Mechanistic insight into asymmetric hetero-Michael addition of α , β unsaturated carboxylic acids catalyzed by multi-functional thioureas

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ABSTRACT: Carboxylic acids and their corresponding carboxylate anions are generally utilized as Brønsted acids/bases and oxygen-nucleophiles in organic synthesis. However, a few asymmetric reactions have used carboxylic acids as electrophiles. Although chiral thioureas bearing both arylboronic acid and tertiary amine were found to promote the aza-Michael addition of BnONH₂ to α , β -unsaturated carboxylic acids with moderate to good enantioselectivities, the reaction mechanism remains to be clarified. Detailed investigation of the reaction using spectroscopic analysis and kinetic studies identified tetrahedral borate complexes, comprising two carboxylate anions, as reaction intermediates. We realized a dramatic improvement in product enantioselectivity with the addition of one equivalent of benzoic acid. In this aza-Michael reaction, the boronic acid not only activates the carboxylate ligand as a Lewis acid, together with the thiourea NH-protons, but also functions as a Brønsted base through a benzoyloxy anion to activate the nucleophile. Moreover, molecular sieves were found to play an important role in generating the ternary borate complexes, which were crucial for obtaining high enantioselectivity as demonstrated by DFT calculations. We also designed a new thiourea catalyst for the intramolecular oxa-Michael addition to suppress another catalytic pathway via a binary borate complex using steric hindrance between the catalyst and substrate. Finally, to demonstrate the synthetic versatility of both hetero-Michael additions, we used them to accomplish the asymmetric synthesis of key intermediates in pharmaceutically important molecules, including sitagliptin and α -tocopherol.

Introduction

Exploiting powerful catalytic methodologies for the asymmetric syntheses of natural products and potential pharmaceuticals from simple materials without using protecting and activating groups is the ultimate goal of synthetic organic chemistry.¹ Catalytic asymmetric Michael additions have attracted much attention regarding their atom economy and green credentials. Accordingly, an increasing number of catalytic asymmetric reactions using activated carboxylic acid derivatives² have been developed and utilized for the construction of complex molecules.³ However, new design concepts for chiral multifunctional catalysts capable of significantly activating substrates and nucleophiles are still needed, especially for reactions using less reactive Michael acceptors, such as α , β unsaturated esters, amides, and carboxylic acids.⁴ The direct Michael addition to α,β -unsaturated carboxylic acids 1 is particularly challenging,⁵ because carboxylic acids generally form inert salts with basic nucleophiles, and the resulting carboxvlate anions are known to be the least reactive electrophiles among carboxylic acid derivatives for both the 1,2- and 1,4additions. Therefore, so far only a few such methods, using enzymes such as ammonia lyases and aminomutases, have been reported,⁶ despite the versatile utilities of adducts 2 (Scheme 1).⁷ At present, no efficient synthetic tools exist for activating free carboxyl groups in a direct catalytic manner. Such a tool would mean that both the protection of 1 to activated adducts **3** using one equivalent of activating auxiliaries (X) and the deprotection of Michael adducts **4** to the corresponding carboxylic acids **2** would not be necessary. This would allow desired products **2** to be obtained in a step- and atom-economical fashion from **1**. In addition, more advanced adducts **5** can be directly synthesized from **2** using established decarboxylative functionalization strategies with photoredox catalysts.⁸

Scheme 1. Concept of this work



Recently, carboxylates have played an important role in ion pair catalysis using chiral thioureas.⁹ This concept has been extended to a variety of asymmetric reactions, but carbox-

