Check for updates<br>Cite this: Chem. Sci., 2019, 10, 3237<br>All publication charges for this article have been paid for by the Royal Society of Chemistry

Received 15th November 2018
Accepted 28th January 2019
DOI: 10.1039/c8sc05096a
rsc.li/chemical-science

# Synthesis of ${ }^{18} \mathrm{~F}$-difluoromethylarenes using aryl boronic acids, ethyl bromofluoroacetate and $\left[{ }^{18} \mathrm{~F}\right]$ fluoride $\dagger$ 

Jeroen B. I. Sap, ${ }^{\text {a }}$ Thomas C. Wilson, ${ }^{\text {a }}$ Choon Wee Kee, (1) ${ }^{\text {a }}$ Natan J. W. Straathof, (10 ${ }^{a}$ Christopher W. am Ende, ${ }^{\text {b }}$ Paramita Mukherjee, ${ }^{\text {b }}$ Lei Zhang, ${ }^{\text {b }}$ Christophe Genicot ${ }^{\text {c }}$ and Véronique Gouverneur (D) *a


#### Abstract

Herein, we report the radiosynthesis of ${ }^{18} \mathrm{~F}$-difluoromethylarenes via the assembly of three components, a boron reagent, ethyl bromofluoroacetate, and cyclotron-produced non-carrier added [ ${ }^{18}$ Fffluoride. The two key steps are a copper-catalysed cross-coupling reaction, and a Mn-mediated ${ }^{18} \mathrm{~F}$-fluorodecarboxylation.


## Introduction

Positron emission tomography (PET) is a molecular imaging technique that requires molecules labelled with a positronemitting radionuclide. Fluorine-18 is a widely used positron emitting radionuclide in part due to its favourable decay properties, and the numerous clinical applications of 2-deoxy-2-[ $\left.{ }^{18} \mathrm{~F}\right]$ fluoro-d-glucose, a radiopharmaceutical prepared from $\left[{ }^{18} \mathrm{~F}\right]$ fluoride. ${ }^{1}$ While radiochemists have in recent years focused their efforts on methods enabling ${ }^{18}$ F-fluorination ${ }^{2}$ and ${ }^{18} \mathrm{~F}$ trifluoromethylation of (hetero)arenes,,$^{2,3}{ }^{18} \mathrm{~F}$-difluoromethylation reactions have been less studied despite the importance of the $\mathrm{CF}_{2} \mathrm{H}$ motif ${ }^{4}$ in radioligand design for drug discovery programmes. In 2013, we reported a $\mathrm{Ag}(\mathrm{I})$-mediated ${ }^{18} \mathrm{~F}$ fluorodecarboxylation of 2-fluoro-2-arylacetic acids with $\left[{ }^{18} \mathrm{~F}\right]$ Selectfluor (bis)triflate leading to $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H} .{ }^{5}$ Subsequently, we disclosed a $\mathrm{Ag}(\mathrm{I})$-mediated halogen exchange reaction using $\left[{ }^{18} \mathrm{~F}\right]$ fluoride. ${ }^{6}$ In 2016, a multi-step method to label $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H}$ from aryl (pseudo)halides was disclosed by Ritter and co-workers. ${ }^{7}$ Later, Liang and co-workers demonstrated that halogen exchange of benzyl (pseudo)halides with $\left[{ }^{18} \mathrm{~F}\right] f l u o r i d e ~ f o l l o w e d ~ b y ~ o x i d a t i v e ~$ benzylic C-H fluorination with Selectfluor afforded $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H}$ with improved molar activity. ${ }^{8}$ Despite these advances, ${ }^{18} \mathrm{~F}$-difluoromethylation remains a challenging problem, especially for structurally complex targets. We initially considered adapting difluoromethylation reactions operating via $\mathrm{C}-\mathrm{H}$

