PHASEGO: A toolkit for automatic calculation and plot of phase diagram

Zhong-Li Liu\textsuperscript{a,*}

\textsuperscript{a}College of Physics and Electric Information, Luoyang Normal University, Luoyang 471022, China

Abstract

The PHASEGO package extracts the Helmholtz free energy from the phonon density of states obtained by the first-principles calculations. With the help of equation of states fitting, it reduces the Gibbs free energy as a function of pressure/temperature at fixed temperature/pressure. Based on the quasi-harmonic approximation (QHA), it calculates the possible phase boundaries among all the structures of interest and finally plots the phase diagram automatically. For the single phase analysis, PHASEGO can numerically derive many properties, such as the thermal expansion coefficients, the bulk moduli, the heat capacities, the thermal pressures, the Hugoniot pressure-volume-temperature relations, the Grüneisen parameters, and the Debye temperatures. In order to check its ability of phase transition analysis, I present here two examples: semiconductor GaN and metallic Fe. In the case of GaN, PHASEGO automatically determined and plotted the phase boundaries among the provided zinc blende (ZB), wurtzite (WZ) and rocksalt (RS) structures. In the case of Fe, the results indicate that at high temperature the...
electronic thermal excitation free energy corrections considerably alter the phase boundaries among the body-centered cubic (bcc), face-centered cubic (fcc) and hexagonal close-packed (hcp) structures.

**Keywords:** Quasi-harmonic approximation; Gibbs free energy; Phase diagram; Thermodynamic properties

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**Program summary**

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*Authors:* Zhong-Li Liu  
*Program Title:* PHASEGO  
*Journal Reference:*  
*Catalogue identifier:*  
*Licensing provisions:* GNU GPL version 3  
*Programming language:* Python (versions 2.4 and later)  
*Computer:* Any computer that can run Python (versions 2.4 and later)  
*Operating system:* Any operating system that can run Python  
*RAM:* 10 M bytes  
*Number of processors used:*  
*Supplementary material:*  
*Keywords:* Quasi-harmonic approximation, Gibbs free energy, Phase diagram, Thermodynamic properties.  
*Classification:* 7.8 Structure and Lattice Dynamics  
*External libraries:* Numpy [1], Scipy [2], Matplotlib [3]  
*Subprograms used:*  
*Nature of problem:* Materials usually undergo structural phase transitions when the environmental pressure and temperature are elevated to enough high values.
The phase transition process obeys the principle of lowest Gibbs free energy. In addition to the static energy, current density functional theory (DFT) calculations can easily give the phonon density of states of lattice vibrations, from which the Helmholtz free energy of phonons are reduced. Then Gibbs free energy can be achieved for the analysis of phase stability and phase transition at high pressure and temperature within the framework of QHA. The problem is to extract the Gibbs free energies from the DFT calculations and automatically analyze the high pressure and temperature phase boundaries among a number of structures.

Solution method: With the help of numerical interpolation techniques, the Gibbs free energy as a function of pressure/temperature at fixed temperature/pressure can be obtained. Then the QHA based phase boundaries can be automatically determined and plotted by scanning the pressure/temperature at fixed temperature/pressure according to the principle of lowest Gibbs free energy.

Restrictions: The restriction is from the QHA which takes partially into account the anharmonic effects.

Unusual features: The phase boundaries among a number of structures can be automatically determined and plotted, which largely improves the efficiency of phase transition analysis. In addition to some basic thermodynamic properties of each single structure, the Hugoniot pressure-volume-temperature relations are also automatically reduced.

Additional comments: This package can treat the phonon density of states data from many packages, such as PHON [4], PHONOPY [5], Quantum ESPRESSO [6], and ABINIT [7].
Running time: The examples provided in the distribution take less than a minute to run.

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1. Introduction

The accurate determinations of phase diagrams, equations of state (EOS), and thermodynamic properties of materials are of primary importance to the materials science, the high pressure science, and the geophysics. The high-pressure experimental techniques, including the diamond-anvil cell (DAC) and shock wave (SW) experiments, are main ways to yield such properties precisely. However, except for high costs there are more or less limitations in these experimental techniques, such as pressure or temperature limit and the very short duration time of the compression process in SW experiment during which it is hard to detect phase transitions in real time. So it very urgently needed that accurate theoretical methods are developed to expand the pressure and temperature regime where experiments can not reach.