ylates that form dual hydrogen bonds with thioureas only work as chirality transmitters and have never been used as electrophiles.¹⁰ In contrast, various arylboronic acids have been developed for the direct condensation of carboxylic acids with amines or alcohols.11 Therefore, we envisioned that the synergistic interaction of carboxylate anions with both thiourea and arylboronic acid would enable direct Michael addition to 1. Accordingly, we have developed the first asymmetric aza-Michael addition^{12a} of O-benzylhydroxylamine 6 to α , β unsaturated carboxylic acid **1a** using catalyst **7a** to afford β amino acid derivative 2a (Eq. 1), and the asymmetric intramolecular oxa-Michael addition^{12b} of phenolic α,β -unsaturated carboxylic acids using dual catalysts of a chiral bifunctional thiourea^{12c,d} and arylboronic acid. However, no spectroscopic evidence was obtained for the reaction mechanism of the aza-Michael addition. Herein, to improve the enantioselectivity and clarify the reaction mechanism, we investigated the following: (i) how catalyst 7 interacts with substrate 1a and nucleophile 6 to furnish (S)-2a as the major enantiomer: (ii) why 4Å molecular sieves (4Å MS) were required to promote the reaction; (iii) why, in some cases, the enantioselectivity was significantly influenced by the β -substituents of substrates 1 (68-86% ee); and (iv) can a green solvent be used instead of volatile CCl₄, which was a major disadvantage of this reaction.



In our previous work, we hypothesized that catalyst 7 would form zwitterionic complex A with carboxylic acid 1 through the acid-base interaction of 1 with the Brønsted basic moiety of 7 (Scheme 2), in which the resulting carboxylate anion is stabilized by both the thiourea and arylboronic acid moieties of the catalyst. The subsequent 1,4-addition of BnONH₂ to the s-trans form of the coordinated carboxylic acid occurred from the top side, assisted by the borate hydroxy group, affording (S)-2a predominantly. However, borate complexes B can also be generated from complex A by a second ligand exchange with another molecule of substrate 1. In both complexes A or **B**, the thiourea can activate the carbonyl oxygen through dual hydrogen-bonding interactions, thereby promoting nucleophilic addition. To identify which complex is more plausible in terms of the nucleophile activation mode, several spectral analyses and kinetic studies were performed to describe the reaction mechanism of the aza-Michael addition. The origin of enantioselectivity was also examined via DFT calculations.

Scheme 2. Possible catalyst–substrate complexes



Furthermore, based on these results, we have rationally designed a new catalyst, and identified efficient additives and environmentally friendly solvents to enhance reaction selectivity. Finally, the optimized reactions were successfully applied to the formal asymmetric synthesis of pharmaceutically important molecules sitagliptin and α -tocopherol.

Result and discussion

To obtain structural information of the catalyst, we first performed X-ray crystallographic analysis. Fortunately, a suitable crystalline solid was obtained by recrystallizing **7b** from acetone-hexane.¹³ As expected,¹⁴ intramolecular hydrogen bonding was observed between one of the hydroxyl groups of the boronic acid and the tertiary amino group. More importantly, the boronic acid and thiourea face each other, which allows simultaneous activation of the resulting carboxylate anion by both the thiourea and arylboronic acid moieties, particularly with the *syn* arrangement of the thiourea NH groups.¹⁵



Figure 1. (a) Structure of complex 7b and acetone in the solid state. (b) ORTEP drawing of 7b•acetone.

We next measured ¹¹B NMR spectra to determine the coordination number of the boron atom of **7b** in solution (Figure 2). The spectrum of **7b** showed a broad single peak at 28 ppm in CDCl₃, indicating that the boron atom exists as a trigonal planar species according to literature values (Figure 2a).¹⁶ This result was in good agreement with the X-ray crystallographic studies above (Figure 1). However, the addition of 4Å MS to the catalyst solution resulted in the appearance of new signals at 4 and 8 ppm, in addition to the original peak (Figure 2b).



Figure 2. ¹¹B NMR titration experiments of catalyst **7b** with crotonic acid **1b** in CDCl₃ (0.033 M): (a) **7b**; (b) **7b** with 4Å MS; (c) **7b** with **1b** (1 equiv) and 4Å MS; (d) **7b** with **1b** (10 equiv) and 4Å MS.

