

Enantio-, Diastereo-, and Regioselective Iridium-Catalyzed Asymmetric Allylic Alkylation of Acyclic β -Ketoesters

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Supporting Information

ABSTRACT: The first regio-, diastereo-, and enantioselective allylic alkylation of acyclic β -ketoesters to form vicinal tertiary and all-carbon quaternary stereocenters is reported. Critical to the successful development of this method was the employment of iridium catalysis in concert with *N*-aryl-phosphoramidite ligands. Broad functional group tolerance is observed at the keto-, ester-, and α -positions of the nucleophile. Various transformations demonstrating the utility of this method for rapidly accessing complex enantioenriched compounds are reported.

he enantioselective synthesis of all-carbon quaternary stereocenters is an enduring challenge faced by organic chemists and a subject of longstanding interest in our laboratory.^{1,2} The generation of enantioenriched all-carbon quaternary centers is complicated by the presence of vicinal tertiary stereocenters due to increased steric demands and the introduction of requisite diastereocontrol. Modern strategies for accessing these highly congested stereochemical dyads have relied primarily on transition metal catalysis,³⁻⁷ notably Pdcatalyzed enolate alkylation cascades,³ Pd-catalyzed trimethylenemethane cycloadditions,⁴ Cu-catalyzed asymmetric Claisen rearrangements,⁵ and Mo⁶- and Ir⁷-catalyzed allylic alkylations. Common to the majority of these reports is the constraint that the nascent quaternary center be formed at a cyclic nucleophile. To date, only two groups have reported success in employing linear nucleophiles to produce vicinal quaternary/tertiary arrays, namely, Trost's communication on the molybdenumcatalyzed allylic alkylation of β -cyanoesters^{6b} and Carreira's recent report on the allylic alkylation of aldehydes using stereodivergent dual catalysis.^{7a} To address these limitations, we have initiated studies investigating the asymmetric allylic alkylation of linear β -ketoesters.

Recently, our group demonstrated the power of iridium-*N*-arylphosphoramidite catalysis⁸ in accessing vicinal all-carbon quaternary and tertiary stereocenters with our report on the regio-, diastereo-, and enantioselective asymmetric allylic alkylation of cyclic β -ketoesters (Scheme 1a).^{7b,9,10} The success of this protocol combined with the virtual absence of reports describing the application of this transformation to acyclic β -ketoesters encouraged our further exploration of iridium catalysts in the domain of this important substrate class. Herein, we report the first highly regio-, diastereo-, and

Scheme 1. Representative Ir-Catalyzed Asymmetric Allylic Alkylation



enantioselective allylic alkylation of acyclic β -ketoesters to forge vicinal tertiary, quaternary centers (Scheme 1b).

Having chosen ethyl 2-methyl-3-oxo-3-phenylpropanoate (1a) and cinnamyl carbonate (2a) as standard coupling partners, several iridacycle complexes¹¹ were investigated at the outset of our studies as shown in Table 1 (entries 1-6). We found that exposure of standard substrates 1a and 2a to a combination of catalytic phosphoramidite ligand L1·[Ir(cod)-Cl]₂ complex¹² and two equiv of NaH in THF at ambient temperature afforded the desired product with good conversion, ee, and regioselectivity but low levels of diastereoselectivity (1:2) (entry 1). Use of either L2 or L4 under these conditions instead favored the reaction pathway yielding the undesired, linear allylic alkylation product in modest conversion (entries 2 and 4). Ligands L5 and $L6^{8b,13}$ gave the branched product in good conversion but with diminished diastereoselectivity and enantioselectivity and protracted reaction times (entries 5 and 6). We were pleased to find that tetrahydroquinoline based ligand $L3^8$ rapidly furnished the desired α -quaternary β -ketoester (3a) in greater than 95% conversion, 95:5 regioselectivity, 13:1 dr, and 99% enantiomeric excess (entry 3). Previous reports demonstrating the marked effect of metal cations over regio-14 and diastereoselectivity^{7b,10e} in iridium-catalyzed allylic alkylations prompted further investigation of both bases and additives (see SI for details). Contrary to our previous findings,^{7b} a sluggish reaction was observed when LiBr was used in place of NaH (entry 7), presumably due to the decreased α -acidity of acyclic β ketoesters relative to cyclic substrates. Use of alkoxide bases in place of NaH, however, resulted in considerably reduced reaction times (entries 8-9). The dramatically diminished

