

Published in final edited form as:

Nat Chem. 2017 December; 9(12): 1269-1275. doi:10.1038/nchem.2816.

Catalytic diastereo- and enantioselective additions of versatile allyl groups to N-H ketimines

Hwanjong Jang, Filippo Romiti, Sebastian Torker, and Amir H. Hoveyda Department of Chemistry, Merkert Chemistry Center, Boston College, Chestnut Hill, Massachusetts, 02467, USA

Abstract

There are countless biologically active organic molecules that contain one or more N-containing moieties and broadly applicable and efficient catalytic transformations that deliver them diastereo-and/or enantioselectively are much sought after. Various methods for enantioselective synthesis of α -secondary amines are available (e.g., from additions to protected/activated aldimines), but those involving ketimines are much less common. There are no reported additions of carbon-based nucleophiles to unprotected/unactivated (or N-H) ketimines. Here, we report a catalytic, diastereo-and enantioselective three-component strategy for merging an N-H ketimine, a monosubstituted allene and B₂(pin)₂, affording products in up to 95% yield, >98% diastereoselectivity and >99:1 enantiomeric ratio. Utility of the approach is highlighted by synthesis of the tricyclic core of a class of compounds that have been shown to possess anti-Alzheimer activity. Stereochemical models, developed with the aid of DFT calculations, which account for the observed trends and levels of enantioselectivity are presented.

Graphical abstract

Reprints and permissions information is available at npg.nature.com/reprintsandpermissions.

Correspondence and requests for materials should be addressed to A.H.H. (amir.hoveyda@bc.edu).

Data availability

X-ray crystallographic data for compounds *rac-***3a**, N-acetyl derivative of **3y** and **9b** are freely available from the Cambridge Crystallographic Data Centre (CCDC 1547738, 1547736 and 1547737, respectively).

Author Contributions

H. J. and F. R. developed the catalytic method and its various applications. S. T. designed and performed the DFT calculations. A. H. H. directed the investigations and composed the manuscript with revisions provided by the other authors.

The authors declare competing financial interests.

Catalytic enantioselective additions of carbanions to ketimines deliver products with a nitrogen-substituted quaternary stereogenic center (α-tertiary amine) but development of these transformations 1,2,3,4 is hardly straightforward. Ketimines are less reactive than aldimines and the reduced size difference between the substituents compared to aldimines makes enantiotopic face differentiation difficult. Catalytic enantioselective additions of allyl moities⁵ to ketimines, while much sought after, remain scarce. One study shows that reactions of allyl–B(pin) (pin = pinacolato) with acyclic N-benzyl ketimines may be promoted by a chiral bis-phosphine—Cu complex (Fig. 1a)⁶, and another deals with reactions of functionalized allylsilanes and tosyl-protected ketimines catalyzed by phosphoramidite-Pd complexes (Fig. 1a)⁷. Other disclosures cover highly activated ketimines, including cyclic sulfonylimines and their reaction with potassium allyltrifluoroborates (with Rh-based catalysts)^{8, 9} and isatin-derived N-Boc-ketimines and their reaction with allylsilanes (with Pd-based complexes and stoichiometric silver fluoride)¹⁰. Other approaches involve either enantiomerically pure ketimines^{11, 12} or enantiomerically pure allyl reagents^{13,14,15}. Our goal was to develop a method that would not require ketimine activation/protection and subsequent unmasking (Fig. 1b). The absence of a protecting group would bypass the intermediacy of E and Z mixtures of ketimine isomers, which can lead to lowering of enantioselectivity. Although preparation of N-H ketimines by condensation of ketones with ammonia followed by reaction with allylboron reagents is known¹⁶, as far as we are aware, diastereo- and/or enantioselective variants have not been introduced.

