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Organocatalytic synthesis of chiral tetrasubstituted allenes from racemic propargylic alcohols

Deyun Qian^{1,2}, LinLin Wu¹, Zhenyang Lin¹ & Jianwei Sun^{1,2,3}

Although chiral allene preparation via formal S_N2' nucleophilic substitutions of enantioenriched propargylic derivatives or metal-catalyzed reactions of racemic propargylic derivatives has attracted considerable attention and found applications in many areas of research, direct use of propargylic alcohols instead of propargylic derivatives for catalytic asymmetric allene synthesis is unknown. Here, we show that a highly enantioselective synthesis of tetrasubstituted allenes from racemic propargylic alcohols has been realized by organocatalysis with good efficiency (up to 96% yield and 97% ee). The intermolecular C-C and C-S bond formation was achieved efficiently with simultaneous stereocontrol over the axial chirality. Furthermore, an adjacent quaternary stereocenter could also be constructed. Mechanistically, the reaction may involve efficient stereocontrol on the propargylic cation by its chiral counter anion or 1,8-conjugate addition of *para*-quinone methides. In sharp contrast to previous central chirality construction, this process employs quinone methides for axial chirality construction.

¹ Department of Chemistry, the Hong Kong University of Science and Technology (HKUST), Clear Water Bay, Kowloon, Hong Kong SAR, China. ² The Hong Kong Branch of Chinese National Engineering Research Center for Tissue Restoration and Reconstruction, the Hong Kong University of Science and Technology (HKUST), Clear Water Bay, Kowloon, Hong Kong SAR, China. ³ HKUST Shenzhen Research Institute, Shenzhen 518057, China. Correspondence and requests for materials should be addressed to Z.L. (email: chzlin@ust.hk) or to J.S. (email: sunjw@ust.hk)

xially chiral allenes are well-known subunits widely present in natural products, bioactive molecules, ligands, organocatalysts, and functional materials as well as versatile chiral building blocks in organic synthesis¹⁻⁵. Their orthogonal cumulative π -systems often lead to complementary and unique reactivity compared with olefins and alkynes. However, the development of efficient methods to access chiral allenes from readily available starting materials has not been able to fulfill the need for their utilization in organic synthesis. Since Maitland and Mills reported the first chiral allene in 1935⁶, substantial endeavors have been made toward the stereoselective synthesis of axially chiral allenes. The well-established approaches mainly rely on either resolution of racemic allenes or chirality transfer from enantioenriched propargylic derivatives (Fig. 1a). In contrast, direct construction of the axial chirality of allenes from achiral or racemic substrates via asymmetric induction by a chiral catalyst remains substantially challenging, though highly attractive.

Recently, some progress has been made in catalyst-controlled asymmetric synthesis of chiral allenes based on metal catalysis^{7–11} and organocatalysis^{12–19}. However, unfortunately these methods are largely limited to the formation of di- or trisubstituted allenes^{1–19}. In particular, organocatalytic approaches have been mainly based on asymmetric protonation or deprotonation (Fig. 1a).

The lack of efficient catalytic strategies^{20–23} for the synthesis of tetrasubstituted chiral allenes is by no means an indication that these molecules are less important. Indeed, as shown in Fig. 1b, tetrasubstituted chiral allenes have been known as key units in useful chiral catalysts²⁴, ligands²⁵, important intermediates toward natural products^{4, 26}, and molecular materials³. In fact, the challenges lie in simultaneous construction of a non-hydrogen- involved bond and establishment of the axial chirality, in which the catalyst must be effective in controlling both events^{8, 16–18}.

