

ARTICLE

DOI: 10.1038/s41467-018-04123-w

OPEN

Stereodivergent synthesis of vicinal quaternaryquaternary stereocenters and bioactive hyperolactones

Haifeng Zheng¹, Yan Wang¹, Chaoran Xu¹, Xi Xu¹, Lili Lin¹, Xiaohua Liu¹ & Xiaoming Feng^{1,2}

Although great success has been achieved in asymmetric Claisen rearrangement for the synthesis of chiral γ , δ -unsaturated carbonyl compounds bearing vicinal tertiary-quaternary stereocenters, the development of asymmetric versions for stereodivergent construction of adjacent quaternary-quaternary stereocenters remains a formidable challenge because of the high steric hindrance. Here we report a catalytic enantioselective dearomatization Claisen rearrangement of allyl furyl ethers catalyzed by chiral N,N'-dioxide-Ni^{II} complex catalysts. A variety of chiral γ , δ -unsaturated carbonyl compounds bearing vicinal quaternary-quaternary stereocenters were obtained with excellent outcomes under mild conditions. Furthermore, we disclosed that by matching the configuration of the catalysts and the alkene unit of the substrates, four stereoisomers of the products could be prepared in excellent yields and stereoselectivities. Finally, the fascination of this strategy was demonstrated by stereo-divergent synthesis of bioactive natural products hyperolactones B, C, and their epimers. A possible catalytic model was proposed to explain the origin of the asymmetric induction.

1

¹ Key Laboratory of Green Chemistry & Technology, Ministry of Education, College of Chemistry, Sichuan University, Chengdu 610064, China. ² Collaborative Innovation Center of Chemical Science and Engineering, Tianjin 300071, China. Correspondence and requests for materials should be addressed to X.L. (email: liuxh@scu.edu.cn) or to X.F. (email: xmfeng@scu.edu.cn)

he Claisen rearrangement of allyl vinyl ethers is one of the most reliable and useful methods for synthesizing γ,δunsaturated carbonyl compounds, valuable intermediates in natural product construction¹⁻⁴. On the basis of chiral Lewis acids⁵⁻¹⁰, Jacobsen's guanidinium salts¹¹, N-heterocyclic carbenes¹², or transition metals^{13–15} catalysts, various of linear and cyclic vinyl units have been transformed via catalytic asymmetric Claisen rearrangement to afford chiral v,δ-unsaturated carbonyl compounds bearing vicinal tertiary-quaternary stereocenters (Fig. 1a). However, the direct and catalytic stereoselective assembly of these compounds with two adjacent quaternaryquaternary stereocenters remains scarce, because of the inherent steric congestion in the formation of C-C bond. To obtain vicinal quaternary-quaternary arrays and accomplish the total synthesis of several natural products, chiral starting materials were usually conducted through this type of reaction. Zhai and co-workers reported the total synthesis of (-)-Jiadifenin, in which the key intermediate was acquired on the basis of Ireland-Claisen rearrangement of chiral acetal precursor 16. Nakamura developed a stereoselective Ireland-Claisen Rearrangement for the synthesis of the CDE ring system of antitumor and chiral building blocks for oxygenated Terpenoids^{17,18}. In regard of catalytic asymmetric methodology, in 2010, the Jacobsen group reported the only catalytic example to produce the vicinal quaternary-quaternary array¹⁹. However, limited substrates were evaluated though good results were obtained, which restricted further application in the total synthesis of natural products. Although great achievement has been made in asymmetric Claisen rearrangement, the direct catalytic stereodivergent accessing to optically pure γ,δ-unsaturated carbonyl compounds bearing two adjacent quaternary stereocenters is yet to be described and would be of great value for

the natural product-oriented asymmetric synthesis. The hyperolactones $A-C^{20,21}$ and (-)-biyouyanagin A^{22} are a family of spirolactone natural products isolated from *Hypericum chinese L*, which possess significant activity against HIV

replication (Fig. 1b). These compounds contain a mutual spirolactone fragment with two chiral vicinal quaternary carbon centers. Besides, the configuration of hyperolactones A, C, and (-)-biyouyanagin A is (S,S), while the configuration of hyperolactone B is (R,S), which varies from the spirocyclic center. Because of their anti-HIV biological activities and unique structures, synthetic chemists have developed many methodologies for their total syntheses^{22–32}. Particularly, Kraus utilized a tandem Claisen rearrangement to realize the synthesis of racemic hyperolactones C^{33} . Though, the yield was only 25% even heating the reaction mixture in a sealed tube at 130 °C for 15 h, which indicated the challenge of the construction of adjacent quaternary stereocenters. Motivated by these initial results, we try to establish a general and efficient asymmetric Claisen rearrangement methodology for the synthesis of the family compounds of hyperolactone. Besides, we envision that by matching the configuration of the chiral catalysts and the substrates, all of the four stereoisomers might be accessible (Fig. 1c).