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^{*a*}Reactions performed with 0.1 mmol of **2a**, 0.2 mmol of **1a** at 0.1 M in THF at 25 °C. ^{*b*}Determined by ¹H NMR and UHPLC-MS analysis of the crude mixture. ^{*c*}Determined by chiral SFC analysis; parenthetical value is the ee of the alternate diastereomer. ^{*d*}Not determined.



diastereoselectivity with KOt-Bu, Cs_2CO_3 , and DABCO indicates that the presence of a lithium cation in the reaction has a pronounced influence over the diastereoselectivity, likely due to formation of a rigid, bidentate-chelated lithium enolate ester (entries 8–11). Ultimately, it was found that LiOt-Bu proved optimal, delivering β -ketoester **3a** with an exceptional branched to linear ratio (93:7), >20:1 diastereoselectivity, and 98% enantioselectivity in only 2 h (entry 8).

With optimized conditions identified, the scope of the reaction with respect to the electrophile was next explored. A highly selective reaction was observed between β -ketoester 1a and various cinnamyl carbonate-derived electrophiles (2) bearing electron-donating substituents about the aryl group, R (Table 2, entries 2-4). 4-Me-, 4-MeO-, and 3-MeOsubstitutions about the aryl ring (substrates 2b-2d) gave the corresponding α -quaternary β -ketoesters (products 3b-3d) in good to excellent yield, dr, ee, and branched to linear ratio (Table 2). Electron deficient aryl substituents at the allyl group (entries 5-7) were also well tolerated, delivering the branched products $3e-3g^{15}$ in good to excellent yield, with outstanding ee and dr, and with only slightly diminished regioselectivities. Interestingly, (4-nitro)-aryl substitution at the allyl carbonate (entry 8, substrate 2h) led to loss of regioselectivity in the reaction, giving equal amounts of products 3h (14:1 dr, 93% ee) and 4h (23% ee). We were pleased to discover, however, that use of heteroaryl-substituted allyl carbonates (substrates 2i and 2j) resulted in smooth reactions and delivered alkylated products 3i and 3j with excellent yield, ee, and regioselectivity and with good to excellent dr (entries 9-10). Finally, we found that sorbyl carbonate 2k was also a suitable participant in the

Table 2. Substrate Scope of Allyl Carbonate Electrophiles^a

*	2	`OCO₂Me	1a [Ir(cod)Cl]₂ L3, TBD LiOt-Bu THF, 25 °C	Ph Me [*] CO ₂ l	Et .	Ph Me CO ₂ Et 4		
entry	2		product (3)		yield (%) ^b	3:4¢	dr of 3c	ee of 3 (%) ^d
1	2a	0 0	ſ	<i>3a</i> : R = H	97	93:7	>20:1	98
2	2b	Ph 👆		<i>3b</i> : R = Me	97	95:5	20:1	>99
3e	2c	EtO ₂ C		<i>3c</i> : R = MeO	85	99:1	>20:1	>99
4	2d	, Ĵ	F.	<i>3d</i> : R = MeO	99	90:10	17:1	>99
5	2e	EtO2C	Me 🚺	<i>3e</i> : R = CI	98	84:16	19:1	99
6	2f	P	ſ	<i>3f</i> : R = Br	98	86:14	14:1	>99
7	2g	Ph 🔨	\sim	<i>3g</i> : R = CF ₃	86	78:22	16:1	99
8	2h	EtO ₂ C		<i>3h</i> : R = NO ₂	78	50:50 [/]	14:1	93
9	2i	o II	ſ	<i>3i</i> : X = S	99	97:3	8:1	95
10	2j	Ph EtO ₂ C	Me x	<i>3j</i> : X = O	93	95:5	13:1	>99
11	2k	Ph EtO ₂ C	Me	3k	76	95:5	6:1	91

^{*a*}Reactions performed under the conditions of Table 1, entry 8. ^{*b*}Combined isolated yield of **3** and **4**. ^{*c*}Determined by ¹H NMR analysis of the crude mixture. ^{*d*}Determined by chiral SFC analysis of the major diastereomer. ^{*e*}Conditions of Table 1, entry 7. ^{*f*}23% ee for the linear product.

reaction, giving the corresponding product (3k) in good yield and dr and with excellent regio- and enantioselectivity (entry 11).

