

Synthetic Methods

Direct Site-Specific and Highly Enantioselective γ -Functionalization of Linear α,β -Unsaturated Ketones: Bifunctional Catalytic Strategy**

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The direct asymmetric γ functionalization of simple unmodified carbonyl compounds has emerged as a powerful, distinct, and atom-economical method offering an efficient entry into functionalized building blocks having high levels of structural complexity.^[1] Since the pioneering studies carried out by the groups of Jørgensen, Chen, Melchiorre, and others, different types of vinylogous processes initiated by dienamine intermediates have been studied extensively.^[2] However, while there is an ever growing number of documents involving catalytic γ -selective activation of enals, the corresponding reaction embracing enones as γ -type nucleophiles remain comparatively rare. To date, the only example of enantioselective γ functionalization of cyclic enones has been reported by Melchiorre and co-workers who use cyclohexenone derivatives as nucleophilic substrates.^[3] Nevertheless, for the γ -site-specific functionalization of linear α,β -unsaturated ketones, which usually act as electrophiles in Michael additions (Figure 1a)^[4] or prefer to selectively direct the reaction toward α alkylation (Figure 1b),^[5] still represents a longstanding challenge (Figure 1c).

Covalent activation facilitated by chiral amines provides a fascinating platform for stereocontrol in the functionaliza-

tion at the γ -position of the carbonyl group through dienamine-type intermediates,^[2] or even at the ϵ -position by trienamine catalysis.^[6] However, as mentioned before, this strategy works well for HOMO-raising activation of γ - or ϵ -type nucleophiles equipped with an aldehyde group.^[2a-i,6a-f] In the case of enones, the covalent catalytic process was not efficient for such a transformation. More importantly, for the ketones bearing an α' hydrogen, the regioselectivity problem must be solved, especially for linear enones.^[5,7] Very recently, Chen and co-workers used trienamine catalysis to perform a highly efficient asymmetric Diels–Alder reaction with linear 2,4-dienones to give an array of useful cyclohexene products.^[8] Still, a “logical and careful design of the dienones” was needed to facilitate the reaction,^[9] as it requires an aryl group at the α' -position to the carbonyl group to suppress reaction at the α' site.

To this end, we envisioned the development of alternative catalytic strategies for the site-selective and stereocontrolled γ functionalization of linear enones.^[10] We have recently reported an asymmetric vinylogous Michael addition of α,β -unsaturated γ -butyrolactams and found that the magnesium is prone to activating the γ hydrogen of the carbonyl group (Figure 2a).^[11] On this basis, we explored the viability of magnesium catalysis to direct γ deprotonation of linear α,β -unsaturated ketones by introducing an appropriately placed base within the ligand to facilitate the γ -activation process (Figure 2b).

To test our hypothesis, we initially investigated a series of representative salen ligands, either bearing a base or not, in our model reaction. As illustrated in Table 1, the nature of the Brønsted base, such as a tertiary amine or even methoxy

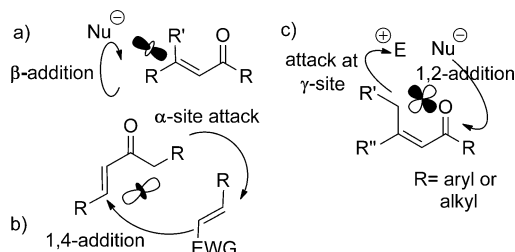


Figure 1. a,b) Previously reported strategies. c) Strategy for γ functionalization of linear enones proposed herein.

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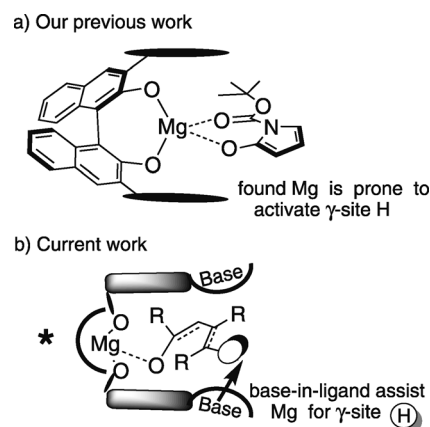
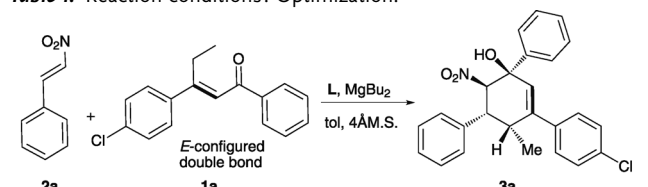


Figure 2. a) Catalyst design reported by our group previously. b) New ligand design reported herein. A base is introduced to assist in the activation of linear enones.

