Investigating the Heaviest Halogen: Lessons

**Learned from Modeling the Electronic Structure** 

of Astatine's Small Molecules

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Abstract

We present a systematic study of electron-correlation and relativistic effects in di-

atomic molecular species of the heaviest halogen astatine (At) within relativistic single-

and multi-reference coupled-cluster approaches and relativistic density functional the-

ory. We establish revised reference ab initio data for the ground states of At<sub>2</sub>, HAt,

AtAu, and AtO<sup>+</sup> using a highly accurate relativistic effective core potential model and

in-house basis sets developed for accurate modeling of molecules with large spin-orbit

effects. Spin-dependent relativistic effects on chemical bonding in the ground state

are comparable to the binding energy or even exceed it in At<sub>2</sub>. Electron-correlation

effects near the equilibrium internuclear separation are mostly dynamical and can be

adequately captured using single-reference CCSD(T). However, bond elongation in

At<sub>2</sub> and, especially, AtO<sup>+</sup> results in rapid manifestation of its multi-reference char-

acter. While useful for evaluating the spin-orbit effects on the ground-state bonding

and properties, the two-component density functional theory lacks predictive power,

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especially in combination with popular empirically adjusted exchange-correlation functionals. This drawback supports the necessity to develop new functionals for reliable quantum-chemical models of heavy-element compounds with strong relativistic effects.

### 1 Introduction

Apart from superheavy elements, the rarest naturally occurring element and the heaviest halogen astatine  $_{85}$ At remains one of the most "enigmatic" in the periodic table. <sup>1,2</sup> The <sup>211</sup>At isotope's unique properties have been attracting much attention in the nuclear medicine community for several decades. As a pure  $\alpha$ -emitter, <sup>211</sup>At is ideally suited for targeted therapy of malignant tumors due to its half-life, linear energy transfer, and the absence of harmful radioactive or toxic decay products or harmful deceleration  $\gamma$ -rays. <sup>3-6</sup> However, <sup>211</sup>At's scarcity due to production complexity and cost keeps its experimental chemistry elusive, thus creating a major obstacle in translating accelerator-generated elemental <sup>211</sup>At to radiopharmaceuticals. <sup>7</sup>

The absence of spectroscopic data even for diatomic species, such as  $At_2$ , HAt, or  $AtO^+$ , leaves state-of-the-art ab initio quantum-chemical calculations the sole source of basic reference data on the equilibrium internuclear distance  $R_e$ , dissociation energy  $D_e$ , and vibrational frequency  $\omega_e$ . As a heavy element, astatine exhibits strong relativistic effects that extend to its valence electrons. The spin-orbit interaction causes substantial energetic and spatial differences within  $5p_{1/2}-5p_{3/2}$ ,  $5d_{3/2}-5d_{5/2}$ , and  $6p_{1/2}-6p_{3/2}$  shells, thereby affecting correlations of the subvalence and valence electrons, which results in a potential departure of the astatine's chemical behavior from other halogens. Methodological complexities of treating the interplay of relativistic and correlation effects lead to substantially varying predictions even across recent advanced models. For example, the CCSD(T) version of a two-component relativistic coupled-cluster calculation (2c-CCSD(T)) of  $At_2$  in Ref. 9 with the X2C Hamiltonian and the complete basis set (CBS) extrapolation across Dyall's acvXz (X = 2, 3, 4) correlates 5d-, 6s-, and 6p-electrons and yields  $D_e(At_2)$ = 0.791 eV. The authors of Ref. 9 mention in-

significant differences between single-reference CCSD and multi-reference Fock-space CCSD (FS CCSD) results, albeit without giving specific numbers, and judge in favor of 2c-CCSD(T) as the method of choice. Using a similar argument, the authors of Ref. 10 conclude that the  $0_g^+$  ground state of At<sub>2</sub> is predominantly single-reference as their multi-reference configuration interaction MRCISD+Q results ( $R_e$ =3.046 Å,  $D_e$ =0.68 eV) agree with earlier four-component (4c-) CCSD(T) calculations ( $R_e$ =3.046 Å,  $D_e$ =0.63 eV). This plane is a specific number of the constant of the configuration interaction of the configuration interaction of the configuration interaction of the configuration interaction of the configuration of the confi

A seminal publication by Sergentu et al. <sup>13</sup> puts to the test conventional quantum-chemical recipes based on Density Functional Theory (DFT) to describe or predict the chemical behavior at the bottom of the periodic table using small molecules and molecular ions of astatine At<sub>2</sub>, HAt, AtF<sub>3</sub>, and AtO<sup>+</sup> as a case in point. While several popular exchange-correlation functional (XCF) approximations such as B3LYP, <sup>14</sup> PBE0, <sup>15,16</sup> or HSE06 <sup>17–19</sup> display reasonable behavior, the authors do not assess their reliability for bond strengths and recognize a gap in the DFT development for heavy-element chemistry. To complicate matters even further, a recent article by Aebersold and Wilson <sup>20</sup> demonstrates that for heavy-element molecules from the An66 database modeled with spin-orbit relativistic density functional theory (RDFT), errors from the standard basis set families are comparable to those from XCFs. The entanglement of these two major sources of errors renders the quality of spin-orbit RDFT models too chaotic to propose reasonable computational recipes.

