Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2018, 9, 5767

Iridium-catalyzed enantioselective direct vinylogous allylic alkylation of coumarins†‡

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Received 7th May 2018 Accepted 3rd June 2018

DOI: 10.1039/c8sc02041h

rsc.li/chemical-science

The first iridium-catalyzed enantioselective vinylogous allylic alkylation of coumarins is presented. Using easily accessible linear allylic carbonates as the allylic electrophile, this reaction installs unfunctionalized allyl groups at the γ -position of 4-methylcoumarins in an exclusively branched-selective manner generally in high yields with an excellent level of enantioselectivity (up to 99 : 1 er).

Introduction

Transition-metal-catalyzed asymmetric allylic substitution (AAS) reactions have emerged as an extremely powerful and versatile method for the synthesis of enantioenriched compounds from easily available starting materials through enantioselective construction of carbon–carbon and carbon–heteroatom bonds. In contrast to initially developed and more commonly used palladium catalysts, iridium-catalyzed AAS reactions enable the synthesis of branched products from unsymmetrical allylic electrophiles through preferential attack of nucleophiles at the more substituted terminus of the π -allyl–Ir intermediate (Scheme 1).

This characteristic of the Ir-catalyzed AAS overcomes the limitations associated with Pd-catalysis with respect to the scope of reaction partners and even allows for the use of achiral (non-prochiral) carbon and heteroatom nucleophiles in reactions with unsymmetrical allylic electrophiles.² Consequently, a wide variety of nucleophiles have been applied³⁻⁵ to highly regioselective (branched-to-linear) and enantioselective allylic alkylation reactions ever since the introduction of Ir-catalysts in 1997.⁶ Despite these developments during the past two decades, the use of vinylogous nucleophiles in Ir-catalyzed asymmetric allylic alkylation (AAA) has received much less attention and only a handful of reports exist (Scheme 2A).⁷⁻⁹



Scheme 1 Transition-metal-catalyzed allylic substitution.

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- \dagger Dedicated to Professor E. J. Corey on the occasion of his 90th birthday.
- ‡ Electronic supplementary information (ESI) available: Experimental details, characterization and analytical data. See DOI: 10.1039/c8sc02041h

In 2014, Hartwig *et al.* reported the first application of a vinylogous nucleophile, namely preformed silyl dienolates, in Ir-catalyzed AAA.⁷ The Jørgensen group combined the concept of dienamine activation with Ir-catalyzed AAA for regio-, diastereo- and enantioselective γ -allylic alkylation of α , β -unsaturated aldehydes.⁸ More recently, the Stoltz group developed

OTroc

Hartwig et al.

OH

Combined with

dienamine catalysis

Jørgensen et al.

MeO₂C

CO₂Me

OCO₂Me

OCO₂Me

Stoltz et al.

(A) Previous reports: Ir-catalyzed enantioselective vinylogous allylic alkylation

(B) Previous reports: enantioselective vinylogous allylic alkylation of coumarins

(C) This work: Ir-catalyzed enantioselective vinylogous allylic alkylation of coumarins

Limitations: structurally & electronically biased allylic electrophiles

Scheme 2 Enantioselective vinvlogous allylic alkylation.

a formal enantioselective $\gamma\text{-allylic}$ alkylation of $\alpha,\beta\text{-unsaturated}$ malonates through a sequential Ir-catalyzed AAA/Cope rearrangement.9

With our own interest in vinylogous nucleophilic reactivity, ¹⁰ we embarked on the development of direct catalytic enantioselective allylic alkylation of other vinylogous nucleophiles.