Current density of functional theory (DFT) has been developed to a powerful tool widely used to calculate and predict all kinds of new properties of materials, and even design new materials theoretically before experimentation [1, 2]. Especially, the DFT has been applied frequently to calculate the high pressure and high temperature (HPHT) dependent properties including the equation of states, thermodynamic properties and phase diagrams under high pressure [3, 4, 5, 6]. As for the HPHT dependent properties, the hydrostatic pressure changes subject to the volume changes of the crystal cell provided that the atoms positions are fully relaxed therein. And the inclusion of temperature effects can be realized either by the quasi-harmonic
approximations (QHA) or molecular dynamics simulations. The \textit{ab intio} molecular dynamics simulations are computationally expensive. While the QHA is relatively cheap and a good solution to count the temperature effects in materials simulations. Nevertheless, one has to gather the necessary information to extract Gibbs free energies for all volumes considered before the calculations of HPHT properties. And then the related properties and even the phase diagrams can be calculated by numerical techniques. This whole process is complex and time-consuming to finish manually. Especially in the construction of phase diagrams for many structures (more than two), the comparison of their Gibbs free energies and the judgment of stable field for each structure are tough work. This motivates us to design a package named \texttt{Phasego} to accomplish all the complicated steps and plot the phase diagrams automatically.

\texttt{Phasego} is designed for the easy implementation of phase transition analysis and plot of phase diagrams. It can also calculate the thermodynamic properties of materials, including the thermal expansion coefficients, the bulk moduli, the heat capacities, the thermal pressures, the Gr"{u}neisen parameters, and the Debye temperatures. For the dynamic response properties of materials, \texttt{Phasego} can automatically find the Hugoniot pressure-volume and pressure-temperature relations according to the Rankine-Hugoniot conditions \cite{7, 8}. All these qualities can be obtained based on the QHA by simply preparing a controlling file. More interestingly, based on the QHA the possible phase boundaries of all the structures provided are analyzed and plotted automatically. So the high pressure and temperature phase diagram can be constructed and plotted easily by \texttt{Phasego}.
2. Theoretical background

2.1. Quasi harmonic approximation

The QHA is a phonon-based model of crystal lattice vibrations used to describe volume-dependent thermal effects, such as the thermal expansion. It starts from the assumption that the harmonic approximation holds for every value of the crystal volume, and then takes into account part of anharmonic effects by varying crystal volume.

In the framework of QHA, the Helmholtz free energy of a crystal is written as

\[ F(V, T) = E_{\text{static}}(V) + F_{\text{zp}}(V, T) + F_{\text{ph}}(V, T), \]  

(1)

where \( E_{\text{static}}(V) \) is the first-principles zero-temperature energy of a static lattice at volume \( V \), and \( F_{\text{zp}} \) is the zero-point motion energy of the lattice given by

\[ F_{\text{zp}} = \frac{1}{2} \sum_{q, j} \hbar \omega_j(q, V). \]  

(2)

The last term is the phonon free energy due to lattice vibrations, and can be obtained from

\[ F_{\text{ph}}(V, T) = k_B T \sum_{q, j} \ln \{1 - \exp \left(-\hbar \omega_j(q, V)/k_B T \right)\}, \]  

(3)

where \( \omega_j(q, V) \) is the phonon frequency of the \( j \)th mode of wavevector \( q \) in the first Brillouin zone.

For a metallic material, the Helmholtz free energy at high temperature includes the electronic thermal excitation free energy,

\[ F(V, T) = E_{\text{static}}(V) + F_{\text{el}}(V, T) + F_{\text{zp}}(V, T) + F_{\text{ph}}(V, T), \]  

(4)
$F_{el}$ can be evaluated by the finite temperature DFT with the help of the Fermi-Dirac smearing [9]. At low temperature, $F_{el}$ is very small and can be neglected, but at high temperature it is large and probably changes final conclusions.

If we write $F_{latt}$ as

$$F_{latt} = F_{zp} + F_{ph},$$

it can also be calculated from the phonon density of states $g(\omega)$ via [10]

$$F_{latt} = \int \left[ \frac{1}{2} \hbar \omega + k_B T \ln(2\sinh \frac{\hbar \omega}{2k_B T}) \right] d\omega g(\omega),$$

where $k_B$ is the Boltzmann constants. The phonon density of states is written as

$$g_j(\omega) = \frac{V}{(2\pi)^3} \int dq \delta[\omega - \omega_j(q)].$$

And the total densities of states are normalized to,

$$\int g(\omega)d\omega = 3nN,$$

where $n$ is the number of atoms in the unit cell and $N$ is the number of the unit cells.

