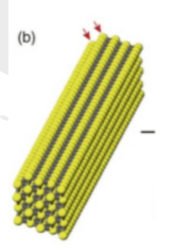
Au Nanowires

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Define Project

Project option 1



Create a bulk gold model, carve out nanowires from the bulk along different crystal directions (e.g. <111>, <100>, <110> etc.

- Compare the energetics of these nanowires after energy minimization.
- Test the elastic properties of these nanowires and try to understand the phenomenon.

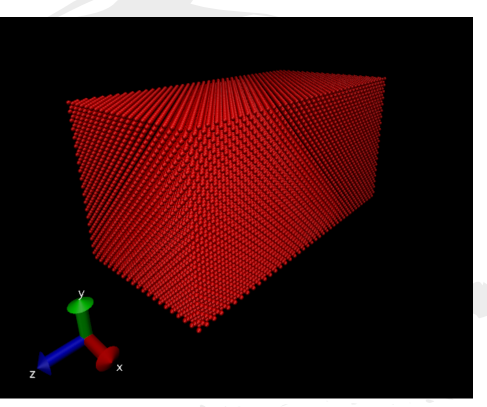
Creating the Bulk

```
# from slides, change later to correct values for Au
#lattice custom 3.2 &
# a1 -0.5 0.5 0.5 &
# a2 0.5 -0.5 0.5 &
# a3 0.5 0.5 -0.5 &
# basis 0.0 0.0 0.0

lattice fcc 4.080

region myRegion block 0 20 0 20 0 40 units lattice
# create a box object from the above defined region create_box 1 myRegion create_atoms 1 box basis 1 1
mass 1 196.967
```

- Created brick of 64,000 atoms
- Energy -251,520eV



Creating 001 NW

```
A = np.loadtxt('
Alength = len(A)
x = A[0:Alength,0] # first column of data
y = A[0:Alength,1] # Second column of data
z = A[0:Alength, 2] # third column of data
#Reshape with x=x.reshape(length,1)
x=x.reshape(Alength,1)
y=y.reshape(Alength,1)
z=z.reshape(Alength,1)
#Set Radius of NW r
 ########### Nanowire <001> ###########
indices2remove=[]
for k in range(Alength):
   if (x[k]-30)**2 + (y[k]-30)**2 > r**2:
       #remove those elements
       storing indices to an array
       lind2rm=len(indices2remove)
       indices2remove[(1+lind2rm):]=[k]
```



```
######## Nanowire <<mark>1</mark>11> ###########
def drange2(start, stop, step):
    numelements = int((stop-start)/float(step))
    for i in range(numelements+1):
            yield start + i*step
for k in range(len(xout)):
    for j in range(k+1,len(xout)):
        if A[k,2] > A[j,2]:
            temp=np.copy(A[j,:])
            A[j,:]=A[k,:]
            A[k,:]=temp
                                                              #NW 2:
indices2remove=[]
zval=[r for r in A[:,2] ] # makes 1D list of zvalues from
C=
for k in range(c.len(zval)):
    for j in range(k+1,len(zval)):
        if zval[k] == zval[j]:
            lindex=len(indices2remove)
            indices2remove[(1+lindex):]=[j]
    zval=np.delete(zval,indices2remove)
    C+=1
```

indices2remove=[]

for k in range(Alength):

if z[k] == lava:

for t in drange2(11,57,161.0/57):

if (x[k]-t)**2 + (y[k]-t)**2 > r**2:

print zval

for lava in zval:

```
Creating 011 & 111 NWs
```

```
#region LLG cylinder y 85 85 10 INF INF units box
#lattice fcc 4.08 orient x 1 0 0 orient y 0 1 -1 orient z 0 1 1
#create atoms 1 region LLG
```

Energy of 001 & 011

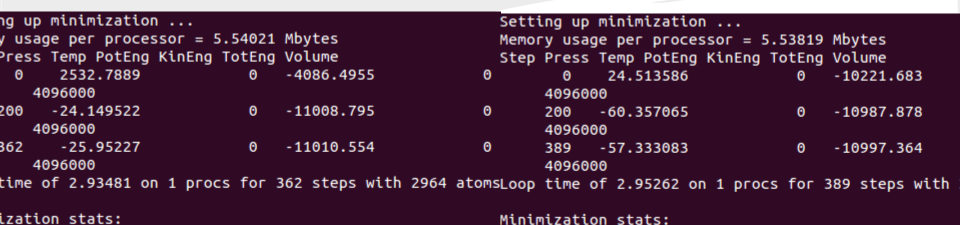
~14eV difference

oping criterion = energy tolerance

rgy initial, next-to-last, final =

-11010.5542193

-4086.49553199



-11010.5543124

Stopping criterion = energy tolerance

Energy initial, next-to-last, final =

-10997.3640976

-109

-10221.6825458

Compression

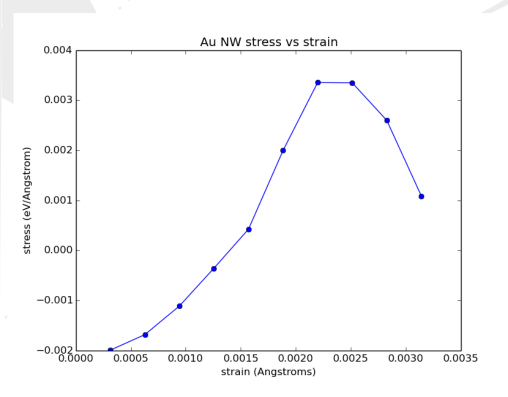
```
variable Nstep loop 10  # reapet 5 times of compression.
label looplable
# -0.05
displace_atoms top move 0 0 -0.09 units box  # compress 0.05 Ansompression step
#displace_atoms bottom move 0 0 0.09 units box  # compress 0.05 //
compression step
fix 1 3 nvt temp 300.0 300.0 1.0
run 2000
next Nstep
jump in.nwcompress001.txt looplable
```