Results

Design plan. The direct transformation of propargylic alcohols or their derivatives to chiral allenes is one of the most straightforward and efficient strategies^{1–5}. Specifically, chirality transfer from enantioenriched propargylic derivatives bearing a suitable leaving group by means of formal S_N2' reaction represents the most studied and reliable approach (Fig. 2a)⁵. However, the required pre-establishment of the central chirality in the starting

material is a major limitation preventing its large scale applications. Racemization is another issue that could occur in many cases. Recently, Ma and co-worker reported a catalytic asymmetric synthesis of chiral 2,3-allenoates from racemic propargylic carbonates⁹. Mechanistically, dynamic kinetic resolution (DKR) of the crucial allenic metal intermediates plays a key role in achieving both high efficiency and enantioselectivity by catalyst control, thereby representing a pioneering study of this type (Fig. 2b). However, this elegant approach is limited to the synthesis of disubstituted allenoates. Indeed, catalytic asymmetric synthesis of fully substituted allenes from similar racemic propargylic derivatives, in particular propargylic alcohol itself, remains unknown. Inspired by this pioneering study and prompted by our continued interest in allene synthesis^{14, 15}, we

a Chirality Transfer (via formal S_N2', well-studied)



b Metal Catalysis by Ma (via DKR of the allenic metal intermediates)







Fig. 2 Chiral allene synthesis from propargylic compounds. **a** Wellestablished formal SN2' reactions from enantioenriched propargylic derivatives. **b** A rare metalyzed-catalyzed example from racemic propargylic derivatives. **c** Our design of an organocatalytic approach from racemic propargylic alcohols



Fig. 1 Strategies for the asymmetric synthesis of allenes and representative axially chiral allenes. **a** Previous approaches to the enantioselective synthesis of allenes, which are largely limited to the formation of di- or trisubstituted allenes. Compared with other methods, organocatalytic approaches still remain underexplored and are largely limited to allenaoates. **b** Useful chiral tetrasubstituted allenes and an allene bearing adjacent quaternary stereocenter



^bDetermined by HPLC with a chiral stationary phase

^cNo reaction ^dCvclohexane/CCl₄ = 1:1

^eRun at 0 °C

^fRun at −20 °C

ga was obtained in 92% isolated yield

envisioned that, in the presence of a chiral Brønsted acid^{27–29}, propargylic alcohols may generate a propargylic cation paired with a chiral counter anion (Fig. 2c)^{30–32}. Subsequent nucleophilic attack may generate an allene in a stereocontrolled trajectory governed by the chiral anion. If successful, this formidably challenging process would provide an attractive approach for efficient synthesis for fully substituted chiral allenes.

Diastereo- and enantioselective C-C bond formation. We began to test our hypothesis using the racemic alcohol 1a as the propargylic cation precursor and 1,3-diketone 2a as the nucleophile. Chiral phosphoric acids and their analogues were employed as potential catalysts in view of their demonstrated versatility in asymmetric induction in chiral anion catalysis^{27–32}. In the presence of the highly acidic N-triflylphosphoramide (R)-A1, the reaction proceeded smoothly at room temperature to provide chiral allene 3a in high yield with good enantioselectivity within a few minutes (Table 1, entry 1). When using (R)-A2 as catalyst, however, the reaction could not take place, indicating the acidity of chiral Brønsted acids plays an important role in our system. These results encouraged us to further evaluate different Brønsted acids for the transformation. As shown in Table 1, the electronic properties and the steric environment as well as the chiral backbone of the catalysts have very strong influence on the diastereoselectivity and enantioselectivity (Table 1, entries 1-6). Catalyst (R)-A1 provided the best results in terms of conversion and enantioselectivity (80% ee) (Table 1, entry 1). After optimizing other reaction parameters, including solvent (Table 1, entries 7-15), temperature (Table 1, entries 16-17), etc. (more details in Supplementary Tables 1–3 of the Supplementary Information), we identified that the axially chiral allene 3a could be obtained in 92% isolated yield with 90% ee and 6.1:1 dr (Table 1, entry 17) when the reaction was run at -20 °C with catalyst (R)-A1 (5 mol%) and solvent CCl₄ (0.05 M). It is noteworthy that the reaction not only furnished the axially chiral allene unit, but also simultaneously established an adjacent all-carbon quaternary stereocenter, which is a significant challenge in asymmetric synthesis^{33, 34}. Notably, quaternary stereocenters are widely found in biologically important allenic natural products (e.g., Grasshopper ketone, Fig. 1b)¹.