Table 1: Reaction conditions: Optimization.



Reaction scheme showing the conversion of nitroalkene **2a** and ketone **1a** (with an *E*-configured double bond) to product **3a** using ligand **L**, MgBu_2 , and toluene at 40 °C for 24 h.

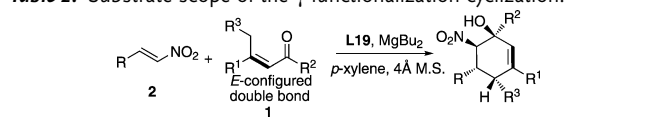
Entry ^[a]	Ligand	Yield ^[b]	d.r. ^[c]	ee ^[d]
1	L1	7	—	—
2	L2	< 5	—	—
3	L3	48	6:1	68
4	L4	52	11:1	85
5	L5	50	11:1	39
6	L12	53	10:1	82
7	L15	62	9:1	80
8	L17	63	9:1	88
9	L19	59	9:1	92
10 ^[e]	L19	64	9:1	98
11 ^[e,f]	L19	72	9:1	98

[a] Reactions were performed with 0.2 mmol of **2a** and 0.3 mmol of **1a** in toluene (2.0 mL) in the presence of **L** (20 mol %), MgBu_2 (20 mol %), and molecular sieves (M.S.) at 60 °C for 24 h. [b] Yield of isolated product. [c] Diastereomeric ratios were determined by ^1H NMR (300 MHz) spectroscopy. [d] Enantiomeric excesses of the major diastereomer were analyzed by HPLC analysis using a chiral stationary phase. [e] The reaction was carried out in *p*-xylene. [f] Use 0.4 mmol of **1a** (2.0 equiv).

group, introduced to a ligand was helpful for the γ -deprotonation process, probably through assistance from the base near the magnesium center of the catalyst (entries 3 and 4 versus 1 and 2). We turned our efforts to the screening and development of salen-type ligands having such an appended base.^[12] This effort led to the identification of the bromine-substituted ligand **L19** as the optimal catalyst, and after optimization of the solvent,^[13] the reaction in *p*-xylene afforded good diastereoselectivity (9:1) and high enantiomeric excess of 98% (entries 9 and 10).^[14] The use of 2.0 equivalents of the ketone **1a** improved the product (**3a**) yield to 72% yield without loss in the enantioselectivity (entry 11).

Having established the optimized reaction conditions, we explored the scope of this new catalytic, stereoselective γ -

Table 2: Substrate scope of the γ -functionalization cyclization.^[a,b,c,d]



Reaction scheme showing the conversion of nitroalkene **2** and ketone **1** (with an *E*-configured double bond) to product **3** using ligand **L19**, MgBu_2 , and *p*-xylene at 60 °C for 24 h.

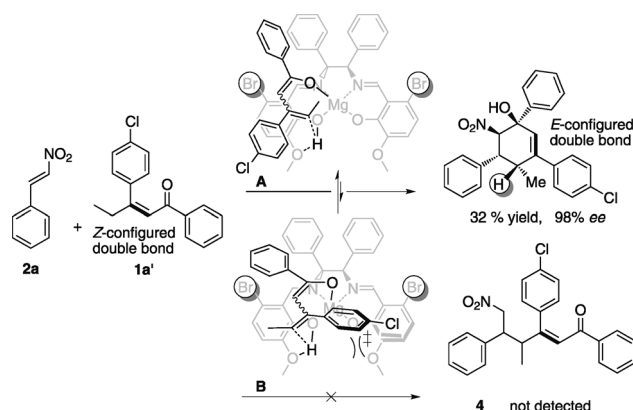
Product	Yield	d.r.	ee
3a	72 %	9:1	98 % ee
3b : R=Me	70 %	10:1	97 % ee
3c : R=F	73 %	10:1	95 % ee
3d : R=Cl	69 %	8:1	98 % ee
3e : R=Br	61 %	8:1	97 % ee
3f : R=OMe	55 %	12:1	96 % ee
3g : R=m-Me	67 %	7:1	98 % ee
3h : R=m-Cl	62 %	9:1	95 % ee
3i : R=o-F	74 %	6:1	95 % ee
3j : X=O	82 %	9:1	98 % ee
3m : X=S	73 %	7:1	97 % ee
3n : R=H	72 %	10:1	98 % ee
3o : R=o-F	71 %	10:1	96 % ee
3p : R=m-Me	75 %	9:1	98 % ee
3q : R=p-OMe	64 %	9:1	99 % ee
3r : R=p-F	72 %	10:1	99 % ee
3s : R=m-OMe	64 %	10:1	91 % ee
3t	57 %	6:1	94 % ee
3u	77 %	6:1	89 % ee
3v	42 %	8:1	92 % ee
3w	53 %	3:1	96 % ee
5 : n=1	60 %	7:1	86 % ee
6 : n=1	63 %	9:1	98 % ee
7 : n=2	42 %	11:1	92 % ee
8	56 %	9:1	95 % ee

