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Organocatalytic asymmetric oxy-Michael addition to γ -hydroxy- α , β unsaturated thioester via hemiacetal intermediates

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We report an asymmetric oxy-Michael addition to a γ -hydroxy- α , β -unsaturated thioester via hemiacetal intermediates in the presence of cinchona-alkaloid-thioureabased bifunctional organocatalysts. This method provides a

 $_{\rm 10}$ novel enantioselective route to β -hydroxy carboxyl compounds, which in turn can be used to synthesise valuable chiral building blocks.

β-Hydroxy carbonyl compounds are important synthetic intermediates, and they exist as structural motifs in a variety of ¹⁵ natural products; hence, considerable efforts have been devoted to their stereoselective synthesis.¹ One of the most notable methods for the enantioselective synthesis of β-hydroxy carbonyls, besides the aldol reaction² and the hydrogenation of βketoesters,³ is the conjugate addition of oxygen-centred ²⁰ nucleophiles to α,β-unsaturated substrates.⁴ Direct hydration of

- α , β -unsaturated substrates by the conjugate addition of water is challenging because of the low nucleophilicity of water, high reaction reversibility, and the difficulty involved in efficient stereochemical control.⁵ Nevertheless, several protocols for ²⁵ stereoselective formal hydration using *O*-nucleophiles bearing a removable group have been developed.^{6–11} Catalytic
- removable group have been developed.^{6–11} Catalytic enantioselective reactions involving the conjugate addition of benzyl alcohol⁸ or allyl alcohol⁹ suffer from drawbacks similar to those observed when using water as the nucleophile. However,
- ³⁰ the use of an oxime^{10a-10c} or hydrogen peroxide^{10d} as a water surrogate is advantageous because of its high nucleophilicity; the labile *N*-*O* or *O*-*O* bond facilitates the subsequent reductive cleavage, leading to free β -hydroxy products. Another efficient route is intramolecularization¹² using boronic acid hemiesters
- ³⁵ generated *in situ* from γ-hydroxy-α,β-unsaturated ketones and boronic acids; the hemiester undergoes intramolecular oxy-Michael addition to form a dioxaborolane, which then affords the corresponding optically active β ,γ-dihydroxy ketone upon oxidative hydration.¹¹ However, in most of the reported
- ⁴⁰ examples, α , β -unsaturated ketones or aldehydes have been used as substrates, and there are very few demonstrations of asymmetric oxy-Michael addition to a higher-oxidation-state substrate such as an α , β -unsaturated carboxylic acid derivative,^{10a,13} which can be an alternative to the acetate aldol ⁴⁵ reaction.¹⁴

We recently reported intramolecular oxy-Michael addition reactions mediated by cinchona-alkaloid-thiourea-based bifunctional organocatalysts.¹⁵ By our protocol, enantioselective oxy-Michael addition to γ-hydroxy- α , β -unsaturated ketones via ⁵⁰ hemiacetal intermediates was realized, and 1,3-dioxolanes bearing an easily removable acetal functionality were obtained (Scheme 1).^{15a} Although the diastereoselectivity of this reaction was only moderate, the absolute configurations at the β -positions of the carbonyl group were consistent in both diastereomers;

⁵⁵ further, this reaction proceeded with high enantioselectivity. Therefore, we sought to apply the abovementioned method to reactions in which carboxylic acid derivatives were used as substrates. Herein, we present a novel asymmetric oxy-Michael addition to a γ-hydroxy-α,β-unsaturated carboxylic acid
⁶⁰ derivative via a hemiacetal intermediate in the presence of bifunctional organocatalysts derived from cinchona alkaloids.^{6,16}



Scheme 1 Oxy-Michael addition via hemiacetal intermediates to γ -hydroxy- α , β -unsaturated ketone via hemiacetal intermediate.