Herein, we report the application of chiral N,N'-dioxide-Ni^{II} complex catalysts in the catalytic asymmetric dearomatization Claisen rearrangement of allyl furyl ethers. A series of γ,δ -unsaturated carbonyl compounds bearing two adjacent quaternary stereocenters were obtained in up to 99% yield, 19:1 dr, and 99% ee under mild reaction conditions. Moreover, the catalytic reaction shows a broad substrate scope, which may lead to the discovery of new bioactive molecules. Furthermore, four stereoisomers could be obtained and stereodivergent synthesis $^{34-48}$ of natural products hyperolactone B, C could be achieved as well.

Results

Optimization of reaction conditions. Initially, we selected compound **1a** with *E*-configuration as the model substrate to optimize the reaction conditions. The classical catalysts of Cu (OTf)₂/**BOX** and Ni(OTf)₂/**BOX** yielded trace amount of the

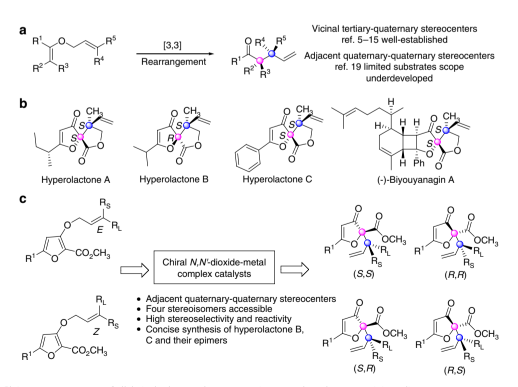


Fig. 1 Asymmetric Claisen rearrangement of allyl vinyl ethers and representative natural products containing adjacent quaternary stereocenters. **a** Previous work for the catalytic asymmetric Claisen rearrangement of allyl vinyl ethers. **b** The structures of natural products hyperolactones A-C and (—)-biyouyanagin A. **c** Our catalytic asymmetric Claisen rearrangement strategy for stereodivergent construction of adjacent quaternary stereocenters

Table 1 Optimization of the reaction conditions^a

Entry	1	Metal source/L	Yield (%) ^b	dr ^c	ee ^d
1	1a	Cu(OTf) ₂ / BOX	trace	-	-
2	1a	Ni(OTf) ₂ / BOX	trace	-	-
3	1a	Yb(OTf) ₃ / L-PiMe₂	47	3:1	33
4	1a	Ni(OTf) ₂ / L-PiMe₂	85	5:1	98
5	1a	Ni(OTf) ₂ / L-PiMe₃	85	5:1	98
6	1a	Ni(OTf) ₂ / L-RaMe₂	90	2:1	94
7	1b	Ni(OTf) ₂ / L-PiMe₂	90	8:1	98
8	1c	Ni(OTf) ₂ / L-PiMe₂	90	5:1	98
9	1b	Ni(BF ₄) ₂ ·6H ₂ O/ L-PiMe₂	96	8:1	98
10 e	1b	$Ni(BF_4)_2 \cdot 6H_2O/L-PiMe_2$	96	10:1	99
11 e	1d	Ni(BF ₄) ₂ ·6H ₂ O/ L-PiMe₂	94	5:1	99
12 e	1e	Ni(BF ₄) ₂ ·6H ₂ O/ L-PiMe₂	90	1.5:1	99

