*0.5
Y(2)=(CON1(3)+CON1(4))*0.5
Y(3)=1.0
CON(1,1)=CON1(5)
CON(1,2)=CON1(6)
CON(2,1)=CON1(6)
CON(2,2)=CON1(7)
TI=(CON1(2)-CON1(1)+CON1(4)-CON1(3))*0.25
CON(3,3)=-((TI-OVMR)/TI)**2
DO 205 K=1,2
CON(K,3)=0.0
DO 200 J=1,2
200 CON(K,3)=CON(K,3)-Y(J)*CON(J,K)
205 CON(3,K)=CON(K,3)
CON(3,3)=CON(3,3)-CON(3,K)*Y(K)
RETURN
END
SUBROUTINE LAPDRW(Y,NPEN,NCQ)
C ***** SUBROUTINE ELIMINATES HIDDEN LINES AND DRAWS VISIBLE LINES *
DIMENSION CB(20),CQ(50,2),QL(4,30,2),SEG(2),Y(3),YN(3),YO(3),Z(3)
COMMON/OLAP/CONIC(7,500),COVER(6,20),KC(20),KQ(30),NCONIC,NCOVER,
1 NCOVER,NQUAD,OVMRGN,QOVER(3,4,30),QUAD(9,600),SEGM(50,2)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAMRK(3,3),AID(3,3)
1 AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 IN,TITL,T,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ORGN(3),PAC(3,5),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCALL
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),V6(3),V6(3,3)
7 ,XLING(3),XO(3),XT(3)

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      NCO=NCOVER+NOOVER
      IF(NCO)200,200,205
    200 RETURN
      ***** CHECK ALL OVERLAPPING ATOMS AND BONDS *****
    205 NPM3=NPN-3
      IF(NPM3) 210,230,230
      ***** SAVE INFORMATION FROM LAST POINT IF PEN IS DOWN *****
    210 YO(1)=YN(1)
      YO(2)=YN(2)
      YO(3)=1.0
      NPO=NPN
      DO 215 K=1,NCO
    215 CQ(K,1)=CQ(K,2)
      IF(NCOVER)230,230,220
    220 DO 225 K=1,NCOVER
      DO 225 J=1,4
    225 QL(J,K,1)=QL(J,K,2)
      ***** EVALUATE CONIC QUADRATIC FORMS AT NEW POINT YN *****
    230 YN(1)=Y(1)
      YN(2)=Y(2)
      YN(3)=1.0
      NPN=NPN
      IF(NCOVER)250,250,235
    235 DO 245 K=1,NCOVER
      Z(1)=YN(1)*COVER(1,K)+YN(2)*COVER(2,K)+COVER(3,K)
      Z(2)=YN(1)*COVER(2,K)+YN(2)*COVER(4,K)+COVER(5,K)
      Z(3)=YN(1)*COVER(3,K)+YN(2)*COVER(5,K)+COVER(6,K)
      CQ(K,2)=Z(1)*YN(1)+Z(2)*YN(2)+Z(3)
      ***** EVALUATE CONIC BILINEAR FORM IF PEN IS DOWN *****
    240 CB(K)= Z(1)*YO(1)+Z(2)*YO(2)+Z(3)
    245 CONTINUE
      ***** EVALUATE LINEAR FORMS AND SIGNATURE FOR QUADRANGLE *****
    250 IF(NCOVER)275,275,255
    255 KCQ=NCOVER
      DO 270 K=1,NCOVER
      T2=3.0
      DO 265 J=1,4
      T1=YN(1)*COVER(1,J,K)+YN(2)*COVER(2,J,K)+COVER(3,J,K)
      IF(T1)260,265,265
    260 T2=T2-I.0
    265 QL(J,K,2)=T1
      KCQ=KCQ+1
      ***** T2=-1 INSIDE, =0 ACROSS ANY EDGE, =1 ACROSS ANY VERTEX *****
    270 CQ(KCQ,2)=T2
      ***** IF PEN IS UP, OMIT ALL SUBSEQUENT CHECKING *****
    275 IF(NPM3)285,280,280
    280 NPN=3
      CALL SCRIBE(YN,NPN)
      RETURN
      ***** CHECK FOR HIDDEN SEGMENT *****
    285 DO 295 K=1,NCO
      IF(CQ(K,1))290,295,295
    290 IF(CQ(K,2))280,295,295
    295 CONTINUE
      ***** FIND ENTRY AND EXIT POINTS ON EACH CONIC *****
    300 NINT=0
      IF(NCOVER)330,330,300
    300 DO 325 K=1,NCOVER
      ***** EVALUATE DISCRIMINANT *****
      T1=CB(K)**2-CQ(K,1)*CQ(K,2)
      IF(T1)325,325,305
    305 T1=SQRT(T1)
      ***** SOLVE QUADRATIC EQUATION *****
      T2=CQ(K,1)-CB(K)
      T3=T2+CQ(K,2)-CB(K)
      IF(ABS(T3)-1.E-5)325,325,310
    310 T4=(T2-T1)/T3
      T5=(T2+T1)/T3
      ***** VALID INTERSECTION IF T4.LT.1 AND T5.GT.0 *****
      IF(T4-1.0)315,325,325
    315 IF(T5)325,325,320
      ***** SAVE VALID CONIC INTERSECTIONS *****
    320 NINT=NINT+1
      SEGM(NINT,1)=T4
      SEGM(NINT,2)=T5
    325 CONTINUE
    330 IF(NCOVER)425,425,335
      ***** FIND ENTRY AND EXIT POINTS FOR EACH QUADRANGLE *****
    335 DO 420 K=1,NCOVER
      I12=0
      KCQ=NCOVER+K
      ***** CHECK FOR SINGLE INSIDE POINT *****
      SEG(1)=CQ(KCQ,1)
      IF(SEG(1))345,340,340
    340 SEG(1)=1.0-CQ(KCQ,2)
      IF(SEG(1)-1.0)350,350,345
      ***** INSIDE POINT FOUND, ONLY ONE INTERSECTION POSSIBLE *****
    345 I12=1
      ***** FIND WHICH EDGES ARE CROSSED BY THE SEGMENT *****
    350 DO 410 J=1,4
      T1=QL(J,K,1)
      T2=QL(J,K,2)
      T3=T1-T2
      IF(T1*T2)355,355,410
      ***** CHECK FOR SEGMENT ON AN EDGE *****
    355 IF(ABS(T3)-1.E-5)420,420,360
      ***** CALCULATE COORDINATES OF INTERSECTION *****
    360 T4=(T1*YN(1)-T2*YO(1))/T3
      T5=(T1*YN(2)-T2*YO(2))/T3
      J1=2*(MOD(J,4)+1)
      IQ=KQ(K)
      ***** IS INTERSECTION WITHIN LIMITS OF QUADRANGLE *****
      1 (QUAD(J1,IQ)-T5)
      IF(ABS(T6)-1.E-4)370,370,365
    365 IF(T6)410,370,370
      ***** CALCULATE FRACTION PARAMETER AND STORE IT *****
    370 T1=T1/T3
      IF(I12-1)375,380,395
      ***** STORE FIRST INTERSECTION *****
    375 I12=1
      GO TO 390
      ***** STORE SECOND INTERSECTION ****
    380 I12=2
      IF(T1-SEG(1))385,405,405
    385 SEG(2)=SEG(1)
    390 SEG(1)=T1
      GO TO 410
      ***** MORE THAN TWO INTERSECTIONS (I.E., QUADRANGLE DIAGONAL) *****
    395 IF(T1-SEG(1))390,410,400
    400 IF(T1-SEG(2))410,410,405

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405 SEG(2)=T1
410 CONTINUE
IF(I12-1)420,420,415
***** STORE FRACTION PARAMETERS *****
C 415 NINT=NINT+1
SEG(NINT,1)=SEG(1)
SEG(NINT,2)=SEG(2)
420 CONTINUE
***** END OF ENTRY-AND-EXIT-POINT CALCULATIONS *****
C 425 IF(NINT-1)430,490,435
***** NO INTERFERENCE FOUND, DRAW ENTIRE SEGMENT *****
C 430 CALL SCRIBE(YN,2)
RETURN
C ***** SORT SEGMENT INTERSECTION LIST *****
C ***** SORTING PROCEDURE BY SHELL,D.L. COMM. ACM 2,30-32 (1959) ***
435 M=NINT
440 M=M/2
445 K=NINT-M
J=1
450 I=J
455 IM=I+M
IF(SEGM(I,1))460,470,470
460 IF(SEGM(IM,1))465,465,485
465 IF(SEGM(I,2)-SEGM(IM,2))485,485,475
470 IF(SEGM(I,1)-SEGM(IM,1))485,485,475
475 DO 480 L=1,2
TL=SEGM(I,L)
480 SEGM(I,L)=SEGM(IM,L)
SEGM(IM,L)=TL
I=I-M
485 J=J+1
IF(I)485,485,455
IF(J-K)450,450,440
***** FIND STARTING POINT P0 AND END POINT P1 *****
C 490 P0=0.0
K=0
495 K=K+1
IF(K=NINT)500,500,515
500 P1=SEGM(K,1)
IF(P1)510,505,505
505 IF(P1-P0)510,510,520
510 P0=AMAX1(P0,SEGM(K,2))
IF(P0-1.0)495,530,525
515 P1=1.0
***** DRAW SEGMENT FROM P0 TO P1 *****
C 520 IF(P0)535,535,530
525 P0=1.0
530 Z(1)=YO(1)*(1.-P0)+YN(1)*P0
Z(2)=YO(2)*(1.-P0)+YN(2)*P0
NPN=3
CALL SCRIBE(Z,NPN)
IF(P0-1.0)535,540,540
535 Z(1)=YO(1)*(1.-P1)+YN(1)*P1
Z(2)=YO(2)*(1.-P1)+YN(2)*P1
NPN=2
CALL SCRIBE(Z,NPN)
IF(P1-1.0)510,540,540
540 RETURN
END
character*(*) function maksym(k,gp)
C *** returns character string representation of symmetry operator
dimension gp(3,4,192)
character*1 xyz(3)
character*5 fract(23)
character*12 part(3)
data fract/'1/24','1/24','1/24','1/24','1/24','13/24','7/12','5/8','2/3',
* '1/724','3/4','19/24','5/6','7/8','11/12','23/24'/
* data xyz/'x','y','z'/
do 200 i=1,3
part(i) = ' '
iff = 0
do 300 j=1,3
if (ifix(gp(i,j,k)).ne. 0) then
if (ifix(gp(i,j,k)) .eq. -1)
part(i) = part(i)(1:iend(part(i))) // '-' // xyz(j)
if (ifix(gp(i,j,k)) .eq. 1 .and. iff .eq. 0)
part(i) = part(i)(1:iend(part(i))) // ' ' // xyz(j)
if (ifix(gp(i,j,k)) .eq. 1 .and. iff .eq. 1)
part(i) = part(i)(1:iend(part(i))) // '+' // xyz(j)
iff = 1
end if
continue
300 gpval = gp(i,4,k)
if(gpval.gt.01 .or. gpval.lt.-.01) then
if (gpval.lt.0.) then
part(i) = part(i)(1:iend(part(i))) // '-'
else
part(i) = part(i)(1:iend(part(i))) // '+'
end if
gpval=abs(gpval)
tfour=1./24.
do 301 mm=1,23
tf=float(mm)*tfour
if (gpval.gt.(tf-.01) .and. gpval.lt.(tf+.01)) iw=mm
301 continue
part(i) = part(i)(1:iend(part(i))) // fract(iw)
end if
200 maksym = part(1) // part(2) // part(3)
return
end
SUBROUTINE MM(X,Y,Z)
MULTIPLY TWO MATRICES
Z(3,3)=X(3,3)*Y(3,3)
C
C DIMENSION X(3,3),Y(3,3),Z(3,3)
X11=X(1,1)
X12=X(1,2)
X13=X(1,3)
X21=X(2,1)
X22=X(2,2)
X23=X(2,3)
X31=X(3,1)
X32=X(3,2)
X33=X(3,3)
Y11=Y(1,1)
Y12=Y(1,2)
Y13=Y(1,3)
Y21=Y(2,1)
Y22=Y(2,2)
Y23=Y(2,3)