The complexes could not be identified by ¹H NMR analysis, but DOSY NMR experiment indicated the dimerization of catalyst **7b** (see Figure S5 in Supporting Information). In fact, a signal for dimeric boron complex of **7b** (calcd. for $C_{42}H_{55}B_2N_6O_3S_2^+$ [M+H]⁺: 777.3958) was observed at 777.3983 in ESI-MS spectrum (see Figure S7 in Supporting Information). The dimeric form of the catalyst is thus considered to be generated by releasing one equivalent of water from two molecules of **7b** in the presence of 4Å MS (Scheme 3). The signal at 28 ppm in ¹¹B NMR spectrum can be assigned to the trigonal boron dimer complex, but the two signals at 4 and 8 ppm suggest that tetrahedral borate complexes, for instance, species with coordinated tertiary amino groups might also be formed. Interestingly, no peak of boroxine of **7b** was detected in the ESI-MS spectrum.

Scheme 3. Dimerization of 7b and complex A in the presence of 4\AA MS



In contrast, when one equivalent of crotonic acid **1b** was added in the presence of 4Å MS, the peak at 28 ppm almost disappeared, with a new peak appearing at 9 ppm (Figure 2c), which could be assigned to B–O–B bridged dimeric forms of complex A,¹⁷ as shown in Scheme 3.¹⁸ With a large excess of crotonic acid (**7b** and **1b** in a 1:10 ratio), another new signal at 4 ppm was only observed (Figure 2d). This signal was assigned to complex **B**, which can be readily formed in the



Figure 3. ¹¹B NMR titration experiments of 7b with maleic acid 8 in CDCl₃ (0.033 M): (a) 7b with 4Å MS; (b) 7b with 8 (0.5 equiv) and 4Å MS; (c) 7b with 8 (1.0 equiv) and 4Å MS; (d) 7b with 8 (2.0 equiv) and 4Å MS.

reaction of complex **A** dimer with another two molecules of carboxylic acid (Scheme 3 and see Figure S6 in Supporting Information). Notably, without 4Å MS, almost no peaks were observed at 4 and 9 ppm, even in the presence of 10 equiv of **1b** (see Figure S3 in Supporting Information). The presence of 4Å MS seems to be essential for the generation of the tetrahedral borate complexes, and complex **B** was presumed to play an important role in the enantioselective Michael addition.¹⁹

To confirm whether two carboxy groups could coordinate to the boron atom of **7b**, we performed ¹¹B NMR titration experiments of **7b** with maleic acid **8** (Figure 3). Interestingly, when 0.5 equiv of **8** was added, only two major peaks at 28 and 3 ppm were observed (Figure 3b), which were distinct from the results in Figure 2c. Increasing the ratio of **7b/8** to 1:1 and 1:2 caused the signal at 28 ppm to disappear, with the same peak at 3 ppm developing (Figure 3c and 3d). These results suggest that both carboxy groups in **8** could be incorporated into borate complexes, as in complex **C**, shown in Figure 4.

We further performed ESI-MS analyses²⁰ of mixtures of **7b/1b** and **7b/8** in the presence of 4Å MS to identify borate complexes **A**–**C**. Despite multiple trials, we could not detect the exact mass signals of complex **A** dimer and double-coordinated complex **B** due to their instability, and only observed a signal for monocoordinated complex **A** at 482.2272, irrespective of the **7b/1b** ratio (see Figures S8 and S9 in Supporting Information). In contrast, when catalyst **7b** and maleic acid **8** were mixed in a 1:1 ratio, the exact mass peak of complex **C** (calcd. for $C_{25}H_{29}BN_3O_5S^-$ [M–H]^{-:} 494.1926) was detected as a major peak at 494.1913 with errors of no more than 5 ppm in the negative region of the spectrum. Complex **C** is thought to be generated by the dehydration of complex **A** dimer consisted of **7b** and **8**.



Figure 4. Detection of complex C by ESI-MS analysis.