^aUnless otherwise noted, the reactions were performed with 1 (0.1 mmol), and metal source/Ligand (1:1, 10 mol%), in DCE (1.0 mL) at 70 °C for 4 h

eAt 35 °C for 4 days

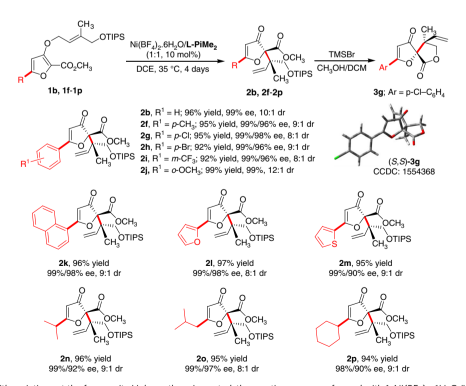


Fig. 2 Substrate scope with variations at the furan unit. Unless otherwise noted, the reactions were performed with 1, Ni(BF₄)₂:6H₂O/L-PiMe₂ (1:1, 10 mol%), in DCE at 35 °C for 4 days. Isolated yield. The dr value was determined by ¹H NMR and HPLC analysis. The ee value was determined by HPLC analysis on chiral stationary phases. The reaction of 2b and 2k were scaled up to 1.0 mmol. The reaction of 2n-2p were performed at 35 °C for 8 days

^cThe dr value was determined by ¹H NMR and HPLC analysis ^dDetermined by HPLC analysis on chiral stationary phases

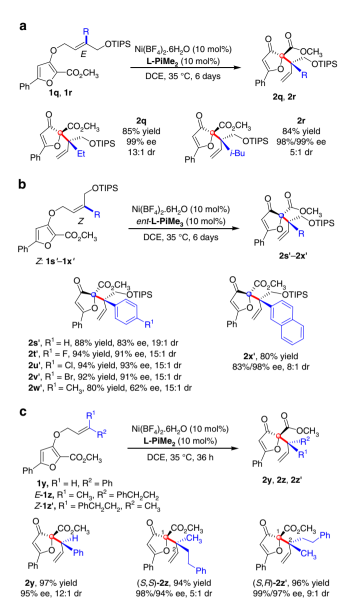


Fig. 3 Substrate scope with variations at alkene unit. **a** Substrate scope of *E*-alkyl substituted alkene. **b** Substrate scope of *Z*-aryl substituted alkene; **c** Substrate scope of mono- and disubstituted alkene. The reactions were performed with 1, Ligand/Ni(BF₄) $_2$ ·6H $_2$ O (1:1, 10 mol%), in DCE at 35 °C for indicated time. Isolated yield. The dr value was determined by ¹H NMR and HPLC analysis. The ee value was determined by HPLC analysis on chiral stationary phases

targeted product 2a even at 70 °C (Table 1, entries 1 and 2)⁵. Next, a range of Lewis acids were examined by complexing with N,N'-dioxide ligand (**L-PiMe**₂) derived from L-pipecolic acid in DCE at 70 °C (For details, see Supplementary Table 1). The reaction proceeded sluggishly in the presence of $Yb(OTf)_3$ (Table 1, entriy 3). Fortunately, $Ni(OTf)_2$ showed excellent catalytic ability and the reaction could complete within 4 h to afford the desired product 2a in 85% yield with 5:1 dr and 98% ee (Table 1, entry 4). Inspired by these results, other N,N'-dioxide ligands were evaluated, however, no better results were obtained (Table 1, entries 5 and 6). Our further attempts to improve the yield and diastereoselectivity focused on the investigation of different silyl protecting group of the substrate 1. Substrates 1b and 1c were synthesized to inspect the reaction (entries 7 and 8). The substrate 1b with bulky TIPS group resulted in the formation of

the corresponding product $2\mathbf{b}$ in 90% yield, 8:1 dr with a comparable ee value (Table 1, entry 7). After variation of the metal counter ion, Ni(BF₄)₂·6H₂O was found to improve the yield to 96% (Table 1, entry 9). Furthermore, we discovered that the reaction temperature even could be reduced to 35 °C by prolonging the reaction time to 4 days and the desired product $2\mathbf{b}$ could be obtained in 96% yield, 10:1 dr and 99% ee (Table 1, entry 10). But further decreasing the reaction temperature led to low conversion. Variation of ester group on furyl unit of the substrate was also explored in the reaction. However, the yield and enantioselectivity were excellent but the dr was moderate (Table 1, entries 11 and 12).

