Microhartree Precision in Density-Functional-Theory Calculations

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To address ultimate precision in density-functional-theory calculations we employ the full-potential linearized augmented plane-wave + local-orbital (LAPW+lo) method and justify its usage as a benchmark method. LAPW+lo and two completely unrelated numerical approaches, multi-resolution analysis (MRA) and linear combination of atomic orbitals, yield total energies of atoms with a mean deviation of 0.9 $\mu$Ha and 0.2 $\mu$Ha, respectively. Spectacular agreement with the MRA is reached also for total and atomization energies of the G2-1 set consisting of 55 molecules. With the example of $\alpha$-iron we demonstrate the capability of LAPW+lo of reaching $\mu$Ha/atom precision also for periodic systems, which allows also for distinction between numerical precision and the accuracy of a given functional.

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Density-functional theory (DFT) \cite{1, 2} is the most widely used method in modern computational condensed-matter physics and chemistry, as reflected by the existence of dozens of implementations, employing diverse numerical schemes. While each of these implementations may be most suitable for a specific type of applications, in essence, all of them solve the Kohn-Sham (KS) equation \cite{2}. Given the diversity of computational tools, it is natural to ask whether they indeed provide the same answers. This question, i.e., how reproducible DFT results are, was recently discussed in the context of a community effort, where the equation of state of 71 elemental solids was calculated using a variety of DFT implementations \cite{3}. It turned out that, after a period of fine-tuning, different DFT codes are now in good overall agreement. Despite the deviations between codes being small on average, discrepancies obtained for certain elements are much more substantial. Moreover, it is not clear, or even not expected that such agreement would be preserved beyond bulk elemental materials. As a matter of fact, the work to be done to explore and guarantee the precision of electronic-structure codes is certainly far from being finished.

Efforts on the improvement of computational methods are immensely alleviated if reliable reference data or reference tools are available. The gold standard for solving the same KS equation of DFT for condensed matter are full-potential all-electron methods, especially those employing (linearized) augmented plane waves with local orbitals (LAPW+lo) \cite{4–6} as basis functions. LAPW+lo is often trusted blindly as the ultimate reference method for validating pseudopotentials or data sets of projector-augmented waves \cite{7–10}. Yet, it has never been shown how precise this method can be in practice. Even more, arguing that the method would depend on parameters which "can influence the results in a more or less erratic way", it was even questioned recently \cite{11} whether LAPW+lo can provide benchmark quality at all.

In this Letter, we use our LAPW+lo implementation in the full-potential all-electron package exciting \cite{12} to show that for absolute total energies outstanding 1 $\mu$Ha/atom precision can be obtained. In order to validate this statement, we first turn to atoms and molecules, since for finite systems one can find other methods that, in principle, are capable of yielding an exact numerical solution of the KS equation. In the second step, we exploit the duality of the LAPW+lo basis for verifying the numerical performance of plane waves using atomic orbitals and vice versa. Making use of this concept, we demonstrate that $\mu$Ha precision is achievable also for periodic systems. Clearly, we can thus distinguish between the accuracy of a DFT functional and the numerical precision of the actual implementation.

Let us recall the Kohn-Sham equation of DFT,

$$\left[-\frac{\nabla^2}{2} + v_{\text{eff}}(r)\right] \psi(r) = \varepsilon_{\text{KS}} \psi(r).$$  \hspace{1cm} (1)

The major source of numerical issues in solving it, is the behavior of the effective potential $v_{\text{eff}}(r)$. While it is very smooth in most of the space, its shape is dominated by the electrostatic contribution in the proximity of nuclei, where it varies rapidly with a divergence at the nuclear sites. As a result, the otherwise well-behaved KS orbitals $\psi(r)$ exhibit cusps at the atomic positions and a nodal structure in their vicinity.