During the course of this investigation, a trend relating regioselectivity and electrophile electron deficiency began to emerge. Specifically, the regioselectivity of the reaction diminished as the electron deficiency of the cinnamyl substituent increased. In order to identify any linear free energy relationship governing the reaction, we performed a linear relationship analysis relating the log of the ratio of branched to linear products, which is proportional to the relative rates of product formation, to the corresponding Brown σ^+ constants (Figure 1).^{16,17} The negative ρ value observed from this plot suggests that as the magnitude of electropositive charge generated at the putative cinnamyl-Ir intermediate¹⁸



Figure 1. Linear relationship analysis of the log of product ratios (3:4) from Table 2 versus Brown σ^+ values.

increases, the reaction pathway yielding the branched allylation product becomes more favorable.¹⁹

Having investigated the reaction substrate scope with respect to the allyl electrophile, we next examined the diversity of nucleophilic coupling partners permitted in the chemistry (Table 3). β -Ketoesters (1) bearing either electron-donating or

Table 3.	Substrate	Scope	of B-	Ketoesters	Nuc	leophiles ^a
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° R¹	R ²	$\begin{array}{c} 2a & O & Ph \\ \hline OR^3 & \underline{L3, TBD} \\ LiOt-Bu \\ THF, 25 \ ^{\circ}C \\ 3 \end{array}$	$ \begin{array}{c} D \\ D \\ D \\ C \\ C \\ \end{array} \begin{array}{c} 0 \\ R^2 \\ R^2 \\ C \\ 3 \end{array} \begin{array}{c} Ph \\ Ph \\ R^2 \\ C \\ R^3 \\ C \\ R^3 \end{array} $		$+ R^{1} \xrightarrow{O}_{R^{2} CO_{2}R^{3}} Ph$ 4		
entry	1	product (<i>3</i>)	yield (%) ^b	3:4 ^c	dr of 3°	ee of 3 (%) ^d	
1	1/	O Ph $3l: \mathbb{R}^3 = \mathbb{E}t$ $\mathbb{H} \xrightarrow{\mathbb{F}} \mathbb{R}^1 = 4-\text{MeO-C}_6 \mathbb{H}_4$	93	93:7	17:1	99	
2	1 <i>m</i>	$\begin{array}{c} \mathbf{R}^{1} & 3m; \mathbf{R}^{3} = \mathbf{M} \mathbf{e} \\ \mathbf{M} \mathbf{e}^{2} \mathbf{CO}_{2} \mathbf{R}^{3} & \mathbf{R}^{1} = 4 - \mathbf{Br} - \mathbf{C}_{6} \mathbf{H}_{4} \end{array}$	92	92:8	17:1	>99	
3	1n	<i>3n</i> : R ² = Et	94	90:10	>20:1	>99	
4	10	<i>30</i> : R ² = Bn	99	90:10	13:1	>99	
5	1p	<i>3p</i> : R ² = allyl	98	95:5	>20:1	>99	
6 ^e	1q	O Ph 3q: R ² = propargyl	84	81:19	13:1	>99	
7'	1r	Ph $rac{1}{R^2 CO_2 Et}$ 3r. R ² = (CH ₂) ₂ COMe	98	93:7	20:1	99	
8	1s	<i>3s</i> : R ² =	88	95:5	7:1	>99	
9	11	3t: R ² = CH ₂ CH ₂ CN	99	95:5	3:1	>99 (>99) ^g	
10	1u	<i>3u</i> : R ² = F	92	96:4	13:1	95	
11	1v	<i>3v</i> : R ² = Cl	96	96:4	>20:1	>99	
12	1w	Me ^x CO ₂ Me	85	90:10	12:1	99	
13	1x	$\begin{array}{ccc} O & Ph & 3x: R^1 = Cy \\ \hline & & & \\ \hline \end{array}$	92	92:8	4:1	96	
14	1y	$\begin{array}{c} R^1 & 3y; R^1 = Me \\ R^2 & CO_2 Me & R^2 = Et \end{array}$	90	93:7	1.5:1	90 (91) ^g	
15	1z	Ph Ph Me ³ CO ₂ t-Bu 3z	95	70:30	6:1	>99	