Substrate scope. With the optimized reaction conditions identified, a wide range of substrates with varied substituents on furan unit were investigated. In all cases, the reaction proceeded smoothly with excellent yields and stereoselectivities (Fig. 2). Either electron-withdrawing (1g-1i) or electron-donating (1f and 1j) substituents on the 5-phenyl ring of the furan unit were tolerated in the reaction, furnishing the corresponding products (2f-2j) in excellent yields, diastereo- and enantioselectivities (up to 99% yield, 12:1 dr and 99% ee). 1-Naphthyl (1k) and heteroaromatic (11 and 1m) substituted furan substrates performed the reaction well, giving the related products (2k-2m) in 94-96% yields, 8:1-9:1 dr and 99% ee. Besides, when the 5-alkyl substituted furan substrates (1n-1p) were conducted in the reaction, the products (2n-2p) could also be obtained in excellent results (up to 96% yield, 9:1 dr and 99% ee) after prolonging the reaction time. To demonstrate the utility of this Claisen rearrangement, preparative-scale synthesis of the products 2b and 2k was carried out. Delightedly, the reactivity and stereoselectivity were maintained, which indicated that this method could tolerate the gramscale chemical production. Furthermore, the absolute configuration of 2g was determined to be (S,S) by the X-ray diffraction of the related spirolactone derivative **3g** (See Supplementary Fig. 76) (CCDC 1554368 (3g)). The compounds 2b and 2n exhibited a similar Cotton effect in their circular dichroism (CD) spectra (See Supplementary Figs. 64-75).

The substrates with variation at the alkene unit were also examined (Fig. 3). Under the optimized reaction conditions, Eethyl substituted substrate 1q and E-isobutyl substituted substrate 1r were transformed into the desired products 2q and 2r in excellent yields and enantioselectivities (Fig. 3a, up to 85% yield and 99% ee). The latter showed relatively lower diastereoselectivity (2q/2r,13:1/5:1dr), which is probably due to the larger steric congestion that affects the stereoselective formation of C-C bond 16,19 . When E-phenyl substituted alkene substrate (E-1 \mathbf{s}) was treated in the reaction, no reaction was detected even raising the reaction temperature to 80 °C, which might be caused by the disfavored 1,3-repulsion interaction between aryl group and furan group in the reaction model (See Supplementary Fig. 80). Inversely, Z-aryl substituted alkene substrates could afford the corresponding diastereo-isomers 2' as the major ones. We found the ligand L-PiMe₃ was superior to L-PiMe₂ in terms of reactivity and stereoselectivity, and the use of Ni(BF₄)₂/ent-L-PiMe₃ catalyst for the reaction of Z-1s' resulted in better formation of the related isomer 2s' compared with Ni(BF₄)₂/L-PiMe₂ catalyst (89% yield, 19:1 dr, 83% ee vs. 75% yield, 9:1 dr, 80% ee). With Ni $(BF_4)_2/ent$ -L-PiMe₃ catalyst, the substrates Z-1s' to Z-1x' could participate in the Claisen rearrangement to afford the desired products 2s'-2x' in excellent yields, stereoselectivities (Fig. 3b, up to 95% yield, 19:1 dr and 93% ee). Halogen substitution on the phenyl ring (Z-1t'-Z-1v') made a significant increase of the enantioselectivity. However, electron-donating functional group on the phenyl ring (Z-1w') decreased the enantioselectivity to

Fig. 4 Synthesis of all four stereoisomers. Unless otherwise noted, the reactions were performed with 1 (0.1 mmol), and cat* or *ent*-cat* (10 mol%), in DCE (1.0 mL) at 35 °C for 4 days. Isolated yield. The dr value was determined by ^{1}H NMR and HPLC analysis. The ee value was determined by HPLC analysis on chiral stationary phases. cat* = Ni(BF₄)₂·6H₂O/L-PiMe₂ (1:1), *ent*-cat* = Ni(BF₄)₂·6H₂O/*ent*-L-PiMe₂ (1:1)

Fig. 5 Stereodivergent synthesis of natural products hyperolactones B, C, (—)-biyouyanagin A, and their epimers. a The deprotection/lactonization transformation of the product 2b, 2b', 2n and 2n'. b Formal synthesis of (—)-biyouyanagin A and epi-(—)-biyouyanagin A

62% ee. In addition, the 2-naphthyl substituent alkene E-1 \mathbf{x}' was found suitable for this reaction, delivering the product $2\mathbf{x}'$ in 82% yield with 8:1 dr and 83% ee. Furthermore, terminal monophenyl substituted alkene substrate E-1 \mathbf{y} could afford the desired product $2\mathbf{y}$ in 97% yield with 95% ee and 12:1 dr in the presence of Ni(BF₄)₂/L-PiMe₂ catalyst. Both E-1 \mathbf{z} and Z-1 \mathbf{z}' with terminal disubstituent were suitable in this reaction to afford the corresponding products $2\mathbf{z}$ and $2\mathbf{z}'$ in excellent results (Fig. 3c, up to 96% yield, 99% ee and 9:1 dr).