[^0]functionalisation. ${ }^{9}$ Whilst this strategy is ideal for (hetero)arenes with innate reactivity leading to site-selective ${ }^{18}$ F-difluoromethylation, substrates that are not reactive or too reactive would be unsuitable, thereby limiting applicability for radioligand synthesis. We therefore opted to develop a method using prefunctionalised aryl boron reagents; these are amenable to ${ }^{18} \mathrm{~F}$ fluorination and ${ }^{18}$ F-trifluoromethylation, ${ }^{10}$ so extension to ${ }^{18} \mathrm{~F}$ difluoromethylation was viewed as a valuable development. Building on our $\mathrm{Ag}(\mathrm{I})$-mediated ${ }^{18} \mathrm{~F}$-fluorodecarboxylation towards $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H},{ }^{5}$ a reaction requiring $\left[{ }^{18} \mathrm{~F}\right]$ Selectfluor (bis)triflate (Scheme 1A), ${ }^{11}$ and on the Mn-mediated fluorodecarboxylation reported by Groves and co-workers, a reaction using $\left[{ }^{18} \mathrm{~F}\right]$ fluoride (Scheme 1B), ${ }^{12,13}$ we envisaged that the ${ }^{18} \mathrm{~F}$-fluorodecarboxylation of 2-fluoro-2-arylacetic acids with $\left[{ }^{18} \mathrm{~F}\right] f l u o r i d e ~ c o u l d ~ a f f o r d ~[~[~ \$ ~ F ~] ~] ~$ $\mathrm{ArCF}_{2} \mathrm{H}$. The beneficial effect of fluorine substitution on radical stabilisation would be favorable for this process. ${ }^{5,14}$ This approach would require a robust method to cross-couple the aryl boron reagent with ethyl bromofluoroacetate followed by hydrolysis to


Scheme 1 (A) Ag(I)-mediated ${ }^{18} \mathrm{~F}$-fluorodecarboxylation with $\left[{ }^{18} \mathrm{~F}\right]$ Selectfluor (bis)triflate. (B) Mn (III)-mediated ${ }^{18} \mathrm{~F}$-fluorodecarboxylation with $\left[{ }^{18} \mathrm{~F}\right] f l u o r i d e ~ t o w a r d s ~\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCH}_{2} \mathrm{~F}$. (C) Synthetic plan towards $\left[{ }^{18} \mathrm{~F}\right]$ $\mathrm{ArCF}_{2} \mathrm{H}$ from boron reagents and ${ }^{18} \mathrm{~F}$ fluoride.
access the carboxylic acid precursor; we gave preference to a coupling methodology applying Cu-catalysis instead of Pd or Ni , a decision driven by guidelines for residual metals in (radio) pharmaceuticals. ${ }^{15}$ The proposed strategy therefore relies on three readily available components, the boron reagent, ethyl bromofluoroacetate, and $\left[{ }^{18} \mathrm{~F}\right]$ fluoride (Scheme 1C). ${ }^{16}$

## Results and discussion

Preliminary experiments demonstrated that the model fluorosubstituted carboxylic acid 1a is amenable to fluorodecarboxylation with fluoride. When an equimolar mixture of 1a and 2a was treated with $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}(2.5 \mathrm{~mol} \%), \mathrm{Et}_{3} \mathrm{~N} \cdot 3 \mathrm{HF}$ (1.2 equiv.) and PhIO (3.3 equiv.) in MeCN at $50^{\circ} \mathrm{C}$, 3 a and $4 \mathbf{4}$ were obtained in $44 \%$ and $20 \%$ yield, respectively. This result indicates that the fluorine-substituted precursor 1a is more reactive than nonfluorinated 2a towards fluorodecarboxylation (Scheme 2A). We verified that product 4a did not undergo fluorination via $\mathrm{C}-\mathrm{H}$ functionalisation under these conditions. ${ }^{17}$ When an excess of 1a ( 1 equiv.) was treated with TBAF ( 0.1 equiv.), PhIO ( 0.5 equiv.) and $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}(0.2$ equiv.) in MeCN, 3 a was obtained in $50 \%$ yield (determined by ${ }^{19} \mathrm{~F}$ NMR based on TBAF consumption) (Scheme 2B). Notably, quantitative fluoride incorporation was observed applying similar reaction conditions to the preformed hypervalent iodine complex $\mathbf{5 a}$ (Scheme 2C). These preliminary data boded well for ${ }^{18} \mathrm{~F}$-labeling with $\left[{ }^{18} \mathrm{~F}\right] f l u o r i d e ~ a s ~ t h e ~$ limiting reagent, and prompted the development of a robust protocol to convert aryl boron reagents into 2-fluoro-2-arylacetic acids.

