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Title: Orthogonal Stability & Reactivity of Aryl Germanes Enables Rapid and Selective (Multi)Halogenations

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Synthetic Methods

multifunctional radiotracer for SPECT and PET imaging via the

ipso-halogenatior

SiR₂ / B(OR)₂ need activation

X⁺ or X

B(OR)₂ or

SnR₃ toxic, less robust & purification issues

B(OH)₂ unstable & difficult to prepare

BPin

I⁺ or Br⁺

○: BPin, SiR₃

Reactivity: [Gel >> Bpin. SiR_o (= selective!)

Rapid & superior functional group tolerance

fluorination

Nucleophilic F (or ¹⁸F) substitutes at I
 SiR₃ lower reactivity in fluorination

Bpin & SiR₃ unstable in

No more iodination-site

fluorination (consumed)

·[l/Br]

introduction of the respective isotope [123I] or [18F].[14]

Established Halogenation Methods

Limited functional group tolerance

Towards Multiple Halogenations - Challenges

R₂Sn

R₂Si

PinB

Strategy A:

Strategy B:

Figure 1. Challenges in halogenation approaches and this work.

SnR₂

[Sn] most reactive

BPin

iodination

fluorination

This Work: Orthogonal stability & reactivity of aryl germanes in halogenation

Unselective halogenations cause challenges in separation of mixtures.

In this context, both halogens are ideally introduced rapidly and with positional selectivity, as the size and electronic effects of the halogens

impact the fate and binding efficiency^[2,3] of a halogenated drug.

Consequently, ideally, two separate and chemically orthogonal

handles are employed, which allow for fully independent

halogenations, irrespective of their relative positioning or the

However, of the currently available 'handles', i.e. B(OH)₂, boronic

esters, SiMe₃ and SnBu₃, the boronic acids are difficult to install, relatively unstable and challenging to purify.^[15] Stannanes show high

reactivities in halogenation and find usage in commercial

radiotracers;^[6d] however, they are also toxic and purification of the toxic by-products is frequently challenging. While aryl boronic esters

or aryl silanes display comparatively greater stability,^[16] this comes

at the expense of their reactivities in ipso-iodination and bromination,

which often require nucleophilic activation or metal catalysis to

facilitate the reaction.^[9b,10d,10e,11a,11b,17] This in turn negatively impacts

GeEt₂

No meta-substitution accessible

direct halogenation

I ow site-selectivity

Example & Uses

synthesis

alters properties

halogen bonding

GeEt₃

IGel uniquely stable in fluorination

presence of additional functionality.

scope and functional group tolerance.

[Ge] non-toxic, highly robust

radioimaging

JUSCL

Orthogonal Stability & Reactivity of Aryl Germanes Enables Rapid and Selective (Multi)Halogenations

Christoph Fricke, Kristina Deckers and Franziska Schoenebeck*

Abstract: While halogenation is of key importance in synthesis and radioimaging, the currently available repertoire is largely designed to introduce a single halogen per molecule. This report makes the selective introduction of several different halogens accessible. We showcase the privileged stability of non-toxic aryl germanes under harsh fluorination conditions (that allow for selective fluorination in their presence), while displaying superior reactivity and functional group tolerance in electrophilic iodinations and brominations, outcompeting silanes or boronic esters under rapid and additive-free conditions. Mechanistic experiments and computational studies suggest a concerted electrophilic aromatic substitution as underlying mechanism.