The objective of this article is threefold. First, we establish new high-level *ab initio* reference data for diatomic At species. Second, we analyze potential multi-reference effects and spin-orbit-correlation interference. Third, based on the new *ab initio* results, we assess several common relativistic DFT approaches and delineate the role of XCF approximations from basis set errors. In addition to the previously mentioned At<sub>2</sub> and HAt, we investigate the AuAt molecule and AtO<sup>+</sup> molecular ion. Astatine-gold interactions attract attention due to the growing interest in gold clusters as At carriers in new therapeutic strategies. <sup>21,22</sup> Also, At may bear chemical similarities to the superheavy element nihonium (113Nh) and serve as a model system to calibrate thermochromatographic experiments on Nh identification and characterization based on its interaction with a gold surface. <sup>23,24</sup> Finally, the AtO<sup>+</sup> molecular cation's chemistry plays a key role in understanding astatine's behavior in common solvents. <sup>25,26</sup>

## 2 Computational details

## 2.1 Effective core potentials and basis sets

The focus on chemical properties justifies the replacement of At and Au core electrons with relativistic effective core potentials (RECPs). In this work, we use accurate small-core shape-consistent RECPs<sup>27</sup> (available on the Petersburg Nuclear Physics Institute Quantum Chemistry Laboratory website<sup>28</sup>). Optimized for valence shells, these RECPs also offer a reasonable accuracy in describing outer-core-valence correlations involving 5s-, 5p-, and 5d-shells of At and Au atoms.

To extrapolate the results of correlated wavefunction calculations to the complete basis set (CBS) limit, we develop triple-, quadruple-, and quintuple- $\zeta$  (respectively, TZ, QZ, and 5Z) correlation-consistent basis sets tailored for the chosen RECP model. Our correlation-consistent basis sets are an adaptation and extension of those accompanying Stuttgart-Cologne energy-adjusted RECPs.<sup>29,30</sup> For both At and Au, we start with the primitive

Gaussian exponents already optimized for the cc-CVnZ-PP $^{31,32}$  families of basis sets. Modifications apply to the contracted s-, p-, and d-functions to account for a different functional form of the RECP operator. In particular, such modifications are necessary for the basis sets to reflect the effects of spin-orbit interaction.  $^{33-35}$  To this end, we decontract the original functions and perform two-component spin-orbit Hartree–Fock calculations for atomic (or ionic) species. These calculations result in atomic spinors, i.e., two-component and generally complex extensions of the usual atomic orbitals, whose coefficients we then use to create new contracted basis functions. The resulting basis sets feature different contractions for  $p_{1/2}$  and  $p_{3/2}$ , as well as  $d_{3/2}$  and  $d_{5/2}$  functions, reflecting the spatial difference between spin-orbit-split p- and d-shells. We do not modify the original sets' uncontracted core-valence correlation functions, polarization, or diffuse Gaussian exponents.

For a statine basis sets, it is convenient to create all  $5s_{1/2}$ ,  $5p_{1/2}$ ,  $5p_{3/2}$ ,  $5d_{3/2}$ ,  $5d_{5/2}$ ,  $6s_{1/2}$ ,  $6p_{1/2}$ , and  $6p_{3/2}$  contracted functions based on the two-component Hartree–Fock calculations for a closed-shell anion At<sup>-</sup>. Working with the open-shell solution for an At atom is more complicated and results in a virtually indistinguishable basis set. The situation with gold is somewhat cumbersome due to a narrower  $5d_{5/2}$ – $6s_{1/2}$  energy gap and the presence of low-lying  $6p_{1/2}$  and  $6p_{3/2}$  virtual spinors. As advised in Ref. 31, it is necessary to derive flexible basis sets from  $5d^{10}6s^1$ ,  $5d^96s^2$ , and  $5d^86s^26p^1$  atomic states. To this end, we implement the following scheme. Core-like  $5s_{1/2}$ ,  $5p_{1/2}$ , and  $5p_{3/2}$  contracted functions represent the corresponding Hartree–Fock spinors for Au<sup>+</sup>. To derive sub-valence  $5d_{3/2}$  and  $5d_{5/2}$  and valence  $6s_{1/2}$  spinors, we use an average-of-configuration open-shell Au<sup>0</sup> solution that covers  $5d^{10}6s^1$  and  $5d^96s^2$  atomic states. Finally,  $6p_{1/2}$  and  $6p_{3/2}$  contractions result from the average-of-configuration open-shell Au<sup>0</sup> solution for the  $5d^86s^26p^1$  state. We calculate all these solutions using the DIRAC19<sup>36</sup> implementation of the relativistic 2c-Hartree–Fock method.

The CBS limit extrapolation procedure in DFT, unlike wavefunction correlated methods, is not well-established.<sup>37</sup> To approach the CBS limit in RDFT calculations, we develop

an adaptation of the universal Gaussian basis set (UGBS). <sup>38</sup> Our modifications include the removal of core-like exponents for heavy atoms with RECPs, adding extra polarization functions, and further augmentation. Without universally accepted basis sets for RDFT calculations, our modified UGBS approach attains sufficient flexibility without developing numerical instabilities, at least for small systems. The systematic development of optimal basis sets for RDFT calculations is beyond the scope of this article. The parameters of all basis sets developed for correlated wavefunction and RDFT calculations are given in the Supplement.

### 2.2 Relativistic coupled-cluster calculations

We perform scalar one-component (1c-) and spin-orbit two-component (2c-) single-reference CCSD(T) simulations using the DIRAC19 program. <sup>36</sup> For At and Au, we employ aug-cc-pwCVnZ-PP (n=3–5) basis sets modified for spin-orbit calculations with shape-consistent RECPs, as described in Sec. 2.1, while for H and O we use the original aug-cc-pVnZ and aug-cc-pwCVnZ (n=3–5) <sup>39–42</sup> sets, respectively. To incorporate outer-core-valence correlation effects involving 5s-,  $5p_{1/2}$ - $5p_{3/2}$ -, and, especially,  $5d_{3/2}$ - $5d_{5/2}$ -shells we keep all 25 electrons of At and 19 electrons of Au outside of the RECPs' small cores active. We find it possible to ignore excitations to virtual spinors with energies above 60 hartree with impunity, thus avoiding unnecessarily time-consuming computations for large basis sets. We evaluate single-reference coupled-cluster binding energies against the sum of individual atomic (or ionic for AtO<sup>+</sup>) energies.