Asymmetric synthesis of four stereoisomers. To examine our initial stereodivergent assumption and the purpose of synthesizing natural products hyperolactones B and C, we prepared the *E*-**1b**, **1n** and *Z*-**1b**', **1n**' and treated them under the optimal reaction conditions (Fig. 4). As expected, the *E*-substrates preferably afforded (*S*,*S*)-configuration products **2b** and **2n**, while, the *Z*-substrates yielded the (*S*,*R*)-configuration products **2b**' and **2n**' (The configuration of the products were confirmed by CD spectra and NOE spectra analysis of hyperolactones, see

Supplementary Figs. 64–79). Besides, by alternating the absolute configuration of N,N'-dioxide ligand **L-PiMe**₂, the corresponding *ent*-products were available in comparable results. Thus, all four stereoisomers of such γ,δ -unsaturated carbonyl compounds containing vicinal quaternary-quaternary centers could be obtained with this kind of easily available chiral catalyst.

Stereodivergent synthesis of hyperolactones. Subsequently, we tried various reagents to realize the deprotection/lactonization transformation of the product **2b** and **2n**. The conditions, such as TBAF, PPTS, and hydrogen fluoride-pyridine solution proved unsuccessful (For details, see Supplementary Table 8). Delightfully, in the presence of TMSBr reagent, the optically active (*S*,*S*)-**2b**, (*S*,*R*)-**2b**', (*R*,*S*)-**2n**' and (*S*,*S*)-**2n** could participate in the transformation well, delivering hyperolactones B, C and *epi*-hyperolactones B, C in excellent yields with the diastereo- and enantioselectivity maintained (For details, see Supplementary Note 11). Accordingly, the access to natural product (—)-biyouyanagin A and its epimer is possible from the key

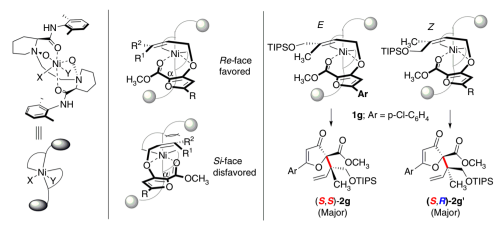


Fig. 6 Proposed stereochemical model. Substrate 1g was selected for the model analysis

reaction intermediates^{26,2729,31} of hyperolactone C or *epi*-hyperolactone C (Fig. 5).

Proposed stereochemical model. On the basis of our previous works^{49–51} and the absolute configuration of the product $2\mathbf{g}$, a possible stereocontrol model was proposed in Fig. 6. The chiral N, N'-dioxide coordinates to the nickel(II) center to form octahedral metal complex in which the chiral ligand occupies four sites. The substrate $1\mathbf{g}$ displays the ancillary solvents or anion and could coordinate to the Lewis acid catalyst via a bidentate manner with the oxygen of ether and carbonyl group of the auxiliary ester group. Due to the steric hindrance between alkene unit and the bulky aniline moiety of ligand, the alkene preferentially approaches α position of furan unit from the Re face. Therefore, the E-alkene substrate $1\mathbf{g}$ affords (S,R)- $2\mathbf{g}'$.

Discussion

In summary, we have developed a catalytic asymmetric dearomatization Claisen rearrangement of allyl furyl ethers for the synthesis of chiral γ , δ -unsaturated carbonyl compounds with two adjacent quaternary-quaternary carbon centers. By adjusting the configuration of the catalysts and alkene units, all four possible product stereoisomers could be prepared in excellent yields and stereoselectivities. In addition, the reaction has a broad substrate scope, which may lead to the discovery of new bioactive molecules. Furthermore, the bioactive natural products hyperolactones B, C, and their epimers could be readily acquired in high efficiency, which are also the key intermediates for the synthesis of natural product (—)-biyouyanagin A and its epimer.

Methods

General procedure for asymmetric Claisen rearrangement. *N,N'*-Dioxide ligand **L-PiMe₂** or *ent*-**L-PiMe₂** or *ent*-**L-PiMe₃** (10 mol%), Ni(BF₄)₂-6H₂O (10 mol%) in CH₂Cl₂ were stirred at 35 °C for 1 h. Then CH₂Cl₂ was removed in *vacuo*. Substrate 1 and DCE were added and the resulting mixture was stirred at 35 °C for 4–8 days. After the reaction completed (monitored by TLC), flash column chromatography was carried out to provide the desired product (Petroleum ether/EtOAc = 20:1 to 10:1).