With the optimized conditions, next we examined the scope of this process. As shown in Fig. 3, a variety of racemic propargylic alcohols bearing electron-rich, electron-neutral, and electron-deficient aryl moieties all smoothly participated in the efficient intermolecular C-C bond formation process. The corresponding tetrasubstituted chiral allenes were all formed in high yield with good diastereoselectivity and good to excellent enantioselectivity (62-96% yield, up to 21:1 dr, 77-97% ee). Heterocycles could also be incorporated into the chiral allene products (**3c**, **3o**, **p**). Cyclic and acyclic 1,3-diketones as well as β -ketoesters were also suitable

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Fig. 3 Allene formation with 1,3-dicarbonyl nucleophiles. Reaction conditions: **1** (0.2 mmol), **2** (0.3 mmol for **3a-3j**; 0.6 mmol for **3k-3p**), (*R*)-**A1** (0.01 mmol), CCl₄ (4.0 mL), -20 °C (**3a-3g**) or 0 °C (**3k-3p**), 0.5-30 h. *Isolated yield. [†]Determined by ¹H NMR analysis. [‡]Determined by HPLC with a chiral stationary phase. [§]Run at room temperature with anhydrous MgSO₄ (200 mg) as additive. [#]Run at -15 °C. [¶]Run at 0 °C. *PMP p*-MeOC₆H₄

nucleophiles (3h-j). It is believed that the 4-methoxyphenyl group $(R^2 = PMP)$ assists the generation and stabilization of the important cation intermediate. It also provides sufficient differentiation from the *gem*-substituent (R^3) in the subsequent enantioselective C-C bond formation. In addition to PMP, other substituents, such as *p*-allyloxyphenyl (3d), *p*-sulfenylphenyl (3e), and *p*-hydroxyphenyl (3k-p) groups, can also serve the same role. The absolute configuration of product 3n was unambiguously confirmed by X-ray crystallography.

Enantioselective C-S bond formation. It is worth noting that in the above scope study allenes bearing a phenol unit 3k-3p were not only smoothly obtained, but also formed with higher reaction rate and generally better enantiocontrol. The difference in reactivity and enantioselectivity was more pronounced with thioacetic acid as nucleophile (Fig. 4, Eq. 1). We reasoned that the mechanisms might be different for these two types of substrates (with or without free hydroxy groups). In the absence of a free hydroxy group, the key intermediate should be a cation bearing a

chiral counter anion (e.g., OX), as initially proposed. In contrast, in the presence of a free hydroxy group, the initially formed cation could induce proton loss to form a neutral para-quinone methide intermediate $(\mathbf{QM})^{35-39}$, which was indeed isolated and fully characterized when the substrate was treated with the catalyst in the absence of nucleophile (for details, see the Supporting Information). The p-QM can be activated by the chiral Brønsted acid catalyst for subsequent nucleophilic attack. Notably, both cases require remarkable remote stereocontrol and regiocontrol, and the latter situation is indeed an asymmetric 1,8-conjugate addition. It is worth noting that although catalytic asymmetric 1,4- and 1,6-conjugate additions have received tremendous attention, efficient catalytic asymmetric 1,8-addition has remained almost unknown and this field is still in its infancy 40-43. Furthermore, to the best of our knowledge, the focus of previous efforts on asymmetric reactions of quinone methides all centered on the establishment of central chirality³⁵⁻³⁹, while this reaction establishes axial chirality with quinone methides.

Encouraged by these results, we next focused on the asymmetric synthesis of hetero-substituted allenes. The presence

of a heteroatom can not only bring new synthetic and biological values, but also lead to potential useful interactions by coordination or hydrogen-bonding when needed. Therefore, the design of new enantioselective synthetic strategies to access hetero-substituted chiral allenes is highly desirable. To the best of our knowledge, examples of catalytic asymmetric synthesis of tetrasubstituted allenes bearing a hetero-substituent are essentially unknown^{1–25, 44}.

