

27-750, A.D. Rollett

Due: 16<sup>th</sup> February, 2020.

Homework 4, Single Slip, and Multiple Slip Crystal Plasticity

Q1. [30 points] *Single Slip: Calculating Schmid Factors*

In a single crystal tensile test on Ni, the orientation of the crystal is given as  $(18.89^\circ, 68.58^\circ, 11.31^\circ)$  in Bunge Euler angles. Assume that an axisymmetric tensile stress is applied along the sample 3 (Z) direction. Note that we apply stress boundary conditions, as discussed in class.

(a) Determine which crystal direction is parallel to the *tensile axis*. Give your answer in the form of  $(hkl)[uvw]$  and simplify the numbers to be single digit integers. You should be able to modify the spreadsheet or program that you wrote in earlier homeworks to produce an answer.

(a) The Miller indices for this crystal orientation are:  $(1\ 5\ 2)\ [3\ -1\ 1]$ , based on a standard conversion of Euler angles to matrix, to Miller indices<sup>†</sup>. Then, obviously, the tensile axis is  $//\ [1\ 5\ 2]$ .

(b) Identify which combination of slip plane and direction is active, and calculate the Schmid factor (to 3 significant figures).

(b)  $(-1\ 1\ 1)\ [1\ 1\ 0]$ ;  $m = 0.490$ .

Note: Ni is fcc and can slip on any  $\{111\}$  plane in any  $\langle 110 \rangle$  direction. You will have to find out which is the most highly stressed slip system, i.e. find the largest Schmid factor. This value of the Schmid factor is what you should use to determine the tensile yield stress because it determines which slip system will be activated first.

(c) Calculate the tensile yield stress (to 3 significant figures) based on a critical resolved shear stress that we will (arbitrarily) set at 100 MPa.

(c) 204 MPa.

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<sup>†</sup> SteveSintay\_Pole\_Figures-26May11.xls; SchmidFactors-2016.xlsx.

	A	B	C	D	E	F	G	H	I	J	K	L	
1	h	k	l	u	v	w	dot product	cos phi	cos lambda	Schmid Factor	ABS(Schmid)		
2	1	1	1	-1	1	0	0	0.843274	0.51639778	0.43546484	0.435464843	0.489897949	
3	-1	1	1	1	0	1	0	0.6324555	0.38729833	0.24494897	0.244948974	largest Abs of Schmid	
4	1	-1	1	1	0	-1	0	-0.210819	-0.1290994	0.02721655	0.027216553		
5	-1	-1	1	1	0	1	0	-0.421637	0.38729833	-0.1632993	0.163299316	critical resolved yield stress (input)	
6	1	1	1	-1	0	1	0	0.843274	0.12909944	0.10886621	0.108866211	100	
7	-1	1	1	0	-1	1	0	0.6324555	-0.3872983	-0.244949	0.244948974		
8	1	-1	1	1	1	0	0	-0.210819	0.77459667	-0.1632993	0.163299316	observed tensile yield stress = tau_crss/max_S Schmid	
9	-1	-1	1	0	1	1	0	-0.421637	0.90369611	-0.3810317	0.381031738	204.12	
10	1	1	1	0	-1	1	0	0.843274	-0.3872983	-0.3265986	0.326598632		
11	-1	1	1	1	1	0	0	0.6324555	0.77459667	0.48989795	0.489897949		
12	1	-1	1	0	1	1	0	-0.210819	0.90369611	-0.1905159	0.190515869		
13	-1	-1	1	1	-1	0	0	-0.421637	-0.5163978	0.21773242	0.217732422		
14	1	5	2										
15	TENSILE AXIS is above						(input)						
16							this column must be all zeros!						

You must submit a copy of the table of results showing how you calculated your answer which must display the maximum Schmid factor and the Miller indices of the activated slip system.

Q2. [30 points] *Compute Slip Matrices; verify 384 non-singular cases out of 792*  
Referring to slides 43 through 55 and #50 in particular, construct all the 5x5 slip matrices and determine which ones are non-singular. Verify that out of the  ${}_5C_{12}$  combinations, only 384 are valid solutions, i.e., have a non-zero determinant.

*Hints:*

Clearly you have to start by making a list of all 12 slip systems in fcc metals based on  $\{111\}\langle 110\rangle$ . Although this problem is deceptively simple, the main programming challenge is to compute a list of all the possible combinations of five slip systems (as opposed to just the number). Fortunately the web has code fragments that will do this (in C++, Fortran etc.), and Matlab has a built-in function.

