# Nano 266 Quantum Mechanical Modeling of Materials Lab 3

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# 1 Question 1

The BCC-HCP transition in iron

## 1.1 Ground State Energy

Ground state energy in BCC Structure: Using energy cutoff of 50 Ry, we do a ground state energy calculation of Fe with bcc structure with K-point setting range [7-13] and lattice parameter from 5.38 to 5.46 (Bohr), from the convergence test as showed in the Figure 1, indicate with K-point set to 11 (K-point number 112), the energy converge to within 1 meV/atom.

### Energy Convergence VS K-point (BCC)

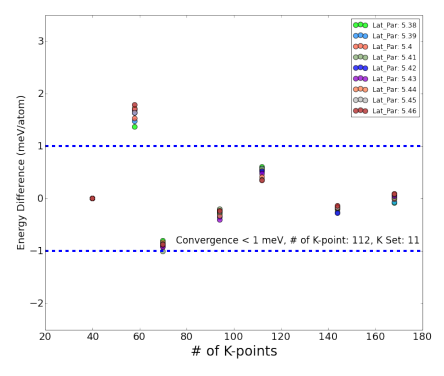


Figure 1: K point convergence test in BCC Fe

Ground state energy in HCP Structure: Using energy cutoff of 50 Ry, we do a ground state energy calculation of Fe in HCP structure with K-point from 7 to 13, and lattice parameter from 4.75 to 4.84 (Bohr), from the convergence test as showd in the Figure 2, Figure 3 and Figure 4, with

K-point setting increases to above 9 (Unique K-point number 72), the energy converge to within 1 meV/atom.

#### Energy Convergence VS K-point (c/a ratio: 1.72)

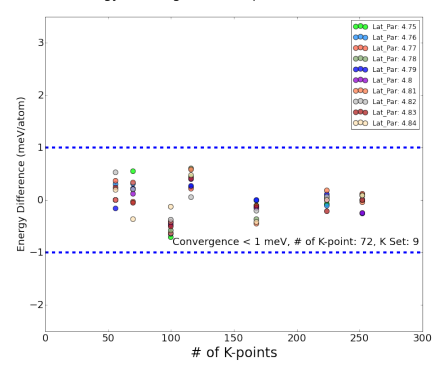


Figure 2: K point convergence test in HCP Fe, c/a ratio: 1.72

## Energy Convergence VS K-point (c/a ratio: 1.73)

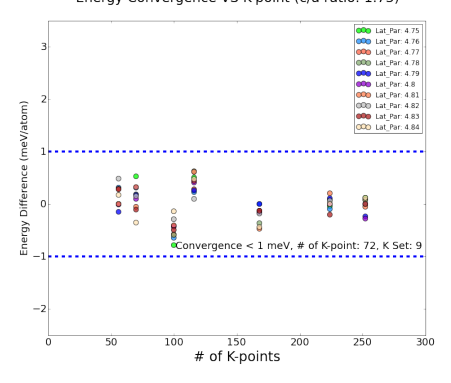


Figure 3: K point convergence test in HCP Fe, c/a ratio: 1.73

Lattice parameter optimization for BCC Using energy cutoff of 50 Ry, K-point set to 11, we do a

## 3 Lat\_Par: 4.75 风 Lat\_Par: 4.76 Lat\_Par: 4.77 000 Lat\_Par: 4.78 Lat\_Par: 4.79 2 Lat Par: 4.8 C Lat Par: 4.81 Energy Difference (meV/atom) 000 Lat\_Par: 4.82 Lat\_Par: 4.83 0 Convergence < 1 meV, # of K-point: 72, K Set: 9 Ō 50 100 150 200 250 300 # of K-points

#### Energy Convergence VS K-point (c/a ratio: 1.74)

Figure 4: K point convergence test in HCP Fe, c/a ratio: 1.74

lattice parameter optimization of BCC structure with lattice parameter between 5.2 to 5.55 with higher sampling point density from 5.3 to 5.4. From the energy vs lattice relationship as showed in the Figure 5, when lattice parameter equals to 5.37 a.u., we reached the lowest energy per atom.

### 1.2 Energy

 $H_2$  **DFT Energy** The total DFT energy of  $H_2$  is -1.176631 Ha, -32.01778 eV.

## 2 Question 2

Geometry optimization and energy of  $N_2$ 

### 2.1 Geometry

 $H_2$  Bond Length From the geometry data, the x, y coordinates of both hydrogen atoms are zero, the Z coordinates are -0.55 and 0.55 Scaling the coordinates by 1.889725 to convert to a.u., the final bond length of  $H_2$  is 1.1 Å.

### 2.2 Energy

 $N_2$  **DFT Energy** The total DFT energy of  $N_2$  is -109.502491 Ha, -2979.71597 eV.

# 3 Question 3

Geometry optimization and energy of  $NH_3$ 

#### 3.1 Non Polarized

 $NH_3$  Bond Length and Bond Angle From the geometry data, the x, y, z coordinates of both hydrogen atoms and N atoms, the bond length between N and H is 1.005 91 Scaling the coordinates by

# **Energy VS Lattice Constant (BCC)**

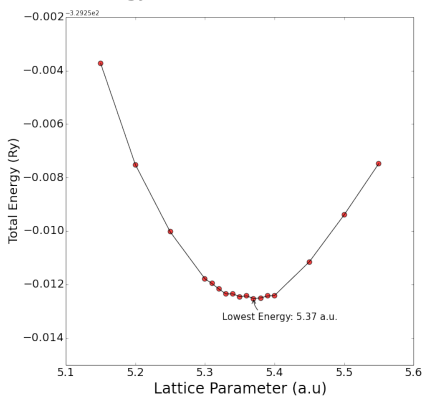


Figure 5: Lattice parameter optimization for BCC

1.889725 to convert to a.u., the final N-H bond length of is 1.9009 Å. The bond angle is 116.18°.

