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Advances in the chemistry of astatine and implications for the development of radiopharmaceuticals

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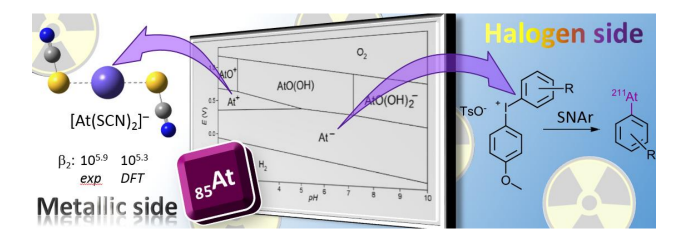
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Conspectus: Astatine (At) is the rarest on Earth of all naturally-occurring elements, situated below iodine in the periodic table. While only short-lived isotopes ($t_{1/2} \le 8.1$ h) are known, ²¹¹At is the object of growing attention due to its emission of high-energy alpha particles. Such radiation is highly efficient to eradicate disseminated tumors, provided that the radionuclide is attached to a cancer-targeting molecule. The interest in applications of ²¹¹At in nuclear medicine translates into the increasing number of cyclotrons able to produce it. Yet, many challenges related to the minute amounts of available astatine are to be overcome in order to characterize its physical and chemical properties. This point is of paramount importance to develop synthetic strategies and solve the labeling instability in current approaches that limits the use of ²¹¹Atlabeled radiopharmaceuticals. Despite its discovery in the 1940s, only the last decade has seen a significant rise in the understanding of astatine's basic chemical and radiochemical properties, thanks to the development of new analytical and computational tools.

In this Account, we give a concise summary of recent advances in the determination of physicochemical properties of astatine, putting in perspective the duality of this element which exhibits both the characteristics of a halogen and of a metal. Striking features were evidenced in the recent determination of its Pourbaix diagram such as the identification of stable cationic species, At^+ and AtO^+ , contrasting with other halogens. Like metals, these species were shown to form complexes with anionic ligands and to exhibit a particular affinity for organic species bearing soft donor atoms. On the other hand, astatine shares many characteristics with other halogen elements. For instance, the At^- species exists in water, but with the least range of $E_{H^-}pH$ stability in the halogen series. Astatine can form molecular interactions through halogen bonding, and it was only recently identified as the strongest halogen-bond donor. This ability is nonetheless affected by relativistic effects, which translate to other peculiarities for this heavy element. For instance, the spin-orbit coupling boosts astatine's propensity to form charge-shift bonds, catching up with the behavior of the lightest halogens (fluorine, chlorine).

All these new data have an impact on the development of radiolabeling strategies to turn ²¹¹At into radiopharmaceuticals. Inspired by the chemistry of iodine, the chemical approaches have sparsely evolved over the past decades and have long been limited to electrophilic halodemetallation reactions to form astatoaryl compounds. Conversely, recent developments have favored the use of the more stable At⁻ species including the aromatic nucleophilic substitution with diaryliodonium salts, or the copper-catalyzed halodeboronation of arylboron precursors. However, it is clear that new bonding modalities are necessary to improve the *in vivo* stability of ²¹¹At-labeled aryl compounds. The tools and data gathered over the past decade will contribute to instigate original strategies for overcoming the challenges offered by this enigmatic element. Alternatives to the C-At bond such as the B-At and the metal-At bonds are typical examples of exciting new axes of research.

Key references

• Guo, N.; Maurice, R.; Teze, D.; Graton, J.; Champion, J.; Montavon, G.; Galland, N. Experimental and Computational Evidence of Halogen Bonds Involving Astatine. *Nat. Chem.* 2018, *10*, 428–434.¹

This article reports the very first experimental characterization of halogen bonds mediated by astatine, quantum chemistry calculations making up for the impossibility of using spectroscopic tools.

Leimbach, D.; Karls, J.; Guo, Y.; Ahmed, R.; Ballof, J.; Bengtsson, L.; Boix Pamies, F.; Borschevsky, A.; Chrysalidis, K.; Eliav, E.; Fedorov, D.; Fedosseev, V.; Forstner, O.; Galland, N.; Garcia Ruiz, R. F.; Granados, C.; Heinke, R.; Johnston, K.; Koszorus, A.; Köster, U.; Kristiansson, M. K.; Liu, Y.; Marsh, B.; Molkanov, P.; Pašteka, L. F.; Ramos, J. P.; Renault, E.; Reponen, M.; Ringvall-Moberg, A.; Rossel, R. E.; Studer, D.; Vernon, A.; Warbinek, J.; Welander, J.; Wendt, K.; Wilkins, S.; Hanstorp, D.; Rothe, S. The Electron Affinity of Astatine. *Nat. Commun.* 2020, *11*, 3824.²

Eighty years after the discovery of astatine, its electronic affinity is finally measured thanks to a state-of-the-art technique of laser photodetachment spectroscopy carried out at CERN.

Sergentu, D.-C.; Teze, D.; Sabatié-Gogova, A.; Alliot, C.; Guo, N.; Bassal, F.; Silva, I.
 D.; Deniaud, D.; Maurice, R.; Champion, J.; Galland, N.; Montavon, G. Advances on the

Determination of the Astatine Pourbaix Diagram: Predomination of AtO(OH)²⁻ over At⁻ in Basic Conditions. *Chem. Eur. J.* 2016, *22*, 2964–2971.³

A significant step towards the completion of the Pourbaix diagram is presented in this study, judiciously combining macroscopic data from radiochemical experiments and information at the molecular scale provided by DFT calculations.

