Ionization potentials in the limit of large atomic number

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By extrapolating the energies of non-relativistic atoms and their ions with up to 3000 electrons within Kohn-Sham density functional theory, we find that the ionization potential remains finite and increases across a row, even as $Z \to \infty$. The local density approximation becomes chemically accurate (and possibly exact) in some cases. Extended Thomas-Fermi theory matches the shell-average of both the ionization potential and density change. Exact results are given in the limit of weak electron-electron repulsion.

A central problem of electronic structure is the calculation of the ground-state energy of the electrons of any atom, molecule, or solid, within the non-relativistic Born-Oppenheimer limit. Density functional theory (DFT) is a popular choice, balancing computational efficiency with useful accuracy. The original DFT was that of Thomas [1] and Fermi [2], TF theory, in which a local density approximation is made for the kinetic energy and the electron-electron repulsion is approximated by the simple Coulomb energy of the charge density. In the 1970’s, Lieb and co-workers [3] showed that the TF energy becomes relatively exact for neutral matter as $Z \to \infty$ in a specific way. The energy, $E$, grows in magnitude as $Z^{7/3}$, where $Z$ is the total charge. For atoms and their ions, the leading corrections in powers of $Z^{-1/3}$ were found by Scott [4], Dirac [5], Schwinger and others [6, 7], as summarized by Englert [8]. These corrections are given exactly by extended Thomas-Fermi (ETF) theory, which includes both the gradient correction for the kinetic energy (one-ninth the von Weisacker functional [9]), and the local density approximation for exchange (LSDX [10]).

However, TF theory and its extensions are insufficiently accurate to predict chemical properties[11]. Modern DFT uses the Kohn-Sham (KS) scheme, in which only a very small fraction of the total energy, the exchange-correlation (XC), needs be approximated. But the idea of asymptotic correctness was recently extended to KS, relating the success of exchange GGA’s such as PBE [12] for total energies to their recovery of the $(Z^1)$ term in the expansion of the exchange energy [13, 14]. The relation between semiclassical and local density approximations [15] contributed to the creation of PBEsol [16], a PBE-like functional that is nearly optimum for solids near equilibrium but not for atoms and molecules, and to revTPSS [17], a nearly optimal semi-local functional for all three kinds of systems.

But total electronic energies are irrelevant to chemistry. Only differences matter, such as the ionization potential of an atom ($I$ is the energy difference between the positive ion and the neutral) or the dissociation energy of a chemical bond. How relevant are asymptotic expansions for these quantities? The asymptotic expansion for $E$ is in powers of $Z^{-1/3}$, so if $I$ remains finite as $Z \to \infty$, the neutral and ion energies must agree for the first seven powers in such an expansion, a truly remarkable balancing act between quantum effects, the Pauli principle, and the Coulomb forces of nuclear attraction and inter-electron repulsion. In this letter, we demonstrate by both calculation and analysis that (i) $I$ has no limit as $Z \to \infty$, but remains column-dependent (ii) that each column has a finite limit; (iii) the local (spin) density approximation (LSD [10]) of KS theory becomes very accurate (if not exact) for $I$ for certain cases; (iv) ETF theory becomes very accurate (if not exact) for the average of $I$ over an entire shell; (v) the shell-averaged difference in density between the neutral and its ion approaches that of TF. We demonstrate these statements in the limit of weak interelectron repulsion.

Our most important results are shown in Fig. 1. We plot $I$ from various calculations, extrapolated to infinite row number, versus the column number for main-group
elements (s and p valence shells). We calculate exchange exactly \( i \), using the optimized effective potential (OEP, which here should be indistinguishable from Hartree-Fock \( [20] \)), extrapolating all values to \( Z \to \infty \). At the exchange level, LSD and PBE are almost exact for \( p \)-valence elements, and are highly accurate but inexact for the \( s \)-valence cases. Furthermore, ETF yields \( S \) a single number (3.15 eV), very close to the \( s \) and \( p \) average (3.02 eV). When correlation is included, gradient effects are slight, and it is in the regime of large electron number that approximate density functionals work best, sometimes exactly \([13, 15, 21]\). We speculate that LSDX on accurate densities becomes almost exact in this limit for \( p \)-shell cases, that ETF is exact for some shell-average, and that our XC results are extremely accurate and practically impossible to calculate with any other method.