The LAPW+lo method meets these properties of $\psi(r)$. The unit cell is partitioned into non-overlapping atomic spheres (or muffin-tin spheres, $MT_\alpha$), centered at the nuclear positions, with index $\alpha$, and the interstitial region ($I$). KS wavefunctions are expanded in terms of atom-like functions, $f_\nu(r_\alpha) = u_\nu(r_\alpha)Y_{\ell m}(r_\alpha)$, and plane waves, respectively:

$$\phi_{G+k\mathbf{r}}(\mathbf{r}) = \left\{ \begin{array}{ll}
\sum_{\nu} A^\alpha_{G+k\mathbf{r},\nu} f_\nu(r_\alpha) & r \in MT_\alpha \\
\frac{1}{\sqrt{V_\Omega}} e^{iGk\mathbf{r}} & r \in I
\end{array} \right..$$ \hspace{1cm} (2)
The coefficients $A_{G+k\nu}^2$ are determined to ensure smoothness of the basis functions at the sphere boundaries. These augmented planewaves are typically complemented by local orbitals:

$$\phi_\mu(r) = \begin{cases} f_\mu^a(r) & r \in MT_\alpha \\ 0 & r \in I \end{cases}$$

(3)

That way the flexibility of the basis is improved which, indeed, has a major impact on results as we demonstrate below. Local orbitals are crucial also for reaching benchmark quality in $GW$ calculations as was pointed out recently [13, 14]. A more detailed introduction to the LAPW+lo method is available in Refs. [12, 15]. The overall size of the basis and the quality in the interstitial region are controlled by the dimensionless parameter $R_{MT}G_{\text{max}}$, where $G_{\text{max}}$ is the maximum length of wavevectors $G + k$ used in the LAPW basis. In other words, $R_{MT}G_{\text{max}}$ can be freely adjusted to make the expansion of wavefunctions in the interstitial region as precise as necessary. In the atomic spheres, the quality of the wavefunctions is governed by the choice of the atomic-like functions (Eqs. 2 and 3).

To illustrate how the LAPW+lo basis can be exploited to reach essentially exact total energies for a given exchange-correlation functional, we consider the oxygen atom. We restrict ourselves to using the local spin-density approximation (LSDA). Still, the same procedure works for any other functional, for which a local KS potential is available, and we present a similar discussion for the generalized-gradient approximation (GGA) in the Supplemental Material [16]. According to the Aufbau principle, the 2p-shell is partially filled, with one $p$-orbital doubly and two others singly occupied. Consequently, this atom is not only magnetic, but its effective potential and hence the electron density are not spherically symmetric. Thus, radial and angular degrees of freedom are entangled, and all wavefunctions formally contain contributions from all angular momenta. We take it into account by introducing local orbitals not only with angular momenta of $\ell$=0 and 1, that are the dominating contributions to the 1s, 2s, and 2p states, but consider also higher values of $\ell$. Their impact on the total energy is shown in Fig. 1. In the classical LAPW formalism, $f_\nu^a(r)$ combines strictly two functions per spherical harmonic for each atom. Local orbitals allow us to correct for all missing features in the pure LAPW representation and are not limited in number. A calculation using local orbitals with $\ell$ up to 1 yields the total energy within an error of $\sim 100$ $\mu$Ha. Adding basis functions with higher angular momenta gradually improves the result, and, at $\ell_{\text{max}} = 4$, this quantity differs from the estimated exact limit by less than $10^{-7}$ Ha. Likewise, we show how the total energy converges with increasing $R_{MT}G_{\text{max}}$. In this case, we observe an exponential decay of the error similarly as in Ref. [12]. Note that the errors due to missing angular degrees of freedom depend on the atomic-sphere radius; the magnitudes shown in Fig. 1 are obtained for $R_{MT} = 1.2 a_0$. It is possible to reduce $R_{MT}$ so far that already at $\ell_{\text{max}} = 1$ the errors are negligible. We find that, at $R_{MT} = 0.5 a_0$, this error is only 2 $\mu$Ha. However, a reduction of $R_{MT}$ to such a small value leads to an enormous increase of the number of LAPWs, making the calculations very expensive.

At this point, it is already clear that the quality of the basis in the atomic spheres is essential for highly accurate results, and the discussed example reflects how to handle systems with substantially asymmetric potentials in the atomic spheres. Such potentials not only occur in a range of spin-polarized systems, but also in systems with short bonds.