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Y31=Y(3,1)
Y32=Y(3,2)
Y33=Y(3,3)
Z(1,1)=X11*Y11+X12*Y21+X13*Y31
Z(2,1)=X21*Y11+X22*Y21+X23*Y31
Z(3,1)=X31*Y11+X32*Y21+X33*Y31
Z(1,2)=X11*Y12+X12*Y22+X13*Y32
Z(2,2)=X21*Y12+X22*Y22+X23*Y32
Z(3,2)=X31*Y12+X32*Y22+X33*Y32
Z(1,3)=X11*Y13+X12*Y23+X13*Y33
Z(2,3)=X21*Y13+X22*Y23+X23*Y33
Z(3,3)=X31*Y13+X32*Y23+X33*Y33
RETURN
END
SUBROUTINE MV(X,Y,Z)
MATRIX * VECTOR
Z(3)=X(3,3)*Y(3)
DIMENSION X(3,3),Y(3),Z(3)
Y1=Y(1)
Y2=Y(2)
Y3=Y(3)
Z(1)=X(1,1)*Y1+X(1,2)*Y2+X(1,3)*Y3
Z(2)=X(2,1)*Y1+X(2,2)*Y2+X(2,3)*Y3
Z(3)=X(3,1)*Y1+X(3,2)*Y2+X(3,3)*Y3
RETURN
END
SUBROUTINE NORM(X,Y,Z,ITYPE)
***** VECTOR PRODUCT Z=X*Y *****
***** ITYPE .GT.0 FOR CARTESIAN, .LE.0 FOR TRICLINIC *****
DIMENSION X(3),Y(3),Z(3),Z1(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWEK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,COMT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WERKV(3,3)
7 ,XING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
DI00=100.
DI100=100000.
IT=IABS(ITYPE)-1
KS=DMOD(DCODE,DI00)
IF(NSYM-KS)105,115,115
105 NG=4
GO TO 300
115 IT=DCODE/DI100
IF(NATOM-IT)125,130,130
125 NG=5
GO TO 300
130 IF(IT)125,125,135
***** CRYSTALLOGRAPHIC SYMMETRY ROTATION *****
135 CALL TMM(PA(1,1,IT),FS(1,1,KS),PAT)
IF(IT-1)160,145,155
***** TRANSFORM TO CARTESIAN SYSTEMS *****
145 CALL TMM(PAT,AAWEK,PAC)
GO TO 175
155 CALL TMM(PAT,AAREV,PAC)
GO TO 175
160 IF(ITYPE)162,155,170
***** TRANSFORM TO TRICLINIC SYSTEM *****
162 DO 165 J=1,9
165 PAC(J,1)=PAT(J,1)

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XD=DIST
10 IF(XD.LT.1.0) GO TO 20
IF(NC.GE.9) GO TO 30
NC=NC+1
XD=XD/10.0
GO TO 10
C-----SET UP FORMAT STATEMENT
20 WRITE (IFMT,25) NC,ND
25 FORMAT('F',11,' ',11,' ',11,' ')
C-----ENCODE DISTANCE AND PUT IT OUT
WRITE (ITXT,IFMT) DIST
CALL SIMBOL(W,W2,HGT,ITEX,THI,NC)
30 RETURN
END
SUBROUTINE PAXES(DCODE,ITYPE)
***** ITYPE .LT.0 FOR COVARIANCE MATRIX IN Q *****
***** ITYPE .GT.0 FOR ELLIPSOID QUADRATIC FORM IN Q *****
C ***** XABSF(ITYPE)=1 BASED ON TRICLINIC COORDINATE SYSTEM *****
C ***** =2 OR 3 FOR WORKING OR REFERENCE CARTESIAN SYSTEMS *****
C ***** CONTRAVARIANT EIGENVECTORS FOR Q IN COLUMNS OF PAC *****
C ***** CHECK ATOM CODE *****
DIMENSION X(3)
REAL*8 DCODE,DI00,DI100K
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWEK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,COMT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WERKV(3,3)
7 ,XING(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
DI00=100.
DI100K=100000.
IT=IABS(ITYPE)-1
KS=DMOD(DCODE,DI00)
IF(NSYM-KS)105,115,115
105 NG=4
GO TO 300
115 IT=DCODE/DI100K
IF(NATOM-IT)125,130,130
125 NG=5
GO TO 300
130 IF(IT)125,125,135
***** CRYSTALLOGRAPHIC SYMMETRY ROTATION *****
135 CALL TMM(PA(1,1,IT),FS(1,1,KS),PAT)
IF(IT-1)160,145,155
***** TRANSFORM TO CARTESIAN SYSTEMS *****
145 CALL TMM(PAT,AAWEK,PAC)
GO TO 175
155 CALL TMM(PAT,AAREV,PAC)
GO TO 175
160 IF(ITYPE)162,155,170
***** TRANSFORM TO TRICLINIC SYSTEM *****
162 DO 165 J=1,9
165 PAC(J,1)=PAT(J,1)

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SUBROUTINE MV(X,Y,Z)
MATRIX * VECTOR
Z(3)=X(3,3)*Y(3)
DIMENSION X(3,3),Y(3),Z(3)
Y1=Y(1)
Y2=Y(2)
Y3=Y(3)
Z(1)=X(1,1)*Y1+X(1,2)*Y2+X(1,3)*Y3
Z(2)=X(2,1)*Y1+X(2,2)*Y2+X(2,3)*Y3
Z(3)=X(3,1)*Y1+X(3,2)*Y2+X(3,3)*Y3
RETURN
END
SUBROUTINE NORM(X,Y,Z,ITYPE)
***** VECTOR PRODUCT Z=X*Y *****
***** ITYPE .GT.0 FOR CARTESIAN, .LE.0 FOR TRICLINIC *****
DIMENSION X(3),Y(3),Z(3),Z1(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWEK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,COMT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SC1,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WERKV(3,3)
7 ,XING(3),XO(3),XT(3)
DO 125 I=1,3
II=MOD(I+3,3)+1
I2=MOD(I+1,3)+1
T1=X(II)*Y(I2)-X(I2)*Y(II)
IF(ITYPE)115,115,105
105 Z(I)=T1
GO TO 125
115 Z(I)=T1
125 CONTINUE
135 CALL MV(BB,Z1,Z)
300 RETURN
END
SUBROUTINE NUMBR(W,W2,HGT,DIST,THI,ND)
CONVERT BOND DISTANCE FOR PLOTTING IN ORTEP
DIMENSION W(3)
CHARACTER*8 IFMT,ITXT
CHARACTER*1 ITEX(8)
EQUIVALENCE (ITEX(1),ITXT)
***** COMPUTE NUMBER OF CHARACTERS FOR OUTPUT
NC=ND+1

