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The Heaviest Possible Ternary Trihalogen Species, IAtBr⁻, Evidenced in Aqueous Solution: An Experimental Performance Driven by Computations

Ning Guo^[a], Dumitru-Claudiu Sergentu^[a,b], David Teze^[a], Julie Champion^[a], Gilles Montavon^[a], Nicolas Galland^{*,[b]} and Rémi Maurice^{*,[a]}

Abstract: Evidencing new chemical species in solution is particularly challenging when one works at ultra-trace concentrations, as it is likely to happen with radioelements such as astatine (Z=85). In this work, quantum mechanical calculations were used to predict the tiny experimental domain in which it is possible to detect the presence of an exotic ternary trihalogen anion, IAtBr⁻, and thus to guide a series of experiments. By analyzing the outcomes of competition experiments, we show that IAtBr exists and can even predominate in aqueous solution. The equilibrium constant associated to the $At^+ + I^- + Br^- \rightleftharpoons IAtBr^-$ reaction is determined to be 10^{7.5±0.2}, which is in fair agreement with the density functional theory predicted one (10^{6.9}). This system not only constitutes the very first example of a ternary trihalogen species that involves the astatine element, but is also the first trihalogen species reported to predominate in solution. Also, we show that the oxidation number of At is zero in this species, as in the other molecules and anions that At⁺ can form with Cl⁻, Br⁻ and I ligands.

Evidencing and characterizing the structures and properties of new chemical compounds usually requires to perform a wide range of complementary experiments, among which spectroscopy ones play a major role. However, in some particular cases, one can only work with quantities of matter that prohibit the use of such golden experiments. This is typically the case with the astatine (At, Z=85) element, which is artificially produced with the help of cyclotron facilities^{[1],[2]}. Things can even become more complicated if only a tiny experimental domain admits the occurrence of the speculated chemical species in sufficient quantity. In this work, we demonstrate that quantum mechanical calculations can a priori guide the experimental study by predicting the experimental domain to be targeted. For this, the expected amounts of the chemical species to be evidenced should be sufficient to result in sizeable changes on the scrutinized "macroscopic" property (here the distribution coefficient of At in biphasic systems). Following this original approach, we show that it becomes possible to identify exotic species that may not have been evidenced otherwise.

It is of fundamental interest for chemists to characterize the chemical species that involve the At element, since some of its compounds may not have any analogue. Indeed, due to relativistic effects, the behavior of At may be very different to its nearest halogen, iodine. For instance, the Pourbaix diagram

[a]	N. Guo, DC. Sergentu, Dr. D. Teze, Dr. J. Champion, Dr. G.
	Montavon, Dr. R. Maurice
	SUBATECH, UMR CNRS 6457
	IN2P3/Mines Nantes/Université de Nantes
	4 rue Alfred Kastler, BP 20722, 44307 Nantes Cedex 3, France
	E-mail: remi.maurice@subatech.in2p3.fr
[b]	DC. Sergentu, Dr. N. Galland
	CEISAM, UMR CNRS 6230
	Université de Nantes
	2 rue de la Houssinière, BP 92208, 44302 Nantes Cedex 3, France
	E-mail: nicolas.galland@univ-nantes.fr
	5
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the document

(*E*-pH) of astatine involves cationic species such as At⁺ and AtO^{+[3],[4],[5]}, while the one of iodine only involves anionic or neutral species. Of course, in some cases, At and I behave in a similar way. Since the ternary trihalogen BrICI[¬] species has been characterized in solution by means of voltametric studies^[6], and its photodissociation behavior in the gas phase studied by means of pump-probe experiments^[7], we hypothesized that the IAtBr[¬] analogue would form in solution. This work aimed at evidencing the realness of this formation, notably though the determination of the equilibrium constant (*K*) associated to the following reaction:

$$At^{+} + I^{-} + Br^{-} \rightleftharpoons IAtBr^{-}$$
⁽¹⁾

However, the presence of the I⁻ and Br⁻ ions can lead to the formation of binary dihalogen molecules and binary trihalogen anions that would compete with the formation of the targeted ternary trihalogen species. These formations are associated with the following reactions:

$$At^+ + X^- \rightleftharpoons AtX$$
 (2)

$$At^{+} + 2X^{-} \rightleftharpoons AtX_{2}^{-} \tag{3}$$

where X is either Br or I, and β_{1,X^-} and β_{2,X^-} are the corresponding equilibrium constants associated with the involvement of one or two X⁻ units, respectively. Note that *K* notations are used for the formations of ternary complexes and β ones for the formations of binary 1:1 and 1:2 ones. Experimental values for the β_{1,X^-} and β_{2,X^-} constants are available in the literature, based on competition (X = Br)^[8] or electromigration (X = I)^[9] methods.