Berdal, M.; Gouard, S.; Eychenne, R.; Marionneau-Lambot, S.; Croyal, M.; Faivre-Chauvet, A.; Chérel, M.; Gaschet, J.; Gestin, J.-F.; Guérard, F. Investigation on the Reactivity of Nucleophilic Radiohalogens with Arylboronic Acids in Water: Access to an Efficient Single-Step Method for the Radioiodination and Astatination of Antibodies. *Chem. Sci.* 2021, *12*, 1258-1468.⁴

The robust ²¹¹At-labeling procedure reported herein facilitates the preparation of radiopharmaceuticals and the transfer to the clinic as ready-to-use kits. Being also applicable to radioiodination, it opens perspectives for radiotheranostic applications.

Introduction

Discovered in 1940 after irradiation experiments of bismuth by accelerated α particles, astatine (Z=85) is the rarest naturally occurring element.⁵ All of its 32 known isotopes exhibit short half-lives (125 ns to 8.1 h), justifying the Greek origin of its name, *astatos*, meaning unstable. The ²¹¹At isotope ($t_{1/2}=7.2$ h) decays to stable ²⁰⁷Pb through a double-branched pathway, producing high-energy α particles (5.9 and 7.5 MeV) of great interest for targeted radionuclide therapy of cancers.⁶ α particles exhibit short path length (<100 μ m), resulting in a higher cytotoxicity for nearby malignant cells and reduced irradiation of surrounding healthy tissues in

comparison to β^- particles, which is particularly appropriate for the treatment of small clusters or isolated cancer cells (**Figure 1**).^{7,8}

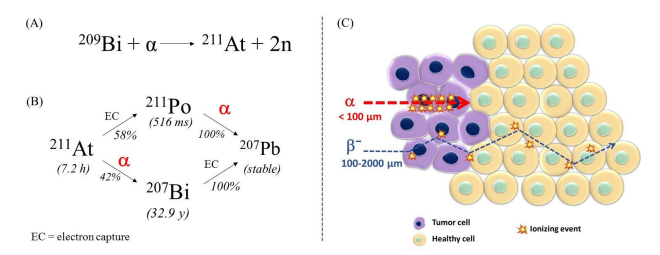


Figure 1. A) Nuclear reaction to produce 211 At from bismuth. B) 211 At decay scheme. C) Simplified view of α vs. β particle track and related ionization density.

Among the eight main radionuclides considered for targeted alpha therapy, ²¹¹At, when coupled to a cancer-targeting agent, appears very promising as illustrated by the first clinical trial reported in 2008 for glioblastoma treatment⁹ and more recently in studies demonstrating long-term low toxicity of ovarian cancer treatment.¹⁰ To expand the number of potential carrier compounds applicable to ²¹¹At, the need of innovative strategies remains to be solved even 80 years after astatine discovery, in particular to improve reactions robustness and *in vivo* labeling stability. The design of optimal astatine-based radiopharmaceuticals has long been hampered by the elusive nature of this element.¹¹ For instance, the measurements of atomic ionization energy and electron affinity, which are fundamental quantities for understanding the subtle mechanisms of bond formation, were only reported in 2013¹² and 2020², respectively. A number of astatine's properties remain thus far inaccessible because the radioelement is available from artificial production at the nanogram scale at best, making usual spectroscopic tools inapplicable to

physical and chemical characterization. Although a number of astatine's characteristics are clearly similar to that of halogens, this element also behaves as a metal in some conditions, which led Visser to qualify it as a "chameleon", ¹³ a term that perfectly reflects its multifaceted character.

With the completion in the late 2000's of the Arronax cyclotron facility, ¹⁴ one of the few accelerators in the world exhibiting adequate characteristic to produce ²¹¹At, ¹⁵ it became obvious, once this radionuclide had become available routinely to us, that new methodologies and new tools were needed to quantitatively evaluate astatine chemistry. It is in this context that we initiated research programs which intended to better understand this element's basic chemical properties, and eventually benefit the radiolabeling chemistry. A decade later, these efforts are beginning to bear fruits as the outline of this element's chemistry has become clearer and radiolabeling reactions of higher efficiency and robustness have been identified, as illustrated in this Account.

I. Pourbaix diagram of astatine - evidences of a metallic behavior

In the early 2000s, astatine was suspected to exist under three different oxidation states in water. ^{13,16–19} However, the exact species involved remained unclear and sometimes inconsistent depending on authors, e.g. At⁺ (**Figure 2**). They were mostly extrapolated from iodine as is still done now. ²⁰ The existence of At(0) was notably unproven, which motivated our initial investigations.

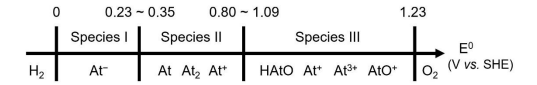


Figure 2. Proposed a statine species from the literature, at pH~0 according to redox conditions. Data from ref. 13, 16-19.

1.1 Construction of the Pourbaix diagram

In order to characterize At species at ultra-trace concentrations, which are unidentifiable by conventional spectroscopic tools, a dual experimental/theoretical approach was followed. In addition to electromobility and chromatographic techniques coupled with radio-detection, we adapted the Schubert's method²¹ to the study of species at trace level.²² The principle is based on the partitioning of total astatine between two immiscible phases, characterized by a distribution ratio (*D*). Variation in *D* when experimental parameters are modified (pH, redox potential, ligand concentration, *etc.*) is indicative of a change in astatine speciation. The analysis of these data, on the basis of a classical equilibrium model, allows one to trace the formed species and to determine the associated thermodynamic constants. The results are then confronted with those of quantum mechanical calculations on speculated species. Since astatine is a heavy element subject to relativistic effects, specific computational approaches were implemented for its study. In particular, the spin-orbit coupling (SOC) can strongly influence astatine chemistry.^{23–28} With the development of two-component relativistic methods in the 2000s, notably in the framework of the density functional theory (DFT), it became possible to determine the structure and

thermodynamic properties of medium-size compounds. As a general rule, the existence of At species is validated if experimental and theoretical data converge towards the same answer.