*What to submit:*

You should submit your code (e.g., python or Matlab script), as well as the list of 384 non-singular cases.

*Answer:* See CalcTaylorMatrices.f90, where I chose to use Fortran to perform the calculations. I was able to find code for computing the combinations and already had a matrix inversion code (that Ben Anglin wrote).

Q3. [45 points] *Bishop-Hill Model: Calculation of Taylor Factors for Multiple Slip*  
Assume that slip occurs on  $\{111\}$  planes and in  $\langle 110\rangle$  directions. For each of the three different strain types, uniaxial tension parallel to the Z axis, plane strain in the +X & -Z,

and simple shear in the XZ sense (see the matrices below), calculate the following quantities.

- (a) the index of the active stress state (from the Bishop-Hill list);
- (b) the Taylor factor (to at least 2 decimal places) for crystals with the following orientations;
- (c) the active multi-slip stress state (as in the values of A, B, C, F, G, & H); for crystals with the following orientations.

- 2.1 (0 0 1) [1 0 0]
- 2.2 (0 8 10) [1 0 0]
- 2.3 (-1 1 0) [1 1 1]
- 2.4 (2 1 3) [3 6 -4]
- 2.5 (1 1 2) [1 1 -1].

The strain tensors for the 3 different strain types are as follows.

Uniaxial tension // z:

$$\begin{bmatrix} -0.5 & 0 & 0 \\ 0 & -0.5 & 0 \\ 0 & 0 & 1 \end{bmatrix}$$

Plane strain compression on +X and -Z:

$$\begin{bmatrix} 1 & 0 & 0 \\ 0 & 0 & 0 \\ 0 & 0 & -1 \end{bmatrix}$$

Simple Shear on XZ:

$$\begin{bmatrix} 0 & 0 & 1 \\ 0 & 0 & 0 \\ 0 & 0 & 0 \end{bmatrix}$$

Question	Orientation	Taylor Factor for Uniaxial Tension //Z	Stress State
2.1a	(0 0 1) [1 0 0]	2.449	1.0, -1.0, 0.0, 0.0, 0.0, 0.0
2.2a	(0 8 10) [1 0 0]	3.585	0.0, 0.0, 0.0, 1.0, 0.0, 0.0

2.3a	(-1 1 0) [1 1 1]	3.674	(to be determined by you)
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Answers are provided (above) for a few of the cases so that you can check your method. The meaning of the Miller indices is the conventional one discussed earlier in the class: (hkl) // sample-Z (=sample-3), [uvw] // sample-X (= sample-1).

The pseudo-code to obtain the Taylor factor is given in the lecture notes.

If you consider uniaxial tension for the three corners of the unit triangle (i.e. 001, 110 and 111), in which two corners is the inverse Schmid factor equal to the Taylor factor, and in which corner are they different?

Answer: See the description in Hosford, Ch. 3. – his definition of the Taylor factor is rather obscure but the von Mises equivalent strain approach seems to work well enough. The Taylor factors appear to be identical with those in the books. The following results were obtained with Bishop\_Hill.f90. If the index to the stress state is >28, it means that negative of the stress state with the same index minus 28 provided the highest dW.

Uniaxial Tension	Orientation	Taylor Factor, M	Index	Stress State (A, B, C, F, G, H)
2.1a	(0 0 1) [1 0 0]	2.449	29	1.0 -1.0 0.0 0.0 0.0 0.0
2.2a	(0 8 10) [1 0 0]	3.585	4	0.0 0.0 0.0 1.0 0.0 0.0
2.3a	(-1 1 0) [1 1 1]	3.674	34	0.0 0.0 0.0 0.5 -0.5 0.5
2.4a	(2 1 3) [3 6 -4]	3.149	5	-0.5 0.5 0.0 0.5 -0.5 0.0
2.5a	(1 1 2) [1 1 -1]	3.062	21	-0.5 0.5 0.0 0.5 0.5 0.0

Plane Strain	Orientation	Taylor Factor	Index	Stress State
2.1b	(0 0 1) [1 0 0]	2.121	1	1.0 -1.0 0.0 0.0 0.0 0.0
2.2b	(0 8 10) [1 0 0]	2.121	2	0.0 1.0 -1.0 0.0 0.0 0.0
2.3b	(-1 1 0) [1 1 1]	3.526	6	0.0 0.0 0.0 0.0 0.0 1.0
2.4b	(2 1 3) [3 6 -4]	3.019	49	-0.5 0.5 0.0 -0.5 -0.5 0.0
2.5b	(1 1 2) [1 1 -1]	3.182	49	-0.5 0.5 0.0 0.5 0.5 0.0