To show that the converged limit in LAPW+lo corresponds to the exact numerical solution we compare them to two other all-electron methods that are expected to deliver highly precise results. The first one is the multi-resolution analysis (MRA) [17]. It recasts the KS equation in the Lippmann-Schwinger integral equation form and solves it iteratively by applying local and non-local operators on trial wavefunctions numerous times. The wavefunctions are stored in an adaptive multi-scale representation, while the integral kernels of the non-local operators are represented in a separable form. Such a numerical approach allows for solving the KS equation efficiently with a guaranteed precision. The MRA is implemented in the MADNESS code [17], which is currently restricted to finite systems. The other alternative method is the linear combination of atomic orbitals (LCAO), for which we use the NWChem package [18]. Although, in the general case, the precision of LCAO for absolute total energies is limited, it was shown how Gaussian-type orbitals can be used for reaching the complete-basis limit for atoms [19] which we employ in this work. In the calculations of molecules, we resort to the augmented correlation-consistent polarized quadruple- and quintuple-$\zeta$ basis sets known also by their acronyms aug-cc-pVQZ and aug-cc-pV5Z [20–
atomization energy

values of the cutoff parameter

an algorithm that predictably converges even at high val-

the KS wavefunctions, all local orbitals and a number of

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other words, we compute isolated atoms and molecules

primarily for studying problems of condensed-matter the-

mental Material [16].)

the LSDA and NWChem (LCAO). All quantities are obtained using the LSDA and given in Ha/atom.

TABLE I. Mean deviation (MD), mean absolute deviation (MAD), and maximum absolute deviation (MaxD) of LAPW+lo total (left columns) and atomization energies (right columns) of the G2-1 molecules with respect to the results obtained with MADNESS (MRA) and NWChem (LCAO). All quantities are obtained using the LSDA.

|        | \( \Delta E_{\text{tot}}^{\text{MRA}} \) | \( \Delta E_{\text{tot}}^{\text{LCAO}} \) | \( \Delta E_{\text{atom}}^{\text{MRA}} \) | \( \Delta E_{\text{atom}}^{\text{LCAO}} \) |
|--------|---------------------------------|-----------------|-----------------|-----------------|
| MD     | \( 0.2 \cdot 10^{-6} \)         | \( 1.2 \cdot 10^{-3} \) | \( 0.4 \cdot 10^{-6} \) | \( 6.1 \cdot 10^{-5} \) |
| MAD    | \( 0.3 \cdot 10^{-6} \)         | \( 1.2 \cdot 10^{-3} \) | \( 0.5 \cdot 10^{-6} \) | \( 9.9 \cdot 10^{-5} \) |
| MaxD   | \( 1.1 \cdot 10^{-6} \)         | \( 7.6 \cdot 10^{-3} \) | \( 1.5 \cdot 10^{-6} \) | \( 2.1 \cdot 10^{-3} \) |

FIG. 2. (Color online) The LSDA energies of atoms calculated using MADNESS (MRA) and NWChem (LCAO). The exciting (LAPW+lo) results are used as the reference. Note the µHa precision throughout.

22], respectively. These basis sets were used by Willand et al. [23] for generating all-electron reference data for benchmarking newly-generated pseudopotentials. The so-obtained atomization energies turned out to be converged to at least 1 kcal/mol (≈ 1.6 \cdot 10^{-3} Ha), which is commonly referred to as chemical accuracy.

MRA and LCAO are designed for calculating finite systems, and thus we make use of them first for comparison with our total energies of atoms and further below for molecules. We employ non-relativistic theory and the LSDA [24]. This choice, however, does not influence the overall conclusions from our work. The total energies of atoms obtained with the three codes are compared in Fig. 2. In all cases, we observe outstanding agreement between LAPW+lo and MRA (LCAO) with a mean absolute deviation of 0.9 µHa (0.2 µHa). (Computational details and the total energies are provided in the Supplemental Material [16].)

The excellent agreement between the different methods is all the more remarkable in view of the very different ways the KS equation is solved. In particular, it concerns the fact that exciting has been developed primarily for studying problems of condensed-matter theory, e. g., it considers electrons in periodic potentials. In other words, we compute isolated atoms and molecules employing periodic boundary conditions. It translates into a huge number of basis functions, and, thus, requires an efficient way to diagonalize the Hamiltonian. To do so, we have implemented a novel approach. It follows the block-Davidson algorithm [25], though with an important modification such that the initial subspace is particularly suitable for LAPW+lo. It consists of an initial guess for the KS wavefunctions, all local orbitals and a number of eigenvectors of the overlap matrix. As a result, we obtain an algorithm that predictably converges even at high values of the cutoff parameter \( R_{\text{MT}} G_{\text{max}} \) when the LAPWs become nearly linearly dependent. Our implementation of this algorithm follows the spirit of Ref. [26] and thus does not require the construction of the entire Hamiltonian and overlap matrices. Details of the implementation will be published elsewhere [27].

