




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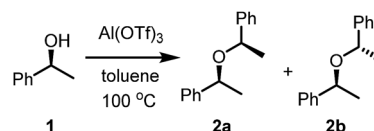
Protecting group-free use of alcohols as carbon electrophiles in atom efficient aluminium triflate-catalysed dehydrative nucleophilic displacement reactions†

Adam Cullen,^a Alfred J. Muller^a and D. Bradley G. Williams ^{*ab}

Benzylic and allylic alcohols are rendered electrophilic without chemical modification by the use of aluminium triflate as catalyst. The reaction succeeds with alcohol, thiol, carbon and nitrogen nucleophiles. When phenols are employed as nucleophiles, *C*-alkylation ensues. An advanced application of the method is demonstrated in the synthesis of 2*H*-chromenes and their N and S analogues.

Introduction

Many of the problems relating to regio- or chemoselective synthesis have not yet been solved.¹ As such, organic synthesis usually requires protecting group strategies to deliver selectivity. In multistep syntheses protection–deprotection strategies are not only critical to the success of the synthesis, they often comprise a significant proportion of the steps involved.¹ Because of these deficiencies, atom economic chemistry remains a touchstone by which the efficiency of synthetic organic chemistry is measured. Alcohols have been used for decades as a source of electrophiles once they have been converted into halides or other leaving groups such as tosylates or triflates.² Alternatively, it is known that propargylic and benzylic alcohols can form stabilised carbocations under appropriate conditions, usually in the presence of aluminium- and gallium triflates, respectively.³ Aluminium triflate has been shown to be a highly active yet selective catalyst for a range of reactions, including some highly atom economic processes where 100% of the atoms are incorporated into the products.^{4–7} Aluminium triflate is believed to exert its function in many cases as a Lewis acid-assisted Brønsted acid.^{8–10} It is also useful as a co-catalyst in the Pd-catalysed hydromethoxycarbonylation reaction, where it takes the place of traditional Brønsted acids.^{6,7} Importantly, it tolerates protic conditions and so can be readily recycled by simply extracting it into water and recovered by subsequent removal of the water.¹¹



Scheme 1 Conversion of (*S*)-1-phenylethanol into di(1-phenylethyl) ether.

When investigating epoxide ring-opening reactions,^{12,13} we became interested in the possibility of employing chiral alcohols as nucleophiles by which to potentially desymmetrise the epoxides. During that study, we employed (*S*)-1-phenylethanol (1) as the nucleophile and obtained the *S,R* (*meso*) and *S,S* diastereomers 2 of di(1-phenylethyl) ether as the major products.¹³ The scrambling of the stereogenic centre implied that a benzylic carbocation had formed in the process, under the action of Al(OTf)₃, and could be captured by a nucleophile present in the reaction mixture (Scheme 1). This manuscript details the direct use of benzylic and allylic alcohols as electrophiles in high yielding C–O, C–S, C–N and C–C bond-forming reactions, where water is the by-product, in intramolecular and intermolecular reactions.

Results and discussion

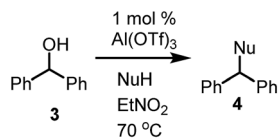
Benzhydrol 3 was subjected to reaction with a range of nucleophiles in the presence of Al(OTf)₃ in nitroethane solvent at 70 °C, affording the corresponding elaborated products in very high yields (Table 1) [nitroethane demonstrates low toxicity to a range of organisms, does not bioaccumulate and readily biodegrades¹⁴]. Notably, the reaction is successful with O, C, N and S nucleophiles, requires no manipulation of the heteroatoms and it can be performed solvent-free without loss of yield or selectivity (Table 1, entries 7 and 11). Furthermore, reactions involving phenols yield *C*-alkylated aromatics as products

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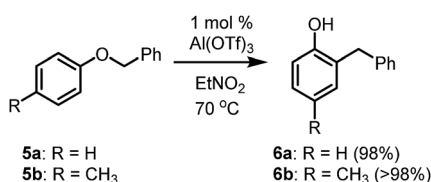
Table 1 Al(OTf)₃-catalysed nucleophilic substitution reactions of benzhydrol with various nucleophiles^a

4a: Nu = OEt; **4b:** Nu = Oallyl; **4c:** Nu = Opropargyl; **4d:** Nu = acetylacetonyl;
4e: Nu = ethyl acetoacetyl; **4f:** Nu = thiophenol; **4g:** Nu = *O*-*l*-menthyl
4h: NuH = phenol, Nu = 4-C₆H₄OH; **4i:** NuH = 2-naphthol, Nu = 2-naphthol-1-yl
4j: NuH = indole, Nu = indol-3-yl
4k: NuH = methyl 2,3-isopropylidene-β-D-ribose, Nu = methyl 2,3-isopropylidene-5-O-β-D-ribose

Entry	Nucleophile	Product	Yield
1	Ethanol	4a	98%
2	Allyl alcohol	4b	94%
3	Propargyl alcohol	4c	>98%
4	Acetyl acetone	4d	96%
5	Ethyl acetoacetate	4e	87%
6	Thiophenol	4f	98%
7 ^a	<i>l</i> -Menthhol	4g	85%
8	Phenol	4h	84%
9	2-Naphthol	4i	98%
10	Indole	4j	80%
11 ^a	Methyl 2,3-isopropylidene-β-D-ribose	4k	68%

^a Solvent free, 40 °C, performed under vacuum (5 mm Hg).