While aryl halides are of utmost importance as key functionalities to enable selective metal-catalyzed C-C or Cheteroatom bond formations.^[1] they are also of importance beyond synthesis, impacting the activities of drugs,^[2,3] material properties (e.g. solubility of nanoribbons)^[4] and (supramolecular) self-assembly via halogen bonding.^[5] Moreover, the use of radioactive isotopes, especially ¹⁸F and ¹²³I, allows for *in vivo* radioimaging via PET and SPECT techniques in the study of biological and physiological processes.^[6] Consequently, there is a significant interest in devising new halogenation strategies that satisfy the needs for efficiency, selectivity, non-toxicity, functional group tolerance as well as rapid speed.^[7] Impressive synthetic advances have been made in recent years, involving approaches of direct C-H functionalization via metal-catalyzed or metal-free (photoredox) halogenation strategies,^[8] halogen exchange (e.g. ArX to ArF)^[9] or the halogenation of suitable precursor functionalities, *i.e.* boronic acid derivatives,^[10] silanes^[11] and stannanes.^[12]

However, the currently available synthetic repertoire was primarily developed for the introduction of a *single halogen* per precursor molecule. By contrast, the development of halogenated materials or drugs would greatly benefit form the ability to introduce multiple halogens late in a synthesis, since especially iodinated building blocks suffer from incompatibility of the C-I bonds with most metal-catalyzed coupling chemistry, which in turn is powerful to connect building blocks to larger molecules.^[13] Moreover, in a radio-halogenation context, the presence of more than one halogen in a molecule, especially I and F, could enable applications as a

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Figure 2. Organogermanes are uniquely stable in fluorination (a) and rapidly reactive in selective iodination/bromination reactions (b-e).

However, in the context of fluorination, all these functional groups are reactive (*i.e.* mostly unproductively consumed due to their instability), especially in the established radio-fluorination methodologies based on KF/cryptand[2.2.2]/ Cu(OTf)2^[18] or Selectfluor (see Figure 2a). An initial fluorination, followed by iodination is therefore not accessible. Conversely, if there is initial iodination at the SnR₃-site, then Bpin or SiMe₃ are not suitable to achieve efficient fluorination thereafter: while for SiMe₃, fluorination is inefficient,^[20] the ¹⁸F installation at BPin is achieved *via* nucleophilic strategies (*e.g.* Cu(OTf)2(py)4 with KF at >100°C), which lead to competing C-I substitution and overall product mixtures (see Figure 1).

As such, there is a need for a new and orthogonal functional group, and we targeted the trialkyl germanium functionality. This non-toxic^[21] class of reagents offers high stability against moisture, air, acids and bases,^[22b] and has shown promise in reactions with molecular halogens.^[19] We recently uncovered that aryl germanes displayed privileged and orthogonal reactivities in metal catalysis over alternative functionalities,^[22,23] and therefore envisioned that potentially orthogonal halogenations might also be feasible, which might unleash access to selective multi-halogenation of molecules.

We initially subjected 4-tolyl germane to the established nucleophilic or electrophilic fluorination methods using KF, cryptand[2.2.2] and Cu(OTf)₂ or Selectfluor (see Figure 2a). Interestingly, the aryl germane fully tolerated these conditions: we recovered 4-tolyl triethyl germane in >99% yield. In stark contrast, the corresponding boron, silane and stannane compounds were fully consumed under these conditions (largely unproductively). There is hence a remarkable stability associated with germanes with respect to harsh fluorination conditions, which uniquely allow for fluorination in their presence. To investigate this further, we prepared the Sn/Gecontaining bifunctional substrate 1 (Figure 2a), and subjected it to

fluorination. Remarkably, the application of the Selectfluor/AgOTfmediated fluorination^[24] protocol resulted in the fully selective fluorination of the SnBu₃-site in the presence of the GeEt₃-site. There was no consumption of the Ge-functionality.

Conversely, the C-GeEt₃ proved to be highly reactive in electrophilic iodination: the iodination of 4-tolyl germane in DMF with *N*-iodosuccinimide (NIS) occurred in good yield within 2 h at room temperature to give **3** (62%), without the need for additives or metal catalysts (Figure 2b).^[25] The corresponding bromination of 4-methoxyphenyl germane proved to be even more facile and was complete within 15 min at room temperature (see SI for details). Consequently, this now offers the opportunity to introduce fluorine and subsequently iodine (or bromine) fully selectively.

