Using forces to accelerate first-principles anharmonic vibrational calculations

Joseph C. A. Prentice\(^1\) and R. J. Needs\(^1\)

\(^1\)TCM Group, Cavendish Laboratory, University of Cambridge, J. J. Thomson Avenue, Cambridge, CB3 0HE, United Kingdom

(Dated: November 29, 2021)

High-level vibrational calculations have been used to investigate anharmonicity in a wide variety of materials using density-functional-theory (DFT) methods. We have developed a new and efficient approach for describing strongly-anharmonic systems using a vibrational self-consistent-field (VSCF) method. By far the most computationally expensive part of the calculations is the mapping of an accurate Born-Oppenheimer (BO) energy surface within the region of interest. Here we present an improved method which reduces the computational cost of the mapping. In this approach we use data from a set of energy calculations for different vibrational distortions of the materials and the corresponding forces on the atoms. Results using both energies and forces are presented for the test cases of the hydrogen molecule, solid hydrogen under high pressure including mapping of two-dimensional subspaces of the BO surface, and the \(\text{bcc}\) phases of the metals Li and Zr. The use of forces data speeds up the anharmonic calculations by up to 40%.

I. INTRODUCTION

Developing an accurate theoretical model for crystalline solids requires going beyond the static lattice model and including the effects of atomic vibrations. Usually this is achieved within the harmonic\(^{1,2}\) or quasiharmonic approximations - the latter includes thermal expansion by making the phonon frequencies volume dependent\(^{1,4}\). In many materials and situations these approximations work very well. However, they may be inaccurate if the vibrational amplitudes are large. Large amplitudes may occur, for example, in materials that contain light elements, materials close to a structural instability, or at high temperatures. Under such conditions it may be necessary to include anharmonic vibrations in the model. The mean-field vibrational-self-consistent-field (VSCF) method proposed by Monserrat, Drummond and Needs\(^5\) provides a sophisticated approach for computing the anharmonic vibrational wavefunction and energy of crystalline solids from first principles. This approach has been applied successfully in several systems\(^6\)–\(^10\).

Within the Born-Oppenheimer (BO) approximation\(^11\) the electronic and nuclear motion can be separated, giving two coupled Schrödinger equations. Application of the VSCF method consists of two main tasks - mapping of the BO energy surface (the “energy landscape”) that the nuclei move through, and self-consistently solving the VSCF equations for the anharmonic vibrational wavefunction and energy. Of these two tasks, the former is by far the most expensive, as it is usually accomplished by performing a large number of single-point density-functional-theory (DFT) calculations and fitting a functional form to the calculated energies. To ensure that the fitted form of the BO surface is accurate, it is necessary to converge the solution of the VSCF equations with respect to the number of mapping points used. Reducing the required number of mapping points reduces the computational cost of mapping the BO surface. The development of a method for increasing the accuracy of the fit for a given number of mapping points would help in reaching this goal.

Here we propose a scheme for improving the accuracy of the fitted functional form of the BO surface by making use of the force data generated by the DFT calculations in addition to the total electronic energy. Accurate forces are readily available within plane-wave basis DFT\(^12\). The gradient of the BO surface at each mapping point can be obtained from the forces, which provides additional data that can be used in the fitting process and therefore provides a better fit with negligible additional cost. We have investigated the effectiveness and cost of the VSCF method with forces (VSCF+f) and without forces (VSCF) for several systems in which anharmonicity is expected to play an important role - the hydrogen molecule, solid hydrogen at 100 GPa, and the \(\text{bcc}\) phases of lithium and zirconium, which exhibit soft modes. The effect of including both 1-D and 2-D subspaces of the BO surface in the calculations for solid hydrogen, corresponding to including coupling between anharmonic phonons, is also considered.

As the lightest element, hydrogen is an excellent test case for the VSCF+f method, as its vibrations have strong anharmonic character, even at low temperatures. Several \textit{ab initio} studies of high pressure phases of hydrogen, with and without anharmonic effects, have been conducted recently to determine the stable structure of the high pressure phases III and IV\(^{13-16}\). In particular, we use as test cases three structures from previous work, labelled by their space group and number of atoms in the primitive unit cell: \(\text{Cmca}-4\), \(\text{Cmca}-12\) and \(\text{C}2\text{}/\text{c}-24\). These are molecular phases arranged in layers; the \(\text{C}2\text{}/\text{c}-24\) phase is a candidate for phase III of hydrogen\(^{15}\). Previous work has shown that anharmonicity has a significant effect on the vibrational energy of these structures\(^{17,18}\). The strongly anharmonic character of the nuclear vibrations in hydrogen also implies that anharmonic effects that cannot be described by simple 1-D mapping of the BO surface could be significant. This provides an opportunity to test the ability of the VSCF+f method to reduce the computational cost of calculations including the mapping of 2-D subspaces of the BO surface. The mapping of 2-D subspaces is expensive in general, and therefore we have chosen the phase of solid hydrogen with the fewest atoms in its unit cell, \(\text{Cmca}-4\), as a test case for applying our improved method to mappings of 2-D subspaces of the BO surface.