We became particularly interested in 4-methylcoumarins as the potential vinylogous nucleophile due to the wide abundance of coumarin derivatives in over 1000 natural products and bioactive targets as well as their utility in the dye industry.11 Although this class of nucleophiles received considerable attention since the introduction of 3-cyano-4-methylcoumarins by Xie et al. in 2010,12 enantioselective allylic alkylation of this potentially useful class of vinylogous nucleophiles remained underexplored until recently. Lautens' group developed the first enantioselective γ-allylic alkylation of 4-methylcoumarins in 2016 through a Rh-catalyzed desymmetrizing ring-opening reaction of oxabicycles (Scheme 2B).13 Very recently, our group14 and subsequently Albrecht et al.15 independently developed the first organocatalytic enantioselective γ-allylic alkylation of 3-cyano-4-methylcoumarins using Morita-Baylis-Hillman carbonates as the allylic electrophile (Scheme 2B). While an excellent level of enantioselectivity has been achieved, the success of these reactions is inherently dependent on the type of allylic electrophile - both structurally and electronically, thereby limiting their scope.

Since Ir-catalyzed allylic alkylation reactions are not constrained by such structural and electronic bias on the allylic electrophile, we believed that Ir-catalysis would provide a general strategy for the synthesis of coumarin derivatives that were previously challenging to access.

The purpose of this communication is to disclose the first Ircatalyzed enantioselective vinylogous allylic alkylation of coumarins (Scheme 2C).

Results and discussion

We began our investigation with the optimization of the catalyst and reaction conditions¹⁶ for a model reaction between 3-cyano-4-methylcoumarin 1a and tert-butyl cinnamyl carbonate 2a in dichloroethane (DCE) at 50 °C (Table 1). A combination of [Ir(COD)Cl]2 and Feringa's phosphoramidite ligand L1,17 pioneered by Hartwig and co-workers, 51 was initially tested. In the absence of any external base to activate 1a, the desired γ -allylated product 3aa was formed exclusively as a single regioisomer with a promising enantioselectivity, although in only 22% yield (entry 1). The tert-butoxide anion, generated in situ from the reaction between an Ir-complex and 2a, was anticipated to be the active base in this reaction.18 In line with the observation reported previously for Ir-catalyzed AAS reactions,3-5 the choice of external base was found to have a profound influence on both the reaction efficacy and enantioselectivity. Extensive exploration of various bases (entries 2-7), including inorganic and organic bases, revealed DABCO to be the optimum,19 affording the product in high vield and enantioselectivity (entry 7). A number of ligands for iridium were then tested under the influence of DABCO (entries 8-11). The use of ligand L2,

Table 1 Ligand screening and reaction optimization^a

$$\begin{array}{c} \text{OBoc} \\ \text{Ne} \\ \text{1a} \\ \text{2a} \\ \end{array} \begin{array}{c} \text{Ilr(COD)CI]}_2 \text{ (3 mol \%)} \\ \text{base (1.0 equiv)} \\ \text{solvent (0.33 M)} \\ \text{3aa} \\ \text{Ph} \\ \end{array} \\ \begin{array}{c} \text{Ph} \\ \text{Ne} \\ (S_a, S, S) - \textbf{L1:} \text{ Ar = Ph} \\ (R_a, S, S) - \textbf{L2:} \text{ Ar = Ph} \\ (S_a, S, S) - \textbf{L3:} \text{ Ar = 2-OMeC}_6 H_4 \\ \end{array} \\ \begin{array}{c} \text{Ph} \\ \text{Ne} \\ (S_a, S, S) - \textbf{L4:} \text{ Ar = 1-Np} \\ \end{array} \\ \begin{array}{c} \text{Ph} \\ \text{Ne} \\ (S_a, S, S) - \textbf{L5:} \end{array}$$

Entry	Ligand	Solvent	Base	t [h]	Yield ^b [%]	er ^c
1	L1	DCE	_	48	(22)	92.5 : 7.5
2	L1	DCE	Cs ₂ CO ₃	48	<5	n.d.
3	L1	DCE	DBU	48	<5	n.d.
4	L1	DCE	<i>i</i> -Pr ₂ NEt	48	72	96:4
5	L1	DCE	Et_3N	48	74	97:3
6	L1	DCE	i-Pr ₂ NH	48	76	97:3
7	L1	DCE	DABCO	48	84	97.5:2.5
8	L2	DCE	DABCO	48	<5	n.d.
9	L3	DCE	DABCO	48	11	98:2
10	L4	DCE	DABCO	48	<5	n.d.
11	L5	DCE	DABCO	48	28	92:8
12	L1	THF	DABCO	48	11	96:4
13	L1	$CHCl_3$	DABCO	48	67	97:3
14	L1	CH_2Cl_2	DABCO	36	88 (86)	98:2