Our research was based on the assumption that the At^- species exists, similarly to halides observed from fluorine to iodine. The first step was to confirm its existence. In non-alkaline and reducing medium, a monocharged anionic species was evidenced by ion exchange chromatography.²⁹ Electromobility measurements strengthened our hypothesis, a negative species being confirmed.³⁰ Molecular dynamics simulations on At^- corroborated the obtained mobility value.³¹ Taken together, these data fully ascertain the existence of At^- . The next step was to study the conversion of At^- when E_H increases. One of the most-controversial species in the literature was species II, found in acidic media in mild redox conditions (**Figure 2**).

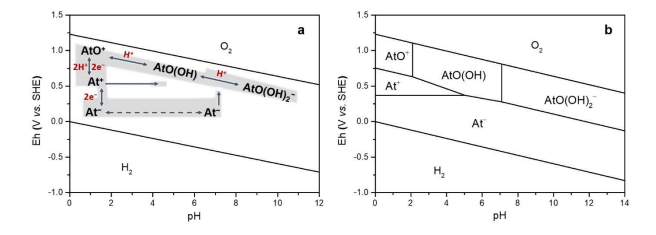


Figure 3. Pourbaix diagram in the range of water stability. a) Shadow zones correspond to the explored E_{H} -pH ranges; b) Proposed diagram for At.

Astatine was suggested to have an oxidation number either of 0 or \pm 1. In acidic conditions, our experimental work revealed that the oxidation of At⁻ to form species II involved an exchange of two electrons, independently of pH (**Figure 3**). ³² Hence, the existence of an At(0) species would

be possible only as At_2 . Such a diatomic species is highly unlikely given the extremely low astatine concentration in studied solutions. Moreover, our DFT calculations predict a standard potential of ~ 0.17 V for the At_2/At^- couple, which is inconsistent with the measured value of 0.36 ± 0.01 V.³² The exchange of two electrons would therefore imply the formation of a At(I) species, namely At^+ . The existence of this stable monatomic cation in aqueous solution contrasts with lighter halogens. This observation is clear evidence of the metallic behavior of astatine.

In acidic conditions and in the presence of a strong oxidizer, there exists another "metallic" species characterized by a positive charge in electromobility experiments³³ and in ion exchange chromatography. The most frequently suggested chemical forms for species III are At^+ , At^{3+} and AtO^+ . Our experiments have shown that the oxidation of At^+ involves an exchange of two electrons and two protons (the variation of the apparent potential is pH-dependent). Therefore, species III cannot be At^+ or At^{3+} and we concluded that AtO^+ is the species in question. At this stage, three oxidation states corresponding to At^- , At^+ and AtO^+ were elucidated at pH between 1 and 2. Under standard conditions (298.15K, 1M), the redox potentials are 0.36 ± 0.01 V and 0.74 ± 0.01 V vs. SHE for the At^+/At^- and AtO^+/At^+ couples, respectively.

In oxidizing conditions and when the pH increases, the conversion of AtO⁺ to a neutral species was observed by ion exchange chromatography.³⁴ The change in speciation was also observed using Schubert's method. Quantitative interpretation of the data showed the exchange of one proton and thus the formation of the AtO(OH) hydrolyzed species. This conclusion was supported by the thermodynamic constants predicted by DFT calculations. A similar strategy was used to highlight the formation with increasing pH of a second hydrolyzed species, AtO(OH)₂⁻, in place of a deprotonation of AtO(OH).³ Conversely, experimental studies have not

been able to reveal hydrolyzed forms of At⁺, testifying to a narrow domain of existence of At⁺ as shown in **Figure 3**.

1.2 Metallic properties of At⁺ and AtO⁺

The interactions between At^+ or AtO^+ , jointly denoted $At(x)^+$, and common inorganic ligands (*e.g.*, $C\Gamma$, Br^- , Γ and SCN^-) were investigated using our experiment/theory approach. Binary species with 1:1 and 1:2 stoichiometries were evidenced, highlighting a metallic character for $At(x)^+$. In particular, relativistic DFT calculations show hypervalent astatine atoms for the 1:2 structures. In addition, an experimental study guided by theoretical calculations has evidenced a ternary trihalogen species: based on the equilibrium constant computed for the formation of the IAtBr $^-$ species ($10^{6.9}$), a narrow domain of existence was anticipated in aqueous solutions. Considering an astatine aqueous solution with finely tuned E_H , pH and Br $^-$ concentration, the experimental results confirmed the formation of a new species when increasing Γ concentration. A thermodynamic model that considers the formation of IAtBr $^-$, with an equilibrium constant of $10^{7.5 \pm 0.2}$, can then describe the experimental data. IAtBr $^-$ is not only the heaviest possible triatomic interhalogen, but it is also the first one declared with a stability domain in aqueous solution.

It also emerges from theoretical calculations that the solvent has an important influence on the chemistry of a the molecular level. Regarding the reaction between AtO⁺ and SCN⁻, and SCN⁻, the solvation completely change the active site within the molecular cation, from the oxygen atom, which controls the reaction in gas phase, to the astatine atom, when solvated (**Figure 4**).

