$C_6$ coefficients and dipole polarizabilities for all atoms and many ions in rows 1-6 of the periodic table

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Abstract

Using time-dependent density functional theory (TDDFT) with exchange kernels we calculate and test imaginary frequency-dependent dipole polarizabilities for all atoms and many ions in rows 1-6 of the periodic table. These are then integrated over frequency to produce $C_6$ coefficients. Results are presented under different models: straight TDDFT calculations using two different kernels; “benchmark” TDDFT calculations corrected by more accurate quantum chemical and experimental data; and “benchmark” TDDFT with frozen orbital anions. Parametrisations are presented for 411+ atoms and ions, allowing results to be easily used by other researchers. A curious relationship, $C_{6,XY} \propto [\alpha_X(0)\alpha_Y(0)]^{0.73}$ is found between $C_6$ coefficients and static polarizabilities $\alpha(0)$. The relationship $C_{6,XY} = 2C_{6,X}C_{6,Y}/[\alpha_X/\alpha_YC_{6,Y} + \alpha_Y/\alpha_XC_{6,X}]$
is tested and found to work well (< 5% errors) in about 80% of cases, but can break down badly (> 30% errors) in a small fraction of cases.

The importance of van der Waals (vdW) forces in physical systems, especially at the nanoscale, is increasingly being recognised (see e.g. Refs. 4–9 and references therein). This renewed interest has come about in part because vdW forces are so vital to binding in layered materials such as graphene. 4–9 Unfortunately, conventional electronic structure techniques like density functional theory in its common approximations (e.g. GGAs10) do not reproduce van der Waals forces11 which can lead to poor predictions for important systems. 6 To remedy this lack, a cornucopia of new approaches has been developed over the past twenty years1,2,12–22 that allow van der Waals forces to be included alongside more conventional density functional approximations. 10,23,24

A number of these approaches - most notably those based around the work of Grimme17–19 and Tkatchenko and Scheffler2,3,20–22,25,26 - are based on atom-in-molecule (AIM) approximations. These (arguably) semi-empirical approximations involve taking free atomic C_6 coefficients and using them to determine binding in more complex bulk and molecular systems. These methods have achieved a great many successes (including in some complex systems9). Although it should be noted that they also have flaws. 7,25,27

A vital component of these AIM approaches is the data set of C_6 coefficients. These must be pre-calculated elsewhere and tabulated for input into the AIM scheme. However, determining accurate C_6 coefficients is difficult. It requires as input the dipole polarizability of a system across a range of (imaginary) frequencies which can then be integrated [see Eq. (1)] to determine the dipole-dipole C_6 coefficient. While static polarizabilities of closed- and some open-shell atoms and small molecules can be calculated very accurately (see e.g. Refs. 28,29 and references therein) using sophisticated quantum chemical techniques, the dynamic polarizabilities are more difficult to evaluate. Reactive open-shell atoms and ions, which dominate chemistry and materials science, represent a particular challenge for such methods. Thus AIM inputs are often least reliable for the elements that play the most
interesting role in materials.

One route around the limitations of quantum chemical approaches is to employ time-dependent density functional theory (TDDFT). In linear response TDDFT the dipole polarizability is obtained directly from the density-density response of an atomic or ionic system, making its evaluation relatively straightforward. Although they are not as efficient as regular DFT calculations, TDDFT calculations are generally more accurate than DFT and more efficient than accurate quantum chemical calculations. They thus provide a middle ground between speed and accuracy, allowing fairly accurate calculations to be carried our in reasonable time even for systems that are essentially intractable for more accurate quantum chemical approaches.

Chu and Dalgarno, in their 2004 paper,\textsuperscript{30} took advantage of the “balanced” nature (in terms of numerical cost and accuracy) of TDDFT to determine accurate $C_6$ coefficients for open-shell atoms, and were able to provide a complete set of coefficients for elements in rows 1-3 of the periodic table. Grimme et al used a similar approach\textsuperscript{19} to determine atomic coefficients for all atoms in Rows 1-7. But “benchmark” quality results for open shell elements in rows 4-7 remain elusive. Open-shell ionic coefficients are even less well studied, even in the upper rows.

This manuscript thus seeks to use TDDFT calculations, loosely based on the basic approach of Chu and Dalgarno, to provide frequency-dependent dipole polarizabilities for all atoms and interesting ions in rows 1-6 of the periodic table. It will further present the results in an easy to use parametrised model, making results readily available for use in AIM calculations.\textsuperscript{1}

\textsuperscript{1}These techniques could, in principle, be employed for row 7 too. But numerical issues and strong relativistic contributions make results for these elements likely to be inaccurate. We thus focus on reliable results for the more common, higher row elements.
1 Theory

The well-known van der Waals (or dispersion) $C_6$ coefficient usually appears in the formula $U_{\text{vdW}} = -C_{6,XY}/D_{XY}^6$ governing a long-range attractive potential between well-separated, localised systems $X$ and $Y$. For spherically symmetric systems, in which the dipole polarizability tensor is a scalar times the identity tensor, $C_6$ can be found through the simplified Casimir-Polder formula

$$C_{6,XY} = \frac{3}{\pi} \int d\omega \alpha_X(i\omega)\alpha_Y(i\omega),$$

which can be derived from many-body perturbation theory on the Coulomb interaction between the two systems. Here the $C_6$ coefficient depends on the frequency dependent dipole polarizability $\alpha_X/Y(i\omega)$, evaluated at imaginary frequency $i\omega$. Thus, knowing $\alpha_X(i\omega)$ for a number of atoms and ions $X$ is sufficient to calculate the $C_6$ interaction between all possible pairs. Furthermore, $\alpha(i\omega)$ can be used to calculate higher order “non-additive” interactions, such as the Axilrod-Teller interaction or higher order dipolar contributions. Note we use atomic units $\hbar = e^2/(4\epsilon_0) = m_e = 1$ (giving energies in Ha and lengths in Bohr radii $a_0$) in equation (1) and in all subsequent equations.

Clearly these dipolar dispersion forces are not the only contribution to the long-range force between such systems. Interactions between free ions are dominated by forces from the Coulomb potential $U_C = q_Xq_Y/D_{XY}$. Additionally, higher order terms, such as $C_8$ coefficients in $U_8 = -C_{8,XY}/D_{XY}^8$ also contribute, and these must be obtained from quadropolar and higher interactions. But the dipolar force plays an important role in all systems, especially in embedded ions where Coulomb forces cancel out over a molecule, leaving induction (which depends on $\alpha(0)$) and dispersion forces as the leading long-range force terms.