Based on the earlier investigations regarding enantioselective additions to aldehydes or ketones 17,18,19 , which were recently extended to N-anisidyl aldimines 20,21 , we envisioned the sequence in Fig. 1b. N-H ketimines would be accessed by addition of an organolithium or a Grignard reagent to a nitrile, many of which are commercially available 22,23 . The ensuing catalytic process would combine an N-H ketimine, a monosubstituted allene and $B_2(pin)_2$ to generate homoallylic amines containing a pair of stereogenic centers and an alkenyl–B(pin) group. A number of biologically active organic molecules would thus become more readily accessible in enantiomerically enriched form; an example would be of the core structure of class of polycyclic compounds shown to possess the ability to reduce beta-amyloid production (Fig. 1b) $^{24, 25}$. Complications typically associated with siteselective removal of protecting/activating units would thus be obviated, particularly when

relatively strong reducing (e.g., for an N-benzyl or an N-tosyl group, Fig. 1a) or oxidizing conditions²⁰ are needed and a stereogenic benzylic C–N bond is present.

Results

Establishing feasibility

We began with the reaction of allene **2a** with N-H ketimine **1a**, obtained from the reaction of benzonitrile with methyllithium (MeLi; 88% yield; Table 1). Diastereoselectivity was complete in every case [>98:2 diastereomeric ratio (d.r.)] but efficiency was catalyst-dependent. There was 60–75% conversion to *rac*-**3a** with Cu complexes derived from triphenylphosphine, tricyclohexylphosphine or racemic 2,2'-bis(diphenylphosphino)-1,1'-binaphthalene (*rac*-binap) (entries 1–3). Evaluation of *N*-heterocyclic carbene (NHC) complexes (entries 4–6) showed that the combination of cyclohexyl-substituted imidazolium salt **4c** and CuCl is the most effective: *rac*-**3a** was obtained in 90% yield and >98:2 d.r. (Table 1, entry 6).

Identifying a chiral catalyst

Several types of Cu complexes were examined (Table 2). Enantioselectivity was minimal with binap [(2,2'-bis(diphenylphosphino)-1,1'-binaphthyl); entry 1, 44:56 enantiomeric ratio (e.r.)]. The desired product was isolated in appreciable yield and e.r. with segphos (**5a**, 51% yield, 18:82 e.r.; entry 2) or josiphos (**5b**, 51% yield, 19.5:80.5 e.r.; entry 3), but less so with the more conformationally flexible **5c** (27%, 69:31, respectively; entry 4).

With imidazolinium salt **6** or sulfonate-bearing **7a** enantioselectivity remained low (entries 5–6). However, when the mesityl (Mes) moiety of **7a** was replaced by a 3,5-diaryl-substituted phenyl moiety (**7b**, entry 7) enantioselectivity increased dramatically: **3a** was obtained in 95:5 e.r. (for X-ray structure of the derived alcohol, see Fig. 3a). The effectiveness of the **7b**-derived catalyst was surprising for several reasons. Enantioselectivity was considerably higher compared to the closely related *N*-mesityl-substituted variant (**7a**; see below for mechanistic analysis). Additionally, while sulfonate-containing NHC–Cu catalysts have been used for enantioselective allylic substitutions^{26,27} and conjugate additions^{28,29,30}, none emerged as optimal for a 1,2-addition.

Applicability

The method has considerable scope (Table 3). At times higher catalyst loading was necessary for high conversion (e.g., **3a**, entry 1 Table 3 vs. entry 7, Table 2) and ketimine:allene ratio was changed to 1:1.5 for better yield (from 1.2:1). Regardless of whether the N-H ketimine had an *ortho* aryl substituent that is electron donating (**3b**), electron withdrawing (**3c**) or relatively sizeable (**3f–g**), products were isolated in 59–95% yield, >98:2 d.r. and 98.5:1.5–99.5:0.5 e.r. Reactions with *meta*-substituted N-H ketimines were similarly efficient and selective (entries 8–10, Table 3). In reactions of ketimines with different *para*-substituted aryl units (see **3k–3m**) e.r. ranged from 92.5:7.5 to 95:5. Products **3n–o** (Table 3, entries 11–12), from reactions with the less electrophilic alkyl-substituted N-H ketimines, were isolated in 38% and 48% yield and 91:9 and 95:5 e.r., respectively. Fluoroaryl-substituted amines **3d–e** (entries 4–5), **3h** (entry 8), and **3l** (entry 12) were

obtained in 64–91% yield and 94:6–98:2 e.r. (>98:2 d.r.); the high yield and e.r. in these transformations, regardless of the position of the fluorine atom, shows that fluorine-metal interaction^{31,32,33} does not exert an adverse influence. Diastereoselectivity was exceptional throughout (>98:2 d.r.).