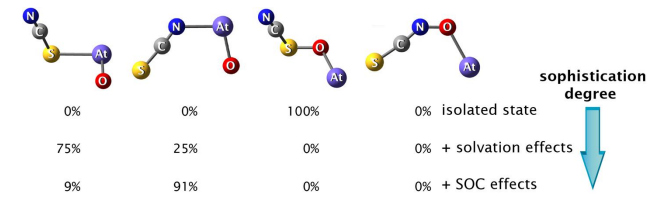


Figure 4. Computed proportions of adducts between AtO⁺ and SCN⁻. Data from ref. 23.

Regarding the non-bonded AtO^+ cation, the solvation is also responsible for a rare phenomenon: a ground-state reversal. Indeed, relativistic quantum mechanical calculations have shown that the closed-shell component of the $^1\Delta$ excited state in the free cation gradually becomes the essentially-spin-singlet electronic ground state when AtO^+ interacts with an increasing number of water molecules. $^{36-39}$ It is worth noting that the proximity between the ground state and the first excited electronic states, responsible for this reversal, is also a common property of metals. In addition, according to recent electronic structure calculations, 40 astatine in condensed state becomes monatomic and simultaneously metallic, conversely to lighter halogens.

II. Astatine also behaves as a halogen

The position of astatine in the periodic table of elements (PTE), as a member of the nonmetallic elements that constitute the group 17 (including fluorine, chlorine, bromine and iodine), suggests that it belongs to the halogen family. This assumption is supported by experimental observations, including some of our recent results among the most significant.

II.1 Astatide and related properties

A characteristic of halogens is their ability to form a stable halide ion in aqueous solution. The construction of the Pourbaix diagram confirmed the existence of the astatide anion, $At^{-29,30}$ In line with the trend observed within the halogen series, the heavier the halogenide, the more restricted the $E_{H^-}pH$ stability range. The specific stability of halide ions derives from the electronic configuration of all halogens. They carry seven valence electrons in their outermost electron shell and tend towards holding one more electron, as in the rare gas next to it in the PTE. This is quantitatively reflected by the electron affinity (EA). The halogens exhibit the largest EAs in the PTE but astatine's EA was still unknown 80 years after its discovery. Thanks to the CERN nuclear-physics facility and our contribution to quantum mechanical calculations to guide the experimental setup, this most fundamental atomic property has recently been measured. Although the value, 2.41578(7) eV, is smaller than those of lighter halogens, it remains larger than measured values for all other elements.

If a high EA reflects the strong tendency to form anions in aqueous solution, its value also impacts several properties of atoms relevant to various chemical concepts. Electronegativity is the most prominent example. Its value for astatine, also relying on the recently determined ionization energy (IE),¹² is presented in **Table 1** together with the softness, hardness, and electrophilicity indices. Taken together, these new data are highly relevant in the context of radiolabeling chemistry that has mainly involved nucleophilic and electrophilic astatine species to form covalent bonds for the preparation of radiopharmaceuticals, but also when studying *in vivo* mechanisms that release At⁻ from the carrier biomolecules.

Table 1. Astatine's atomic properties derived from the high-precision measurements of EA and IE, compared to those of iodine. Data from ref. 2, 12.

Property	Definition	At	Ι
electronegativity	$\chi = \frac{IE + EA}{2}$	5.87 eV	6.76 eV
hardness	$\eta = \frac{IE - EA}{2}$	3.45 eV	3.70 eV
softness	$S = \frac{1}{2\eta}$	0.145 eV^{-1}	0.135 eV ⁻¹
electrophilicity	$\omega = \frac{\chi^2}{2\eta}$	4.99 eV	6.17 eV

II.2 Astatine as the strongest halogen-bond donor

A noticeable property specific to halogens is their ability to form so-called halogen-bond (XBs). In short, XBs are attractive and highly directional non-covalent interactions formed between an electrophilic region of the halogen atom X (called σ-hole) in a R–X compound, and a nucleophilic site of a Lewis base. Halogen bonding is recognized as playing an important role in many fields related to molecular recognition, including medicinal, supra-molecular, and materials chemistry, apart from its role in catalysis. 41 Fluorine, chlorine, bromine and iodine are XB donors with an ability that increases with increasing atom polarizability and decreasing atom electronegativity, i.e., F < Cl < Br < I. Astatine was expected from early theoretical studies to be the strongest XB donor atom, ^{42,43} despite the persistent lack of experimental data. ⁴¹ We reported in 2018 the first characterization of XBs involving At through the formation of adducts between astatine monoiodide (AtI) and nine Lewis bases. The agreement between the measured equilibrium constants and the computed ones for the formation of At-mediated XB complexes allowed us to unambiguously identify each adduct (Figure 5). This work and the next ones confirmed the highest donating ability of a tatine, especially compared to iodine. 44 Practical interests of this high ability to form XB exist for the improvement of ²¹¹At-labeling strategies.