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C ***** WAS INPUT FOR REAL OR RECIPROCAL CELL *****
135 IF(A(1)-1.)135,150,150
DO 140 J=1,9
TI=AA(J,1)
AA(J,1)=BB(J,1)
BB(J,1)=TI
TI=A(J)
A(J)=B(J)
B(J)=TI
***** WRITE OUT CELL PARAMETERS *****
140 FORMAT(1H010X,22HDIRECT CELL PARAMETERS/1H 15X,1H414X,1HB14X,1HC14
1X,5HALPHA10X,4HBETA11X,5HGAMMA)
145 FORMAT(1H 10X,F9.5,2F15.6,3F15.3/1H 48X,6HCOSINE,F12.8,2F15.8)
147 FORMAT(1H010X,26HRECIPROCAL CELL PARAMETERS/1H 15X,2HA*13X,2HB*13X
1,2HC*13X,6HALPHA*9X,5HBETA*10X,6HGAMMA*)
150 IF (NOUT.GE.0)
&WRITE (NOUT,143)
IF (NOUT.GE.0)
&WRITE (NOUT,145) (A(I),I=1,3), (A(I),I=7,9), (A(I),I=4,6)
IF (NOUT.GE.0)
&WRITE (NOUT,147)
IF (NOUT.GE.0)
&WRITE (NOUT,145) (B(I),I=1,3), (B(I),I=7,9), (B(I),I=4,6)
***** STORE STANDARD VECTORS *****
CALL AXES(AID,AID(1,2),REFV,0)
CALL MM(AA,REFV,AAREV)
DO 160 I=1,3
DO 160 J=1,3
AAWK(J,I)=AAREV(J,I)
Q(J,I)=REFV(I,J)
160 WRKV(J,I)=REFV(J,I)
***** READ AND WRITE SYMMETRY TRANSFORMATIONS *****
C
171 FORMAT(1H0 10X,24HSYMMETRY TRANSFORMATIONS/1H 14X,3HNO.12X,13HTRANS
FORMED X18X,13HTRANSFORMED Y18X,13HTRANSFORMED Z)
173 FORMAT(1I,1F4.10,3F3.0,2(F15.10,3F3.0))
175 FORMAT(1H 13X,I2,3(F13.6,F4.0,2H X,F4.0,2H Y,F4.0,2H Z))
176 FORMAT(1H 13X,I2,5X,a)
177 FORMAT(1H1.10X,18A4)
IF (NOUT.GE.0)
&WRITE (NOUT,171)
LINES=14
DO 190 I=1,96
LINES=MOD(LINES+1,56)
READ (IN,107)CARD
write (NED,107) card
if (iflag.eq.0)READ(card,173)IS,(TS(J,I),(FS(K,J,I),K=1,3),J=1,3)
read (card,1771) is
format(i1)
1771 ipart=1
do 1772 jk=1,3
do 1772 kl=1,24
sympart(jk)(kl:kl)=' '
jk=2
if (card(jk:jk).eq.' ') go to 1776
1772
1773 lm=1
do 1774 kl=jk,80
if(card(kl:kl).eq.' ' .or. card(kl:kl).eq.' ') then
&WRITE (NOUT,207)
jk=kl
go to 1775
end if
sympart(ipart)(lm:lm)=card(kl:kl)
lm=lm+1
continue
ipart=ipart+1
jk=jk+1
if (jk.lt.80) go to 1773
do 1777 isymp=1,3
call tepsym(sympart(isymp),i,isymp)
continue
1777
do 178 j=1,3
fsym(j,4)=ts(j,i)
do 178 k=1,3
fsym(j,k)=fs(k,j,i)
178 IF(LINES)185,180,185
180 IF (NOUT.GE.0)
&WRITE (NOUT,177) (TITLE(J),J=1,18)
IF (NOUT.GE.0)
&WRITE (NOUT,171)
C *** ORTEP II symmetry output
C 185 IF (NOUT.GE.0)
C &WRITE (NOUT,175) I,(TS(J,I),(FS(K,J,I),K=1,3),J=1,3)
C 185 if (nout.ge.0) WRITE (NOUT,176) I,makeSym(1,fsym)
C ***** NON-CRYSTALLOGRAPHIC HELIX-SYMMETRY INPUT *****
186 IF(FS(3,3,I)-5.)188,186,186
TI=FS(1,3,I)/FS(3,3,I)
TS(3,I)=TS(3,I)+TI
TI=AMOD(TI*FS(2,3,I),1.)*6.28318531
TI=COS(TI)
TI=SIN(TI)
DO 187 J=1,9
VT(J,1)=AID(J,1)
VT(1,1)=T2
VT(2,2)=T2
VT(2,1)=-TI
VT(1,2)=TI
CALL MM(VT,O,PAC)
CALL MM(AAREV,PAC,FS(1,1,I))
188 IF(IS)195,190,195
190 CONTINUE
NG=1
CALL EREPT(0.D0,0)
I=96
195 NSYM=I
196 ISW=IS
NATOM=0
C ***** POSITIONAL AND THERMAL PARAMETERS *****
207 FORMAT(11H0 NO. ATOM 8X,1HX10X,1HY10X,1HZ13X,3HB118X,3HB228X,3HB33
18X,3HB128X,3HB138X,10HB23 TYPE)
209 FORMAT(1H 13.1X,A6,3F11.6,5X,6F11.6,F5.0)
210 FORMAT(1H 13.1X,A6,3F11.6,5X,2F11.6,4F11.0,F5.0)
211 FORMAT(A6,3X,6F9.0)
213 FORMAT(1I,F8.0,5F9.0,7X,F2.0)
LINES=LINES+2
IF(LINES-56)220,215,215
215 LINES=1
GO TO 225
220 IF (NOUT.GE.0)
&WRITE (NOUT,207)
if (isw.eq.2) then

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      iu=18
      call gtafill(iu)
    end if
  C ***** MAXIMUM NUMBER OF ATOMS EQUALS MAXATM *****
  DO 225 NATOM=NATOM+1
  DO 245 I=NATOM,MAXATM
  LINES=MOD(LINES+1,56)
  IF(ISW.EQ.1) GO TO 226
  ***** CALL SPECIAL PURPOSE READIN ROUTINE *****
  CALL readin(iu,CHEM(I),IDENT(1,I),IDENT(2,I),P(1,I),P(2,I),P(3,I),
  1,IT1,IS,PA(1,I),PA(2,I),PA(3,I),PA(1,2,I),PA(2,2,I),
  2,PA(3,2,I),PA(1,3,I))
  IT1=IT1+1
  K=IT1+1
  GO TO 229
226 continue
  READ (IN,107)card
  write (NED,107) card
  READ (card,211)CHEM(I),V1(1),V1(2),P(J,I),J=1,3),T1
  IDENT(1,I)=V1(1)
  IDENT(2,I)=V1(2)
  K=1.*T1
  IF(FLOAT(K-1)-T1)227,228,227
227 K=1
228 continue
  READ (IN,107)card
  write (NED,107) card
  READ (card,213)IS,(PA(J,I),J=1,7)
229 IF(LINES)230,230,232
230 IF (NOUT.GE.0)
  &WRITE (NOUT,177)(TITLE(J),J=1,18)
  &WRITE (NOUT,207)
232 IF(PA(3,1,I)-10000.)235,234,234
234 IF (NOUT.GE.0)
  &WRITE (NOUT,210) I,CHEM(I),P(J,I),J=1,3),P(J,I),J=1,7)
  GO TO 238
235 IF (NOUT.GE.0)
  &WRITE (NOUT,209) I,CHEM(I),P(J,I),J=1,3),P(J,I),J=1,7)
238 GO TO (244,239,241,242,244),K
  ***** TYPE 1 POSITIONAL PARAMETERS (ANGSTROMS) *****
  DO 240 J=1,3
  P(J,I)=P(J,I)/A(J)
  GO TO 244
  ***** TYPE 2 POSITIONAL PARAMETERS, STANDARD CARTESIAN *****
  V1(1)=P(1,I)
  V1(2)=P(2,I)
  GO TO 243
  ***** TYPE 3 POSITIONAL PARAMETERS *****
  ***** CYLINDRICAL COORDINATES REFERRED TO STANDARD CARTESIAN *****
242 T2=P(2,I)*.01745329252
  V1(1)=V1(1)+P(1,I)*COS(T2)
  V1(2)=V1(2)+P(1,I)*SIN(T2)
243 V1(3)=P(3,I)
  CALL VM(V1,Q,P(1,I))
244 IF(IS)246,245,246
245 CONTINUE
  if (isw.eq.2) close(iu)
  NG=2
  CALL ERENT(0.D0,0)
  I=MAXATM

```

```

246 NATOM=I
  ***** CONVERT TEMP FACTOR COEF TO STANDARD TYPE ZERO *****
  NG1=0
  DO 450 I=1,NATOM
  TI=PA(I,I,I)
  interim fix for IBM AIX
  K9=7
  K=1.+PA(K9,1,I)
  IF(TI)255,250,255
250 TI=.1
  GO TO 405
255 T6=.0506605918
  GO TO(270,260,265,270,260,400,405,270,260,270,450),K
  ***** TYPE 1 *****
260 DO 262 J=4,6
262 PA(J,I,I)=PA(J,I,I)*.5
  GO TO 270
  ***** TYPES 2 AND 3 (BASE 2 SYSTEMS) *****
265 T6=.351152464
  IF(K-4)270,260,270
  ***** TYPES 0 THROUGH 5 *****
270 IF(PA(2,1,I))400,400,272
272 DO 300 J=1,3
  DO 300 I=J,3
  T2=T6
  IF(K-5)285,275,275
  IF(K-6)280,280,281
  ***** TYPES 4 AND 5 *****
280 T2=B(J)*B(L)*T2*.25
  GO TO 285
  ***** TYPES 8 AND 9 (U(I,J) TENSOR SYSTEMS) *****
281 T2=B(J)*B(L)
  IF(K-11)285,282,282
  ***** TYPE 10, (CARTESIAN TENSOR SYSTEM) *****
282 T2=1.0
285 IF(J-L)290,287,290
287 VT(J,J)=T2*PA(J,I,I)
  GO TO 300
290 M=J+L+1
  VT(J,L)=T2*PA(M,I,I)
  VT(L,J)=VT(J,L)
300 CONTINUE
  ***** FIND PRINCIPAL AXES *****
  IF(K-11)310,305,305
305 CALL MM(VT,O,PAC)
  CALL MM(REFV,PAC,VT)
310 CALL MM(VT,AA,DA)
  CALL EIGEN(DA,RMS,PAT)
  ***** ARE EIGENVALUES POSITIVE *****
320 IF(RMS(1))325,325,320
325 NG=3
330 NG1=1
  CALL ERENT(DBLE(I)*D100K+55501.D0,0)
  ***** 3 EQUAL EIGENVALUES, USE REFERENCE VECTORS *****
340 T3=SIGN(SORT(ABS(RMS(1)+RMS(2)+RMS(3))/3.),RMS(1))
  DO 345 J=1,3
  DO 342 K=1,3
342 PA(J,K,I)=REFV(J,K)
345 EV(J,I)=T3
  GO TO 450

```

```

350 IF(NG+6)340,340,352
C ***** TWO EQUAL EIGENVALUES *****
352 N=NG+5
CALL UNITY(PAT(1,N),V1,-1)
DO 354 K=1,3
IF(ABS(VMV(V1,AA,REFV(1,K)))-.58)356,354,354
354 CONTINUE
356 CALL MM(AA,DA,VT)
DO 359 K=1,3
L=MOD(N+K-2,3)+1
DO 358 J=1,3
358 PA(J,L,I)=DA(J,K)
359 EV(L,I)=SIGN(SQRT(ABS(VMV(DA(1,K),VT,DA(1,K))))),RMS(L))
GO TO 450
C ***** MAKE EIGENVECTORS 1 ANGSTROM LONG *****
360 CALL AXES(PAT(1,1),PAT(1,3),PA(1,1,1),-1)
370 NG=0
C ***** SORT EIGENVALUE = RMS DISPLACEMENT *****
DO 375 J=1,3
T2=RMS(J)
375 EV(J,I)=SIGN(SQRT(ABS(T2)),T2)
GO TO 450
C ***** TYPE 6 (ISOTROPIC TEMP FACTOR) *****
400 T1=SQRT(T1*1.266515E-2)
C ***** TYPE 7 (DUMMY SPHERE OR ELLIPSOID OF REVOLUTION) *****
405 IF(PA(2,1,I))409,409,406
C ***** ELLIPSOID OF REVOLUTION FOR PASS OR PALE *****
406 EV(1,I)=T1
EV(2,I)=PA(2,1,I)
EV(3,I)=PA(2,1,I)
GO TO 411
C ***** SPHERE FOR PEAK OR PIT, OR A GENERAL SPHERE ATOM *****
409 DO 410 J=1,3
410 EV(J,I)=T1
411 IF(PA(3,1,I))430,430,415
C ***** FIRST DEFINED VECTOR FOR SPHERE OR CRITICAL POINT *****
415 DO 417 J=1,2
TD=PA(J+2,1,I)
CALL ATOM(TD,VT(1,J))
IF(NG)416,417,416
416 CALL ERENT(TD,0)
GO TO 430
417 CONTINUE
CALL DIFV(VT(1,2),VT(1,1),V1)
T11=SQRT(VMV(V1,AA,V1))
DO 418 J=1,3
C ***** SECOND DEFINED VECTOR FOR SPHERE OR CRITICAL POINT *****
418 V1(J)=V1(J)/T11
DO 420 J=3,4
TD=PA(J+2,1,I)
IF(TD.EQ.0.0)GO TO 422
CALL ATOM(TD,VT(1,J))
IF(NG)419,420,419
419 CALL ERENT(TD,0)
GO TO 430
420 CONTINUE
CALL DIFV(VT(1,4),VT(1,3),V2)
T11=SQRT(VMV(V2,AA,V2))
DO 421 J=1,3
421 V2(J)=V2(J)/T11