In addition to these spectroscopic analyses, we performed kinetic studies using classical initial rate kinetics for the reaction shown in Eq. 1. Using an excess of 6, the reaction rate became greater, as the concentration of 1a was increased gradually from 5.0 mM to 20 mM (Figure 5a). The reaction rates vs concentration plot reveals a linear relationship for the initial phase of the reaction (Figure 5b), pointing to first order kinetics in substrate 1a. Regarding the nucleophile 6, the kinetic measurements similarly suggest first order rate dependence in the initial phase (see Figures 5c-d). Additional kinetic data indicate that the reaction rate is reduced after the initial period (see Figures S10a and S11a). This latter kinetic behavior is likely due to product inhibition, which could be supported by further experiments. Addition of product 2a to the reaction mixture retarded the reaction significantly (Figure 5e).²¹ Finally, the reaction was found to be first order with respect to the catalyst 7b, as illustrated by the overlay of conversion



Figure 5. Kinetic studies of the reaction of carboxylic acid 1a and BnONH₂ 6 catalyzed by 7b with 4Å MS in CCl₄. (a) ($[1a]_0-[1a]$) vs. time using an excess (10 equiv) of 6 (first 0.5 h). [\blacktriangle : 20 mM; \triangle : 15 mM; •: 10 mM; o: 5.0 mM of 1a]. (b) (d[1a]/dt) vs. [1a]. (c) ($[6]_0-[6]$) vs. time using an excess (10 equiv) of 1a (first 2.0 h). [\bigstar : 20 mM; \triangle : 15 mM; •: 10 mM; o: 5.0 mM of 6]. (d) (d[6]/dt) vs. [6]. (e) [1a] vs. time with and without product 2a [•: standard conditions; \bigstar : with 2a added at time = 0; :: with 2a mixed with catalyst 7b for 4 hours prior to reaction]. (f) [1a] vs. time [7b] ([•: 5 mol%; \bigstar : 10 mol%; o: 15 mol% in [7b]).

curves on the [1a] vs time [7b] plot (Figure 5f).²²

A plausible reaction mechanism emerging from the results of NMR titration and kinetic experiments is shown in Scheme 4. The molecular sieves play a crucial role in the formation of dimeric catalyst, which readily reacts with carboxylic acid 1 to form a dimer of tetra-coordinated borate complex A and subsequently complex **B**. This latter process is accompanied by the release of water as well. In contrast, the dimerization of neither 7b nor complex A can occur in the absence of 4Å MS. In complex **B**, one of the coordinated carboxylate ligands is thought to form hydrogen bonds with the thiourea NH groups. The second coordinated carboxylate may act as a Brønsted base that binds and activates the nucleophile via H-bonding interactions. Based on the first order kinetics in both substrates (1 and 6), we anticipate that complex **B** is the resting state of the catalytic cycle in the initial phase of the reaction, and the rate-determining C-N addition step takes place via the assistance of an additional carboxylic acid (see transition state TS in Scheme 4). The reaction likely proceeds via proton transfer from the NH₂ group to the sp² carbon of the adduct to give intermediate int. This species is analogous to complex B, but it involves a product carboxylic acid. The catalytic cycle is finally completed by subsequent ligand-exchange by substrate 1 to furnish the desired product 2.

Scheme 4. Plausible reaction mechanism



Table 1. Effect of carboxylic acids as additives and reoptimization of reaction conditions^a



Entry	Catalyst	Additive (equiv)	Solvent	Yield (%) ^b	ee (%) ^c
1	7a	none	CCl ₄	84	73
2	7b	none	CCl ₄	77	76
3	7a	none	Cl ₂ C=CCl ₂	47	48
4	7b	none	Cl ₂ C=CCl ₂	73	55
5	7a	none	4-CF ₃ C ₆ H ₄ Cl	31	55
6	7b	none	4-CF ₃ C ₆ H ₄ Cl	57	69
7	7b	PhCO ₂ H (1.0)	CCl ₄	85	97
8	7b	PhCO ₂ H (1.0)	Cl ₂ C=CCl ₂	77	95
9	7b	PhCO ₂ H (1.0)	4-CF ₃ C ₆ H ₄ Cl	53	92
10	7b	AcOH (1.0)	Cl ₂ C=CCl ₂	6	26
11	7b	$4-MeOC_{6}H_{4}CO_{2}H(1.0)$	Cl ₂ C=CCl ₂	69	88
12	7b	4-CF ₃ C ₆ H ₄ CO ₂ H (1.0)	Cl ₂ C=CCl ₂	67	90
13	7b	$2-MeOC_{6}H_{4}CO_{2}H(1.0)$	Cl ₂ C=CCl ₂	80	85
14	7b	PhCO ₂ H (0.5)	Cl ₂ C=CCl ₂	77	86
15	7b	PhCO ₂ H (2.0)	Cl ₂ C=CCl ₂	69	94
16	7b	PhCO ₂ H (5.0)	Cl ₂ C=CCl ₂	40	91