Additions to heterocyclic N-H ketimines afforded products in high e.r. (cf. **3p–3r**, Fig, 2a), although efficiency was slightly lower. Synthesis of amines **3s–3u** shows that different monosubstituted allenes may be used, but the size of the allene substituent did impact efficiency (41–78% yield, >98:2 d.r., up to >99:1 e.r.).

The method extends beyond methyl-substituted substrates. Ketimines bearing an n-alkyl or unsaturated alkyl group (e.g., 3v-x, Fig. 2a) and the *iso*-propyl-containing ketimine precursor to 3y were converted to the desired amines efficiently and with exceptional diastereoselectivity. Nevertheless, e.r. varied depending on the substituent. Whereas 3v was formed in 94:6 e.r., there was gradual diminution in enantioselectivity as the side chain became longer (e.g., 88:12 e.r. for *n*-butyl-substituted 3w, 85.5:14.5 e.r. for pentenylsubstituted 3x). More enantioselective was the reaction that afforded *iso*-poropyl-containing 3y (96:4 e.r.); the X-ray structure secured for the N-acetyl derivative of 3y (Fig. 2a) indicates that there is no reversal in stereochemical induction (see Fig. 3a for corroborative X-ray data). A possible rationale for these selectivity trends will be provided below. Various compounds of interest, such as those derived from intramolecular hydroamination that afford heterocyclic derivatives^{34, 35}, may be accessed through functionalization of compounds such as 3x. Nonetheless, there are limitations. Reactions of ketimines that contain an α - or β alkoxy or a benzyl group are inefficient, likely due to facile decomposition (enamine formation and β-elimination, respectively). The same applies to additions to trifluoromethylsubstituted ketimines (decomposition to unidentified products). There was no reaction with phenyl-tert-butyl N-H ketimine.

Utility

Catalytic enantioselective addition to ketimine **8** followed by oxidation of the C–B bond gave β-amino ketone **9a**, the product of a Mannich-type addition, in 70% overall yield, yield and without any loss in d.r. (Fig. 2b). The efficiency with which **9a** was obtained is higher than those shown in Table 3 and Fig. 2a, indicating that there might be some decomposition during purification and that yields may be improved if the alkenyl–B(pin) moiety is modified. The absolute stereochemistry of the product was confirmed through X-ray structure of primary alcohol **9b**. It merits note that catalytic enantioselective enolate additions to *N*-activated ketimines (e.g., *N*-phosphinoylketimines^{36,37} or those derived from α-ketoesters³⁸ or diethyl ketomalonate³⁹) are limited in scope (see the Supplementary Information for additional references).

We then investigated the possibility of application to enantioselective synthesis of the core structure of the aforementioned anti-Alzheimer compounds (Fig. 2c). NHC–Cu-catalyzed protodeboration¹⁷ of enantiomerically enriched alkenyl–B(pin) amine **3c** furnished **10** in 72% yield. Synthesis of α -olefin **10** by a related route and with a sterically less demanding Cu–H complex [vs. Cu–B(pin) addition/protodeboration] would present a chemoselectivity

issue (competitive reaction with ketimine⁴⁰); moreover, we find that the presence of a B(pin) group is critical to high enantioselectivity (see Fig. 3). Thiourea generation and removal of the silyl group afforded alcohol 11 in 71% overall yield. The cyclic ether was formed by treatment of 11 with 10 mol % CuI and 20 mol % 8-hydroxyquinoline (110 °C, 24 h)⁴¹, affording oxepane 12 in 78% yield. Oxidative cleavage of the vinyl group, reduction of the resulting aldehyde and subjection of the resulting primary alcohol to triflic anhydride $(-20 \, ^{\circ}\text{C}, 2 \, \text{h})^{25}$ delivered 13 in 75% yield after recrystallization (this compound is unstable towards a variety of chromatography procedures). The aryl ring in 13 may be functionalized site selectively according to formerly reported procedures^{42,43,44} (see the Supplementary Information for extended bibliography).