While the previous standard conditions with catalyst A1 could not lead to good enantioselectivity for the intermolecular C-S



Fig. 4 The rate difference between **1i** and **1i**' with thioacetic acid nucleophile and the proposed intermediates. When $R \neq H$, the reaction may proceed via ion pair OX, while when R=H, the para-quinone methide **QM** activated by hydrogen bonding might be involved

bond formation with thioacid nucleophiles, fortunately, we were able to improve the reaction selectivity after considerable optimization efforts. As shown in Table 2, other additional acid catalysts were evaluated. Among them, the spirocyclic acid B1 provided the best combination of catalytic activity and asymmetric induction (73% vield, 81% ee, entry 4). Further solvent screening indicated that toluene can slightly improve the yield and enantioselectivity. However, the reactivity was shut down in Et₂O, presumably due to competing binding of ether with the catalyst (entry 6). Next, we found that the use of 3 Å molecular sieves could further improve the enantioselectivity (entry 8), while in drastic contrast, 4 Å molecular sieves resulted in almost no product formation (entry 9). Finally, evaluation of other parameters, such as concentration and temperature, identified the optimal conditions that gave both excellent enantioselectivity and good yield (entry 11).

With the optimized conditions, we next examined the scope of the C-S bond formation process. As shown in Table 3 and Fig. 5, a wide range of chiral tetrasubstituted allenes bearing a sulfur substituent can be prepared with good efficiency and excellent enantioselectivity. Various substituents, including aryl, alkenyl, alkyl, and hetero substituents, at different positions of the substrates did not affect the good reactivity and stereocontrol (51-96% yield, 88-94% ee). Notably, steric hindrance on the alkyne, where the nucleophile approaches, did not affect the reaction $(5\mathbf{u}, \mathbf{v})^8$. Substitution with heterocycles proved suitable $(5\mathbf{f}, \mathbf{g})$. Furthermore, the mild conditions can tolerate a range of functional groups, such as ethers $(5\mathbf{b}, 5\mathbf{w}-\mathbf{x})$, aromatic and aliphatic halides $(5\mathbf{c}, \mathbf{d}, 5\mathbf{r})$, silanes and silyl-protected alcohols $(5\mathbf{k}, \mathbf{l}, 5\mathbf{q})$, olefins $(5\mathbf{o})$, alkynes $(5\mathbf{p}, 5\mathbf{w})$, nitriles $(5\mathbf{s})$, etc. Furthermore, this protocol can also be applied to late-stage