Lithium and zirconium both have body-centered cubic (\(\text{bcc}\)) phases that are unstable at 0 K, and are stabilised at...
finite temperatures. This property is shared with other elements such as titanium and hafnium. In the case of Li, the structure at zero temperature is a close-packed \( R^9 \) structure, with the phase transition to \( bcc \) occurring at around 70 K\(^{19-21}\). Zirconium, on the other hand, has an hexagonal \( hcp \) structure at room temperature, with the transition to \( bcc \) occurring at around 1366 K\(^{22}\). In both materials, the \( bcc \) phase is unstable at low temperatures due to the presence of soft modes, which are dynamically stabilised at higher temperatures. Soft modes inherently have strong anharmonic character, and an accurate description of anharmonicity is required to treat these modes and the phase transition to the \( bcc \) structure. Several first principles vibrational studies including anharmonicity have been reported for Li, Zr and other similar elements\(^{19,22-27}\). The well-known presence of significant anharmonic effects in the \( bcc \) phase of these materials makes them suitable test cases for the VSCF+\( f \) method.

The rest of this work is organised as follows: In Sec. II we outline the VSCF method and describe our implementation of it. In Sec. III we describe the improved VSCF+\( f \) method and the testing of it, which is the focus of the rest of this work. In Sec. IV we compare results calculated using the VSCF+\( f \) and standard VSCF methods. Applications to molecular and high pressure solid hydrogen are reported in Sec. IV.A, mappings of 2-D subspaces of the BO surface in the \( Cmca-4 \) hydrogen phase in Sec. IV.B, and the \( bcc \) phases of Li and Zr in Sec. IV.C. Finally, in Sec. V we summarise our results. All equations are in Hartree atomic units, with \( \hbar = |e| = m_e = 4\pi\varepsilon_0 = 1 \).

II. THE VSCF METHOD

We work within the BO approximation in which the electronic and nuclear motions are separated out, which leads to the vibrational equation\(^5\)

\[
\left( -\frac{1}{2m_\alpha}\nabla^2 + E_{el}(\mathbf{R}) \right) \psi_{vib}(\mathbf{R}) = \hat{H}_{vib} \psi_{vib}(\mathbf{R}) = E_{vib} \psi_{vib}(\mathbf{R}).
\]

(1)

where \( p \) and \( \alpha \) label unit cells in the system and atoms within a unit cell respectively, with \( m_\alpha \) the mass of the \( \alpha \)\( \text{th} \) atom. \( \mathbf{R} \) is a collective vector of all the nuclear positions, \( \psi_{vib}(\mathbf{R}) \) is the vibrational wavefunction, and \( E_{vib} \) is the vibrational energy of the system. The electronic energy for a given \( \mathbf{R}, E_{el}(\mathbf{R}) \), acts as the potential in the vibrational Hamiltonian \( \hat{H}_{vib} \), and is generally known as the BO surface\(^5\).

Typically, this equation is solved approximately using the harmonic approximation by expanding \( E_{el}(\mathbf{R}) \) up to quadratic order in the displacement co-ordinates \( x_{p\alpha} = r_{p\alpha} - r_{p\alpha}^0 \). Here, \( r_{p\alpha} \) and \( r_{p\alpha}^0 \) are, respectively, the displaced and equilibrium nuclear positions. We calculate the harmonic potential by determining the matrix of force constants \( \frac{\partial^2 E_{el}(\mathbf{R}^0)}{\partial x_{p\alpha} \partial x_{p'\alpha'}^0} \), using a finite differences method\(^{28}\), and transforming to reciprocal space to give the dynamical matrix:

\[
D_{ia;ja'}(q) = \frac{1}{N_p \sqrt{m_\alpha m_\alpha'}} \sum_{p,p'} \frac{\partial^2 E_{el}(\mathbf{R}^0)}{\partial x_{p\alpha} \partial x_{p'\alpha'}^0} e^{i(q \cdot (\mathbf{R}_{p\alpha} - \mathbf{R}_{p'\alpha'}))}.
\]

(2)

\( \mathbf{R}_{p\alpha} \) is the position vector of the \( p^\text{th} \) unit cell, \( N_p \) is the number of unit cells in the system, and \( i, j \) run over the Cartesian directions. The eigenvectors, \( u_{nq;\alpha,i} \), and eigenvalues, \( \omega_{nq} \), of \( D_{ia;ja'}(q) \) can then be found, with \( n \) labelling the phonon branch index\(^5\).

\( \hat{H}_{vib} \) is then re-expressed in terms of harmonic normal or phonon co-ordinates given by:

\[
u_{nq} = \frac{1}{\sqrt{N_p}} \sum_{p,\alpha,i} \sqrt{m_\alpha} e^{-i(q \cdot \mathbf{R}_{p\alpha} - w_n i \alpha)}.
\]

Within the harmonic approximation, this gives a set of non-interacting simple harmonic oscillators of frequencies \( \omega_{nq} \). In the VSCF method, the principal axes approximation\(^{29}\) is used to include anharmonic effects. Assuming that anharmonicity is a perturbation to the harmonic approximation, and therefore a description of the BO surface in terms of many 1-D subspaces is a good approximation, we expand the BO surface as a series of \( N-D \) subspaces, using a basis given by the harmonic normal modes\(^5\),

\[
E_{el}(\mathbf{u}) = E_{el}(\mathbf{0}) + \sum_{n,q} V_{nq}(u_{nq}) + \frac{1}{2} \sum_{n,q,n',q'} V_{nq:n'q'}(u_{nq},u_{n'q'}) + \cdots
\]

(4)

\( \mathbf{u} \) is a collective mapping amplitude vector. The re-expression of \( E_{el}(\mathbf{R}) \) allows the mapping of the BO surface along the directions corresponding to the harmonic phonons and builds up an expression for \( E_{el}(\mathbf{R}) \), by means that will be discussed below. With this expression in hand, the energy can be minimised, using a Hartree product of 1-D states \( |\phi_{nq}(u_{nq})\rangle \) as a trial wavefunction. Solving the resulting VSCF equations gives a set of anharmonic vibrational eigenstates, with their associated energy eigenvalues and wavefunctions\(^5,30\). A perturbation theory can be constructed on these states, providing a further correction to the energy, and a partition function can be constructed, allowing the anharmonic free energy to be calculated at any finite temperature\(^5\).