Table 1. Thermodynamic constants at zero ionic strength for the reactions of the At⁺ cation with Br⁻ and l⁻ ligands (from the literature).

x	Br⁻	I ^{- [a]}
$\log \beta_{1, X^-}$	$3.0 \pm 0.2^{[8]}$	6.0 ^[9]
$\log \beta_{2,X^-}$	4.1 ± 0.3 ^[8]	9.2 ^[9]

[a] Extrapolated from the values at 0.5 M ionic strength (6.2 and 9.4, respectively) to zero ionic strength through a truncated Davies equation^[10].

Quantum mechanical calculations can be used as a costeffective tool for assessing the stability of IAtBr⁻ with respect to the other dihalogen and trihalogen species of interest. To accurately predict equilibrium constants by means of relativistic density functional theory (DFT), one can make use of ligandexchange reactions in order to take profit of error cancellations^[3,5,8,11]. Since experimental Log β_{2,X^-} values are available for X = Br and X = I, we consider here the following reaction:

$$\frac{1}{2}\text{AtBr}_2^- + \frac{1}{2}\text{AtI}_2^- \rightleftharpoons \text{IAtBr}^- \tag{4}$$

The computation of the associated equilibrium constant, K_1 , readily leads to the Log K value of interest as follows:

$$\log K = \frac{1}{2} \log \beta_{2, Br^{-}} + \frac{1}{2} \log \beta_{2, I^{-}} + \log K_{1}$$
(5)

The B3LYP exchange-correlation functional^[12] was used, solvation free energies were estimated by means of a conductor-like polarizable continuum model (CPCM)[13], and other computational details that typically lead to accurate complexation constants in the field of astatine chemistry were chosen^[3,5,8,11] (see Supporting Information for more details and ref 11 for an extensive benchmark). A Log K1 value of 0.2 was computed, leading to a predicted Log K value of 6.9. Using the latter combined with the equilibrium constants reported in Table 1, we can elaborate a predictive speciation diagram of At in aqueous phase with the presence of both the I⁻ and Br⁻ anions (see Figure 1). It becomes clear from this prediction that a predominance domain does exist for the IAtBr species, but that it should be rather small. Indeed, according to the theoretical prediction, IAtBr could be a predominant species when the concentrations of I^- is within the ~10⁻⁴ to ~10⁻² M range meanwhile the Br one is within the ~0.1 to ~1 M one (see Figure 1). This severely restricts the experimental domain to be covered. Additional experimental constraints may further be considered.

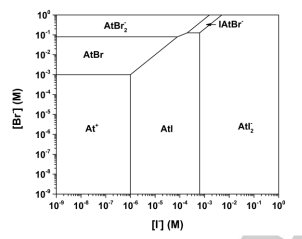


Figure 1. Predicted speciation diagram of At as a function of the initial concentrations of both the Br⁻ and I⁻ anions, based on the parameters displayed in Table 1 and the DFT predicted Log K value.

A competition method was used to experimentally investigate the speciation of At in the aqueous phase. In this work, we consider the distribution coefficient (D) between two liquid phases, one aqueous and the other organic (the solvent being the toluene)^[14]. Although a change in the D value arising from a change in experimental conditions must indicate a change in speciation, an apparent constant D value may hide some composition changes in each phase. Therefore, one must find conditions that lead to sizeable changes in D to evidence new species. Simulations of the distribution coefficient as a function of the initial I⁻ concentration at fixed Br⁻ concentrations were conducted (see Figure 2). Four Br⁻ concentrations were considered, the largest value corresponding to the limit of validity of the truncated Davies equation, and the smallest one being out of the predicted predominance domain of IAtBr⁻. We then select experimental conditions for which significant deviations appear between the model curves obtained by only considering the binary species (dash lines) and the ones obtained by also considering the ternary anion (solid lines). Therefore, we have retained the 0.1 and 0.05 M initial concentrations for the Br ligand, and we vary the I one between 5×10^{-5} and 2.5×10^{-4} M (*i.e.* we start near the beginning of the expected deviation and we end close to the maximum of it).

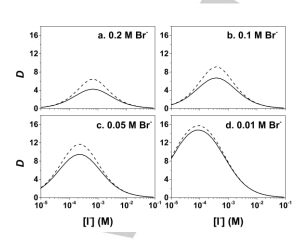


Figure 2. Simulation curves of the biphasic system: distribution coefficient (*D*) of astatine as a function of the I^- initial concentration at fixed Br⁻ concentrations (a.: 0.2 M, b.: 0.1 M, c.: 0.05 M, d.: 0.01 M), based on the parameters displayed in Table 1 and the DFT predicted Log *K* value. Dash lines correspond to the simulations that only deal with the binary 1:1 and 1:2 species, while solid lines are used for the simulations that also account for the formation of the IAtBr⁻ species in the aqueous phase.

