Equations of state and stability of MgSiO$_3$ perovskite and post-perovskite phases from quantum Monte Carlo simulations


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Equations of state and stability of MgSiO$_3$ perovskite and post-perovskite phases from quantum Monte Carlo simulations

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We have performed quantum Monte Carlo (QMC) simulations and density functional theory (DFT) calculations to study the equations of state of MgSiO$_3$ perovskite (Pv) and post-perovskite (PPv), up to the pressure and temperature conditions of the base of Earth’s lower mantle. The ground state energies were derived using QMC and the temperature dependent Helmholtz free energies were calculated within the quasi-harmonic approximation and density functional perturbation theory. The equations of state for both phases of MgSiO$_3$ agree well with experiments, and better than those from generalized gradient approximation (GGA) calculations. The Pv-PPv phase boundary calculated from our QMC equations of states is also consistent with experiments, and better than previous LDA calculations. We discuss the implications for double crossing of the Pv-PPv boundary in the Earth.

I. INTRODUCTION

The accurate description of electronic correlation effects is one of the main challenges in theoretical condensed matter physics. Quantum Monte Carlo (QMC) [1–5] simulations can describe these correlation effects while maintaining a high computational efficiency [6]. A number of recent studies demonstrate the growing ability of the QMC method to accurately describe ground state properties of complex solids [7–14]. This large basis of previous work provides the motivation to apply QMC calculations to solid silicate perovskite (Pv) and post-perovskite (PPv) (MgSiO$_3$) in order to derive equations of state that are more accurate than those that have been previously obtained with density functional theory (DFT).

The Pv-PPv phase transition is particularly important because Pv is the dominant phase in Earth’s lower mantle [15]. Pv was the only known phase under lower mantle conditions until a phase transition to PPv at a pressure of 125 GPa and temperature of 2500 K was discovered in 2004 [16, 17]. The post-perovskite phase is believed to exist in Earth’s thin, core-mantle boundary layer, known as D″. The discovery of MgSiO$_3$ PPv has attracted considerable attention as it offers a possible explanation for many of the unusual properties of the D″ layer, such as the inhomogeneous seismic discontinuity observed a few hundred kilometers above the core-mantle boundary, anomalous seismic anisotropy, and ultra-low velocity zones [16–21]. Some quantitative estimates of these anomalies were made by Wentzcovitch et al. [22].

Many computations [17, 23–31] based on DFT [32, 33] have reported the equations of state of Pv and PPv. However, DFT results are dependent on the choice of exchange-correlation functional [10, 34, 35] since the exact exchange-correlation functional is unknown. Generally, DFT with the local density approximation (LDA) provides a good P-V relationship for MgSiO$_3$ perovskite [23, 24], but underestimates the Pv-PPv transition pressures [17, 27]. In contrast, whereas DFT with the generalized gradient approximation (GGA) provides a better prediction of the Pv-PPv transition pressure, it overestimates the zero pressure lattice volume in the equation of state [17]. The ~10 GPa difference between the LDA and GGA predictions of the phase transition pressure [27] makes a difference in depth of about 150 km, according to the preliminary reference Earth model (PREM) [36]. The discrepancy among DFT calculations, although relatively small for many applications, are significant with regard to geophysical modelling. The position of the Pv to PPv boundary is crucial for interpreting seismic data from the base of the mantle, to understand if this transition is sufficient to explain most lower mantle heterogeneity, or if there must also be partial melt, compositional heterogeneity, etc. In particular, double crossing of the Pv-PPv boundary, [37, 38] or more generally, through the two-phase region, [39] may give indication of temperature and compositional variations, which are crucial for interpreting the seismological data, [40] and as input into geodynamic modelling [41].