2.2. Fitting equation of state

For a crystal structure, when obtained the Helmholtz free energies at different volumes and fixed temperature, one can derive the analytical function of $F(V)$ at certain temperature by fitting EOS. The types of the EOS in Phasego package are: Murnaghan [11], Birch [12], 3rd-order Birch-Murnaghan [13], 4rd-order Birch-Murnaghan [13], Vinet [14], Vinet Universal [15], 3rd-order Natural strain [16], 4rd-order Natural strain [16], and the
3rd and 4rd Polynomial. Then the pressure $P$ is obtained by

$$P = -\left(\frac{\partial F}{\partial V}\right)_T.$$  \hfill (9)

One can calculate Gibbs free energy $G$ as a function of temperature ($T$) and pressure ($P$) via,

$$G(T, P) = F(V, T) + PV.$$ \hfill (10)

Once the Gibbs free energy is known, the phase stabilities and transitions can be determined by comparing the Gibbs free energies of different structures at fixed pressure or temperature.

### 2.3. The derivation of thermodynamic properties

Other thermodynamic properties can be reduced numerically according to the thermodynamics relations [17]. The thermal pressure $P_{\text{th}}$ at fixed volume is

$$P_{\text{th}} = P(V, T) - P(V, 0).$$ \hfill (11)

The enthalpy $H$ does not include thermal effects and can be written as

$$H = E_{\text{static}}(V) + PV.$$ \hfill (12)

The volume thermal expansion coefficient $\alpha$ can be derived via

$$\alpha = \frac{1}{V} \left(\frac{\partial V}{\partial T}\right)_P,$$ \hfill (13)

and the constant temperature bulk modulus is calculated from

$$B_T = -V \left(\frac{\partial P}{\partial V}\right)_T.$$ \hfill (14)

The entropy at constant volume is

$$S = -\left(\frac{\partial F}{\partial T}\right)_V.$$ \hfill (15)
Then the constant volume heat capacity can be calculated from

\[ C_V = T \frac{\partial S}{\partial T}_V = -T \left( \frac{\partial^2 F}{\partial T^2} \right)_V. \]  

(16)

Hence, the thermodynamic Grüneisen parameter \( \gamma_{th} \) is derived via

\[ \gamma_{th} = \frac{\alpha B_T V}{C_V}. \]  

(17)

The adiabatic bulk modulus is written as

\[ B_S = B_T \left( 1 + \gamma_{th} \alpha T \right), \]  

(18)

and the constant pressure heat capacity is calculated from

\[ C_P = C_V \left( 1 + \gamma_{th} \alpha T \right) = C_V + \alpha^2 B_T VT. \]  

(19)

The Hugoniot pressure-volume and pressure-temperature relations are reduced according to the Rankine-Hugoniot conditions [7, 8],

\[ \frac{1}{2} P_H (V_0 - V_H) = (E_H - E_0), \]  

(20)

where \( E_H \) is the molar internal energy along the Hugoniot, and \( E_0 \) and \( V_0 \) are the molar internal energy and volume at zero pressure and room temperature, respectively. Because \( P_H \) and \( E_H \) are both the functions of temperature, temperature \( T \) can be found by numerically solving Eq.20 at fixed volume \( V_H \).

The internal energy, entropy, and constant volume heat capacity of lattice vibrations can be directly achieved from the phonon density of states (normalized to \( 3nN \)) via [10],

\[ E = \frac{\hbar}{2} \int_0^{\omega_{max}} \omega \coth \frac{\hbar \omega}{2k_B T} g(\omega)d\omega, \]  

(21)
\[ S = k_B \int_0^{\omega_{\text{max}}} \left[ \frac{\hbar \omega}{2k_BT} \coth \frac{\hbar \omega}{2k_BT} - \ln \left( 2 \sinh \frac{\hbar \omega}{2k_BT} \right) \right] g(\omega) d\omega, \] (22)

and

\[ C_V = k_B \int_0^{\omega_{\text{max}}} \left( \frac{\hbar \omega}{2k_BT} \right)^2 \mathrm{csch}^2 \left( \frac{\hbar \omega}{2k_BT} \right) g(\omega) d\omega, \] (23)

respectively.