The relationship between polarizabilities and dispersion coefficients has been exploited indirectly in various methods. But it has been more rarely (if ever) exploited directly in atom-in-molecule approaches. In part this is likely due to a lack of viable
frequency-dependent data for atoms and ions. We will thus outline, in the remainder of this theory section, how to calculate polarizabilities that can be used directly in van der Waals calculations using (11) or related formulae.

1.1 TDDFT Methodology

At the heart of our results are linear response TDDFT calculations using the all electron numerical method described in Refs. 32–35. In summary, we calculate linear response functions using time-dependent density functional theory in an ensemble averaged (where appropriate) exact exchange groundstate\textsuperscript{34} and using an equivalent ensemble averaged\textsuperscript{32} version of the Petersilka, Gossman and Gross\textsuperscript{36} kernel. The radial exchange hole kernel\textsuperscript{32} is also employed for a further test of accuracy. We do not explicitly include any relativistic calculations. However, for our final benchmarks we do include them implicitly via corrections based on accurate reference static dipole polarizabilities (see Sec. 1.2).

To calculate polarizabilities we employ a post-DFT linear response formalism. In this approach, we first calculate the groundstate of the system using a DFT approximation. Once the groundstate properties are calculated, we use them to determine the response function $\chi$ governing the small change in densities to a small change to the potential of form $\Delta v(r; t) = \Delta v(r)e^{i\omega t}$\textsuperscript{2}. From $\chi$ we can calculate the spherically averaged dipole polarizability using

$$\alpha(i\omega) = \int drdr'zz'\chi(r, r'; i\omega)$$

and, if desired, $C_6$ coefficients can be evaluated using the Casimir-Polder formula (11). We note that $\alpha$ depends on electron number $N$ and nuclear charge $Z$. In our calculations, the dependence on $N$ is via the occupation factors, while the dependence on $Z$ is via the external potential $v_{\text{ext}}(r) = -Z/r$.

\textsuperscript{2}Conventionally $\chi$ is defined as the response to periodic $e^{-i\omega t}$. Thus we refer to the response to imaginary frequency $i\omega$ when considering the real exponential $e^{i\omega t}$.
In our calculation we start from groundstate Kohn-Sham properties [notably potentials $v_s(r)$, occupation factors $f_i$, orbitals $\phi_i(r)$ and densities $n(r)$] calculated in ensemble DFT. The use of ensembles allows us to properly account for atomic symmetries despite working from spherical and spin-unresolved groundstates in which $n(r) = n(r)$ and $n_\uparrow(r) = n_\downarrow(r) = n(r)/2$. We employ the LEXX approximation adapted to general open shell systems with $d$ and $f$ orbitals. For atoms and ions in Rows 4-6 we make a further simplifying approximation: that the shells fill “trivially” according to Hund’s rules. This can lead to energies that are higher than other fillings, but for reasons discussed in Appendix A we feel they are a more appropriate starting point.

In LEXX theory, the groundstate energy is approximated by an orbital energy functional

$$E_0[n] = \sum_i f_i t_{s,i} + \int v_{\text{ext}}(r)n(r)dr$$

$$+ \sum_{ij} (F_{ij}^H P_{ij} - \frac{1}{2} F_{ij}^x Q_{ij})$$

(3)

where the pair occupation factors $F_{ij}$ are determined by the orbital occupation factors $f_i$. Here $\sum_i f_i = N$ is the total number of electrons. The energy terms are

$$t_{s,i} = -\frac{1}{2} \int dr \nabla^2 \rho_i(r,r')|_{r=r'}$$

(4)

$$P_{ij} = \frac{\int dr dr' }{2|r-r'|} n_i(r)n_j(r')$$

(5)

$$Q_{ij} = \frac{\int dr dr' }{2|r-r'|} \rho_i(r,r')\rho_j(r',r)$$

(6)

where $\rho_i(r,r') = \phi_i^*(r)\phi_i(r')$ and $n_i(r) = \rho_i(r,r)$. In cases where all orbitals are equally occupied (i.e. where $f_i = 2$ or 0 for all orbitals) $F_{ij} = f_i f_j$ and (3) is identical to conventional EXX theory.

The Kohn-Sham potential $v_s$ is found using the Krieger, Li and Iafrate (KLI) approxi-
information to the optimized effective potential. Here we write

\[ v_s(r) = v_{\text{ext}}(r) + v_{\text{Hxc}}[\{\phi_i\}, \{f_{ij}, F_{ij}^{H}, F_{ij}^{X}\}](r) \]  

(7)

and determine \( v_{\text{Hxc}} \) using the orbitals and occupation factors. The orbitals themselves obey

\[ \left[-\frac{1}{2} \nabla^2 + v_s(r) - \epsilon_i\right] \phi_i(r) = 0 \]

(8)

so that we need to iterate to self-consistency. LEXX+KLI should (and we have found no evidence to the contrary) yield good approximations for the potential, orbital and density of atoms, at least up to relativistic effects.

To calculate the polarizability we need the density response function \( \chi \) from linear-response\(^\text{38} \) time dependent DFT (TDDFT). To find \( \chi \) we solve

\[ \chi(r, r'; i\omega) = \chi_0 + \chi_0 \ast f_{\text{Hxc}} \ast \chi, \]  

(9)

where the spin-symmetry allows us to ignore spin. Here the convolution \( \ast \) indicates an integral over the interior space variable such that \([f \ast g](r, r') = \int d r_2 f(r, r_2) g(r_2, r').\)

Equation (9) requires two inputs. Firstly, it needs the non-interacting response function \( \chi_0(r, r'; i\omega) \), which governs the change in density at \( r \) in response to changes in the Kohn-Sham potential \( v_s \) at \( r' \). For our calculations, we determine \( \chi_0 \) using

\[ \chi_0(r, r'; i\omega) = \text{2Re} \sum_i f_i \rho_i(r, r') G(r, r'; \epsilon_i - i\omega). \]

(10)

Here the KS Greens function \( G(r, r'; \epsilon_i - i\omega) = \text{2Re} \sum_j \frac{\rho_j(r', r)}{\epsilon_j - \epsilon_i + i\omega} \) obeys

\[ \left[-\frac{1}{2} \nabla^2 + v_s - \Omega\right] G(r, r'; \Omega) = -\delta(r - r'). \]

(11)

In our implementation we use the effective one dimensionality of the Hamiltonian to solve \( G \)
directly using (11) rather than using the sum form. This allows us to calculate very accurate Greens functions using a shooting method and the cusp condition.