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II. COMPUTATIONAL METHODS

A. Quantum Monte Carlo

A rigorous discussion of QMC methods has been reported in previous publications [4, 5]. Here, we briefly outline the main choices we make within the methodology. We employ two types of QMC sequentially in order to extract the ground state properties of a system. The first is known as variational Monte Carlo (VMC), and we employ two types of QMC sequentially in order to extract the ground state properties of a system. The first is known as variational Monte Carlo (VMC), where the fixed form of the trial wavefunction, is evolved in imaginary time until it exponentially decays to the ground state. The statistical wavefunction, constructed from the optimized trial VMC wavefunction, is evolved in imaginary time propagation, and satisfies the cusp condition. The Jastrow parameters are optimized by minimizing a combination of variance of the VMC energy and the energy itself [42].

VMC by itself is generally not accurate enough due to the fixed form of the trial wavefunction. In a second method, diffusion Monte Carlo (DMC), a statistical representation of the wavefunction is evolved according to a version of the Schrödinger equation which has been transformed to an imaginary time diffusion equation. The statistical wavefunction, constructed from the optimized trial VMC wavefunction, is evolved in imaginary time until it exponentially decays to the ground state. The DMC method is very efficient at projecting out the time-averaged bound on the total energy. In MgSiO$_3$, an insulating, closed shell system, we expect this approximation to be very good, and in general to be independent of structure or compression, so even if there is a small shift in the total energy from inexact nodes, it should be very close in each structure studied. The agreement we discuss below with experiment is post-hoc evidence of this. The second approximation is related to the use of pseudopotentials, discussed more below. This would be no worse than the use of pseudopotentials in DFT, except that there is an additional “locality approximation” which can make results sensitive to the pseudopotentials used. This approximation can be controlled and tested, and again should give similar errors to each structure for the MgSiO$_3$ system. Again, the evidence is that this is usually a very small error.

B. Pseudopotentials

While great care is taken to generate pseudopotentials, they are constructed within the mean field treatment rather than a many-body framework. A recent paper which systematically applied diffusion Monte Carlo to calculate properties of solids concluded that the determining factor on the accuracy of the method was the fidelity of the pseudopotentials used [13]. This conclusion echoed results of earlier studies on geophysics with QMC which found the effects of the pseudopotential approximation to be large [8]. One possible strategy to mitigate this error is to perform all electron calculations as was done in recent work on boron nitride [11]. This approach is however impractical for this study due to the extreme computational demands posed by all electron calculations within QMC for heavier elements. As an alternative, we have endeavored to test the pseudopotentials used in this paper as rigorously as possible in an attempt to validate their use in DMC.

Our testing methodology has three parts. First we have tested the Mg and O pseudopotentials by comparing LAPW calculations performed with the ELK code [43] to pseudopotential calculations performed with quantum espresso [44]. In calculations of the equilibrium lattice constant and bulk modulus we find excellent agreement with the all electron results giving 4.219 Å and 156.6 GPa and the pseudopotential results giving 4.206 Å and 159.9 GPa. While this test is not conclusive in terms of stating that the pseudopotential will be accurate for QMC calculations, it does preclude corrections of the type applied in earlier work [8].

The second test we applied was to calculate the electron affinity and ionization potential for each of the pseudopotentials used and to compare the results to experiment as was shown to be useful in a recent paper on Ca$_2$CuO$_3$ [45]. Here we use the Slater-Jastrow form for the trial wavefunction with a single Slater determinant. The spin state for the neutral atom, anion and cation

\[
\Psi_T(R) = D^\uparrow D^\downarrow e^J, \tag{1}
\]

The up- and down-spin Slater determinants, $D^\uparrow$ and $D^\downarrow$, are obtained from DFT calculations. The Slater determinant fixes the nodal surface of the calculation, which is used in the so-called fixed-node approximation. The Jastrow factor, $J$, is the exponential of a sum of parameterized one-body and two-body terms that are a function of particle separation, and satisfies the cusp condition. The Jastrow parameters are optimized by minimizing a combination of variance of the VMC energy and the energy itself [42].

\[
\Psi_{\text{DMC}} = \lim_{\Delta t \to 0} \prod_{j=1}^{N} e^{-H \Delta t} \Psi_T, \tag{2}
\]

where $H$ is the Hamiltonian, $\Delta t$ is step size used for imaginary time propagation, and $N$ corresponds to the number of projections. In both VMC and DMC, the space of electron configurations is simultaneously explored with an ensemble of independent configurations, which follow a random walk. In DMC, the walk is guided by an importance-sampled wave-function for efficiency. Once configurations have equilibrated, averages of their energies can be accumulated and analyzed statistically. This allows QMC methods to be massively parallelized in computations.
TABLE I. Electron affinity (EA) and ionization potential (IP) from QMC and experiment given in eV. The electron affinity of Si and O is taken from Ref. 47, while the negative electron affinity of Mg is from Ref. 48. The ionization potentials are taken from Ref. 49.