 a Reaction conditions: 3 mol% [Ir(COD)Cl]₂, 6 mol% ligand, 0.24 mmol of **1a**, 0.2 mmol of **2a** and 0.2 mmol of base in 0.6 mL solvent. The catalyst was prepared via n-PrNH₂ activation. b Yields were determined by 1 H-NMR spectroscopy with mesitylene as the internal standard. Isolated yields are given in the parentheses. c The enantiomeric ratio (er) was determined by HPLC analysis on a chiral stationary phase; n.d. = not determined. DCE = 1,2-dichloroethane.

a diastereomer of **L1**, failed to catalyze the reaction (entry 8). Ligand **L3**, an *ortho*-methoxy-derivative of **L1**, introduced by Alexakis *et al.*²⁰ and generally known to act as a superior ligand compared to **L1** in many AAS reactions, furnished **3aa** with improved er but in poor yield (entry 9). No improvement in yield or enantioselectivity was observed with **L4** or **L5** (entries 10 and 11), and **L1** remained the ligand of choice. A solvent screening (entries 12–14) at this point revealed dichloromethane as the optimum, providing the product in 86% isolated yield and with 98: 2 er (entry 14).

With the optimum ligand and reaction conditions (Table 1, entry 14) in hand, we chose to explore the scope and limitations of this direct vinylogous allylic alkylation protocol. We were pleased to note that the efficacy displayed by the Ir/L1 combination for the reaction between 1a and 2a under the optimum reaction conditions is indeed a general phenomenon and could be extended to other substrate combinations. As shown in Table 2, 3-cyano-4-methylcoumarin (1a) underwent facile allylic alkylation with an assortment of allylic carbonates (2a–u). Arylsubstituted allylic carbonates with diverse steric and electronic demands on the aryl ring (2a–i) were well tolerated, providing the products (3aa–ai) in high yields with excellent

Table 2 Scope of the enantioselective allylic alkylation with regard to allylic carbonates a

enantioselectivities (Table 2A, entries 1–9). When the substituent was present in either the *meta*- or *para*-position of the phenyl ring, products were formed with similar yields and enantioselectivities. In accordance with previously reported AAS reactions with L1,²¹ the *ortho*-substituent on the phenyl ring led to slow reaction, giving the product with moderate yield and er (entry 10). This adverse effect of *ortho*-substituent is clearly evident from the comparative outcome of the reactions with dichloro-substituted cinnamyl carbonates 2k and 2l (entries 11 and 12). We were also interested in incorporating pharmaceutically relevant heterocycles into our products, and found that allylic carbonates bearing dioxolane (2m), pyridine (2n), furan (2o) and thiophene (2p) provided the products in good to high yields with excellent er (entries 13–16).

The scope of our protocol is not limited to (hetero)aromatic allylic carbonates. As illustrated by the examples in Table 2B, products derived from allylic carbonates containing a linear (2q) and a branched alkyl group (2r) as well as benzyl (2s) were obtained with good enantioselectivities, albeit in moderate

yield. In addition, alkenyl-substituted allylic carbonates (2t and 2u) participated in this reaction and generated the single regioisomeric products (3at and 3au), out of three possible regioisomers, in good yield with good to high er.

The scope of the reaction with respect to other cyanocoumarin derivatives was examined next (Table 3). A number of substituted cyanocoumarins bearing either electron donating (e.g. Me and OMe) or electron withdrawing (e.g. F, Cl, and Br) groups were found to be equally suited under our standard reaction conditions, affording γ -allylated products (3ba-ga) with excellent yields and enantioselectivities. Disubstituted cyanocoumarin 1h also participated in this reaction with equal efficiency.