To understand why local functionals become accurate in this limit, begin with total energies of neutral atoms, whose large-\( Z \) expansion is

\[
E_Q(Z) = -c_q^{(0)} Z^{7/3} + 0.5 Z^2 - c_q^{(2)} Z^{5/3} + \ldots \tag{1}
\]

where \( E_Q(Z) \) is the energy of an atom with atomic number \( Z \) and charge \( Q \), and the \( c_j^{(i)} \) are coefficients depending on the degree of ionization, \( q = Q/Z \). We use atomic units throughout. The neutral coefficients were derived via semiclassical arguments of Schwinger and Englert \([6, 7]\).

The TF energy is exactly \( -c_q^{(0)} Z^{7/3} \). The second term \( \xi \) comes from the \( s \)-orbitals at the nucleus and must be treated quantum mechanically. The third term is derivable in ETF theory \([6]\), of which \( 2/11 \) arises from the gradient correction to the kinetic energy, and \( 9/11 \) from LSDX. When treated in terms of the potential, the divergence at the nucleus is avoided \([8]\).

The extension of these ideas to \( I \) has proven more difficult. Terms of higher order than those shown in Eq. (1) oscillate \( i \) with \( Z \), as a precursor to the periodic variation of chemical properties that is missed by ETF, but well-described in KS DFT. The oscillations in \( I \) dominate over trends with \( Z^{-1/3} \). While numerous studies exist in the literature \([22]\) for fixed electron number \( N \) with \( Z \to \infty \), we are interested in \( I(Z) = E_I(Z) - E_0(Z) \) as \( Z \to \infty \). Within TF theory, Lieb proved \([23]\) that \( I \) does not grow with \( Z \), and by considering \( c_q^{(0)} \) as \( q \to 0 \), Englert showed \( I^{TF} \to 3a^{-2/3}/7a \approx 1.29 \text{ eV} \), where \( \Lambda = 32.729416 \) is a known constant \([8]\), and \( a = (9\pi^2/128)^{1/3} \). Even this simple result requires explanation, because \( \mu \), the chemical potential, is zero for the neutral atom in TF theory, suggesting \( I \) should be too. But the TF energy is the smooth envelope of \( E_Q(Z) \) as a function of \( q \), whereas the true energy consists of line segments between integer values \([24]\). Thus \( \mu = \partial E/\partial q = -I \) for the exact system, but the TF energy behaves as \( q^{7/3} \) for small \( q \). So \( \mu_{TF} = 0 \), but the better value of \( I^{TF} \) is the energy difference \([8]\) with \( Q = 1 \).

Next we discuss KS DFT, in which the (non-interacting) kinetic energy is not approximated, but is found exactly from the KS orbitals. We perform KS self-consistent calculations for atoms and ions up to 2938 electrons using LSD and PBE XC functional approximations, as well as the exact OEP exchange. These were done using the Engel code \([20]\), but with tightened convergence criteria and maximum numbers of orbitals, and a logarithmic radial grid with 800 points. In Fig. 2 we show \( I \) versus \( Z^{-1/3} \) for each main-group column of the periodic table. In all cases, the behavior is almost linear as a function of \( Z^{-1/3} \) for all \( Z \gtrsim 169 \), so we extrapolated these curves using a parabolic fit in \( Z^{-1/3} \) and found the ionization energy for \( Z \to \infty \) as shown in Fig. 1. The spherical approximations of the density (LSD, PBE) and of the potential (OEP), used in the Engel code (see Ref. \([19]\)), give errors less that 0.1 eV for \( I \). We use electronic configurations based on the aufbau principle and Madelung rule \([25]\). For the noble gases, \( Z = n(n^2 + 6n + 14)/6 - \Delta(n)(n^2 + 1) \), where \( n \) is the row number and \( \Delta(n) = 0 \) for even and 1 for odd rows.