<table>
<thead>
<tr>
<th></th>
<th>QMC EA</th>
<th>Expt EA</th>
<th>QMC IP</th>
<th>Expt IP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mg</td>
<td>unbound</td>
<td>unbound</td>
<td>7.591±0.013</td>
<td>7.64624</td>
</tr>
<tr>
<td>O</td>
<td>1.372±0.013</td>
<td>1.4611134</td>
<td>13.681±0.026</td>
<td>13.61806</td>
</tr>
<tr>
<td>Si</td>
<td>1.430±0.013</td>
<td>1.3896210</td>
<td>8.228±0.012</td>
<td>8.15169</td>
</tr>
</tbody>
</table>

are determined from spin polarized DFT calculations and are kept fixed in the QMC. While this is not likely to produce highly accurate electron affinities or ionization potentials, large errors greater than approximately 0.1 eV would be a cause of concern. In this case however, the pseudopotentials prove to be rather accurate as shown in Table I.

The third test we applied was to calculate binding curves of the molecules MgO, O$_2$ and SiO. These energy vs. separation curves were then fitted to a Morse potential and the resulting atomization energies, bond lengths and vibrational frequencies are compared to experiment where possible. From the example of MgO case in Fig. 1, we could see the fitting of Morse potential to the QMC energy is pretty good. It should again be noted that an attempt to make these calculations as accurate as possible would use a more sophisticated wavefunction containing for instance a multidental expansion [46]. However, the performance of the single Slater-Jastrow trial wavefunction is highly relevant as this is the form used in calculation of the properties of the solid phases. In this case we find excellent agreement with experiments (Table II), leading us to conclude that these pseudopotentials are accurate for use in calculating the perovskite to post-perovskite phase transition pressure.

![MgO energy vs separation from DMC](image)

**FIG. 1.** Energy vs separation curve for MgO calculated using DMC.

TABLE II. Equilibrium bond length ($r_e$) in angstroms, vibrational frequency ($\omega_0''$) in cm$^{-1}$ and atomization energy ($D_0$) in eV calculated with QMC and fit to a Morse potential compared to experiments. Note that the vibrational frequency of SiO is difficult to obtain experimentally due to issues in isolating SiO at low temperatures.

<table>
<thead>
<tr>
<th></th>
<th>QMC $r_e$</th>
<th>Expt $r_e$</th>
<th>QMC $\omega_0''$</th>
<th>Expt $\omega_0''$</th>
<th>QMC $D_0$</th>
<th>Expt $D_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>MgO</td>
<td>1.7519±0.0018</td>
<td>1.749 [50]</td>
<td>782.3±32.5</td>
<td>785.2183±0.0006 [51]</td>
<td>2.14±0.02</td>
<td>2.56±0.22 [52]</td>
</tr>
<tr>
<td>O$_2$</td>
<td>1.1978±0.0007</td>
<td>1.208 [50]</td>
<td>1480.5±11.4</td>
<td>1580.161±0.009 [51]</td>
<td>4.89±0.02</td>
<td>5.117 [53]</td>
</tr>
<tr>
<td>SiO</td>
<td>1.5097±0.0007</td>
<td>1.51 [50]</td>
<td>788.9±22.6</td>
<td>1241.54388±0.00007 [51]</td>
<td>7.98±0.42</td>
<td>8.24 [53]</td>
</tr>
</tbody>
</table>

It is possible to make high quality pseudo potentials for correlated systems [54]. However, such pseudopotentials would not be applicable to the DFT computations we use to generate our trial functions, so one would have to use different pseudopotentials to generate the trial functions. Put in this way, one could say that the problem is not with pseudo potentials per se, but in ones that are useable for generation of trial functions, or that the ultimate problem is with the trial functions. In principle the trial functions could be parametrized and optimised variationally, [55], or backflow [56], or other nodal variations could be used, but such computations have not yet been possible for complex solids such as we study here.