III. IMPROVEMENTS TO THE VSCF METHOD

In order to solve the VSCF equations we require the form of \( E_{el}(\mathbf{u}) \). In previous work using this method, this is typically obtained using DFT to calculate \( E_{el}(\mathbf{u}) \) for various values of \( \mathbf{u} \) (i.e., various sets of atomic positions), and then fitting a functional form to these DFT energy results\(^{6,8,10}\). To find the form of the 1-D terms \( V_{nq}(u_{nq}) \), calculations are performed at several mapping points along the direction given by the appropriate harmonic phonon; to find the form of the 2-D terms
\[ V_{\text{anh}}(u_{nq}, u_{nq}', q_i), \] calculations must be done at points on a grid, and so on. However, these calculations rapidly become computationally expensive as we include more terms in the expansion of the BO surface, and therefore an increasingly large number of calculations is required. This restricts us to including very few or none of the 2-D or higher dimensional terms for all but the smallest systems.

Even when 2-D and higher terms are neglected, the mapping of the BO surface is by far the most computationally expensive part of using the VSCF method. In order to obtain an accurate fit to the BO surface, and therefore an accurate solution to the VSCF equations, it is necessary to converge the anharmonic correction to the energy, \( \Delta E_{\text{anh}} \), with respect to the number of mapping points used per mapping direction, which introduces further computational expense. Reducing the computational cost of the mapping is therefore extremely desirable in order to speed up the calculations. One way to reduce this cost would be to try and reduce the number of mapping points required to reach convergence, by utilising information other than the energy from the DFT calculations, such as the calculated forces on the atoms.

The gradient of the BO surface at a given value of \( \mathbf{u} \) can be calculated from the forces on the atoms in the corresponding atomic configuration

\[
\frac{\partial E_\alpha(\mathbf{u})}{\partial u_{n\alpha}} = -\sum_{p1;\mathbf{q}} \frac{1}{\sqrt{N_p}} c_\mathbf{q}^p \mathbf{R}_p \cdot \mathbf{w}_{q1;\mathbf{c}_\alpha}, \tag{5}
\]

where \( f_{p1;\mathbf{q}} \) is the force on the atom labelled by \( p, \alpha \) in Cartesian direction \( i \). Information about the forces can then be fed into the fitting procedure, which then increases the accuracy of the fit. In previous work, the DFT energy values have typically been fitted using a cubic spline\(^5\). In this work we investigate using cubic and quintic splines in the fitting procedure, to see whether either generally produces a more accurate fit. Utilising a seventh order (heptic) spline was also tested, but this approach suffered from overfitting and performed significantly worse than the cubic and quintic splines. For this reason, heptic splines will not be discussed further in this work.

Utilising forces to improve the mapping of the 1-D terms in the expansion of the BO surface in equation (4) is a simple extension of the usual splining procedure, but the task becomes more complex when trying to improve the mapping of the higher dimensional terms. The only higher dimensional terms considered here are 2-D. A functional form for the BO surface is found by using a series of 1-D splines along one direction, with the co-ordinate \( u_1 \), and then using the results to fit a spline along the second direction, with the co-ordinate \( u_2 \). Typically this would require knowledge of the cross derivative \( \frac{\partial^2 E_\alpha(\mathbf{u})}{\partial u_1 \partial u_2} \) at each sampling point\(^31\). We avoid this requirement by using a simple cubic spline to obtain the form of the function \( \frac{\partial E_\alpha(\mathbf{u})}{\partial u_1} \mid_{x_2} (u_1) \). This represents the gradient along the direction defined by \( u_2 \) as a function of \( u_1 \), for a fixed value of \( u_2 = x_2 \). Obtaining this function allows forces to be used in the fit along \( u_2 \). To ensure the accuracy of the calculations including 2-D terms, we converge the correction to the anharmonic vibrational energy due to these terms, \( \Delta E_{2-D} \), with respect to the number of mapping points used per direction.

Although including the force data in the fitting process should reduce the number of calculations required to obtain convergence, the accuracy of the forces themselves must be considered. If a variational method is used to minimise the total energy in the DFT calculations, the energy itself will be correct to second order errors in the charge density. However, the error in the forces is instead linear with the error in the charge density\(^12\), and calculations must be converged to within a strict tolerance to obtain accurate forces. This requirement could potentially cancel out the reduction in computational cost gained by reducing the number of mapping points if the convergence tolerance is too strict, and our results include tests to determine whether this is true or not. These tests showed that this issue did not negatively affect the speed-up obtainable with the VSCF+f method, with both methods breaking down at the same level of energy convergence.

IV. RESULTS

All DFT calculations were performed using version 8.0 of the plane-wave DFT code CASTEP\(^32\) and ultrasoft pseudopotentials\(^33\) generated `on-the-fly`. Throughout, the ratio between the fast Fourier transform (FFT) grid used for the electronic density and that used for the Kohn-Sham states (the ‘grid scale’ parameter in CASTEP) is 2.0. The local density approximation (LDA) was used for the exchange-correlation functional in the hydrogen calculations\(^34\), while the PBE functional was used for lithium and zirconium\(^35\). Previous work on solid hydrogen has shown that, while the exact quantitative results of DFT calculations are strongly dependent on the choice of functional, the qualitative results are similar for most functionals\(^36\), and that the LDA is a reasonable choice for the purpose of tests on solid hydrogen. The PBE functional has been used successfully in several previous studies of lithium, zirconium and other similar materials\(^19\,24\). Calculations of the harmonic normal modes and energy were performed by determining the matrix of force constants via a finite differences method\(^28\), before diagonalization of the dynamical matrix. Atomic displacements of 0.01 bohr were used.

