Ground-state properties of metallic solids from \textit{ab initio} coupled-cluster theory

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Metallic solids are a challenging target for wavefunction-based electronic structure theories and have not been studied in great detail by such methods. Here, we use coupled-cluster theory with single and double excitations (CCSD) to study the structure of solid lithium and aluminum using optimized Gaussian basis sets. We calculate the equilibrium lattice constant, bulk modulus, and cohesive energy and compare them to experimental values, finding accuracy comparable to common density functionals. Because the quantum chemical “gold standard” CCSD(T) (CCSD with perturbative triple excitations) is inapplicable to metals in the thermodynamic limit, we test two approximate improvements to CCSD, which are found to improve the predicted cohesive energies.

\textbf{Introduction.} Ab initio wavefunction-based electronic structure theories are being increasingly applied to periodic solids \cite{1–13}, where they can be used as predictive tools on their own or to guide the choice of functionals in more affordable density functional theory (DFT) \cite{14, 15} calculations. To date, most applications of these methods are to semiconducting or insulating systems. Extending to metals is a challenge because many of the most successful wavefunction-based methods employ finite-order perturbation theory, whose correlation energy typically diverges in the thermodynamic limit \cite{3, 16–18}. Coupled-cluster (CC) theory \cite{19} is particularly promising in this regard, because even its lowest-order nontrivial truncation to single and double excitations (CCSD) includes a number of important, canonical classes of diagrams, including ladder diagrams (important at low density) and ring diagrams (important at high density and necessary to remove the aforementioned divergence) \cite{20–24}. Despite extensive application to the ground state of the uniform electron gas (UEG) \cite{17, 18, 21–23, 25–31}, CC theory has seen limited application to atomistic metals \cite{8, 32–34}.

Here, we apply CCSD to study the structural and energetic properties of two simple metals, body-centered cubic (BCC) lithium and face-centered cubic (FCC) aluminum. We address two of the key technical hurdles associated especially with the \textit{ab initio} study of metals, namely the removal of basis set error and finite-size error. To address basis set error, we use system-specific Gaussian-type orbital (GTO) basis sets \cite{35, 36} that are optimized to lower the total energy and to lower the condition number of the overlap matrix. We demonstrate the success of this approach by comparing our results to those obtained with plane-wave basis sets. To address finite-size error, we employ relatively dense Brillouin zone samplings with twisted boundary conditions and subsequent extrapolation. We then address the CCSD error by two approximate methods: adding the correlation energy due to perturbative triples \cite{37} with a coarse Brillouin zone sampling or scaling the CCSD correlation energy by a non-empirical factor determined by the UEG.

\textbf{Methods.} For BCC lithium, we use an isotropic primitive cell containing two atoms. For FCC aluminum, we use two unit cells: an anisotropic primitive cell containing two atoms and an isotropic cubic cell with four atoms. Except where indicated, calculations were performed with the lattice constants $a = 3.5$ Å (Li) and $a = 4.05$ Å (Al), which are close to the experimental values. All calculations are performed at zero temperature using PySCF \cite{6, 38, 39} with libcint \cite{40}, and GTO-based calculations were performed with Gaussian density fitting \cite{41–43}. We use GTH pseudopotentials \cite{44–47}, correlating three electrons per atom, for both lithium and aluminum.

The original GTO basis sets designed for use with GTH pseudopotentials \cite{36} were not optimized for correlated calculations and furthermore only contain $s$, $p$, and $d$ functions. Therefore in this work, we re-optimize the GTH-DZVP (DZ), GTH-TZV2P (TZ), and GTH-QZV3P (QZ) basis sets; for aluminum, we also added $f$ functions to the TZ and QZ basis sets, which were found to be important in our testing. In periodic solids, increasing the size of the basis set by brute force frequently leads to linear dependencies, quantified by an overlap matrix with large condition number, and concomitant numerical issues. Following similar works \cite{36, 48–51}, here we optimize these basis functions for each solid by minimizing the cost function