General procedure for deprotection/cyclization reaction. To a solution of the product 2 (0.1 mmol) in CH₃OH/CH₂Cl₂ (1.5 mL, 2:1, v/v), TMSBr (0.4 mmol, 4.0 equiv.) was added, the reaction mixture was stirred at 30 °C for 8–24 h. Then the reaction was quenched by saturated aq. Na₂CO₃ (1 mL) and diluted with water (1 mL). The product was extracted with ethyl acetate (3 × 10 mL). The combined organic extracts were washed with brine (20 mL) and concentrated in vacuo. The product was purified by column chromatography on silica gel (hexane/EtOAc = 10/1 as eluent).

Data availability. The X-ray crystallographic coordinates for structures reported in this article have been deposited at the Cambridge Crystallographic Data Centre (CCDC). Crystal data and structure refinement for **3g** was displayed in Supplementary Information. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via https://www.ccdc.cam.ac.uk/ under deposition number 1554368. All other data is available from the corresponding author upon reasonable request.

Received: 15 January 2018 Accepted: 5 April 2018 Published online: 17 May 2018

References

- Ito, H. & Taguchi, T. Asymmetric Claisen rearrangement. Chem. Soc. Rev. 28, 43–50 (1999).
- Nubbemeyer, U. Recent advances in asymmetric [3,3]-sigmatropic rearrangements. ChemInform 7, 0961–1008 (2003).
- Ilardi, E. A., Stivala, C. E. & Zakarian, A. [3,3]-Sigmatropic rearrangements: recent applications in the total synthesis of natural products. *Chem. Soc. Rev.* 38, 3133–3148 (2009).
- Rehbein, J. & Hiersemann, M. Claisen rearrangement of aliphatic allyl vinyl ethers from 1912 to 2012: 100 years of electrophilic catalysis. Synthesis 45, 1121–1159 (2013).
- Abraham, L., Czerwonka, R. & Hiersemann, M. The catalytic enantioselective Claisen rearrangement of an allyl vinyl ether. *Angew. Chem. Int. Ed.* 40, 4700–4703 (2001).
- Marié, J.-C. et al. Enantioselective synthesis of 3,4-chromanediones via asymmetric rearrangement of 3-allyloxyflavones. J. Org. Chem. 75, 4584–4590 (2010).
- Tan, J. J., Cheon, C. H. & Yamamoto, H. Catalytic asymmetric Claisen rearrangement of enolphosphonates: construction of vicinal tertiary and allcarbon quaternary centers. *Angew. Chem. Int. Ed.* 51, 8264–8267 (2012).
- Liu, Y. B. et al. Nickel(II)-catalyzed asymmetric propargyl and allyl Claisen rearrangements to allenyl- and allyl-substituted β-ketoesters. Angew. Chem. Int. Ed. 53, 11579–11582 (2014).
- Li, J. et al. Gold(I)/chiral N,N'-dioxide-nickel(II) relay catalysis for asymmetric tandem intermolecular hydroalkoxylation/Claisen rearrangement. Angew. Chem. Int. Ed. 56, 885–888 (2017).
- Fu, J. K. et al. Gold-catalyzed rearrangement of allylic oxonium ylides: efficient synthesis of highly functionalized dihydrofuran-3-ones. *Angew. Chem. Int. Ed.* 52, 4198–4202 (2013).
- Uyeda, C. & Jacobsen, E. N. Enantioselective Claisen rearrangements with a hydrogen-bond donor catalyst. J. Am. Chem. Soc. 130, 9228–9229 (2008).
- Kaeobamrung, J., Mahatthananchai, J., Zheng, P. G. & Bode, J. W. An enantioselective Claisen rearrangement catalyzed by N-heterocyclic carbenes. J. Am. Chem. Soc. 132, 8810–8812 (2010).
- Linton, E. C. & Kozlowski, M. C. Catalytic enantioselective Meerwein-Eschenmoser Claisen rearrangement: asymmetric synthesis of allyl oxindoles. J. Am. Chem. Soc. 130, 16162–16163 (2008).
- Geherty, M. E., Dura, R. D. & Nelson, S. G. Catalytic asymmetric Claisen rearrangement of unactivated allyl vinyl ethers. J. Am. Chem. Soc. 132, 11875–11877 (2010).
- Wu, H. M., Zi, W. W., Li, G. G., Lu, H. J. & Toste, F. D. Gold(I)-catalyzed desymmetrization of 1,4-dienes by an enantioselective tandem alkoxylation/ claisen rearrangement. *Angew. Chem. Int. Ed.* 54, 8529–8532 (2015).

