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Acid-mediated decarboxylative C–H coupling between arenes and O-allyl carbamates†

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Treatment of O-allyl N-tosyl carbamates with aromatic compounds in the presence of $Cu(OTf)_{2}$ or TMSOTf as promoters affords N-substituted 1-arylpropan-2-amines, 1,2-diarylpropanes, 1,1-diarylpropanes, or indanes, depending on the nature of the promoter and of the aryl substrates. A full mechanistic rational allowing appreciation of the outcome of these novel C–H based cascades is proposed. An initial acid promoted decarboxylative/deamidative Friedel–Crafts allylation takes place. After protonation of the allylated arene, evolution of the resulting cation may follow different paths depending on the nature of the arene partner and of the allyl moiety in the carbamate.

Introduction

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In the last century, carbocation-mediated or -catalysed reactions have revolutionised the history of organic chemistry, giving rise to a set of fundamental reactions such as C–C bond formations, eliminations, and rearrangements. 1 In particular, the Friedel–Crafts (FC) alkylation is one of the most powerful organic reactions that allows the selective C–H functionalization of aromatics.² Although the FC alkylation has long been studied in many variations, its implementation in cascade reactivities that allow the multiple functionalizations of unsaturated substrates represents an extraordinary way to discover still unexplored reactivity patterns.³ **RESEARCH ARTICLE**
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In the frame of our long-term study dedicated to carbamates as precursors for metal-catalysed cyclisations, 4 we investigated in particular the behaviour of O-allyl N-tosyl carbamates. These compounds have been shown to undergo exo-⁵ or endo-trig⁶ cyclisations involving further functionalisation of the carbon double bond. Moreover, allyl carbamates can also allow decarboxylative $O \rightarrow N$ allylic rearrangement affording selectively anti-Markovnikov hydroamination products.⁷ In this work, we show that O-allyl carbamates behave as C3 1,2-, 1,1-, and 1,3-dication equivalents, allowing the generation of 1-aryl-

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propan-2-amines, 1,2- and 1,1-diarylpropanes, as well as indane structures, in the presence of $Cu(OTf)_2$ or TMSOTf as acid-promoters (Scheme 1).

Results and discussion

We started our study reacting carbamate 1a in mesitylene, as aromatic reaction partner and solvent, in the presence of copper $\left(\Pi\right)$ triflate as the promoter (Table 1, entry 1). After 6 hours at 100 °C, the reaction provided a mixture of the arylated N-tosylamide 2 (36% isolated), together with a big amount of $TsNH₂$ (4) arising from the degradation of 1a. Unchanged starting material was found working at lower temperature, and using a catalytic amount of $Cu(OTf)_2$ (entries 2 and 3). Although using an excess of copper salt at 130 °C led to an increased yield of 2, the formation of 4 could not be

Scheme 1 Different reactivities in the acid promoted coupling between arenes and O-allyl N-tosyl carbamates observed in the present study (bold bonds refer to forming bonds).

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Table 1 Optimization of reaction conditions^a