\begin{equation}
    \text{cost} = E_{\text{HF}} + E_{c}^{(2)} + \gamma \ln(\text{cond}(S))
\end{equation}

where $E_{\text{HF}}$ is the Hartree-Fock (HF) energy, $E_{c}^{(2)}$ is the second-order M\o{}ller-Plesset perturbation theory (MP2) \cite{52} correlation energy, $S$ is the periodic overlap matrix of the GTO basis, and $\gamma = 10^{-4} E_{h}$. For this basis set optimization, we sampled the Brillouin zone with a uniform mesh \cite{53} of $N_{k} = 2^{3}$ $k$-points (Li) or $N_{k} = 1^{3}$ $k$-points (Al), including the $\Gamma$ point; with these boundary conditions, the system is gapped and thus MP2 provides a well-defined and computationally affordable correlation energy. At this level of theory, the exponents and contraction coefficients of the GTO basis functions were optimized in an approximately alternating fashion. The optimization was started from the original GTH basis set (with additional $f$ functions in TZ and QZ, for Al) and the final basis functions may represent a local minimum of the cost function. To avoid biasing the atomic structure, the cost function \eqref{eq:cost} was averaged over three lattice parameters approximately spanning the range used in later calculations. See the Supporting Materials (SM) for further details about pseudopotentials and our optimized basis sets.
FIG. 1. Basis set convergence of MP2 (faint, open symbols) and CCSD (solid, closed symbols) correlation energies $E_c$ per atom for (a) lithium and (b) aluminum. Results were obtained using plane waves (PW) of increasing energy and using Gaussian type orbitals (GTO) and optimized GTOs, at the DZ, TZ, and QZ level. The Brillouin zone was sampled only at the $\Gamma$ point.

To demonstrate the impact of basis set optimization, in Fig. 1 we show the basis set convergence of the MP2 and CCSD correlation energy of lithium and aluminum at the $\Gamma$ point of their primitive cells, comparing GTO and plane wave (PW) results (for this figure only, the GTO results were evaluated using a PW basis set to compute the occupied bands and an approximate PW resolution of the original and optimized GTO bases to compute the virtual bands [49, 54, 55]). For Li, we see that GTO optimization increases the correlation energy by almost a factor of two, i.e., about 20 $mE_h$. A crude extrapolation suggests that the optimized QZ results recover about 80–90% of the correlation energy in the basis set limit. With this small $k$-point mesh, we can perform PW calculations with a reasonably large number of orbitals, but for Li these results converge slowly due to the inclusion of the core $1s$ electrons. An MP2 calculation with our optimized QZ basis set with 67 bands recovers more correlation energy than one with a PW basis set containing 1203 orbitals, highlighting the immense computational savings afforded by GTO basis sets. With very large PW basis sets, we begin to see the onset of $N^{-1}_{\text{bands}}$ convergence to a basis set limit in good agreement with that of the optimized GTO basis sets. For Al, the core electrons are not explicitly treated in the calculations and the PW calculations converge faster. Again, we see the benefit of basis set optimization as well as the addition of $f$ functions. The QZ result captures about 90% of the correlation energy in the basis set limit.

In addition to recovering a greater amount of electron correlation, our basis sets were optimized to reduce their numerically problematic linear dependencies. Indeed, with our largest QZ basis, the condition number of the overlap matrix decreases from about $10^{13}$ to $10^4$ for Li and from about $10^{16}$ to $10^7$ for Al. For both Li and Al, the qualitative similarity between the MP2 and CCSD correlation energies justifies our use of the former when optimizing the GTO basis set and suggests good transferability to other correlated methods [49]. Henceforth, we use these optimized GTO basis sets. Testing (not shown) indicates that these system-specific optimized GTO basis sets perform similarly to the transferable, correlation-consistent basis sets recently developed by two of us [55].