Simple Shear	Orientation	Taylor Factor	Index	Stress State		
2.1c	(0 0 1) [1 0 0]	4.243	5	0.0 0.0	0.0 1.0	0.0 0.0
2.2c	(0 8 10) [1 0 0]	3.313	5	0.0 0.0	0.0 1.0	0.0 0.0
2.3c	(-1 1 0) [1 1 1]	1.732	2	0.0 0.0	1.0 0.0	1.0 0.0
2.4c	(2 1 3) [3 6 -4]	2.541	13	1.0 0.0	-1.0 0.0	0.0 0.0
2.5c	(1 1 2) [1 1 -1]	2.0	1	1.0 0.0	-1.0 0.0	0.0 0.0

Bishop & Hill stress states (Fortran style); factor of  $\sqrt{6}$  *not* included:

```

stress(1,1)=1.
stress(1,2)=-1.
stress(1,3)=0.
stress(1,4)=0.
stress(1,5)=0.
stress(1,6)=0.
! # 1
stress(2,1)=0.
stress(2,2)=1.
stress(2,3)=-1.
stress(2,4)=0.
stress(2,5)=0.
stress(2,6)=0.
! # 2
stress(3,1)=-1.
stress(3,2)=0.
stress(3,3)=1.
stress(3,4)=0.
stress(3,5)=0.
stress(3,6)=0.
! # 3
stress(4,1)=0.
stress(4,2)=0.
stress(4,3)=0.
stress(4,4)=1.
stress(4,5)=0.
stress(4,6)=0.
! # 4
stress(5,1)=0.
stress(5,2)=0.
stress(5,3)=0.
stress(5,4)=0.
stress(5,5)=1.
stress(5,6)=0.
! # 5
stress(6,1)=0.
stress(6,2)=0.
stress(6,3)=0.
stress(6,4)=0.
stress(6,5)=0.
stress(6,6)=1.
! # 6
stress(7,1)=0.5
stress(7,2)=-1.

```

```
stress(7,3)=0.5
stress(7,4)=0.
stress(7,5)=0.5
stress(7,6)=0.
! # 7
stress(8,1)=0.5
stress(8,2)=-1.
stress(8,3)=0.5
stress(8,4)=0.
stress(8,5)=-0.5
stress(8,6)=0.
! # 8
stress(9,1)=-1.
stress(9,2)=0.5
stress(9,3)=0.5
stress(9,4)=0.5
stress(9,5)=0.
stress(9,6)=0.
! # 9
stress(10,1)=-1.
stress(10,2)=0.5
stress(10,3)=0.5
stress(10,4)=-0.5
stress(10,5)=0.
stress(10,6)=0.
! # 10
stress(11,1)=0.5
stress(11,2)=0.5
stress(11,3)=-1.
stress(11,4)=0.
stress(11,5)=0.
stress(11,6)=0.5
! # 11
stress(12,1)=0.5
stress(12,2)=0.5
stress(12,3)=-1.
stress(12,4)=0.
stress(12,5)=0.
stress(12,6)=-0.5

stress(13,1)=0.5
stress(13,2)=0.
stress(13,3)=-0.5
stress(13,4)=0.5
stress(13,5)=0.
stress(13,6)=0.5

stress(14,1)=0.5
stress(14,2)=0.
stress(14,3)=-.5
stress(14,4)=-0.5
stress(14,5)=0.
stress(14,6)=0.5

stress(15,1)=0.5
stress(15,2)=0.
stress(15,3)=-0.5
stress(15,4)=0.5
stress(15,5)=0.
stress(15,6)=-0.5

stress(16,1)=0.5
stress(16,2)=0.
stress(16,3)=-0.5
stress(16,4)=-0.5
```

stress(16,5)=0.  
stress(16,6)=-0.5

stress(17,1)=0.  
stress(17,2)=-0.5  
stress(17,3)=0.5  
stress(17,4)=0.  
stress(17,5)=0.5  
stress(17,6)=0.5