Table 2 Optimization of the reaction conditions using thioacetic acid 4a as nucleophile								
	н	Ph OH Ph Ph + AcSH 1i (0.05 mmol)	B*H (5 mol%) Solvent (0.1 M) rt HO	Ph SAc Ph 5a				
		A1: R = 2,4,6 X = NHTF A2: R = 2,4,6 X = 0.4 X	$\frac{1}{Pr_3C_6H_2}$, $\frac{1}{Pr_3C_6H_2}$, = OH	,4,6- [/] Pr ₃ C ₆ H ₂ P ^O X 4.6- [/] Pr ₃ C ₆ H ₂				
		R A7: R = 9-ant X = OH	hryl B1: X = NH	Tf; B2 : X = OH				
Entry	B*H	R A7: R = 9-ant X = OH	hryl 52. B1: X = NH	Tf; B2 : X = OH Yield (%) ^a	ee (%) ^b			
Entry	<u>B*H</u> A1	A7: R = 9-ant R A7: R = 9-ant X = OH Solvent CH2Cl2	hryl B1: X = NH t (h) 0.5	Tf; B2 : X = OH Yield (%) ^a 90	ee (%)^b 43			
Entry 1 2	<u>B*H</u> A1 A2	A7: R = 9-ant R A7: R = 9-ant X = OH X = OH CH2Cl2 CH2Cl2 CH2Cl2 CH2Cl2	hryl B1: X = NH t (h) 0.5 18	Tf; B2 : X = OH Yield (%) ^a 90 62	ee (%)^b 43 57			
Entry 1 2 3	B*H A1 A2 A6	A7: R = 9-ant X = OH Solvent CH ₂ Cl ₂ CH ₂ Cl ₂ CH ₂ Cl ₂ CH ₂ Cl ₂	hryl B1: X = NH t (h) 0.5 18 12	Tf; B2 : X = OH Yield (%) ^a 90 62 40	ee (%)^b 43 57 7			
Entry 1 2 3 4	B*H A1 A2 A6 B1	A7: R = 9-ant X = OH Solvent CH ₂ Cl ₂ CH ₂ Cl ₂	hryl B1: X = NH 0.5 18 12 0.3	Trf; B2 : X = OH Yield (%) ^a 90 62 40 73	ee (%)^b 43 57 7 81			
Entry 1 2 3 4 5	B*H A1 A2 A6 B1 B2	$\begin{array}{c} \textbf{A7: } \textbf{R} = 9\text{-ant}\\ \textbf{X} = 0\text{H}\\ \end{array}$	hryl B1 : X = NH t (h) 0.5 18 12 0.3 24	Tf; B2 : X = OH 90 62 40 73 51	ee (%)^b 43 57 7 81 75			
Entry 1 2 3 4 5 6	B*H A1 A2 A6 B1 B2 B1	$\begin{tabular}{ c c c c c } \hline A7: R = 9-ant \\ X = OH \\ \hline \\ $	hryl B1 : X = NH t (h) 0.5 18 12 0.3 24 12	Tf; B2: X = OH 90 62 40 73 51 <5	ee (%) ^b 43 57 7 81 75 -			
Entry 1 2 3 4 5 6 7	B*H A1 A2 A6 B1 B2 B1 B1 B1	R $A7: R = 9-ant X = OH$ $X = OH$ CH_2CI_2 CH_2CI	hryl B1 : X = NH 0.5 18 12 0.3 24 12 1	Yield (%) ^a 90 62 40 73 51 <5	ee (%) ^b 43 57 7 81 75 - 85			
Entry 1 2 3 4 5 6 7 8 ^c	B*H A1 A2 A6 B1 B2 B1 B1 B1 B1 B1 B1	$\begin{tabular}{ c c c c c } \hline A7: R = 9-ant \\ X = OH \\ \hline \\ $	hryl B1 : X = NH t (h) 0.5 18 12 0.3 24 12 1 5	Yield (%) ^a 90 62 40 73 51 <5	ee (%) ^b 43 57 7 81 75 - 85 89			
Entry 1 2 3 4 5 6 7 8 6 9 d	B*H A1 A2 A6 B1 B2 B1 B1 B1 B1 B1 B1 B1 B1	$\begin{tabular}{ c c c c c } \hline A7: R = 9-ant \\ X = OH \\ \hline \\ $	hryl B1 : X = NH t (h) 0.5 18 12 0.3 24 12 1 5 12	Тf; B2: X = OH Уield (%) ^a 90 62 40 73 51 <5 93 87 <5	ee (%)^b 43 57 7 81 75 - 85 85 89 -			
Entry 1 2 3 4 5 6 7 8 c 9 d 1 0 c e	B*H A1 A2 A6 B1 B2 B1 B1 B1 B1 B1 B1 B1 B1 B1	$\begin{tabular}{ c c c c c } \hline A7: R = 9-ant \\ X = OH \\ \hline X = OH \\ \hline \\ $	hryl B1 : X = NH t (h) 0.5 18 12 0.3 24 12 1 5 12 6	Yield (%) ^a 90 62 40 73 51 <5	ee (%) ^b 43 57 7 81 75 - 85 89 - 90			

^dRun 4 Å MS (15 mg)