C. Quasi Harmonic Phonon free energies

Whereas the static crystal energy can be obtained by DFT [32, 33] or QMC [1–3] calculations, it is currently intractable to calculate the phonon frequencies from quantum Monte Carlo simulations. Therefore, in
our results we combine static QMC energies with vibra-
tional energies from density functional perturbation the-
ory (DFPT) calculations. The accuracy of QMC static
energy plus DFPT vibrational energies has been shown
to be an improvement over using DFT plus DFPT for
the silica phases \[10\]. Once the Helmholtz free energies
are obtained for several lattice volumes at various tem-
peratures, the temperature dependent equation of state
and other thermodynamic properties of interest are de-
determined.

The Helmholtz free energy is a function of lattice vol-
ume and temperature \( F (V, T) \). Using the Vinet equa-
tion of states \[57, 58\], the Helmholtz free energy is
\[
F = F_0 + \frac{4K_0 V_0}{(K'_0 - 1)^2} - \frac{2K_0 V_0}{(K'_0 - 1)^2} \left\{ 5 + 3K'_0 \left[ \left( \frac{V}{V_0} \right)^{1/3} - 1 \right] - 3 \left( \frac{V}{V_0} \right)^{1/3} \right\} \exp \left\{ -\frac{3}{2} (K'_0 - 1) \left[ \left( \frac{V}{V_0} \right)^{1/3} - 1 \right] \right\}
\]
where \( F_0, V_0, K_0 \) and \( K'_0 \) are the Helmholtz free energy,
lattice volume, bulk modulus and its pressure deriv-
tive respectively, under zero pressure. Within the quasi-
harmonic approximation (QHA), the Helmholtz free en-
ergy is given by \[59, 60\],
\[
F = E + TS = E_{\text{static}} + \frac{1}{2} \sum_{k,i} h \omega_{k,i} + k_B T \sum_{k,i} \ln \left[ 1 - \exp \left( \frac{-h \omega_{k,i}}{k_B T} \right) \right]
\]
where \( E \) is the internal energy, \( S \) is the entropy, \( E_{\text{static}} \)
is the static energy, \( h \) is the Planck constant/\(2\pi \), \( \omega_{k,i} \)
is the angular frequency of a phonon with wave vector \( k \)
the \( i \)-th band, \( k_B \) is the Boltzmann constant and
\( T \) is the absolute temperature. In the quasi-harmonic
approximation, \( E_{\text{static}} \) and \( \omega_{k,i} \) are independent of \( T \)
and are determined only by the atomic positions and
lattice parameters at zero temperature.

D. Computational details

The pseudopotentials used for all DFT, DFPT and
QMC calculations in this work were generated with the
OPIUM code \[61\] using the WC exchange correlation
functional \[35\]. The core radii of the pseudopotentials
are as follows: 1.2(1s), 1.2(2s) for magnesium; 1.3(1s),
1.3(2s) for oxygen; and 1.7(1s), 1.7(2s), 1.7(2p) for silici-
on.

Our DFT equations of state are constructed from the
energies of seven different volumes in both the Pv and
PPv phases. We used the plane-wave pseudopoten-
tial DFT code, PWSCF \[44\], to relax the atomic po-
sition, obtain the static DFT energy, and extract the
single-particle orbitals for the QMC wavefunction at
each volume. The seven volumes correspond to con-
tinuous pressure simulations at -20, -10, 0, 50, 100, 150,
and 200 GPa. These calculations used the Wu-Cohen
(WC) \[35\] exchange correlation approximation, a plane-
wave energy cutoff of 300 Ry and Monkhorst-Pack k-
point meshes of \( 6 \times 6 \times 6 \) and \( 12 \times 6 \times 6 \) for the 20
atoms unit cell of perovskite and post-perovskite re-
spectively, which converged the total energy to tens of
milli-Ry/MgSiO\(_3\) accuracy. At each volume above, the
phonon frequencies and temperature dependent vibra-
tional energies were calculated with ABINIT \[62\] using
density functional perturbation theory within the quasi-
harmonic approximation. These calculations used
q-point meshes of \( 4 \times 4 \times 4 \) for the 20 atom unit cell of
perovskite and \( 4 \times 4 \times 2 \) for the 10 atom unit cell of
post-perovskite, which ensured calculated phonon free
energies were converged to tens of milli-Ry/MgSiO\(_3\).