Finite-size errors are especially problematic for metals due to shell-filling effects, which can be alleviated with twisted boundary conditions [18, 30, 34, 56–60]. In Fig. 2, we show the finite size convergence of the HF energy and CCSD correlation energy for Li and Al in the optimized DZ bases. HF calculations were performed with up to $N_k = 8^3$ (Li) and $N_k = 7^3$ (Al) $k$-points, and a Madelung constant correction was used to eliminate the leading-order $N^{-1/3}_k$ finite-size error [53, 61, 62] due to nonlocal exchange. All calculations were performed using a twisted boundary condition defined by the Baldereschi point [56, 58, 60], which was found to yield smoother convergence to the thermodynamic limit (TDL) than calculations without a twist angle. Results for Li obtained by averaging over four Chadi-Cohen twist angles [57] (not shown) were found to give very similar results.

In this manner, the TDL of the HF energies can be estimated to an accuracy of about 1 $mE_h$. For Li, CCSD calculations were performed with up to $N_k = 4^3$ $k$-points; with that mesh...
they were performed using a truncated basis of MP2 natural orbitals and corrected based on results obtained with smaller meshes. Extrapolation assuming finite-size errors that scale as $N_k^{-1}$ suggests an extrapolation uncertainty of about 1 m$E_h$. For Al, due to the larger number of atoms in a cubic unit cell, our largest mesh has $N_k = 3^3$ $k$-points. For better extrapolation, we also performed calculations using an anisotropic primitive cell, with up to $N_k = 5 \times 5 \times 2$. Except for two meshes, all results from both cell choices are found to lie roughly on a straight line and reliable extrapolation can be performed to estimate the HF energy in the TDL. With the anisotropic cell, two meshes ($4 \times 4 \times 2$ and $8 \times 8 \times 4$) yield HF energies that are too high, which may correspond to incorrect HF solutions due to the challenge of minimization in metals at zero temperature; somewhat surprisingly, the correlation energy associated with the higher-energy HF solution at $N_k = 4 \times 4 \times 2$ is in line with the other correlation energies. In all future calculations on Al, we use the isotropic cell for HF energies and the anisotropic cell for correlation energies, facilitating extrapolation. The total energies that we calculate using our largest meshes differ from the extrapolated results by about 1 m$E_h$ (Li) and 3-4 m$E_h$ (Al), and the extrapolated results have an uncertainty of about 1 m$E_h$ or less.

Henceforth, we assume basis set corrections and finite-size corrections are independent and additive. To our DZ results obtained with increasingly large $N_k$, we add basis set corrections determined by calculations with smaller values of $N_k$.

For the HF energy, we assume that the QZ result is near the complete basis set (CBS) limit. For the correlation energy, the CBS limit is estimated via $X^3$ extrapolation [65] of TZ and QZ results, where $X = 3, 4$ is the cardinality. See the SM for further details about our composite corrections.

Results. In Fig. 3, we show the equation of state (EOS) of Li and Al as a function of the cell volume, where the cohesive energy, which is defined with respect to a single atom, was counterpoise-corrected by surrounding the atom with ghost atoms and their basis functions [66–68]. Single-atom calculations were performed in a spin-unrestricted manner. For Al, in sequential calculations from small to large volumes, we found that using the converged HF density matrix of the previous volume was essential in obtaining a smooth curve. The HF EOS was calculated by performing TDL extrapolation at each lattice constant. The correlation energy contribution to the CCSD EOS was calculated using our largest $k$-point meshes and then rigidly shifted by a finite-size correction calculated at lattice parameters 3.5 Å (Li) and 4.05 Å (Al), which are close to the experimental values.

In Fig. 3(a) and (b), we show CCSD results in the QZ basis and CBS limit, which only differ by about 0.2 m$E_h$ for Li and 2 m$E_h$ for Al, indicating the good performance of our optimized basis sets. By fitting our data to a Birch-Murnaghan equation [63, 64], we extract the lattice constant, bulk modulus, and cohesive energy. These properties are listed in Tab. I and compared to experimental values. The experimental results have been corrected for zero-point motion (ZPM) using the ZPM correction obtained in Ref. 64 with the HSE06 functional [69–71], although other functionals yield similar corrections.