stress(18,1)=0.  
stress(18,2)=-0.5  
stress(18,3)=0.5  
stress(18,4)=0.  
stress(18,5)=-0.5  
stress(18,6)=0.5

stress(19,1)=0.  
stress(19,2)=-0.5  
stress(19,3)=0.5  
stress(19,4)=0.  
stress(19,5)=0.5  
stress(19,6)=-0.5

stress(20,1)=0.  
stress(20,2)=-0.5  
stress(20,3)=0.5  
stress(20,4)=0.  
stress(20,5)=-0.5  
stress(20,6)=-0.5

stress(21,1)=-0.5  
stress(21,2)=0.5  
stress(21,3)=0.  
stress(21,4)=0.5  
stress(21,5)=0.5  
stress(21,6)=0.

stress(22,1)=-0.5  
stress(22,2)=0.5  
stress(22,3)=0.  
stress(22,4)=-0.5  
stress(22,5)=0.5  
stress(22,6)=0.

stress(23,1)=-0.5  
stress(23,2)=0.5  
stress(23,3)=0.  
stress(23,4)=0.5  
stress(23,5)=-0.5  
stress(23,6)=0.

stress(24,1)=-0.5  
stress(24,2)=0.5  
stress(24,3)=0.  
stress(24,4)=-0.5  
stress(24,5)=-0.5  
stress(24,6)=0.

stress(25,1)=0.  
stress(25,2)=0.  
stress(25,3)=0.  
stress(25,4)=0.5  
stress(25,5)=0.5  
stress(25,6)=-0.5

```
stress(26,1)=0.
stress(26,2)=0.
stress(26,3)=0.
stress(26,4)=0.5
stress(26,5)=-0.5
stress(26,6)=0.5
```

```
stress(27,1)=0.
stress(27,2)=0.
stress(27,3)=0.
stress(27,4)=-0.5
stress(27,5)=0.5
stress(27,6)=0.5
```

```
stress(28,1)=0.
stress(28,2)=0.
stress(28,3)=0.
stress(28,4)=0.5
stress(28,5)=0.5
stress(28,6)=0.5
```

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+++++
```

```
program BishopHill
```

```
! gfortran -gdwarf-2 -O0 -o BishopHill Bishop_Hill.f90
! solve the Bishop Hill model, adr 28 iv 01
! updated for crystal orientation, 21 iv 05
! formatted to f90, x 11
```

```
implicit none
```

```
real scalar
```

```
real rtmp
```

```
real :: stress, strain, dw, dwmax
integer :: index, i, j, k, l
dimension stress(28,6), strain(6)
real :: a(3,3), hkl(3), uvw(3), t1(3)
real :: eps(3,3), eps_xtal(3,3)
real :: strain_xtal(6)
real :: rtmp1, rtmp2, tayf
real :: evm, enorm
real :: sqr3
real :: tayf2
integer :: iq
real :: phi1, Phi, phi2
real :: d1, d2, d3, atranspose(3,3)
integer :: iopt, ior, kerr
character :: nomen*1 = 'B'
```

```
! code::
```

```
sqr3 = sqrt(3.)
write(*,*) 'Welcome to the Reid version of the BH algorithm'
print *, 'Adapted to use the vM strain as the reference'
write(*,*) 'Enter the strain as de11, de22, 2*de23, 2*de31, 2*de12'
read(*,*) strain(1), strain(2), strain(4), strain(5), strain(6)
strain(3) = 0. - strain(1) - strain(2)
! incompressibility
```



```

write(*,*) 'Full strain vector: '
print"(2x,6(2x,f7.3))", (strain(i),i=1,6)

eps(1,1) = strain(1)
eps(2,2) = strain(2)
eps(3,3) = strain(3)
eps(2,3) = strain(4)/2.
eps(3,1) = strain(5)/2.
eps(1,2) = strain(6)/2.
eps(3,2) = strain(4)/2.
eps(1,3) = strain(5)/2.
eps(2,1) = strain(6)/2.
!   evm = sqrt((2./9.*((strain(1)-strain(2))**2 +
! 1  (strain(2)-strain(3))**2 + (strain(3)-strain(1))**2)) +
! 2  ((eps(1,2)**2 + eps(1,3)**2 + eps(2,3)**2)/3.))
evm = (2./sqrt(3))*sqrt(0.5*(eps(1,1)**2 + eps(2,2)**2 + eps(3,3)**2 &
+ eps(2,1)**2 + eps(3,1)**2 + eps(3,2)**2 + &
eps(1,2)**2 + eps(1,3)**2 + eps(2,3)**2))
print>(' eps vM = ',f7.3)",evm