^a Unless otherwise noted, the reaction was carried out using **1c** (1.0 mmol), **6** (1.0 equiv), catalyst **7a** or **7b** (0.1 equiv), and 4Å MS at room temperature for 24 h. ^b Isolated yield after treatment with TMSCHN₂. ^c Estimated by chiral HPLC after treatment with TMSCHN₂.

The observed product inhibition implies that the resting state shifts from complex **B** to intermediate **int** as the reaction proceeds. Due to additional noncovalent interactions between the product and the catalyst, the **int** state may indeed be thermodynamically more stable than complex **B** and therefore it may become more populated at higher conversions.²³

The proposed mechanism is consistent with the kinetic data and underlines the importance of the second coordinated carboxylate unit in the present stereoselective transformation. The revealed mechanism suggests that the addition of different unreactive carboxylic acids, such as benzoic acid, as Brønsted acids/bases might improve the reaction rate and enantioselectivity by altering the 1,4-addition and/or proton transfer steps.

To test this hypothesis, we investigated the effect of a variety of carboxylic acids as additives, and the reoptimization of solvent and catalyst. Using the asymmetric synthesis of sitagliptin, an anti-diabetic drug,^{3b,24} as the aim, the reaction of *O*benzylhydroxylamine **6** with fluorinated carboxylic acid **1c** was screened using a range of carboxylic acid additives in several solvents (Table 1). When the reaction was performed with **7a** or **7b** without any additives in various halogenated solvents (CCl₄, Cl₂C=CCl₂, 4-CF₃C₆H₄Cl), **7b** always gave better enantioselectivities for desired product **2c** than **7a** (entries 1-6). We then examined the effect of additive with **7b** in halogenated solvents. To our delight, adding benzoic acid (1 equiv) significantly enhanced the *ee* of **2c**, irrespective of the solvent used (entries 7–9). For further screening, tetrachloroethylene was selected as a solvent to examine types and amounts of additives. Electron-rich and electron-poor benzoic acids, and AcOH, did not surpass benzoic acid in terms of enantioselectivity (entries 10–13), but did support the proposed reaction mechanism via complex **B** (Scheme 4), where electron-rich and less-bulky additives competed with 1 to form three ternary complexes **B**, such as 7b•1₂, 7b•1•(additive), and 7b•(additive)₂, composed of two carboxylates. Consequently, the predominant generation of homodicarboxy borate complex 7b•(additive)₂ lowered the yield of **2**, and hetero-dicarboxy borate complex 7b•1•(additive) resulted in either higher or lower ee values for **2**, depending on the additive. Increasing the amount of benzoic acid additive led to a lower yield, for the same reason as described above, together with a slight improvement in selectivity (entries 14–16), in agreement with the proposed mechanism.

With optimized reaction conditions in hand, we further explored the substrate scope, as shown in Scheme 5. Notably, the enantioselectivities of products **2d-2i** (90-94% ee) were greatly improved by adding benzoic acid in comparison with the previous outcomes (68-86% ee), while the product yields were not significantly affected.

Scheme 5. Improved substrate scope of optimized conditions



 $^{\rm b}$ The results with ${\bf 7a}$ in the absence of ${\rm PhCO_2H}$ were indicated in the parentheses

To gain insight into the origin of enantioselectivity, we carried out DFT calculations for the stereogenic C-N bond formation step of the reaction.²⁵ The particular system we considered involves catalyst **7b**, reactants **1b** and **6**, and benzoic acid as the additive. Based on the results of kinetic analysis (i.e. first order kinetic dependence on both acid **1** and nucleophile **6**), we assumed that the C-N addition step is coupled with the protonation process, which occurs via the involvement of an extra carboxylic acid (benzoic acid in the present model).