[a] Reactions were performed with 0.2 mmol of **2a** and 0.4 mmol of **1a** in *p*-xylene (2.0 mL) in the presence of **L19** (20 mol %), MgBu_2 (20 mol %) and 4 Å molecular sieves (200 mg) at 60 °C for 24 h. [b] Yield of isolated product. [c] Diastereomeric ratios were determined by ^1H NMR (300 MHz) spectroscopy. [d] Enantiomeric excesses of the major diastereomer were analyzed by HPLC analysis using a chiral stationary phase.

functionalization protocol. The representative results are listed in Table 2. Nitroalkenes bearing a variety of aryl groups with either electron-donating or electron-withdrawing substituents, or heteroaryl groups were well tolerated in this reaction, and the corresponding cyclization products were obtained with high *ee* values and moderate to good yields and diastereoselectivity (**3a–m**). However, aliphatic nitroalkenes could not undergo this transformation; the desired products were not detected under our standard reaction conditions. Furthermore, the linear β,β -disubstituted α,β -unsaturated ketones **1** bearing different aryl groups at either the β - or α' -positions were also compatible, thereby leading to the expected products (**3n–3t**). Notably, the presence of different substituents at the γ -position of the enones **2** were all tolerated, thus giving excellent *ee* values and moderate

yields (**3u**, **3v**, **3w**). We then turned our attention to substrates bearing both *Z*- γ - and *E*- γ' -hydrogen atoms to furnish the cyclization product **5** and the bicyclic compounds **6** and **7**. The benzocyclic ring systems **8** could also be obtained with excellent *ee* values and moderate yields. However, the ketones bearing different aliphatic substitutes at the β site and simple disubstituted enones were not tolerated under the present catalytic reaction and only resulted in complex mixtures.^[15]

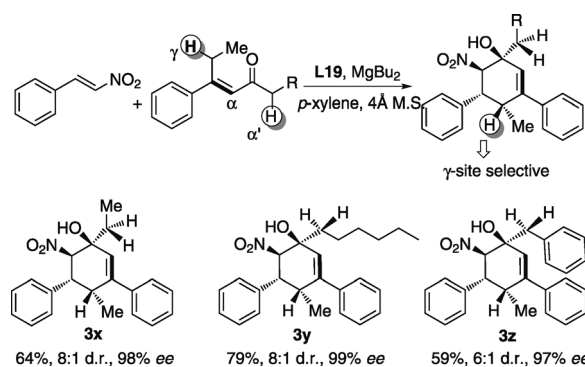
To gain insight into the activation mode of the γ -deprotonation-defining step of the reaction, we employed the (*Z*)- α,β -unsaturated ketone **1a'** under the standard, aforementioned reaction conditions (Scheme 1). We were surprised to observe that the cyclization product **3a** was also



Scheme 1. The cyclization reaction using the (*Z*)- α,β -unsaturated ketone **1a'**

formed with excellent *ee* value and the *Z*-configured double bond in **1a'** resulted in an *E*-configured double bond in **3a**. The product **4**, which maintains the *Z*-configured double bond was not observed. The result suggests that a dienol transition state should exist and the *Z*-configured double bond is transformed into an *E*-configured double bond during the γ -deprotonation process (**A**, Scheme 1). The dienol transition state might be more compatible with the catalysis structure, and a concerted reaction might also proceed smoothly through this favored configuration. Notably, the yield in this transformation was lower, probably owing to the isomerization of the double bond during γ -deprotonation process.

Finally and importantly, an experiment to probe site selectivity was carried out by using β,β -disubstituted α,β -unsaturated linear ketones containing α' hydrogen atoms (Scheme 2). We were pleased to find complete γ selectivity in preference to the traditional α' selectivity under our catalytic system (Scheme 2; **3x**, **3y**, **3z**).^[16] The highly efficient site selectivity was no doubt determined by the presence of the bifunctional base within the ligand for the magnesium catalysis: 1) Magnesium is a superior metal for activating the γ hydrogen atoms of carbonyl groups, 2) the methoxy group introduced into the ligand might act as a Brønsted base to assist in the γ -deprotonation process, and 3) compared to the α' site, the γ site might be better positioned to capture the



Scheme 2. Site selectivity demonstration for substrates bearing both α' and γ hydrogen atoms.