write(*,*)

print * , 'Enter orientation as (hkl)[uvw], =0, or as 3 Bunge angles, =1?'
read * , iq

if ( iq == 0 ) then
write(*,*) 'Enter (hkl) // sample.3 as 3 numbers'
read(*,*) hkl(1),hkl(2),hkl(3)
write(*,*) ('Plane HKL = ',3(f8.1))" hkl
call vecnorm(hkl)
10 write(*,*) 'Enter [uvw] // sample.1 as 3 numbers'
read(*,*) uvw(1),uvw(2),uvw(3)
write(*,*) ('Plane UVW = ',3(f8.1))" uvw

call vecnorm(uvw)
rtmp=scalar(hkl,uvw)
if(abs(rtmp).gt.1e-4) then
write(*,*) 'Sorry, uvw is not perp. to hkl; try again'
goto 10
endif
call vecpro2(hkl,uvw,t1)
call vecnorm(t1)
do 20, i=1,3
a(i,1)=uvw(i)
a(i,2)=t1(i)
a(i,3)=hkl(i)
!   a(i,3)=t1(i)
!   a(i,2)=hkl(i)
20 enddo
! standard arrangement, not Canova matrix!
elseif ( iq == 1 ) then
print * , 'Enter three Bunge angles in degrees:'
read * , phi1 , Phi, phi2
iopt = 2
call euler(atranspose,iopt,nomen,phi1 , Phi, phi2,ior,kerr)
do i = 1,3
do j = 1,3
a(i,j) = atranspose(j,i)
end do
end do
else
print * , 'invalid choice, stopping ...'
stop

```

```

end if

print*,' Orientation Matrix: '
do i=1,3
  print"([' ,3(1x,f7.3),' ])", (a(i,j),j=1,3)
enddo

do i = 1,3
  do j = 1,3
    eps_xtal(i,j) = 0.
    do k=1,3
      do l = 1,3
        eps_xtal(i,j) = eps_xtal(i,j) + a(i,k)*a(j,l)*eps(k,l)
      enddo
    enddo
  enddo
enddo

print*,' Strain in Xtal coords: '
do i=1,3
  print"([' ,3(1x,f7.2),' ])", (eps_xtal(i,j),j=1,3)
enddo

evm = (2./sqrt(3))*sqrt(0.5*(eps_xtal(1,1)**2 + eps_xtal(2,2)**2 + &
  + eps_xtal(3,3)**2 + &
  eps_xtal(2,1)**2 + eps_xtal(3,1)**2 + eps_xtal(3,2)**2 + &
  eps_xtal(1,2)**2 + eps_xtal(1,3)**2 + eps_xtal(2,3)**2))
print"(' eps[xtal] vM = ',f7.3)",evm

enorm = sqrt((eps_xtal(1,1)**2 + eps_xtal(2,2)**2 + eps_xtal(3,3)**2 + &
  eps_xtal(2,1)**2 + eps_xtal(3,1)**2 + eps_xtal(3,2)**2 + &
  eps_xtal(1,2)**2 + eps_xtal(1,3)**2 + eps_xtal(2,3)**2))
!print"(' eps[xtal] vM = ',f7.3)",evm

strain_xtal(1) = eps_xtal(1,1)
strain_xtal(2) = eps_xtal(2,2)
strain_xtal(3) = eps_xtal(3,3)
strain_xtal(4) = eps_xtal(2,3) + eps_xtal(3,2)
strain_xtal(5) = eps_xtal(3,1) + eps_xtal(1,3)
strain_xtal(6) = eps_xtal(1,2) + eps_xtal(2,1)

print*
print*,'Strain in crystal axes: '
print"(2x,6(2x,f7.3))", (strain_xtal(i),i=1,6)