^ec = 0.05 M ^fRun at -5 °C



Fig. 5 Formation of allenes **5w**-z. Reaction was carried out with **1** (0.3 mmol), **4** (0.6 mmol), catalyst **B1** (5 mol%), and 3 Å MS (90 mg) in solvent (6.0 mL, CCl₄ for **5w**, **x** and CH₂Cl₂ for **5y**, **z**) at -5 °C. *Run without 3 Å MS. [†]Run with **A7** at -15 °C

installation of a chiral allene unit to an estrone-derived complex molecule (5 h), in which high diastereomeric excess was observed as a result of stereocontrol by the chiral catalyst. In addition to thioacetic acid, thiobenzoic acid (BzSH) can also serve as a suitable nucleophile with slight modification of the conditions. For example, with catalyst (R)-A7, the reaction of 1 u with BzSH smoothly afforded the corresponding chiral allene 5z with high efficiency and enantioselectivity (90% ee). It is remarkable that the stereogenecity was efficiently relayed over eight chemical bonds. It is worth noting that the excellent generality of the process also benefits largely from the compatible in situ generation of the unstable p-QMs.

Product transformations. To further demonstrate the utility of the chiral allene products from our protocol, we carried out a preparative-scale synthesis of allene **5v** and some transformations. To our delight, comparably high efficiency and enantioselectivity (0.6 g, 90% yield, 92% ee) were observed (Fig. 6a, Eq. 2). Moreover, catalyst **B1** could be recycled without loss of catalytic activity. These results suggest that our reaction is amenable to



Fig. 6 Representative product transformations and mechanistic studies. **a** Triflation of the free hydroxy group (Eq. 2), transformation of the thioacetate moiety to a sulfone unit (Eq. 3), and NIS-mediated cyclizations of the chiral tetrasubstituted allenes proceed via the axial-to-central chirality transfer (Eqs. 4-5). **b** Important molecules bearing chiral indene or indane units. **c** Control experiments and possible mechanism. The reactions suggest that the 1,8-addition to *p*-QM is rate-determining (Eq. 6) and the conjugate addition step is irreversible (Eq. 7). The product stereochemistry was purely determined by the chiral catalyst, regardless of the absolute configuration of substrate (Eqs. 8-9), which rules out the possible S_N2' pathway (via **TS1**). **d** Kinetic study indicated that the reaction is first-order in catalyst, suggesting that one catalyst molecule is involved in the rate-determining transition state

Table 3 Allenes 5a-v formation with thioacetic acid								
HO HO 1 (racemic) HO HO HO HO HO HO HO H HO H HO H HO H		(S)- B1 (5 mol%) Solvent (0.05 M), <i>t</i> –5 °C, 3A MS		Ph R ¹ Ho 5				
Entry	R ¹	<i>t</i> (h)	5	Yield	ee			
	N		J	(%)	(%)			
1	Ph	12	5a	82	94			
2^{a}	$(p-MeO)C_6H_4$	3	5b	75	91			
3 ^b	$(p-F)C_6H_4$	18	5c	83	92			
4 ^b	$(p-Cl)C_6H_4$	24	5d	90	92			
5 ^b	$(3,5-Me_2)C_6H_3$	15	5e	93	92			
6 ^a	2-Furyl	1.5	5f	75	91			
7 ^a	3-Thienyl	3	5g	88	90			
8	H H H H H H	10	5h	76	89 de			
9	Cyclohexenyl	10	5i	70	91			
10	-(Me)C=CH ₂	6	5j	72	88			
11	TMS	36	5k	70	92			
12 ^c	TES	10	51	90	90			
13	<i>n</i> Bu	3	5m	95	92			
14	-(CH ₂) ₂ Ph	5	5n	98	94			
15	Allyl	6	50	92	93			
16 ^c	-(CH ₂) ₆ C≡C	12	5p	64	91			
17°	-(CH ₂) ₂ OTBS	12	5q	82	93			
18	-(CH ₂) ₃ Cl	6	5r	96	93			
19°	-(CH ₂) ₃ CN	7	5 s	78	92			
20	Cyclopropyl	12	5t	85	92			
21	<i>i</i> Pr	10	5u	90	92			
22	<i>t</i> Bu	12	5v	89	94			