Secondly, we need to evaluate the Hartree, exchange and correlation kernel

\[ f_{Hxc}(r, r'; i\omega) = \frac{\delta v_{Hxc}(r)}{\delta n(r')} |_{i\omega} = \frac{\delta v_{Hxc}(r')}{\delta n(r)} |_{i\omega}, \]  

which is not known exactly and must be approximated. In this work we employ two different approximations for the kernel \( f_{Hxc} \). For most calculations we use the Petersilka, Gossman, Gross approximation to the kernel adapted for LEXX. The resulting kernel takes the form

\[ f_{Hxc}^{PGG} = \sum_{ij} \left[ F_{ij}^{H} n_{i}(r)n_{j}(r') - \frac{1}{2} F_{ij}^{x}\rho_{i}(r,r')\rho_{j}(r',r) \right] n(r)n(r'), \]  

For additional tests we also employ the radial exchange hole (RXH) kernel\(^{32}\) similarly adapted to open-shell atoms. Both kernels are more accurate than the popular random phase approximation (RPA) due to their inclusion of dynamic screening effects and consequent reduction in self interaction errors (see e.g. Refs. 32,35).

Finally, we can calculate \( \chi(r, r'; i\omega) \) for a given \( N \) and \( Z \) using (9) and its inputs. \( \alpha(i\omega) \) follows from (2). More formally, we calculate the response \( \chi[v_{s}, \{ f_{i} \}, f_{Hxc}](r, r'; i\omega) \), as a functional of any spherically symmetric effective potential \( v_{s} \), any set of occupation factors \( \{ f_{i} \} \) obeying \( \sum_{i} f_{i} = N \), and any kernel \( f_{Hxc} \). In the two approximations presented here, \( f_{Hxc} \) is itself entirely determined by \( v_{s} \) and \( \{ f_{i} \} \), allowing us to drop the dependence on the kernel.

1.1.1 Numerical convergence

All calculations are carried out using a bespoke radial code which, together with the LEXX\(^{34}\) approximation, is designed to perform highly accurate calculations without problems with basis set convergence. The code makes use of the spherical symmetry of the Kohn-Sham potential and employs a radial grid to calculate orbitals and Greens functions using a shooting
method. Details of the method are described in Refs. \textsuperscript{32}–\textsuperscript{35} together with some details on convergence and accuracy.

As a result, our calculations are limited only by the number of radial grid points $N_r$ (we set $N_r = 576$ for larger atoms), the number of spherical harmonics included (we include up to $L = 10$), and the cutoff radius of the grid $r_m$ (typically $r_m \geq 24$ $a_0$ for larger atoms - at this radius tiny e.g. $n(r_m) \ll 10^{-12}$). We thus expect our calculations to be very well converged, with estimated numerical errors of no more than 1% (as an absolute worst case - we suspect most atoms and ions will be well within 0.5% of their converged values). For example, we have tested Lutetium ($Z = N = 71$ - a likely “bad” case due to its open $p$-shell and large number of electrons) against increases in the abscissae number and grid radius and found variation of under 0.2% in the $C_6$ coefficient and static polarizability (e.g. the $C_6$ coefficient ranges from 2546.2 for $N_r = 576$ and $r_m = 30.0$ to 2547.1 for $N_r = 640$ and $r_m = 24.0$, and 2547.7 for $N_r = 640$ and $r_m = 30.0$).

### 1.2 Corrected TDDFT calculations

While the TDDFT calculations described above give moderately accurate polarizabilities (often within 10% of high level theories) and $C_6$ coefficients (often within 20%), they cannot meet quantum chemical accuracy. To obtain true benchmarks we thus need to correct the TDDFT results using higher level calculations or experimental data.

Let us first consider the main sources of fundamental, methodological errors in our approach (as opposed to numerical errors, which we estimate to be under 1%, see Sec. 1.1.1). In error free calculations, the real frequency poles of $\chi$ and $\chi_0$ are respectively the excitation energies $E_k - E_l$ (for dipole transition excitations only), and the excitation energies of the Kohn-Sham system $\epsilon_k - \epsilon_l$ (again for dipole transitions only). Furthermore, the set of full transitions $E_k - E_l$ is related to the set of Kohn-Sham, transitions $\epsilon_k - \epsilon_l$ with corrections depending on $f_{Hxc}$. This relationship can be approximated as $E_k - E_l \approx \epsilon_k - \epsilon_l + \langle kl | f_{Hxc} | kl \rangle$. Methodological errors thus appear in two ways: i) via the Kohn-Sham orbitals $\phi_i$ and their
energies $\epsilon_i$ which are inaccurate due to errors in $v_s$ [see Eqs (10) and (11)]; and ii) via the approximate kernels which introduces additional errors to $\chi$ via the screening equation (9).

These errors influence the quality of the polarizability mostly via the lowest energy transition, a fact that we will exploit. Using perturbation theory we can write

$$\alpha(i\omega) \approx \sum_{kl} \frac{d_{kl}}{(E_k - E_l)^2 + \omega^2}$$

(14)

where $\sum d_{kl} = N$. Setting $\omega = 0$ in (14) shows that the static polarizability $\alpha(0)$ is dominated by the lowest energy transitions provided the dipole factor $d_{kl}$ is sufficiently large (which has been confirmed for many atoms and ions\textsuperscript{28}). $\alpha(0)$ is consequently more susceptible to errors in the energy difference than $\alpha(i\omega > 0)$. We thus expect that sensibly correcting for errors in $\alpha(0)$ will go a long way to correcting errors in $\alpha(i\omega)$ for all $\omega$.

Following Chu and Dalgarno\textsuperscript{30} we perform frequency rescaling to improve our frequency dependent polarizabilities and $C_6$ coefficients. We replace the raw TDDFT polarizability $\alpha^{\text{TDDFT}}$ by

$$\alpha(i\omega) \approx S^2 \alpha^{\text{TDDFT}}(iS\omega)$$

(15)

where $S = \sqrt{\alpha^{\text{Ref.}}(0)/\alpha^{\text{TDDFT}}(0)}$ is our scaling factor. This form guarantees $\alpha(i\omega \to \infty) \to N/\omega^2$ and also ensures that

$$\alpha(0) = \alpha^{\text{Ref.}}(0).$$

(16)

where $\alpha^{\text{Ref.}}$ is a reference benchmark static polarizability. Under the assumption that the low frequency polarizability is dominated by the smallest transition energy this is equivalent to correcting a poor TDDFT lowest transition energy using higher level data.