β site of the electrophiles after complex formation with the metal center.

In summary, we have described the first examples of direct site- and enantioselective γ functionalization of linear β,β -disubstituted α,β -unsaturated ketones, thus leading to a variety of optically active cyclohexene frameworks which are not easily accessible using other methodologies. Success of the activation mode, site-specific selectivity, and asymmetric induction relies on the bifunctional base within the ligand. The development of this strategy in other, still challenging γ -functionalization reactions is currently ongoing.

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- [1] For reviews on vinylogous processes of carbonyl compounds, see: a) J.-L. Li, T.-Y. Liu, Y.-C. Chen, *Acc. Chem. Res.* **2012**, *45*, 1491; b) G. Casiraghi, L. Battistini, C. Curti, G. Rassu, F. Zanardi, *Chem. Rev.* **2011**, *111*, 3076; c) S. F. Martin, *Acc. Chem. Res.* **2002**, *35*, 895; d) S. E. Denmark, J. R. Heemstra, Jr., G. L. Beutner, *Angew. Chem.* **2005**, *117*, 4760; *Angew. Chem. Int. Ed.* **2005**, *44*, 4682.
- [2] a) L. Albrecht, G. Dickmeiss, C. F. Weise, C. Rodríguez-Esrich, K. A. Jørgensen, *Angew. Chem.* **2012**, *124*, 13286; *Angew. Chem. Int. Ed.* **2012**, *51*, 13109; b) L. Albrecht, G. Dickmeiss, F. C. Acosta, C. Rodríguez-Esrich, R. L. Davis, K. A. Jørgensen, *J. Am. Chem. Soc.* **2012**, *134*, 2543; c) S. Bertelsen, M. Marigo, S. Brandes, P. Dinér, K. A. Jørgensen, *J. Am. Chem. Soc.* **2006**, *128*, 12973; d) J.-L. Li, T.-R. Kang, S.-L. Zhou, R. Li, L. Wu, Y.-C. Chen, *Angew. Chem.* **2010**, *122*, 6562; *Angew. Chem. Int. Ed.* **2010**, *49*, 6418; e) J.-L. Li, S.-L. Zhou, P.-Q. Chen, L. Dong, T.-Y. Liu, Y.-C. Chen, *Chem. Sci.* **2012**, *3*, 1879; f) G. Bergonzini, S. Vera, P. Melchiorre, *Angew. Chem.* **2010**, *122*, 9879; *Angew. Chem. Int. Ed.* **2010**, *49*, 9685; g) C. Cassani, P. Melchiorre, *Org. Lett.* **2012**, *14*, 5590; h) K. Liu, A. Chougnet, W.-D. Woggon, *Angew. Chem.* **2008**, *120*, 5911; *Angew. Chem. Int. Ed.* **2008**, *47*, 5827; i) R. M. de Figueiredo, R. Fröhlich, M. Christmann, *Angew. Chem.* **2008**, *120*, 1472; *Angew. Chem. Int. Ed.* **2008**, *47*, 1450; j) A. G. Nigmatov, E. P. Serebryakov, *Russ. Chem. Bull.* **1993**, *42*, 213.
- [3] a) G. Bencivenni, P. Galzerano, A. Mazzanti, G. Bartoli, P. Melchiorre, *Proc. Natl. Acad. Sci. USA* **2010**, *107*, 20642; b) D.