C ***** CHECK FOR NEARLY PARALLEL UNIT VECTORS *****
IF(ABS(VMV(V1,AA,V2))-.LT.0.9)GO TO 429
C ***** SUBSTITUTE BEST REFERENCE VECTOR *****
422 T22=1.0
J22=0
DO 424 J=1,3
T11=ABS(VMV(V1,AA,REFV(1,J)))
IF(T22.LE.T11)GO TO 424
T22=T11
J22=J
424 CONTINUE
DO 425 J=1,3
425 V2(J)=REFV(J,J22)
429 CALL AXES(V1,V2,PA(1,1,1),-1)
GO TO 450
C ***** REFERENCE VECTORS FOR SPHERE *****
430 DO 435 J=1,9
435 PA(J,1,I)=REFV(J,1)
450 NG=0
C ***** WRITE OUT RMS VALUES *****
LINES=LINES+2
IF(LINES-56)458,458,455
455 LINES=-1
GO TO 460
458 IF (NOUT.GE.0)
&WRITE (NOUT,461)
460 DO 465 I=1,NATOM
LINES=MOD(LINES+1,56)
IF(LINES)465,462,465
461 FORMAT(10HONO.ATOM 8X,1HX10X,1HX10X,1HZ13X,7HRMSD 1 4X,7HRMSD 2 4
1X,7HRMSD 3 )
462 IF (NOUT.GE.0)
&WRITE (NOUT,177)(TITLE(J),J=1,18)
IF (NOUT.GE.0)
&WRITE (NOUT,461)
463 FORMAT(1H ,I3,1X,A6,3F11.6)
465 IF (NOUT.GE.0)
&WRITE (NOUT,209)I,CHEM(I),(P(J,I),J=1,3),(EV(J,I),J=1,3)
1)
IF(NG1)999,999,470
470 CALL EXIT(NG)
999 RETURN
END
C *****GENERAL INITIALIZATION OF PRIME PARAMETERS*****
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BR(3,3),BRDR,CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,ITILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCALI
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),VRKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
INTEGER*2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
BEDR=0.5
C *****CALCULATE CONSTANTS*****

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DO 2950 I=1,5
2950 CONT(I)=SORT(1./.(2.*(1.+COS(3.141593 /2.**I))))
C
DISP= .005
dls=0.
FORE=.866
ITILT=0
LATM=0
MAXATM=505
NGD=0
NG=0
DO 3000 J=1,3
3000 ORGN(J) = 0.0
RES(1)=.75
RES(2)=.5*res(1)
RES(3)=.25*res(2)
SCAL1=1.0
SCAL2=1.54
SCL=1.54
DO 3005 I=1,3
3005 SYMB(I,I)=1.
SYMB(I+1,1)=0.
SYMB(I+5,1)=0.
TAPER=.375
THETA=0.0
VLEW=0.0
XLNG(1)=10.5
XLNG(2)=8.0
XO(1)=5.25
XO(2)=4.0
XO(3)=0.0
***** INITIATE OVERLAP ROUTINES *****
CALL LAP500(0)
RETURN
END
SUBROUTINE PROJ(DP,X,XO,VIEW,I1,I2,I3)
***** 3D CARTESIAN TO 2D PLOTTER COORDINATES *****
DIMENSION D(3,129),DP(2,129),X(3),XO(3)
T3=VLEW-X(3)
DO 145 I=I1,I2,I3
145 DP(1,I)=TI-XO(I)
DP(2,I)=T3+XO(2)
RETURN
END
120 T4=VIEW/(T3-D(3,I))
T1=TI*T4
T2=T2*T4
135 DP(1,I)=TI-XO(I)
145 DP(2,I)=T2+XO(2)
RETURN
END
SUBROUTINE RADIAL(ND)
***** GENERATE ELLIPSE FROM TWO CONJUGATE VECTORS *****
***** ORTHONORMAL VECTORS PRODUCE 8-128 SPOKED CIRCLE *****
***** ND DENOTES NUMBER OF SUBDIVISIONS (1 TO 5) *****
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAWRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR_CD(8,20)
2 ,CONT(5),D(3,130),DA(3,3),DE(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TITLIT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,INOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,3),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
SUBROUTINE READIN(IU,CHEM,IDI,ID2,X1,X2,X3,IT,IS,B1,B2,B3,B4,
1 ,integer*2 id1,id2
character*1 chain
character*3 res
character*4 atom
character*6 rec
character*8 chem
b1=1
b2=0
b3=0
b4=0
b5=0
b6=0
btype=7.
id1=0
id2=0
it=2
***** read the pdb file *****
read (iu,201) rec,iserno,atom,res,chain,id2,x1,x2,x3,occ,tf
format(a6,i5,lx,a4,lx,a3,lx,a1,i4,4x,3f8.0,2f6.0)
id1=9
if (atom.eq.' N ') id1=1
if (atom.eq.' CA ') id1=2
if (atom.eq.' C ') id1=3
if (atom.eq.' O ') then
id1=4
bi=.15
chem=atom(2:4)//res
is=0
read (iu,202,end=203) rec
format(a6)
backspace(iu)
return
203 is=1
return
C
***** read the pdb file *****
read (iu,201) rec,iserno,atom,res,chain,id2,x1,x2,x3,occ,tf
format(a6,i5,lx,a4,lx,a3,lx,a1,i4,4x,3f8.0,2f6.0)
id1=9
if (atom.eq.' N ') id1=1
if (atom.eq.' CA ') id1=2
if (atom.eq.' C ') id1=3
if (atom.eq.' O ') then
id1=4
bi=.15
chem=atom(2:4)//res
is=0
read (iu,202,end=203) rec
format(a6)
backspace(iu)
return
203 is=1
return

```