Figure 6. Acid-assisted C-N bond formation transition states leading to (*S*) and (*R*) product enantiomers in the aza-Michael reaction of 6 and 1b. Relative stabilities are given in kcal/mol with respect to $7b\cdot1b\cdot(PhCO_2) + 6 + PhCO_2H$. H-bonding interactions are indicated as dotted lines. The developing C-N bond is shown as a blue dotted line. All C–H hydrogen atoms are omitted for clarity. The C atoms of the reacting partners are highlighted in green.

Transition states corresponding to the addition of nucleophile 6 to ternary complex $7b\cdot 1b\cdot (PhCO_2)$ (complex **B** in Scheme 4) in the presence of a benzoic acid were explored computationally and the most stable structures leading to (S) and (R) product enantiomers are depicted in Figure 6. In intermediate $7b \cdot 1b \cdot (PhCO_2)$, the electrophile is attached to the catalyst via a covalent B-O bond and its orientation is fixed by a double Hbond formed with the thiourea. The benzoyloxy ligand is anchored via an intramolecular O-H···O bond, such that the oxygen atom that activates the nucleophile is positioned above the Si face of the crotonate double bond. This structural arrangement in 7b·1b·(PhCO₂) enables nucleophile activation (via N-H…O(benzoate) H-bonding interaction) only for Si-face attack of the crotonate (see **TS-S** in Figure 6). In this transition state, the benzoic acid is bound to the second NH group of the nucleophile via the carbonyl moiety, whereas the acidic OH group is oriented towards the proton acceptor carbon atom of the crotonate. The calculations reveal that the C-N addition and the proton transfer processes are indeed coupled, as transition state TS-S represents a concerted asynchronous mechanism, wherein the acid-assisted C-N addition occurs first and then followed by a double proton transfer event (concerted protonation of the substrate by the acid OH and deprotonation of the NH₂ group by the acid carbonyl).²⁶ An analogous transition state was identified computationally for the Re-face attack of the crotonate as well, however, in that transition state (TS-R in Figure 6), the nucleophile is not activated due to the lack of N-H…O(benzoate) interaction. The revealed mechanism, which identifies the irreversible C-N addition/protonation event as the enantioselectivity-determining step, is in accordance with the kinetic data and it also accounts well for the observed enantioselectivity (the barriers calculated for the C-N bond formation step for the Si and Re pathways are 24.5 and 27.1 kcal/mol, respectively).

Scheme 6. Synthetic application to sitagliptin



To demonstrate the applicability of the established aza-Michael addition, the asymmetric synthesis of sitagliptin was carried out on a gram scale (Scheme 6). In this case, 4-CF₃C₆H₄Cl (PCBTF), which has been used in chemical manufacture due to its high boiling point and recyclability, was adopted as the solvent instead of CCl₄ and Cl₂C=CCl₂. The reaction of **1c** (1.08 g) with BnONH₂ (1.2 equiv) in the presence of (*S*,*S*)-**7b**, an enantiomer of **7b**, gave the desired ester **2c-[OMe]** (1.46 g) in two steps without suffering from a decrease in enantioselectivity (91% ee). Following to the procedure developed by Ishihara,¹¹⁰ the obtained adduct **2c** was converted into corresponding amide **10** by catalytic amidation with bicyclic amine **D** without using conventional condensing reagents. Successive subjection of amine **10** to Pd/C-mediated hydrogenolysis and Boc-protection furnished the known *N*-Boc derivative of sitagliptin in good yield. The synthetic compound was identical to an authentic sample by a comparison of spectral data.^{24c}