A noteworthy caveat of 2c-CCSD(T) calculations for  $At_2$  in DIRAC19 is generating the ground-state 2c-Hartree–Fock reference. Similar to  $I_2$ ,  $^{43}$  a plain Kramers-restricted 2c-Hartree–Fock calculation for  $At_2$  converges to a spurious excited state, leading, in turn, to an equally spurious 2c-CCSD(T) solution. Using the 2c-Hartree–Fock solution for  $At_2^{2-}$  at a large R(At-At) as the guess drives the self-consistent procedure for  $At_2$  to the actual ground state at internuclear separations around  $R_e$ . Other cases do not seem to suffer from this problem.

Due to substantial bond lengths in At compounds (about 3 Å in At<sub>2</sub>), their ground states may display some multi-reference character already near equilibrium. We perform multi-reference calculations within the relativistic Fock-space CCSD (FS CCSD) method as implemented in the EXP-T program.  $^{44,45}$  We choose closed-shell systems of  $At_2^{2-}$ ,  $HAt^{2-}$ ,  $AuAt^{2-}$ , and  $AtO^-$  to generate vacuum states and and solve the FS CCSD equations in the (0h,0p), (1h,0p), and (2h,0p) Fock-space sectors, the last one corresponding to neutral  $At_2$ , HAt, AuAt, and cationic  $AtO^+$ . The energy denominator shifting technique allows the FS CCSD procedure to converge in difficult cases. We keep the denominator shifts as small as possible to facilitate convergence without compromising separability. This FS CCSD scheme can describe the atomization of  $At_2$ , HAt, and AuAt correctly, allowing us to calculate bond strengths relative to the molecular energies at large (20 bohr) internuclear separations. We describe the issues with the  $AtO^+$  case in Sec. 3.4. We include spin-orbit interaction in all FS CCSD calculations.

## 2.3 Relativistic density functional theory calculations

We perform 2c-RDFT calculations using the collinear spin-orbit formalism <sup>46</sup> as implemented in the NWChem <sup>47</sup> program or the non-collinear <sup>48–50</sup> one available in the TURBOMOLE <sup>51</sup> suite. By design, the non-collinear approach is invariant with respect to the molecule's orientation, albeit at a higher computational cost than the collinear approximation. The magnitude of the spin-orbit interaction in Au or At is not enough to cause a significant dependence of the molecular energy on the quantization axis choice in the collinear approach and in well-behaved cases such as At<sub>2</sub> or HAt, both versions of 2c-RDFT yield identical results. We find AtAu simulations difficult to converge with the collinear method and rely on the TURBOMOLE non-collinear 2c-RDFT approach. Additionally, we use TURBOMOLE's 2c-RDFT for simulations with range-separated XCFs due to more efficient implementation and a wider range of available functionals.

Our reference RDFT calculations employ the PBE0<sup>15,16</sup> XCF approximation, which is (almost) free of empirically fitted parameters and has already demonstrated its reliability in heavy- and superheavy-element molecular modeling. <sup>23,24,52,53</sup> Based on the  $R_e$  for each molecule, we generate a set of internuclear separations to model the potential energy surface within a  $\pm 0.5$  Å range of  $R_e$  with a 0.1 Å step. We also utilize these grids for scanning the energy curves with coupled-cluster and RDFT methods. The molecular cation AtO<sup>+</sup> being a potentially more difficult case, we describe the details separately in Sec. 3.4.

We simulate the binding energy curves and compute the ground-state spectroscopic constants  $R_e$ ,  $D_e$ , and  $\omega_e$  using the following representatives of the main XCF families: generalized-gradient approximation (GGA) Becke-88–Perdew-86, <sup>54–56</sup> meta-GGAs TPSS <sup>57,58</sup> and SCAN, <sup>59</sup> global hybrids PBE0, <sup>15,16</sup> B3LYP, <sup>14</sup> TPSSh, <sup>60</sup> PW6B95, <sup>61</sup> and M06-2X, <sup>62</sup> and range-separated hybrids HSE06 <sup>17–19</sup> and  $\omega$ B97X-D. <sup>63</sup>

#### 2.4 Composite approach

We generally follow the Feller–Peterson–Dixon approach  $^{12,64}$  that treats electron-correlation effects calculated within a scalar-relativistic approximation and spin-orbit effects on molecular properties as additive.  $^{53,64-66}$  In particular, we are interested in approximating 2c-CCSD(T) binding-energy  $E_b$  profiles using 1c-CCSD(T) and the spin-orbit contribution to the binding energy from RDFT. The formal expression for the composite binding energy of a diatomic molecule AB is

$$E_b(AB) = E_{1c-CCSD(T)}(AB) - \left(E_{1c-CCSD(T)}(A) + E_{1c-CCSD(T)}(B)\right) + \Delta SO(E_b), \quad (1)$$

where

$$\Delta SO(E_b) = E_{2c-RDFT}(AB) - (E_{2c-RDFT}(A) + E_{2c-RDFT}(B)) - - [E_{1c-RDFT}(AB) - (E_{1c-RDFT}(A) + E_{1c-RDFT}(B))].$$
(2)

1c-RDFT calculations are performed without the spin-orbit RECP operator and are operationally equivalent to non-relativistic DFT.

## 3 Results and discussion

## 3.1 At<sub>2</sub>

We summarize the equilibrium properties of the At<sub>2</sub> molecule in Table 1. Additionally, we present the binding energy  $E_b$  profiles near equilibrium (Fig. 1), the spin-orbit RDFT contributions  $\Delta SO(E_b)$  to the binding energy as a function of the bond length (Fig. 2), and juxtapose 1c-CCSD(T), 2c-CCSD(T), and Feller-Peterson-Dixon-style composite profiles combining 1c-CCSD(T) with the RDFT  $\Delta SO(E_b)$  contribution (Fig. 3).