Stereochemical models

The results of DFT calculations are in agreement with the high diastereoselectivities (see the Supplementary Information for details). We then evaluated the role of the chiral NHC ligand that contains a pendant sulfonate moiety on enantioselectivity (Fig. 3a). The computational errors for modeling a charged species notwithstanding, we propose a similar steric and electronic environment as suggested formerly vis-à-vis enantioselective allylic substitutions effected by the same catalyst class²⁷. The sulfonate group is probably situated in the rear (**I**– II); this would allow for the large 3.5-bis-(2.4.6- $(i-Pr)_3$ -phenyl)phenyl moiety to obstruct the right side of the complex, and causes the sizeable B(pin) moiety to be situated in the less occupied left/front quadrant in I (Fig. 3a). In II, which would lead to the minor enantiomer, there is steric repulsion between the B(pin) group and the chiral ligand's N-aryl group, and thus the energy barrier would be higher (7.4 kcal/mol less favored than I). Consistent with the above analysis (Fig. 3a) the high energy of \mathbf{II} and the steric pressure involving the B(pin) moiety is reflected in a considerably widened CNHC_Cu_C1_C2 dihedral angle (177.2° vs. 151.3° in I). Calculations on the system containing the more diminutive NHC-Cu complex derived from 7a point to energetically similar processes (energy difference of 0.6 kcal/mol between III and IV; Fig. 3b), which is in agreement with the lower e.r. obtained when the NHC-Cu complex derived from 7a is involved (55:45 vs. 95:5 e.r., Table 2).

The stereochemical model offers a rationale for why reactions with imines containing longer linear alkyl groups (e.g., n-butyl or 4-pentenyl) are less enantioselective (Fig. 2c); these lower e.r. might partly arise from an increase in attractive London dispersion forces between the 3,5-bis-(2,4,6-(i-Pr)₃-phenyl)phenyl group and the substrate's alkyl chain^{45,46,47}. It is however more plausible that higher conformational mobility of the alkyl chains disrupt N \rightarrow Na chelation [less optimal C^{NHC}-Cu-C₁-C₂ dihedral angle in **I** (151.3°) vs. in **I** (131.8°)]. The smaller energy difference between anionic structures **V** and **VI** (2.2 kcal/mol, Fig. 2a) supports the notion that enantioselectivity would probably be lower without a sodium bridge, and that reaction via **VI** is likely the most competitive pathway versus the involvement of the most favored **I**. The C^{NHC}-Cu-C₁-C₂ and N-Cu-C₁-C^{NHC} dihedral angles of **V** and **VI** are close to the optimal parameters in **I**, implying that some strain induced by N \rightarrow Na association in **I** is released in **V**. With the less flexible isopropyl substituent in **3y** the aforementioned chelation may remain intact, allowing for higher enantioselectivity (96:4 e.r.).

Conclusions

The catalytic method introduced here puts forth an expeditious strategy for synthesis of α -tertiary amines in high diastereo- and enantiomeric purity, thus providing a solution to an important and persisting problem in catalytic enantioselective synthesis. There are no more than a small number of catalytic enantioselective protocols that allow access to such coveted N-containing compounds, yet none involves an unprotected/unactivated imine. These investigations provide the first step towards development of a series of catalytic enantioselective reactions involving N-H ketimines and other types of readily available and versatile carbon-based nucleophiles, protocols that render a range of chiral drug candidates with one or more α -tertiary amine moieties much more accessible. Finally, this study further expands the utility of sulfonate-containing chiral NHC ligands, previously utilized in catalytic enantioselective conjugate additions ⁴⁸, allylic substitutions ⁴⁹ as well as copperboryl additions to alkenes ⁵⁰ and allenes ⁵¹ and copper-hydride additions to alkenes ⁵², to include allyl additions to ketimines.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This research was supported by a grant from the National Institutes of Health (GM-57212). H. J. was supported as a LaMattina Graduate Fellow in Chemical Synthesis. We thank F. Meng and J. Lee for helpful discussions.