For our benchmarks we utilize a variety of different sources (Refs. 29, 39, 43) to find static polarizabilities that we consider to be optimal reference values. These are discussed
in Section 1 of the Supplementary Material. For most ions and Row 6 elements accurate dipole data are not available. In cases where data are not available we use: a) for neutral atoms we use an average scaling factor obtained from known cases in the same row; b) for ions we use the scaling coefficient of the atom with the same number of electrons.

We also produce a second set of data based on rescaled TDDFT with the PGG kernel that utilises a “minimal chemistry” model described in Appendix B. This allows us to extend our results to double anions and give a (hopefully) more realistic depiction of embedded open shell single anions while maintaining the presumed good results for atoms and cations. These polarizabilities are not intended to be compared to experimental data. But they provide a potentially more realistic starting point for atom-in-molecule approaches like vdW functionals and semi-classical molecular modelling.

## 2 Benchmarking tests

In our first test we check the accuracy of the rescaling of TDDFT results using equation (15). In Table 1 we compare $C_6$ coefficients from this approach with the same $C_6$ coefficients calculated using accurate quantum chemical theories. As can be seen, the agreement is generally very good, with the worst case (Ne) off by 8.4% (we cannot explain this difference, but note that both PGG and RXH kernels give similar results), and the second worst case (Cr) only 5.5% off the reference results despite our use of a different symmetry state. The $C_6$ coefficients obtained from our approach are also similar to those found by Chu and Dalgarno.

By using different TDDFT kernels we are able to provide a second test of the rescaling that includes systems for which quantum chemical data are unavailable. Our test is based on the following idea: if rescaling can give good dynamic polarizabilities by correcting TDDFT calculations using static polarizabilities, then rescaled $C_6$ coefficients of different species should be largely independent of the exchange-correlation kernel employed in the TDDFT
Table 1: Comparison of our scaled $C_6$ values with high-level wave-function method reference values. Results in $\text{Ha}a_0^6$.

| Species | Ref. value | This work | Err % | Ref. |
|---------|------------|-----------|------|-----|
| He      | 1.46       | 1.47      | 0.9  | 44  |
| Li      | 1396       | 1408      | 0.9  | 44  |
| Be$^+$  | 68.8       | 70.3      | 2.2  | 44  |
| Be      | 213        | 214       | 0.3  | 44  |
| Ne      | 6.38       | 6.91      | 8.4  | 44  |
| Na      | 1562       | 1566      | 0.2  | 44  |
| Mg$^+$  | 155        | 155       | 0.5  | 44  |
| Mg      | 630        | 629       | -0.1 | 44  |
| Ar      | 64.3       | 67.4      | 4.8  | 44  |
| K       | 3906       | 3914      | 0.2  | 44  |
| Ca$^+$  | 541        | 554       | 2.3  | 44  |
| Ca      | 2188       | 2232      | 2.0  | 44  |
| Cu      | 250        | 264       | 5.5  | 44  |
| Kr      | 130        | 136       | 4.8  | 44  |
| Rb      | 4667       | 4660      | -0.1 | 44  |
| Sr$^+$  | 776        | 790       | 1.9  | 44  |
| Sr      | 3149       | 3230      | 2.6  | 44  |
| Ag      | 342        | 341       | -0.3 | 44  |
| Xe      | 286        | 302       | 5.6  | 44  |
| Cs      | 6733       | 6657      | -1.1 | 44  |
| Ba$^+$  | 1293       | 1296      | 0.3  | 44  |
| Ba      | 5380       | 5543      | 3.0  | 44  |
| MAE     |             |           | 2.2  |     |
| ME      |             |           | 2.0  |     |
Figure 1: Normalised RXH $C_6$ coefficients versus PGG values. The dotted lines indicate a difference of 20\% between the two methods. Note that the worst outliers are all elements for which the rescaling parameter was interpolated based on neighbouring elements.
calculations. To test this hypothesis, we carried out calculations using two kernels: that of Petersilka-Gossman and Gross,\textsuperscript{36} and the “radial exchange hole” kernel of Gould.\textsuperscript{32} These two approaches give very different dipole polarizabilities before scaling, and can thus provide a clear test of the rescaling.

The results of our tests are plotted in Figure 1, in which rescaled RXH atomic $C_6$ coefficients are shown as a function of their PGG values. If all points fell perfectly on the straight line this would indicate that the rescaling was perfect, as two sets of inputs would have yielded the same set of $C_6$ coefficients after scaling. As it is, the values are all within 20\% of each other, with the worst cases all being elements for which accurate reference polarizability values were not available and the rescaling parameters had to be interpolated based on other elements in the same row.

We thus conclude that starting with a moderately accurate TDDFT kernel, and then rescaling based on static dipole polarizabilities, is a very accurate approach for determining dynamic polarizabilities. Not only do we get good agreement with more accurate approaches, but we also get agreement across three TDDFT approaches: that of Chu and Dalgarno,\textsuperscript{30} and the two kernels tested here.

3 Results

We show in Tables 2-4 static polarizabilities and same-species $C_6$ coefficients using the “benchmark” data set of rescaled atoms and ions. These are arranged by shell structure to make for easy access. As discussed previously, these are in good agreement with external reference data and, for the most part, are in good agreement with RXH values. The exceptions show no more than a 15\% difference, suggesting that this is a good estimate for the worst-case accuracy of our approach.

In addition to the tabulated static polarizabilities and $C_6$ coefficients, we also provide readily usable data for all cases considered here. Rather than provide tabulated frequency
dependent polarizability data for all 411 atoms and ions (or 443 for the minimal chemistry model), we instead provide parameters $a_{c,X}$ and $\Omega_{c,X}$ where $c = 1, 2$ for a two Lorentzian model

$$\alpha_X(i\omega) \approx \sum_{c=1,2} \frac{a_{c,X}}{\omega^2 + \Omega_{c,X}^2},$$

(17)

for species $X$ with nuclear charge $Z$ and $N$ electrons. Eq. (17) can then be used to accurately reproduce the polarizabilities for arbitrary frequency.

This two-Lorentzian representation of polarizabilities can be used to reproduce all $C_6$ coefficients.