A. Hydrogen

As the lightest element, hydrogen is a good material in which to test the VSCF+f method. Its vibrational motions explore the BO surface out to large amplitudes due to its low mass, which introduces significant anharmonic character. The method was first applied to molecular hydrogen before moving onto solid hydrogen at 100 GPa. All hydrogen calculations were performed at zero temperature and only the zero-point energy was considered.

A plane-wave cut-off energy of 800 eV was used for the calculations on molecular hydrogen, with a \( 5 \times 5 \times 5 \) Monkhorst-Pack k-point grid\(^37\). As CASTEP uses periodic boundary con-
FIG. 1: Convergence of anharmonic correction to the energy at 0 K for H₂, \( \Delta E_{\text{anh}} = E_{\text{anh}} - E_{\text{har}} \), with respect to the number of mapping points used per mapping direction, for the basic VSCF method, as well as the VSCF+f method, fitting with both cubic and quintic splines. The inset shows the results at low numbers of mapping points on a different energy scale.

It is necessary to make the unit cell large enough to prevent the molecule from interacting with its periodic images. The frequencies of the harmonic phonon modes were converged with respect to the size of the cubic unit cell, resulting in a converged lattice constant of 8 Å. The distance between the atoms was allowed to relax, using the Broyden-Fletcher-Goldfarb-Shanno (BFGS) method\(^{18}\) to converge the forces on the atoms to within 0.001 eV Å\(^{-1}\), before single-point energy calculations were conducted.

Once the harmonic calculations were completed, a range of different numbers of mapping points per direction from 7 to 27 were used to map the BO surface of the hydrogen molecule. The performance of the basic VSCF method was compared to the VSCF+f method using both cubic and quintic splines in the fitting process. This was repeated with the energy convergence tolerance of the calculations set to \(10^{-10}, 10^{-6}, 10^{-4}\) and \(10^{-2}\) eV per SCF cycle, to test whether this affected the accuracy of the forces, and therefore the accuracy of the VSCF+f fitting relative to the normal fitting procedure. Fig. 1 shows the convergence of the anharmonic correction to the zero point energy, \( \Delta E_{\text{anh}} = E_{\text{anh}} - E_{\text{har}} \), with respect to the number of mapping points for the three different methods. Here the energy convergence tolerance was set to \(10^{-6}\) eV per SCF cycle. It can be seen that including the forces in the fitting process significantly improves the convergence of \( \Delta E_{\text{anh}} \), with the quintic spline fit performing even better than the cubic spline. This suggests that including forces in the fitting process can significantly improve the efficiency of the VSCF method, and that utilising a quintic spline allows fitting of the BO surface even more accurately for the same number of DFT calculations, especially for low numbers of mapping points. An almost identical set of results was found for energy convergence tolerances from \(10^{-4}\) eV up to \(10^{-10}\) eV per SCF cycle, with the VSCF+f method converging more rapidly with the number of mapping points. For an energy convergence tolerance of \(10^{-2}\) eV per SCF cycle, both the VSCF and VSCF+f methods failed to converge with less than 27 mapping points per direction. This shows that for a range of energy convergence tolerances, the VSCF+f fitting method is still able to outperform the basic VSCF method and map the BO surface accurately at a lower computational cost. Using a quintic spline improves the quality of the fit still further.

With the results from the hydrogen molecule in mind, we turn our attention to the case of high pressure solid hydrogen. Three different structures of solid hydrogen were considered at a pressure of 100 GPa - \( \text{Cmca}-4, \text{Cmca}-12 \) and \( \text{C2/c}-24 \), as previously described. Fig. 2 gives a view of these three structures, showing that they are molecular in nature. These structures have all been studied previously with DFT over a range of pressures\(^{13}\). Again, once harmonic calculations were completed, the convergence of the anharmonic correction to the zero point energy per atom with respect to the number of mapping points was calculated for a single unit cell of each structure. An plane-wave cut-off energy of 1000 eV and an energy convergence tolerance of \(10^{-6}\) eV per SCF cycle was used throughout, with Monkhorst-Pack grids of size \(28 \times 28 \times 16, 18 \times 18 \times 4 \) and \(16 \times 8 \times 16\) for the \(\text{Cmca}-4, \text{Cmca}-12 \) and \( \text{C2/c}-24 \) structures respectively. Fig. 3 shows the convergence of \( \Delta E_{\text{anh}} \) per atom for all three structures for the basic VSCF and the improved VSCF+f quintic spline methods. Again, convergence was reached with fewer numbers of mapping points per direction using the VSCF+f method than with the VSCF method, especially for the \( \text{Cmca}-12 \) and \( \text{C2/c}-24 \) structures. In the latter case, including forces in the fitting reduces the computational cost by around 40%.

This further implies that the VSCF+f method is robust and improves on the efficiency of the basic VSCF method.

B. Two-dimensional subspaces

A natural next test of the VSCF+f method is to apply it to mapping 2-D subspaces of the BO surface. This poses a more significant challenge than the 1-D terms considered up to now, as interpolating data in two dimensions is required.