Our 1c-CCSD(T) scalar relativistic results for  $R_e$ ,  $D_e$ , and  $\omega_e$  closely reproduce those from the most recent all-electron calculations by Vasiliu, Peterson, and Dixon<sup>12</sup> and corroborate the "chemical accuracy" preservation within the chosen RECP model.<sup>27</sup>

Our relativistic 2c-CCSD(T) results agree with the CBS-extrapolated all-electron X2C CCSD(T) ones<sup>9</sup> and support the reliability of the RECP model<sup>27</sup> for treating spin-dependent relativistic effects on valence and outer-core electrons. Apart from the Hamiltonian model, small differences likely arise from the CBS extrapolation scheme and the correlations of 5s- and 5p-electrons included in this work. We observe, for example, that the TZ-to-QZ extrapolation lowers  $D_e$  by about 0.005 eV compared to the QZ-to-5Z scheme. In contrast, the inclusion of 5s- and 5p-electrons in the correlated calculation strengthens the bond by about 0.003 eV. The magnitude of all these effects is below the "chemical accuracy" threshold of 1 kcal/mol. All At<sub>2</sub> coupled-cluster data presented here correspond to the QZ-to-5Z CBS extrapolation scheme.

Several sources<sup>9,10,13</sup> claim an essentially single-reference character of the At<sub>2</sub> electronic ground state. Our relativistic FS CCSD calculations aim to test this hypothesis directly and revise the underlying logic in Refs. 9,10,13. The authors of Ref. 9 suggest that single-

reference CCSD and multi-reference FS CCSD results are in close agreement. This statement contradicts our findings. In particular, we find that single-reference 2c-CCSD predicts  $D_e = 0.551$  eV, almost 30% smaller than our FS CCSD value. In Ref. 10, the authors make a stronger argument in favor of the single-reference ground state based on the agreement between their MRCISD+Q results for  $R_e$ ,  $D_e$ , and  $\omega_e$  and the corresponding values in Ref. 67 obtained within the 4c-CCSD(T) formalism. However, the authors of Ref. 10 indicate that they use the same basis set as in Ref. 67, although the former (aug-cc-PVTZ), unlike the one in Ref. 67, has an extra set of diffuse functions. Also, Visscher et al. revisited 10 their earlier results 67 to account for substantial correlation effects involving 5d-electrons. Our multi-reference relativistic FS CCSD calculations yield  $D_e = 0.780$  eV, which is very close to the single-reference CCSD(T) result (0.777 eV). However, compared to CCSD(T), FS CCSD suggests a shorter  $R_e$ , larger  $\omega_e$ , and a somewhat different shape of the  $E_b$  profile (see Table 1 and Fig. 1).

We have analyzed the determinantal structure of the  $0_g^+$  ground-state FS CCSD solution in the TZ-quality basis near equilibrium and compared it with similar results for I<sub>2</sub>. In Fig. 4, we give the coefficients above the  $10^{-4}$  threshold of the determinants contributing to the (2h, 0p)-sector normalized model vector, except for the dominant one. In the range from  $R_e - 0.1\text{Å}$  to  $R_e + 0.2\text{Å}$ , such contributions to the At<sub>2</sub>  $0_g^+$  ground state are markedly larger than in I<sub>2</sub>. Thus, while viewing the At<sub>2</sub>'s ground state as mostly single-reference is qualitatively appropriate, a quantitative description of the At–At bond may not be sufficient at the relativistic single-reference CCSD(T) or multi-reference FS CCSD level alone. A definitive analysis of the At<sub>2</sub>'s ground state would require relativistic CCSDT and FS CCSDT calculations, which are not feasible yet.

All examined XCF approximations reasonably reproduce the reference  $R_e$  and  $\omega_e$  coupledcluster data. However, the  $D_e$  estimates vary wildly. Pure functionals (B88P86, TPSS, and SCAN) and heavily parameterized hybrids (M06-2X and  $\omega$ B97XD) give the least satisfactory results. The only XCFs that yield "chemically accurate" results are HSE06, PW6B95, and, marginally, PBE0. Refs. 10,13,68 recognize B3LYP as a safe choice of an XCF approximation for the 2c-RDFT description of the  $At_2$  ground state. Our calculations indicate that it underestimates  $D_e$  by about 2 kcal/mol. In contrast to observations in Ref. 13, TPSSh is far from a safe choice as it overestimates  $D_e$  by almost 0.2 eV, or 5 kcal/mol.

Fig. 2 demonstrates a strong dependence of  $\Delta SO(E_b)$  on the internuclear distance. For the five representative XCFs, the  $\Delta SO(E_b)$  curves are virtually parallel within a 0.1 eV energy window, which is much narrower than the range of 2c-RDFT data range for the  $E_b$  profiles. Relative  $\Delta SO(E_b)$  insensitivity to the XCF choice makes RDFT a reliable tool for extracting the energy component responsible for the spin-orbit effects and their interference with electronic correlations. For the composite scheme shown in Fig. 3, we choose  $\Delta SO(E_b)$  evaluated with HSE06 since this XCF approximation yields the best 2c-RDFT approximation of 2c-CCSD(T) data. The addition of the  $\Delta SO(E_b)$  term to 1c-CCSD(T) results closely reproduces the 2c-CCSD(T) energy profile by causing not only a ca. 1 eV weakening of the At—At bond and its elongation by almost 0.3 Å. It is worth stressing that the  $\Delta SO(E_b)$  effect on  $D_e$  exceeds the  $D_e$  value itself, and the inclusion of spin-dependent relativistic effects in the valence shells is critical for a meaningful description of astatine's molecular properties.