Reaction was carried out with 1 (0.3 mmol), thioacetic acid 4a (0.6 mmol), catalyst B1 (5 mol %), and 3 Å MS (90 mg) in solvent (6.0 mL, toluene for 5a-h and CH_2Cl_2 for others, or otherwise noted) at -5 °C

^bB1 (7.5 mol%) and thioacetic acid (0.75 mmol)

^cRun without 3 Å MS

large-scale application. Triflation of the free hydroxy group in 5v efficiently provided the corresponding triflate compound, which is poised for further functionalizations, such as cross-coupling⁴⁵. The thioacetate moiety can also be smoothly converted to a sulfone unit (7, Eq. 3). Thus, another family of useful activated allenes, allenyl sulfones, can be made in their enantioenriched

form⁴⁶. Furthermore, the tetrasubstituted allene products can be transformed to other useful molecules, demonstrating the ability of axial-to-central chirality transfer^{1–5}. For example, iodocyclization of allene **3b** in the presence of NIS and MeOH furnished chiral spirocyclic molecule **8** (Eq. 4). In addition, chiral indene derivative **9** can be obtained from allene **5o**. Notably, chiral indene/indane is an important subunit in medicinally important molecules (Fig. 6b)^{47–49}.

Mechanistic studies. To gain further insight into the reaction mechanism, we carried out some control experiments. Firstly, subjecting the fully characterized pure QM to the standard conditions led to successful formation of chiral allene 5a in essentially the same yield and enantiomeric excess as the standard protocol from 1i (Fig. 6c, Eq. 6). Indeed, in the standard reaction, we could observe rapid consumption of 1i followed by accumulation and slow disappearance of QM. These observations suggest that the 1,8-addition to p-QM is rate-determining. Moreover, when the allene product 5a (either racemic or enantioenriched) was treated with catalyst B1, no change in enantiopurity was observed, indicating that the conjugate addition step is irreversible (Eq. 7). Furthermore, the enantiopure tertiary alcohol substrate 1i was subjected to the standard conditions (Eqs. 8-9). At partial conversion, the recovered substrate 1i remained enantiopure, indicating the first step (dehydration) is irreversible, presumably due to the elimination of water by molecular sieves. Without molecular sieves, essentially the same results were obtained (1i: >99%) ee) along with some enone by products generated through the classical Meyer–Schuster rearrangement $^{\rm 50}.$ Moreover, it was found that the product stereochemistry was purely determined by the chiral catalyst, regardless of the absolute configuration of substrate 1i (Eqs. 8-9). This observation is consistent with the intermediacy of p-QM. It also rules out the possible $S_N 2'$ pathway (via TS1), in which DKR might be operative¹⁻⁵. Finally, kinetic study indicated that the reaction is first-order in catalyst, suggesting that one catalyst molecule is involved in the ratedetermining transition state (Fig. 6d). Thus, we proposed a possible transition state (TS2, Fig. 6c), in which the catalyst plays a bifunctional role for activation of both partners and remote stereocontrol (see the Supplementary Note 4 for preliminary DFT calculations).

Discussion

In summary, we have developed an efficient and unified method for catalytic asymmetric synthesis of chiral tetrasubstituted allenes from racemic propargylic alcohols. It is not only an early demonstration of highly enantioselective chiral ion-pair catalysis involving a formal propargylic carbocation as well as catalytic asymmetric 1,8-conjugate addition, but also a new addition to the small family of catalytic syntheses of chiral hetero- and tetrasubstituted allenes. The chiral Brønsted acid catalysts of proper choice not only allow in situ generation of the reactive intermediates, thereby simplifying substrate preparation and expanding functional group compatibility, but also ensure excellent remote stereocontrol. Control experiments provided important insights into the reaction mechanism. The enantioenriched allene products are also precursors to other useful chiral molecules.