(Table 1, entries 8 and 9). It is possible that the reaction initially yields the analogous *O*-alkylated product which converts into the *C*-alkylated thermodynamic product. To test this hypothesis, analogues **5** (Scheme 2) of the putative benzyl ether intermediates were subjected to reaction with Al(OTf)₃ under identical reaction conditions, affording the *C*-alkylated products in quantitative (>98%) yield. This result supports the notion that a kinetically produced intermediate converts into the thermodynamically preferred *C*-alkylated product. Interestingly, the *ortho* product was the exclusive product for both substrates (Scheme 2) in this reaction. While this is expected for the *p*-cresol-derived ether **5b**, it was somewhat unexpected for the phenol-derived substrate **5a**. In contrast, the product of the reaction of phenol and benzhydrol is the *para* product **4h** (Table 1, entry 8). The difference in outcome may be accounted for by the relative stability of the diphenylmethyl carbocation that forms (compared to benzyl cation), allowing it to migrate to the less hindered *para* position of the phenol ring. This is in contrast to *O*-diphenylmethyl trichloroacetimidate which affords *O*-alkylated aromatics from 2-naphthol and several

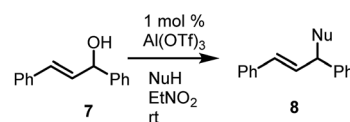
**Scheme 2** Conversion of benzyl ethers into *C*-alkylated products.

phenols.¹⁵ PdCl₂ (10 mol%) as catalyst in dichloroethane also affords *O*-alkylated aromatics from benzhydrol.¹⁶ Usefully, ribose derivative **4k** can be selectively deprotected of the diphenylmethane group in high yield (90%) by a simple reaction involving Al(OTf)₃ and 2 equivalents of EtOH. This compares favourably to PdCl₂-catalysed (10 mol%) deprotection of monosaccharides with EtOH (10 equivalents) in dichloroethane.

trans-1,3-Diphenylprop-2-en-1-ol (**7**) was subjected to a similar set of reactions (Table 2). Once again, high yields of products were obtained and the reaction also tolerated a range of nucleophiles, including O, C, N and S nucleophiles (Table 2). As was the case with benzhydrol, phenols and indole underwent *C*-alkylation while thiophenol underwent *S*-alkylation in otherwise smooth transformations. The reactions were performed at ambient temperature, affording very high yields of products across the range of nucleophiles employed.

Reactions with benzhydrol or *trans*-1,3-diphenylprop-2-en-1-ol did not proceed with unprotected anilines, acetanilide or their *p*-tosylamide analogues. This is consistent with other work¹⁷ with electron-deficient amines, where higher catalyst loadings (1–5 mol%) and (often) microwave heating were required before successful substitution reactions were observed. However, successful *N*-alkylation could be achieved in the present instance when using *p*-toluenesulphonyl amide (Scheme 3), with which the C–N bond formation proceeded quite readily to produce the *N*-alkylated products in quantitative (>98%) yields. Nitroethane remained the solvent of choice.

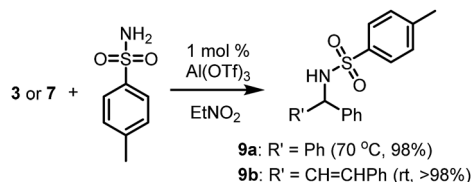
We next turned our attention to the synthesis of chromenes *via* intramolecular cyclisation reactions. The chromene substructure is present in a wide range of physiologically active molecules which demonstrate anti-HIV,¹⁸ antitumour,¹⁹ antibacterial²⁰ and antiprotozoal²¹ activity. Significant efforts have been made towards the synthesis of the *2H*-chromene motif,²² including sugar-derived chiral systems.²³ In addition, there has

Table 2 Al(OTf)₃-catalysed nucleophilic substitution reactions of **7** with various nucleophiles

8a: Nu = OEt; **8b:** Nu = Oallyl; **8c:** Nu = Opropargyl; **8d:** Nu = acetylacetonyl
8e: Nu = ethyl acetoacetyl; **8f:** Nu = thiophenol; **8g:** NuH = phenol, Nu = 4-C₆H₄OH
8h: NuH = 2-naphthol, Nu = 2-naphthol-1-yl; **8i:** NuH = indole, Nu = indol-3-yl