Initially, we employed the optimized conditions reported in our previous work of the reaction with γ -hydroxy- α , β -unsaturated ketones as substrates (Table 1).^{15a} The reaction of γ -hydroxy- α , β unsaturated ester 1a with cyclohexanecarboxaldehyde (2a) in the presence of quinidine-based bifunctional catalyst 4a (Fig. 1) did 70 not proceed at all, presumably because of the poor electrophilicity of the substrate (Table 1, entry 1). Phenyl ester 1b afforded the desired products, albeit in very low yield (Table 1, entry 2). In order to increase the electrophilicity of the substrate, we used some thioesters as substrates (Table 1, entries 3–6).¹⁷ Thioester 75 1c afforded the corresponding product in higher yield than did the abovementioned esters (Table 1, entry 3). Benzenethiol ester 1d also gave the desired products in low yield, but when thioesters bearing bulkier aryl groups were employed, side reactions were suppressed to a great extent. 2,6-Dimethylbenzenethiol ester 1e ⁸⁰ was identified as the best substrate in terms of the product yield (Table 1, entries 4–6).

We next optimized the reaction conditions using **1e** as the substrate (Table 2). After screening a number of solvents, we

found that cyclopentyl methyl ether (CPME) was the optimum solvent in terms of both yield and stereoselectivity (Table 2, entries 1–5). Further modification of other conditions such as the amount of **2a**, concentration, and reaction time helped improve the yield to a practical level with only a slight degrees in the

- ⁵ the yield to a practical level with only a slight decrease in the enantioselectivity (Table 2, entry 5). Catalyst screening showed that **4c** (Fig. 1) efficiently catalyzes the reaction to afford opposite enantiomers of the products in good yield and with high enantioselectivity (Table 2, entry 8).
- ¹⁰ Table 1 Optimization of substrates^{*a,b*}

		4a (10 mol %) CPME, 25 °C, 24 h R ⁻		RX 3'
	1b: R 1c: R	K = PhO 1e: R' = K = BnS 1f : R' =	= 2,6-dimethyl = 2,4,6-triisopropyl	
Entry	1	Yield $(\%)^{c,d}$	<i>ee</i> (%) (3 , 3 ')	dr (3/3')
1	1a	0 (97)		
2	1b	13 (87)	96, 84	3.0
3	1c	18 (69)	94, 61	3.6
4	1d	7 (<1)	97, 87	3.7
5	1e	62 (38)	97, 84	4.3
6	1f	28 (60)	93, 85	4.0

^{*a*} Reactions were run using **1** (0.25 mmol), **2a** (0.25 mmol), and **4a** (0.025 mmol) in CPME (0.5 mL). ^{*b*} CPME = cyclopentyl methyl ether. ^{*c*} Isolated yields. ^{*d*} Values in parentheses show the starting material ¹⁵ recovery.



Fig. 1 Bifunctional organocatalysts derived from cinchona alkaloids.

Table 2 Optimization of reaction conditions^{*a,b*}

Ars + OH + O						
Entry	Catalyst	Solvent	Yield $(\%)^{c,d}$	<i>ee</i> (%) (3ea, 3ea')	<i>dr</i> (3ea/3ea')	
1	4a	CPME	62 (38)	97, 84	4.3	
2	4a	THF	20 (78)	96, 76	4.3	
3	4a	Et ₂ O	65 (12)	97, 87	3.9	
4	4a	benzene	60 (<1)	94, 84	3.4	
5	4 a	CH_2Cl_2	43 (1)	95, 73	4.1	
6^e	4 a	CPME	90 (8)	96, 81	4.4	
7^e	4b	CPME	89 (1)	95, 74	3.5	
8^e	4 c	CPME	90 (6)	-94, -59	4.2	
9^e	4 d	CPME	90 (6)	-91 -47	41	

²⁰ ^a Reactions were run using **1e** (0.25 mmol), **2a** (0.25 mmol), and the catalyst (0.025 mmol) in the solvent (0.5 mL). ^b CPME = cyclopentyl methyl ether. ^c Isolated yields. ^d Values in parentheses show the starting material recovery. ^e Reactions were run using 0.50 mmol of **2a** in 0.25 mL of CPME for 48 h.