```

end
SUBROUTINE recycle
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,CONT(5),P(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TILT,KD(5,20),LATM,NATOM,NCD,NJ,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,5),PAT(3,3),O(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCI,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,VT(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XLNG(3),XO(3),XT(3)
INTEGERS 2 IDENT
COMMON /PARMS/ CHEM(505),EV(3,505),P(3,505),PA(3,3,505)
1 ,IDENT(2,505),MAXATM
logical featur
featur=.false.
if (ifcat.eq.1 .or. ifeat.eq.2) .and.
& ((mod(nj2,10).eq.5 .or. mod(nj2,10).eq.6)) featur=.true.
& ***** OBTAIN PROBLEM PARAMETERS *****
D10K=10000.
D100K=100000.
IF (NOUT.GE.0)
&WRITE (NOUT,20)
20 FORMAT(1H0 9X,82H FROM ATOMS TO ATOMS WITH RADIUS OR
1 , IF A BOX, WITH SEMIDIMENSIONS /11X,46HCODE (MIN MAX) (MIN
2 MAX) 7X,IH8X,IH8X,IHC)
IF(AIN(1)-D10K)100,100,101
100 ITOM1=AIN(1)
SYITOM=55501.
GO TO 103
101 ITOM1=AIN(1)/D100K
SYITOM=DMOD(AIN(1),D100K)
102 IF(DABS(AIN(2))-D10K)103,103,104
103 ITOM2 = DABS(AIN(2))
SYITO2=SYITOM
GO TO 105
104 ITOM2=DABS(AIN(2))/D100K
SYITO2=DMOD(DABS(AIN(2)),D100K)
105 ITAR1=AIN(3)
IF (ITAR1)108,108,110
108 ITAR1=1
110 ITAR2=AIN(4)
DMAX=AIN(5)
IF(DMAX)115,115,120
115 DMAX=4.
AIN(5)=DMAX
120 DMX=DMAX*DMAX
TEM=.01
KFUN=NJ*100+MOD(NJ2,10)
K=NJ*100+NJ2
I0=SYITOM
I02=SYITO2
LATOM=LATM
121 FORMAT(1H0,10X,2I3,I5,I4,I5,2I4,18X,3F9.3/1H )
IF (NOUT.GE.0)
&WRITE (NOUT,121)K,ITOM1,I0,ITOM2,I02,
1,ITAR1,ITAR2,(AIN(J),J=5,7)
124 FORMAT(1H ,15X,2I5,I8,I5,2F9.3)
IF(NCD)130,130,125
125 IF (NOUT.GE.0)
&WRITE (NOUT,124) ((KD(J,I),J=1,4),(CD(J,I),J=1,2),I=1,
1NCD)
130 DO 135 J=1,4
W(1,J)=99.
135 W(2,J)=-99.

```

C

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if (ndraw.eq.2 .or. ndraw.eq.3) call getpap
c *** find 201(203) instruction
do 545 i=1,nque
if (que(i)(7:9).eq.'201' .or. que(i)(7:9).eq.'203') then
next=i
go to 570
end if
545 continue
c *** ZERO ATOMS ARRAY AND RETURN TO EXECUTE NEXT INSTRUCTION ***
570 LATM=0
DO 580 I=1,500
ATOMID(I)=0.
DO 580 J=1,3
580 ATOMS(J,I)=0.
RETURN
END
SUBROUTINE SCRIBER(Y,NPEN)
DIMENSION Y(2),YO(2)
c ***** SUBROUTINE WHICH LINKS WITH THE PLOTTER-SPECIFIC SUBROUTINES
IF(NPEN-3)210,205,205
c ***** KEEP TRACK OF COORDINATES FOR LAST PEN-UP LOCATION *****
205 YO(1)=Y(1)
YO(2)=Y(2)
NPO=0
RETURN
c ***** CALL MECHANICAL PLOTTER PLOTTING SUBROUTINE *****
210 IF(NPO)225,220,225
220 CONTINUE
CALL PLOT(YO(1),YO(2),3)
225 CONTINUE
CALL PLOT(Y(1),Y(2),2)
NPO=1
RETURN
END
SUBROUTINE SEARC
DIMENSION NW(6),DX(3),S1D(200),S2(200),U(3),V(3),W(2,4),WW(2,3)
DIMENSION X(4),Y(3),Z(3)
REAL*8 DZMIN,DZMAX,DZSTO,S1D,TD1,TD2,TD3,TD4,D10K,D100K
REAL*8 TD
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
CHARACTER*8 CHEM
COMMON NG,A(9),AA(3,3),AREV(3,3),AAMRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)

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inum=0
DO 153 KI=ITAR1,ITAR2
  i=ki
136 if (featur) then
  inum=inum+1
  if (ident(ifeat,i).ne.ki) go to 154
  end if
  TD=FLOAT(I)*D100K
  CALL ATOM(TD,X)
  IF(NG)140,145,140
140 CALL ERENT(TD,KFUN)
  GO TO 600
145 X(4)=X(1)-X(2)
  DO 155 J=1,4
  TEM=X(J)
  IF(W(2,J)-TEM)148,150,150
148 W(2,J)=TEM
150 IF(TEM-W(1,J))152,155,155
152 W(1,J)=TEM
155 CONTINUE
154 if (featur .and. inum.lt.natom) go to 136
153 CONTINUE
  KFUN2=MOD(KFUN,10)
  GO TO (165,165,160,156,165,165),KFUN2
  ***** FIND PARALLELEPIPED WHICH ENCLOSES TRICLINIC BOX *****
156 DO 158 J=1,3
158 DX(J)=AIN(J+4)
  GO TO 170
  ***** FIND PARALLELEPIPED WHICH ENCLOSES RECTANGULAR BOX *****
160 DO 162 J=1,3
  DX(J)=0.
  DO 162 I=1,3
  T9=AIN(I+4)
162 DX(J)=DX(J)+ABS(REFY(J,I))*T9)
  GO TO 170
  ***** FIND PARALLELEPIPED WHICH ENCLOSES DMAX SPHERE *****
165 T1=1.-A(4)*A(4)-A(5)*A(5)-A(6)*A(6)+2.*A(4)*A(5)*A(6)
  DO 168 J=1,3
168 DX(J)=SQRT((1.-A(J+3)**2)/T1)*DMAX/A(J)
  ***** START SEARCH AROUND REFERENCE ATOMS *****
170 LIST=0
  LAST=0
  M1=ITOM1
  M2=ITOM2
  IF(KFUN2-5)186,172,172
  ***** CONVOLUTE AND RELITERATIVE CONVOLUTE INSTRUCTIONS *****
172 IF(LATM)174,174,176
  ***** FAULT, NO ENTRIES IN ATOMS LIST *****
174 NG=12
  CALL ERENT(0,D0,KFUN)
  GO TO 600
  ***** CHECK FOR REFERENCE ATOMS IN ATOMS LIST *****
176 IF(LATM-LAST)600,600,177
177 LIST=LAST
  LAST=LATM
178 LIST=LIST+1
180 TDI=ATOMID(LIST)
  IF(LAST-LIST.LE.0 .OR. AIN(8).EQ.0.D0) GO TO 184
  ***** FIND SMALLEST ATOM NUMBER IN REMAINDER OF ATOMS LIST *****
LISTP1=LIST+1
DO 182 J=LISTP1,LIST
  IF(TD1.LE.ATOMID(J)) GO TO 182
DO 181 I=1,3
  T1=ATOMS(I,J)
  ATOMS(I,J)=ATOMS(I,LIST)
  TD1=ATOMID(J)
  ATOMID(J)=ATOMID(LIST)
  ATOMID(LIST)=TD1
181 ATOMS(I,J)=ATOMS(I,LIST)
  ATOMS(I,LIST)=T1
  ATOMID(LIST)=TD1
182 CONTINUE
184 ITOM=TD1/D100K
  if (featur) then
  & ident(ifeat,atom).gt.itom2) go to 178
  else
  IF(ITOM.LT.ITOM1 .OR. ITOM.GT.ITOM2) GO TO 178
  end if
  SYITOM=DMOD(TD1,D100K)
  SYIT02=SYITOM
  M1=ITOM
  M2=ITOM
  ***** SET INITIAL RUN PARAMETERS *****
186 M2=AMOD(SYITOM,100.)
  M5=AMOD(SYITOM/100.,1000.)
  M3=M5/100
  M4=MOD(M5/10,10)
  M5=MOD(M5,10)
  ***** SET TERMINAL RUN PARAMETERS *****
  N2=AMOD(SYIT02,100.)
  N3=N5/100
  N4=MOD(N5/10,10)
  N5=MOD(N5,10)
  ***** START SEARCH AROUND REFERENCE ATOMS *****
  DO 500 L5=M5,N5
  DO 500 L4=M4,N4
  DO 500 L3=M3,N3
  DO 500 L2=M2,N2
  DO 500 ITOM=M1,M2
  TD3=DBLE(ITOM)*D100K+DBLE(L3*10000+L4*1000+L5*100+L2)
  CALL ATOM(TD3,Y)
  IF(NG)188,190,188
188 CALL ERENT(TD3,KFUN)
  GO TO 500
  ***** K-SYMMETRY EQUIVALENT POSITION *****
190 NUM=0
  DO 400 K=1,NSYM
  ***** SUBTRACT SYMMETRY TRANSLATION FROM REFERENCE ATOM *****
  DO 192 J=1,3
  U(J)=Y(J)-NS(J,K)
  ***** DETERMINE LIMITING CELLS TO BE SEARCHED *****
  ***** FIRST, MOVE THE BOX THROUGH THE SYMMETRY OPERATION *****
  DO 200 J=1,3
  DO 200 L=1,2
  WW(L,J)=0.0
  DO 200 I=1,3
  TEM=FS(I,J,K)
  IF(TEM)194,200,196
  N=MOD(L,2)+1
  GO TO 198

```

```

196 N=L
198 WW(L,J)=WW(L,J)+N(N,I)*TEM
200 CONTINUE
C ***** CHECK FOR MIXED INDEX TRANSFORMATION *****
DO 215 J=1,2
TEM=FS(1,J,K)
IF(TEM+FS(2,J,K))215,201,215
201 IF(TEM)203,215,207
203 WW(1,J)=W(2,4)*TEM
WW(2,J)=W(1,4)*TEM
GO TO 215
207 WW(1,J)=W(1,4)*TEM
WW(2,J)=W(2,4)*TEM
215 CONTINUE
C ***** MOVE 4 CELLS AWAY THEN MOVE BACK UNTIL PARALLELEPIPED AROUND
C REF ATOM AND BOX AROUND TRANSFORMED ASYM UNIT INTERSECT *****
N=0
DO 235 J=1,3
DO 225 I=1,2
N=N+1
TT=(U(J)-WW(I,J))*FLOAT(I*2-3)-DX(J)
TEM=5.0
221 TEM=TEM-1.0
IF(TEM+TT)225,225,221
225 NW(N)=TEM*FLOAT(I*2-3)+5.
***** IF NO POSSIBILITY OF A HIT, GO TO NEXT SYMMETRY OPER *****
IF(NW(N)-NW(N-1))400,235,235
235 CONTINUE
LL=NW(1)
LU=NW(2)
ML=NW(3)
MU=NW(4)
NL=NW(5)
NU=NW(6)
***** L CELL TRANSLATIONS IN X *****
DO 396 I=LL,LU
V(1)=U(1)+FLOAT(I-5)
***** M CELL TRANSLATIONS IN Y *****
DO 396 M=ML,MU
V(2)=U(2)+FLOAT(M-5)
***** N CELL TRANSLATIONS IN Z *****
DO 396 NN=NL,NU
V(3)=U(3)+FLOAT(NN-5)
***** I = TARGET ATOM *****
inum=0
DO 396 KI=ITAR1,ITAR2
I=KI
244 IF (feat) then
inum=inum+1
i=inum
if (ident(ifeat,i).ne.ki) go to 395
end if
DO 250 J=1,3
TEM=0.0
DO 245 II=1,3
TEM=TEM+FS(II,J,K)*P(II,I)
***** SEE IF WITHIN PARALLELEPIPED*****
TEM=TEM-V(J)
IF(DX(J)-ABS(TEM))395,250,250
250 X(J)=TEM
GO TO (255,255,252,265,255,255),KFUN2

```

```

C ***** SEE IF WITHIN MODEL BOX *****
252 CALL VM(X,AAREV,V1(2))
DO 253 J=2,4
IF(AIN(J+3)-ABS(V1(J)))395,253,253
253 CONTINUE
GO TO 265
C ***** SEE IF WITHIN SPHERE *****
DSQ=VM(X,AA,X)
IF(DMX-DSQ)395,256,256
256 IF(DSQ-.0001)258,260,260
258 IF(KFUN-402)395,260,260
260 TEM=SQRT(DSQ)
IF(AIN(8))/265,265,261
C *****SELECT ONLY FIRST ASYMMETRIC UNIT ENCOUNTERED *****
261 IF(LATM)265,265,262
262 DZMIN=DSLE(1)*D100K
DZMAX=DZMIN+D100K
DO 264 J=1,LATM
DZSTO=ATOMID(J)
IF(DZSTO-DZMIN)264,263,263
263 IF(DZMAX-DZSTO)264,264,395
264 CONTINUE
C ***** SELECT VECTORS ACCORDING TO CODES IF ANY *****
265 IF(INCL.LE.0) go to 277
C IF LOGC=0, screening conditions are ORED
C IF LOGC=1, screening conditions are ANDED
LOGC=AIN(8)
268 DO 275 J=1,NCD
NORG=ITOM
NTAR=I
IF (KD(5,J).EQ.1) THEN
NORG=IDENT(1,ITOM)
NTAR=IDENT(1,I)
END IF
IF (KD(5,J).EQ.2) THEN
NORG=IDENT(2,ITOM)
NTAR=IDENT(2,I)
END IF
IF (LOGC.EQ.0) THEN
IF (KD(2,J).GT.0) THEN
IF (NORG.LT.KD(1,J) .OR. NORG.GT.KD(2,J)) go to 275
END IF
IF (KD(4,J).GT.0) THEN
IF (NTAR.LT.KD(3,J) .OR. NTAR.GT.KD(4,J)) go to 275
END IF
IF (CD(2,J).GT.0.) THEN
IF (TEM.LT.CD(1,J) .OR. TEM.GT.CD(2,J)) go to 275
END IF
GO TO 277
END IF
IF (LOGC.EQ.1) THEN
IF (KD(2,J).GT.0) THEN
IF (NORG.LT.KD(1,J) .OR. NORG.GT.KD(2,J)) go to 276
END IF
IF (KD(4,J).GT.0) THEN
IF (NTAR.LT.KD(3,J) .OR. NTAR.GT.KD(4,J)) go to 276
END IF
IF (CD(2,J).GT.0.) THEN
IF (TEM.LT.CD(1,J) .OR. TEM.GT.CD(2,J)) go to 276

