Bulk aluminum at high pressure: A first-principles study

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(Received 20 October 2007; revised manuscript received 21 March 2008; published 16 May 2008)

The behavior of metals at high pressure is of great importance to the fields of shock physics, geophysics, astrophysics, and nuclear materials. We study here bulk crystalline aluminum from first principles at pressures up to 2500 GPa—soon within reach of laser-based experimental facilities. Our simulations use density-functional theory and density-functional perturbation theory in the local-density and generalized-gradient approximations. Notably, the two different exchange-correlation functionals predict very similar results for the fcc→hcp, fcc→bcc, and hcp→bcc transition pressures, around 175, 275, and 380 GPa, respectively. In addition, our results indicate that core overlaps become noticeable only beyond pressures of 1200 GPa. From the phonon dispersions of the fcc phase at increasing pressure, we predict a softening of the lowest transverse acoustic vibrational mode along the [110] direction, which corresponds to a Born instability of the fcc phase around 725 GPa.

DOI: 10.1103/PhysRevB.77.172102 PACS number(s): 62.50.-p, 62.20.D—

First-principles calculations have proven useful to the fields of geophysics,1 astrophysics,2 and nuclear materials.3 Aluminum, being cubic close packed and having no d-shell electrons, is a prototype for theoretical predictions and understanding the high-pressure behavior of simple metals.4 Currently, the National Ignition Facility5 at LLNL is expected to achieve shock compression6 of metals up to 2000 GPa. This new facility may provide rapid advances to high-pressure physics and could partner very successfully with theoretical studies.

The equation of state (EOS) and phase stability of aluminum were first studied from first principles in the early 1980s.7–9 In all cases, the predicted phase sequence was fcc→hcp→bcc, but predictions greatly differed in the transition pressures. Several other calculations within the local-density approximation (LDA)10 or the generalized-gradient approximation (GGA)11,12 have since been performed, with a predicted static (i.e., without the phonon contribution) fcc→hcp transition pressure of 205±20 GPa (Ref. 10) in LDA and 170 (Ref. 11) and 192 GPa (Ref. 12) in GGA. These discrepancies are more notable for the hcp→bcc transition pressure: 565±60 GPa (Ref. 10) in LDA versus 360 GPa (Ref. 11) in GGA, leaving significant uncertainties open. Theoretical work on the vibrational properties of aluminum also suggests for the fcc→hcp transition a transition pressure higher than the static one.11,12 Elastic properties13,14 and the absolute strength under tension15 have also been calculated; the latter results are of particular interest as they demonstrate the important role vibrational modes play in determining mechanical stability and suggest that shear failure modes are inherent in aluminum.

Experimentally, the equation of state at high pressures was studied by shock compression16 at pressures above the predicted maximum for the fcc→hcp phase boundary,16 but a transition was not observed. However, recent diamond anvil cell experiments observed a fcc→hcp transition at 217±10 GPa,17 highlighting the difficulty in achieving thermodynamic equilibrium in shock compression.

In this Brief Report, we report first-principles calculations of aluminum under hydrostatic compression up to 2500 GPa. In order to assess mechanical stability under shock, we also calculate the vibrational properties in the fcc phase and determine the elastic constants from the slopes of the phonon dispersions (i.e., the sound velocities).

The equations of state in the fcc, bcc, and hcp phases have been calculated with density-functional theory within both LDA18 and GGA.19 Calculations have used the QUANTUM-ESPRESSO package.20 We use plane-wave basis sets and pseudopotentials and both 3 electron (3e) norm-conserving pseudopotentials,21 with the 3s and 3p electrons in the valence and nonlinear core corrections, and 11 electron (11e) ultrasoft pseudopotentials22 where the 2s and 2p electrons, usually frozen in the core, are explicitly included in the valence. The inclusion of the 2s and 2p electrons in the valence is essential to investigate the relevance of inner core electrons at high pressure. The plane-wave cutoffs for the wave functions are 25 and 100 Ry for the 3e and 11e pseudopotentials, respectively, and 150 and 800 Ry for the charge density. Brillouin zone integrations have been performed using a cold smearing14 of 0.02 Ry over shifted Monkhorst–Pack meshes of order 16×16×16 for the fcc, 22×22×22 for the bcc, and 16×16×10 for the hcp phases. The large sizes of the k-point meshes are necessary to obtain fully converged transition pressures. The data for the total energy as a function of volume have been fitted to the Birch third order EOS23 near equilibrium to obtain equilibrium volumes and bulk moduli. The data for the equation of state have been determined from calculations performed at around 50–100 different volumes over a pressure range 0–2500 GPa. Finally, the vibrational properties have been computed using density-functional perturbation theory (DFPT).24 The dynamical matrices have been calculated on a 4×4×4 q-point mesh and Fourier interpolation has been used to evaluate the phonon frequencies on finer grids.

The role of the inner core electrons is of primary concern at very high pressures. Under normal conditions, there is not sufficient overlap between the core and valence shell electrons to question the frozen-core approximation,25 but at the pressures considered here core overlaps may become significant. To study the validity of the frozen-core approximation, we first compared the equations of state for different phases using both the 3e and 11e pseudopotentials. We report in
Table I the equilibrium lattice parameters and bulk moduli in the fcc phase at zero pressure, and in Fig. 1(a) the relative enthalpies of fcc and bcc Al with respect to the fcc phase, up to 2500 GPa. These results show that (1) there is little difference between the LDA and GGA predictions, hinting at a broad applicability of density-functional theory in either approximation, and (2) that the role of the core electrons starts to become noticeable only around 1200 GPa, even if already at zero pressure the cores of the 3e pseudopotential start to overlap.27 The equations of state for aluminum in the fcc, bcc, and hcp phases, using the 11e GGA pseudopotentials, are shown in Fig. 1(b). Although the 11e and 3e calculations give consistent results up to 1200 GPa, the calculated transition pressures can vary, particularly for the hcp→bcc transition. This could easily derive from the fact that enthalpy differences between these three phases are only a few mRy [see Fig. 1(b)], and so, even at full computational convergence of all parameters, small effects (e.g., core-state relaxations), which could shift the calculated enthalpy by less than a mRy, can significantly affect the calculated transition pressures. On the other hand, core electrons seem to have a negligible effect in determining the equilibrium volume, bulk modulus, and even phonon dispersions (see below).