Using the optimized reaction conditions and 4a as a catalyst, we subsequently investigated the reactions of some other aldehydes and ketones 2 (Table 3).¹⁸ Although aryl aldehydes

were much less reactive,¹⁹ some aliphatic aldehydes **2b-2d** gave the corresponding products in high yields and with good 30 enantioselectivity (Table 3, entries 2–4). Pivalaldehyde (2d) proved to be a good counterpart and gave excellent enantioselectivity for both diastereomers (Table 3, entry 4). An electoron-deficient ketone such as 2e could also be employed in the reaction; however, the enantioselectivity was only moderate 35 in this case (Table 3, entry 5). Although the use of a symmetric ketone or aldehyde would circumvent the generation of diastereomers, the reaction using acetone (2f) was sluggish (Table 3, entry 6), and the yield obtained from cyclohexanone (2g) was low despite the excellent enantioselectivity (Table 3, ⁴⁰ entry 7). On the other hand, aqueous formaldehyde (2h) afforded the product in acceptable yield, but the enantioselectivity was unsatisfactory, probably because of the presence of water (Table 3, entry 8).

Table 3	Ontimization	of aldehydes	and	ketones	2^{a}
1 abic 5	Optimization	of alucityues	anu	Retones	4

-	Ars 1e	∽ ^{0H ⁺ R}	$\frac{1}{2} \frac{1}{R^2} \frac{4}{C}$	a (10 mol %) PME, 25 °C,	48 h Ars 3	Ar = 2,6-dim	3' ethylphenyl
_	Entry	\mathbf{R}^1	\mathbb{R}^2	2	Yield $(\%)^c$	ee (%) (3 , 3')	dr (3/3')
	1	Су	Н	2a	90	96, 81	4.4
	2	Et	Н	2b	99	95, 88	3.4
	3	<i>i</i> -Pr	Н	2c	99	96, 87	3.8
	4	t-Bu	Н	2d	73	99, 97	3.5
	5	CF_3	Ph	2e	99	69,72	1.1
	6	CH_3	CH_3	2f	<5	N. D.	
	7	-(CH ₂) ₅ -		2g	31	99	
	8^d	Н	Н	2h	86	72	

^{*a*} Reactions were run using **1e** (0.25 mmol), **2** (0.5 mmol), and **4a** (0.025 mmol) in CPME (0.5 mL). ^{*b*} CPME = cyclopentyl methyl ether. ^{*c*} Isolated yields. ^{*d*} Reaction was run using aqueous formaldehyde (37% solution, 0.5 mmol).

To demonstrate the utility of the proposed method, we extended the optimized reaction to the asymmetric syntheses of some β -hydroxy carboxyl compounds (Scheme 2). Oxy-Michael addition to **1e** using **2d** as the source of oxygen-centred nucleophile in the presence of 13 mol % **4a** on 2-mmol scale ⁵⁵ afforded the products **3ed** and **3ed'** in 3.8:1 diastereomeric ratio, with excellent enantioselectivity. Subsequent treatment of the diastereomixture of **3ed** and **3ed'** with titanium tetrachloride led to the generation of a free β , γ -dihydroxy product **5** with high optical purity while keeping the thioester group intact. ⁶⁰ Alternatively, treatment of the diastereomixture with *p*toluenesulfonic acid in aqueous medium gave β -hydroxy- γ butyrolactone **6**, a versatile chiral synthetic intermediate²⁰ that could be transformed into (*L*)-carnitine (**7**), an important bioactive agent, via a reported procedure.²¹

Taking advantage of the thioester functionality, we carried out functional group transformations of **3ed**, and found that the chiral acetal moiety was unaffected after the transformations (Scheme 3).^{15a,22} Reduction of **3ed** with lithium aluminium hydride afforded the corresponding primary alcohol **8** quantitatively without loss of optical purity. Besides, Liebeskind–Srogl cross coupling enabled the replacement of the arylthio group of **3ed** to give ketone **9**, indicating that these thioester products can be easily transformed into various chiral ketones.²³



Scheme 2 Application of proposed protocol to asymmetric syntheses of β -hydroxy carboxyl compounds.



Scheme 3 Transformations of the thioester group of 3ed.

In conclusion, we have developed a novel asymmetric oxy-Michael addition reaction to the α , β -unsaturated carboxylic acid derivative. The use of a suitable γ -hydroxy- α , β -unsaturated thioester allowed for enantioselective oxygen induction via

- ¹⁰ hemiacetal formation, and subsequent deacetalization afforded valuable optically active β -hydroxy carboxyl compounds. Further studies on the application of this methodology to the asymmetric syntheses of various chiral materials, including natural products, are currently underway in our laboratory.
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