 $NH_3$  Total DFT Energy Total DFT energy of  $NH_3$  is  $-56.55168\,\mathrm{Ha}$ ,  $-1538.850329\,\mathrm{eV}$ .

#### 3.2 Polarized

 $NH_3$  Bond Length and Bond Angle From the geometry data, the x, y, z coordinates of both hydrogen atoms and N atoms, the bond length between N and H is 1.018 01 Scaling the coordinates by 1.889 725 to convert to a.u., the final N-H bond length of is 1.018 Å. The bond angle is 107.68°.

 $NH_3$  Total DFT Energy Total DFT energy of  $NH_3$  is  $-56.55168\,\mathrm{Ha}$ ,  $-1538.850329\,\mathrm{eV}$ .

# 4 Formation enthalpy of $NH_3$

To get the right formation enethalpy of  $NH_3$ , we first reset the basis set of  $H_2$  and  $N_2$  in Q1 and Q2 with polarization functions. After rerun  $H_2$  and  $N_2$  calculations, we get the thermal correction and DFT energy of each parts as table show below.

Compound	Energy (Ha)	Correction (kcal/mol)	Enthalpy H (kcal/mol)
$H_2$	-1.17663	8.436	-729.903782
$N_2$	-109.56056	5.580	-68744.00271
$NH_3$	-56.57363	24.042	-35476.07972

Given by the formula:  $0.5N_2 + 1.5H_2 > NH_3$ , the calculated formation enthalpy is -38.59 kJ/mol.NIST (2015), which is not far from the experimental data of NIST source: http://cccbdb.nist.gov/, -45.95 kJ/mol.

### 5 Effect of Functional Choice and Basis Set

In this section, we enumerated all 36 differenct conditions with different functional choice and basis set. Only basis set with polarization function are used. We use HF, PBE and B3LYP(B3) funtionals with 6-31G\* and 6-311G\* basis set, for each compound calculation, there are two stages, i.e. Geometry optimization stage and DFT energy stage. The experimental formation enthalpy for  $NH_3$  is  $-45.95 \pm 0.35kJ/mol$  from NIST website.NIST (2015). The bond length of N-H is 1.017Å based on Wiki (2015) data. The bond angle is 107.8° based on wiki Wiki (2015) data also.

#### 5.1 Overview

From the Figure ??, we could find that with difference functional set, the deviation of formation enthalpy various a lot. But the combination of different functional set provides us with enough data point that fall in the accepted deviation band, the band width is  $\pm 2\%$ . The best functional set combination is using PBE function for both stages calcuation, using 6-31+G\* as basis set for geometry stage optimization and using 6-311+G\* for DFT energy calculation with calculated formation energy -46.18 kJ/mol, with only 0.5% deviation from experimental value.

#### 5.2 B3LYP Function and Effect

If we use B3LYP functional as first stage function, to minimize the deviation from experimental data, we should keep using B3LYP as second stage function from Figure ??.

#### 5.3 HF Function and Effect

If we use HFexch functional as first stage function, to minimize the deviation from experimental data, B3LYP should still be considered as second stage function Figure ??.

#### 5.4 PBE Function and Effect

If we use PBE functional as first stage function, to minimize the deviation from experimental data, B3LYP should still be considered as second stage function Figure ??.

#### 5.5 Function and Geometry Angle Prediction

Using different functional, the N-H bond predicted are not much different and all pretty close to the data we found from online source wiki Wiki (2015), as show in Figure ??. The blue horizontal line is the experimental angle data we found from wiki.

#### 5.6 Function and Bond Length Prediction

Using different functional, the N-H bond length predicted are not much different and all pretty close to the data we found from online source wiki Wiki (2015), as show in Figure ??. The blue horizontal line is the experimental angle data we found from wiki. The B3LYP functional use 6-31+G\* (optimization stage) and 6-311+G\* (energy stage) gives the best geometry prediction with highest accuracy.

#### 5.7 CPU wall time and effectiveness

If we could accept 2% as deviation, from Figure  $\ref{eq:condition}$ , using HF functional for geometry optimization and PBE for DFT energy calculation might be the most efficient calculation strategy with relatively good accuracy within 2%.

#### 5.8 Basis set effect

In this section, we will investigate the effect of 6-31+G\* and 6-311+G\* basis set in second stage (DFT energy calculation) calculation, from the below bar chart ??, using 6-311+G\* as second stage function, the predicted formation energy is more close to experimental value with decreased standard deviation also.

Row content	Formation Enth	nalpy(kJ/mol)	
$6-31+G^*$ mean value	-39.640336		
6-31+G* standard deviation	6.651545		
$6-311+G^*$ mean value	-38.815557		
6-311+G* standard deviation	5.669244		
Row content	Wall Time (s)	N-H Angle	N-H Length
$6-31+G^*$ mean value	26.588889	107.623333	1.015263
6-31+G* standard deviation	13.591214	0.456645	0.010470
$6-311+G^*$ mean value	26.588889	107.623333	1.015263
6-311+G* standard deviation	13.282392	0.456645	0.010470

# 6 Question 6

For the reaction to happen, we need to break N-N triple bond, the atomic energy of each N is -54.6 Ha. Compared with energy of N2, -109.53 Ha, the dissociation energy of N2 is 0.33 Ha = 8.98 ev, 207.08 kcal/mol. This is also the reaction barrier of reaction.

# References

NIST (2015). Ammonia thermochemistry data.

Wiki (2015). Wiki ammonia thermochemistry data.