To understand in detail the results shown in Fig. 1 which are also tabulated in Table 1, we begin at the exchange level. Both PBE and LSD exchange are almost identical to the OEP values for the \( p \)-group elements, with a maximum difference between them of 0.02 eV, and of either from OEP of 0.08 eV. This is not so for the alkalis and alkali earths, presumably because they have only one or two electrons outside a closed shell, with accompanying self-interaction error of approximate functionals. The ionization of \( p \)-elements involves removing electrons from a full (or almost full) shell with \( \sim (n + 2)^2/2 \) electrons, where \( n \) is the row number for even \( n \), and rows \( n \) and \( n + 1 \) have the same structure.

In fact, Englert also showed that the TF result is not correct as \( Z \to \infty \). The terms of \( O(Z^{5/3}) \) in Eq. (1) also yield a finite contribution, which is included in ETF, yielding an \( I \) of 3.15 eV, very close to the average over
TABLE I: Extrapolated ionization potentials I (eV) of main group elements. Mean absolute differences (m.a.d.) are taken relative to OEP for X, and PBE for XC. The last two columns show the electron affinity \( A \) (eV) (estimated as \( I - 1/\langle r \rangle \) in atomic units) and the average radius \( \langle r \rangle \) (bohr) of the ionization density, in the \( Z \to \infty \) limit, using PBE. For ETF, \( I=3.15 \text{ eV}, \langle r \rangle = 5.6 \text{ Å}, \) and \( A = 0.58 \text{ eV}. \)

| group | \( X \) | \( XC \) | \( A \) | \( \langle r \rangle \) |
|-------|-------|-------|------|---------|
| I | 1.56 | 1.66 | 1.42 | 1.90 | 1.77 | -0.15 | 14.13 |
| II | 1.77 | 1.89 | 1.65 | 2.41 | 2.27 | -13.56 |
| \( s \)- m.a.d. | 0.13 | 0.24 | 0 | 0.13 | 0 | -13.85 |
| \( p \)- m.a.d. | 0.05 | 0.05 | 0 | 0.08 | 0 | -8.82 |
| \( p \)- avg | 3.47 | 3.46 | 3.47 | 4.23 | 4.14 | 1.10 | 9.45 |

Both \( s \) and \( p \)-shell values (3.02 eV).

To check this is no accident, consider the simpler system of atoms with an infinitesimal electron-electron repulsion, \( \lambda \), sometimes called Bohr atoms. The orbitals are hydrogenic, requiring no self-consistency and simplifying the integrals [26]. One finds that \( I_{\text{TF}} \) is exact for large \( Z \) at \( \lambda = 0 \). In Fig. 3 we show the exchange correction (divided by \( \lambda \)) to \( I \) for LSDX applied to the TF density (yielding \( 8(23)/3 \approx 0.2360 \), to the exact densities (each averaged over entire shells), and exactly. All three match as \( Z \to \infty \), but a small error remains if, e.g., just the \( s \)-shell is used. Thus we speculate that, for real atoms, LSDX (in a KS calculation) matches the average over the entire shell as \( Z \to \infty \).

Next, we discuss the DFT calculations with correlation, which remains finite as \( Z \to \infty \) and varies across a row. The differences between PBE and LSD are relatively small, giving greater confidence in both. The maximum deviation between them for \( p \)-elements is 0.14 eV, comparable to the deviations of these functionals at the exchange-only level from OEP for the alkali and alkali earths. Thus the gradient corrections are not vanishing, suggesting that while both calculations are accurate, neither is exact. The PBE average, 3.61 eV, is our best estimate of a universal ionization potential, defined as the limit of \( I \) averaged over the \( n \)-th shell, as \( n \to \infty \).