Our composite results, while generally consistent with those in the most recent study by Vasiliu et al. 12 and only negligibly deviating from their  $\omega_e = 116.5 \text{ cm}^{-1}$  and  $R_e = 2.986 \text{ Å}$ , differ noticeably from  $D_e = 19.7 \text{ kcal/mol}$ , or 0.854 eV. 12 We cannot determine the exact cause of this discrepancy as it is unclear which internuclear separations were used to find  $D_e$  at the 1c-CCSD(T) level and the spin-orbit correction. However, it appears to us that the 1c-CCSD(T)  $D_e = 43.97 \text{ kcal/mol}$  corresponds to R = 2.8321 Å, which agrees with our results very well. At that same distance, the cited spin-orbit contribution to the binding energy of -26.98 kcal/mol, again agreeing with our value of -27.03 kcal/mol for that distance. However, the final binding energy in Ref. 12 uses the spin-orbit contribution of -24.278 kcal/mol. In our calculations, this value occurs around R = 2.940 Å, quite close to  $R_e = 2.986 \text{ Å}$  in Ref. 12. We surmise that  $D_e = 19.7 \text{ kcal/mol}$  results from combining the

1c-CCSD(T)  $D_e = 43.97$  kcal/mol value at R = 2.8321 Å with the spin-orbit contribution of -24.278 kcal/mol at R = 2.986 Å. Therefore, had the authors of Ref. 12 used the 1c-CCSD(T)  $D_e$  at R = 2.986 Å, it would have been closer to 41 kcal/mol, and the final result, to  $D_e = 17.7$  kcal/mol, which lines up with our findings.

Table 1: Ground-state properties of the  ${\rm At}_2$  molecule. This work's coupled-cluster results are extrapolated to the CBS limit.

Method	$R_e$ , Å	$D_e$ , eV	$\omega_e,  \mathrm{cm}^{-1}$
1c-CCSD(T)	2.817	1.909	162.9
$1\text{c-CCSD}(T)^{12}$	2.832	1.907	157.5
2c-CCSD(T)	2.957	0.777	119.4
$2c\text{-CCSD}(T)^9$	3.006	0.791	110.0
FS CCSD	2.858	0.780	132.1
B88P86	3.042	1.024	106.4
TPSS	3.017	1.074	110.7
SCAN	2.962	1.179	122.7
PBE0	2.972	0.828	120.8
B3LYP	3.043	0.705	108.7
TPSSh	2.995	0.973	115.5
PW6B95	2.989	0.751	116.6
M06-2X	2.935	0.370	129.1
$\omega \mathrm{B}97\mathrm{XD}$	2.984	0.504	122.5
HSE06	2.989	0.798	117.8
$1$ c-CCSD(T) + $\Delta$ SO	2.965	0.792	116.9
$1$ c-CCSD(T) + $\Delta$ SO <sup>12</sup>	2.986	0.854	116.5

#### 3.2 HAt

Table 2 contains our summary of the HAt ground-state properties, and Figs. 5, 6, and 7 visualize, respectively,  $E_b$  profiles,  $\Delta SO(E_b)$  as a function of R(H---At), and the juxtaposition of composite  $1\text{c-CCSD}(T) + \Delta SO(E_b)$  results and 2c-CCSD(T) data. Unlike  $At_2$ , HAt has a clear-cut single-reference ground state  $0^+$  near equilibrium: the FS CCSD solution's dominating determinant has a weight of 0.9999, and the next-largest contribution is only 0.0045. The multi-reference FS CCSD solution accounts for some dynamical-correlation effects, which single-reference 2c-CCSD(T) captures more systematically. Our 1c- and 2c-

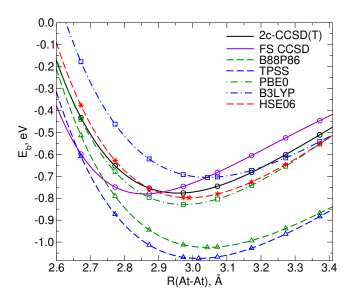


Figure 1: At<sub>2</sub> ground-state binding energy  $(E_b)$  profiles obtained using relativistic 2c-CCSD(T) and FS CCSD, and 2c-RDFT approaches.

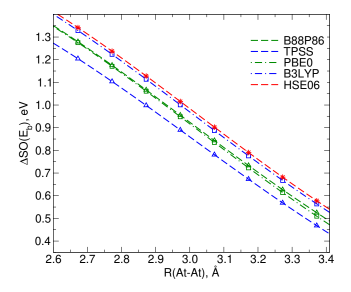


Figure 2: The ground-state spin-orbit contribution to the binding energy,  $\Delta SO(E_b)$ , of At<sub>2</sub> calculated using 1c- and 2c-RDFT with various XCF approximations.

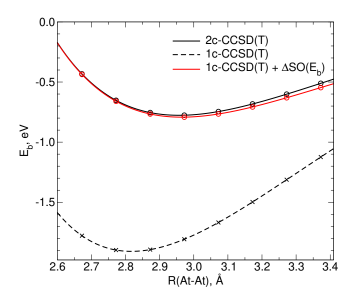


Figure 3: The composite approximation to the 2c-CCSD(T) ground-state  $E_b$  profile of At<sub>2</sub> obtained from 1c-CCSD(T) and HSE06  $\Delta$ SO(E<sub>b</sub>).

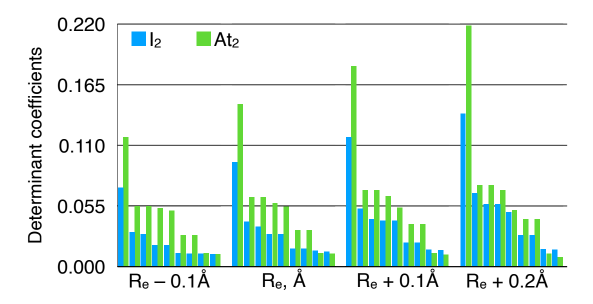


Figure 4: Coefficients of the determinants, excluding the dominant one, that contribute to the (2h, 0p)-sector FS CCSD solutions for the  $0_g^+$  ground states of  $I_2$  and  $At_2$ . The coefficients are given in the vicinity of the respective  $R_e$  values as calculated using the TZ-quality basis sets.