^{*a*}Reactions performed under the conditions of Table 1, entry 8. ^{*b*}Combined isolated yield of **3** and **4**. ^{*c*}Determined by ¹H NMR analysis of the crude mixture. ^{*d*}Determined by chiral SFC analysis of the major diastereomer. ^{*e*}4 mol % of $[Ir(cod)Cl]_2$ and 8 mol % of L3 were used. ^{*f*}The reaction was run at 0.5 mmol scale. ^{*g*}Ee for the minor diastereomer.

electron-withdrawing aryl substituents (R^1) at the ketone fared very well in the reaction, delivering products 31 and 3m in excellent yield, dr, ee, and branched to linear ratio (entries 1 and 2). Gratifyingly, a wide variety of functional groups are readily permitted at the α -position (R²), including alkyl, benzyl, allyl, propargyl, keto, and heteroaryl groups (substrates 1n-1s, entries 3-8, respectively). The products of these reactions (products 3n-3s, respectively) were obtained with excellent ee and regioselectivities and in good to excellent dr and yield. To the best of our knowledge, substrate 1q represents the first example of a nucleophile bearing propargyl substitution to undergo Ir-catalyzed allylic substitutions.²⁰ Nitrile-containing substituents were tolerated in the reaction as well (substrate 1t), and α -quaternary β -ketoester 3t was furnished in excellent yield, ee, and regioselectivity, albeit with diminished dr (3:1). We were pleased to learn that use of α -halogenated nucleophiles (substrates 1u and 1v) also resulted in an efficient

and selective reaction as α -fluoro and α -chloro β -ketoesters **3u** and **3v** were obtained in excellent yield, dr, ee, and regioselectivity. In addition to aryl ketones, cyclohexenyl β ketoester **1w** was found to deliver the corresponding product **3w** in excellent yield, dr, ee, and branched to linear ratio with no detectable products resulting from competitive bimolecular Michael addition. Although the use of alkyl β -ketoesters **1x** and **1y** provided the desired products (**3x** and **3y**, respectively) with excellent yield, ee, and regioselectivity, we were disappointed to find that the diastereoselectivities were diminished considerably. Lastly, we found that the use of a sterically hindered ester moiety (**1z**) gave an efficient and highly enantioselective reaction but with a concurrent loss in regio- and diastereoselectivity (entry 15).

In order to exhibit the utility of our method for generating interesting and useful chiral building blocks, a number of selective transformations were carried out on products obtained in the course of our studies (Scheme 2). Aldol condensation of

Scheme 2. Derivatization of β -Ketoester Products^{*a*}



^{*a*}Conditions: (a) **3r** (R = CH₂CH₂COMe), pyrrolidine, AcOH, *t*-BuOMe, reflux, 95% yield. (b) **3q** (R = propargyl), $Co_2(CO)_{8}$, CH₂Cl₂, 25 °C, then Me₃NO·2H₂O, >20:1 dr, 99% yield. (c) **3p** (R = allyl), Hoveyda–Grubbs II (10 mol %), CH₂Cl₂, 40 °C, 96% yield.

 β -ketoester **3r** yielded γ -quaternary cyclohexenone **5**. Pauson– Khand cyclization of progaryl-substituted β -ketoester **3q** smoothly delivered bicycle **6**.^{20b} Finally, ring closing metathesis of diallyl β -ketoester **3p** cleanly furnished cyclohexene **7**.

In summary, the first enantioselective catalytic allylic alkylation of linear β -ketoesters to generate vicinal quaternary and tertiary stereocenters in high yield, dr, ee, and regioselectivity has been reported. The process hinges on the use of an Ir-*N*-aryl-phosphoramidite catalyst. A variety of substitution patterns at the allyl electrophile and β -ketoester are well tolerated in the chemistry. A number of transformations were carried out on reaction products to demonstrate the value this method holds for the rapid generation of highly functionalized chiral building blocks. Studies utilizing this method toward the synthesis of complex biologically active natural products are underway in our laboratory.

ASSOCIATED CONTENT

S Supporting Information

Experimental procedures, characterization data, X-ray analysis. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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