```

```

end if
if (j.eq.ncd) go to 277
end if
275 CONTINUE

276 GO TO 395
277 TD=D100K*DBLE(I)+DBLE((1110-L*100-M*10-NN)*100+K)
IF(KFUN-402)278,325,325
***** DETERMINE CORRECT POSITION IN SORTED VECTOR TABLE *****
278 IF(NUM)317,317,279
279 DO 315 I=1,NUM
IT=S2(II)-TEM
IF(ABS(TT)-0.0001)297,297,281
281 IF(TT)315,297,283
***** MOVE LONGER VECTORS TOWARD END OF TABLE *****
283 IF(200-NUM)287,287,289
287 NUM=199
289 IJ=NUM
DO 295 J=II,NUM
SID(IJ+1)=SID(IJ)
S2(IJ+1)=S2(IJ)
295 IJ=IJ-1
GO TO 319
***** CHECK FOR DUPLICATE VECTORS IF DISTANCES ARE EQUAL *****
297 CALL ATOM(SID(II),Z)
DO 305 J=1,3
IF(ABS(X(J)+Y(J)-Z(J))-0.0001)305,305,315
305 CONTINUE
315 CONTINUE
GO TO 395
***** STORE THE RESULT IN VECTOR TABLE *****
317 II=NUM+1
319 NUM=NUM+1
SID(II)=TD
S2(II)=TEM
320 IF(KFUN-106)395,325,325
***** STORE RESULT IN ATOMS TABLE *****
325 DO 330 J=1,3
330 VI(J)=X(J)+Y(J)
CALL STOR(TD)
395 if (featur .and. inum.lt.natom) go to 244
396 CONTINUE
400 CONTINUE
C ***** PRINT OUT DISTANCES *****
421 FORMAT(1H0,10X,20VECTORS FROM ATOM (I3,1H,15,1H)6X,8HTO ATOMSI4,
18H THROUGH,I4)
I0 = DMOD(TD3,D100K)
IF (NOUT.GE.0)
&WRITE (NOUT,421)ITOM,I0,ITAR1,ITAR2
423 DO 435 I=1,NUM
TD2=SID(I)
I1=TD2/D100K
I2=TD2-DBLE(I1)*D100K
CALL ATOM(TD2,Z)
IF(I-1)432,432,434
427 FORMAT(1H 13X,2(A6,1X),.39X,1H(I3,1H,15,1H)3F7.4,7X,3HD =F6.3)
429 FORMAT(1H 13X,2(A6,1X),.2(3H (I3,1H,15,1H)3F7.4,3X),4X,3HD =F6.3)
432 IF (NOUT.GE.0)
&WRITE (NOUT,429)CHEM(ITOM),CHEM(I1),ITOM,I0,(Y(J),J=1,3)
1),I1,I2,(Z(J),J=1,3),S2(I)
GO TO 435
&WRITE (NOUT,427)CHEM(ITOM),CHEM(I1),I1,I2,(Z(J),J=1,3),
1S2(I)
434 IF (NOUT.GE.0)
&WRITE (NOUT,427)CHEM(ITOM),CHEM(I1),I1,I2,(Z(J),J=1,3),
1S2(I)
435 CONTINUE
***** CALCULATE ANGLES ABOUT REF ATOM IF CODE IS 102 *****
C
437 IF(KFUN-102)500,451,500
441 FORMAT(1H0,10X,18ANGLES AROUND ATOM,I5)
451 IF (NOUT.GE.0)
&WRITE (NOUT,441)ITOM
L=NUM-1
IF(L)500,500,457
DO 465 I=1,L
L=NUM-I
I1=TD2/D100K
I2=TD2-DBLE(I1)*D100K
CALL ATOM(TD2,X)
CALL DIFV(X,Y,U)
CALL MV(AA,U,V2)
M=I+1
DO 465 J=M,NUM
TD4=SID(J)
J1=TD4/D100K
J2=TD4-DBLE(J1)*D100K
CALL ATOM(TD4,Z)
CALL DIFV(Z,Y,V)
F=ARCCOS(VV(V,V2)/(T3*S2(J)))
F1=SORT(VMV(V3,AA,V3))
460 FORMAT(1H 13X,3(A6,1X),7X,3(2H (I3,1H,15,1H)),12X,3HD =F6.3,7X,3HA
1=F6.2)
IF (NOUT.GE.0)
&WRITE (NOUT,460)CHEM(I1),CHEM(ITOM),CHEM(J1),I1,I2,ITOM
1,I0,J1,J2,F1,F
465 CONTINUE
495 CONTINUE
500 CONTINUE
IF(LAST-LIST)505,505,178
505 IF(KFUN-6)600,176,600
600 IF(KFUN-106)610,605,610
605 LATM=LATOM
610 RETURN
END
SUBROUTINE SIMBOL(W,W2,HGT,ITXT,THT,N)
DIMENSION W(3),ITXT(72),DS(10),DC(10)
DIMENSION IPTR(64),NKNT(64),IXYT(349)
DIMENSION IPTR(90),NKNT(90),IXYT(556)
CHARACTER*1 ITXT
common /ns/ ipf,ndraw,norient,nvar
common /trfac/ xtrans,ytrans
DATA IPTR
1/312,32,41,54,47,20,20,54,64,70,103,108,17,115,118,128
2,140,128,140,194,166,102,208,120,27,211,216,0,0,0,0,237
3,0,150,328,301,168,223,92,328,88,188,180,1,82,180,82,30
4,128,239,244,252,269,277,258,284,192,289,76,76,11,6,14,153
5,350,360,370,378,388,398,407,421,428,436,444,450,455,465,472,484
6,494,504,510,521,526,533,536,541,546,550/
DATA NKNT

```

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1/ 16, 9, 12, 8, 7, 6, 10, 6, 5, 7, 3, 5, 4, 9
2, 7, 12, 10, 12, 6, 3, 5, 5, 7, 0, 0, 0, 2
3, 0, 7, 9, 11, 12, 14, 10, 4, 4, 8, 5, 6, 2, 5, 2
4, 9, 5, 8, 13, 8, 9, 11, 5, 16, 12, 11, 12, 3, 5, 3, 13
5, 10, 10, 8, 10, 10, 9, 14, 7, 8, 8, 6, 5, 10, 7, 9, 10
6, 10, 6, 11, 5, 7, 3, 5, 4, 7/
C @ A B C D E F G H I J K L M N O
C P Q R S T U V W X Y Z
C 0 1 2 3 4 5 6 7 8 9 : ; < = > ?
C a b c d e f g h i j k l m n o p
C q r s t u v w x y z
DATA IXYT
1/44,48,46,26,66,24,64,99,66,26,68,26,64,24,66,28,29,22,62,69
2,29,26,56,26,22,62,29,62,99,22,69,22,25,65,25,28,39,59,68,62
3,63,65,56,26,56,67,68,59,29,22,52,63,68,68,59,39,28,23,32,52
4,63,65,55,22,29,26,66,69,62,32,52,42,49,39,59,35,36,46,45,35
5,99,42,32,33,43,42,31,69,58,53,62,62,37,38,49,58,25,24,33,43
6,64,29,23,32,52,63,69,29,22,25,69,99,47,62,22,29,45,69,62,29
7,22,46,62,69,47,69,99,68,59,39,28,23,32,52,63,68,99,44,62,22
8,29,59,68,67,56,26,56,65,62,49,44,99,32,43,52,32,99,44,46,56
9,67,68,59,39,28,29,69,49,42,99,23,53,64,65,56,36,27,38,68,25
A,65,45,63,27,45,23,67,29,38,33,22,56,67,68,59,39,28,27,36,56
B,65,63,52,32,25,36,29,42,69,29,47,42,47,69,29,69,22,62,99
C,36,56,38,28,29,39,38,99,69,22,99,53,63,62,52,53,15,75,38,49
D,42,32,52,28,39,59,68,66,24,22,62,28,39,59,68,67,56,36,56,65
E,63,52,32,23,28,39,59,68,29,24,64,54,59,52,42,62,23,32,52,63
F,65,56,26,29,69,68,43,42,23,32,52,63,68,59,39,28,26,35,55,66
G,24,64,54,53,57,56,66,26,36,37,33,66,57,47,36,35,44,54,65,67
H,58,38,27,24,33,53,64,57,49,59,57,99,37,29,39,37,22,32,12,22
I,23,21,22,31,13,22,33,11,22
c 349 a,62,67,66,57,37,26,23,32,52,63
b,22,29,26,37,57,66,63,52,32,23
c,63,52,32,23,26,37,57,66
d,62,69,66,57,37,26,23,32,52,63
e,63,52,32,23,26,37,57,66,65,35
f,32,36,26,46,36,38,49,59,68
g,64,67,66,57,37,26,24,33,53,64,62,51,31,22
c 420 h,22,29,26,37,57,66,62
i,32,52,42,47,37,99,48,49
j,32,41,51,62,67,99,68,69
k,22,29,24,57,35,62
l,32,52,42,49,39
m,22,27,26,37,46,42,46,57,66,62
c 458 n,22,27,26,37,57,66,62
o,63,52,32,23,26,37,57,66,63,99,67,45
p,21,27,26,37,57,66,64,53,33,24
q,61,67,66,57,37,26,24,33,53,64
r,22,27,26,37,57,66
c 500 82 s,23,32,52,63,64,55,35,26,37,57,66
c t,42,48,46,36,56
u,42,49,47,27,67
v,67,62,63,52,32,23,27
w,27,42,67
x,27,22,45,62,67
y,27,32,46,52,67
z,22,67,99,27,62