CCSD(T) results (QZ-to-5Z CBS extrapolation) agree perfectly with, respectively, the CBS-extrapolated all-electron scalar and composite CCSD(T) data in Ref. 12. The deviation from earlier findings<sup>11,69</sup> is most likely due to the basis set quality and core-valence correlation effects, as suggested in Ref. 11.

Out of all XCF approximations tried, only B3LYP gives a "chemically accurate"  $D_e$  value with respect to 2c-CCSD(T), although PW6B95 and PBE0 deviate by less than 1.4 kcal/mol. Notably, HSE06 already gives a 3.6 kcal/mol error in  $D_e$  despite its remarkable accuracy for At<sub>2</sub>. At the same time, the HSE06 estimate of the  $\Delta$ SO( $E_b$ ) contribution differs from the CCSD(T) one by less than 0.2 kcal/mol, making the 2c-CCSD(T) and 1c-CCSD(T)+ $\Delta$ SO( $E_b$ ) binding energy profiles practically indistinguishable, as illustrated in Fig. 7. Although the spin-orbit effect on  $D_e$ (HAt) is not as dramatic as in At<sub>2</sub>, it still amounts to almost 30% of the resulting binding energy.

Table 2: Ground-state properties of the HAt molecule. This work's coupled-cluster results are extrapolated to the CBS limit.

Method	$R_e$ , Å	$D_e$ , eV	$\omega,  \mathrm{cm}^{-1}$
1c-CCSD(T)	1.690	3.186	2169
$1\text{c-CCSD}(T)^{12}$	1.692	3.181	2167
$2c\text{-CCSD}(T)^{69}$	1.718	2.446	1994
2c-CCSD(T)	1.721	2.462	1990
$4c\text{-CCSD}(T)^{11}$	1.739	2.281	1966
FS CCSD	1.728	2.667	1892
B88P86	1.745	2.616	1911
TPSS	1.737	2.619	1955
SCAN	1.726	2.588	1988
PBE0	1.724	2.403	2013
B3LYP	1.734	2.426	1960
TPSSh	1.730	2.585	1990
PW6B95	1.723	2.405	2002
M06-2X	1.712	2.194	2046
$\omega \mathrm{B}97\mathrm{XD}$	1.725	2.335	2019
HSE06	1.727	2.305	2002
$1$ c-CCSD(T) + $\Delta$ SO	1.722	2.461	1979
$1c\text{-CCSD(T)} + \Delta SO^{12}$	1.722	2.485	1997

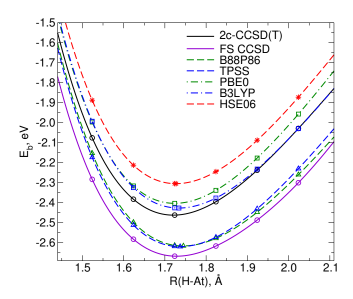


Figure 5: HAt ground-state binding energy  $(E_b)$  profiles obtained using relativistic 2c-CCSD(T) and FS CCSD, and 2c-RDFT approaches.

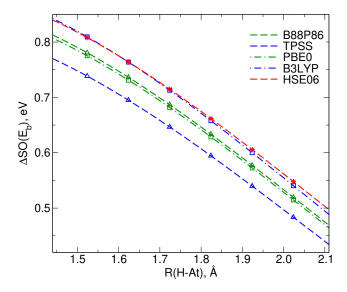


Figure 6: The ground-state spin-orbit contribution to the binding energy,  $\Delta SO(E_b)$ , of HAt calculated using 1c- and 2c-RDFT with various XCF approximations.

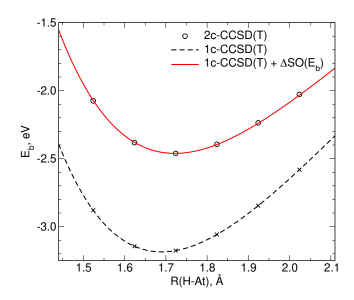


Figure 7: The composite approximation to the 2c-CCSD(T) ground-state  $E_b$  profile of HAt obtained from 1c-CCSD(T) and HSE06  $\Delta$ SO( $E_b$ ).

#### 3.3 AtAu

The AtAu results are presented in Table 3 and Figs. 8, 9, and 10. As in HAt, the ground state 0<sup>+</sup> in the relativistic FS CCSD approximation is predominantly single-reference with the leading determinant's coefficient of 0.981. However, a substantial number of determinants with coefficients' magnitudes above 10<sup>-3</sup> indicates a significant role of dynamical correlations beyond the 2c-CCSD level of theory. It suggests 2c-CCSD(T) as a preferred method to describe the ground state of AtAu.

The only 2c-RDFT calculation that approaches "chemical accuracy" in reproducing the 2c-CCSD(T)  $D_e$  value employs TPSSh. PBE0, B88P86, TPSS, and HSE06 yield only marginally acceptable results, while other XCF approximations behave unsatisfactorily. Additionally, we find M06-2X calculations too problematic to converge with either 2c-RDFT implementation and do not present any data for this XCF.

Considering At<sub>2</sub>, HAt, and AtAu, it is problematic to pinpoint a single XCF approximation that makes 2c-RDFT and 2c-CCSD(T) data universally consistent. A mostly non-empirical PBE0 approximation demonstrates an overall acceptable behavior in all three cases. However, the extrapolation of the agreement between 2c-RDFT/PBE0 and 2c-CCSD(T) re-

sults for diatomic molecules onto more complex structures requires caution. <sup>53,70</sup> The range-separated HSE06 approximation, as in the At<sub>2</sub> and HAt cases, yields the  $\Delta SO(E_b)$  estimate closest to CCSD(T), resulting in an overall reasonable agreement between 2c-CCSD(T) and 1c-CCSD(T)+ $\Delta SO(E_b)$  results.