The accuracy of our QMC calculations is determined
by three classes of approximations that are necessary
for computational efficiency of fermionic calculations:
finite simulation cell size effects, pseudopotentials, and
the fixed node approximation \[63\]. For accurate QMC
results, one must reduce the error introduced by these
approximations such that the end result is converged.
The effectivity of pseudopotentials in QMC calculations
was checked in the previous section. Here, we discuss
how the other approximations are mitigated.

In any simulation we are forced to simulate a true
solid with a simulation cell that is subject to periodic
boundary conditions. Finite-size errors arise from both
one-body effects due to discrete k-point sampling of the
Brillouin zone and two-body effects from spurious elec-
tron correlation in the periodic cells. We minimize the
one-body errors by using twist averaged boundary con-
ditions. We average over eight twists, allowing us to
improve our sampling of the Brillouin zone. The two-
body errors are minimized by using the Model Periodic
Coulomb (MPC) interaction \[63-65\], which corrects the
potential energy for the spurious correlation effects. We
then use the scheme of Chiesa et al. \[66\] to correct the
kinetic energy two-body effects. While applying these
techniques, we then perform our calculations in three
different supercell sizes of 40, 80, and 120 atoms, and
we fit an extrapolation to infinite cell size.

The final approximation we will discuss is that of
the nodal surface to handle the fermion sign problem.
QMC samples a positive definite probability function
constructed from a antisymmetric wavefunction which
has positive and negative regions. Unchecked, sampling
the probability in this way will lead to a bosonic ground
state as positive and negative contributions cancel out
and the odd-parity solution becomes swamped in sta-
tistical noise. In order to circumvent this problem, ab-
sorbing barriers are placed between all nodal pockets in
configuration space. This can only be done if the nodes
are fixed to a known location at the start of DMC (we
use the nodes from DFT), which is called the fixed-node
approximation. The size of the fixed-node error is gener-
ally assumed to be small, and, for small systems, can be checked with backflow optimization of the single-particle orbital coordinates, but this is too computationally expensive for the systems studied here.

In order to ensure electron correlation was treated uniformly across all of our calculations, we fixed the Jastrow parameters in DMC simulations for all volumes and supercell sizes to values obtained from optimizing the Jastrow for the smallest volume and supercell size. In addition, for computational efficiency, a b-spline basis set is used to represent the single particle orbitals centered on a grid of points. The b-spline basis set provides an order-N speed up in the calculation, where N is the number of atoms, but doubles the memory requirement relative to an analytic, plane-waves basis. The mesh size of this grid is decreased until the total energy is converged to tenths of milli-Ry/MgSiO$_3$. We use a b-spline mesh factor of 0.8.

For the wavefunction optimization part of our calculations, a combination of energy and variance minimization was used in a series of twenty optimizations in which the VMC total energy was determined to a one-sigma statistical accuracy of 0.05 eV/MgSiO$_3$ and the fluctuation among the VMC energies after each optimization became less than 0.1 eV/MgSiO$_3$. The Jastrow factor which gave both lowest total energy and smallest variance was chosen for use in the subsequent DMC calculations, a combination of energy and variance minimization became less than 0.1 eV/MgSiO$_3$.

The VMC total energy was determined to a one-sigma statistical accuracy of 0.05 eV/MgSiO$_3$. We use a b-spline mesh factor of 0.8.