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Y,27,43,67,31
Y,27,44,67,21,21
Y,27,42,31,42,67
Z,62,22,67,27,99,35,55/
DATA RAD/0.01745329/
if (ndraw.eq.9) then
write (npf,21) n,nvar,w(1)+xtrans,w(2)+ytrans,8.*hgt,tht
format('TXP',i2,lx,i2,' ',4(1x,f10.6))
write (npf,22) (itxt(k),k=1,n)
format(80a1)
return
end if
C-----TEST FOR SPECIAL CASE OF CENTERED SYMBOL
IF(N.LE.0) GO TO 400
C-----SET UP TABLE OF INCREMENTS BASED ON HGT AND THT
IF(THT.EQ.0.0) GO TO 120
TH=RAD*THT
ST=SIN(TH)
CT=COS(TH)
GO TO 130
120 ST=0.0
130 D=HGT/7.0
DST=D*ST
DCT=D*CT
DS(1)=-DST
DC(1)=-DCT
DO 145 I=2,10
DS(I)=DS(I-1)+DST
DC(I)=DC(I-1)+DCT
145 CONTINUE
C-----START LOOP THROUGH THE N CHARACTERS OF ITXT
XO=0.0
YO=0.0
DO 370 J=1,N
ITXTJ=ICHAR(ITXT(J))
if (itxtj.ge.97.and.itxtj.le.122) then
ichs=itxtj-32
go to 221
end if
C-----MASK IT TO SIX BITS AND ADD ONE. PICK UP POINTER AND COUNTER
220 ICH=MOD(ITXTJ,64)+1
221 IP=IPTR(ICH)
NK=NKNT(ICH)
C-----TEST FOR SPACE OR UNDEFINED CHARACTER
IF(NK.EQ.0) GO TO 360
C-----START LOOP THROUGH SEGMENTS OF CHARACTER. LIFT PEN INITIALLY
IPEN=3
DO 350 K=1,NK
IXY=IXYT(IP)
C-----LIFT PEN IF SPECIAL INDICATOR IS FOUND
IF(IXY.NE.99) GO TO 300
IPEN=3
GO TO 340
300 IX=IXY/10
IY=IXY-10*IX
DX=XO+DC(IX)-DS(IY)
DY=YO+DC(IY)+DS(IX)
CALL DRAW(W,DX,DY,IPEN)
C-----PUT PEN DOWN TO DRAW NEXT SEGMENTS
IPEN=2

```



```

C
      Z(3,3)=X(3,3)*Y(3,3)
      DIMENSION X(3,3),Y(3,3),Z(3,3)
      X11=X(1,1)
      X12=X(1,2)
      X13=X(1,3)
      X21=X(2,1)
      X22=X(2,2)
      X23=X(2,3)
      X31=X(3,1)
      X32=X(3,2)
      X33=X(3,3)
      Y11=Y(1,1)
      Y12=Y(1,2)
      Y13=Y(1,3)
      Y21=Y(2,1)
      Y22=Y(2,2)
      Y23=Y(2,3)
      Y31=Y(3,1)
      Y32=Y(3,2)
      Y33=Y(3,3)
      Z(1,1)=X11*Y11+X21*Y21+X31*Y31
      Z(1,2)=X12*Y11+X22*Y21+X32*Y31
      Z(1,3)=X13*Y11+X23*Y21+X33*Y31
      Z(2,1)=X11*Y12+X21*Y22+X31*Y32
      Z(2,2)=X12*Y12+X22*Y22+X32*Y32
      Z(2,3)=X13*Y12+X23*Y22+X33*Y32
      Z(3,1)=X11*Y13+X21*Y23+X31*Y33
      Z(3,2)=X12*Y13+X22*Y23+X32*Y33
      Z(3,3)=X13*Y13+X23*Y23+X33*Y33
      RETURN
END
      subroutine uinput(in,rout)
      user input routine
      common /ns/ npf,ndraw,norient,nvar
      common /df1/ infile,ldraw,lorient,iout,ext,atomfi,fpaplen
      character*60 fname,txtans,infile,atomfi
      character*4 ext
      character*1 answer
      call dflts
      iflag=0
      c *** get the input file name and open the file or "exit" ***
      if (iargc().eq.1) then
        call getarg(1,fname)
        open(in,file=fname,status='old',err=l10)
        go to 135
      end if
      110 fname=infile
        ipos=index(fname,' ')
        write (*,115) fname(1:ipos-1)
      115 format(' Enter instruction set file name or "exit" [',a,']: ', $)
      read (*,120) txtans
      120 format(a)
        * if (txtans(1:4).eq.'exit' .or. txtans(1:4).eq.'EXIT')
          * call exiting(0)
        open(in,file=fname,status='old',err=l25)
        go to 135
      125 ipos=index(fname,' ')
        fname=fname(1:ipos-1)
        write (*,130) fname(1:ipos-1)
C
      end if
      if (twoDIG) then
        inum=ksign * (m * 10 + k)
      else
        inum=ksign * k
      end if
      ts(kk,num)=float(inum)/float(iden)
    end if
  c *** look for and interpret decimal style fraction
  n=index(txt,',')
  if (n.gt.0) then
    get post decimal point portion
    k=i-char(txt(n-1:n+1))-48
    m=i-char(txt(n+2:n+2))-48
    if (m.ge.0 .and. m.le.9) then
      ts(kk,num)=float(k) * .1 + float(m) * .01
    else
      ts(kk,num)=float(k) * .1
    end if
  end if
  c
  get sign
  ksign=1
  if (n-1.ge.1) then
    if (txt(n-1:n-1).eq.'-') ksign=-1
  end if
  if (n-2.ge.1) then
    if (txt(n-2:n-2).eq.'-') ksign=-1
  end if
  ts(kk,num)=float(ksign) * ts(kk,num)
  end if
  c
  interpret xyz portion of symmetry operation
  do 303 i=1,24
    if (txt(i:i).eq.'X') then
      fs(1,kk,num)=1
      if (i.ge.2) then
        end if
        if (txt(i-1:i-1).eq.'-') fs(1,kk,num)=-1.
      end if
    end if
    if (txt(i:i).eq.'Y') then
      fs(2,kk,num)=1
      if (i.ge.2) then
        end if
        if (txt(i-1:i-1).eq.'-') fs(2,kk,num)=-1.
      end if
    end if
    if (txt(i:i).eq.'Z') then
      fs(3,kk,num)=1
      if (i.ge.2) then
        end if
        if (txt(i-1:i-1).eq.'-') fs(3,kk,num)=-1.
      end if
    end if
  303 continue
  return
end
SUBROUTINE TMM(X,Y,Z)
Z = TRANSPOSED (TRANSPOSE(X) * (Y) )
C

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```

130 format(/' ',a,'" does not exist')
go to 110
c *** determine where ortep drawing should go ***
135 write (*,140) idraw
140 format(' Drawing to (1) Screen, (2) Postscript file, (3) HPGL file
', or (0) Omit [' ,il,'] : ', $)
read (*,145) answer
145 format(a)
if (answer .eq. ' ') then
ndraw=idraw
else
read (answer,155) ndraw
end if
if (ndraw.lt.0. or .ndraw.gt.3. and.ndraw.ne.9)) then
write(6,*) 'invalid selection'
go to 135
end if
if (ndraw.eq.0. or.ndraw.eq.1.or.ndraw.eq.9) go to 149
go to 1451
c *** need to get this information if printing from editor
entry getpap
iflag=1
c *** determine orientation of drawing ***
1451 write (*,1452) iorient
1452 format(' (1) Portrait or (2) Landscape orientation [' ,il,'] : ', $)
read (*,145) answer
if (answer .eq. ' ') then
iorient=iorient
else
read (answer,155) norient
end if
if (norient.lt.1.or.norient.gt.2) then
write(6,*) 'invalid selection'
go to 1451
end if
c *** determine paper length for postscript landscape
if (ndraw.eq.2. and.norient.eq.2) then
write (*,1453) fpaplen
format(' How tall is printer page in inches? [' ,f5.2,'] : ', $)
read (*,120) txtans
if (txtans(1:1) .ne. ' ') read (txtans,1454) fpaplen
format(f10.0)
nvar=fpaplen*1000.
end if
c *** if called from recycle, return there
if (iflag.eq.1) return
c *** determine where ortep output should go ***
149 write (*,150) iout
150 format(' Text output to (1) File, (2) Screen, or (0) Omit [' ,il,
', ] : ', $)
read (*,145) answer
if (answer .eq. ' ') then
nout=iout
else
read (answer,155) nou
end if
155 format(il)
c *** set output unit number ***
nout=-4
if (nou .eq. 1) nout=4
if (nou .eq. 2) nout=6
130 format(/' ',a,'" does not exist')
if (nout .eq. 4) then
ipos=index(fname,' ')
if(ipos .ne. 0) then
fname=fname(1:ipos-1)//ext
go to 160
end if
ipos=index(fname,' ')
fname=fname(1:ipos-1)//ext
ipos=index(fname,' ')
write (*,165) fname(1:ipos-1)
format(' Enter output file name [' ,a,'] : ', $)
read (*,120) txtans
if (txtans(1:1) .ne. ' ') fname=txtans
open(nout,file=fname,status='old',err=170)
go to 175
170 open(nout,file=fname,status='new')
end if
175 continue
return
c *** get file name of an external file containing atomic parameters
entry gtafil(iu)
210 fname=atomfi
ipos=index(fname,' ')
write (*,215) fname(1:ipos-1)
format(' Enter atom parameter file name or "exit" [' ,a,'] : ', $)
read (*,120) txtans
if (txtans(1:4).eq.'exit' .or. txtans(1:4).eq.'EXIT')
* call exiting(0)
if (txtans(1:1) .ne. ' ') fname=txtans
open(iu,file=fname,status='old',err=225)
return
225 ipos=index(fname,' ')
write (*,130) fname(1:ipos-1)
go to 210
c *** ask user about using editor
entry go2edr
write (*,303)
303 format(/' Edit instruction set? (Y)es or (N)o [N] : ', $)
303 format(/' (1) Save drawing as Postscript file'//,
&' (2) Save drawing as HPGL file'//,
&' (3) Redraw structure on screen'//,
&' (4) Edit instruction set'//,
&' [Quit] : ', $)
read (*,304) answer
304 format(a)
if (answer.eq.'1') then
ndraw=2
call recycle
else
if (answer.eq.'2') then
ndraw=3
call recycle
else
if (answer.eq.'3') then
ndraw=1
call recycle
else
if (answer.eq.'4') call editr
end if
end if

```