According to the Debye approximation, the specific heat can be expressed as,

\[ C_V = 9N_A k_B \left( \frac{T}{\Theta_D} \right)^3 \int_0^{\Theta_D/T} \frac{x^4 e^x}{(e^x - 1)^2} dx \] (24)

where \( \Theta_D \) is Debye temperature and \( N_A \) is the Avogadro constant. So, at a fixed volume the Debye temperature \( \Theta_D \) at each temperature \( T \) can be obtained from Eqs. 23 and 24.

3. Description of the package and Input/output files

3.1. Installation requirements

The installation of PHASEGO package is very easy. PHASEGO is based on Python, but to increase the computation speed and realize the numerical interpolation and extrapolation, it also uses Numpy and Scipy python libraries.

The following packages are required:

- Python 2.6 or later.
- NumPy.
- Scipy.
- Matplotlib.
Matplotlib is used for the automatic plot of the HPHT phase diagram. In the Ubuntu system, just simply execute `sudo apt-get install python python-numpy python-scipy python-matplotlib`. In the Windows systems, one can install `pythonxy` package for all the necessary libraries.

### 3.2. Installation

There are two methods to install PHASEGO. When one has the python setuptools module installed, the first method is to execute `python setup.py install` in the PHASEGO root directory as the administrator. As a common user, one can also specify a directory to install using additional “`--prefix=/path/to/install/`”. If one has no python setuptools, then the second method to install is: first put the following `PATH` and `PYTHONPATH` environment variable in user’s `~/.bashrc` file:

```bash
export PATH=$PATH:/path/to/phasego/src
export PYTHONPATH=$PYTHONPATH:/path/to/phasego/src
```

or in `~/.cshrc` file:

```csh
setenv PYTHONPATH /path/to/phasego/src:$PYTHONPATH
setenv PATH /path/to/phasego/src:$PATH
```

Then, execute `chmod +x /path/to/phasego/src/phasego`. Finally execute `source ~/.bashrc` or `source ~/.cshrc`. 
3.3. Run

Just type **phasego** to run the phase analysis in your work directory contained a controlling file **phasego.in**. All the controlling arguments are detailed below.

3.4. Input files

The input files needed by **Phasego** package are the static volume-energy data files and the phonon density of states files for each volume. The structure information can be obtained from the crystal structure prediction codes, such as our recently developed **Muse** package [18] and the **calypso** package [19]. All the files are placed in the **inp-phasename** directory for each single structure, where **phasename** is the name of this structure named by the user in the controlling file **phasego.in**. The common prefix name of the static volume-energy data files for all structures are specified in the **phasego.in** file, e.g., **ve-**. The full file names of volume-energy data files are this prefix name plus “T”, where T is the temperature at which the volume-energy data are calculated. To include the electronic thermal excitation effects (especially for metallic materials), the energies can be calculated within the framework of finite temperature DFT. There should be a number of volume-energy data files, e.g., **ve-0, ve-50, ve-100, ve-150**, and so on. For nonmetallic materials, T equals to 0 and there is only one file, e.g., **ve-0**. The phonon dos file is named as **ph.dos-volume**, where **volume** is exactly the same value in the volume-energy file. **ph.dos-** is the common prefix name specified in **phasego.in** for all the structures. The density of states is normalized to $3nN$, where $n$ is the number of atoms in the unit cell and $N$ is the number of the unit cells. The frequencies are in cm$^{-1}$ and dos are in states/cm$^{-1}$.
The following input file example is for the phase transition analysis of GaN at HPHT. The main input file is named as `phasego.in`, in which the lines started with ‘#’ are neglected. Each argument and its value(s) are placed in the same line, in any order and with any number of blank lines. The values of each argument are separated by whitespace and placed after ‘=’. The arguments are all self-explained in their comment line(s).

```plaintext
###========================================================
###=================== Phasego input file ===================
###========================================================
### Lines started with "#" are neglected.
### Parameter and value(s) are placed in the same line, in any
### order with any number of blank lines.
###
### # Note:
###
### # A. volumes in Bohr$^3$, energies in Ry.
### # B. Num_atoms is the number of atoms in the unit cell for
### #   energy and phonon dos calculations.
### # C. dos files: normalized to 3*N, and frequencies in cm$^{-1}$
### #   and dos in states/cm$^{-1}$.
###
### # EOS Names (number):
### # 1. Murnaghan: Murnaghan EOS (F. D. Murnaghan, Am. J. Math.
### #       49, 235 (1937))
```
# The name of EOS used for fitting

Eos_Name = 1

# The user-defined name for each corresponding structure. The
# ve and phonon dos files are placed in each inp-name dir. The
# output files are placed in each out-name dir.