A typical DMC simulation used 300-400 electron configurations and collected statistics over 25,000 Monte Carlo steps. The first 5000 steps were used to equilibrate the simulation. The total energy of each supercell was obtained by averaging the energies of the remaining 20,000 steps over 8 twists. The standard error $\delta$ of the total energy was obtained by $\delta = \sqrt{\sigma^2/N}$, where $\sigma$ is the energy variance of block samples and $M$ is the uncorrelated samples [67]. The DMC time-step was determined by converging the total energy with respect to changes in the time-step. Our convergence tests found that a time-step of 0.001 Ha$^{-1}$ is sufficient for 0.05 eV/MgSiO$_3$ accuracy. The VMC and DMC simulations were preformed with the GPU version of QMCPACK [67–69].

III. RESULTS AND DISCUSSION

A. Enthalpy and Volume

In our simulations, both the MPC corrected and uncorrected QMC total energies as a function of simulation cell size are quite linear (Fig. 2). All QMC results hereafter are from DMC simulations unless stated otherwise. Two linear equations were used to fit the MPC-corrected and uncorrected total energies synchronously. The equations are $E_{N}^{\text{cor}} = E_{\text{inf}}^{\text{cor}} + k_{\text{cor}}N/N$ for MPC corrected energies and $E_{N}^{\text{uncor}} = E_{\text{inf}}^{\text{uncor}} + k_{\text{uncor}}N/N$ for uncorrected energies, where $k$ is the slope and $N$ is the number of MgSiO$_3$ formula. $E_{\text{inf}}^{\text{cor}}$ and $E_{\text{inf}}^{\text{uncor}}$ were kept equal to each other during the fitting process using the least squares method and they are our final infinite size QMC total energy, which is the static energy in Eq. 4. The error of the infinite size energy was taken to be the same as that of the largest supercell size case.

![Fig. 2. The extrapolation to infinite size QMC total energy using the MPC-corrected and uncorrected finite size total energies. All the total energies are the 8 twists averaged results. $N_{\text{MgSiO}_3}$ is the number of formula units in the supercell used in the QMC simulation. Here the volume is 41.19 Å$^3$/MgSiO$_3$ for $P_v$ and 41.15 Å$^3$/MgSiO$_3$ for PPv.](image)

At a given temperature, the Vinet equation of state (Eq. 3) was used to fit the Helmholtz free energies as a function of volume. The fittings both for DFT and QMC calculations are quite good as shown in Fig. 3. The predicted equilibrium volume, bulk modulus and its pressure derivative from QMC simulations at 300 K are all in good agreement with experimental results (Table III) both for $P_v$ and PPv. This indicates QMC is better than GGA because all the equilibrium volumes predicted by this and previous GGA calculations are larger than experimental data, and the bulk modulus predicted by GGA are smaller than experiments. With the increase of temperature, the bulk modulus decreases while it’s derivative on pressure increases (Fig. 4).

![Fig. 3. QMC and DFT Helmholtz free energies of MgSiO$_3$ $P_v$ and PPv at 300 K, the error bars of QMC results are covered by the symbols. The dashed and solid lines are the fit of the Vinet equation of state.](image)
TABLE III. Equation of state parameters of pervoskite and post-perovskite. The uncertainties of QMC thermodynamic quantities were propagated from the errors of QMC static energies via a linearized Taylor expansion.