Table 1 Optimization of reaction conditions ^a Me					of it (4.0 equiv.) gladly brought about the selective formation of 3 in 89% yield (entry 15). A similar result was also obtained			
	promoter mesitylene temp., time 1a	ŅHTs `Me Me Me 2	Me Me Me Me Me 3	$+$ TsNH ₂ 4		using DCE as solvent (entry 16). hydrocarbons (Table 2).	Among the range of branched amines, N-substituted phen- ethylamine derivatives have been widely studied in recent years for their value in organic, bioorganic, and medicinal chemistry. ⁸ We thus set out to test the decarboxylative aryla tion/hydroamination described above with other aromatic	
Entry	Promoter (equiv.)	Temp. $({}^{\circ}C)$	Time (h)	Product(s) $(\%$ yield) ^b			Accordingly, reacting allyl carbamate $1a$ in p -xylene under the conditions of Table 1, entry 5 $\left[\text{Cu(OTf)}_{2}\right]$ (4 equiv.), chloro benzene ⁹ /H ₂ O (96/4 v/v), 130 °C, 4.0 h] afforded the corres	
$\mathbf{1}$ $\mathbf{2}$ 3 $\overline{4}$ $\sqrt{5}$ $\,$ 6 $\,$ $\overline{7}$ $\, 8$ 9 10 11 12 13 14 15	$Cu(OTf)_{2}(1.0)$ $Cu(OTf)_{2}(1.0)$ $Cu(OTf)_{2}(0.1)$ $Cu(OTf)_{2}(4.0)$ $Cu(OTf)_{2}(4.0)^{c}$ PTSA (4.0) $H_2SO_4(1.0)$ AgOTf (4.0) TfOH(0.05) TfOH $(0.05)^e$ $TfOH (0.05)^J$ $BF_3 \cdot Et_2O(1.0)$ $BF_3 \cdot Et_2O(4.0)$ TMSOT $f(0.05)$ TMSOT $f(4.0)$	100 50 100 130 130 130 ^d 130 130 130 130 130 100 ^d 130 ^d 130 130	6.0 24 24 3.0 4.0 1.5 4.0 1.5 4.0 4.0 4.0 0.5 1.5 3.0 3.0	$2(36) + 4(43)$ S.M. S.M. $2(85) + 4(11)$ $2(77) + 4(22)$ $2(59) + 4(18)$ $2(25) + 4(14)$ $2(25) + 4(31)$ $2(69) + 4(26)$ $2(58) + 4(41)$ degr. products $2(21) + 3(63)$ $2(15) + 3(72)$ $2(23) + 4(41)$ 3(89)			ponding arylated 2-tosylaminopropane 5 as the sole product in good yield (Table 2, entry 1). The reaction carried out in o -xylene, m -xylene, or toluene gave the corresponding aryla tion/hydroamination products as mixtures of two regioisomers $6a/6b$, $7a/7b$, and $8a/8b$ (entries 2-4). The less electron-rich benzene also provided the expected arylation/hydroamination product 9^{10} in acceptable yield (entry 5). Finally, the heavily alkylated arenes durene and 1,3,5-triethylbenzene gave the	
16	TMSOTf $(4.0)^g$ ^{<i>a</i>} Reaction conditions: 1a (1.0 equiv.), mesitylene (0.25 M), oil bath as heat source. ^b Isolated yields. ^c Chlorobenzene/H ₂ O as solvent (96/4 v/v) (0.4 M) with mesitylene (5.0 equiv.). ^d MW irradiation at 300 W. e Chlorobenzene as the solvent (0.25 M) with mesitylene (5.0 equiv.).	80	4.0	3(79)			Table 2 Synthesis of the N-substituted phenethylamines ^a $Cu(OTf)_{2}$ (4.0 equiv.) aryl hydrocarbon chlorobenzene/H ₂ O (5 equiv.) 130 °C, 4 h	NHTs `Me
	f DMF as the solvent (0.25 M) with mesitylene (5.0 equiv.). g DCE (0.25 M) with mesitylene (5.0 equiv.).				Entry	Aryl hydrocarbon	Product(s) $(\%$ yield) ^b	
					$\mathbf{1}^c$	p -Xylene	Me Me	
	avoided (entry 4). Carrying out the reaction in chlorobenzene in the presence of mesitylene (5.0 equiv.) and H_2O (96/4 v/v) allowed to obtain 2 and 4 in 77% and 22% yields, respectively				2	o -Xylene	NHTs Ńе 5 (79%) Me	

 a Reaction conditions: 1a (1.0 equiv.), mesitylene (0.25 M), oil bath as heat source. $\frac{b}{c}$ Isolated yields. $\frac{c}{c}$ Chlorobenzene/H₂O as solvent (96/4 v/v) (0.4 M) with mesitylene (5.0 equiv.). ^d MW irradiation at 300 W. eChlorobenzene as the solvent (0.25 M) with mesitylene (5.0 equiv.). f DMF as the solvent (0.25 M) with mesitylene (5.0 equiv.). g DCE (0.25 M) with mesitylene (5.0 equiv.).

avoided (entry 4). Carrying out the reaction in chlorobenzene in the presence of mesitylene (5.0 equiv.) and $H₂O$ (96/4 v/v) allowed to obtain 2 and 4 in 77% and 22% yields, respectively (entry 5). Although this protocol does not represent an improvement in terms of yields, it shows that it is possible to work in the presence of a solvent other than the aromatic reaction partner itself.

Different promoters were then tested, using either conventional heating or microwave irradiation. However, p-toluenesulfonic acid, H_2SO_4 and silver(I) triflate behaved analogously to $Cu(OTf)_2$ (Table 1, entries 6–8). Assuming that the above described reaction conditions involved the in situ generation of TfOH, we also tested a catalytic amount of this acid in different solvents (entries 9–11). Working in mesitylene or in chlorobenzene with 5.0 equivalents of mesitylene, the recovery of $TsNH₂$ was still high, whereas the use of DMF gave only intractable degradation products. The use of $F_3B \cdot OEt_2$ disclosed a new reactivity involving the double arylation to the allyl moiety. Indeed, treatment of 1a with a stoichiometric or an excess amount of F_3B ·OEt₂ under microwave irradiation furnished a mixture of 2 and the 1,2-diarylpropane 3 (entries 12 and 13). Switch to TMSOTf as the acid promoter was then considered. While the use of a catalytic amount of this promoter afforded a mixture of 2 and 4 (entry 14), an excess amount

Table 2 Synthesis of the N -substituted phenethylamines $e^{\frac{1}{2}}$

^a Reaction conditions: 1a (1.0 equiv.), Cu(OTf)₂ (4.0 equiv.), aryl hydrocarbon (5.0 equiv.), chlorobenzene/H₂O as solvent (96/4 v/v) (0.4 M), 130 °C, 4 h. δ Isolated yields. Isomeric ratios calculated from the H-NMR of the crude reaction mixture. ^c Scale up: performing the reaction with 5 mmol of 1a at 130 °C, after 6 h 5 was obtained in 71% yield.

corresponding cascade products 10 and 11 in good yields (entries 6 and 7).