Mapping the BO surface in two dimensions is expensive, and can only feasibly be done for small systems. Here, we focus on the \( \text{Cmca}-4 \) structure of solid hydrogen discussed previously, which possesses twelve potential mapping directions, corresponding to the twelve harmonic phonon modes, labelled with numbers from 1 to 12. Three of these modes are acoustic, and thus have zero frequency. To minimise the computational cost further, we consider only two of the many 2-D subspaces in this system – those corresponding to the directions described by the harmonic modes 4 and 5, and 4 and 7. The harmonic frequencies of modes 4, 5, and 7 are 69.4, 74.0 and 114 meV, respectively, and the displacement patterns corresponding to each of these mapping directions can be found in the Supplemental Material. These subspaces were chosen by conducting a preliminary mapping of all 2-D subspaces with a low number of mapping points, and taking only those with significant corrections to the 1-D description of the BO

\(^{18}\) H. F. van Duijn, M. J. Frisch, C. C. van Lenthe, J. P. Perdew, and J. A. Pople, J. Chem. Phys. 96, 6517 (1992).

\(^{13}\) K. Okada, Y. M. Gao, H. Nomura, and A. Okazaki, J. Phys. Chem. B 120, 5125 (2016).
surface. The subspaces where the mapping entered parts of energy minima corresponding to structures significantly lower in energy than the Cmca-4 structure were also neglected. The two subspaces presented here were then chosen as being representative of those remaining. The same cut-off energy and Monkhorst-Pack grid was used as in the calculations of Sec. IV.A, but an energy convergence tolerance of $10^{-10}$ eV was used to ensure accurate forces.

Fig. 4 shows the results of tests including the mapping of 2-D subspaces in the Cmca-4 solid hydrogen structure. The two rows of figures correspond to the two subspaces mapped. The left-hand column shows the BO surface mapped in the relevant subspace, and the right-hand column shows the convergence of the correction to the energy due to 2-D terms $\Delta E_{2-D}$ with respect to the number of mapping points used per mapping direction. All energies were again calculated at zero temperature. The convergence graphs show that utilising forces in the mapping of the BO surface in two dimensions improves the results relative to the converged final value, especially for small numbers of mapping points, although the improvement is not as pronounced as in the 1-D case. This could be due to the small size of the energy scales in question – the energies shown are all smaller than 1 meV per atom, which is around the finest energy scale that such anharmonic calculations can reasonably be assumed to be accurate to. The small size of the corrections due to 2-D terms compared to those seen for 1-D terms shows that the neglect of such higher-order terms in the BO surface expansion of equation (4) is justified. The ability of the VSCF+f method to show improvement even at such small energy scales again demonstrates its capabilities, even in cases including mapping of 2-D subspaces of the BO surface.

C. Lithium and zirconium

Finally, we consider two systems in a very different regime from molecular or solid hydrogen - the metals lithium and zirconium, in their bcc phases. The main anharmonic contributions to the vibrational energy in these systems arise from soft modes, which necessarily have some quartic character. Here, we focus our attention on mapping the BO surface along the direction defined by these soft modes, and how utilising the VSCF+f procedure can improve this mapping. In addition to calculations at zero temperature, the effect of anharmonicity on the free energy at finite temperature is also considered, as the bcc phase only becomes stable experimentally at 70 K in Li and 1366 K in Zr. Thermal expansion effects are not included in these calculations.

For the calculations in lithium and zirconium, we used the recently introduced non-diagonal supercells method to reduce the computational cost of sampling the vibrational Brillouin zone. Typically, diagonal supercells are used to sample the vibrational BZ, with an $N \times N \times N$ sampling grid requiring an $N^3$ unit cells. The

FIG. 2: Views of the structures of solid hydrogen considered at 100 GPa. Both of the Cmca structures are shown looking along the $x$-axis, while the C2/c-24 structure is shown looking along the $y$-axis. As these are layered structures, atoms in inequivalent layers are denoted by different colours. Green, purple, translucent green and translucent purple denote the first, second, third and fourth inequivalent layers,

FIG. 3: Convergence of the anharmonic correction to the energy at 0 K, $\Delta E_{anh}$, of the Cmca-4, Cmca-12 and C2/c-24 phases of solid hydrogen at 100 GPa, with respect to the number of mapping points used per mapping direction. The convergence of the basic VSCF method as well as the VSCF+f method of fitting with a quintic spline are shown. The inset shows the results at low numbers of mapping points on a different energy scale.
non-diagonal supercells method allows such an $N \times N \times N$ sampling to be done using a series of non-diagonal supercells each containing a maximum of $N$ unit cells, significantly reducing the computational cost. However, as the aim of this work is to consider methods for improved fitting of the BO surface, rather than to conduct high-accuracy calculations on these well-studied materials, we did not attempt to completely converge our results with respect to the sampling of the vibrational BZ. Even with the non-diagonal supercell method the computational cost increases rapidly with increasing sampling grid size. Instead, we used a $8 \times 8 \times 8$ sampling of the vibrational BZ as a compromise between accuracy and speed.

As lithium and zirconium are metallic, it is necessary to use partial band occupancies to eliminate discontinuities in the energy during the SCF minimisation. This is done by artificially giving the Kohn-Sham quasiparticles a finite temperature to smear out their energy levels. In these calculations, a smearing width of 0.2 eV, corresponding to 2320 K, is used. The calculated DFT energies do depend on the size of this smearing, but the effect on the overall shape of the BO surface is small, meaning the results of the vibrational calculations are largely unaffected by the choice of smearing width. The Monkhorst-Pack grids in all supercells used had a spacing of 0.025 Å$^{-1}$, corresponding to a $16 \times 16 \times 16$ grid in the unit cell. An energy cut-off of 1500 eV was used in all calculations. To obtain these parameter values, the harmonic vibrational energy was converged to within 1 meV with respect to the Monkhorst-Pack grid size and energy cut-off. This constitutes a somewhat stronger convergence criterion than that used in other work$^{46}$, as it is important to ensure the harmonic results are accurate before using them as a basis for further calculations.

