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Thermodynamics of spin crossover in ferropericlase: an improved LDA + U_{sc} calculation

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Abstract

We present LDA + U_{sc} calculations of high-spin (HS) and low-spin (LS) states in ferropericlase (fp) with an iron concentration of 18.75%. The Hubbard parameter U is determined self-consistently with structures optimized at arbitrary pressures. We confirm a strong dependence of U on the pressure and spin state. Static calculations confirm that the antiferromagnetic configuration is more stable than the ferromagnetic one in the HS state, consistent with low-temperature measurements. Phonon calculations guarantee the dynamical stability of HS and LS states throughout the pressure range of the Earth mantle. Compression curves for HS and LS states agree well with experiments. Using a non-ideal mixing model for the HS to LS states solid solution, we obtain a crossover starting at \sim 45 GPa at room temperature and considerably broader than previous results. The spin-crossover phase diagram is calculated, including vibrational, magnetic, electronic, and non-ideal HS–LS entropic contributions. Our results suggest the mixed-spin state predominates in fp in most of the lower mantle.

1. Introduction

Ferropericlase (fp) is the second most abundant mineral in the Earth's lower mantle. It may be responsible for up to \sim 20 vol% of this region [1]. It is a solid solution (Mg_{1-x}Fe_x)O of MgO and FeO in the rocksalt-type (B1) crystal structure, with $x_{\rm Fe}=0.15-0.20$ in the lower mantle. Its high-pressure electronic properties, spin state, and phase stability are critical to understanding the properties of and processes taking place in the mantle. In particular, iron in fp undergoes a pressure-induced spin-crossover from a high spin (HS) state with S=2 to a low spin (LS) state with S=0. This spin-crossover has attracted extensive research interest because it can have critical geophysical consequences, e.g., a density increase [2], a bulk modulus softening [3], thermoelastic anomalies [4, 5], etc.

Experiments have reported an HS–LS crossover pressure in the range of 40-70 GPa at room or lower temperatures [6–10], with x_{Fe} around 0.20. High temperature leads to an increase in the spin-crossover pressure range and in the crossover onset pressure. This is caused by a mixed HS–LS state (MS) [2, 7, 11] caused entropic contributions. Because of the strongly correlated nature of the 3d electrons in Fe and the large supercells used to study the fp solid-solution by first-principles, the HS–LS crossover diagram has been challenging. Tsuchiya *et al* [2] first performed local density approximation (LDA) + U calculations to study the HS and LS crossover with HS/LS configuration entropy and magnetic entropy. Later [4] extended LDA + U calculations to include vibrational effects based on a virtual-crystal model. The model was also used to calculate thermodynamic anomalies in fp [2]. While LDA + U can reasonably address the electronic structure of the correlated 3d electrons of iron, its performance on the crossover pressure highly depends on the U value [12–17]. It has been argued that the complete dependence of U on pressure/volume, structure, spin state, or even pseudopotential should be taken into account if one is to make predictions of phase transitions at extreme conditions.

In this work, we employ a recent implementation of the self-consistent calculation of the Hubbard parameter based on density-functional perturbation theory (DFPT) [18]. Using LDA + U_{sc} calculations, we compute the spin crossover diagram for fp with $x_{Fe} = 0.1875$. This calculation differs from previous ones from our group by computing the HS and LS states' vibrational spectra vs volume and going beyond the ideal solid-solution model by computing the excess free energy due to HS–LS interaction in the MS state.

In the next section, we describe the computational details of the first-principle calculations. In section 3, we first analyze the electronic structure of fp with $x_{Fe} = 0.03125$. In section 4, we present static calculations of the spin-crossover of fp with $x_{Fe} = 0.1875$ at T = 0 K. In section 5, we compute the phonon spectra and consider the finite temperature effect on the spin crossover by including various entropic contributions and non-ideal mixing effects. We summarize all findings in section 6.