References

- Riant O, Hannedouche J. Asymmetric catalysis for the construction of quaternary carbon centres: nucleophilic addition on ketones and ketimines. Org. Biomol. Chem. 2007; 5:873–888. [PubMed: 17340001]
- Shibasaki M, Kanai M. Asymmetric synthesis of tertiary alcohols and α-tertiary amines via Cucatalyzed C–C bond formation to ketones and ketimines. Chem. Rev. 2008; 108:2853–2873.
 [PubMed: 18570481]
- Kobayashi S, Mori Y, Fossey JS, Salter MM. Catalytic enantioselective formation of C–C bonds by addition to imines and hydrazones: A ten-year update. Chem. Rev. 2011; 111:2626–2704. [PubMed: 21405021]
- Clayden J, Donnard M, Lefranc J, Tetlow DJ. Quaternary centres bearing nitrogen (α-tertiary amines) as products of molecular rearrangements. Chem. Commun. 2011; 47:4624–4639.
- 5. Yus M, González-Gómez JC, Foubelo F. Catalytic enantioselective allylation of carbonyl compounds and imines. Chem. Rev. 2011; 111:7774–7854. [PubMed: 21923136]
- 6. Wada R, Shibuguchi T, Makino S, Oisaki K, Kanai M, Shibasaki M. Catalytic enantioselective allylation of ketoimines. J. Am. Chem. Soc. 2006; 128:7687–7691. [PubMed: 16756326]
- 7. Trost BM, Silverman SM. Enantioselective construction of highly substituted pyrrolidines by palladium- catalyzed asymmetric [3+2] cycloaddition of trimethylenemethane with ketoimines. J. Am. Chem. Soc. 2010; 132:8328–8238.
- 8. Luo Y, Hepburn HB, Chotsaeng N, Lam HW. Enantioselective rhodium-catalyzed nucleophilic allylation of imines with allylboron reagents. Angew. Chem. Int. Ed. 2012; 51:8309–8313.
- 9. Hepburn H, Lam HW. The isomerization of allylrhodium intermediates in the rhodium-catalyzed nucleophilic allylation of cyclic imines. Angew. Chem. Int. Ed. 2014; 53:11605–11610.

 Nakamura S, Hyodo K, Nakamura M, Nakane D, Masuda H. Catalytic enantioselective allylation of ketimines by using palladium pincer complexes with chiral bis(imidazoline)s. Chem. Eur. J. 2013; 19:7304–7309. [PubMed: 23633426]