- 1) https://www.youtube.com/watch? v=jE8En6FZjQU&feature=youtu.be
- 2) https://www.youtube.com/watch? v=tzJn2lCvGKQ&feature=youtu.be
- 3) https://www.youtube.com/watch?v=7jjGF-hXtms&feature=youtu.be
- 4) https://www.youtube.com/watch? v=liWQjjuVCx4&feature=youtu.be

Video Links

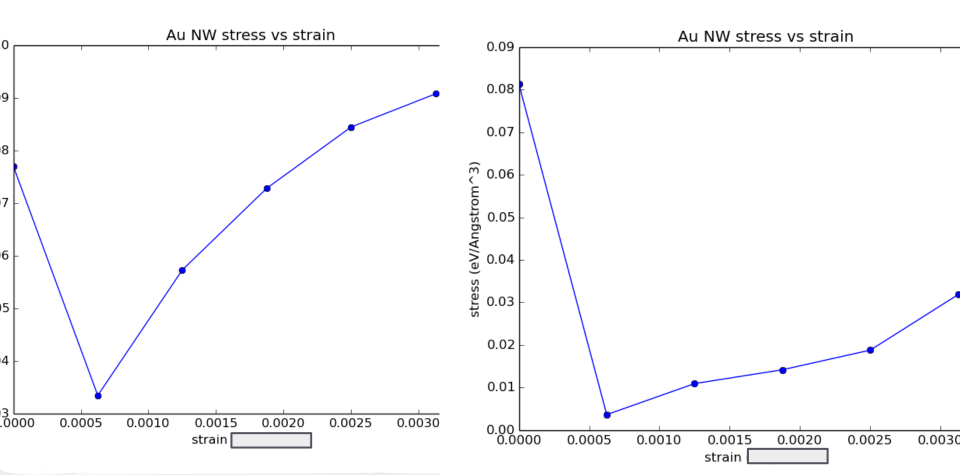


Young's Modulus



```
time_of_loop_in_inputfile=
 # 1st NW
a=np.loadtxt('outputforce.txt')
|b=[]
for k in range(len(a)):
    if a[k,0]%time_of_loop_in_inputfile ==
        b.append(a[k,:])
B=np.array(b)
orint B
del b
Γ=
area=np.pi*r**
#calculate stress
stress=B[:,1]/(np.pi*r**2)
eprate=abs(-0.05)
deltaL=eprate/time of loop in inputfile
L=
strain=B[:,0]*deltaL/L
```

Moduli r=10,ed=0.1, 001(left) & 011(right)

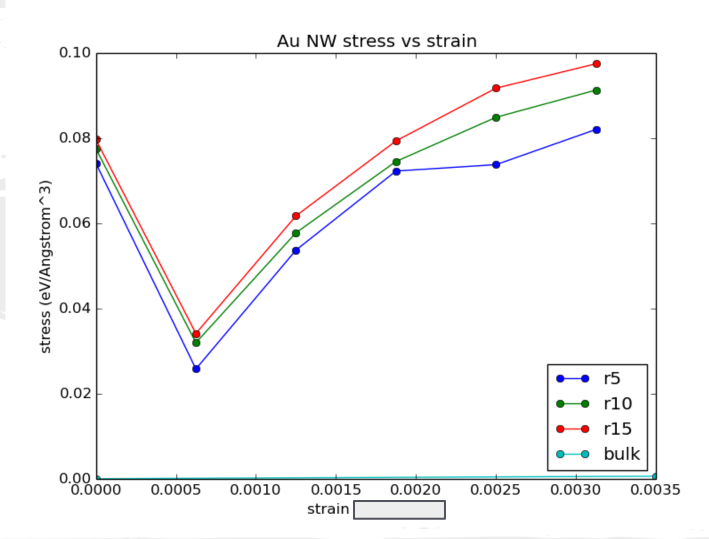


Initial Results in eV/Angstrom^3

```
e4=(stress[4]-stress[1])/(strain[4]-strain[1]) = 2.2388042
e6=(stress[6]-stress[4])/(strain[6]-strain[4]) = 4.6673040
enwavg=3.4530541
ebulk = 0.1715986331
```

- Bulk Au has E=29GPa or 0.172eV/A³
- NW Au has E=583.5GPa
- Ratio Enwavg/Ebulk = 20.12850
- Next steps
 - Compare with a different radii NWs
 - Test orientations <011>, <111>

Comparing
Youngs
Modulus
for different
size radii



Extras

- 1) https://www.youtube.com/watch?
 v=oSw3CaI2u6U&feature=youtu.
 be
- 2) https://www.youtube.com/watch?
 v=dpQ3-waEUT4&feature=youtu.
 be