Our LDA and GGA results are consistent with previously reported GGA calculations,11,12 discrepancies arise with the LDA results reported in Ref. 10, which predict 205±20 and 565±60 GPa for fcc→hcp and hcp→bcc transition pressures, respectively. This discrepancy could arise from Ref. 10 using only 10–15 points to fit the equation of state: As reported there, this approximation could significantly affect transition pressures due to the aforementioned small enthalpy differences between competing structures. We also observe that all parameters of the calculation, and particularly the k-point sampling of the Brillouin zone, need to be carefully converged.

Although the pressure that we obtain for the fcc→hcp transition, 175 GPa, is consistent with previous works,11,12 this result is lower than the experimental value of 217 GPa. As suggested in Refs. 11 and 12, this discrepancy could arise from excluding the phonon contribution to the free energy—a hypothesis that should be thoroughly tested, but that is beyond the scope of this Brief Report.

It should be noted that in our simulations, the hcp phase was always fully relaxed to identify the optimal, equilibrium c/a ratio; this is shown in Fig. 2(a). Comparison with experiment at 292 GPa finds agreement in the c/a ratio to within 0.1% and well within experimental uncertainty.17 At 222 GPa, the predicted value of the c/a ratio differs from experiment by 1%. Since experiments observe a region be-
between 217 and 260 GPa in which the fcc and hcp phases coexist, this discrepancy is made more reasonable considering that the system might be out of equilibrium. We also show in Fig. 2(b) the equilibrium volumes for the different phases as a function of pressure. Volume differences between the phases are −0.055, −0.104, and −0.040 Å³ for the fcc → hcp, fcc → bcc, and hcp → bcc transitions, respectively, corresponding to volume changes of 0.6(6)%, 1.4(0)%, and 0.6(1)%. We note that in our phase sequence, and in those discussed in literature,7–9 only the fcc, hcp, and bcc phases are considered. As a brief self-check, we performed variable-cell relaxations at 1000 GPa using a four-atom unit cell and five random distinct perturbations. The bcc structure was always found.

In order to estimate the dynamical response of aluminum under compression, we calculated both the phonon dispersions of the fcc phase and the cubic elastic constants (these were derived from the sound velocities, i.e., the slope of the phonon dispersions around Γ) as a function of pressure. Experiments have shown that the fcc phase may exist at pressures beyond the equilibrium transition pressure, and any mechanical instabilities that may lead to mechanical failure are relevant for high-pressure experiments.

We calculated the phonon dispersions up to 1150 GPa using DFPT. For aluminum, this method has been shown to accurately reproduce experimental values at P=0.24 Our calculations were performed with the 3e GGA pseudopotential, but compared at selected points in the Brillouin zone with 11e GGA calculations at pressures up to 1200 GPa. As shown in Fig. 3, the discrepancies between the 3e and 11e results are at most of the order of 3%–4% at the highest pressure, and much smaller below that. The phonon dispersions are shown in Fig. 4(a) and we highlight the appearance, with increasing pressure, of a distinct softening of the lowest energy mode in the [110] direction. This starts to become evident at approximately 400 GPa, and is complete at 725 GPa, as highlighted in Fig. 4(b). Since the slope of the dispersion curves is directly related to the elastic constants [Eq. (1)], we can extract the stiffness tensor from the vibrational modes near Γ; in our case, the Born28 criterion for stability is

\[ \frac{1}{2m} \left( \frac{\partial E}{\partial k_{110}} \right)^2 = c_{11} - c_{12} \geq 0. \]

As Fig. 5 shows, the stiffness against shear deformation, \( \Delta c = c_{11} - c_{12} \), decreases above 400 GPa and goes to zero around 725 GPa, resulting in a Born instability. These results complement existing studies of the properties of bulk aluminum13,15 and suggest another shear failure mode, supporting previous studies suggesting shear failure modes to be inherent to bulk aluminum.13,15 More advanced treatments of mechanical stability including such effects of anharmonic
modes\textsuperscript{24} and internal shear stresses created by loading\textsuperscript{29–31} need to be considered in relation to the specific experimental setup before reliable maximum stable pressures can be definitively determined.

Funding for this project has been provided by the U.S. Department of Energy Contract No. DE-FG02-05ER46253. The authors would like to thank A. Dal Corso, I. Dabo, Y.-S. Lee, B. Wood, and J. Garg for useful discussions.

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\bibitem{heller1999} A. Heller, Science and Technology Review (4 July/August 1999); https://lasers.llnl.gov/.
\bibitem{core-radius} The core radius of the 3\ensuremath{\varepsilon} pseudopotential is 1.43 Å; the half-bond length at 0 GPa is also 1.43 Å, decreasing to 0.93 Å at 1200 GPa. RRK\textsuperscript{21} pseudo-wave-functions closely reproduce all-electron wave functions even for radii smaller than the matching radius, partially explaining the good results obtained with the 3\ensuremath{\varepsilon} pseudopotential.
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