Indeed, an At-mediated XB may explain the robust radiolabeling of bis-(nido-carboranylmethyl)benzene derivatives previously considered as coupling reagents.⁴⁵

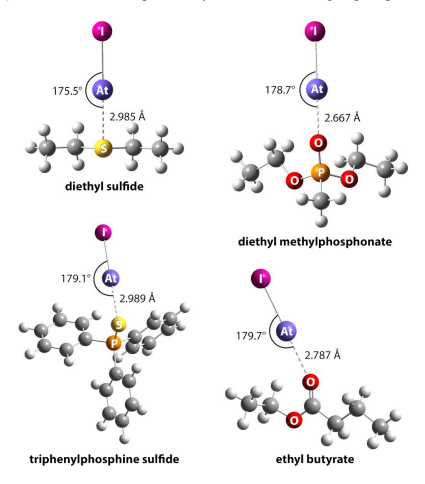


Figure 5. Calculated structures (at the 2c-PW6B95/aug-cc-pVDZ-PP level of theory) of the complexes between a statine monoiodide and Lewis bases. The interaction distances, shorter than the sum of van der Waals radii, and the interaction angles, close to 180°, fulfill the geometrical characteristics of XB interactions. Data from ref. 1.

It is worth noting that some astatinated compounds may form weaker XB interactions than their iodinated counterparts, e.g., At_2 vs. I_2 . 26,46,47 Quantum mechanical calculations have evidenced that relativistic effects are responsible for this unexpected behavior. 26,27,46 Indeed, the

electrophilicity at the astatine σ -hole decreases upon the inclusion of SOC, making At a weaker XB donor.

II.3. Charge-shift bonding

Alongside the two traditional bond families, covalent and ionic, charge-shift (CS) bonds have recently been proposed as a third bonding modality. 48 For a few σ-bonds of the general type A-X, the bonding itself owes its entire origin to large and dynamic fluctuations of the bonding electron-pair: $^-A|X^+ \leftrightarrow A-X \leftrightarrow ^+A|X^-$. This CS mechanism is connected to the lone-pair bond-weakening effect (LPBWE), i.e. repulsions between the bonding electrons and the lonepairs adjacent to the bond, which have the same symmetry as the bond (here σ). The optimal covalent-ionic mixing results in a tremendous resonance energy. CS bonding was first characterized in homopolar and heteropolar bonds involving fluorine. 49 Indeed, electronegative and/or lone-pair-rich elements such as halogens are prone to form CS bonds. Among the theoretical tools for chemical bond analysis, the topological approaches based on the electronic localization function (ELF) and the electron density (QTAIM) are able to reveal the characteristics of CS bonds. 48 Within these topological approaches, the 3D real-space is divided into mutually disjoint electronic volumes (or basins) that have chemical significance in terms of bonding, lone pairs, core regions, etc. Thanks to recent developments that extended the ELF and QTAIM topological analyses to spin-orbit-coupled wavefunctions, 50-52 characteristics of CS bonding have been evidenced for some bonds involving astatine. For instance, (i) the hypothetical At₂ diatomic molecule do not satisfy the standard QTAIM classification (Figure 6) because of a depleted electron density at midbond (ho_{cp}), associated to a small positive second

derivative $(\nabla^2 \rho_{cp})$, ^{53,54} and, (ii) regarding heteropolar bonds with carbon atoms or in the AtI species, the electron distribution between the bonding and the valence non-bonding ELF basins displays specific features of LPBWE (revealed by SOC, **Figure 6**). ⁵³ The astatine propensity to form CS bonds, greater than iodine, resembles the behavior of the lightest halogens, fluorine and chlorine.

QTAIM descriptors

	$ ho_{cp}$ (a.u.)	$ abla^2 ho_{cp}$ (a.u.)	delocalization index
covalent bonding	> 0.2	< 0	~ formal bond order
ionic bonding	< 0.1	>0	~0
At ₂ :	0.05	0.03	0.78

ELF electron populations vs. Lewis pictures

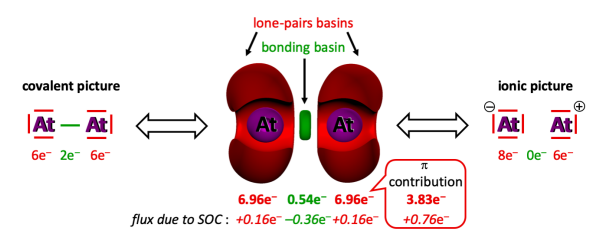


Figure 6. Compared QTAIM and ELF topological analyses of the At₂ species. Data from ref. 52, 53.

Detailed ELF and QTAIM topological analyses have shown that the inclusion of SOC in relativistic calculations emphasizes the LPBWE for bonds involving a statine. ^{27,46,47,50–54} It turns to an enhanced propensity of a statine to form CS bonds, resulting in a locally decreased

electrophilicity at the astatine σ -hole in R-At compounds. Hence, the weakened XB donating ability previously assigned to the relativistic behavior of astatine electron, is eventually connected to the CS character of At-bonds. ^{27,47}

Overall, the multifaceted character of astatine discussed in this section offers many potential options regarding the preparation of ²¹¹At-labeled radiopharmaceuticals, from covalent halogen chemistry to metal-ligand coordination, without neglecting the propensity of astatine to form strong XB interactions.