Table 3: Ground-state properties of the AtAu molecule. This work's coupled-cluster results are extrapolated to the CBS limit.

Method	$R_e$ , Å	$D_e$ , eV	$\omega$ , cm <sup>-1</sup>
1c-CCSD(T)	2.544	2.572	186.3
2c-CCSD(T)	2.577	1.926	168.7
FS CCSD	2.563	2.013	179.3
B88P86	2.630	2.002	151.4
TPSS	2.617	2.007	155.6
SCAN	2.566	2.207	169.7
PBE0	2.609	1.853	159.1
B3LYP	2.650	1.663	147.9
TPSSh	2.611	1.970	158.0
PW6B95	2.617	1.794	156.1
$\omega \mathrm{B97XD}$	2.627	1.676	156.3
HSE06	2.619	1.840	156.7
$1$ c-CCSD(T) + $\Delta$ SO	2.580	1.983	167.3

## $3.4 \text{ AtO}^+$

While AtO<sup>+</sup> is one of the most studied a statine species, <sup>71-75</sup> some of its basic properties, such as  $D_e$  in the ground state, remain difficult to model. The most reliable approaches to incorporate dynamical correlation, spin-orbit interaction, and possible multi-reference character employ advanced perturbative techniques (NEVPT2) and spin-orbit configuration interaction. <sup>71,74</sup> Relativistic FS CCSD is also capable of capturing these effects near equilibrium progressing to the (2h, 0p) Fock-space sector <sup>71</sup> from the the closed-shell AtO<sup>-</sup> vacuum. Towards dissociation, however, the appropriate closed-shell vacuum state corresponds to AtO<sup>3-</sup>, and the FS CCSD model of AtO<sup>+</sup> would require access to the (4h, 0p) Fock-space sector. Due to the unavailability of such FS CCSD implementations, it is impossible to evaluate  $D_e$  at this level of theory.

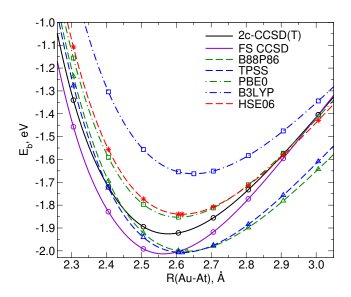


Figure 8: AtAu ground-state binding energy  $(E_b)$  profiles obtained using relativistic 2c-CCSD(T) and FS CCSD, and 2c-RDFT approaches.

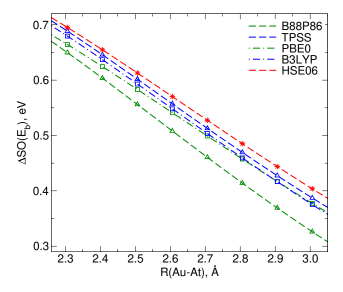


Figure 9: The ground-state spin-orbit contribution to the binding energy,  $\Delta SO(E_b)$ , of AtAu calculated using 1c- and 2c-RDFT with various XCF approximations.

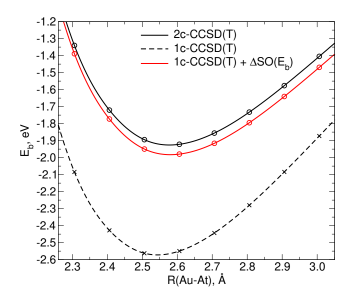


Figure 10: The composite approximation to the 2c-CCSD(T) ground-state  $E_b$  profile of AtAu obtained from 1c-CCSD(T) and HSE06  $\Delta$ SO( $E_b$ ).

A feasible alternative to multi-reference approaches is 2c- or 4c-CCSD(T) with a relativistic closed-shell Hartree–Fock solution as the reference determinant for the  $0^+$  ground state. 71 However, the application of Dirac-Coulomb 4c-CCSD(T) in Refs. 71 and 74 does not allow to estimate  $D_e$  due to problems in finding the ground-state CCSD(T) energy of an oxygen atom with the Dirac-Coulomb Hamiltonian. Our 2c-CCSD and 2c-CCSD(T) approaches within the RECP model avoid this issue. At dissociation, we treat the oxygen atom non-relativistically and compute its energy in the  ${}^{3}P$  ground state using the Coulomb Hamiltonian. These results are summarized in Table 4. A more fundamental problem with this CCSD(T) approach is the assumption of the ground state's single-reference character. According to Ref. 71, the  $T_1$  diagnostic value at R(At-O) = 2.0 Å is 0.03, which agrees with 0.037 in our calculations and justifies the applicability of a single-reference method. At the same time, the analysis of the (2h, 0p) FS CCSD wavevector's composition at this distance suggests the departure from a clear-cut single-reference case. Further elongation of R(At-O) to 2.1 Å leads to a steep increase of  $T_1$  to 0.1, consistent with the manifestation of the ground state's multi-reference character. In Fig. 11, we illustrate the change in the determinantal composition of the (2h, 0p) FS CCSD wavevector in the vicinity of  $R_e$ . Due to these indications of the single-reference CCSD(T) unreliability, we do not continue the 2c-CCSD(T) binding energy profile beyond 2.1 Å. Also, due to the rapid onset of multi-reference effects, we resort to a finer R(At-O) grid with a 0.01 Å increment and the TZ-to-QZ CBS extrapolation scheme to keep the calculations' volume reasonable.