Our results for Li show that any differences in the anharmonic vibrational energy arising from the three different fitting methods are negligible, down to scales of 0.1 meV. Because of this we do not present the results for Li here, although we do discuss them. The negligible difference between fitting methods arises from the fact that, apart from along the direction defined by the soft modes present, the vibrational properties of Li are described well by the harmonic approximation, despite its low mass$^{41}$. This means that a very good fit to the BO surface along most modes can be found with small numbers of mapping points (as in the standard finite displacement method for calculating harmonic frequencies), and so all three methods agree very well. Even in the case of the soft modes, which must necessarily contain some anharmonic character, the double-well structure is not very pronounced, with the overall BO surface appearing essentially quadratic. To see this double-well structure, where the two minima are very close to
FIG. 5: Convergence with respect to the number of mapping points used per mapping direction of contribution of 1-D terms corresponding to the total anharmonic vibrational energy of the bcc phase of zirconium at 0 K. The convergence of the basic VSCF method as well as the VSCF+f method using both a cubic spline and a quintic spline are shown.

FIG. 6: The shape of the BO surface as mapped along one of the soft modes of zirconium; 'puc' stands for 'per unit cell'.

The soft modes in Zr are more numerous, and mapping the BO surface along the directions defined by them gives much more pronounced double-well structures than in Li, meaning that the structure is not dynamically stable at zero temperature. An example of the pronounced double-well structure of the BO surface mapped along one of the soft modes in Zr is shown in Fig. 6. Our results can be used to calculate the temperature at which the bcc phase is stabilised dynamically, by calculating the internal energy at a range of temperatures and finding where it becomes positive. The results are shown in Fig. 7 for a range of mapping points and the three fitting methods used. At these higher temperatures, it can be seen that the differences between the three fitting methods, visible at zero temperature in Fig. 5, are much less significant – the differences are mostly washed out by the overall vibrational energy increasing. Our calculations predict that the bcc structure of Zr should become dynamically stable above about 520 K, which is significantly lower than the observed transi-
tion temperature of 1366 K. This disagreement could again potentially be caused by the incomplete convergence with respect to the vibrational BZ sampling, higher-order terms in the BO surface expansion, thermal expansion effects, or errors inherent in DFT itself.

V. CONCLUSIONS AND FUTURE WORK

In summary, we have shown that the efficiency of the vibrational self-consistent field method proposed in Ref. 5 can be significantly improved by using both energy and force data from DFT calculations when mapping the BO energy surface. Tests of this method on molecular and high-pressure solid hydrogen, including the contribution of 2-D subspaces of the BO surface to the energy, as well as on lithium and zirconium in the bcc structure, show that the VSCF+f method agrees well with the basic VSCF method, but significantly reduces the computational cost involved.

Our results show that the VSCF+f method allows us to perform accurate calculations of anharmonic corrections to the harmonic phonon model for a significantly reduced computational cost compared to the basic VSCF method. The next step will be to apply the VSCF+f method to new systems in which large anharmonicities are of interest.

VI. ACKNOWLEDGEMENTS

The authors would like to thank Bartomeu Monserrat for useful discussions and comments on the manuscript. Calculations were performed on the Cambridge High Performance Computing Service facility and the ARCHER facility of the UK National Supercomputing Service. R.J.N. acknowledges financial support from the Engineering and Physical Sciences Research Council (EPSRC) of the U.K. under Grant No. EP/J017639/1. Computational resources were provided by the Archer facility of the U.K.’s national high-performance computing service (for which access was obtained via the UKCP consortium, EPSRC Grant No. EP/K014560/1). Supporting research data may be freely accessed at https://doi.org/XX.XXXXX/XXX.XXXX, in compliance with the applicable Open Data policies.