We next explored our protocol in the halogenation towards *meta*iodobenzyl guanidine (MIBG) derivatives, which are used (as their stannane analoges) as tumor therapeutics or for diagnostic imaging *via* SPECT tomography (see Figure 2e).^[26] To our delight, the iodination with NIS, and bromination with NBS proceeded in high yields and short reaction times. We also performed a halogenation with a mixture of *N*-chlorosuccinimide (NCS) and NaBr (as radiolabeling generally relies on nucleophilic isotopes). Under these conditions, we successfully obtained [Br]MIBG **10a** in 78% yield after 60 min at 80 °C.

We next investigated the potential chemoselectivity of aryl germanes relative to boronic esters and silanes. Interestingly, despite its privileged robustness towards harsh fluorination conditions, the germane is much more reactive in iodination and bromination than the corresponding boronic ester derivatives or silanes, which are unreactive when employing similarly mild reaction conditions (see Figure 2c/d). Our intermolecular competitions of halogenating aryl germanes *versus* traditional reagents showcased high chemoselectivity for germane functionalization over boronic acids,



boronic esters and silanes (see Figure 2c). Even substrates containing competing boronic ester (Bpin) or silane (SiMe₃) functionalities showed exclusive and high reactivity at the C-GeEt₃ bond, leaving the Bpin- or SiMe₃ moieties untouched (see Figure 2d; **7a/b** and **8a/b**).

As such, the germanes show privileged robustness in fluorination, but superior reactivity in iodination and bromination, which allows for the selective introductions of the halogen couples I/Br, or F/I or F/Br.

To facilitate wider applications, knowledge about functional group tolerance in the respective iodination and halogenation of the germanium-functionality is imperative. We therefore next assessed the scope of halogenation of organogermanes.

To our delight, a broad range of electron-rich aryl germanes was halogenated in good to excellent yields (3, 11-15, see Table 1). Especially trimethoxy-phenyl- (14) or naphtyl-derivatives (15) are often challenging to halogenate selectively, as they are prone to undergo multiple – and often unselective – competing direct C-H-halogenation reactions.

Pleasingly, also electron deficient aryl germanes, such as prehalogenated substrates were halogenated in excellent yields (16-19). This feature allows for the construction of multiply halogenated scaffolds for diverse synthetic purposes. Even sterically demanding o,o-disubstituted (20 and 21) and heterocyclic scaffolds (22-25), whose boronic acid derivatives show a high tendency to decompose,^[15] underwent halogenation in high yields. Especially pleasing is the selective halogenation of 3-thiophenyl substrates (23 and 25), as the 2-thiophenyl-position is the more nucleophilic site and hence usually preferentially targeted in reactions with electrophiles.

Since electrophilic halogenation strategies can suffer from drawbacks such as strong oxidizing behavior or high affinity to react with *e.g.* alkenes, alkynes^[27] or α -acidic ketones^[28] in competing pathways,^[30] we further investigated the generality of our method with an additive screen.^[31] We tested a variety of potentially sensitive additives (Figure 3, top; see SI for details) and pleasingly found that owing to the privileged reactivity of the Ge-functionality, numerous basic, nucleophilic and electrophilic additives were fully tolerated in the halogenation.





Yields of isolated products are given. ^aYields determined by quantitative ¹H NMR using mesitylene as internal standard. ^bPerformed at 50 °C and with prolonged reaction time (see SI for details).

We tested 52 additives in total – including 16 heterocyclic, 24 carbonyl-containing and 11 N- or O-protected compounds; 44 thereof were not affected by the halogenation and recovered after the reaction (in > 66%).



Figure 3. Additive screen to test functional group tolerance. Number of reactions with high yield (>66%), medium yield (34-66%) and low yield (<34%).