Prior to study ternary media that a priori include At⁺, I⁻ and Br⁻ ions, we start by revisiting the binary At⁺/Br⁻ and At⁺/I⁻ systems to get accurate and consistent (with the same experimental procedure) values of the β_{1, X^-} and β_{2, X^-} constants. The aqueous phase consists in 0.1 M of HCIO₄ (pH≈1, E≈0.6V), in which At⁺ is the astatine dominant species, in the absence of complexing agent, according to its Poubaix diagram^{[3],[4],[5]}. The modelling of the At⁺/Br⁻ system with adjustable $\log \beta_{1, Br^-}$ and $\log \beta_{2, Br^-}$ parameters (see Supporting Information) leads to the values displayed in Table 2. These values are in a good agreement with the previously published data^[8]. Table 2 also gives the thermodynamic constants associated with the At+/I- binary system, which were derived from the modelling of the At distribution coefficient as a function of I⁻ concentration (see Figure 3). The experimental data shows no signature of any speciation change when the initial I⁻ concentration is below 10⁻⁸ M. The increase of D value between 10^{-8} and 10^{-5} M implies the formation of a new species that is extracted in toluene, and a plateau is reached between 10^{-5} and 10^{-3} M. The D value then decreases, which suggests the formation of another species that is not extracted in toluene. The experimental data are successfully modelled by considering the formation of 1:1 and 1:2 complexes, and the determined thermodynamic constants are found to be in good agreement with the electromigration data.[9]

X ⁻	Br⁻	ſ
$\log \beta_{1,X^-}$	2.7 ± 0.2	6.1± 0.2
$\log \beta_{2, X^-}$	3.8 ± 0.3	8.8± 0.2

Table 2. Thermodynamic constants at zero ionic strength for the reactions of the At⁺ cation with Br⁻ and l⁻ ligands (this work).

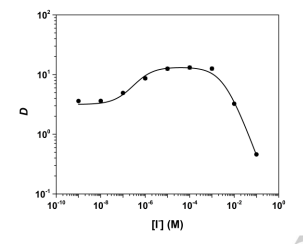


Figure 3. At^{*}/l⁻ binary system: distribution coefficient as a function of the initial l⁻ concentration. The model curve obtained with the Log β_{1, l^-} and Log β_{2, l^-} values reported in Table 2 is displayed in solid line.

We now continue by analysing the data obtained for the ternary At⁺/I⁻/Br⁻ system. As shown in Figure 4, the D value increases with the I⁻ concentration for both the 0.05 and 0.1 M initial Br⁻ concentrations. First, we tried to model the data by only considering binary 1:1 and 1:2 species with the parameter values that are given in Table 2. As can be seen, the model curves (a for 0.05 M of Br and c for 0.1 M of Br) cannot explain at all the experimental data. Therefore, the formation of at least another species must be considered. Following the previous prediction, we then further consider the formation of the IAtBr species via reaction 1. As can be seen in Figure 4 (curve b for 0.05 M of Br and curve d for 0.1 M of Br), a Log K value of 7.5 leads to a satisfactory modelling of the experimental data. Furthermore, the fairly good agreement between the experimental value of Log K (7.5 ± 0.2) and the theoretical one (6.9) prompt us to confirm the existence of the IAtBr⁻ species. The introduction of the IAtBr species in the model leads to an effective decrease in the estimated Atl population (see Figure S2). Since this species can be extracted in toluene, contrary to the anionic IAtBr one, the formation of the IAtBr species translates in Figure 4 in a decrease in the modelled distribution coefficient.



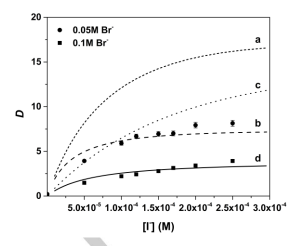


Figure 4. At⁺/[⁻/Br⁻ ternary system: distribution coefficient as a function of the initial I⁻ concentration for two initial Br⁻ concentrations (0.05 M of Br⁻, circles, and 0.1 M of Br⁻, squares). The model curves obtained with only the values presented in Table 2 are displayed in short dash line (a. 0.05 M of Br⁻) and dot line (c. 0.1 M of Br⁻), and the ones obtained by further considering a Log *K* value of 7.5 are displayed in dash line (b. 0.05 M of Br⁻) and solid line (d. 0.1 M of Br⁻).