- 16. Yang, Y., Fu, X. N., Chen, J. W. & Zhai, H. B. Total synthesis of (-)-Jiadifenin. Angew. Chem. Int. Ed. 51, 9825–9828 (2012).
- Akahori, Y., Yamakoshi, H., Hashimoto, S. & Nakamura, S. Stereoselective synthesis of the CDE ring system of antitumor Saponin Scillascilloside E-1. Org. Lett. 16, 2054–2057 (2014).
- Akahori, Y., Yamakoshi, H., Sawayama, Y., Hashimoto, S. & Nakamura, S. Synthesis of chiral building blocks for oxygenated Terpenoids through a simultaneous and stereocontrolled construction of contiguous quaternary stereocenters by an Ireland-Claisen rearrangement. J. Org. Chem. 79, 720–735 (2014).
- Uyeda, C., Rötheli, A. R. & Jacobsen, E. N. Catalytic enantioselective Claisen rearrangements of O-allyl ß-ketoesters. *Angew. Chem. Int. Ed.* 49, 9753–9756 (2010).
- Aramaki, Y., Chiba, K. & Tada, M. Spirolactones, hyperolactone A-D from hypericum Chinense. *Phytochemistry* 38, 1419–1421 (1995).
- Crockett, S. L., Schühly, W., Belaj, F. & Khan, I. A. Hyperolactone C. Acta Crystallogr. Sect. E 60, o2174–o2176 (2004).
- Tanaka, N. et al. Biyouyanagin A, an anti-HIV agent from Hypericum chinense L. var. salicifolium. Org. Lett. 7, 2997–2999 (2005).
- Ichinari, D., Ueki, T., Yoshihara, K. & Kinoshita, T. First total synthesis of (±)-hyperolactone A. Chem. Commun. 0, 1743–1744 (1997).
- Ueki, T., Ichinari, D., Yoshihara, K., Morimoto, Y. & Kinoshita, T. Synthesis and absolute configuration of (+)-hyperolactone B. *Tetrahedron Lett.* 39, 667–668 (1998).
- Ueki, T. et al. Synthesis of hyperolactones A and C. J. Heterocycl. Chem. 38, 165–172 (2001).
- Nicolaou, K. C., Sarlah, D. & Shaw, D. M. Total synthesis and revised structure of biyouyanagin A. Angew. Chem. Int. Ed. 46, 4708–4711 (2007).
- Nicolaou, K. C. et al. Total synthesis, revised structure, and biological evaluation of biyouyanagin A and analogues thereof. J. Am. Chem. Soc. 130, 11114–11121 (2008).
- Hodgson, D. M., Erickson, D., Angrish, S. P., Kloesges, J. & Lee, C. H. Consecutive alkene cross-metathesis/ oxonium ylide formationrearrangement: synthesis of the anti-HIV agent hyperolactone C. Org. Lett. 10, 5553–5556 (2008).
- Nicolaou, K. C. et al. Design, synthesis, and biological evaluation of a biyouyanagin compound library. PNAS 108, 6715–6720 (2011).
- Hodgson, D. M. & Man, S. Synthesis of the anti-HIV agent (-)-hyperolactone C by using oxonium ylide formation-rearrangement. *Chem. Eur. J.* 17, 9731–9737 (2011).
- Du, C., Li, L. Q., Li, Y. & Xie, Z. X. Construction of two vicinal quaternary carbons by asymmetric allylic alkylation: total synthesis of hyperolactone C and (-)-biyouyanagin A. Angew. Chem. Int. Ed. 48, 7853-7856 (2009).
- Wu, Y. Y., Du, C., Hu, C. C., Li, Y. & Xie, Z. X. Biomimetic synthesis of hyperolactones. J. Org. Chem. 76, 4075–4081 (2011).
- Kraus, G. A. & Wei, J. Q. A direct synthesis of hyperolactone C. J. Nat. Prod. 67, 1039–1040 (2004).
- 34. Schafroth, M. A., Zuccarello, G., Krautwald, S., Sarlah, D. & Carreira, E. M. Stereodivergent total synthesis of Δ^9 Tetrahydrocannabinols. *Angew. Chem. Int. Ed.* **53**, 13898–13901 (2014).
- Krautwald, S. & Carreira, E. M. Stereodivergence in asymmetric catalysis. J. Am. Chem. Soc. 139, 5627–5639 (2017).
- Lin, L. L. & Feng, X. M. Catalytic strategies for diastereodivergent synthesis. Chem. Eur. J. 23, 6464–6482 (2017).
- Zhan, G., Du, W. & Chen, Y. C. Switchable divergent asymmetric synthesis via organocatalysis. Chem. Soc. Rev. 46, 1675–1692 (2017).
- Krautwald, S., Sarlah, D., Schafroth, M. A. & Carreira, E. M. Enantio- and diastereodivergent dual catalysis: α-allylation of branched aldehydes. *Science* 340, 1065–1068 (2013).
- Schindler, C. S. & Jacobsen, E. N. A new twist on cooperative catalysis. Science 340, 1052–1053 (2013).
- Huo, X. H., He, R., Zhang, X. & Zhang, W. B. An Ir/Zn dual catalysis for enantio- and diastereodivergent α-allylation of α-hydroxyketones. J. Am. Chem. Soc. 138, 11093–11096 (2016).
- Jiang, X. Y., Beiger, J. J. & Hartwig, J. F. Stereodivergent allylic substitutions with aryl acetic acid esters by synergistic iridium and Lewis base catalysis. J. Am. Chem. Soc. 139, 87–90 (2017).
- Cruz, F. A. & Dong, V. M. Stereodivergent coupling of aldehydes and alkynes via synergistic. J. Am. Chem. Soc. 139, 1029–1032 (2017).