- 11. Tang TP, Ellman JA. Asymmetric synthesis of β-amino acids derivatives incorporating a broad range of substitution patterns by enolate additions to *tert*-butanesulfinyl imines. J. Org. Chem. 2002; 67:7819–7832. [PubMed: 12398509]
- Zhao Y-S, Liu Q, Tian P, Tao J-C, Lin G-Q. Copper-catalyzed asymmetric allylation of chiral Ntert-butanesulfinyl imines: dual stereocontrol with nearly perfect diastereoselectivity. Org. Biomol. Chem. 2015; 13:4174–4178. [PubMed: 25758913]
- Chen JL-Y, Aggarwal VK. Highly diastereoselective and enantiospecific allylation of ketones and imines using borinic esters: Contiguous quaternary stereogenic centers. Angew. Chem. Int. Ed. 2014; 53:10992–10996.
- Rabbat PMA, Valdez SC, Leighton JL. Phenol-directed enantioselective allylation of aldimines and ketimines. Org. Lett. 2006; 8:6119–6121. [PubMed: 17165944]
- Perl NR, Leighton JL. Enantioselective imidazole-directed allylation of aldimines and ketimines. Org. Lett. 2007; 9:3699–3701. [PubMed: 17685538]
- Dhudshia B, Tiburcio J, Thadani AN. Diastereoselective allylation and crotylation of *N*-unsubstituted imines derived from ketones. Chem. Commun. 2005:5551–5553.
- 17. Meng F, Jang H, Jung B, Hoveyda AH. Cu-catalyzed chemoselective preparation of 2-(pinacolato)boron-substituted allylcopper complexes and their in situ site-, diastereo-, and enantioselective additions to aldehydes and ketones. Angew. Chem. Int. Ed. 2013; 52:5046–5051.
- Meng F, Haeffner F, Hoveyda AH. Diastereo- and enantioselective reactions of bis(pinacolato)diboron, 1,3-enynes, and aldehydes catalyzed by an easily accessible bisphosphine— Cu complex. J. Am. Chem. Soc. 2014; 136:11304–11307. [PubMed: 25089917]
- 19. Meng F, McGrath KP, Hoveyda AH. Multifunctional organoboron compounds for scalable natural product synthesis. Nature. 2014; 513:367–374. [PubMed: 25230659]
- Yeung K, Ruscoe RE, Rae J, Pulis AP, Procter DJ. Enantioselective generation of adjacent stereocenters in copper-catalyzed three-component coupling of imines, allenes and diboranes. Angew. Chem. Int. Ed. 2016; 55:11912–11916.
- Liu RY, Yang Y, Buchwald SL. Regiodivergent and diastereoselective CuH-catalyzed allylation of imines with terminal alkenes. Angew. Chem. Int. Ed. 2016; 55:14077–14080.
- Hou G, Gosselin F, Li W, McWilliams JC, Sun Y, Weisel M, O'Shea PD, Chen C, Davies IW, Zhang X. Enantioselective hydrogenation of N–H imines. J. Am. Chem. Soc. 2009; 131:9882–9883. [PubMed: 19569686]
- Tran DN, Cramer N. syn-Selective rhodium(I)-catalyzed allylation of ketimines proceeding through a directed C–H activation/allene addition sequence. Angew. Chem. Int. Ed. 2010; 49:8181–8184.
- 24. Thompson, LA., et al. Compounds for the reduction of beta-amyloid production. United States Patent US. 2013/0131051 A1.
- 25. Butler CR, et al. Discovery of a series of efficient, centrally efficacious BACE1 inhibitors through structure- based drug design. J. Med. Chem. 2015; 58:2678–2702. [PubMed: 25695670]
- 26. Gao F, Carr JL, Hoveyda AH. A broadly applicable NHC–Cu-catalyzed approach for efficient, site-and enantioselective coupling of readily accessible (pinacolato)alkenylboron compounds to allylic phosphates and applications to natural product synthesis. J. Am. Chem. Soc. 2014; 136:2149–2161. [PubMed: 24467274]
- 27. Shi Y, Jung B, Torker S, Hoveyda AH. N-Heterocyclic carbene–copper-catalyzed group-, site-, and enantioselective allylic substitution with a readily accessible propargyl(pinacolato)boron reagent: Utility in stereoselective synthesis and mechanistic attributes. J. Am. Chem. Soc. 2015; 135:8948–8964
- 28. Brown KM, May TL, Baxter CA, Hoveyda AH. All-carbon quaternary stereogenic centers by enantioselective Cu-catalyzed conjugate additions promoted by a chiral N-heterocyclic carbene. Angew. Chem. Int. Ed. 2007; 46:1097–1100.

29. Dabrowski JA, Villaume MT, Hoveyda AH. Enantioselective synthesis of quaternary carbon stereogenic centers through copper-catalyzed conjugate additions of aryl- and alkylaluminum reagents to acyclic trisubstituted enones. Angew. Chem. Int. Ed. 2013; 52:8156–8159.