Methods

General information. Reagents were purchased at the highest commercial quality and used without further purification, unless otherwise stated. Anhydrous dichloromethane, toluene, diethyl ether, and tetrahydrofuran were purified by the Innovative[®] solvent purification system. ¹H, ¹³C, and ¹⁹F nuclear magnetic resonance (NMR) spectra were collected on a Bruker AV 400 MHz NMR spectrometer using residue solvent peaks as an internal standard (¹H NMR: CDCl₃ at 7.26 ppm, CD₂Cl₂ at 5.32 ppm, acetone-*d*₆ at 2.05 ppm; ¹³C NMR: CDCl₃ at 77.0 ppm, CD₂Cl₂ at 54.0 ppm, acetone-*d*₆ at 29.8 ppm). Mass spectra were collected on an

^aRun in CCl₄ solvent

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Agilent GC/MS 5975 C system, a MALDI Micro MX mass spectrometer, or an API QSTAR XL System. Infrared spectra were recorded on Bruker TENSOR 27 spectrometer and reported in terms of frequency of absorption (cm⁻¹). Optical rotations were measured on JASCO P-2000 polarimeter with $[\alpha]_D$ values reported in degrees; concentration (*c*) is in 10 mg/mL. The enantiomeric excess values were determined by chiral high-performance liquid chromatography (HPLC) using an Agilent 1200 LC instrument. Flash column chromatography was performed on silica gel. NMR and high performance liquid chromatography (HPLC) spectra are supplied for all compounds: see Supplementary Figs 1–218. See Supplementary Method and Notes 1–2 for the characterization data of compounds 1–9.

All the racemic products were prepared using the same substrates with the racemic phosphoric acid derived from BINOL as catalyst.

General procedure for the synthesis of axially chiral allenes 3. At room temperature, an oven-dried 10-mL vial was charged with a mixture of the alcohol substrate 1 (0.2 mmol), the carbon nucleophile 2 (0.3 mmol, 1.5 equiv), and CCl₄ (3.8 mL). The reaction mixture was cooled to -20 °C (for 3a-j) or 0 °C (for 3k-p) unless otherwise specified, and a solution of the catalyst (*R*)-A1 (10 µmol, 5 mol%) in CCl₄ (0.2 mL) was added in one portion. The reaction mixture was stirred at the same temperature. The progress was monitored by thin layer chromatography. Upon completion, the crude mixture was directly subjected to silica gel flash chromatography (5–10% EtOAc in hexanes) to afford the pure product 3.

General procedure for the synthesis of axially chiral allenes 5. At room temperature, an oven-dried 10-mL vial was charged with a mixture of the alcohol substrate 1 (0.3 mmol), 3 Å molecular sieves (90 mg), and the solvent (6.0 mL, toluene for **5a-g** and DCM for others, or noted otherwise). The reaction mixture was cooled to -5 °C, and a solution of the catalyst (*S*)-**B1** (0.015 mmol, 5 mol%) in the same solvent (0.2 mL) and thioacetic acid **4a** (0.6 mmol, 42.9 uL) were added in one portion. The reaction progress was monitored by thin layer chromatography. Upon completion, the reaction mixture was directly subjected to silica gel column flash chromatography (10% EtOAc and 1% AcOH in hexanes, the column was pretreated with 10% AcOH in hexanes) to afford the desired product **5**.

Data availability. The X-ray crystallographic coordinates for structures reported in this article have been deposited at the Cambridge Crystallographic Data Centre (CCDC), under deposition number of CCDC 1498348. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via http://www.ccdc.cam.ac.uk/data_request/cif. All other data is available from the authors upon reasonable request.

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Author contributions

D.Q. conceived the project, performed the experiments, and wrote the paper. L.W. and Z.Y. performed preliminary DFT calculations showing the role of the catalyst in the relevant bond activation. J.S. conceived and directed the project and wrote the paper.

Additional information

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