Table 2: Static polarizabilities and $C_6$ coefficients from the “benchmark” data set for all neutral atoms. We note that the small error in Hydrogen’s $C_6$ coefficient of 13/2 is an artefact of the two-Lorentzian model. Results in $a_0^3$ for polarizabilities and $Hao_6^6$ for $C_6$ coefficients.

| ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ |
|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|
| He  | 4.90   | 0.51  | Be  | 3.84  | 1.17  | Li  | 164   | 1410 | Na  | 163   | 99.2 | C  | 11.7  | 47.9 | N  | 7.25  | 25.7 |
| B   | 20.5   | 55.9  | O   | 3.43  | 1.47  | F   | 3.60  | 10.2 | Ne  | 2.67  | 6.91 | Ar  | 4.50  | 0.51 | He  | 3.84  | 1.17 |

Table 3: Static polarizabilities and $C_6$ coefficients from the “benchmark” data set for all cations. Results in $a_0^3$ for polarizabilities and $Hao_6^6$ for $C_6$ coefficients.

| ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ | ID  | $a(0)$ | $C_6$ |
|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|-----|--------|-------|
| He$^+$ | 0.294 | 0.109 | Li$^+$ | 0.193 | 0.079 | Be$^+$ | 24.5 | 70.3 | B$^+$ | 9.67 | 25.2 | C$^+$ | 5.66 | 13.3 | N$^+$ | 3.68 | 8.02 |
| Mg$^{2+}$ | 35.0 | 155 | Al$^{3+}$ | 19.6 | 89.7 | Si$^+$ | 18.2 | 94.8 | P$^{5+}$ | 14.3 | 74.1 | S$^+$ | 11.9 | 62.2 | Cl$^+$ | 9.27 | 46.1 |
| Ca$^{2+}$ | 75.5 | 554 | Sc$^{3+}$ | 60.0 | 531 | Ti$^{4+}$ | 44.6 | 286 | V$^{6+}$ | 36.5 | 217 | Cr$^{6+}$ | 38.2 | 230 | Mn$^{2+}$ | 25.8 | 132 |
| Ni$^{2+}$ | 18.4 | 81.5 | Cu$^{2+}$ | 17.6 | 75.9 | Zn$^{2+}$ | 17.9 | 91.7 | Ga$^{2+}$ | 15.2 | 71.8 | Ge$^{2+}$ | 18.5 | 108 | As$^{5+}$ | 16.4 | 102 |
| Sr$^{2+}$ | 90.2 | 790 | Y$^{3+}$ | 88.1 | 1020 | Zr$^{2+}$ | 62.0 | 574 | Nb$^{5+}$ | 55.3 | 512 | Mo$^{6+}$ | 47.6 | 424 | Tc$^{6+}$ | 32.7 | 246 |
| Pd$^{2+}$ | 31.5 | 248 | Ag$^{1+}$ | 20.0 | 115 | Cd$^{2+}$ | 23.1 | 155 | In$^{3+}$ | 20.2 | 127 | Sn$^{2+}$ | 29.1 | 237 | Sb$^{5+}$ | 20.0 | 115 |
| Ba$^{2+}$ | 121 | 1300 | La$^{3+}$ | 94.2 | 1180 | Ce$^{3+}$ | 89.1 | 1080 | Pr$^{4+}$ | 93.4 | 1160 | Nd$^{4+}$ | 89.6 | 1080 | Pm$^{5+}$ | 85.7 | 1010 |
| Sm$^{3+}$ | 81.8 | 941 | Eu$^{4+}$ | 78.1 | 876 | Gd$^{5+}$ | 67.2 | 698 | Tb$^{6+}$ | 71.8 | 769 | Dy$^{7+}$ | 68.6 | 717 | Ho$^{8+}$ | 65.8 | 672 |
| Er$^{8+}$ | 63.0 | 630 | Tm$^{9+}$ | 60.4 | 590 | Yb$^{10+}$ | 58.0 | 555 | Lu$^{11+}$ | 74.6 | 808 | Tb$^{12+}$ | 84.6 | 1220 | Yb$^{12+}$ | 84.6 | 1220 |
| Hf$^{4+}$ | 47.5 | 443 | Ta$^{5+}$ | 42.0 | 388 | W$^{6+}$ | 37.1 | 333 | Re$^{7+}$ | 33.4 | 290 | Os$^{8+}$ | 30.4 | 256 | Ir$^{9+}$ | 27.9 | 228 |
| Pt$^{6+}$ | 25.8 | 204 | Au$^{7+}$ | 24.0 | 184 | Hg$^{8+}$ | 17.5 | 114 | Ti$^{10+}$ | 17.0 | 109 | Ta$^{11+}$ | 20.2 | 190 | Ru$^{12+}$ | 22.4 | 236 |
Table 4: Static polarizabilities and $C_6$ coefficients from the “benchmark” data set for all anions. Results in $a_0^3$ for polarizabilities and Ha$^a_6$ for $C_6$ coefficients.

| ID | $\alpha(0)$ | $C_6$ ID | $\alpha(0)$ | $C_6$ ID | $\alpha(0)$ | $C_6$ ID | $\alpha(0)$ | $C_6$ ID | $\alpha(0)$ | $C_6$ ID |
|----|-------------|----------|-------------|----------|-------------|----------|-------------|----------|-------------|----------|
| H^- | 216 | 2400 | | | | | | | | |
| Li^- | 1180 | 35170 | | | | | | | | |
| B^- | 32.9 | 232 | C^- | 15.5 | 81.7 | N^- | 8.04 | 33.1 | O^- | 5.40 | 19.1 | F^- | 15.0 | 73.5 |
| Na^- | 1310 | 42690 | | | | | | | | |
| Al^- | 109 | 1550 | Si^- | 59.6 | 698 | P^- | 33.7 | 322 | S^- | 24.1 | 206 | Cl^- | 30.3 | 276 |
| K^- | 2090 | 87490 | | | | | | | | |
| Sc^- | 134 | 1860 | Ti^- | 110 | 1380 | V^- | 94.5 | 1100 | Cr^- | 83.5 | 908 | Mn^- | 74.7 | 762 | Fe^- | 66.5 | 640 |
| Co^- | 59.8 | 545 | Ni^- | 54.2 | 470 | Cu^- | 500 | 9440 | | | | | | | |
| Zn^- | 845 | 17790 | Ga^- | 103 | 1430 | Ge^- | 63.3 | 796 | As^- | 40.0 | 443 | Se^- | 31.7 | 332 |
| Rb^- | 2110 | 92240 | | | | | | | | | |
| Y^- | 175 | 3080 | Zr^- | 134 | 2160 | Nb^- | 114 | 1680 | Mo^- | 100 | 1390 | Tc^- | 90.6 | 1170 | Ru^- | 82.4 | 1010 |
| Rh^- | 75.9 | 881 | Pd^- | 70.5 | 779 | Ag^- | 501 | 9800 | | | | | | | |
| Cd^- | 863 | 20230 | In^- | 133 | 2240 | Sn^- | 89.1 | 1410 | Sb^- | 59.6 | 859 | Te^- | 49.1 | 684 | I^- | 61.7 | 925 |
| Ce^- | 2480 | 118890 | La^- | 729 | 19310 | Ce^- | 4.02 | 46.5 | Pr^- | 4.88 | 62.7 | Nd^- | 5.57 | 77.0 | Pm^- | 6.50 | 97.9 |
| Sn^- | 7.38 | 119 | Eu^- | 8.08 | 138 | Gd^- | 8.08 | 198 | Dy^- | 10.5 | 210 | Ho^- | 11.1 | 230 |
| Er^- | 11.6 | 246 | Tm^- | 12.0 | 260 | | | | | | | | | | |
| Lu^- | 13.8 | 325 | Hf^- | 236 | 4170 | Ta^- | 206 | 3640 | W^- | 295 | 5350 | Re^- | 368 | 6710 | Os^- | 378 | 6840 |
| Ir^- | 381 | 6830 | Pt^- | 383 | 6810 | Au^- | 387 | 6840 | | | | | | | | |
| Ti^- | 356 | 6820 | Pb^- | 135 | 2310 | Bi^- | 103 | 1710 | Po^- | 73.1 | 1140 | At^- | 54.5 | 801 |