- Bastida, Y. K. Liu, X. Tian, E. Escudero-Adán, P. Melchiorre, *Org. Lett.* **2013**, *15*, 220.
- [4] a) B. L. Feringa, *Acc. Chem. Res.* **2000**, *33*, 346; b) A. W. Hird, A. H. Hoveyda, *J. Am. Chem. Soc.* **2005**, *127*, 14988; c) M. d'Augustin, L. Palais, A. Alexakis, *Angew. Chem.* **2005**, *117*, 1400; *Angew. Chem. Int. Ed.* **2005**, *44*, 1376; d) R. Shintani, Y. Tsutsumi, M. Nagaosa, T. Nishimura, T. Hayashi, *J. Am. Chem. Soc.* **2009**, *131*, 13588; e) Y. Tanaka, M. Kanai, M. Shibasaki, *J. Am. Chem. Soc.* **2010**, *132*, 8862; f) C. Mazet, E. N. Jacobsen, *Angew. Chem.* **2008**, *120*, 1786; *Angew. Chem. Int. Ed.* **2008**, *47*, 1762; g) X. Feng, J. Yun, *Chem. Eur. J.* **2010**, *16*, 13609; h) D. Zhao, L. Mao, L. Wang, D. Yang, R. Wang, *Chem. Commun.* **2012**, *48*, 889; i) K. Kikushima, J. C. Holder, M. Gatti, B. M. Stoltz, *J. Am. Chem. Soc.* **2011**, *133*, 6902.
- [5] a) M. D. Pierce, R. C. Johnston, S. Mahapatra, H. Yang, R. G. Carter, P. H.-Y. Cheong, *J. Am. Chem. Soc.* **2012**, *134*, 13624; b) L.-Y. Wu, G. Bencivenni, M. Mancinelli, A. Mazzanti, G. Bartoli, P. Melchiorre, *Angew. Chem.* **2009**, *121*, 7332; *Angew. Chem. Int. Ed.* **2009**, *48*, 7196; c) G. Bencivenni, L.-Y. Wu, A. Mazzanti, B. Giannichi, F. Pesciaoli, M.-P. Song, G. Bartoli, P. Melchiorre, *Angew. Chem.* **2009**, *121*, 7336; *Angew. Chem. Int. Ed.* **2009**, *48*, 7200.
- [6] a) Z.-J. Jia, H. Jiang, J.-L. Li, B. Gschwend, Q.-Z. Li, X. Yin, J. Grouleff, Y.-C. Chen, K. A. Jørgensen, *J. Am. Chem. Soc.* **2011**, *133*, 5053; b) Z.-J. Jia, Q. Zhou, Q.-Q. Zhou, P.-Q. Chen, Y.-C. Chen, *Angew. Chem.* **2011**, *123*, 8797; *Angew. Chem. Int. Ed.* **2011**, *50*, 8638; c) H. Jiang, B. Gschwend, L. Albrecht, S. G. Hansen, K. A. Jørgensen, *Chem. Eur. J.* **2011**, *17*, 9032; d) Y. Liu, M. Nappi, E. Arceo, S. Vera, P. Melchiorre, *J. Am. Chem. Soc.* **2011**, *133*, 15212; e) Y. Liu, M. Nappi, E. C. Escudero-Adán, P. Melchiorre, *Org. Lett.* **2012**, *14*, 1310; f) C. Ma, Z.-J. Jia, J.-X. Liu, Q.-Q. Zhou, L. Dong, Y.-C. Chen, *Angew. Chem.* **2013**, *125*, 982; *Angew. Chem. Int. Ed.* **2013**, *52*, 948.
- [7] a) N. Momiyama, N. Yamamoto, H. Yamamoto, *J. Am. Chem. Soc.* **2007**, *129*, 1190; b) H. Sundén, I. Ibrahim, L. Eriksson, A. Córdova, *Angew. Chem.* **2005**, *117*, 4955; *Angew. Chem. Int. Ed.* **2005**, *44*, 4877.
- [8] X.-F. Xiong, Q. Zhou, J. Gu, L. Dong, T.-Y. Liu, Y.-C. Chen, *Angew. Chem.* **2012**, *124*, 4477; *Angew. Chem. Int. Ed.* **2012**, *51*, 4401.
- [9] E. Arceo, P. Melchiorre, *Angew. Chem.* **2012**, *124*, 5384; *Angew. Chem. Int. Ed.* **2012**, *51*, 5290.
- [10] For an alternative activation mode for direct γ functionalization of enals, see: J. Mo, X. Chen, Y. R. Chi, *J. Am. Chem. Soc.* **2012**, *134*, 8810.
- [11] L. Lin, J. L. Zhang, X. J. Ma, X. Fu, R. Wang, *Org. Lett.* **2011**, *13*, 6410.
- [12] For pioneering examples employing a bifunctional base to assist with catalysis, see: a) Y. Tanaka, M. Kanai, M. Shibasaki, *J. Am. Chem. Soc.* **2008**, *130*, 6072; b) I. Fujimori, T. Mita, K. Maki, M. Shiro, A. Sato, S. Furusho, M. Kanai, M. Shibasaki, *J. Am. Chem. Soc.* **2006**, *128*, 16438; c) Y. Hamashima, D. Sawada, M. Kanai, M. Shibasaki, *J. Am. Chem. Soc.* **1999**, *121*, 2641.
- [13] See the Supporting Information for details of the reaction optimization with respect to the ligands and solvents.
- [14] See the Supporting Information for details of the X-ray analysis.
- [15] See the Supporting Information for some unsuccessful examples of the transformation.
- [16] The enone bearing α' , γ , and γ' hydrogen atoms could also generate the desired cyclization product with a lower *ee* value:

