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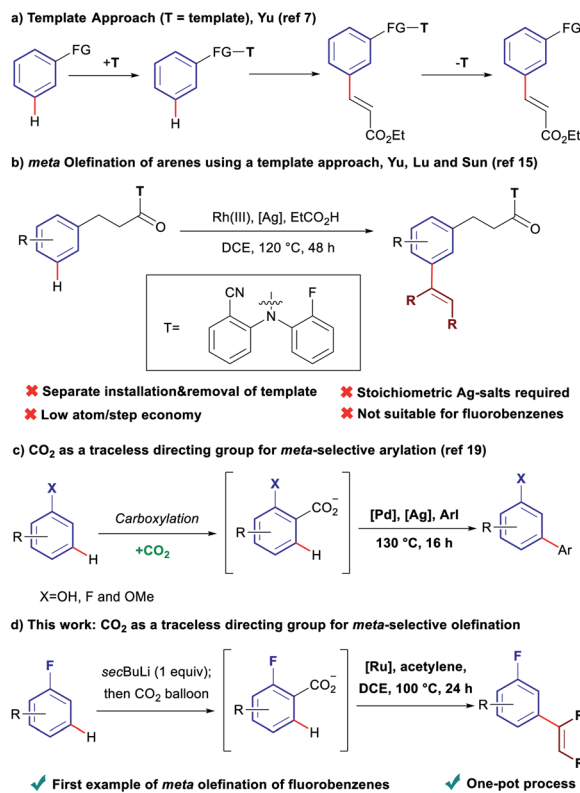
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## Introduction

Over the last decades, direct C–H bond activation has emerged as a powerful tool providing a wide variety of novel disconnections simplifying access to, and accelerating the synthesis of complex molecules.<sup>1</sup> Aryl olefins are synthetically important motifs, as useful intermediates in synthesis,<sup>2</sup> and also due to their widespread presence in bioactive molecules and pharmaceuticals.<sup>3</sup> Consequently, large efforts have been devoted to the development of C–H olefination methodologies.<sup>4</sup> While a large number of strategies have been developed for the olefination of C–H bonds *ortho* to a directing group,<sup>5</sup> comparatively few exist for those in *meta* positions.<sup>6</sup> To date, the only available ‘direct’ strategy for *meta*-olefination, involves the use of the U-shaped directing groups pioneered by the group of Yu (Scheme 1a).<sup>7</sup> This approach has been used to perform *meta*-olefinations on derivatives of benzyl alcohols,<sup>8</sup> *N*-methyl anilines,<sup>9</sup> phenyl acetic acids,<sup>10</sup> benzyl sulfonyl esters,<sup>11</sup> benzoic acids,<sup>12</sup> aromatic carbonyls<sup>13</sup> and aryl boronic acids,<sup>14</sup> using alkenes as coupling partners. The major drawback of this approach arises from the need to install the large U-shaped directing group, covalently bound, and its subsequent removal after the C–H olefination, as separate synthetic steps. In addition, stoichiometric toxic Ag(I)-salts are required as terminal oxidants in these oxidative couplings. A recent report has expanded the applicability of this strategy to the Rh(III)-catalysed *meta*-olefination of hydrocinnamic acid derivatives using alkynes as coupling partners (Scheme 1b).<sup>15</sup> However, despite it being a redox neutral process, it still requires over

three equivalents of Ag(I)-salts as an additive. Furthermore, in addition to the main *meta*-olefination product, 5–10% of the sometimes difficult to separate *ortho* and *para* olefination products are typically obtained. Additionally, the U-shaped directing group strategy is only applicable to aromatics containing a group that can be easily derivatised.



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Scheme 1 Comparison of the template approach and the traceless directing group relay strategy for the *meta* olefination of arenes.



Fluoroarenes are recurring structural motifs in pharmaceuticals, agrochemicals, organic materials and other biologically relevant compounds.<sup>16</sup> Approximately 30% of pharmaceuticals and 40% of agrochemicals currently contain at least one fluorine atom, usually at the aromatic ring.<sup>16</sup> Thus, the direct C–H functionalisation of monofluorobenzenes can provide straightforward access to valuable materials (Fig. 1).<sup>3b–d</sup> While monofluoroarenes generally present low reactivity towards direct C–H olefination, a number of examples have been reported that use the arene as cosolvent to achieve *ortho*, *para*-selective olefination.<sup>17</sup> Some pioneering methods for direct olefination using the fluoroarene as the limiting reagent have been reported by Yu *et al.*, but mixtures of *ortho*, *meta* and *para* substitution are obtained.<sup>18</sup> However, *meta*-selective olefination has never been achieved. Importantly, the U-shaped directing group strategy cannot be applied to this class of substrates. We have previously shown that CO<sub>2</sub> can be used as a traceless directing group for the *meta*-selective arylation of phenols, fluorobenzenes and anisoles (Scheme 1c).<sup>19</sup> The process relies on the easy carboxylation of these aromatic substrates, to install a temporary carboxylate directing group. The carboxylate can then direct the arylation before it is cleaved, in a tandem process, thus allowing a one-pot *meta*-arylation to proceed. However, the CO<sub>2</sub> traceless directing group approach has never been demonstrated on any other type of functionalisation. Herein we report the first example of a *meta*-olefination of fluorobenzenes (Scheme 1d). This ruthenium-catalysed process involves the *in situ* installation and removal of a carboxylate, from CO<sub>2</sub>, uses alkynes as coupling partners and avoids the need for stoichiometric use of Ag(I)-salts.

## Results and discussion

We have previously developed an optimised protocol for the lithiation/carboxylation of fluoroarenes suitable for combination in a one-pot process with a Pd-catalysed tandem arylation/protodecarboxylation, leading to the *meta*-arylation of fluoroarenes.<sup>19c</sup> In 2016, three methods for the Ru-catalysed tandem *ortho*-olefination/protodecarboxylation of benzoic acids by hydroarylation of alkynes were reported by Hartwig and Zhao,<sup>20</sup> Ackermann<sup>21</sup> and Gooßen.<sup>22</sup> Miura and co-workers have also

**Table 1** Optimisation of catalyst. Yields were determined by <sup>19</sup>F NMR analysis using 1-bromo-4-fluorobenzene as an internal standard

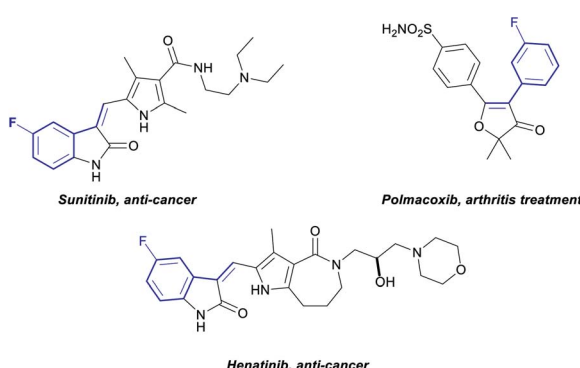
			[Ru] (10 mol %), LiOAc (2.0 equiv), AcOH (3.0 equiv) DCE, 100 °C, 24 h, under N <sub>2</sub>	
	1a	2a	2.0 equiv	3aa
Entry	[Ru]			3aa (yield%)
1 <sup>a</sup>	Ru( <i>p</i> -cymene)(CO <sub>2</sub> Mes) <sub>2</sub>			81
2 <sup>b</sup>	Ru( <i>p</i> -cymene)(CO <sub>2</sub> Mes) <sub>2</sub>			60
3	Ru( <i>p</i> -cymene)(CO <sub>2</sub> Mes) <sub>2</sub>			86
4	[Ru( <i>t</i> BuCN) <sub>6</sub> ][BF <sub>4</sub> ] <sub>2</sub>			0
5	[Ru(C <sub>6</sub> H <sub>6</sub> )(OPiv) <sub>2</sub> ]			25
6	[Ru(C <sub>6</sub> Me <sub>6</sub> )(OAc) <sub>2</sub> ]			93

<sup>a</sup> Without LiOAc and AcOH. <sup>b</sup> Without AcOH.

reported numerous methods for the *ortho*-olefination/protodecarboxylation of benzoic acids using acrylates and styrenes as coupling partners.<sup>23</sup> We envisaged these methods could be ideally adapted to operate in combination with the directed *ortho*-metalation/carboxylation approach to furnish the desired *meta*-olefination of fluorarenes.

We started our investigation by probing the decarboxylative olefination of the fluorotoluic acid **1a** with diphenyl acetylene (**2a**), using Ackermann's protocol (Table 1, entry 1).<sup>21</sup> To evaluate the effect of the installation of the carboxylic acid using *ortho*-lithiation during the desired one-pot process we tested the addition of 2 equiv. of LiOAc (entry 2), revealing a significant negative effect in reactivity. Gratifyingly, addition of 3 equiv. of AcOH efficiently reversed the effect of the presence of the Li-salt (entry 3), providing a method to ensure compatibility of the protocol with the carboxylation step. In previous work on Ru-catalysed *ortho*-arylation of polyfluorobenzenes we observed an inhibitory effect of coordinated *p*-cymene, leading to the development of the arene-free Ru-precatalyst [Ru(*t*BuCN)<sub>6</sub>][BF<sub>4</sub>]<sub>2</sub>.<sup>24</sup> However, the use of this catalyst in the olefination reaction led to no product formation (entry 4), suggesting the  $\eta^6$ -coordinated arene is essential for reactivity towards olefination. Accordingly, the weaker coordinating benzene-complex led to poor reactivity (entry 5), whereas the highly coordinating C<sub>6</sub>Me<sub>6</sub>-bearing Ru complex gave an improved yield, an effect which has previously been observed by Gooßen.<sup>25</sup>

We then moved to optimize the full one-pot protocol, starting from *ortho*-fluorotoluene (**4a**, Table 2). Carboxylation of the fluoroarene was observed to occur in nearly quantitative conversion using *sec*BuLi at –78 °C for 30 min, followed by quenching with CO<sub>2</sub>. Subsequent addition to the same flask of AcOH (3 equiv.), alkyne **2a** and 5 mol% Ru(C<sub>6</sub>Me<sub>6</sub>)(OAc)<sub>2</sub> in DCE led to the formation of the *meta*-olefinated product **3aa** in an excellent 85% yield (entry 1). The use of 4 equiv. or 5 equiv. of AcOH led to reduced yields (entries 2 and 3). Examination of other organic acids also led to lower yields (entries 4–6),