Names_of_Strs = rocksalt wurtzite zencblende
# Number of atoms used for ve and phonon dos data of each str.
Num_atoms = 2 4 2

# The prefix name of volume-energy data files for each str.
# The full name is the prefix name plus "T", where T is
# temperature to take into account the electronic thermal
# excitation by finite temperature DFT. If including the
# electronic thermal excitation free energy, the temperature
# start, end, and interval should be the same as Tdata below.
VE_data_File_Name = ve-

# Phonon dos file prefix name, i.e., it plus the volume value
# in ve file is the full name.
Ph_Dos_File_Base_Name = ph.dos-

# Temperature data (K): start, end, interval
Tdata = 0 6000 100

# Pressure data (GPa): start, end, interval
Pdata = 0 40 1

# If calculate thermal properties, yes or no
If_Incl_Phonon = yes
# If include electronic thermal excitation free energy by
# finite temperature DFT, for metals
If_Incl_Electronic_Excitation = no

# If analyze the potential phase transition P-T points between
# these strs, yes or no
If_Analyze_Phase_Transition = yes

# If plot phase diagram using matplotlib, yes or no
If_Plot = yes

# If analyze Hugoniot PTV, yes or no
If_Hugoniot = yes

# If calculate Debye Temperature, yes or no
If_Calc_Debye_Temp = yes

### 3.5. Output files

The output files for single phase analysis are placed in the `out-phasename` directory of each structure. The Alpha.dat file contains the thermal expansion coefficients data as the function of temperature at fixed pressures. The B_S.dat file has the adiabatic bulk modulus data as the function of temperature at fixed pressures. The B_T_P.dat and B_T_T.dat files contain the isothermal bulk moduli at fixed pressures and temperatures, respectively. The C_P.dat file contains the outcomes of heat capacities at fixed pressures. The C_V_P.dat and C_V_V.dat files have the constant volume heat
capacities at fixed pressures and volumes, respectively. The DebyeT.dat file has the Debye temperatures as the function of temperature at fixed volumes. The Enthalpy.dat is the data file of the enthalpy as a function of pressure. The Entropy_V.dat and Entropy_P.dat collect the total entropies at fixed volumes and pressures, respectively. The fittedC_V_V.dat includes the fitted $C_V$ at fixed volumes. The fittedE_V.dat is the data file of the fitted internal energy and pressure. The FittedHelmFreeE_T.dat file contains the fitted Helmholtz free energies at fixed temperatures. The gamma_P.dat has the thermodynamic Grüneisen parameters as the function of temperature at fixed pressures. The G_P.dat and G_T.dat files include the Gibbs free energies as functions of temperature and pressure, respectively. The HugoniotPTV.dat is the output file of the Hugoniot P-T-V data. The PV_T.dat file contains the pressure-volume data at fixed temperatures. The ThermalP_T.dat and ThermalP_V.dat have the thermal pressure data at fixed pressures and volumes, respectively. The VT_P.dat is the output file of the volume-temperature data at fixed pressures. The C_V-ph-direct.dat, Entropy-ph_V.dat and E-ph_V.dat are the constant volume heat capacities, entropy and internal energy of lattice vibrations directly from the phonon density of states, respectively.

The phase transition data and the automatically plotted phase diagram figures are placed in the Phase-PT directory. The P-T.dat file contains the transition pressures at fixed temperatures, which are obtained by canning pressure at each fixed temperature. Similarly, the T-P.dat file contains the transition temperatures at fixed pressures obtained by scanning temperature at each fixed pressure. They can be plotted by other plot softwares. The
phase boundaries are indexed by “|” and the phase transitions are labelled by “—>”.

4. Two examples: phase analyses of GaN and Fe

To test the ability of PHASEGO package, I performed phase analyses for the semiconductor GaN and the metal Fe. PHASEGO can automatically determine the phase boundaries and plot the phase diagrams for the several competing structures. The total time to run is less than a minute both for the two cases. For GaN, the thermodynamic properties of the wurtzite phase are also presented and discussed below.