2. Methods

DFT + U calculations were performed using the simplified formulation of Dudarev *et al* [19] as implemented in the quantum ESPRESSO code [20, 21]. The LDA was used for the exchange-correlation functional with spin polarization. The projector-augmented wave (PAW) data sets from the high-accuracy version of PSlibrary [22] were employed with valence electronic configurations $3s^23p^63d^64s^2$, $2s^22p^63s^2$ and $2s^22p^4$ for Fe, Mg, and O, respectively. A kinetic-energy cutoff of 100 Ry for wave functions and 600 Ry for spin-charge density and potentials were used. In all cases, atomic orbitals were used to construct occupation matrices and projectors in the LDA + U scheme. A cubic supercell of B1 structure with 64 atoms was constructed for the current study, i.e., $(Fe_xMg_{1-x})_{32}O_{32}$. The $2 \times 2 \times 2$ k-point mesh was used for Brillouin zone integration. Structure optimization was performed by relaxing atom positions with a force convergence threshold of 0.01 eV/Å. The convergence threshold of all self-consistent field (SCF) calculations was 1×10^{-9} Ry.

The Hubbard correction [23] was applied to Fe-3d states. The Hubbard parameter U was computed using DFPQ [18] implemented in the quantum ESPRESSO code. The convergence threshold for the response function is 1×10^{-6} Ry. An automated iterative scheme was employed to obtain the self-consistent U_{sc} parameter while simultaneously optimizing the structure and desired spin state: starting from an empirical U of 4.3 eV, the energies of all possible occupation matrices for a spin state were computed. There are five possible occupation matrices corresponding to the HS state of ferrous iron with $3d^6$ configuration (S=2), while there are ten possibilities for the LS state (S = 0). The electronic configuration, i.e., occupation matrix, with the lowest energy, was selected for further structural optimization of lattice parameters and atomic positions. Then a new U parameter is recalculated for further structural optimization. The process continued until mutual convergence of structure and U is achieved for a convergence threshold of 0.01 eV for the U parameter and the convergence criteria mentioned above for structural optimizations. Only the lowest energy configuration was adopted in subsequent calculations. Finite temperature effects on the static DFT energy were included using the Mermin functional with the Fermi-Dirac smearing [24, 25]. The temperature-dependent electronic entropy was obtained from 0 to 4500 K every 500 K and then interpolated. The scheme used here was described in reference [26]. We also computed the excess free energy from non-ideal mixing of HS and LS states which is described together with the ideal solution model in section 5.

With large unit cells containing 64 atoms, phonon calculations were performed using the finite-displacement method, using Phonopy code [27] with force constants computed by quantum ESPRESSO. Vibrational density of states (VDOSs) were obtained using a q-point 20 \times 20 \times 20 mesh. The vibrational contribution to the free energy was calculated using the quasiharmonic approximation [28] with the qha code [29].

3. Electronic structure fp with $x_{Fe} = 0.031 25$ (fp3)

To first have a clear picture of the electronic structure of fp, we consider only one Mg substitution by Fe in the 64-atom supercell, i.e., FeMg₃₁O₃₂, $x_{Fe} = 0.031$ 25 (fp3 hereafter), as shown in figure 1(a). In this case, there is no Fe–Fe interaction so that the energy levels of the 3d orbitals in the ferrous Fe can be well-identified. Ferrous Fe (Fe²⁺) with 3d6 electronic configurations has six O neighbors in octahedral coordination. The octahedral crystal field splits the fivefold d-orbital degeneracy, producing a doublet with e_g symmetry and a triplet with t_{2g} symmetry. Because the t_{2g} orbitals are pointing away from the oxygen neighbors, the t_{2g} orbitals have lower energy than the e_g orbitals, shown in figure 1(c). In the HS state at low pressures, following Hund's rule, five of six electrons occupy five spin-up orbitals, and the remaining minority electron fills one of the t_{2g} orbitals, as shown in figure 1(a). The Fe–O octahedron is Jahn–Teller (JT) distorted in this electronic configuration, i.e., it has two short and four long bonds. At P = 0 GPa, the difference between the short and long bonds is 2.4% in figure 1(a). The JT distortion, in turn, causes further energy splitting within the e_g and t_{2g} levels so