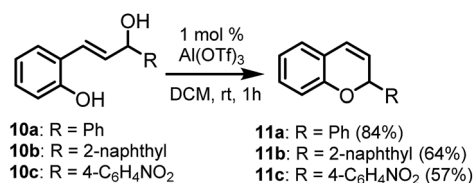
Entry	Nucleophile	Product	Yield
1	Ethanol	8a	85%
2	Allyl alcohol	8b	84%
3	Propargyl alcohol	8c	96%
4	Acetyl acetone	8d	98%
5	Ethyl acetoacetate	8e	98%
6	Thiophenol	8f	86%
7	Phenol	8g	82%
8	2-Naphthol	8h	81%
9	Indole	8i	87%



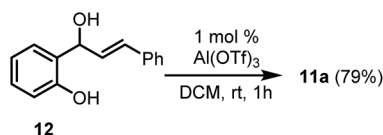


Scheme 3 Reactions of activated alcohols with *p*-toluenesulfonyl amide.

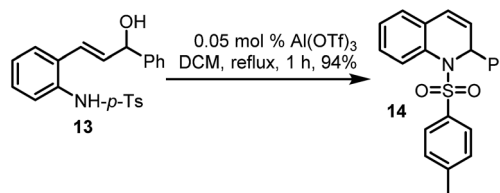
been continuing interest in heteroatom analogues of chromenes such as dihydroquinolines²⁴ and thiochromenes,²⁵ which are also associated with biological activity. To set up the intramolecular cyclisation that would build the chromene motif in our case, it was necessary to synthesise phenylpropenols **10**, which was accomplished in four steps in good yields (see ESI for details†). Propenols **10** were subjected to the action of 1 mole percent Al(OTf)₃ in nitroethane at ambient temperature in an effort to produce the corresponding 2*H*-chromenes (Scheme 4). Upon addition of the Al(OTf)₃ to the solution of the propenols in nitroethane a deep purple colour ensued immediately, presumably the consequence of carbocation formation, and the starting alcohol was fully consumed (by tlc analysis) within a few minutes. Al(OTf)₃ is known to produce carbocations in solution from retinyl acetate, for example, which has been used to probe the likely mechanism by which Al(OTf)₃ functions.⁸ Unfortunately, the propenol reaction produced an intractable mixture of products. To tame the reactivity of the system, the reaction was performed in dichloromethane, which allowed the desired 2*H*-chromenes to be prepared in acceptable to good yields in a dehydrative *endo* cyclisation step (Scheme 4). 2*H*-Chromene **11a** could also be prepared from **12** (Scheme 5; see ESI† for details on the synthesis of **12**), a regioisomer of **10a**, under the same conditions in comparable yield for the *endo*-cyclisation step of its regioisomer of **10a**. This anticipated ability of the reaction to proceed as desired regardless of the position of the leaving OH group imparts significant flexibility in the synthesis protocols that might lead to the functionalised diphenylpropenol substrates. In a previous study, this type of intramolecular cyclisation reaction of phenylpropenols in the



Scheme 4 Synthesis of 2*H*-chromenes in Al(OTf)₃-catalysed cyclisation reactions.



Scheme 5 Synthesis of 2*H*-chromene from an isomeric substrate.



Scheme 6 Synthesis of dihydroquinoline in an Al(OTf)₃-catalysed dehydrative cyclisation.

presence of 5 mole percent of a (*t*-Bu)₂P(2-C₆H₄-Ph)-AuCl/AgOTf catalyst at elevated temperatures has been noted.²⁶ The Au catalyst tolerates some functional groups but fails outright in some instances.

Finally, propenol **13** could also be made to undergo Al(OTf)₃-catalysed dehydrative cyclisation, to afford the desired dihydroquinoline **14** in a pleasing yield of 94% (Scheme 6), using only 0.05 mol% of the catalyst. While the use of DCM is less desirable, the overall high yields on the one hand, and the fact that this method enables alcohols to be used directly in nucleophilic displacement reactions, cannot be overlooked. Further investigations will most likely unveil other benign solvents which are compatible with this chemistry.

Conclusions

Al(OTf)₃ catalyses the direct nucleophilic displacement of OH groups in 'activated' systems with various O, N, S and C nucleophiles. This efficient process allows the ready synthesis of a wide range of product types in very high yields with water as the by-product. Furthermore, the displacement reaction is usefully employed in the synthesis of 2*H*-chromenes and dihydroquinolines in *endo* cyclisation reactions. This facile and high yielding dehydrative procedure is expected to be useful in the synthesis of a wide range of organic materials and biologically active molecules. We are currently investigating the applications of this innovation to other ring structures.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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