stress(1,1)=1.
stress(1,2)=-1.
stress(1,3)=0.
stress(1,4)=0.
stress(1,5)=0.
stress(1,6)=0.
! # 1
stress(2,1)=0.
stress(2,2)=1.
stress(2,3)=-1.
stress(2,4)=0.
stress(2,5)=0.
stress(2,6)=0.
! # 2
stress(3,1)=-1.
stress(3,2)=0.
stress(3,3)=1.

```

stress(3,4)=0.  
stress(3,5)=0.  
stress(3,6)=0.  
! # 3  
stress(4,1)=0.  
stress(4,2)=0.  
stress(4,3)=0.  
stress(4,4)=1.  
stress(4,5)=0.  
stress(4,6)=0.  
! # 4  
stress(5,1)=0.  
stress(5,2)=0.  
stress(5,3)=0.  
stress(5,4)=0.  
stress(5,5)=1.  
stress(5,6)=0.  
! # 5  
stress(6,1)=0.  
stress(6,2)=0.  
stress(6,3)=0.  
stress(6,4)=0.  
stress(6,5)=0.  
stress(6,6)=1.  
! # 6  
stress(7,1)=0.5  
stress(7,2)=-1.  
stress(7,3)=0.5  
stress(7,4)=0.  
stress(7,5)=0.5  
stress(7,6)=0.  
! # 7  
stress(8,1)=0.5  
stress(8,2)=-1.  
stress(8,3)=0.5  
stress(8,4)=0.  
stress(8,5)=-0.5  
stress(8,6)=0.  
! # 8  
stress(9,1)=-1.  
stress(9,2)=0.5  
stress(9,3)=0.5  
stress(9,4)=0.5  
stress(9,5)=0.  
stress(9,6)=0.  
! # 9  
stress(10,1)=-1.  
stress(10,2)=0.5  
stress(10,3)=0.5  
stress(10,4)=-0.5  
stress(10,5)=0.  
stress(10,6)=0.  
! # 10  
stress(11,1)=0.5  
stress(11,2)=0.5  
stress(11,3)=-1.  
stress(11,4)=0.  
stress(11,5)=0.  
stress(11,6)=0.5  
! # 11  
stress(12,1)=0.5  
stress(12,2)=0.5

stress(12,3)=-1.  
stress(12,4)=0.  
stress(12,5)=0.  
stress(12,6)=-0.5  
!  
stress(13,1)=0.5  
stress(13,2)=0.  
stress(13,3)=-0.5  
stress(13,4)=0.5  
stress(13,5)=0.  
stress(13,6)=0.5  
!  
stress(14,1)=0.5  
stress(14,2)=0.  
stress(14,3)=-.5  
stress(14,4)=-0.5  
stress(14,5)=0.  
stress(14,6)=0.5  
!  
stress(15,1)=0.5  
stress(15,2)=0.  
stress(15,3)=-0.5  
stress(15,4)=0.5  
stress(15,5)=0.  
stress(15,6)=-0.5  
!  
stress(16,1)=0.5  
stress(16,2)=0.  
stress(16,3)=-0.5  
stress(16,4)=-0.5  
stress(16,5)=0.  
stress(16,6)=-0.5  
!  
stress(17,1)=0.  
stress(17,2)=-0.5  
stress(17,3)=0.5  
stress(17,4)=0.  
stress(17,5)=0.5  
stress(17,6)=0.5  
!  
stress(18,1)=0.  
stress(18,2)=-0.5  
stress(18,3)=0.5  
stress(18,4)=0.  
stress(18,5)=-0.5  
stress(18,6)=0.5  
!  
stress(19,1)=0.  
stress(19,2)=-0.5  
stress(19,3)=0.5  
stress(19,4)=0.  
stress(19,5)=0.5  
stress(19,6)=-0.5  
!  
stress(20,1)=0.  
stress(20,2)=-0.5  
stress(20,3)=0.5  
stress(20,4)=0.  
stress(20,5)=-0.5  
stress(20,6)=-0.5  
!  
stress(21,1)=-0.5

```
stress(21,2)=0.5
stress(21,3)=0.
stress(21,4)=0.5
stress(21,5)=0.5
stress(21,6)=0.
!
stress(22,1)=-0.5
stress(22,2)=0.5
stress(22,3)=0.
stress(22,4)=-0.5
stress(22,5)=0.5
stress(22,6)=0.
!
stress(23,1)=-0.5
stress(23,2)=0.5
stress(23,3)=0.
stress(23,4)=0.5
stress(23,5)=-0.5
stress(23,6)=0.
!
stress(24,1)=-0.5
stress(24,2)=0.5
stress(24,3)=0.
stress(24,4)=-0.5
stress(24,5)=-0.5
stress(24,6)=0.
!
stress(25,1)=0.
stress(25,2)=0.
stress(25,3)=0.
stress(25,4)=0.5
stress(25,5)=0.5
stress(25,6)=-0.5
!
stress(26,1)=0.
stress(26,2)=0.
stress(26,3)=0.
stress(26,4)=0.5
stress(26,5)=-0.5
stress(26,6)=0.5
!
stress(27,1)=0.
stress(27,2)=0.
stress(27,3)=0.
stress(27,4)=-0.5
stress(27,5)=0.5
stress(27,6)=0.5
!
stress(28,1)=0.
stress(28,2)=0.
stress(28,3)=0.
stress(28,4)=0.5
stress(28,5)=0.5
stress(28,6)=0.5
!
dwmax=0.
index=0
do 100, i=1,28
  dw = 0.
  dw = dw - strain_xtal(1)*stress(i,2)
  dw = dw + strain_xtal(2)*stress(i,1)
  dw = dw + strain_xtal(4)*stress(i,4)
```