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Most heterocycles, amides, lactones, and even activated alkenes, alkynes or α -acidic ketones as additives gave high yields of the desired halogenation on the germanium site, leaving the additive untouched. However, additives containing silyl-protected alcohols or acidic protons (*e.g.* R-OTMS or R-NH₂) underwent side reactions with the electrophilic halogenation reagent thus substantially lowering the yields.



Figure 4. (a) Experimental LFER analysis and computational study^[29] of the bromination using NBS. (b) Comparison of transition state energies for aryl germanes and aryl silanes. Free energies in (a) and (b) computed at the CPCM (DMF) M06/6-311++G(d,p) (SDD)// ω B97XD/def2SVP level of theory. To account for charged intermediates, geometry optimizations were performed with an implicit solvent model. Free energies are given in kcal mol⁻¹.

In stark contrast, when we performed an analogous additive compatibility test with PhSiMe₃ and PhBpin (see Figure 3, bottom), we found that while PhGeEt₃ fully tolerated alkyne **26** and alkene **27** as additives in bromination with NBS, these functionalities were consumed in the corresponding reactions with PhBpin and PhSiMe₃, and no bromination of BPin or SiMe₃ took place.

To gain insight on the origins of high reactivity of Ge in halogenation we next performed mechanistic investigations, combining a set of experimental and computational investigations (Figure 4). A linear free energy relationship (LFER) analysis of the reaction with a ρ_{σ} value of -4.8 indicated the build-up

of a positive charge in the transition state and hence supported the hypothesized pathway $via S_EAr$ activation of the C-Ge bond.

In line with this, our computational studies indicated that germanium is halogenated in a *concerted* electrophilic aromatic substitution (in the gas-phase and under implicit solvation optimization using CPCM solvation model). The generally assumed Wheland intermediate was not observed.^[32] An activation free energy barrier of $\Delta G^{\ddagger} = 21.9$ kcal mol⁻¹ was calculated^[29] for PhGeEt₃, which is in line with the high experimental reactivity observed for ArGeEt₃ at room temperature. For comparison, the corresponding aryl silane is predicted to react with a barrier of $\Delta G^{\ddagger} = 25.2$ kcal mol⁻¹, in line with the exclusive selectivity for Ge-functionalization.

We further determined the activation barriers for other substituted germanes and plotted these against the corresponding σ -parameter. A linear correlation between the electronic σ -parameter^[33] of the *para*-substituent and the corresponding activation barrier for the concerted transition state was observed.

In conclusion, we developed an operationally simple, rapid and widely applicable halogenation method for the selective introduction of iodine and bromine *via* concerted electrophilic aromatic substitution at germanium. While the aryl germane is superior in reactivity over silanes or boronic esters, it displays unique robustness towards fluorination conditions, which unleashes the possibility for chemoselective and orthogonal introduction of *multiple different halogens* (*i.e.* F/I, I/Br or F/Br) with complete positional control and wide functional group tolerance.

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Conflict of interest

The authors declare no conflict of interests.

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Synthetic Methods

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Orthogonal Stability & Reactivity of Aryl Germanes Enables Rapid and Selective (Multi)Halogenations $F \xrightarrow{f^{+} \text{ or } F^{-}} GeEt_{3} \xleftarrow{F^{+} \text{ or } F^{-}} GeEt_{3} \xrightarrow{I^{+} \text{ or } Br^{+}} \underbrace{I^{+} \text{ or } Br^{+}}_{\bigcirc: \text{ BPin, SiR}_{3}} \underbrace{I^{+} \text{ or } Br^{+}}_{O: \text{ BPin, SiR}_{3}} \underbrace{I^{+} \text{ or } Br^{+}}_{O:$

[Ge] non-toxic, highly robust

Reactivity: [Ge] >> Bpin, SiR₃ (= selective!)
Rapid & superior functional group tolerance

Non-toxic aryl germanes are shown to be of privileged stability under harsh fluorination conditions, while displaying superior reactivity and functional group tolerance in electrophilic iodinations and brominations, outcompeting silanes or boronic esters under rapid and additive-free conditions.