```

end if
return
end
SUBROUTINE UNITY(X,Z,IATYPE)
DIMENSION X(3),Y(3),Z(3)
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,COMT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TITLIT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,3),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
Y(1)=X(1)
Y(2)=X(2)
Y(3)=X(3)
IF(IATYPE)125,125,105
105 T1=SQRT(Y(1)*Y(1)+Y(2)*Y(2)+Y(3)*Y(3))
GO TO 145
125 T1=SQRT(Y(1)*Y(1)+AA(1,1)+Y(2)*Y(2)+AA(2,2)+Y(3)*Y(3)+AA(1,3)+AA(3,1)+Y(2)*Y(2)+Y(3)*Y(3)+AA(2,3)+AA(3,2))+Y(3)*Y(3)+AA(3,2,3))
145 IF(T1)155,155,175
155 NG=5
GO TO 300
175 Z(1)=Y(1)/T1
Z(2)=Y(2)/T1
Z(3)=Y(3)/T1
300 RETURN
END
SUBROUTINE VMV(X,Z)
TRANSPOSED VECTOR * MATRIX
Z(3)=Y(3)*X(3,3)
DIMENSION X(3,3),Y(3),Z(3)
Y1=Y(1)
Y2=Y(2)
Y3=Y(3)
Z(1)=X(1,1)*Y1+X(2,1)*Y2+X(3,1)*Y3
Z(2)=X(1,2)*Y1+X(2,2)*Y2+X(3,2)*Y3
Z(3)=X(1,3)*Y1+X(2,3)*Y2+X(3,3)*Y3
RETURN
END
FUNCTION VMV(X1,O,X2)
TRANSPOSED VECTOR * MATRIX * VECTOR
VMV=X1(3)*Q(3,3)*X2(3)
DIMENSION X1(3),Q(3,3),X2(3)
VMV=X1(1)*X2(1)*Q(1,1)+X2(2)*Q(1,2)+X2(3)*Q(1,3))
& +X1(2)*X2(2)*Q(2,1)+X2(2)*Q(2,2)+X2(3)*Q(2,3)
& +X1(3)*X2(3)*Q(3,1)+X2(3)*Q(3,2)+X2(3)*Q(3,3)
RETURN
END
FUNCTION VV(X,Y)
TRANSPOSED VECTOR * VECTOR
VV=X(3)*Y(3)
DIMENSION X(3),Y(3)
VV=X(1)*Y(1)+X(2)*Y(2)+X(3)*Y(3)
RETURN
END
SUBROUTINE XYZ(DQA,X,IATYPE)

```

```

C ***** IATYPE .GT.0 CART. COORD. FROM ATOM CODE WORD *****
C ***** XABSF(IATYPE) .LE.2 FOR WORKING SYSTEM *****
C ***** XABSF(IATYPE) .GT.2 FOR REFERENCE SYSTEM *****
C ***** IATYPE .LE.0 USES TRICLINIC COORD. XT *****
DIMENSION X(3)
REAL*8 DQA
REAL*8 AIN,ATOMID
CHARACTER*4 TITLE,TITLE2
COMMON NG,A(9),AA(3,3),AAREV(3,3),AAWRK(3,3),AID(3,3)
1 ,AIN(140),ATOMID(500),ATOMS(3,500),BB(3,3),BRDR,CD(8,20)
2 ,COMT(5),D(3,130),DA(3,3),DP(2,130),DISP,EDGE,FORE,FS(3,3,96)
3 ,IN,TITLIT,KD(5,20),LATM,NATOM,NCD,NG,NJ2,NOUT,NSR,NSYM
4 ,ORGN(3),PAC(3,3),PAT(3,3),Q(3,3),REFV(3,3),RES(4),RMS(5),SCAL1
5 ,SCAL2,SCL,SYMB(3,3),TAPER,THETA,TITLE(18),TITLE2(18),TS(3,96)
6 ,VIEW,V(3,4),V1(4),V2(3),V3(3),V4(3),V5(3),V6(3),WRKV(3,3)
7 ,XING(3),XO(3),XT(3)
IT=IABS(IATYPE)-2
NG1=NG
NG=0
IF(IATYPE)10,10,5
5 CALL ATOM(DQA,XT)
IF(NG)30,10,30
10 T1=0.
DO 15 J=1,3
T2=XT(J)-ORGN(J)
V1(J)=T2
15 T1=T1+ABS(T2)
20 NG=NG1
30 DO 35 J=1,3
35 X(J)=0.
GO TO 300
40 IF(IT)45,45,60
***** RELATIVE TO WORKING SYSTEM *****
45 DO 55 I=1,3
T1=0.
DO 50 J=1,3
50 T1=T1+V1(J)*AAWRK(J,I)
55 X(I)=T1*SCAL1
GO TO 300
C ***** RELATIVE TO REFERENCE SYSTEM *****
60 DO 70 I=1,3
T1=0.
DO 65 J=1,3
65 T1=T1+V1(J)*AAREV(J,I)
70 X(I)=T1*SCAL1
300 RETURN
END
C ***** DUMMY SCREEN OUTPUT (MAY BE REPLACED WITH SCREEN DRIVER CODE) *****
C *****
C subroutine initsc
C return
C end
C subroutine penwsc(penw)
C return
C end
C subroutine colrsc(icolor)

```



```

call pgrrect(7.5,11.1,8.2,8.5)
call pgscli(1)
call pgtxt(7.6,8.3,'Hit <RETURN> or <ENTER> key')

call ppend
return
end

subroutine curssc
character ch
character*21 str
integer pgcurs
character*6 label,alabel
character*9 tomid,atomid
common /trfac/ xtrans,ytrans
common /ns/ npf,ndraw,norient,nvar

call pgsfs(1)
call pgscli(1)
call pgsch(1.)

c *** get cursor position
1 junk = pgcurs(x,y,ch)
if (ch.eq.'x'.or.ch.eq.'X') return
if (ch.eq.'d'.or.ch.eq.'D') return
if (x.ge.10.4.and.x.le.11. .and. y.ge.8.2 .and. y.le.8.5) return
if (ichar(ch).eq.13) return

c *** initial values for variables
xpt = x
ypt = y
adiffx = .0625
adiffy = .0625
odiffx = adiffx
odiffy = adiffy
atomid = ' '
alabel = ' '
iflag = 0
nflag = 0

rewind(npf)

2 read(npf,3,end=4) label,tomid,xx,yy
3 format(11x,a6,3x,a9,4x,2f8.0)
difix = abs(xx-xpt)
diffy = abs(yy-ypt)
if (diffx.le.adiffx .and. diffy.le.adiffy) nflag=nflag+1
if (diffx.le.odiffx .and. diffy.le.odiffy) then
atomid = tomid
alabel = label
odiffx = diffx
odiffy = diffy
end if
go to 2

4 if (nflag.eq.0) write(str,5)
if (nflag.eq.1) write(str,6) alabel,atomid
if (nflag.gt.1) write(str,7) alabel,atomid

5 format('Not near atom center')
6 format(a6,1x,a9)
7 format(a6,1x,a9,' + ??')

c *** erase rectangle
call pgscli(0)
call pgsfs(1)
call pgrrect(0.,2.8,8.2,8.5)
c *** redraw empty rectangle
call pgscli(1)
call pgsfs(2)
call pgrrect(0.,2.8,8.2,8.5)

c *** print atom information in rectangle
call pgtxt(0.1,8.3,str)
go to 1
end

c *** end of PGLOT specific routines
c *****
c ***** HPGL FILE OUTPUT
c *****
subroutine inithp
common /ns/ npf,ndraw,norient,nvar
character ESC
character*10 fname
do 11 i=1,999
write (fname, 10) i
10 format('REP',i3.3,'.PRN')
open(unit=npf,file=fname,status='old',err=12)
close(npf)
11 continue
12 open(unit=npf,file=fname,status='new')
write (*,13) fname
13 format('/', ' HPGL file name: ',a)
ESC=char(27)
write (npf,21) ESC
write (npf,22) ESC
write (npf,23)
if (norient.eq.2) write (npf,24)
21 format(a1,'E')
22 format(a1,'%0B')
23 format('IN:','SP1:','PW.15:')
24 format('RO90:;')
return
end
subroutine colrhp(icolor)
common /ns/ npf,ndraw,norient,nvar
c *** set plot color
c *** in OKREP icolor=0 => black
c *** plotter pen l=black
icol=icolor

```

```

if (icolor.eq.0) icol=1
write (npf,21) icol
21 format('SP',i1,'')
return
end

subroutine penwhp(penw)
common /ns/ npf,ndraw,norient,nvar
if (penw.eq.0.) then
penw=.15
else
penw=penw*.0252
end if
write (npf,21) penw
21 format('PW',f5.2,'')
return
end

subroutine penhp(x,y,ipen)
common /ns/ npf,ndraw,norient,nvar
common /trfac/ xtrans,ytrans
ix = nint((x + xtrans) * 1000.)
iy = nint((y + ytrans) * 1000.)

if (ipen.eq.2) write (npf,101) ix,iy
101 format('PD',i4,'',i4,';')
if (ipen.eq.3) write (npf,102) ix,iy
102 format('PU',i4,'',i4,';')
return
end

subroutine endhp
common /ns/ npf,ndraw,norient,nvar
character ESC
ESC=char(27)
write (npf,31)
31 format('PU',/,',',SP0',/,',',PG;',/,',',IN;')
34 format(a1,'%0A')
write (npf,34) ESC
write (npf,35) ESC
35 format(a1,'E')
close(npf)
return
end

c *** end of HPGL specific routines
c *****
c *****
c *** POSTSCRIPT FILE OUTPUT
c *****
subroutine initps
common /ns/ npf,ndraw,norient,nvar
common /ps/ ixmin,ixmax,iymin,iymax,ixt,iyt
character*10 fname

c *** initialize variables to calculate bounding box

```

```

ixmin=20000
ixmax=0
iymin=20000
iymax=0

do 11 i=1,999
write (fname, 10) i
10 format('TEP',i3.3,'.PRN')
open(unit=npf,file=fname,status='old',err=12)
close(npf)
11 continue
12 open(unit=npf,file=fname,status='new')
write (*,13) fname
13 format(/,' Postscript file name: ',a)

ixt=0
iyt=0
write (npf,21)
write (npf,22)
write (npf,23)
if (norient.eq.2) then
write (npf,24)
iyt=nvar
else
write (npf,25)
end if
write (npf,26)
write (npf,27)
write (npf,28)
write (npf,29)
write (npf,30)
write (npf,31)
write (npf,32) ixt,iyt
if (norient.eq.2) write (npf,33)
write (npf,34)
write (npf,35)
21 format('!PS-Adobe-3.0 EPSF-3.0')
22 format('%%Creator: ORTEP-III')
23 format('%%BoundingBox: (atend)',/,',',%%Pages: 1')
24 format('%%Orientation: Landscape')
25 format('%%Orientation: Portrait')
26 format('%%BeginProlog')
27 format('/m {moveto} def')
28 format('/l {lineto} def')
29 format('%%EndProlog',/,',',%%page: 1 1')
30 format('%%BeginPageSetup')
31 format('0.072 0.072 scale')
32 format('16,1x,16,1 translate')
33 format('0 setgray 1 setlinecap 5 setlinewidth')
35 format('%%EndPageSetup')
return
end

subroutine colprc(icolor)
common /ns/ npf,ndraw,norient,nvar
write (npf,101)
101 format('stroke')
if (icolor.eq.0) write (npf,1)
if (icolor.eq.1) write (npf,1)
if (icolor.eq.2) write (npf,2)

```