```

dw = dw + strain_xtal(5)*stress(i,5)
dw = dw + strain_xtal(6)*stress(i,6)
!   dW = -Bde11 + Ade22 + 2Fde23 + 2Gde31 + 2Hde12
if(dw.gt.dwmax) then
  dwmax=dw
  index=i
  print*,'dw,i ',dw,' ',i
endif
if((-1.*dw).gt.dwmax) then
  dwmax=-1.*dw
  index=i+28
  print*,'dw,i ',dw,' ',i
endif
100 end do

print*
write(*,*) 'Index of the multiple slip stress state = ',index
if(index.gt.28) then
  index = index-28
  write(*,*) 'Index of the +/- stress state = ',index
endif
print*,'Stress #  A   B   C   F   G   H'
write(*,120) (stress(index,j),j=1,6)
120 format(6x,6(2x,f6.1))

rtmp1 = 0.
do i = 1,6
  rtmp1 = rtmp1 + stress(index,i)**2
enddo
! magnitude of the stress
rtmp2 = 0.
do i = 1,6
  rtmp2 = rtmp2 + strain_xtal(i)**2
enddo
! magnitude of the strain in xtal coords.
!   print*,'dwmax,rtmp1,rtmp2',dwmax,sqrt(rtmp1),sqrt(rtmp2)

!   tayf = 2.* dwmax / sqrt(rtmp2)
tayf = dwmax / evm
! This is the definition of Taylor factor that allows for multi-axial
! strain or stress states, as found in LApp

print*
print*('The Taylor factor = ',f7.3,' *sqrt(6)'),tayf
print*('          = ',f7.3)',tayf*sqrt(6.)

! these next few lines were for curiosity about other normalizations of strain
!   tayf2 = dwmax / enorm

!   print*
!   print*('The Taylor factor = ',f7.3,' *sqrt(6)'),tayf2
!   print*('          = ',f7.3)',tayf2*sqrt(6.)

call exit(0)
end program BishopHill

!
! -----
!
subroutine euler(a,iopt,nomen,d1,d2,d3,ior,kerr)
!           Last revision 20nov90 UFK

```

```

! common a(3,3),grvol(1152),epsga(5),ist1,ist2,sqr3,sqrh,ident(3,3)
! SPECIAL VERSION WITHOUT COMMON BLOCK
real a(3,3),d1,d2,d3,th,sth,cth,sph,cph,sps,cps,ps,ph,dth,dph,dps
character nomen
save kor
real pi
parameter ( pi = 3.14159265 )
real rad
parameter ( rad = 57.29578 )

! CODE::