In order to reach the limit of an isolated molecule (or atom) in LAPW+lo calculations, it is necessary to ensure that there is no artificial interaction between the periodic images of molecules in neighboring unit cells. It is especially critical for polarizable molecules with permanent dipoles, as their interaction energy scales as \( d^{-3} \) [28], where \( d \) is the distance between adjacent molecules. We eliminate this slow decay by truncating the Coulomb potential [29]. Such an approach is particularly important for molecules like LiH. The truncation of the Coulomb potential allows for a reduction of the unit-cell dimensions from 80 Å to 16 Å. Thus, the size of the LAPW+lo basis reduces from \( 10^8 \) to \( 10^6 \) making the total-energy calculation feasible. Note that even \( 10^6 \) basis functions is an unusually large problem size in comparison to typical LAPW+lo calculations.

Equipped with this methodology, we turn to the second benchmark, which is the G2-1 set [30]. This set contains 55 molecules, consisting of 2–8 atoms for which a variety of experimental data is available. Thus it provides an excellent opportunity for benchmarking methods of DFT and quantum chemistry. Here, we use it for comparing different computational methods. We consider fixed geometries according to the data published in Refs. [31, 32].

Table I summarizes deviations of the MRA and LCAO energies from those obtained with exciting. The complete list of total energies can be found in the Supplemental Material [16]. The agreement between LAPW+lo and MRA is spectacular for both absolute total energies and atomization energies. The average and maximum deviations in total energy from the other methods amount to 0.3 µHa/atom and 1.1 µHa/atom, respectively, consistent with the results for atoms shown above. Similarly, we obtain 0.5 µHa/atom and 1.5 µHa/atom for the average and maximum deviations in atomization energies, respectively. The excellent agreement between the highly
converged LAPW+lo and MRA calculations allows us to argue that these two methods provide essentially exact answers. The obtained discrepancy, thus, can be considered as the error estimate of the two methods.

As argued above, the employed LCAO basis sets are not supposed to yield exact answers. Indeed, the average absolute deviations in the total (atomization) energies of LAPW+lo from LCAO calculations are three (two) orders of magnitude larger compared to those from the MRA. Still, with the only exception of the SO₂ molecule (error of 1.9·10⁻³ µHa (1.2 kcal/mol)), the LCAO atomization energies are within the chemical-accuracy limit of 1.0 kcal/mol.

The high precision of the LAPW+lo method obtained in calculations of atoms and molecules suggests a similar performance for solids. While it would be desirable to benchmark against other methods also for periodic systems, we are not aware of implemented alternative approaches that are expected to be exact. On the other hand, the nature of the LAPW+lo method opens a door for “self-validation” through the partitioning of space that introduces two very different ways of representing wavefunctions. More specifically, if the atomic-sphere volume is reduced, a certain region that was previously described by atomic-like orbitals is now described by planewaves. We argue that if such a rearrangement does not introduce a change in the total energy, the complete-basis limit has been reached.

We demonstrate the performance of the LAPW+lo method for periodic systems with the example of ferromagnetic α-iron. This material presents numerical challenges as reflected in Ref. 3, where the corresponding results show scattering well above the average over the 71 elements of the Periodic Table. The need for a precise and thus unique result given a certain functional is also motivated by the fact that all-electron calculations are commonly taken as a reference for benchmarking pseudopotentials, as it was done in a recent DFT study of thermoelastic properties of iron [46].