<table>
<thead>
<tr>
<th></th>
<th>$F_0$ (eV/MgSiO$_3$)</th>
<th>$V_0$ (Å$^3$/MgSiO$_3$)</th>
<th>$K_0$ (GPa)</th>
<th>$K_0'$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Pv</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-3113.61(3)</td>
<td>40.36(8)</td>
<td>270(14)</td>
<td>3.9(4)</td>
<td>QMC, Static, this work</td>
</tr>
<tr>
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<td>40.88(10)</td>
<td>258(15)</td>
<td>4.0(4)</td>
<td>QMC, 300 K, this work</td>
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<tr>
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<td>41.20</td>
<td>239.4</td>
<td>4.1</td>
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<tr>
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<tr>
<td>-3109.54</td>
<td>40.78</td>
<td>232</td>
<td>3.86</td>
<td>GGA, Static [30]</td>
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<td>-3109.54</td>
<td>40.58-40.83</td>
<td>246-272</td>
<td>3.65-4.00</td>
<td>Exp. [15, 70-77]</td>
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<tr>
<td><strong>PPv</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-3113.38(3)</td>
<td>40.51(8)</td>
<td>232(9)</td>
<td>4.1(3)</td>
<td>QMC, Static, this work</td>
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<td>-3113.00(3)</td>
<td>41.08(9)</td>
<td>221(10)</td>
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<td>4.7</td>
<td>GGA (WC), 300 K, this work</td>
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<td>231.9</td>
<td>4.43</td>
<td>LDA, 300 K [17]</td>
</tr>
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<td>200.0</td>
<td>4.54</td>
<td>GGA, 300 K [17]</td>
</tr>
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<td>222</td>
<td>4.2</td>
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<tr>
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<td>4.41</td>
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<td>4.05</td>
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<td>4.19</td>
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<td>209</td>
<td>4.4</td>
<td>GGA (PW91), Static [31]</td>
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<tr>
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<td>40.55-41.23</td>
<td>219-248</td>
<td>4.0(fixed)</td>
<td>Exp. [77-79]</td>
</tr>
</tbody>
</table>

B. P-V-T equation of state

The thermal equation of state can be calculated by $P = -(\partial F/\partial V)_T$ from the Helmholtz free energy Eq. 3. The comparison between the computed thermal equations of state and previous experimental data for both Pv and PPv phases are figured in Fig. 5. The shading of the QMC curves in Fig. 5 indicates the width of standard deviation of volume as a function of pressure caused by the statistical errors of QMC energies. These comparisons indicate our QMC simulations and LDA calculations predicted a better P-V-T relationship than GGA calculations for both Pv and PPv. The LDA calculations for PPv were taken from Ref. [29] where the comparison with experiments was not checked.

We also calculated the volume differences between MgSiO$_3$ pervoskite and post-perovskite phases as a function of pressure and compared them with some available experimental data in Fig. 6. The comparison indicates that our QMC results are in good agreement with experiments [84]. At lower mantle conditions, our Pv-PPv volume difference is much closer to experiment than DFT.

C. Phase boundary

In thermodynamics, the Gibbs free energy $G$ is defined as $G = F + PV$. At a fixed temperature, a phase transition occurs when Gibbs free energy of the current phase becomes greater than that of another phase with the change of pressure. Because of the uncertainty of static total energy from QMC simulations, we could only predict a range of transition pressure as shown in Fig. 7. Due to the fact Gibbs free energy differences between MgSiO$_3$ Pv and PPv are very small, the range of transition pressure from QMC simulations looks somehow wide. In spite of that, the predicted one-sigma range of transition pressure by QMC simulations still has obvious deviation from that predicted by DFT calculations for MgSiO$_3$ Pv and PPv phases. In static state, we obtained a Pv-PPv phase transition pressure of 91.2 GPa from GGA results and, at a one sigma intervals, 101.0±4.6 GPa from QMC results. Again, we see the transition pressure predicted by this DFT calculation is different from previous DFT studies. At 2000 K, we obtained a Pv-PPv phase transition pressure of 107.1 GPa from DFT (GGA) results and, at a one sigma interval, 117.5±4.8 GPa from QMC results.

At any temperature in the range of 0 to 4500 K, the Pv-PPv transition pressure predicted by our DFT computations with the WC exchange-correlation func-
FIG. 4. Vinet equation of state parameters based on QMC static energies and DFPT vibrational energies as a function of temperature. (a), (b), (c) and (d) are for perovskite. (e), (f), (g) and (h) are for post-perovskite. The shading of the curves indicate one-sigma width of the uncertainties.

D. Thermodynamic properties

The thermal pressure is defined as

\[ P_{th}(V, T) = P(V, T) - P(V, 0) = -(\partial F_{th}/\partial V)_T \] (5)

where \( F_{th} \) is the thermal free energy (the third term in Eq. 4). For either phase of Pv and PPv, the averaged thermal pressure over volume as a function of temperature is quite linear at temperatures larger than 1000 K (Fig. 9). The slopes of the linear parts of thermal pressure curves are 7.74 MPa K\(^{-1}\) for Pv and 7.65 MPa K\(^{-1}\) for PPv.