The cross-coupling of arylboronic acids and ethyl bromofluoroacetate has been reported using an excess of boron reagent under Ni or Pd catalysis, but has not been accomplished under Cu catalysis. ${ }^{18-22}$ Initial studies reacting [1, $1^{\prime}$-biphenyl]-4-

C


Scheme 2 (A) Competition studies evaluating the effect of fluorine substitution on fluorodecarboxylation. (B) Reaction with sub-stoichiometric fluoride. (C) Reaction of iodine(III) complex 5a with substoichiometric fluoride. Yields of isolated products. $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}=\mathrm{Mn}(I I)$ meso-tetra(2,4,6-trimethylphenyl)porphyrin chloride. ${ }^{\text {a }}$ Yield determined by ${ }^{19}$ F NMR using $\alpha, \alpha, \alpha$-trifluorotoluene as internal standard.
ylboronic acid $\mathbf{6 a}$ (2 equiv.) with ethyl bromofluoroacetate (1 equiv.) in the presence of 1,10-phenanthroline ( $\mathbf{L} 1,20 \mathrm{~mol} \%$ ), $\mathrm{CuI}(20 \mathrm{~mol} \%)$ and $\mathrm{Cs}_{2} \mathrm{CO}_{3}$ ( 2 equiv.) in dioxane ( 0.2 M ) under $\mathrm{N}_{2}$ at $100^{\circ} \mathrm{C}$ afforded 7a in $7 \%$ yield (Table 1, entry 1). When $2,2^{\prime}: 6^{\prime}, 2^{\prime \prime}$-terpyridine ( $\mathbf{L} 2$ ) was used as the ligand, the yield was significantly improved to $58 \%$ yield (Table 1, entry 2). When the stoichiometry was altered to 1 equivalent of $\mathbf{6 a}$ and 2 equivalents of ethyl bromofluoroacetate in the presence of $4,4^{\prime}, 4^{\prime \prime}$-tri-tert-butyl-2, $2^{\prime}: 6^{\prime}, 2^{\prime \prime}$-terpyridine (L3) in toluene instead of dioxane 7 a was obtained in $63 \%$ yield (Table 1, entry 3). Further optimisation increasing the concentration led to the optimal protocol consisting of treating $\mathbf{6 a}(0.1 \mathrm{mmol})$ with ethyl bromofluoroacetate ( 0.2 mmol ), $\mathrm{Cs}_{2} \mathrm{CO}_{3}(0.2 \mathrm{mmol})$, $\mathrm{CuI}(20 \mathrm{~mol} \%)$ and $\mathbf{L} 3(20 \mathrm{~mol} \%)$ in toluene $(0.4 \mathrm{M})$ at $100{ }^{\circ} \mathrm{C}$. Under these reaction conditions, 7 a was isolated in $82 \%$ yield (Table 1, entry 4). A one-pot sequence involving cross-coupling followed by hydrolysis with MeOH and aqueous $\mathrm{K}_{2} \mathrm{CO}_{3}$ afforded 8a isolated in $75 \%$ yield (Table 1, entry 5). In the absence of ligand and/or copper source (Table 1, entries 6, 7), no product formation was observed. Furthermore, no reaction was observed with $\mathrm{CuCl}_{2}$ (Table 1, entry 8), or when the reaction solvent was DMF or DMSO (Table 1, entry 9).

These optimised conditions gave access to a range of 2-fluoro-2-arylacetic acids (Scheme 3). The reaction is broad in scope and tolerates various functional groups, for example alkyl $8 \mathrm{c}-8 \mathrm{e}$ and $8 \mathrm{~s}-8 \mathrm{u}$, alkoxy $8 \mathrm{f}, 8 \mathrm{~g}$, trifluoromethyl 8 h , bromo 8 p , 8q, iodo $8 \mathbf{r}$, and aldehyde $8 \mathbf{i}$ all performed well. Substrates featuring heterocycles such as dibenzofuran $\mathbf{8 j}$, pyridine $\mathbf{8 k}$,

Table 1 Optimisation of the Cu-catalysed cross-coupling of aryl boronic acid 6a with ethyl bromofluoroacetate towards ester 7a and the corresponding carboxylic acid $8 \mathrm{a}^{a}$


| Entry | Solvent | Cu-source | Ligand | Product | Yield $^{b}$ |
| :--- | :--- | :--- | :--- | :--- | :--- |
| $1^{c}$ | Dioxane (0.2 M) | CuI | $\mathbf{L 1}$ | $7 \mathbf{7 a}$ | $7 \%$ |
| $2^{c}$ | Dioxane (0.2 M) | CuI | $\mathbf{L 2}$ | $7 \mathbf{a}$ | $58 \%$ |
| 3 | Toluene (0.2 M) | CuI | $\mathbf{L 3}$ | $7 \mathbf{a}$ | $63 \%$ |
| $4^{d}$ | Toluene (0.4 M) | CuI | $\mathbf{L 3}$ | $7 \mathbf{a}$ | $82 \%{ }^{e}$ |
| $5^{d}$ | Toluene (0.4 M) | CuI | $\mathbf{L 3}$ | $\mathbf{8 a}$ | $75 \%{ }^{e, f}$ |
| $6^{d}$ | Toluene (0.4 M) | CuI | - | $7 \mathbf{a}$ | $0 \%$ |
| $7^{d}$ | Toluene (0.4 M) | - | - | $7 \mathbf{a}$ | $0 \%$ |
| $8^{d}$ | Toluene (0.4 M) | CuCl $_{2}$ | $\mathbf{L 2}$ | $7 \mathbf{a}$ | $0 \%$ |
| $9^{d}$ | DMF or DMSO (0.2 M) | CuI | $\mathbf{L 3}$ | $7 \mathbf{a}$ | $0 \%$ |

${ }^{a}$ Screening reactions performed on 0.1 mmol scale. ${ }^{b}$ Yield determined by ${ }^{19} \mathrm{~F}$-NMR using $\alpha, \alpha, \alpha$-trifluorotoluene as internal standard. ${ }^{c} 2$ equiv. of $\mathbf{6 a}$ and 1 equiv. of ethyl bromofluoroacetate. ${ }^{d} 1$ equiv. of $\mathbf{6 a}$, and 2 equiv. of ethyl bromofluoroacetate. ${ }^{e}$ Yield of isolated product. ${ }^{f}$ Onepot procedure towards $8 \mathbf{8 a}$.


Scheme 3 Scope of Cu-catalysed cross-coupling. The reactions were performed on a 0.3 mmol scale. Conditions: Cul ( $20 \mathrm{~mol} \%$ ), L3 (20 mol\%), aryl boronic acid (1 equiv.), ethyl bromofluoroacetate (2 equiv.), $\mathrm{Cs}_{2} \mathrm{CO}_{3}$ (2 equiv.), toluene ( 0.4 M ) at $100^{\circ} \mathrm{C}$ for 18 h then onepot hydrolysis with $\mathrm{K}_{2} \mathrm{CO}_{3}$ (10 equiv.), $\mathrm{MeOH} / \mathrm{H}_{2} \mathrm{O}$ (1:1), 5 h . ${ }^{\text {a }} \mathrm{Hy}$ drolysis performed as a subsequent step with $\mathrm{K}_{2} \mathrm{CO}_{3}$ (5 equiv.). ${ }^{\mathrm{b}}$ Reaction run on 5 mmol scale. All yields are of isolated products.
triazole 81, and pyrazoles $\mathbf{8 m}, \mathbf{8 n}$ are also suitable coupling partners applying our optimised protocol affording the desired products in $40 \%$ to $70 \%$ yield. Additionally, this cross-coupling


Scheme 4 (A) Competition experiment subjecting equimolar amount of $9 a$ and $5 a$ to $\left[{ }^{18} \mathrm{~F}\right] f l u o r o d e c a r b o x y l a t i o n . ~(B) ~ C o m p e t i t i o n ~ e x p e r i-~$ ment reacting equimolar amount of $1 a$ and $3 a$ with PIDA.
chemistry afforded $\mathbf{8 0}$, a derivative of fenofibrate, in $72 \%$ yield. Finally, the reaction was amenable to scale-up to 5 mmol (Scheme 3, 8m).

The key ${ }^{18}$ F-fluorodecarboxylation step was studied next (Table 2). We started our investigation applying protocol A that consists of reacting in one-pot $\mathbf{8 b}(0.11 \mathrm{mmol})$ with $\mathrm{PhIO}(0.33$ $\mathrm{mmol}), \mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}(2 \mathrm{mg})$ and $\left[{ }^{18} \mathrm{~F}\right]$ TEAF ( $20-30 \mathrm{MBq}$ ) in MeCN $(600 \mu \mathrm{~L})$ at $50^{\circ} \mathrm{C}$; this protocol led to only traces of $\left[{ }^{\mathbf{1 8}} \mathbf{F}\right] \mathbf{3 b}$ (Table 2, entry 1 ). When the loading of PhiO ( 0.02 mmol ) and MeCN $(300 \mu \mathrm{~L})$ was reduced, $\left[{ }^{18} \mathbf{F}\right] \mathbf{3 b}$ was obtained in $6 \% \pm 1 \%$ radiochemical conversion (RCC) (Table 2, entry 2 ). Similar results were obtained in DMF (Table 2, entry 3). Reducing the stoichiometry of $\mathbf{8 b}$ led to a significant increase in RCC $(22 \% \pm 7 \%)$ (Table 2, entry 4). When applying protocol B which consists of mixing 8b with PhIO, a process generating complex $\mathbf{5 b}$, prior to the addition of $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}(2 \mathrm{mg})$ and $\left[{ }^{18} \mathrm{~F}\right]$ TEAF $(20-30 \mathrm{MBq})$

Table 2 Optimisation studies for the ${ }^{18} \mathrm{~F}$ Ifluorodecarboxylation of 8 b


| Entry | Starting material (mmol) | Protocol | Solvent | PhIO (mmol) | $\operatorname{RCC}^{a, b}(n=2)$ |
| :---: | :---: | :---: | :---: | :---: | :---: |
| 1 | 8b (0.11) | A | MeCN ${ }^{\text {c }}$ | 0.33 | $3 \% \pm 1 \%$ |
| 2 | 8b (0.11) | A | MeCN ${ }^{d}$ | 0.02 | $6 \% \pm 1 \%$ |
| 3 | 8b (0.11) | A | $\mathrm{DMF}^{\text {d }}$ | 0.02 | $7 \% \pm 2 \%$ |
| 4 | 8b (0.055) | A | $\mathrm{DMF}^{\text {d,e }}$ | 0.02 | 22\% $\pm 7 \%$ |
| 5 | 5b (0.014) | B | $\mathbf{D M F}^{\text {d,e }}$ | - | $\mathbf{4 0 \%} \pm 10 \%{ }^{f}$ |
| 6 | 5b (0.014) | B | $\mathrm{DMF}^{\text {d,e }}$ | - | $0 \% \pm 0 \%^{g}$ |
| 7 | 8b (0.014) | A | MeCN ${ }^{d}$ | 0.02 | $0 \% \pm 0 \%^{h}$ |
| 8 | 5b (0.014) | B | $\mathrm{DMF}^{\text {d,e }}$ | - | $0 \% \pm 0 \%^{i}$ |

${ }^{a}$ Radiochemical conversion. ${ }^{b} n=$ number of reactions. ${ }^{c} 600 \mu \mathrm{~L}$ of MeCN. ${ }^{d} 300 \mu \mathrm{~L}$ of $\mathrm{MeCN} .{ }^{e} \mathrm{MeCN}$ removed at $100^{\circ} \mathrm{C}$ after dispensing $\left[{ }^{18} \mathrm{~F}\right]$ TEAF. ${ }^{f}(n=10) .{ }^{g}$ Reaction temperature $=100^{\circ} \mathrm{C} .{ }^{h}$ Catalyst is Mn(tmp)OTs. ${ }^{i}$ No Mn Catalyst.
and DMF ( $300 \mu \mathrm{~L}$ ), a drastic improvement was observed, and [ ${ }^{18}$ F] $] \mathbf{3 b}$ was obtained in $40 \% \pm 10 \%$ RCC $(n=10)$ (Table 2, entry 5). When the reaction was run at $100{ }^{\circ} \mathrm{C}$, the formation of [ ${ }^{18} \mathbf{F}$ ] 3b was not observed (Table 2, entry 6). No ${ }^{18}$ F-labelled product was obtained when $\mathrm{Mn}(\mathrm{tmp})$ OTs was used as catalyst, or in the absence of $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}$ (Table 2, entries 7 and 8 ).

The fluorine substituent is advantageous for ${ }^{18} \mathrm{~F}$ fluorodecarboxylation as demonstrated with a competition experiment subjecting equimolar amount of pre-formed hypervalent iodine(III) complexes 9a and 5a to ${ }^{18}$ F-fluorination with $\left[{ }^{18} \mathrm{~F}\right]$ TEAF, $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}$ at $50{ }^{\circ} \mathrm{C}$ in DMF. Difluoromethylarene $\left[{ }^{18} \mathrm{~F}\right] 3 \mathrm{a}$ was the only product observed in the crude reaction mixture (Scheme 4A). Furthermore, an additional competition experiment


Scheme 5 Scope of ${ }^{18}$ F]fluorodecarboxylation applying protocol B: ${ }^{a} \mathrm{ArCHFCO}_{2} \mathrm{H}(0.028 \mathrm{mmol}), \mathrm{PhlO}\left(0.5\right.$ equiv.), $\mathrm{MeCN}(1 \mathrm{~mL}), 50{ }^{\circ} \mathrm{C}$, 10 min then addition of $\left[{ }^{18} \mathrm{~F}\right]$ TEAF $(20-30 \mathrm{MBq}) \mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}(2 \mathrm{mg}), \mathrm{DMF}$ $(300 \mu \mathrm{~L}), 50^{\circ} \mathrm{C}, 20 \mathrm{~min} .{ }^{\mathrm{b}} \mathrm{ArCHFCO}_{2} \mathrm{H}(0.014 \mathrm{mmol})$, $\mathrm{PhIO}(0.5$ equiv.), $\mathrm{MeCN}(1 \mathrm{~mL}), 50^{\circ} \mathrm{C}, 10 \mathrm{~min}$ then addition of $\left[{ }^{18} \mathrm{~F}\right] \mathrm{Mn}(\mathrm{tmp}) \mathrm{F}(841 \mathrm{MBq})$ DCE $(300 \mu \mathrm{~L}), 60^{\circ} \mathrm{C}, 20 \mathrm{~min}$.
showed that the iodine(iiI) complex $\mathbf{5 a}$ is formed preferentially to 9a (Scheme 4B). Fluorine substitution therefore facilitates the two steps of the process leading to fluorodecarboxylation.

Protocol B was applied to a selection of arenes using 20-30 MBq of $\left[{ }^{18} \mathrm{~F}\right]$ fluoride (Scheme 5). Ether, alkyl, aldehyde, ketone, pyridine, triazole, pyrazole, dibenzofuran motifs were all tolerated. The highest RCCs were obtained for electron rich arenes. $\left[{ }^{18} \mathbf{F}\right] 3$ o derived from a boronic acid analogue of fenofibrate was successfully labelled in $23 \% \pm 4 \%(n=4)$. The boronic acid derivative of the COX-II inhibitor ZA140 $6 z$ was transformed into the labelled difluoromethylated product $\left[{ }^{\mathbf{1 8}} \mathbf{F}\right] 3 \mathrm{z}$ in $15 \% \pm$ $2 \% \operatorname{RCC}(n=3)$.

The ${ }^{18}$ F-fluorodecarboxylation of $\mathbf{5 b}$ performed with 841 MBq of $\left[{ }^{18} \mathrm{~F}\right]$ fluoride required further optimisation. For this experiment, $\left[{ }^{18} \mathrm{~F}\right]$ fluoride was captured on an anion exchange cartridge then eluted using a solution of $\mathrm{Mn}(\mathrm{tmp}) \mathrm{Cl}$ in methanol, resulting in $85 \%{ }^{18}$ F-recovery. Lowering the starting material stoichiometry to 0.007 mmol of $5 \mathbf{b}$ and changing the solvent from DMF to DCE afforded the cartridge-purified $\left[{ }^{18} \mathbf{F}\right] \mathbf{3 b}$ in a decay corrected RCY of $12 \%$ and a molar activity of $3.0 \mathrm{GBq} \mu \mathrm{mol}^{-1}$ in a total synthesis time of 30 minutes. ${ }^{23}$

Pleasingly, ${ }^{18} \mathrm{~F}$-fluorodecarboxylation also enabled access to the $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArOCF}_{2} \mathrm{H}$ motif. The only known route to label this motif was reported by our group, and required a multi-step synthesis of the ArOCHFCl precursors which were themselves prepared from ArOCHFCO $2 \mathrm{H}_{2}{ }^{24}$ The reaction of estrone (1.0 equiv.) with ethyl bromofluoroacetate ( 1.5 equiv.) and $\mathrm{K}_{2} \mathrm{CO}_{3}$ ( 2.5 equiv.) in DMF ( 2 mL ) at room temperature followed by a subsequent hydrolysis with aqueous NaOH ( 2.5 equiv.) in $1: 1 \mathrm{H}_{2} \mathrm{O} / \mathrm{Et}_{2} \mathrm{O}$ afforded the precursor required for fluorodecarboxylation. ${ }^{18} \mathrm{~F}$-labelling applying protocol B afforded $\left[{ }^{18} \mathbf{F}\right] 11 \mathrm{a}$ in $21 \% \pm 6 \%$ RCC $(n=3)$.

## Conclusions

In summary, a novel method was developed to transform aryl boronic acids to $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H}$. Prior to labelling, the crosscoupling with ethyl bromofluoroacetate was accomplished under Cu catalysis followed by in situ hydrolysis. The radioisotope ${ }^{18} \mathrm{~F}$ is then introduced in the last step applying a Mn-mediated fluorodecarboxylation with readily available $\left[{ }^{18} \mathrm{~F}\right]$ fluoride. This study has unveiled three key features for this last transformation. Firstly, the fluorine substituent on the carboxylic acid precursor is advantageous for fluorodecarboxylation; secondly, the benefit of preforming the hypervalent iodine complex prior to ${ }^{18}$ F-fluorination; and thirdly, we have established that Mn-mediated fluorodecarboxylation enables access to $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArOCF}_{2} \mathrm{H}$ in addition to $\left[{ }^{18} \mathrm{~F}\right] \mathrm{ArCF}_{2} \mathrm{H}$.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was supported by Pfizer, and the Engineering and Physical Sciences Research Council (EP/N509711/1) (studentship
to J. B. I. S.). We also acknowledge the financial support from the Cancer Research UK (C5255/A16466) (T. C. W.), The Agency for Science, Technology and Research (A*STAR, Singapore) (fellowship to C. W. K.), and UCB (N. J. W. S).

## References

1 (a) S. M. Ametamey, M. Honer and P. A. Schubiger, Chem. Rev., 2008, 108, 1501; (b) P. M. Matthews, E. A. Rabiner, J. Passchier and R. N. Gunn, Br. J. Clin. Pharmacol., 2012, 73, 175.
2 (a) P. W. Miller, N. J. Long, R. Vilar and A. D. Gee, Angew. Chem., Int. Ed., 2008, 47, 8998; (b) Z. Gao, Y. H. Lim, M. Tredwell, L. Li, S. Verhoog, M. Hopkinson, W. Kaluza, T. L. Collier, J. Passchier, M. Huiban and V. Gouverneur, Angew. Chem., Int. Ed., 2012, 51, 6733; (c) B. H. Rotstein, N. A. Stephenson, N. Vasdev and S. H. Liang, Nat. Comтип., 2014, 5, 4365; (d) E. L. Cole, M. N. Stewart, R. Littich, R. Hoareau and P. J. H. Scott, Curr. Top. Med. Chem., 2014, 14, 875; (e) M. Tredwell, S. M. Preshlock, N. J. Taylor, S. Gruber, M. Huiban, J. Passchier, J. Mercier, C. Genicot and V. Gouverneur, Angew. Chem., Int. Ed., 2014, 53, 7751; (f) A. V. Mossine, A. F. Brooks, K. J. Makaravage, J. M. Miller, N. Ichiishi, M. S. Sanford and P. J. H. Scott, Org. Lett., 2015, 17, 5780; (g) C. N. Neumann, J. M. Hooker and T. Ritter, Nature, 2016, 534, 369; (h) S. Preshlock, M. Tredwell and V. Gouverneur, Chem. Rev., 2016, 116, 719; (i) M. K. Narayanam, G. Ma, P. A. Champagne, K. N. Houk and J. M. Murphy, Angew. Chem., Int. Ed., 2017, 56, 13006; (j) X. Deng, J. Rong, L. Wang, N. Vasdev, L. Zhang, L. Josephson and S. Liang, Angew. Chem., Int. Ed., 2018, DOI: 10.1002/anie.201805501.
3 (a) M. Huiban, M. Tredwell, S. Mizuta, Z. Wan, X. Zhang, T. L. Collier, V. Gouverneur and J. Passchier, Nat. Chem., 2013, 5, 941; (b) D. van der Born, C. Sewing, J. D. M. Herscheid, A. D. Windhorst, R. V. Orru and D. J. Vugts, Angew. Chem., Int. Ed., 2014, 53, 11046; (c) T. Rühl, W. Rafique, V. T. Lien and P. J. Riss, Chem. Comтип., 2014, 50, 6056.
4 (a) N. A. Meanwell, J. Med. Chem., 2011, 54, 2529; (b) Y. Zafrani, D. Yeffet, G. Sod-Moriah, A. Berliner, D. Amir, D. Marciano, E. Gershonov and S. Saphier, J. Med. Chem., 2017, 60, 797; (c) C. D. Sessler, M. Rahm, S. Becker, J. M. Goldberg, F. Wang and S. J. Lippard, J. Am. Chem. Soc., 2017, 139, 9325; (d) N. A. Meanwell, J. Med. Chem., 2018, 61, 5822; (e) D. Rageot, T. Bohnacker, A. Melone, J. B. Langlois, C. Borsari, P. Hillmann, A. M. Sele, F. Beaufils, M. Zvelebil, P. Hebeisen, W. Löscher, J. Burke, D. Fabbro and M. P. Wymann, J. Med. Chem., 2018, 61, 10084; (f) G. W. Rewcastle, S. A. Gamage, J. U. Flanagan, R. Frederick, W. A. Denny, B. C. Baguley, P. Kestell, R. Singh, J. D. Kendall, E. S. Marshall, C. L. Lill, W.-J. Lee, S. Kolekar, C. M. Buchanan, S. M. F. Jamieson and
P. R. Shepherd, J. Med. Chem., 2011, 54, 7105; (g) F. Jeppsson, S. Eketjall, J. Janson, S. Karlström, S. Gustavsson, L. L. Olsson, A. C. Radesäter, B. Ploeger, G. Cebers, K. Kolmodin, B. M. Swahn, S. von Berg, T. Bueters and J. Fälting, J. Biol. Chem., 2012, 287, 41245.

5 S. Mizuta, I. S. Stenhagen, M. O'Duill, J. Wolstenhulme,
A. K. Kirjavainen, S. J. Forsback, M. Tredwell, G. Sandford, P. R. Moore, M. Huiban, S. K. Luthra, J. Passchier, O. Solin and V. Gouverneur, Org. Lett., 2013, 15, 2648.
6 S. Verhoog, L. Pfeifer, T. Khotavivattana, S. Calderwood, T. L. Collier, K. Wheelhouse, M. Tredwell and V. Gouverneur, Synlett, 2016, 27, 25.

7 H. Shi, A. Braun, L. Wang, S. H. Liang, N. Vasdev and T. Ritter, Angew. Chem., Int. Ed., 2016, 55, 10786.

8 G. Yuan, F. Wang, N. A. Stephenson, L. Wang, B. H. Rotstein, N. Vasdev, P. Tang and S. H. Liang, Chem. Commun., 2017, 53, 126.
9 (a) Y. Fujiwara, J. A. Dixon, R. A. Rodriguez, R. D. Baxter, D. D. Dixon, M. R. Collins, D. G. Blackmond and P. S. Baran, J. Am. Chem. Soc., 2012, 134, 1494; (b) T. T. Tung, S. B. Christensen and J. Nielsen, Chem.-Eur. J., 2017, 23, 18125; (c) R. Sakamoto, H. Kashiwagi and K. Maruoka, Org. Lett., 2017, 19, 5126.

10 T. C. Wilson, T. Cailly and V. Gouverneur, Chem. Soc. Rev., 2018, 47, 6990.
11 H. Teare, E. G. Robins, A. Kirjavainen, S. Forsback, G. Sandford, O. Solin, S. K. Luthra and V. Gouverneur, Angew. Chem., Int. Ed., 2010, 49, 6821.
12 X. Huang, W. Liu, J. M. Hooker and J. T. Groves, Angew. Chem., Int. Ed., 2015, 54, 5241.
13 X. Huang, W. Liu, H. Ren, R. Neelamegam, J. M. Hooker and J. T. Groves, J. Am. Chem. Soc., 2014, 136, 6842.

14 W. R. Dolbier, Chem. Rev., 1996, 96, 1557.
15 Source: http://www.ich.org/products/guidelines/quality/ article/quality-guidelines.html, accessed on 20/09/18.
16 Our attempts to assemble one-pot the aryl boron reagent, ethyl bromofluoroacetate and $\left[{ }^{18} \mathrm{~F}\right] f l u o r i d e$ were not fruitful. Details in ESI. $\dagger$
17 See the ESI. $\dagger$
18 Y. Wu, H.-R. Zhang, Y.-X. Cao, Q. Lan and X.-S. Wang, Org. Lett., 2016, 18, 5564.
19 C. Guo, X. Yue and F. L. Qing, Synthesis, 2010, 11, 1837.
20 Y. M. Su, G. S. Feng, Z. Y. Wang, Q. Lan and X. S. Wang, Angew. Chem., Int. Ed., 2015, 54, 6003.
21 T. Xia, L. He, Y. A. Liu, J. F. Hartwig and X. Liao, Org. Lett., 2017, 19, 2610.
22 A. Fahandej-Sadi and R. J. Lundgren, Synlett, 2017, 28, 2886.
23 All radiochemical yields (RCYs) are decay corrected.
24 T. Khotavivattana, S. Verhoog, M. Tredwell, L. Pfeifer, S. Calderwood, K. Wheelhouse, T. L. Collier and V. Gouverneur, Angew. Chem., Int. Ed., 2015, 54, 9991.


[^0]:    ${ }^{a}$ Chemistry Research Laboratory, Department of Chemistry, Oxford University, OX1 3TA Oxford, UK. E-mail: veronique.gouverneur@chem.ox.ac.uk; Tel: +44 (0)1865 285002
    ${ }^{b}$ Pfizer Inc., Medicine Design, Eastern Point Road, Groton, Connecticut 06340, and 1 Portland Street, Cambridge, Massachusetts 02139, USA
    ${ }^{c}$ Global Chemistry, UCB New Medicines, UCB Biopharma Sprl, 1420 Braine-L'Alleud, Belgium
    $\dagger$ Electronic supplementary information (ESI) available. See DOI: 10.1039/c8sc05096a