We propose for the above double coupling reactions the following mechanism, shown in the case of carbamate 1a (Scheme 2). First, we assume that triflic acid is formed in situ from Cu(OTf)₂ (or TMSOTf) and water.¹¹ Following protonation at the carbonyl oxygen atom of the carbamate function generates the activated O-allyl carbamate I, which undergoes a decarboxylative/deamidative FC alkylation by attack of arene to give intermediate III passing through an allyl carbenium ion II. Subsequent Markovnikov protonation of III generates the allyl carbenium ion IV. At this point, the nature of the additive directs the reaction path. In the presence of copper triflate, the extruded N-tosylamide can coordinate the metal, generating the amido copper-complex V, which selectively attacks the carbenium ion IV, affording the hydroamination product $2.^{12}$ Alternatively, when TMSOTf is used, intermediate IV undergoes a second FC allylation, providing the 1,2-diarylpropane 3. Finally, in the presence of excess triflic acid product 2 suffers a deamidative substitution by mesitylene via an incipient or discrete carbenium ion, to afford the diarylated product 3. Organic Chemistry Frontiers

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Corollary experiments using the O-allyl N-4-chlorophenyl carbamate 1b in place of 1a were carried out next (Scheme 3). Running the reaction in mesitylene/ H_2O (98/2 v/v) in the presence of $Cu(OTf)_2$ as the promoter gave the diarylated product 3, and not the corresponding aniline derivative. However, carrying out the same reaction in the presence of exogenous

H2NTs (2.0 equiv.) afforded a mixture of the arylated/hydroaminated derivative 2 and the diarylated derivative 3. These results corroborate the above proposed mechanism and show that tosylamine is a competent nitrogen nucleophile to trap the carbenium ion **IV** when $Cu(OTf)_2$ is used as the promoter.

The scope of the reaction was evaluated next, keeping the promoting system $Cu(OTf)_2/H_2O$ in chlorobenzene (Table 3). On the one hand, reacting O-allyl N-2-nosylcarbamate 1c with 5 equiv. of mesitylene (entry 1), and carbamate 1a with mesityl bromide (entry 2), 4-methylanisole (entry 3), 4-bromoanisole (entry 4) gave the corresponding hydroaminated products 12–15 in fair to good yields. On the other hand, strongly activated arenes such as 1,3,5-trimethoxybenzene gave the diarylated product 16 as the only product (entry 5).

To have a better knowledge of the behaviour of this C–H cascade as a function of the adopted reaction protocol and the nature of the engaged reaction partners, other tests were undertaken. Accordingly, the C–H coupling between variously substituted N-tosyl carbamates in the presence of the Cu $(OTf)₂/H₂O$ system and 5.0 equivalents of arene was next tested (Scheme 4). In the event, each isomeric carbamate 1d–f converged toward the same corresponding arylation/hydroamination product, namely 17 when reacted with mesitylene and 18 when reacted with *p*-xylene.

Such a reactivity suggests the convergence of 1d, 1e and 1f toward the common allylic carbenium ion VI, which, after the

Scheme 2 Proposed mechanisms for the acid-mediated decarboxylative C–H coupling between arenes and O-allyl carbamates.

Scheme 3 Cu(OTf)₂/H₂O mediated decarboxylative C-H couplings between mesitylene and O-allyl N-4-chlorophenyl carbamate. Reaction condition involves the use of 100 μ L of H₂O for 1 mmol of 1b.

Table 3 Other C-H couplings with heterosubstituted arenes⁸

 a Reaction conditions: N-protected O-allyl carbamate (1.0 equiv.), Cu (OTf)₂ (4.0 equiv.), arene (5.0 equiv.), chlorobenzene/H₂O as solvent (96/4 v/v) (0.4 M), 130 °C, 4 h. ^b Isolated yields. ^c The reaction was carried out in mesitylene as solvent (0.25 M).

Scheme 4 Reaction with Cu(OTf)₂ with substituents on the allylic chain of the N-tosyl carbamates.

Scheme 5 Proposed mechanism for the Cu(OTf)₂ promoted transformation of 1d–f into 17 and 18.