Comparing to experiment, we find that the magnitude of
the CCSD cohesive energy is too small by about 10 mE₂/h (Li) and 20 mE₂/h (Al), which our testing suggests is mostly due to insufficient correlation in the solid rather than the single atom. Therefore, we tested two approximate improvements to CCSD. First, we tested CCSD(T), which applies a perturbative correction to the correlation energy due to triple excitations. The energy of the atom was calculated in the usual manner, but the CCSD(T) energy of the solid must be calculated in a modified form because otherwise it diverges in the TDL [17] due to the contribution of low-energy excitations with vanishing momentum transfer. Because the long-range part of the Coulomb interaction is already treated with properties given in Tab. I. The cohesive energies are significantly improved over CCSD or CCSD(T)SR. For Li, the magnitude of the cohesive energy is overestimated by 4 mE₂/h (7% error), although structural properties are notably worse than those from CCSD or CCSD(T)SR. For Al, the cohesive energy is almost perfectly predicted, but structural properties are marginally improved (bulk modulus) or slightly worse (lattice parameter).

**Discussion and conclusions.** For Li, our results can be compared to previous ones in the literature, which are included in Tab. I. First, our HF results are in good agreement with those reported in Ref. 68, which were calculated using an optimized DZ basis with the CRYSTAL package [85]. This agreement confirms a consistent starting point for correlated calculations. To our knowledge, there are no reports of periodic CCSD calculations of the equation of state of BCC Li (a recent work [34] used periodic CCSD to estimate the
energy difference between FCC and BCC Li). However, in Ref. 32, an incremental CCSD scheme was applied based on finite clusters. Although that work found a lattice constant in good agreement with ours, it found a cohesive energy that was significantly different (−0.060 $E_h$ compared to our −0.051 $E_h$) and in much better agreement with experiment. As discussed in detail in that work, the application of incremental schemes to metallic systems is delicate and nontrivial, and this might be responsible for the disagreement. Based on our investigations, we find no evidence for errors on the order of 0.01 $E_h$. Instead, we believe that this level of accuracy is expected for CCSD based on its known performance for the UEG [18, 28], where it underestimates the correlation energy at metallic densities by about 10-20%.

Finally, it is natural to compare CC methods to DFT, which is significantly more affordable. In Fig. 4, we compare the properties of Li and Al predicted by CCSD, CCSD(T)$_{SR}$, and CCSD-SAC to those predicted by HF and by common functionals, as reported in Ref. 64. We compare to a few popular functionals of increasing sophistication, including the local density approximation (LDA) [15], the generalized gradient approximations (GGAs) PBE [86] and PBEsol [87], the meta GGA M06-L [88], and the screened hybrid HSE06 [69–71]. Overall, the CC results are comparable to those from GGAs but worse than the hybrid HSE06, which performs extremely well for these two materials.

Looking forward, it will be interesting to apply CC methods to less uniform metallic systems, such as metal surfaces including adsorbates or chemical reactants. For these problems, we expect to see an increased advantage of CC over DFT, due to the greater variations in the electron density and the importance of dispersion interactions. More broadly, our work has emphasized the need for wavefunction-based methods that improve upon CCSD without the use of perturbation theories that diverge for metals. Beyond more systematic investigation of the two methods proposed here, other possibilities include the use of spin-component scaling [89–91], regularization [92–94], or screened interactions [17, 95], although these approaches typically introduce empirical parameters. Alternatively, the full or limited inclusion of non-perturbative triple excitations [96–98] is a promising ab initio route towards chemical accuracy in metallic solids.

**Data Availability.** After acceptance, data will be made openly available. To ask for further data, a reasonable request can be sent to the authors.

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