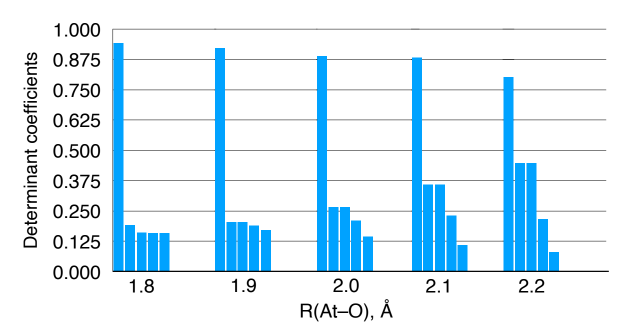


Figure 11: The first five largest coefficients of the determinants, including the dominant one, that contribute to the (2h, 0p)-sector FS CCSD solution for the  $0^+$  ground state of AtO<sup>+</sup>. These coefficients result from calculations using the TZ-quality basis sets.

Among the four species studied here, AtO<sup>+</sup> is the most difficult case for 2c-RDFT, especially in predicting  $D_e$ . Provided that 2c-CCSD(T) yields reliable ground-state spectroscopic constants, only B3LYP reproduces these results accurately, including a deviation in  $D_e$  below 1 kcal/mol. The only other marginally acceptable XCF approximations are PBE0 and PW6B95. In Fig. 12, we show the  $E_b$  profiles near equilibrium calculated using 2c-CCSD(T) (TZ-to-QZ CBS extrapolation) and 2c-RDFT with B3LYP and PBE0.

Spin-orbit interaction in  $AtO^+$  causes it to be a relativistic closed-shell system in its ground state instead of a spin-triplet in the scalar relativistic approximation.<sup>71</sup> Due to this spin-orbit-induced qualitative change in the ground state's electronic structure, the composite  $1c\text{-CCSD}(T)+\Delta SO(E_b)$  approach is hardly applicable to  $AtO^+$ .

Table 4: Ground-state properties of the  ${\rm AtO^+}$  molecular ion. This work's coupled-cluster results are extrapolated to the CBS limit.

Method	$R_e$ , Å	$D_e$ , eV	$\omega,  \mathrm{cm}^{-1}$
uc-SOCI/NEVPT2 <sup>74</sup>	1.893	2.30	799
$4c$ -CCSD $^{71}$	1.903	_	730
2c-CCSD	1.891	1.664	747.1
$4c\text{-CCSD}(T)^{71}$	1.930		676
2c-CCSD(T)	1.919	2.320	699.0
B88P86	1.934	3.166	683.0
TPSS	1.930	2.881	688.1
SCAN	1.913	2.619	715.4
PBE0	1.892	2.226	742.6
B3LYP	1.915	2.283	706.4
TPSSh	1.915	2.452	709.9
PW6B95	1.894	2.219	737.0
M06-2X	1.871	1.488	781.1
$\omega \mathrm{B}97\mathrm{XD}$	1.883	2.044	760.0
HSE06	1.897	2.105	733.8

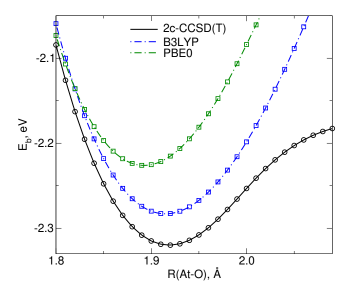


Figure 12: AtO<sup>+</sup> ground-state binding energy  $(E_b)$  profiles obtained using relativistic 2c-CCSD(T) and 2c-RDFT approaches.

## 4 Conclusions

Predictive modeling of a statine compounds' chemical properties poses challenges to high-level ab initio and DFT methods. Relativistic CCSD(T) approach in combination with accurate RECPs and be spoke basis sets allows us to provide revised reference data on the ground-state spectroscopic constants of  $At_2$ , HAt, AtAu, and  $AtO^+$  via the equal-footing treatment of large relativistic effects, especially spin-orbit interaction, and dynamical electronic correlations. Spin-orbit effects on chemical binding in a statine compounds are critically important: they are responsible for about 30% of binding energy in HAt and AtAu, exceed  $D_e$  in  $At_2$ , and fundamentally alter the electronic structure of  $AtO^+$ . Additionally, it is worth noting that the lack of suitable and generally adopted basis sets for spin-orbit calculations remains an issue for routinely modeling heavy elements' electronic structure and properties.

Relativistic Fock-space CCSD approach reveals some electronic-structure features beyond a typical dynamical-correlation case in  $At_2$  and, in particular,  $AtO^+$ . However, modeling the full length of binding-energy curves for  $At_2$  and  $AtO^+$  that would account for spin-orbit, dynamical-correlation, and multi-reference effects remains beyond reach for currently available relativistic coupled-cluster techniques.

Relativistic DFT is indispensable for modeling a statine systems more complex than diatomic molecules. It offers a computationally affordable procedure to simulate spin-free and spin-dependent relativistic and electronic-correlation effects. However, modern relativistic DFT extensions rely on XCF approximations derived or parameterized for non-relativistic systems, which often compromises the results' quality. The combination of non-relativistic XCF approximations with common basis set families compounds such simulations' errors even further. We disentangle these error sources using modified and extended UGBSs and reaching the complete basis set limit. For At<sub>2</sub>, HAt, AtAu, and AtO<sup>+</sup>, it appears problematic to indicate XCF approximations that would behave consistently well in reproducing ab initio coupled-cluster data, PBE0 being a marginal exception, likely owing to its mostly non-empirical character. Empirically fitted XCF tend to behave least satisfactorily. Spinorbit effects on molecular properties evaluated using relativistic DFT are less sensitive to the XCF choice, which allows for acceptable composite schemes based on scalar-relativistic coupled cluster and spin-orbit DFT approaches. These observations concerning relativistic DFT urge the development of XCF approximations and basis sets suitable for heavy-element molecular systems with large relativistic effects, especially spin-orbit interaction.

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# Supporting Information Available

All basis sets developed in this work are available online.

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