---

1 D.C. Wallace, *Thermodynamics of Crystals* (John Wiley & Sons, New York, 1972).
2 A.A. Maradudin, E.W. Montroll, G.H. Weiss, and I.P. Ipatova, *Theory of Lattice Dynamics in the Harmonic Approximation*, 2nd ed. (Academic Press, New York, 1971).
3 P. Pavone, K. Karch, O. Schütt, D. Strauch, W. Windl, P. Giannozzi, and S. Baroni, “Ab initio lattice dynamics of diamond,” *Phys. Rev. B* **48**, 3156–3163 (1993).
4 S. Baroni, S. de Gironcoli, A. Dal Corso, and P. Giannozzi, “Phonons and related crystal properties from density-functional perturbation theory,” *Rev. Mod. Phys.* **73**, 515–562 (2001).
5 B. Monserrat, N.D. Drummond, and R.J. Needs, “Anharmonic vibrational properties in periodic systems: energy, electron-phonon coupling, and stress,” *Phys. Rev. B* **87**, 144302 (2013).
6 B. Monserrat, N.D. Drummond, C.J. Pickard, and R.J. Needs, “Electron-phonon coupling and the metallization of solid helium at terapascal pressures,” *Phys. Rev. Lett.* **112**, 055504 (2014).
7 B. Monserrat, R.J. Needs, and C.J. Pickard, “Temperature effects in first-principles solid state calculations of the chemical shielding tensor made simple,” *J. Chem. Phys.* **141**, 134113 (2014).
8 N.D. Drummond, B. Monserrat, J.H. Lloyd-Williams, P. López Ríos, C.J. Pickard, and R.J. Needs, “Quantum Monte Carlo study of the phase diagram of solid molecular hydrogen at extreme pressures,” *Nat. Commun.* **6**, 7794 (2015).
9 E.A. Engel, B. Monserrat, and R.J. Needs, “Vibrational effects on surface energies and band gaps in hexagonal and cubic ice,” *J. Chem. Phys.* **145**, 044703 (2016).
10 J.C.A. Prentice, B. Monserrat, and R.J. Needs, “First-principles study of the dynamic Jahn-Teller distortion of the neutral vacancy in diamond,” *Phys. Rev. B* **95**, 014108 (2017).
11 M. Born and R. Oppenheimer, “Zur Quantentheorie der Molekeln,” *Ann. Phys.* **389**, 457–484 (1927).
12 M.C. Payne, M.P. Teter, D.C. Allan, T.A. Arias, and J.D. Joannopoulos, “Iterative minimization techniques for ab initio total-energy calculations: molecular dynamics and conjugate gradients,” *Rev. Mod. Phys.* **64**, 1045–1097 (1992).
13 C.J. Pickard, M. Martinez-Canales, and R.J. Needs, “Density functional theory study of phase IV of solid hydrogen,” *Phys. Rev. B* **85**, 214114 (2012).
14 J. Chen, X. Ren, X.-Z. Li, D. Alifè, and E. Wang, “On the room-temperature phase diagram of high pressure hydrogen: An ab initio molecular dynamics perspective and a diffusion Monte Carlo study,” *J. Chem. Phys.* **141**, 024501 (2014).
15 C.J. Pickard and R.J. Needs, “Structure of phase III of solid hydrogen,” *Nat. Phys.* **3**, 473–476 (2007).
16 K.A. Johnson and N.W. Ashcroft, “Structure and bandgap closure in dense hydrogen,” *Nature* **403**, 632–635 (2000).
17 M. Borinaga, I. Errea, M. Calandra, F. Mauri, and A. Bergara, “Anharmonic effects in atomic hydrogen: Superconductivity and lattice dynamical stability,” *Phys. Rev. B* **93**, 174308 (2016).
18 B. Monserrat, R.J. Needs, E. Gregoryanz, and C.J. Pickard, “Hexagonal structure of phase III of solid hydrogen,” *Phys. Rev. B* **94**, 134101 (2016).
19 O. Hellman, I.A. Abrikosov, and S.I. Simak, “Lattice dynamics of anharmonic solids from first principles,” *Phys. Rev. B* **84**, 180301 (2011).
20 A.W. Overhauser, “Crystal structure of lithium at 4.2 K,” *Phys. Rev. Lett.* **53**, 64–65 (1984).
21 A.M. Schaeffer, W. Cai, E. Olejnik, J.J. Molaison, S. Sinogeikin, A.M. dos Santos, and S. Deemyad, “Boundaries for martensitic transition of Li under pressure,” *Nat. Commun.* **6**, 8030 (2015).
22 G. Grimvall, B. Magyari-Köpe, V. Ozoliņš, and K.A. Persson, “Lattice instabilities in metallic elements,” *Rev. Mod. Phys.* **84**, 945–986 (2012).
23 N. Antolin, O.D. Restrepo, and W. Windl, “Fast free-energy calculations for unstable high-temperature phases,” *Phys. Rev. B* **86**, 054119 (2012).
24 P. Souvatzis, O. Eriksson, M.I. Katsnelson, and S.P. Rudin, “Entropy driven stabilization of energetically unstable crystal structures explained from first principles theory,”
25. W. Petry, A. Heiming, J. Trampenau, M. Alba, C. Herzig, H.R. Schober, and G. Vogl, “Phonon dispersion of the bcc phase of group-IV metals. I. bcc titanium,” Phys. Rev. B 43, 10933–10947 (1991).

26. A. Heiming, W. Petry, J. Trampenau, M. Alba, C. Herzig, H.R. Schober, and G. Vogl, “Phonon dispersion of the bcc phase of group-IV metals. II. bcc zirconium, a model case of dynamical precursors of martensitic transitions,” Phys. Rev. B 43, 10948–10962 (1991).

27. J. Trampenau, A. Heiming, W. Petry, M. Alba, C. Herzig, W. Miekeley, and H.R. Schober, “Phonon dispersion of the bcc phase of group-IV metals. III. bcc hafnium,” Phys. Rev. B 43, 10963–10969 (1991).

28. K. Kunc and R.M. Martin, “Ab Initio force constants of GaAs: A new approach to calculation of phonons and dielectric properties,” Phys. Rev. Lett. 48, 406–409 (1982).

29. J. O. Jung and R. B. Gerber, “Vibrational wave functions and spectroscopy of (H$_2$O)$_n$, $n = 2, 3, 4, 5$: Vibrational self-consistent field with correlation corrections,” J. Chem. Phys. 105, 10332 (1996).

30. J.M. Bowman, “Selfconsistent field energies and wavefunctions for coupled oscillators,” J. Chem. Phys. 68, 608–610 (1978).

31. W.H. Press, S.A. Teukolsky, W.T. Vetterling, and B.P. Flannery, Numerical Recipes in FORTRAN: The Art of Scientific Computing, 2nd ed. (Cambridge University Press, Cambridge, 1992).