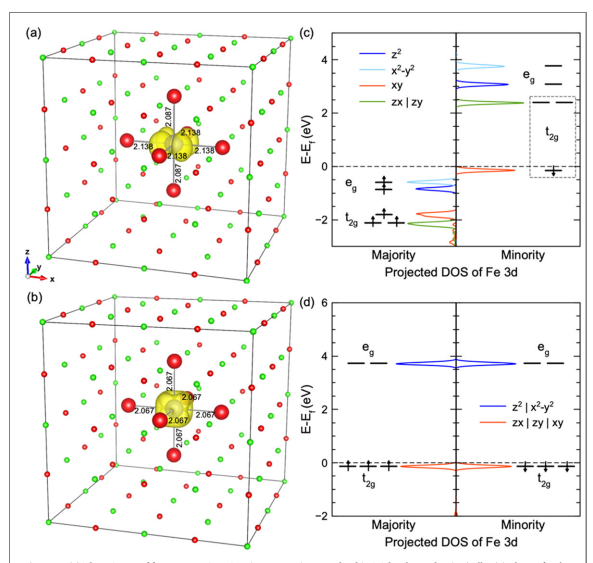


Figure 1. (a) The HS state of fp3 at P=0 GPa. Gray is Fe, green is Mg and red is O. The charge density (yellow) is shown for the Fe minority electron occupying the d_{xy} orbital. T Fe–O bond lengths in angstrom (Å) are shown inside the octahedron. Mg and O without Fe–O bonds are drawn smaller for clarity. (b) The LS state of fp3 at P=0 GPa. The charge density (yellow) is shown for the occupied t_{2g} orbitals. (c) and (d) Projected density of state (DOS) for Fe 3d orbitals in HS and LS fp3 at P=0 GPa. The schematic shows the energy splitting of e_g doublet and t_{2g} triplet.

that one can see a slight energy difference between d_z^2 and $d_x^2_{-y}^2$, as well as d_{xy} and d_{zx} (d_{zy}). Ferrous Fe can also exhibit the LS state with all six electrons in the t_{2g} orbitals, as shown in figure 1(d). Because the occupied t_{2g} orbitals have cubic symmetry, as shown in figure 1(b), there is no JT distortion in the equilibrated LS state. The volume of the FeO octahedron in the LS state is smaller than that in the HS state. Comparing the lattices at 0 GPa in figures 1(a) and (b), the octahedron volume of the LS is 7.4% smaller than the one with Fe²⁺ in the HS state. Both HS and LS of fp3 are insulators.

4. Spin crossover of fp18 at T = 0 K

We now perform static calculations to study the spin crossover in fp at T = 0 K. For a pyrolitic mantle composition, the fp volume fraction should be around 0.15–0.20 [1]. Here we focus on the spin crossover in the Fe₆Mg₂₆O₃₂ lattice, i.e., $x_{Fe} = 0.1875$ (fp18 hereafter), while some results on fp3 are also included for comparison. To construct the supercell structure, we distribute 6 Fe uniformly by occupying the face centers and edge centers, as shown in figure 2. Because fp is a solid solution of FeO and MgO, a uniform distribution is more relevant to the real situation. Moreover, it has been shown that different types of Fe configurations have only a small effect on the spin crossover [30]. With $x_{Fe} = 0.1875$, one would expect the exchange interaction between Fe ions in the HS state to be sizable because of the small Fe–Fe distance. Here we consider both the ferromagnetic (FM) and antiferromagnetic (AFM) configurations with spins aligned as in figures 2(a) and (b).

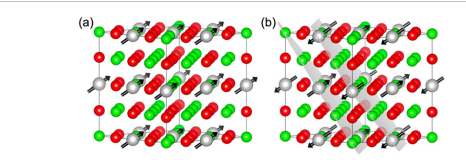


Figure 2. Supercell structure of fp18 with (a) FM and (b) AFM configurations. The [111] planes are indicated in (b). However, in our collinear spin calculations, the direction of the spin magnetic moment is not relevant.

Considering the AFM spin configuration in FeO-B1 [17], we assign opposite magnetic moments in the AFM configuration alternating in neighboring [111] planes.

Figure 3(a) shows the volume-dependent energy for fp18 HS states with FM and AFM configurations and LS state. The HS states have lower energy at large volumes (low or negative pressures) than the LS state. The AFM configuration has the lowest energy of all. Therefore, at ambient conditions, the ground state is the HS with AFM magnetic order. This is consistent with the experimental measurement that AFM is the ground state below the Néel temperature [10]. With decreasing volume, the LS state energy decreases w.r.t. to that of the HS states. This is mainly because the energy splitting between t_{2g} and e_g increases with increasing pressure (see supplementary figure S1 (https://stacks.iop.org/EST/4/014008/mmedia)), leading to the spin crossover. Figure 3(b) shows the self-consistent Hubbard parameters for the different spin states. The U_{sc} values of the LS state are systematically higher than the HS states regardless of volume. This trend is similar to fp3 and previous studies of Hubbard parameters of HS and LS states in the FeO system [17] (see supplementary figure S2). The self-consistent U value of HS-FM and HS-AFM also shows a slight difference. The energy-volume data are fitted by the third-order Birch-Murnaghan (BM) equation of state (EoS) using the least squares method. The fitted results are shown in the supplementary table S1. The enthalpies are obtained from the fitted BM-EoS and are shown in figure 3(c). Based on the enthalpy difference, the transition pressure from HS to LS can be identified. In fp18, the transition pressures are 60 GPa for the FM state and 66 GPa for the AFM state. By performing similar calculations with fp3, we find the HS-LS transition pressure in fp3 is 53 GPa. Therefore the spin crossover pressure increases with an increasing iron concentration in fp. This transition pressure in fp18 is in good agreement with the experimental measurement of fp with $x_{\text{Fe}} = 0.17$ at room temperature [8], which is ~60 GPa. A change in the distribution of iron in the supercell of fp18 changes the transition pressure by a few GPa only [29], but it gives rise to a distribution of transition pressures.