coefficients using analytic solutions of the Casimir-Polder formula (1) which take the form (here $X \equiv Z_X, N_X$ and $Y \equiv Z_Y N_Y$)

$$C_{6,XY} = \sum_{cc'} \frac{3a_{c,X}a_{c',Y}}{2\Omega_{c,X}\Omega_{c',Y}[\Omega_{c,X} + \Omega_{c',Y}]}.$$  \hfill (18)

Results are within a few percent compared of similar calculations carried out on the raw data (see Appendix C for details). Our two-Lorentzian model thus provides a useful and compact representation for all dynamic polarizabilities, allowing reconstruction of the 84000+ different-species $C_6$ coefficients. These data are included in the supplementary material, tabulated for the “benchmark” set and included in ascii files for all data sets.

We also test interactions of rare gas atoms with halides and the alkali cation Na$^+$, to demonstrate the method’s capabilities on heteronuclear $C_6$ coefficients. We compare results from our data set against results from three references, and are thus able to cover elements in all of the six rows considered here. Results are presented in Table 5. Agreement is generally excellent, with a few cases substantially worse than the mean absolute relative error of 3.7%.

The worst relative errors all involve Na$^+$ and rare gases. However, the absolute errors for these cases are small since neither Na$^+$ nor rare gas atoms are very polarizable and the $C_6$ coefficients are consequently small. Also, the data from Ref. 48 may not be as accurate as
Table 5: Interaction of rare gas atoms with hailde atoms and ions and Na+. Results in Ha\(\alpha\)\(_6\).

|                        | \(C_6\) [Ref] | \(C_6\) [this work] | % Err | Ref. |
|------------------------|---------------|----------------------|-------|------|
| \(\text{Br}^+\)–He    | 12.0          | 11.3                 | -5.8  | 47   |
| \(\text{Br}^-\)–He    | 15.0          | 15.8                 | 5.1   | 47   |
| \(\text{Br}^+\)–He    | 27.0          | 24.0                 | -11.0 | 47   |
| \(\text{Br}^-\)–Ne    | 23.0          | 23.7                 | 3.2   | 47   |
| \(\text{Br}^-\)–Ne    | 31.0          | 32.8                 | 5.8   | 47   |
| \(\text{Br}^+\)–Ne    | 48.0          | 49.4                 | 2.9   | 47   |
| \(\text{Br}^-\)–Ar    | 78.0          | 78.3                 | 0.4   | 47   |
| \(\text{Br}^-\)–Ar    | 110           | 111                  | 0.9   | 47   |
| \(\text{Br}^-\)–Ar    | 174           | 174                  | -0.3  | 47   |
| \(\text{Na}^+\)–He    | 1.79          | 1.39                 | -22.2 | 48   |
| \(\text{Na}^+\)–Ne    | 3.22          | 3.15                 | -2.2  | 48   |
| \(\text{Na}^+\)–Ar    | 10.4          | 8.71                 | -15.9 | 48   |
| \(\text{F}^-\)–He    | 9.37          | 9.81                 | 4.7   | 49   |
| \(\text{F}^-\)–Ne    | 19.4          | 20.5                 | 5.6   | 49   |
| \(\text{F}^-\)–Ar    | 66.4          | 69.0                 | 3.8   | 49   |
| \(\text{F}^-\)–Kr    | 95.2          | 98.9                 | 3.9   | 49   |
| \(\text{F}^-\)–Xe    | 143           | 148                  | 3.7   | 49   |
| \(\text{F}^-\)–Rn    | 170           | 172                  | 1.4   | 49   |
| \(\text{Cl}^-\)–He    | 19.1          | 18.3                 | -4.0  | 49   |
| \(\text{Cl}^-\)–Ne    | 39.4          | 37.8                 | -4.1  | 49   |
| \(\text{Cl}^-\)–Ar    | 138           | 131                  | -4.8  | 49   |
| \(\text{Cl}^-\)–Kr    | 198           | 190                  | -4.3  | 49   |
| \(\text{Cl}^-\)–Xe    | 299           | 286                  | -4.3  | 49   |
| \(\text{Cl}^-\)–Rn    | 357           | 333                  | -6.7  | 49   |
| \(\text{Br}^-\)–He    | 24.0          | 24.0                 | 0.2   | 49   |
| \(\text{Br}^-\)–Ne    | 49.4          | 49.4                 | -0.0  | 49   |
| \(\text{Br}^-\)–Ar    | 174           | 174                  | -0.3  | 49   |
| \(\text{Br}^-\)–Kr    | 251           | 251                  | 0.0   | 49   |
| \(\text{Br}^-\)–Xe    | 380           | 380                  | 0.1   | 49   |
| \(\text{Br}^-\)–Rn    | 452           | 443                  | -1.9  | 49   |
| \(\text{I}^-\)–He    | 32.1          | 32.1                 | -0.0  | 49   |
| \(\text{I}^-\)–Ne    | 65.9          | 65.8                 | -0.2  | 49   |
| \(\text{I}^-\)–Ar    | 233           | 233                  | -0.0  | 49   |
| \(\text{I}^-\)–Kr    | 336           | 338                  | 0.5   | 49   |
| \(\text{I}^-\)–Xe    | 510           | 514                  | 0.7   | 49   |
| \(\text{I}^-\)–Rn    | 608           | 599                  | -1.5  | 49   |
more modern calculations would allow. The next worst case is Br\(^-\) with He when compared
with the reference data of Ref. 47. Oddly, we get almost perfect agreement for this case with
the (presumably less accurate) TDMP2 results of Ref. 49.