Having experimentally determined the Log K value, we can now build the At speciation diagram as a function of both the initial concentrations of I and Br (see Figure 5). The IAtBr species can actually predominate for concentrations of I around $10^{-5}-10^{-2}$ M and Br⁻ around 0.1-1 M. This species, being the heaviest possible ternary trihalogen one, deserves some more attention from a chemical point of view. Of course, no structural information can be experimentally obtained on this species. Therefore, we discuss its relativistic DFT structure. All the structures of the trihalogen species considered in this work, i.e. AtBr₂, Atl₂, and IAtBr, are linear, the At atom being located in between the other two halogen ones. Therefore, the IAtBrspecies is analogous to the ternary trihalogen BrICI⁻ species, for which the heavier halogen, I, is located in between the two lighter ones^[7]. The At-X distances are 2.91 Å (AtBr2-), 3.15 Å (Atl₂⁻), and 2.92 and 3.14 Å (IAtBr⁻) at the considered relativistic DFT level. One should stress that spin-orbit coupling must be taken into account for obtaining accurate geometries, At-X distances being affected by up to ~0.1 Å in these systems. To determine the oxidation numbers (ONs) of At in these species, and also in the AtBr and AtI ones, as well as the ON of I in BrICI, we have estimated the number of valence p electrons of the heaviest halogen with complete active space configuration interaction (CASCI) calculations for which previously obtained complete active space self-consistent field (CASSCF) active orbitals were localized (see Supporting Information for more details). In all these species, more than 4.8 valence p electrons are obtained for the heaviest halogen, meaning that the actual wave function is much closer to the neutral limit for this halogen (5 p electrons) than to the first ionized one (4 p electrons). In other words, the ON of At in AtBr, AtI, AtBr2-, AtI2- and IAtBr- is zero (note that the ON of I in BrICI⁻ is actually also zero), contrary to what was previously assumed for the Atl and Atl₂

cases^[9]. Since these species were formed from the At⁺ moiety, one can consider that the formations of the molecules and ions considered in this work lead to a change in the At ON. This reveals that the wave functions in these species are not dominated by ionic configurations (of the forms At⁺ X⁻ and X⁻ At⁺ Y⁻, respectively), but rather by covalent ones (of the forms At -X and X-At Y⁻ and X⁻ At-Y, respectively). Note that, as expected, test calculations showed that the inclusion of spin-orbit coupling hardly affects the number of At *p* electrons, and thus, cannot change our conclusions on the At ON.

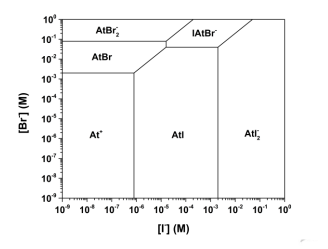


Figure 5. Experimental speciation diagram of At as a function of the initial concentrations of both the Br^- and I^- anions, based on the parameters displayed in Table 2 and the experimentally determined Log *K* value of 7.5.

In the present work, we have shown that adequate computational predictions can lead to the experimental discovery of an exotic chemical species, namely IAtBr⁻. Of course, it is of high interest for chemists to know if other ternary trihalogen species that involve the At element can exist or even predominate in solution. Using the *in silico* methodology introduced in this work, we have assessed the possibility of forming the IAtCl⁻ anion in the ternary At⁺/l⁻/Cl⁻ case (see Supporting Information). Based on this prediction, we conclude that IAtCl⁻ can be formed in solution. However, considering the extremely small-predicted domain of predominance, the experimental evidence of its existence may not be reachable in

practice. Therefore, it is not yet confirmed if any other ternary trihalogen species involving At can predominate in solution.

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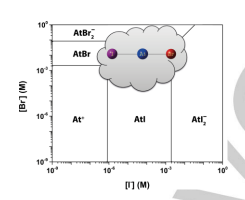
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Entry for the Table of Contents

COMMUNICATION

Relativistic density functional theory is used by Guo *et al.* in their Communication on page xxx ft. to predict the tiny experimental domain in which it may be possible to detect the heaviest possible trihalogen species, IAtBr⁻. Competition experiments are then performed at ultra-trace concentrations, for (i) confirming the theoretical prediction and (ii) determining experimentally the associated reaction equilibrium constant.



Ning Guo, Dumitru-Claudiu Sergentu, Dr. David Teze, Dr. Julie Champion, Dr. Gilles Montavon, Dr. Nicolas Galland* and Dr. Rémi Maurice*

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The heaviest possible ternary trihalogen species, IAtBr⁻, evidenced in aqueous solution: An experimental performance driven by computations