32. S.J. Clark, M.D. Segall, C.J. Pickard, P.J. Hasnip, M.I.J. Probert, Keith Refson, and M.C. Payne, “First principles methods using CASTEP,” Z. Kristallogr. 220, 567–570 (2005).

33. D. Vanderbilt, “Soft self-consistent pseudopotentials in a generalized eigenvalue formalism,” Phys. Rev. B 41, 7892–7895 (1990).

34. W. Kohn, A.D. Becke, and R.G. Parr, “Density functional theory of electronic structure,” J. Phys. Chem. 100, 12974–12980 (1996).

35. J.P. Perdew, K. Burke, and M. Ernzerhof, “Generalized gradient approximation made simple,” Phys. Rev. Lett. 77, 3865–3868 (1996).

36. S. Azadi and W.M.C. Foulkes, “Fate of density functional theory in the study of high-pressure solid hydrogen,” Phys. Rev. B 88, 014115 (2013).

37. H.J. Monkhorst and J.D. Pack, “Special points for Brillouin-zone integrations,” Phys. Rev. B 13, 5188–5192 (1976).

38. J. Nocedal and S.J. Wright, Numerical Optimization, 2nd ed. (Springer-Verlag, Berlin, 2006).

39. J.H. Lloyd-Williams and B. Monserrat, “Lattice dynamics and electron-phonon coupling calculations using nondiagonal supercells,” Phys. Rev. B 92, 184301 (2015).

40. Kurt Lejaeghere, Gustav Bihlmayer, Torbjörn Björkman, Peter Blaha, Stefan Blügel, Völker Blum, Damien Caliste, Ivano E. Castelli, Stewart J. Clark, Andrea Dal Corso, Stefano de Gironcoli, Thierry Deutsch, John Kay Dewhurst, Igor Di Marco, Claudia Draxl, Marcin Dulak, Olle Eriksson, José A. Flores-Livas, Kevin F. Garrity, Luigi Genovese, Paolo Giannozzi, Matteo Giantomassi, Stefan Goedecker, Xavier Gonze, Oscar Gránás, E. K. U. Gross, Andris Gulans, François Gygi, D. R. Hamann, Phil J. Hasnip, N. A. W. Holzwarth, Diana Iuşan, Dominik B. Jochem, François Jollet, Daniel Jones, Georg Kresse, Klaus Koepernik, Emine Küçükbenli, Yaroslav O. Kvashnin, Inka L. M. Locht, Sven Lubeck, Martijn Marsman, Nicola Marzari, Ulrike Nitzsche, Lars Nordström, Taisuke Ozaki, Lorenzo Paulatto, Chris J. Pickard, Ward Poelman, Matt I. J. Probert, Keith Refson, Manuel Richter, Gian-Marco Rignanese, Santanu Saha, Matthias Scheffler, Martin Schlipf, Karlheinz Schwarz, Sangeeta Sharma, Francesca Tavazza, Patrik Thunström, Alexandre Tkatchenko, Marc Torrent, David Vanderbilt, Michel J. van Setten, Veronique Van Speybroeck, John M. Wills, Jonathan R. Yates, Guo-Xu Zhang, and Sangeeta Sharma, “Reproducibility in density functional theory calculations of solids,” Science 351, aaad3000 (2016).

41. S.H. Taole, H.R. Glyde, and R. Taylor, “Anharmonic properties of Li,” Phys. Rev. B 18, 2643 (1978).
Supplementary Information: Using forces to accelerate first-principles anharmonic vibrational calculations

Joseph C. A. Prentice        R. J. Needs
November 29, 2021

1 Pseudopotentials

All density functional theory calculations were performed using CASTEP version 8.0, and its own “on-the-fly” ultrasoft pseudopotentials. The definition strings for the pseudopotentials were:

- H:
  \[1|0.6|1|6|10|10(qc=8)\]
- Li:
  \[1|1.0|14|16|18|10U:20(qc=7)\]
- Zr:
  \[3|2.1|7|8|9|40U:50:41:42\]

2 Equilibrium unit cell configurations

The unit cells for the structures used in this work, containing the atoms at their equilibrium positions, are given in the .cif files H2.cif, cmca4.cif, cmca12.cif, c2c24.cif, Li.cif and Zr.cif.

3 Harmonic mode displacement patterns

The displacement patterns corresponding to the mapping directions used in the mapping of 2-D subspaces of the BO surface of the Cmca-4 structure of solid hydrogen are given below. They correspond to the displacement patterns of harmonic modes with frequencies of 69.4, 74.0 and 114 meV, labelled as 4,
5 and 7 respectively. Each row shows the displacement of a H atom in the three Cartesian directions, in the same order as the atoms are listed in the file cmca4.cif.

- Direction 4:

  \[
  \begin{array}{ccc}
  x & y & z \\
  0.00 & 0.75 & -1.00 \\
  0.00 & 0.75 & 1.00 \\
  0.00 & -0.75 & 1.00 \\
  0.00 & -0.75 & -1.00 \\
  \end{array}
  \]

- Direction 5:

  \[
  \begin{array}{ccc}
  x & y & z \\
  0 & 1 & 0 \\
  0 & -1 & 0 \\
  0 & 1 & 0 \\
  0 & -1 & 0 \\
  \end{array}
  \]

- Direction 7:

  \[
  \begin{array}{ccc}
  x & y & z \\
  1 & 0 & 0 \\
  -1 & 0 & 0 \\
  -1 & 0 & 0 \\
  1 & 0 & 0 \\
  \end{array}
  \]