FC reaction, undergoes a regioselective Cu-promoted hydroamination (Scheme 5).

The coupling of N-tosyl carbamates 1a, and 1d–e was also tested using TMSOTf as acid promoter in DCE at 80 °C (Table 4). In this case, reaction of $1a$ with *p*-xylene gave an inseparable mixture made of the 1,2-diarylated product 19, the 1,1-diarylated product 20, and indane 21 (entry 1). Carbamates 1d and 1e converged again toward a mixture of the 1,1-diarylated product 22 and the indane 23 when reacted with p -xylene (entry 2), and toward a mixture of the 1,2-diarylated product 24 and 1,3-diarylated product 25 when reacted with mesitylene (entry 3).

The reactivity observed with the above experiments deserves further remarks. First, the formation of 19, 20, and 21 (Table 4, entry 1) can be rationalized as follows (Scheme 6, top part). After the first FC allylation of p-xylene, Markovnikov protonation and interception of the resulting carbenium ion VII by a second molecule of p-xylene generates 19. However, p-xylene – less nucleophilic than mesitylene – allows a competitive VII-to-VIII 1,2-hydride shift that takes place before the second FC reaction, generating 20. At this point, the benzylic carbocation VIII and the styrene IX could react together through a formal $\begin{bmatrix} 3 & + & 2 \end{bmatrix}$ cycloaddition giving the Wheland intermediate **X** which can evolve into 21.¹³ Pescando Article.

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The outcome of entry 3 in Table 4 needs further observations, too. Indeed, in this case, the protonation after the first

^a Reaction conditions: 1a (1.0 equiv.), TMSOTf (4.0 equiv.), arene (5.0 equiv.), DCE (0.4 M), 80 °C, 4 h. ^b Isolated yields. Isomeric ratios calculated from the ¹H-NMR of the crude reaction mixture. c_1 **19** + 20 isomeric ratio: 1/1. ^{*d*} The relative configuration of indane structures 21 and 23 was determined by NOESY experiment.

Scheme 6 Proposed mechanisms associated to the reactions of Table 5. Top part: entry 1, from 1a to 19, 20, and 21. Bottom part: entry 3, from 1d or 1e to 24 and 25.

Table 5 Reaction with TMSOTf with α, α -dimethyl substituted O-allyl N -tosyl carbamates a

 a Reaction conditions: 1a (1.0 equiv.), TMSOTf (4.0 equiv.), arene (5.0 equiv.), DCE (0.4 M), 80 °C, 4 h. b Isolated yields. Isomeric ratios calculated from the 1 H-NMR of the crude reaction mixture. c 1 : 1 isomeric ratio. a^d 28 + 29 in 1 : 1 ratio.

FC reaction is non-regioselective, and the very nucleophilic mesitylene intercepts the two resulting carbenium ions to give a mixture of 24 and 25. In this case, the very reactive arene does not let the time for a homobenzylic-to-benzylic cation 1,2-hydride shift (Scheme 6, bottom part).

Scheme 7 Proposed mechanisms for the generation of the indane structures 26, 27a and 27b and of hydrindacene 29 from carbamate 1g.

The α , α -dimethyl substituted O-allyl N-tosyl carbamate 1g was investigated next (Table 5). Thus, treatment of this carbamate with durene under the usual TMSOTf conditions gave indane 26 in 87% yield (entry 1). Analogous treatment with mesitylene afforded an inseparable mixture of the two isomeric indanes 27a and 27b in 89% yield (entry 2), while reaction with p -xylene gave an inseparable 1:1 mixture of the indane 28 and the hydrindacene 29 in 93% yield (entry 3).

Here, in contrast to the previous cases, a one-to-one coupling between the carbamate and the arene takes place, generating indane structures. This is likely due to the fact that after the first FC allylation on the less substituted allyl terminus $XI₁¹⁴$ the aromatic ring is favourably biased to undergo a second, intramolecular, FC reaction. It should be noted that in the cases of mesitylene and durene, 1,2-methyl migrations on the aryl moiety¹⁵ take place - before or after the first FC reaction – opening the way to the final intramolecular FC reaction (Scheme 7). However, a second carbocation species could attack the highly reactive indane 28 providing the hydrindacene 29. 16

Conclusions

In summary, this work shows for the first time that O-allyl carbamates are ideal C3 dication equivalents. Acidic conditions expected to generate in situ TfOH enable decarboxylative FC/ hydromidation sequences or twofold FC alkylations, most of which proceed in synthetically useful yields. According to the nature of the arene partner and of the allyl moiety in the carbamate, the mechanisms can take different paths, all of them mechanistically justified. These results offer the chemist a rich palette of synthetic opportunities, further enriching the

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domains of cascade reactions in general, and FC, and hydroaminations in particular.

Conflicts of interest

There are no conflicts to declare.

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