The fp18 compression curves obtained from the zero-kelvin EoS are shown in figure 4. The curves of HS-FM and HS-AFM states are almost overlapped here. The experimental measurement at room temperature with $x_{\text{Fe}} = 0.17$ fp [8] is also included for comparison. The calculated zero-kelvin compression curves are close but systematically smaller than the experimental data. This result is reasonable because no temperature effect is included yet. As discussed later, vibrational effects at finite temperature further improve the agreement with experiments.

Figure 5 shows the projected DOS of Fe-3*d* for fp18. Both HS and LS DOS are qualitatively similar to that of fp3 in figure 1. However, a larger Fe concentration caused stronger crystal field splitting, leading to complicated energy levels of different orbitals. Nevertheless, both HS and LS states remain to be the insulating states. The gaps are around 2 eV and almost independent of the pressure increasing.

5. Finite temperature effect on the spin crossover

Phonon calculations are performed with LDA + U_{sc} for all calculated volumes of both HS- and LS-fp18. Figure 6 shows examples of three VDOS from low to high pressures. All other VDOS are shown in supplementary figure S3. With increasing pressure, the phonon frequencies are shifted toward higher energies. No imaginary frequency is found in either HS or LS state up to 100 GPa. This is consistent with the recent phonon calculations in $x_{Fe} = 0.0625$ fp with density functional perturbation theory [30]. Therefore, both HS and LS states of fp with iron concentrations lower than 0.1875 are dynamically stable.

With the inclusion of vibrational entropy and electronic entropy described by the Mermin functional in static free energy calculations [24, 25], quasiharmonic calculations are performed to compute the free energy and EoS at finite temperatures. With the inclusion of thermal electronic excitation effects on the static free

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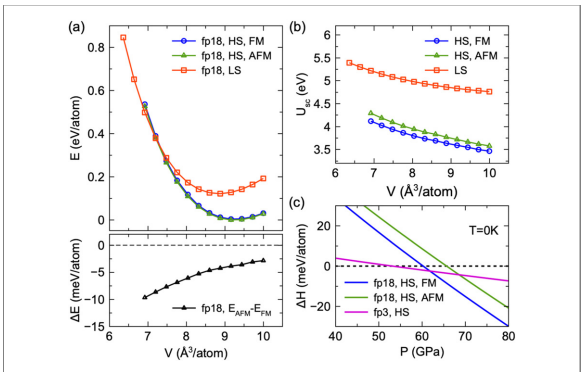
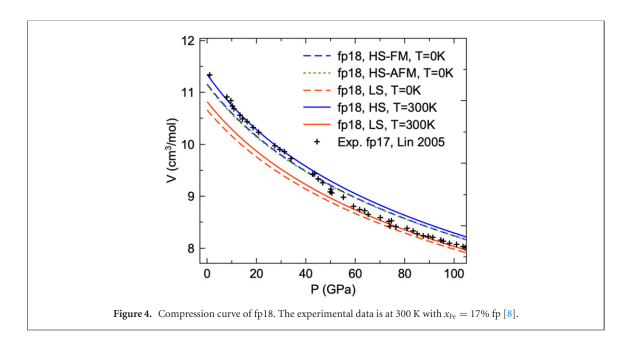


Figure 3. (a) Upper panel shows the E vs V curves for the three states of fp18. The lower panel shows the energy difference between the HS–AFM and HS–FM. (b) The self-consistent Hubbard parameters U for different spin states. (c) The enthalpy difference between HS and LS ($\Delta H = H_{\rm HS} - H_{\rm LS}$).