**Fig. 1** Commercially available pharmaceuticals containing a *meta*-alkenyl fluoroarene.

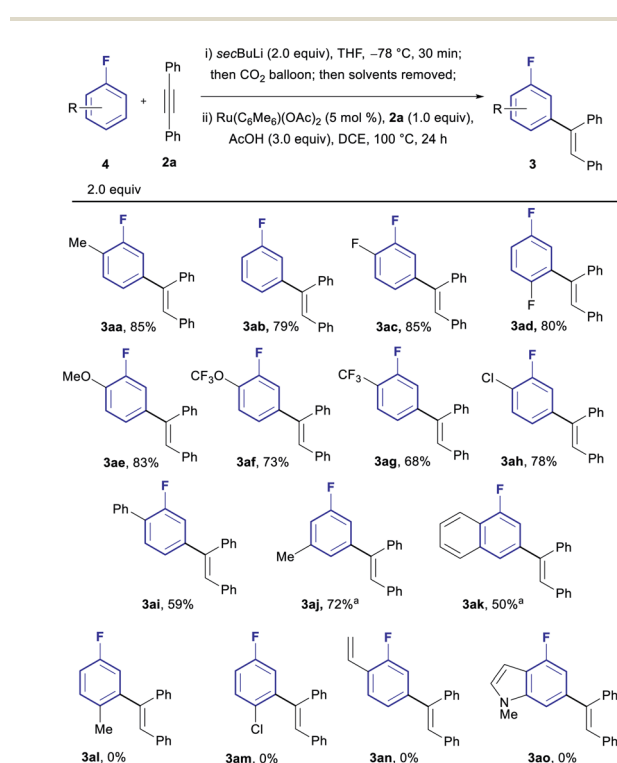


**Table 2** Optimisation of one-pot protocol. Yields were determined by  $^{19}\text{F}$  NMR analysis using 1-bromo-4-fluorobenzene as an internal standard

Entry	Acid (equiv.)	<b>4aa</b> (yield%)
		2.0 equiv
1	AcOH (3)	85
2	AcOH (4)	75
3	AcOH (5)	72
4	PivOH (3)	1
5	<i>i</i> BuCOOH (3)	71
6	TFA (3)	5

revealing AcOH as the optimal acid to facilitate this one-pot process.

With the optimised conditions in hand, we investigated the generality of the process with regards to the fluoroarene core (Scheme 2). Substitution patterns in *ortho*, *meta* or *para* positions were all tolerated, albeit only the relatively small F-atom was compatible in *para* (**3ad**). When larger groups were installed in the *para* position such as Me and Cl, no reactivity could be observed (**3al** and **3am**). Furthermore, the reaction is in all cases completely selective towards mono-olefination and towards the *meta* position, with no traces of neither

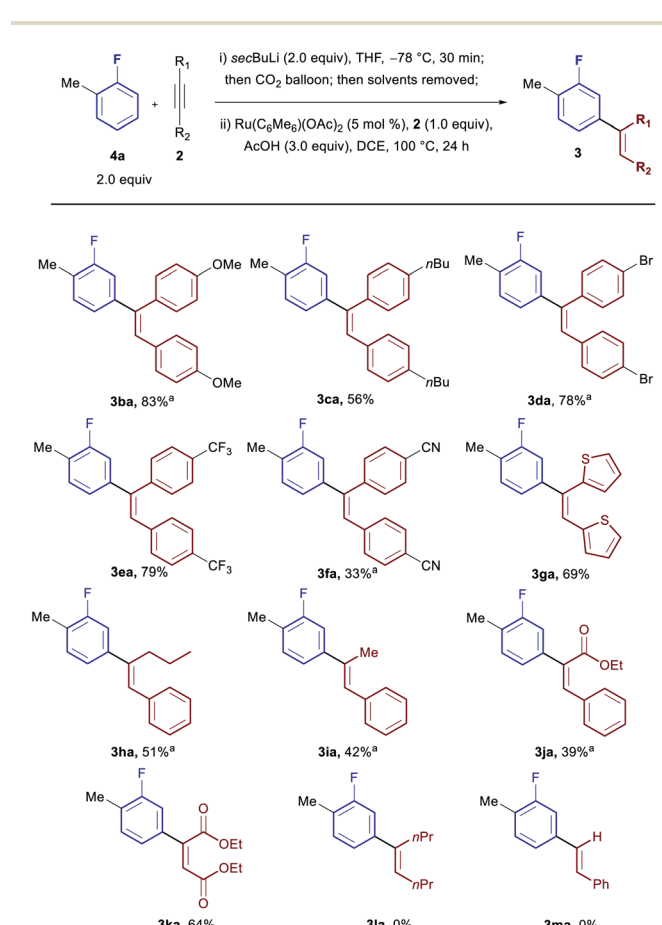


**Scheme 2** Scope in fluoroarene core. <sup>a</sup>10 mol% catalyst used.

bisolefination nor other regioisomers observed by NMR and GCMS analysis of the reactions, even for simple fluorobenzene (**3ab**). Both electron withdrawing groups (**3ac**, **3ad**, **3ag** and **3ah**) and donating groups (**3aa**, **3ae**, **3af** and **3aj**) were compatible with the procedure. Chloroarenes (**3ah**) were also tolerated with no traces of de-halogenated products. Biaryl and naphthyl-based aromatic systems were also suitable substrates (**3ai** and **3ak**).