III. Radiolabeling chemistry with a statine-211: improving methods and in vivo stability

III.1. Astatoaryl compounds

The assumption that astatine would behave similarly to iodine has historically led to radiolabeling strategies consisting in transpositions from well-known radioiodination procedures, with some successes, but also unexpected results. At⁻ and At⁺ are viewed as a typical halogen species in classical nucleophilic and electrophilic reactions used in aryl labeling protocols as discussed in detail elsewhere (**Figure 7**).⁶

Halogen exchange
$$\begin{array}{c} 211 \text{At} \\ \text{CuSO}_4, \, \text{SnSO}_4, \\ \text{gentisic acid, citric acid} \\ 60 \, \text{min, } 120^{\circ}\text{C} \\ \end{array}$$

$$\begin{array}{c} 211 \text{At} \\ \text{RCY} = 67\text{-}80\% \\ \end{array}$$

$$\begin{array}{c} 111 \text{At} \\ \text{OH} \\ \text{Halodediazotation} \\ \text{Halodeprotonation} \\ \end{array}$$

$$\begin{array}{c} 211 \text{At} \\ \text{OH} \\ \text{1h, } 50^{\circ}\text{C} \\ \end{array}$$

$$\begin{array}{c} 211 \text{At} \\ \text{Na}_2 \text{So}_2 \text{Os} \\ \text{H}_2 \text{O} \\ \text{10-15 min, } 100^{\circ}\text{C} \\ \end{array}$$

$$\begin{array}{c} 111 \text{At} \\ \text{RCY} = 68\% \\ \end{array}$$

$$\begin{array}{c} 111 \text{At} \\ \text{RCY} = 68\% \\ \end{array}$$

$$\begin{array}{c} 111 \text{At} \\ \text{Chloramine-T} \\ \text{EtOH/CHCl}_3/\text{AcOH} \\ \text{25 min, r.t.} \\ \end{array}$$

$$\begin{array}{c} 211 \text{At} \\ \text{RCY} = 74\% \\ \end{array}$$

Figure 7. Conventional ²¹¹At-labeling approaches discussed in ref. 6.

Conversely, the direct radiolabeling of proteins is a striking example of how astatine can demonstrate unanticipated outcomes. Indeed, early attempts consisted in adapting the well-known radioiodination of proteins by electrophilic aromatic substitution on tyrosine residues (**Figure 8a**),⁵⁵ but results were somewhat disappointing. Although radiochemical yields (RCYs) were high (>80%), the labeling stability was low, resulting in a fast release of astatine from the protein.⁵⁶ Initially attributed to the supposedly weak At-tyrosine bond, the origin of this instability was latter assumed to arise from the formation of a weak At-sulfur bond with cysteine

residues.⁵⁷ However, this interpretation deserves to be reconsidered if we look closely at the experimental conditions. In light of the Pourbaix diagram at near neutral pH and oxidizing conditions, AtO⁺ or an hydrolyzed form of AtO⁺ is most probably the involved species that binds to proteins. Our DFT calculations support interactions with thiol-containing molecules via the oxygen atom in AtO⁺, leading to a O–S covalent bond.⁵⁸

The strategies developed after these initial attempts consisted in two-step procedures, *i.e.* the synthesis of stable astatoaryl prosthetic groups followed by their conjugation to the protein. The main approach developed by Wilbur and Zalutsky uses N-succinimidyl-[211 At]astatobenzoate ([211 At]SAB) - obtained by electrophilic astatodemetallation of the organotin precursor - which is then conjugated to the amino groups of lysines residues (**Figure 8b**). The method has become a gold standard, applied in the first clinical trial with 211 At, and that we also used in our early investigations. But we faced limits, in particular reproducibility issues mostly due to the lack of robustness of the required electrophilic species, owing to (i) its restricted E_H -pH range of existence, in addition to (ii) radiolysis issues leading over time to changes in astatine speciation. Furthermore, another limit is the suboptimal bioconjugation step yield (50% to 75% at best) which lowers the overall procedure efficiency. From these observations, we started a series of investigations aiming at improving the access to 211 At-labeled antibodies.

a) Direct radiohalogenation

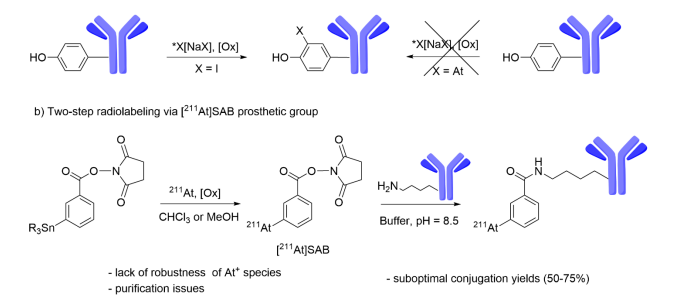


Figure 8. Early developed protocols for ²¹¹At-labeling of proteins.

Ionic liquid organotin precursors. A critical step in conventional [211At]SAB radiosynthesis is the purification to remove unreacted astatine, oxidizer and, more importantly, toxic stannylated precursor/side products. This is usually performed using HPLC, but at the cost of a lengthy step and considerable retention of radiolabeled product in the chromatographic system. Simplified purification steps can be achieved using resin supported precursors, as demonstrated with the preparation of *meta*-[211At]astatobenzylguanidine for the treatment of neuroendocrine tumors. With the perspective of automated [211At]SAB radiosynthesis, we developed a further simplified approach based on the use of an ionic liquid supported organotin compound. The precursor, obtained in 3 steps from an imidazolium hexafluorophosphate moiety, led to a shorter radiolabeling procedure thanks to faster purification step by filtration on a silica cartridge instead of usual HPLC. Furthermore, the negligible activity retention observed contributed to improved RCYs for the preparation of [211At]SAB and corresponding radioimmunoconjugates (Figure 9).

Figure 9. Preparation of [²¹¹At]SAB using an ionic liquid supported precursor.