energy, the compression curve of both HS and LS at 300 K agrees very well with room-temperature experimental measurements [8], as shown in figure 4. To obtain the HS-LS phase boundary at finite temperature, we further consider the non-ideal mixing of HS and LS states and its contribution to the free energy. Using n to represent the fraction of LS states, the total free energy of an ideal solid solution of HS and LS can be written as

$$G_{\text{ideal}}(P, T, n) = (1 - n) G_{\text{HS}}(P, T) + nG_{\text{LS}}(P, T) + G_{\text{mix}}^{\text{ideal}}(n),$$
 (1)

where $G_{\rm HS/LS}$ is the molar Gibbs free energy of the pure HS/LS states ('/' indicates 'or'), i.e.,

$$G_{\mathrm{HS/LS}}(P,T) = G_{\mathrm{HS/LS}}^{\mathrm{stat+vib}}(P,T) + G_{\mathrm{HS/LS}}^{\mathrm{mag}},\tag{2}$$

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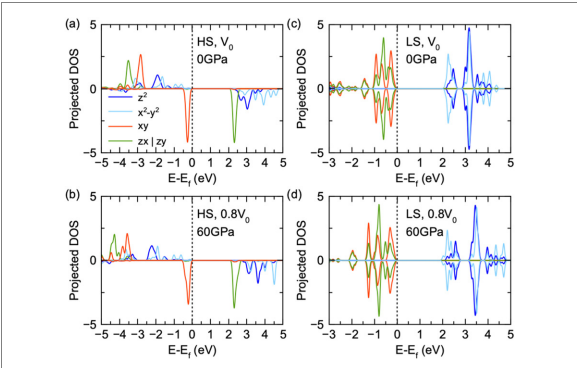


Figure 5. Projected density of state (DOS) of the Fe 3*d* orbitals in fp18 with (a) HS at 0 GPa; (b) HS at 60 GPa; (c) LS at 0 GPa; (d) LS at 60 GPa.

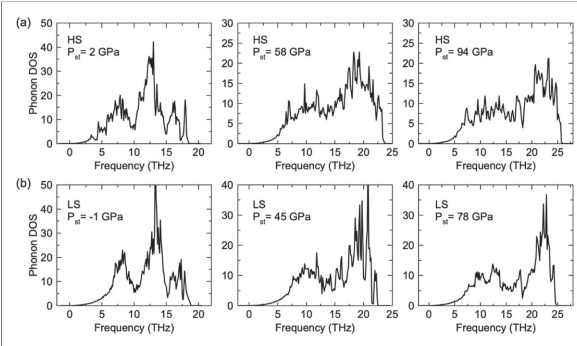


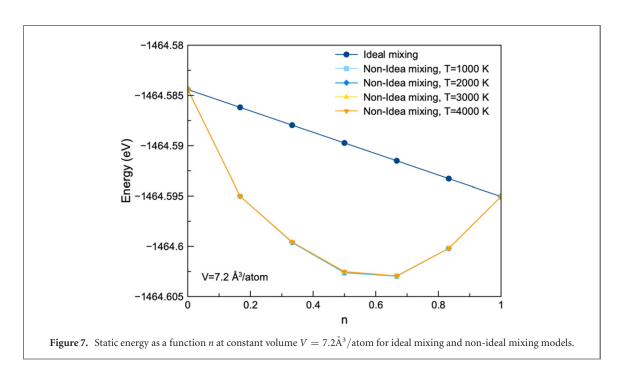
Figure 6. VDOS for (a) HS and (b) LS fp18 with three volumes. The upper and lower panels correspond to the same volume. Their static pressures at zero kelvin are indicated in the figure.

where $G_{\mathrm{HS/LS}}^{\mathrm{stat+vib}}\left(P,T\right)$ is the Gibbs free energy containing static and vibrational contribution and $G_{\mathrm{HS/LS}}^{\mathrm{mag}}$ is the magnetic contribution. $G_{\mathrm{HS/LS}}^{\mathrm{mag}}$ is a purely entropic contribution that one can estimate approximately as

$$G^{\text{mag}} = -k_{\text{B}} \mathsf{T} x_{\text{Fe}} \ln \left[m \left(2S + 1 \right) \right], \tag{3}$$

where *S* and *m* are the spin and electronic configuration (orbital) degeneracies of iron. In HS, S = 2 and m = 3. In LS, S = 0 and m = 1. The ideal free energy of mixing is given by the mixing entropy as

$$G_{\text{mix}}^{\text{ideal}}(n) = -TS_{\text{mix}} = -k_{\text{B}}Tx_{\text{Fe}} [n \ln n + (1-n)\ln(1-n)].$$
 (4)