Figure 2: Dispersion coefficients for pair of species \(X\) and \(Y\) (neutral atoms from first six
rows of PT) plotted against \(\alpha_{XY}(0) \equiv \sqrt{\alpha_X(0)\alpha_Y(0)}\) (red crosses) compared with best fit
function (blue line). Units are \(a_0^3\) for polarizabilities and \(Ha_0^6\) for \(C_6\) coefficients. Inset shows
the same as the main plot but zooms into the most populated range of polarizabilities and
coefficients.

Finally, we use our comprehensive data set to test for broad relationships between
\(C_6\) coefficients and static dipole polarizabilities \(\alpha(0)\). Based on units, one predicts that
\(C_{6,XY} \propto \alpha_X(0)\alpha_Y(0)\) might be a reasonable approximation for such a relationship. However,
after testing all 3741 possible pairs of neutral atoms (plotted in Figure 2) we find that

\[ C_{6,XY} \approx \Xi \left[ \alpha_X(0)\alpha_Y(0) \right]^{0.73 \pm 0.01}, \quad (19) \]

\( \Xi = (1.5 \pm 0.1) \left[ \text{Ha} a_{1.62} \right], \) is a better power law fit, showing a somewhat surprising reduction in the scaling coefficient from the expected product form. Here our parameter error bars are crudely estimated by comparing more limited fits with only same-species atomic and ionic coefficients, and all neutral atom pairs using our PGG and RXH data sets.

The constant prefactor \( \Xi \) varies considerably depending on the data set selected, and Figure 2 makes it clear that the actual results have a much wider range of values. By contrast, the exponent \( 0.73 \pm 0.01 \) remains essentially constant. This suggests a certain universality of the power law relationship between polarizabilities and \( C_6 \) coefficients. We suspect that it might be possible to improve the quality of this fit by improving the dependence on \( \alpha_x(0) \) and \( \alpha_y(0) \) separately.

Along these lines, we test the quality of the relationship

\[ C_{6,XY} = \frac{2C_{6,XX}C_{6,YY}}{\alpha_X(0)C_{6,YY} + \alpha_Y(0)C_{6,XX}}, \quad (20) \]

found\(^{150}\) from a one-Lorentzian model \( \alpha(i\omega) = \alpha(0)/(1 + \omega^2/\eta^2) \) (equivalent to the \([1,0]\) Padé approximation for the polarizability). This has previously been used (e.g. Ref. \( ^{20} \)) to derive different-species coefficients from same-species data. We find that (20) is generally very accurate, giving answers within 5% of our two-Lorentzian model \( ^{18} \) for almost 80% of cases (from 84000 pairs of atoms, anions and cations). However in 2.0% (\( \approx 1700 \)) of cases it is more than 20% out and in 0.2% (\( \approx 200 \)) of cases it is more than 30% out.

It should be noted that most of these worst case examples involve 2+ or 3+ cations interacting with an anion, and thus give rise to small absolute errors. None of the cases with > 20% errors involve two neutral atoms. Some notable bad cases examples are Na\(^+\) with Cs (-21%), and Al\(^-\) with many anions in Row 6 (\( \approx -24\% \)).
4 Conclusions

Using TDDFT with the PGG\textsuperscript{36} kernel we calculated all-electron dipole polarizabilities of all atoms and many cations and anions from rows 1-6 (1 ≤ Z ≤ 86) of the periodic table. We also performed calculations using an alternative RXH\textsuperscript{32} kernel; and using a variant method designed to approximate more realistic environmental effects for atoms in molecules, which yields different coefficients for open-shell anions. We argue that these dipole polarizabilities can provide a rough benchmark (likely within 15% for all species, with better results expected in some cases) for imaginary frequency dipole polarizability calculations of $C_6$ coefficients. They are almost certainly of sufficient quality to be used in atom-in-molecule (AIM) approaches, be they classical, semi-classical, or semi-empirical.

Our polarizabilities were parametrised using a two-Lorentzian model\textsuperscript{17} that can be used to reproduce $C_6$ coefficients within 5% (and typically <1%) of the value obtained by full quadrature via the Casimir-Polder formula\textsuperscript{(1)}. These parameters were tabulated for all atoms and ions investigated, and are included in the supplementary materials.\textsuperscript{46} These were used to calculate homonuclear-isoelectronic $C_6$ coefficients and some heteronuclear coefficients.

We finally used our data to study the dependence of $C_6$ coefficients on polarizabilities, finding $C_{6,XY} \approx \Xi[\alpha_X(0)\alpha_Y(0)]^{0.73\pm0.01}$, with error bars indicating the spread of best-fit parameters found on different data sets. While the prefactor $\Xi$ was found to vary considerably depending on the data set used to make the fit, the exponent varied much less suggesting it is a more universal quantity. Similarly, we tested the relationship $C_{6,XY} = 2C_{6,X}C_{6,Y} / [\alpha_X/\alpha_YC_{6,Y} + \alpha_Y/\alpha_XC_{6,X}]$ sometimes used to relate homonuclear-isoelectronic coefficients and $C_6$ for pairs of unlike species (atoms or ions). We found that it gave errors of less than 5% in 80% of cases, but in 0.2% of cases gave very poor results with > 30% errors.

In future we also hope to use our approach to study atoms and ions in Row 7, after developing techniques to deal with relativistic effects and fixed core approximations. We also aim to explore other environmental effects to better understand how embedded atoms...
behave compared to their free counterparts, and thus to improve and extend the “minimal chemistry” model presently used to determine some cation and all double cation dipole polarizabilities.

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**Supporting Information Available**

We include with this manuscript supporting information including:

A PDF document that includes: a) further details of the benchmark data set; b) several additional tables of chief results; c) a description of the accompanying ascii files;

Several ascii files presenting chief results in a readily usable format.

This material is available free of charge via the Internet at [http://pubs.acs.org/](http://pubs.acs.org/).