Aryliodonium salts chemistry. Bringing further the idea to move away from astatodestannylation and unstable electrophilic astatine species required, we investigated the use of the At species and turned our attention to the nucleophilic aromatic substitution of diaryliodonium salts. The reactivity of a statide revealed an unexpected higher reactivity than anticipated from the trend observed with lighter halogens, allowing to develop radiolabeling protocols at a significantly lower temperature and with a broader scope of solvents than radiofluorine and radioiodine.⁶⁴ The reaction was applied to the radiosynthesis of [²¹¹At]SAB with the advantage of significantly improved reproducibility and efficiency as well as simplified purification step using silica cartridges instead of HPLC. 65 The precursor was then used successfully for the ²¹¹At-labeling of a monoclonal antibody (mAb) in a preclinical study on the treatment of multiple myeloma. 66 Other prosthetic groups for bioconjugation by click chemistry were also obtained in high RCYs thanks to a high selectivity of substitution towards the aryl position of interest (Figure 10). The astatinated benzylazide and tetrazine compounds led to significantly improved bioconjugation yields (> 90%), in comparison with the initial [211At]SAB strategy when applied to a mAb respectively preconjugated to strained alkyne and transcyclooctene click chemistry partner moieties.⁶⁷

Figure 10. Nucleophilic ²¹¹At-labeling of diaryliodonium salts.

Arylboron chemistry. Whereas aryliodonium salts and click chemistry allowed us to improve both the efficiency and the reproducibility of mAb astatination, they still resulted in time consuming two-step procedures. A one-step approach using mAbs pre-conjugated with the precursor has been reported by Lindegren's group, but it remains based on the use of electrophilic astatodestannylation reaction.⁶⁸ With the aim to reach the optimal method, we continued our investigations on the use of At⁻. The copper assisted radiohalogenation of arylboron precursors (derived from the Chan-Evans-Lan cross-coupling reaction) has recently

86%

13%

been reported by other groups. In particular, it was shown that a broad scope of arylboronic esters could be efficiently astatinated at room temperature in organic solvents.⁶⁹ Pushing the investigation further, we found that the reaction, applied to arylboronic acids, could remain highly efficient in aqueous solution even at precursor concentration as low as 250 µM, provided that a ligand such as 1,10-phenanthroline is added to keep the catalyst active. These conditions, now compatible with proteins, were then applied to a multiple myeloma targeting mAb preconjugated with an arylboronic acid moiety. An efficient and fast procedure was set up, which outperforms the previously reported ones in terms of RCY, specific activity and protocol duration (Figure 11).⁴ It is furthermore applicable to radioiodination, allowing the development of radiotheranostic pharmaceuticals based on ¹²³I or ¹²⁴I for imaging the lesions, and on ¹³¹I or ²¹¹At to treat them.⁷⁰ In addition, the long-term storability of the pre-conjugated mAb observed (> 1 year) opens possibility to develop radiolabeling kits for easy development in a clinical setting.

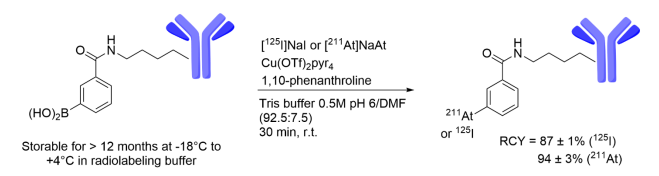


Figure 11. One-step radiohalogenation of a mAb preconjugated to an arylboronic acid moiety.

III.2. Weakness of astatoaryl compounds and alternatives.

The stability of the carbon-halogen bond decreases with increasing halogen size and increasing carbon hybridization state. We confirmed this trend for astatinated compounds by computing descriptors including bond enthalpies and bond orders,⁷¹ which supports the initial choice to

favor astatoaryl compounds. Yet, a number of *in vivo* studies have highlighted the progressive dissociation of the aryl-At bond, at a significantly higher rate than radioiodinated analogues, especially with small biomolecules or antibody fragments that are rapidly metabolized. This is also true when mAbs are internalized within targeted cells.⁷² Searching for potential deastatination mechanisms led us to consider the role of lysosomes found within cells and associated to acidic and oxidizing conditions. Model experiments including incubation of 3-ethylastatobenzoate in oxidizing, Fenton-like conditions, evidenced a significantly faster release of free astatine compared to the radioiodinated analogue, suggesting that oxidation plays an important role in deastatination (**Figure 12**). This hypothesis is supported by the calculated IE of halobenzoates, suggesting a facilitated oxidation of the astatinated derivative, and the lower C-At bond dissociation energy (BDE) for the resulting oxidized form, compared to the iodine analogue.⁷³

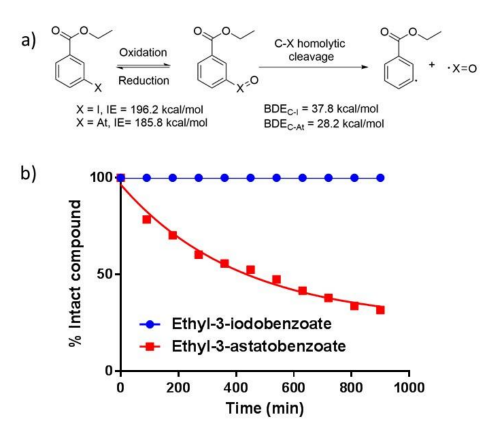


Figure 12. a) Proposed oxidative mechanism leading to C-At bond dissociation *in vivo*. b) Stability of halobenzoate derivatives submitted to Fenton-like oxidative conditions. Data from ref. 73.