The aspherical density distribution due to the magnetic moment, as discussed above, requires particular care. In order to handle the anisotropy of the potential in the atomic spheres we introduce local orbitals with angular momenta ℓ up to 6. Using this setup and a sufficiently high LAPW cutoff, we vary \( R_{MT} \) in the range of 1.4–2.1 \( a_0 \). Note that such a variation of the atomic sphere corresponds to a change in its volume by a factor of three. Remarkably, the total energy stays within the 1 µHa range. This is the case for any considered volume within 10% deviation from the equilibrium volume of the primitive unit cell. Thus, we argue that the 1 µHa precision has been achieved also for this case. The outstanding agreement is obtained for the entire energy-versus-volume dependence, as demonstrated by the Birch-Murnaghan equation of state based on 21 data points within ±5% of volume change, shown in in Table II. As soon as convergence is reached in terms of the planewave cutoff, we obtain extremely stable values of the equilibrium volume \( V_0 \), the bulk modulus \( B_0 \), and its pressure derivative \( B' \). The former two fluctuate only in their sixth, the latter one in its fifth decimal place. A polynomial fit considering a wider range of volumes (±10%) exhibits the same stability (see Supplemental Information [16]).

Using the same settings as above, we obtain also highly precise values for the equilibrium lattice constant and bulk modulus from scalar-relativistic PBE calculations, shown in Fig. 3 together with data from the literature.

Declaring the scattering of the calculations (wider than the experimental ones) do not allow for conclusions about the exact result unless a highly reliable reference calculation is available. Strikingly, our reference value obtained in this work is located far from the middle of the cloud of the PBE data from Refs. [3] and [39], and farthest away.

**TABLE II.** Equilibrium volume, \( V_0 \) (in atomic units), bulk modulus, \( B_0 \) (in GPa), and its pressure derivative, \( B' \), for \( \alpha \)-iron as obtained from a fit of non-relativistic LSDA results to the Birch-Murnaghan equation of state. \( \Delta E^{tot} \) (in µHa) is relative to the value in the first row. All results above the separating line are fully converged.

| \( R_{MT} \) | \( R_{MT} G_{max} \) | \( V_0 \) (in Å³) | \( B_0 \) (in GPa) | \( B' \) | \( \Delta E^{tot} \) (in µHa) |
|-----|----------------|--------------|---------------|-----|-----------------|
| 1.4 | 14             | 71.3298      | 236.296       | 4.5992 | 0.0             |
| 1.8 | 14             | 71.3299      | 236.295       | 4.5994 | 0.4             |
| 2.1 | 14             | 71.3302      | 236.295       | 4.5994 | 0.8             |
| 2.1 | 12             | 71.3297      | 236.298       | 4.5998 | 3.8             |
| 2.1 | 10             | 71.3243      | 236.337       | 4.5968 | 95.0            |
| 2.1 | 8              | 71.1087      | 239.443       | 4.6320 | 2253.0          |

FIG. 3. (Color online) Bulk moduli and lattice constants of \( \alpha \)-iron. Yellow triangles correspond LAPW+lo calculations, red circles to results by other methods taken from Ref. [3]. Empty and blue triangles represent older LAPW(+/lo) calculated circles to results by other methods taken from Ref. [3]. Lines indicate results obtained in this work. Diamonds correspond to experimental data [33–45].
from experiment. Notably, our results allow for comparing performance of two functionals as we illustrate by discussing PBE and PW91 [47]. The spread of data obtained with the two GGAs implies that, without reference data, a distinction between the accuracy of a given functional and the numerical precision introduced by a specific implementation would not be possible. Indeed, besides a few exceptions [48], these two GGAs were often considered synonymous. Our calculations clearly show that PW91 yields a smaller lattice constant and a larger bulk modulus than PBE.

In conclusion, we have challenged the numerical accuracy of the LAPW+lo method. In order to demonstrate its capability, calculations for atoms and molecules have been benchmarked against two completely unrelated, highly precise methods. The differences in absolute total energies are on average 1 µHa/atom. Furthermore, we have shown that we reach the same precision also for solids. The presented results allow us to claim that, once properly converged, LAPW+lo is an essentially exact method for DFT calculations. Overall, this work presents also a justification for using LAPW+lo as a reference method, backing up its reputation as the gold-standard method of DFT for condensed matter. The ability to reach the complete-basis limit will be indispensable for benchmarking less precise methods and for quality control of data collections. Furthermore, it opens perspectives towards reliably computing numerically sensitive quantities, like magnetisation anisotropy, weak non-covalent interactions, relative stabilities of isomers or polymorphs etc., where high precision is crucial.

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