The other major descriptor of chemistry is the electron affinity \( A(Z) = E_0(Z) - E_{-1}(Z) \). Within LSD or PBE, the first negative atomic ion of energy \( E_{-1}(Z) \) has no stable solution, but \( A(Z) \) can still be estimated [27] via a charged conductor model, in which \( I - A = 1/\langle r \rangle \), and \( \langle r \rangle \) is the centroid of the added charge. Define the radial ionization density as

\[
\Delta n_R(Z,r) = 4\pi r^2(n_{\text{atom}}(r) - n_{\text{con}}(r)),
\]

which integrates to 1. Then choose \( \langle r \rangle = \int_0^\infty dr \Delta n_R(Z,r) \). Table I shows PBE \( Z \to \infty \) limits for \( I, \langle r \rangle \), and \( A \). Averaging over \( s \) and \( p \), our best estimate for a universal value of \( A \) is 0.78 eV.

We next extrapolate the ionization density via

\[
\Delta n_R(Z,r) \approx \beta \Delta n_R(Z_0,\beta r) + \gamma d[\Delta n_R(Z_0,r)]/dr,
\]

which correctly integrates over \( r \) to 1. Here \( Z_0 = 2935 \), \( \beta = 1 + b(Z^{-1/3} - Z_0^{-1/3}) \), and \( \gamma = c(Z^{-1/3} - Z_0^{-1/3}) \), with fit parameters \( b = 5 \), and \( c = 2 \). Finally, we also averaged over the 6 \( p \)-shell curves, to find the results shown in Fig. 4. The TF solution for the infinitesimally charged ion has a finite size [8, 23], i.e.,

\[
r_c = \lim_{Z \to \infty} r_0(Z) = aZ^{2/3} \approx 9.0588 \text{ bohr} \approx 4.8 \text{ Å}.
\]
Beyond this radius, $\Delta n_{ETF}^{1}(r) \propto 1/r^d$. The maximum of this curve is about 0.1830 at $r = 8.855$ bohr. The agreement between the extrapolated $p$-shell densities and the TF theory is remarkably good, but not exact, while the extrapolated alkali ionization density is very different. We speculate that averaging over an entire shell would yield perfect agreement, as we find numerically for the Bohr atom.

Finally, we justify why such large atomic numbers (larger by a factor of 10 than those of Ref. [28]) are needed to get these results. Because of the scaling with $Z^{-1/3}$, even $Z = 125$ only makes $Z^{-1/3} = 0.2$, while $Z > 1000$ brings $Z^{-1/3}$ below 0.1, making the extrapolation much more reliable. In Fig. 5 we show accurate ionization densities for the 8-th column of the extended table at finite $Z$. The scaling of the TF ionization density is quite different from that of the exact solutions: Before extrapolation, even at $Z = 2935$, the TF ionization density agrees much better with that of the alkalics, not the $p$-shell average. For the same reasons, having HF energies for only $Z \lesssim 100$, Englert erroneously concluded that $T^{ETF}$ was the limit of the alkalics, not the shell-average (see Fig. 4-8 and its discussion of Ref. [8]).

Thomas-Fermi theory produces the first term of Eq. (1) and extended TF yields an average $Z \rightarrow \infty$ limit for the ionization energy, but no periodic variation of chemical properties and no binding of atoms to form molecules or solids. Within non-relativistic KS theory, any reasonable approximation to the XC energy with the correct uniform-density limit for exchange will produce the total-energy expansion of Eq. (1) and a finite column-dependent $Z \rightarrow \infty$ limit for the ionization energy. It appears that LSD is extremely accurate and possibly exact in certain cases for $I$. We have shown this for shell-averages in the limit of weak inter-electron repulsion. But in that case, the last shell is spread throughout the entire atom, and average gradients contributing to ionization vanish as $Z \rightarrow \infty$, which is not true for real atoms.

Thus we have established that, in the large-$Z$ limit, the periodic table becomes perfectly periodic. Moreover, local approximations appear to become exact, even for energy differences that are relatively vanishingly small in this limit. These are new, numerically relevant, exact conditions that approximate functionals should satisfy.

All conclusions are based upon numerical calculations and extrapolation. Proving them rigorously is a challenge to mathematical physics. This work was supported by DOE grant DE-FG02-08ER46496 at Irvine, and NSF (Grants DMR-0501588 and DMR-0854769) at Tulane.

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