After successfully demonstrating the scope of the reaction with cyanocoumarins, we wondered whether the reactivity of the coumarin derivative could be retained by replacing the cyano group with other α-substituents. To our delight, when CN (in 1a) was replaced with an amide group (CONH₂), the vinylogous allylic alkylation reaction indeed took place to furnish the corresponding y-allylated product 3ia with a similar level of yield and enantioselectivity (Table 4, entry 2). Not only amide but also related esters (1j and 3k) could be employed as substrates and afforded the products (3ja and 3ka) in high yield and with excellent er (entries 3 and 4). However, α-carboxylatocoumarin (11) failed to react in the desired fashion and instead resulted in 4-methylcoumarin 1m in 52% yield through decarboxylation (entry 5). Similarly, unsubstituted 4-methylcoumarin (1m) itself remained unreacted under our standard reaction conditions even after 48 h (entry 6).

Table 3 Scope of the enantioselective allylic alkylation with respect to coumarins^a

^a Reaction conditions: 3 mol% [Ir(COD)Cl]₂, 6 mol% L1, 0.24 mmol of 1a, 0.2 mmol of 2 and 0.2 mmol of DABCO in 0.6 mL CH₂Cl₂. The catalyst was prepared *via n*-PrNH₂ activation. Yields correspond to the isolated product after chromatographic purification. er was determined by HPLC analysis on a chiral stationary phase.

^a Reaction conditions: 3 mol% $[Ir(COD)Cl]_2$, 6 mol% L1, 0.24 mmol of 1, 0.2 mmol of 2 and 0.2 mmol of DABCO in 0.6 mL CH_2Cl_2 . The catalyst was prepared *via n*-PrNH₂ activation. Yields correspond to the isolated product after chromatographic purification. er was determined by HPLC analysis on a chiral stationary phase.

Table 4 Effect of α -substituent on coumarins in enantioselective allylic alkylation^a

Entry	X	t [h]	3	Yield ^b [%]	er ^c
1	CN (1a)	36	3aa	86	98:2
2	$CONH_2$ (1i)	36	3ia	81	98:2
3	CO_2Et (1j)	48	3ja	80	97.5:2.5
4	CO_2Bn (1k)	48	3ka	85	98:2
5	CO_2H (11)	42	3la	<5 ^d	_
6	H (1m)	48	3ma	<5	_

^a Reaction conditions: 3 mol% [Ir(COD)Cl]₂, 6 mol% L1, 0.24 mmol of 1, 0.2 mmol of 2a and 0.2 mmol of DABCO in 0.6 mL CH₂Cl₂. The catalyst was prepared *via n*-PrNH₂ activation. ^b Yields correspond to the isolated product after chromatographic purification. ^c er was determined by HPLC analysis on a chiral stationary phase. ^d 1m was isolated in 52% yield.

These studies clearly indicate that the presence of an electron withdrawing group at the α -position of 4-methylcoumarin is necessary to exert its vinylogous reactivity. This prerequisite is certainly a limitation of our protocol and prevents the direct vinylogous allylic alkylation of α-unsubstituted 4-methylcoumarin. Our attempts to access this motif (3ma) through a one-pot sequential Ir-catalyzed γ-allylic alkylation/deallylative decarboxylation of allyl esters (1n and 1o) proved futile and led only to the formation of 1m (Scheme 3A). Similarly, an attempted intramolecular decarboxylative migratory allyl transfer²² of 1n under Ir-catalysis also resulted in deallylative decarboxylation to generate 1m. Finally, a two-step sequence consisting of ester hydrolysis followed by decarboxylation delivered the desired α-unsubstituted γ-allylcoumarin 3ma in overall 60% yield from 3ja without any erosion of enantiopurity (Scheme 3B).