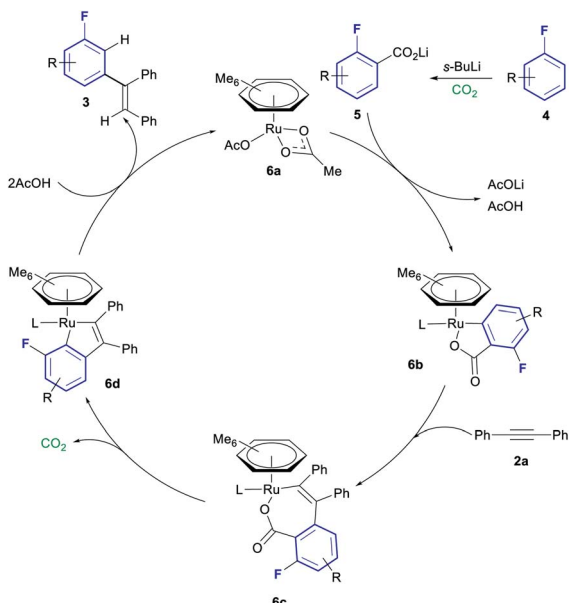
Subsequently we investigated the scope with respect to the alkyne coupling partner (Scheme 3). Both electron donating (**3ba** and **3ca**) and electron withdrawing groups (**3da**, **3ea** and **3fa**) were reactive giving excellent yields. Heterocyclic moieties were also tolerated (**3ga**). While bisalkyl acetylenes were incompatible with the procedure (**3la**), unsymmetrical alkyl, aryl-acetylenes led to completely regioselective addition at the carbon adjacent to the alkyl group (**3ha** and **3ia**). Diesters and unsymmetrical ester, aryl-acetylenes were also tolerated offering a handle for further functionalisation (**3ja** and **3ka**) with ethyl phenyl propiolate preferentially forming the  $\alpha$ -aryl ester (**3ja**). No product was observed when terminal acetylenes were used (**3ma**).

This new *meta*-olefination methodology can be easily scaled up with, for example, **3aa** being formed in 70% yield (1.10 g) without any changes to the protocol.



**Scheme 3** Scope in acetylene. <sup>a</sup>10 mol% catalyst used.





**Scheme 4** Plausible mechanism for the Ru catalysed *meta* olefination of fluoroarenes.

A plausible mechanism for this transformation is shown in Scheme 4, based on the mechanistic studies performed by Hartwig and Zhao.<sup>20</sup> *ortho*-Lithiation and carboxylation of fluoroarene 4 affords lithium benzoate 5. *ortho*-C–H activation of lithium benzoate 5 with ruthenium complex 6a affords cyclo-metallated complex 6b. Insertion of alkyne 2a into the Ru–C of 6b forms complex 6c, which can in turn decarboxylate to form the 5-membered metallocycle in complex 6d. Protonation of this complex with 2 equiv. of AcOH liberates the final product 3 and reforms complex 6a, thus closing the catalytic cycle.

## Conclusions

In conclusion, we have developed the first example of a methodology for the *meta*-selective olefination of fluoroarenes. The natural *ortho*, *para*-reactivity of this class of substrates has been overcome by employing CO<sub>2</sub> as a traceless directing group, that can be installed, used to control reactivity and then seamlessly removed in a one-pot process. Good to excellent yields can be obtained with a variety of functional groups and substitution patterns in both fluoroarene and alkyne, and in all cases complete *meta*-regioselectivity is observed.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

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