Having clarified the mechanism of asymmetric aza-Michael addition and having optimized the reaction conditions in terms of additive and solvent, we next attempted to synthesize optically active chroman derivatives, which are common in natural products and biologically active compounds (Scheme 7). Among these, chromans bearing tetrasubstituted chiral carbon centers, such as tocopherols,²⁷ MDL-73404,²⁸ and clusifoliol,²⁹ have attracted much attention from synthetic chemists due to their unique structure and biological activities. Indeed, several elegant asymmetric syntheses toward α -tocopherol have been reported using intramolecular *O*-alkylation,³⁰ oxidative C-O bond formation,³¹ and an oxa-Michael addition.^{4e,32} However, constructing the chroman scaffold while incorporating the tetrasubstituted carbon center in a stereocontrolled manner remains challenging.³³ In particular, there have been no successful examples of asymmetric oxa-Michael addition to B.Bdisubstituted- α , β -unsaturated carboxylic acid surrogates due to the weak nucleophilicity of the oxygen nucleophile and the low reactivity of the Michael acceptor. We envisioned that our catalysts 7 would recognize the carboxylic acid and phenolic OH of substrate 16 through the thiourea and arylboronic acid, as in complex **B** in Scheme 4, to facilitate stereoselective transformation into desired product 15.

Scheme 7. Strategy for the construction of chroman scaffolds with chiral tetrasubstituted carbon centers



Based on this hypothesis, we first investigated the reaction of α , β -unsaturated carboxylic acid **16a** as a model substrate (Scheme 8). In fact, the reaction of **16a** under dual catalytic conditions previously reported by our group^{12b} gave desired product **15a**, albeit with a low yield and ee. However, when 10 mol% of **7b** was used in PCBTF at 50 °C, the reaction proceeded slowly to afford the desired product in 45% yield with 50% ee after 3 days. To our delight, catalysts **7c** and **7d**, bearing electron-withdrawing groups CF₃ and SF₅, respectively, on the aromatic ring, improved both the yields (94-99%) and enantioselectivities (70-74% ee), while the corresponding urea catalyst **7e**, bearing 3,5-CF₃ substituents, resulted in a lower yield and selectivity. Notably, 4Å MS were again important for achieving high enantioselectivities for the tetrasubstituted chiral carbon center in product 15. For instance, when the reaction of 16a was performed without 4Å MS in the presence of 10 mol% 7c, the opposite enantiomer, (R)-15a, was obtained with 70% ee, albeit in a low yield (28% yield after 3days). We further explored the effect of aryl substituent in the catalysts on the enantioselectivity to suppress the undesired pathway to the (R)-isomer. Of these, catalyst 7f, which possessed an ortho-methyl group on the aromatic ring, afforded the best results (99% yield, 91% ee). However, our additional experiments demonstrated that there is no strong correlation between the reaction rate (or yield) and the electron density of aryl groups. We suspect that the solubility of these catalysts in PCBTF is important for good yields, while rational explanation of these results is still needed (see page S17 in Supporting Information).

Scheme 8. Optimization of reaction conditions



Similar to the aza-Michael addition, the enantioselectivity was further improved to 98% ee when 1 equiv of benzoic acid was added to the reaction mixture (Eq. 2). In contrast, the addition of *ortho*-phthalic acid (1.0 equiv) completely shut down the reaction, affording no desired product (Eq. 3). Double-coordination of the carboxylate to the boron center of **7f** seemed to play a crucial role in the enantioselectivity, as seen for the intermolecular aza-Michael addition. These results strongly suggested that our catalytic system using ternary borate complexes would be generally applicable to a range of asymmetric reactions.



Taking all the results into account, we proposed a plausible reaction mechanism, as shown in Scheme 9. Substrate **16a** and