! *** this subroutine calculates the euler angles associated with the
! transformation matrix a(i,j) if iopt=1 and viceversa if iopt=2
! *** Note that a is sample (rows) in terms of crystal (columns);
! -- opposite of standard g (e.g.Bunge) - this is Canova's
! *** Note that in this version, the Euler angles are defined symmetrically:
! so that interchanging phi and psi means transposing a.
! ("Kocks" nomen: defined going from +X to +Y in both COD and SOD)
! *** However, other angle conventions are translated, according to
! nomen="K" - Kocks (as internally) -- also sometimes "N"...
! "R" - Roe      (Psi=psi, Phi=180-phi)
! "B" - Bunge    (phi1=90+psi, Phi=Theta, phi2=90-phi)
! any other - Canova (Theta first, phiC=90+phi, omega=90-psi)
! *** Note: only in symm. notation does a point with all Euler angles
! between 0 and 90 deg appear in the +x,+y quadrant!
! If you want to see an individual point in this quadrant and are:
! using Roe, the third angle must be between 90 and 180;
! Bunge, first ;
! Canova, second .
! *** Input and output Euler angles d1,d2,d3 in degrees
!
goto(5,20),iopt
5 if(abs(a(3,3)) .ge. 0.999) goto 10
th=acos(a(3,3))
sth=sin(th)
!
if((abs(a(2,3)/sth).lt.1e-35).and. &
& (abs(a(1,3)/sth).lt.1e-35)) then
ps=pi/4.
else
ps=atan2(a(2,3)/sth,a(1,3)/sth)
endif
!
if((abs(a(3,2)/sth).lt.1e-35).and. &
& (abs(a(3,1)/sth).lt.1e-35)) then
ph=pi/4.
else
ph=atan2(a(3,2)/sth,a(3,1)/sth)
endif
!cccc if it bombs out here, both a-components in one arg. are zero
! (this should not be possible, but has happened, probably fixed)
! ADR: i 01 - above is the fix!!
!
go to 15
!
10 if((abs(a(1,2)/sth).lt.1e-35).and. &
& (abs(a(1,1)/sth).lt.1e-35)) then
ps=pi/4.
else
ps=0.5*atan2(a(1,2),-a(1,1))
endif

```

```

!
ph=-ps
! The above still have the problem that they give too many
! equivalents of the same grain in DIOROUT and density file. Therefore:
if(kerr.eq.1.and.kor.ne.ior) then
  print*,NOTE: grain',ior,' has Theta < 1 deg.: sometimes problems'
  print*
  kor=ior
endif
!
15 dth=th*rad
dph=ph*rad
dps=ps*rad
d1=dps
d2=dth
if(nomen.eq.'K'.or.nomen.eq.'k'.or.nomen.eq.'N') then
  d3=dph
elseif(nomen.eq.'R'.or.nomen.eq.'r') then
  d3=180.-dph
elseif(nomen.eq.'B'.or.nomen.eq.'b') then
  d1=dps+90.
  d3=90.-dph
else
  d1=dth
  d2=dph+90.
  d3=90.-dps
endif
if(d1.ge.360.) d1=d1-360.
if(d3.ge.360.) d3=d3-360.
if(d1.lt.0.) d1=d1+360.
if(d3.lt.0.) d3=d3+360.
return
! *****
20 dth=d2
dps=d1
if(nomen.eq.'K'.or.nomen.eq.'k') then
  dph=d3
elseif(nomen.eq.'R'.or.nomen.eq.'r') then
  dph=180.-d3
elseif(nomen.eq.'B'.or.nomen.eq.'b') then
  dps=d1-90.
  dph=90.-d3
else
  dth=d1
  dph=d2-90.
  dps=90.-d3
endif
ph=dph/rad
th=dth/rad
ps=dps/rad
sph=sin(ph)
cph=cos(ph)
sth=sin(th)
cth=cos(th)
sps=sin(ps)
cps=cos(ps)
a(1,1)=-sps*sph-cph*cps*cth
a(2,1)=cps*sph-cph*sps*cth
a(3,1)=cph*sth
a(1,2)=sps*cph-sph*cps*cth
a(2,2)=-cph*cps-sph*sps*cth
a(3,2)=sth*sph

```



```
a(1,3)=sth*cps
a(2,3)=sps*sth
a(3,3)=cth
return
end subroutine euler
```

```
!
```

---

```
subroutine vecnorm(vec)
real vec(3),rnorm
rnorm=0.
do 10, i=1,3
  rnorm=rnorm+vec(i)**2
10 end do
if(rnorm.le.0.0) return
rnorm=sqrt(rnorm)
do 20, i=1,3
  vec(i)=vec(i)/rnorm
20 end do
return
end subroutine vecnorm
```

```
! -----
```

```
function scalar(a,b)
! return SCALAR PRODUCT of A and B
real scalar,a(3),b(3)
scalar=0.
do 100, i=1,3
  scalar=scalar+a(i)*b(i)
100 end do
return
end function scalar
```

```
!
```

---

```
subroutine vecpro2(v1,v2,vout)
! vector product
real v1(3),v2(3),vout(3)
vout(3)=v1(1)*v2(2)-v1(2)*v2(1)
vout(1)=v1(2)*v2(3)-v1(3)*v2(2)
vout(2)=v1(3)*v2(1)-v1(1)*v2(3)
return
end subroutine vecpro2
```