**A Transition metal and lanthanide atoms and ions**

Some of the transition metal and lanthanide atoms and ions represent very difficult cases for single-determinant theories like DFT and HF. In certain cases the $s$ and $d$ orbitals (or $d$ and $f$ orbitals) are so close to degenerate that the aufbau principle can be violated in the lowest energy configuration for DFT calculations (see e.g. Ref. 51 or Ref. 52 for
discussion in a Hartree-Fock context). Despite the LEXX approximation including some multi-determinental characteristics via the ensemble DFT formalism it too is affected by these issues. Furthermore, the route of explicit symmetry breaking employed by Johnson et al is not available in our radial code.

This poses a particular challenge for polarizability problems as the effective Kohn-Sham gap $\epsilon_l - \epsilon_h$ (for lowest unoccupied $f_l = 0$ and highest occupied $f_h > 0$ KS orbitals) can become negative. Since this gap appears in the denominator of terms contributing to $\chi_0$ it leads to unphysical small frequency (real or imaginary) polarizabilities. In certain cases it gives negative bare polarizabilities (i.e. calculated via (2) except with $\chi_0$ rather than $\chi$). While these problems are somewhat mitigated in the interacting response $\chi$ by the screening kernel $f_{\text{Hxc}}$ (e.g. our approach returns positive static polarizabilities for all species tested), it still contributes substantial errors to polarizabilities.

We are thus left with a dilemma: do we choose the lowest energy groundstate or choose a state that does not give negative transitions but is in a less realistic electronic configuration? We thus decide to simplify matters by filling the orbitals according to $(n+1)s^2(n)d^m$, where $n = 3$ or 4 (or equivalent for $d$ and $f$ orbitals). This leads to some differences from free atom calculations, especially in Cr and Mo which have clearly lower energies in their $(n+1)s^1(n)d^5$ states. However, as these elements usually appear as embedded ions we expect the practical effect to be minimal.

## B Minimal chemistry model

The main purpose of this manuscript is to report reference polarizabilities and $C_6$ coefficients for free-standing atoms and ions (for ions with $N \leq Z+1$). However, polarizabilities are often desired for their utility in embedded atom/ion models - such as for high-level calculations using van der Waals dispersion corrections (e.g. Refs. 17–22), or for (semi-)classical force-field models. We thus also report a slightly modified set of reference coefficients for this
Atoms and ions “embedded” in a larger system such as a molecule or material can behave very differently to their free-standing counterparts. Most notably, the surrounding environment of an anion has a substantial effect on the behaviour of its outermost electron(s). They are only very weakly bound in the free-standing case, with an asymptotic effective potential going to zero as \( r^{-3} \). When embedded, the other electrons and nuclei will introduce an effective confining potential, which can be approximated [see e.g. Ref. 53] by a positive power law function of \( r \) such as \((r/r_a)^{\sigma_a}\) where \( r_a \) is an effective embedding radius and \( \sigma_a \) governs the sharpness. Clearly the energetics of the outermost orbital and unoccupied orbitals are very different in both cases.

But the same electrons that are most sensitive to the embedding environment, namely electrons in the outermost electronic shell(s), are the ones that contribute the most to the polarizability. Thus the polarizability of an embedded system is highly sensitive to its environment and care must be taken in considering what “anions” should be used in embedding theories. This is especially pertinent for open shell systems which are likely to be the most sensitive to the environment.

To account for embedding, we thus approximate embedded anions using a minimal chemistry model chosen to ensure that the “free” ions behave as closely as possible to their embedded counterparts without taking into account the full details of the chemical environment. To this end, we carry out polarizability calculations using a frozen orbital model, in which the polarizabilities \( \alpha \equiv \alpha[v_s[v_Z],\{f_i\}] \) are treated (in TDDFT) as a functional of the interacting potential \( v_s \) via \( \chi[v_s,\{f_i\},f_{\text{Hxc}}](r,r';i\omega) \), discussed in detail just before Sec. 1.1.1. This allows us to introduce some embedding effects via our choice of \( v_s \) arising from external potential \( v_Z = -Z/r \).

Our minimal chemistry model involves determining \( v_s \) from the following rules:

1. For neutral atoms and cations, find the atomic potential self-consistently using LEXX theory, Hund’s rules and the aufbau principle, (see Appendix A for further details on
Rows 4-6) then rescale the polarizabilities using (15);

2. For single anions $A^{-}$ of halides ($N = 10, 18, 36, 54, 86, Z = N - 1$) calculate the ionic potential self-consistently using LEXX theory and then rescale (like for neutral atoms);

3. For the remaining single anions $A^{-}$ use the self-consistent LEXX potential of the neutral atom and do not rescale;

4. For all double anions $A^{2-}$ use the self-consistent LEXX potential of the neutral atom and do not rescale;

This method produces an alternative set of frequency dependent anionic polarizabilities that, we feel, may better reflect the reality of embedded anions. This alternative data set is provided in the supplementary data.46

C Two-Lorentzian polarizability model

From equation (14)

$$\alpha (i\omega) \approx \sum_{kl} \frac{d_{kl}}{(E_k - E_l)^2 + \omega^2}$$

it is clear that the imaginary frequency dependence of $\alpha$ is a sum over Lorentzian functions $d/(\omega^2 + \Omega^2)$. In a typical atom one Lorentzian will typically have a denominator (the square of the lowest excitation energy + $\omega^2$) that is significantly smaller than the other terms for small $\omega$, and thus dominates the $C_6$ coefficient. When combined with the known limit $\lim_{\omega \to \infty} \alpha (i\omega) = N/\omega^2$ (i.e. $\sum d_{kl} = N$) this suggests that a reduced number of Lorentzians should be sufficient to represent the imaginary frequency dipole polarizabilities.

In fact, our results suggest that $\alpha (i\omega)$ for atoms/ions can be approximated by just two Lorentzians with minimal loss of accuracy. Thus, for every atom and ion with nuclear charge
Z and electron number N we write

\[ \alpha_X (i\omega) \approx \sum_{c=1,2} \frac{a_{c,X}}{\omega^2 + \Omega_{c,X}^2} \]  

without any great loss of accuracy. Here the parameters \( a_1, \Omega_1 \) and \( \Omega_2 \) are varied to minimize \( \int (\alpha - \alpha^{\text{Numeric}})^2 d\omega \) while \( a_2 = N - a_4 \) is kept fixed to ensure that the polarizability has the correct asymptote.

We note that Figari et al.\(^{54,55}\) have carefully studied similar pseudospectral methods to the one employed here. They showed that four Lorentzians are generally sufficient for very high accuracy and that careful treatment of the frequencies \( \Omega_c \) can further improve results. However, given the merely “moderate” quality of our inputs, we feel that such an analysis would not offer meaningful benefits for our present work.

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