Their phonon frequencies and phonon density of state were calculated using the density functional perturbation theory (DFPT) [20, 21], as implemented in the QUANTUM-ESPRESSO package [22]. The exchange-correlation functional used is generalized gradient approximation (GGA)
parametrized by PBE [23] and the pseudopotential is ultrasoft pseudopotential [24]. The Fe pseudopotential is the newest version from the recently developed pseudopotential library [25]. We carefully tested on \( \mathbf{k} \) and \( \mathbf{q} \) grids, the kinetic energy cutoff, and other technical parameters to ensure good convergence of phonon frequencies. The kinetic energy cutoff and the energy cutoff for the electron density were chosen to be 50 Ryd., 500 Ryd for GaN, and 40 Ryd. and 400 Ryd. for Fe in both total energy and phonon dispersion calculations, respectively. We applied the Marzari-Vanderbilt [26] smearing method with the smearing width of 0.03 Ryd. For the electronic thermal excitation in Fe, the smearing with the Fermi-Dirac function was applied to set the electronic temperature.

![Graph](image)

Fig. 2: (a) The thermal pressure of wurtzite-GaN as the function of temperature at different pressures. (b) The Debye temperature of wurtzite-GaN as the function of temperature at fixed volumes.

4.1. Thermodynamics of GaN

The thermodynamic properties of its zinc blende, wurtzite and rocksalt structures are first calculated before phase diagrams calculations. Figure1a
shows the thermal expansion coefficients of wurtzite-GaN as the function of
temperature at fixed pressures 0, 10, 20, and 30 GPa. The thermal expansion
coefficient increases with temperature at fixed pressure. But it decreases with
pressure at fixed temperature. The isothermal bulk modulus as a function
of temperature at fixed pressure is illustrated in Fig.1b, which shows its
decreasing behaviour with temperature at fixed pressure.

![Graph showing the thermal expansion coefficients of wurtzite-GaN.]

The thermal pressure of wurtzite-GaN as a function of temperature at
fixed atomic volumes 72.07, 76.71, and 81.35 a.u.\(^3\) are reduced and shown
in Fig.2a. The thermal pressure increases with temperature as fixed vol-
ume. But unusually the thermal pressure of wurtzite-GaN decreases with
decreasing volume. The Debye temperatures of wurtzite-GaN (Fig.2b) ex-
hibit dramatic drops before 50 K and then increase quickly before 300 K,
and finally converge to constants at fixed volumes.

Manually reducing the Hugoniot pressure-temperature-volume relations
is a time-consuming work. For the PHASEGO package, it is very easy to
implement. As an example, the Hugoniot pressure-volume and pressure-temperature relations of wurtzite-GaN are calculated and presented in Fig. 3a.

Fig. 4: (a) The QHA based high pressure and temperature phase diagram of GaN from the automatic analysis and plot of phase transition. (b) The QHA based high PT phase diagram of Fe with (red symbols) and without (black symbols) electronic thermal excitation free energy corrections.

4.2. Phase diagrams of GaN and Fe

Figure 4a is the automatic plot of the phase diagram of GaN. One only needs provide the necessary volume-energy and phonon density of state data for different structures covering the pressure range of interest. The PHASEGO package will gather the Gibbs free energy information and judge the stable field of each structure, and finally plot the phase diagram and label the phase boundary between two structures automatically. The triple point is also clearly shown in the phase diagram. For metallic materials, the electronic thermal excitation free energies at high temperature can not be neglected. In the case of Fe, the results of automatic phase transition analyses with and
without electronic thermal excitation free energy corrections are shown in Fig. 4b. One notes that the electronic thermal excitation free energies alter the phase boundaries much obviously at high temperature.

5. Conclusions

I described here the PHASEGO package which implements the automatic calculation and plot of phase diagram based on the QHA. The QHA theory and the numerical derivation process of the thermodynamic properties are also detailed. In order to test the validity and efficiency of the PHASEGO package, I performed the phase transition analyses and thermodynamic properties calculation for GaN. For the metallic materials, I took Fe as an example, and found the electronic thermal excitation free energy can not be neglected at high temperature and it changes the phase diagram of Fe obviously.

6. ACKNOWLEDGMENTS

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