Equation (4) gives the free energy of mixing of the ideal solution of HS and LS states, where n is the LS

fraction. By minimizing the free energy in equation (1) with respect to n, one obtains

$$n_{\text{ideal}} = \frac{1}{1 + m\left(2S + 1\right) \exp\left[\frac{\Delta G_{\text{LS-HS}}(P,T)}{k_{\text{B}}Tx_{\text{E}_{\text{A}}}}\right]},\tag{5}$$

where $\Delta G_{LS-HS}(P, T) = G_{LS}(P, T) - G_{HS}(P, T)$.

We now examine the effect of non-ideal mixing of HS and LS states. In this case, different HS/LS configurations are considered for a single Mg/Fe atomic arrangement in fp18 and several n values. The single Mg/Fe configuration in the $(Mg_{26}Fe_6)O_{32}$ supercell is sampled for $n=\frac{1}{6},\frac{2}{6},\frac{3}{6},\frac{4}{6}$ and $\frac{5}{6}$. These configurations are listed in supplementary table S2 and supplementary figure S4. The static energy ε_i of the ith non-equivalent atomic configuration is computed in a large pressure range, using the consistent Hubbard parameters of fp18 shown in figure 3(b). The non-ideal mixing energy can be obtained by making a Boltzmann ensemble average for all non-equivalent arrangements. Then

$$E_{\text{non-ideal}} = \sum_{i} g_i p_i \varepsilon_i, \tag{6}$$

where g_i is the multiplicity of the ith non-equivalent atomic configuration and p_i is the Boltzmann factor as $p_i = \frac{e^{\epsilon_i/k_{\rm B}T}}{\sum_i g_i e^{\epsilon_i/k_{\rm B}T}}$. Figure 7 shows that the results from non-ideal mixing deviate from the one in the ideal mixing model, indicating that the non-ideal mixing effect is relatively significant when the HS and LS states have similar static energies at the same volume. The excess energy can be obtained by calculating the energy difference between ideal and non-ideal models as $E_{\rm ex}(n) = E_{\rm non-ideal}(n) - E_{\rm ideal}(n)$. As can be seen in figure 7, the temperature dependence of $E_{\rm ex}(n)$ is insignificant. We then include an excess free energy term, $G_{\rm ex}(P,T,n)$, of non-ideal mixing in equation (1),

$$G_{\text{non-ideal}}(P, T, n) = G_{\text{ideal}}(P, T, n) + G_{\text{ex}}(P, T, n).$$

$$(7)$$

Here we assume the excess free energy is mainly contributed by the static part so that we keep the vibrational contributions the same as the one in the ideal mixing model, i.e. $G_{\rm ex}(P,T,n)\approx H_{\rm ex}(P,T,n)$, where the T-dependence is negligible as in $E_{\rm ex}(n)$. The excess enthalpy can be obtained by fitting $E_{\rm non-ideal}$ with 3rd BM-EOS, obtaining pressure and adding the $P_{\rm non-ideal}V$ term. Similar to solving the ideal mixing model, $n_{\rm non-ideal}$ can be obtained by minimizing equation (6) with respect to n, which leads to

$$f(P,T,n) = \Delta G_{\text{LS-HS}}(P,T) + \frac{\partial H_{\text{ex}}(P,T,n)}{\partial n} + k_{\text{B}} T x_{\text{Fe}} \ln \left[\frac{n}{1-n} \left(m \left(2S+1 \right) \right) \right] = 0.$$
 (8)

We *numerically* solve equation (8) for $n_{\text{non-ideal}}$ by first fitting $H_{\text{ex}}(n)$ with polynomial functions at each pressure and temperature, as shown in supplementary figure S5. The obtained $n_{\text{non-ideal}}(P, T)$ are plotted as a function of pressure and temperature in figure 8.