The thermal expansivity \( \alpha \) is calculated from thermal pressure as

\[ \alpha = (\partial P_{th}/\partial T)_V/K_T \] (6)

where \( K_T = -V(\partial P/\partial V)_T \) can be obtained from the thermal equation of state. In the next part of this section, the related thermal equation of state is derived based on QMC static energies and DFPT vibrational energies. The obtained thermal expansivities in this work fall in the region of previous models which were derived from experimental data (Fig. 10).

The Grüneisen ratio \( \gamma \) is calculated by

\[ \gamma = \frac{\alpha K_T V}{C_V} \] (7)

where \( C_V \) is the constant volume heat capacity obtained from phonon calculations. The Grüneisen ratios both
FIG. 5. Thermal equations of state of (a) perovskite and (b) post-perovskite at three temperatures. The static LDA results of Pv are taken from Ref. [23], and the finite temperature LDA results of Pv are taken from Ref. [24]. The LDA results of PPv are taken from Ref. [29]. The experimental results of Pv are taken from Refs. [23, 71, 76, 77, 80–85] at 300 K and from Refs. [81, 82, 84] at 2000 K. The experimental results of PPv are taken from Refs. [16, 77–79, 84] at 300 K and Refs. [79, 84] at 2000 K. The shading width of QMC results indicates two-sigma statistic errors at the given pressure.

for Pv and PPv fall in the region of previous models (Fig. 11).

IV. CONCLUSION

We have presented QMC computations of MgSiO₃ equations of state and stability for both perovskite and post-perovskite. Our results showed that QMC not only gives good equations of state but also a reasonable Pv-PPv phase boundary for MgSiO₃ under lower mantle temperature conditions. For this iron free silicate, the predicted QMC Pv-PPv phase boundary may have a double-crossing of the geotherm, which would lead to a second Pv phase region just above the core mantle boundary. However, we could not conclude that this double-crossing will exist in the lower mantle, since the presence of Fe could change the Pv-PPv phase boundary dramatically [30, 77]. The accuracy of QMC in this three component system has been demonstrated. It indicates that it is possible to further study the equations of state of the iron bearing silicate (Fe,Mg)SiO₃ using QMC simulations.

V. ACKNOWLEDGMENTS

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FIG. 8. Phase diagram of MgSiO$_3$ under lower mantle pressure conditions. The black solid line and gray shading area showing the one-sigma error band show the P$_V$-P$_{Pv}$ boundaries from our DFT (WC) and QMC computations respectively. The shading of QMC results indicates one-sigma statistic errors. The DFT C1, C2, and C3 curves are taken from Ref. [17], Ref. [27] and Ref. [29] respectively. The pressure in the D$''$ layer of the lower mantle falls between the vertical dash line and solid line of the core-mantle boundary. The experimental results are taken from Ref. [17]. The average geotherm is from Ref. [90].

FIG. 9. The thermal pressures averaged over volume from 27.8 to 44.5 Å$^3$/MgSiO$_3$ as a function of temperature.

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FIG. 10. Temperature and pressure dependence of the thermal expansion. (a) and (b) are for perovskite. (c) and (d) are for post-perovskite. The shading of the curves indicate one-sigma width of the uncertainties. The references for all the models and DFT calculations are [93](a,M1), [84](a,M2; c,M1; d,M11), [83](a,M3; b,M4), [74](a,M4); [25](a,LDA, [94](b,M1), [95](b,M2), [96](b,M3), [24](b,LDA, [79](c,M2; d,M2), [97](c,M3), and [29](c,LDA).
FIG. 11. Grüneisen ratio as a function of pressure. (a) is for perovskite and (b) is for post-perovskite. The shading of the curves indicate one-sigma width of the uncertainties. The references for all the models and DFT calculations are [98](a,M1 and M2), [29]LDA in a and b, and [79](b,M1).