- Denmark, S. E. & Ghosh, S. K. The first catalytic, diastereoselective, and enantioselective crossed-aldol reactions of aldehydes. *Angew. Chem., Int. Ed.* 40, 4759–4762 (2001).
- Kim, H. Y., Li, J. Y., Kim, S. & Oh, K. Stereodivergency in catalytic asymmetric conjugate addition reactions of glycine (ket)imines. *J. Am. Chem. Soc.* 133, 20750–20753 (2011).
- Molinaro, C. et al. Catalytic, asymmetric, and stereodivergent synthesis of non-symmetric β,β-diaryl-α-amino acids. J. Am. Chem. Soc. 137, 999–1006 (2015).
- Wang, L., Liu, X. F. & Deng, L. Dual-function cinchona alkaloid catalysis: catalytic asymmetric tandem conjugate addition-protonation for the direct creation of nonadjacent stereocenters. J. Am. Chem. Soc. 128, 3928–3930 (2006).
- Mechler, M. & Peters, R. Diastereodivergent asymmetric 1,4-addition of oxindoles to nitroolefins by using polyfunctional nickel-hydrogen-bondazolium catalysts. *Angew. Chem. Int. Ed.* 54, 10303–10307 (2015).
- Engl, O. D., Fritz, S. P. & Wennemers, H. D. Stereoselective organocatalytic synthesis of oxindoles with adjacent tetrasubstituted stereocenters. *Angew. Chem. Int. Ed.* 54, 8193–8197 (2015).
- Liu, X. H., Lin, L. L. & Feng, X. M. Chiral N,N'-dioxides: new ligands and organocatalysts for catalytic asymmetric reactions. Acc. Chem. Res. 44, 574–587 (2011).
- Liu, X. H., Lin, L. L. & Feng, X. M. Chiral N,N'-dioxide ligands: synthesis, coordination chemistry and asymmetric catalysis. Org. Chem. Front. 1, 298–302 (2014)
- Liu, X. H., Zheng, H. F., Xia, Y., Lin, L. L. & Feng, X. M. Asymmetric cycloaddition and cyclization reactions catalyzed by chiral N,N'-dioxide-metal complexes. Acc. Chem. Res. 50, 2621–2631 (2017).

Acknowledgements

We appreciate the National Natural Science Foundation of China (Nos. 21432006, 21625205) for financial support. Dedicated to professor George A. Kraus for his initial contribution on the dearomatization Claisen rearrangement of allyl furyl ethers.

Author contributions

H.F.Z. performed experiments and prepared the Supplementary Information and Manuscript. Y.W. and C.R.X. took part in the reaction development and synthesized several substrates. X.X. repeated some experiments. L.L.L. helped with modifying the paper and Supplementary Information. X.H.L. and X.M.F. conceived and directed the project.

Additional information

Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-018-04123-w.

Competing interests: The authors declare no competing interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

Publisher's note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing,

adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2018