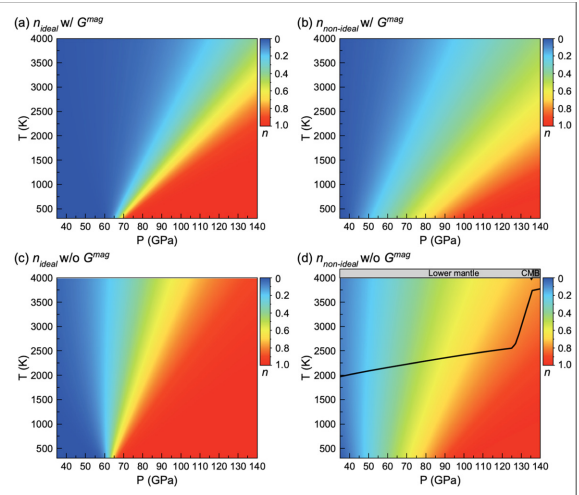


Figure 8. Temperature-dependent spin-crossover ranges with fp18 based on the different models. (a) and (b) are ideal and non-ideal mixing models with G^{mag} ; (c) and (d) are ideal and non-ideal mixing models without G^{mag} . The black line indicates the geotherm [31, 32].

We now analyze how the magnetic entropy contribution and non-ideal mixing affect the temperature dependence of the HS-LS spin-crossover pressures in figure 8. Comparing the results from ideal and nonideal mixing models (figure 8(a) vs 8(b), or 8(c) vs 8(d)), the HS-LS mixing range is much broader in the non-ideal mixing models. At room temperature, the crossover pressure range is \sim 5 GPa in the ideal mixing model, while it is ∼30 GPa in the non-ideal mixing model. This MS pressure range broadens further at higher temperatures. In principle, the non-ideal mixing model should be closer to the real situation than the ideal mixing model. The wide crossover range agrees better with the experimental data [7]. By comparing figures 8(a) and (c) (or figures 8(b) and (d)), we find G^{mag} significantly increases the Clapeyron slope of the spin crossover range. The slope of the phase boundary in figure 8(d) is more similar to the previous experimental data from reference [7] than others. This might occur because the current G^{mag} is an approximate analytical estimate of the largest possible contribution of the magnetic entropy. In reality, the local spin magnetic moment at high temperatures should not be as large as S = 2 for the HS state. Therefore, this magnetic entropic effect is likely overestimated. Taking these factors into account, figure 8(d) might represent the most realistic situation for the HS-LS spin cross in fp18. In figure 8(d), the spin crossover starts at ~ 45 GPa. High temperatures do not significantly affect the HS fraction at ~45 GPa, which agrees with the experimental data [7]. We include the mantle geotherm for a pyrolytic composition [31] in figure 8(d). The spin-crossover starts at \sim 45 GPa, 2100 K and ends at ~115 GPa, 2400 K along the geotherm. These results indicate the MS state should predominate in fp in most of the lower mantle. fp near the core-mantle boundary should be mainly in the LS state. We note that thermal electronic excitation effects described using the Mermin functional do not significantly affect the phase boundary, as shown in supplementary figure S6. This is expected as both HS and LS are insulators.

6. Summary

In summary, we revisited the HS–LS crossover in fp by performing LDA + U_{sc} calculations. The Hubbard parameters U are determined self-consistently (U_{sc}) using DFPT. The U_{sc} parameter depends on pressure, spin state, electronic and atomic configuration, etc, and varies by 1–2 eV. The AFM configuration is found to be the ground state at low temperatures, consistent with experiments. The energy difference between FM and AFM configurations is less than 10 meV/atom at T=0 K, and magnetic ordering has a relatively minor impact on the spin crossover pressure in static calculations for $x_{Fe}=0.1875$. Phonon spectra are computed for both HS and LS states. No imaginary frequencies are found in any case from 0 to 120 GPa, confirming phonon stability in fp in the entire pressure range of the Earth's mantle. Quasiharmonic free energy calculations offer *ab initio* compression curves for HS and LS states in good agreement with experimental data at room temperature. The HS–LS phase diagram is obtained by including all finite-temperature effects, i.e., vibrational, magnetic, electronic, and non-ideal HS–LS mixing contributions. The non-ideal HS–LS solid-solution mixing model gives a crossover starting at \sim 45 GPa at room temperature and is considerably broader than previous calculations. The magnetic entropy is found to affect the Clapeyron slope of the HS–LS crossover significantly. Considering these effects, the mixed spin state is predicted to predominate the fp throughout most of the lower mantle.

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Data availability statement

All data that support the findings of this study are included within the article (and any supplementary files).

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