Whereas reported modulations of the astatoaryl structure have not yet clearly improved *in vivo* stability, this issue may be addressed by using interactions other than the covalent C–At bond. Exploiting the metallic properties of electrophilic astatine was pioneered by Ludwig *et al.* through complexation studies with thioether ligands, ⁷⁴ as well as subsequent attempts with well-known metal chelators such as DTPA. However, none have yet led to *in vivo* stable products, their biodistribution patterns in mice being nearly identical to sodium astatide. The most successful alternative to date was developed by Wilbur's group. It relies on the formation of a covalent B–At bond with boron clusters. ⁷² In particular the decaborate moiety drastically reduced

in vivo release of At compared to astatoaryl (reduced activity uptake in target organs of free astatine such as thyroid (neck), stomach, lungs or spleen in **Figure 13**). Our DFT calculations allowed to establish a relationship between the B–At and C–At bond enthalpies and *in vivo* stability, the higher the bond enthalpy the lower the content of released astatine being trapped by the thyroid.⁷⁶ However, this chemistry may impair the pharmacokinetics of the carrier compounds, probably due to the double negative charge on the boron cluster.⁷⁷

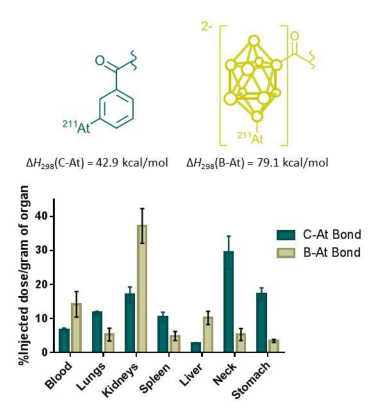


Figure 13. Boron cluster vs astatoaryl stability. Data from ref. 76, 78.

Astatine being a soft base like iodine (Table 1), investigations of the interaction between astatide and soft metal cations recently emerged. Pruszyńsky reported the investigation of astatide as a ligand of rhodium(III) and iridium(III) metal centers trapped within a polythioether chelating agent (**Figure 14**). However, applications on the radiolabeling of relevant biomolecule has not yet proven more stability than the use of astatoaryl prosthetic groups.⁷⁹ Looking for

softer metallic centers, we used rhodium(I) complexed within a *N*-heterocyclic carben ligand (NHC), and we introduced a tatide as a ligand by nucleophilic substitution of a chloride, leading to a compound that was highly stable in human blood serum. ⁸⁰ The impact of the nature of the metal and of NHC ligands on the metal-At bond stability is currently under investigation.

Figure 14. Complexes with a tatine-metal covalent bond.

Conclusion

Over the past decade, we have taken research on astatine from an all-empirical work, relying on extrapolations from lighter halogens, to a new era where computer assisted experimentation has revealed unanticipated findings. If the construction of the Pourbaix diagram has confirmed both metallic and halogen behavior of astatine, its expected superiority in the field of halogen-bond interactions is also ambiguous. Just as Janus presents two faces (one turned to the past, the other to the future), astatine seems to have two antinomic characters with charge-shift bonding that hinders halogen bonding. This duality stands as a whole for astatine, unquestionably the last naturally occurring element of the halogen family, which however exhibits simultaneously a metallic character. Is the final option to merge astatine into the metalloid family? Based on current knowledge, this question probably cannot be definitively answered. Fortunately, the growing community having access to this rare radioelement will certainly accelerate discoveries in the coming decade and contribute in solving the puzzle of astatine.

Even if insufficiently defined, astatine's multifaceted character is fortunate because it allows as many possibilities to improve radiolabeling chemistry as illustrated with the novel approaches discussed herein. These methods, in particular the ready-to-use arylboron chemistry, should ease the clinical transfer of alpha emitters-based radiopharmaceuticals by lifting logistical obstacles currently encountered. With an increasing knowledge on astatine chemical properties allowing for the development of rationally designed labeling procedures, it makes no doubt that applications on ²¹¹At-based cancer therapy will be facilitated in the near future.

Biographies

François Guérard received his Ph.D. in chemistry from the University of Nantes (France) in 2010. He then joined the Brechbiel's group (National Cancer Institute, USA) to develop ⁸⁹Zr labeling chemistry. In 2017, he was appointed CNRS research associate at Inserm (Nantes), in the Nuclear Oncology Group where he leads a research program on ²¹¹At-labeling chemistry.

Clémence Maingueneau received her Ph.D. in chemistry from the University of Caen Normandie (France) in 2019. Currently, she is a postdoctoral fellow in the Nuclear Oncology Group (Inserm, Nantes). Her research interests focus on ²¹¹At-labeling chemistry.

Lu Liu received her Ph.D. in analytical chemistry and Radiochemistry from IMT Atlantique (Nantes, France) in 2020. Her research focus on the fundamental aspects of astatine chemistry, including its speciation and reactivity in solutions.

Romain Eychenne received his Ph.D. in chemistry from the University of Toulouse (France) in 2017. He is currently a research engineer at the Arronax cyclotron facility, with a focus on radiopharmaceutical chemistry and especially ²¹¹At-labeling chemistry.

Jean-François Gestin received his Ph.D. in chemistry from the University of Nantes (France) in 1990 after spending one year at CEA (France) and two years at Brookhaven National Laboratory

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Gilles Montavon received his Ph.D. in radiochemistry from the University of Paris-Sud in 1996. Since 1998, he is a CNRS researcher in the radiochemistry group at Subatech laboratory where he leads a team interested in the behavior of radioelements at ultra-trace concentrations for medical and environmental issues.

Nicolas Galland received his Ph.D. in physical chemistry from the University of Bordeaux (France) in 2002. Since 2003, he is an associate professor at University of Nantes (France). His research interests include modeling non-covalent interactions (hydrogen and halogen bonds) and the development of novel tools for bonding analysis in compounds containing heavy atoms.

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