The practicality of our enantioselective direct vinylogous allylic alkylation protocol is established by carrying out a gram-

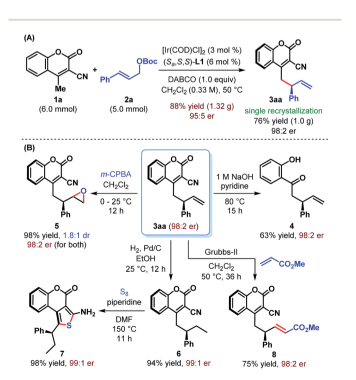
(A) OBoo [Ir(COD)CI]₂ (3 mol %) (S. S.S)-L1 (6 mol %) DABCO (1.0 equiv) R = Ph (1n) CH₂Cl₂ (0.33 M) R = H(10)50 °C (B) . LiOH·H₂O (5.0 equiv) THF/H₂O (1:1) 0 - 25 °C, 24 h 2. toluene, 120 °C 48 h 3ia 3ma (97.5:2.5 er) 60% vield (2 steps) 97.5:2.5 er

Scheme 3 Attempted and successful synthesis of α -unsubstituted γ -allylcoumarin.

scale synthesis of **3aa**, which gave the product in 88% yield but with somewhat diminished enantioselectivity (Scheme 4A). However, a single recrystallization restored the enantiopurity of **3aa** to 98: 2 er.

We realized that the ability to transform the existing functionalities present in the products may further increase their synthetic potential. Accordingly, a base-mediated retro-Knoevenagel condensation/hydrolysis of 3aa was carried out to furnish α -allylated o-hydroxyacetophenone 4 in 63% yield (Scheme 4B). Treatment of 3aa with m-CPBA provided the corresponding epoxide 5 in excellent yield but with poor diastereoselectivity (1.8:1 dr). Selective hydrogenation of the terminal double bond of 3aa was possible under Pd/C and resulted in 6 in 94% yield. A base-catalyzed cyclization of 6 with sulfur furnished tricyclic aminothiophenocoumarin 7. This structural motif is known for its presence in antifungal agents.23 The absolute stereochemistry of 3aa, 6 and 7 has previously been confirmed by Waldmann et al.24 and in turn established the absolute configuration of our allylated products. Finally, an olefin cross-metathesis of 3aa with methyl acrylate in the presence of Grubbs' 2nd generation catalyst occurred smoothly to give terminally functionalized olefin 8 in 75% yield. In all these cases, the reactions proceeded with complete conservation of stereochemical integrity.

A tentative catalytic cycle based on the literature precedence²⁵ is depicted in Scheme 5 and involves the intermediacy of the iridacycle intermediate **A**. Ligand dissociation from the coordinatively saturated species **A** is likely to generate species **B** having 16 valence electrons at Ir. Coordination of allylic carbonate (2) to **B** followed by oxidative addition-decarboxylation gives the π -allyl-Ir intermediate **C**. An



Scheme 4 Gram-scale synthesis and synthetic elaboration of γ -allylcoumarin 3aa.

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Scheme 5 Tentative catalytic cycle for the Ir-catalyzed enantioselective allylic alkylation of coumarins.

enantiodetermining nucleophilic addition of dienolate D to C then produces complex E, product dissociation from which completes the catalytic cycle. The sense of stereoinduction was found to be the same as predicted by the Hartwig model.25b

Conclusions

In conclusion, we have developed the first Ir-catalyzed enantioselective vinylogous allylic alkylation of coumarins. Our protocol does not require preactivation of 4-methylcoumarins and installs an unfunctionalized allyl group using easily accessible linear allylic carbonates as the allylic electrophile. The resulting Ir/phosphoramidite-catalyzed direct vinylogous allylic alkylation reaction produces γ-allylcoumarins in an exclusively branched-selective manner generally in high yields with an excellent level of enantioselectivity. An enantioselective synthesis of α-unsubstituted γ-allylcoumarin as well as synthetic elaboration of γ -allylcyanocoumarin to a diverse range of products have also been demonstrated. Future efforts in this direction from our laboratory would focus on the study of other classes of vinylogous nucleophiles.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research is funded by the Science and Engineering Research Board (SERB), New Delhi [Grant No. EMR/2016/ 005045]. R. S. thanks the Council of Scientific and Industrial Research (CSIR), New Delhi for a doctoral fellowship.

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