catalyst **7f** form complex **E** via acid–base interactions, where the linear s-cis conformer would be favorable. In the presence of molecular sieves, another carboxylic acid (substrate or additive) coordinates to the borate complex as a second ligand to give complex F. Although TS-II, adopting the s-trans conformer, seems unfavorable compared to s-cis TS-I, the interaction of the second carboxylate with the phenolic OH group in **16a** makes the 1,4-addition via **TS-II** relatively fast, furnishing product (S)-15a, whose absolute configuration is in good agreement with the experimental results. In this case, the oxa-Michael addition and protonation probably takes place in a stepwise manner, because of the bulkiness of 2-tolyl group of 7f. Even in the absence of molecular sieves, the reaction proceeds via TS-I, where intramolecular hydrogen bonding between the phenolic OH and carbonyl group of the substrate promotes the Michael addition from the Re face without participation of the borate carboxvlate. However, distinct from the cyclization via TS-II, the reaction occurs very slowly because reactive complex \mathbf{F} is not sufficiently produced in the absence of 4Å MS. Moreover, the effect of the ortho-methyl substituent in catalyst **7f** can be neatly explained by its repulsion of the phenol ring in the substrate in TS-I, affording (S)-15a in higher ee through the predominant reaction via **TS-II**.

Scheme 9. Possible mechanism explaining the effect of 4Å MS and the *ortho*-Me group in catalyst 7f



Finally, the established oxa-Michael addition was applied to the formal asymmetric synthesis of tocopherol (Scheme 10). Starting from known lactone 17,³⁴ Weinreb amide 19 was synthesized via sulfonamide 18 by successive treatment of 17 with Tf₂O and MeONHMe/AlMe₃. After converting 19 to its corresponding MOM ether, the obtained product was subjected to methylation with MeMgBr to afford ketone 20, which was then transformed into α,β -unsaturated carboxylic acid (E)-16b in 72% yield using the Horner-Wadsworth-Emmons reaction and acid hydrolysis. As expected, the key intramolecular oxa-Michael reaction of 16b took place in an enantioselective manner, giving cyclized adduct 15b in good yield. However, the product could not be isolated in a pure form due to contamination of the additive. Oxa-Michael adduct 15b was converted into known compound 22 by treatment with LiAlH₄ via reduction of the carboxy group together with deprotection of the triflate group. Compound 22 had moderate to high ee values depending on the solvent and additive, as shown in Scheme 10. Under the best conditions (1.0 equiv of crotonic acid additive in Cl₂C=CCl₂ at 50 °C), target compound **22** was obtained in 70% yield (over two steps) with 92% ee. The absolute configuration of **22** was determined to be *S* by comparing its specific optical rotation value with the literature value.^{27b} As obtained chromanethanol **22** has been successfully transformed into α -tocopherol^{27a} and MDL-73404,^{28b} we had achieved a concise formal asymmetric synthesis of these molecules in only eight steps involving the asymmetric oxa-Michael addition using our original thiourea catalysts.



Scheme 10. Asymmetric synthesis of key intermediate 22 toward chroman derivatives

Conclusion

We have developed multifunctional chiral thiourea catalysts comprised of a tertiary amine and arylboronic acid. These hybrid catalysts (7) can form 1:2 complexes containing two equivalents of homo- or hetero-carboxylic acids, which concurrently activate the carboxy group of the coordinated substrate and heteroatoms of nucleophiles through multiple hydrogen bonding interactions, in a similar manner to enzymes. The resulting ternary complexes accelerate asymmetric intermolecular aza-Michael addition of N-benzylhydroxylamine and asymmetric intramolecular oxa-Michael addition of phenol. The products of both reactions can be efficiently applied to the asymmetric synthesis of pharmaceutically important molecules on either a gram-scale or via short-step routes. Spectral data and kinetic studies showed that molecular sieves play a critical role in the double coordination of carboxylic acids to catalysts, leading to the formation of catalytically active ternary borate complexes. These ternary borate complexes, comprising thiourea, arylboronic acid, and carboxylates, function as both Brønsted and Lewis acids, and Brønsted base, which could be supported by computations. We think that this concept will help to develop a novel catalytic system for the selective activation of carboxylic acids, for which research is now underway in our laboratory.

ASSOCIATED CONTENT

Supporting Information

The supporting information is available free of charge on the ACS Publication website at DOI:

Part I: experimental procedures, characterization data, ¹¹B NMR studies, ESI-Mass spectra, kinetic studies, copies of ¹H, ¹³C NMR spectra, and copies of HPLC analyses (PDF)

Part II: mechanistic studies using DFT calculation (PDF)

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Notes

The authors declare no competing financial interest.

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