- 30. Peese KM, Gin DY. Asymmetric synthetic access to the hetisine alkaloids: Total synthesis of (+)-nominine. Chem. Eur. J. 2008; 14:1654–1665. [PubMed: 18046691]
- 31. Takemura H, Nakashima S, Kon N, Yasutake M, Shinmoyozu T, Inazu T. A study of C–F•••M⁺ interaction: Metal complexes of fluorine containing cage compounds. J. Am. Chem. Soc. 2001; 123:9293–9298. [PubMed: 11562211]
- 32. Yamazaki T, Kawashita S, Kitazume T, Kubota T. Diastereoselective alkylation of glycinates by assistance of intramolecular potassium•••fluorine interactions. Chem. Eur. J. 1999; 15:11461–11464.
- Sazarin Y, Liu B, Maron L, Carpentier J-F. Discrete, solvent-free alkaline-earth metal cations: Metal•••fluorine interactions and ROP catalytic activity. J. Am. Chem. Soc. 2011; 133:9069–9087. [PubMed: 21545119]
- Julian LD, Hartwig JF. Intramolecular hydroamination of unbiased and functionalized primary aminoalkenes catalyzed by a rhodium aminophosphine complex. J. Am. Chem. Soc. 2010; 132:13813–13822. [PubMed: 20839807]
- 35. Musacchio AJ, Nguyen LQ, Beard GH, Knowles RR. Catalytic olefin hydroamination with aminium radical cations: A photoredox method for direct C–N bond formation. J. Am. Chem. Soc. 2014; 136:12217–12220. [PubMed: 25127420]
- 36. Du Y, Xu L-W, Shimizu Y, Oisaki K, Kanai M, Shibasaki M. Asymmetric reductive Mannich reactions to ketimines by a C(I) complex. J. Am. Chem. Soc. 2008; 130:16146–16147. [PubMed: 18998691]
- 37. Hayashi M, Iwanaga M, Shiomi N, Nakane D, Masuda H, Nakamura S. Direct asymmetric Mannich-type reaction of α-isocyanoacetates with ketimines using cinchona alkaloid/copper(II) catalysts. Angew. Chem. Int. Ed. 2014; 53:8411–8415.
- 38. Wieland LC, Vieira EM, Snapper ML, Hoveyda AH. Ag-catalyzed diastereo- and enantioselective vinylogous Mannich reactions of α-ketoimine esters. Development of a method and investigation of its mechanism. J. Am. Chem. Soc. 2009; 131:570–576. [PubMed: 18980303]
- 39. Kano T, Song S, Kubota Y, Maruoka K. Highly diastereo- and enantioselective Mannich reactions of synthetically flexible ketimines with secondary amine organocatalysts. Angew. Chem. Int. Ed. 2012; 51:1191–1194.
- 40. Lipshutz BH, Shimizu H. Copper(I)-catalyzed asymmetric hydrosilylations of imines at ambient temperature. Angew. Chem. Int. Ed. 2004; 43:2228–2230.
- Niu J, Guo P, Kang J, Li Z, Xu J, Hu S. Copper(I)-catalyzed aryl bromides to form intermolecular and intramolecular carbon–oxygen bonds. J. Org. Chem. 2009; 74:5075–5078. [PubMed: 19476328]
- 42. Wang JL, et al. The novel benzopyran class of selective cyclooxygenase-2 inhibitors. Part 2: The second clinical candidate having a shorter and favorable human half-life. Bioorg. Med. Chem. Lett. 2010; 20:7159–7163. [PubMed: 20709553]
- Shavnya A, Coffey SB, Smith AC, Mascitti V. Palladium-catalyzed sulfination of aryl and heteroaryl halides: Direct access to sulfones and sulfonamides. Org. Lett. 2013; 15:6226–6229. [PubMed: 24256546]
- 44. Ye X-Y, et al. Synthesis and structure–activity relationship of dihydrobenzofuran derivatives as novel human GPR119 agonists. Bioorg. Med. Chem. Lett. 2014; 24:2539–2545. [PubMed: 24755425]
- 45. Wagner JP, Schreiner PR. London Dispersion in Molecular Chemistry Reconsidering Steric Effects. Angew. Chem. Int. Ed. 2015; 54:12274–12296.
- Chen L, Ren P, Carrow BP. Tri(1-adamantyl)phosphine: Expanding the Boundary of Electron-Releasing Character Available to Organophosphorus Compounds. J. Am. Chem. Soc. 2016; 138:6392–6395. [PubMed: 27164163]
- Albers L, Rathjen S, Baumgartner J, Marschner C, Müller T. Dispersion-Energy-Driven Wagner-Meerwein Rearrangements in Oligosilanes. J. Am. Chem. Soc. 2016; 138:6886–6892. [PubMed: 27195490]

48. Slutskyy Y, Jamison CR, Lackner GL, Müller DS, Dieskau AP, Untiedt NL, Overman LE. Short enantioselective total syntheses of *trans*-clerodane diterpenoids: Convergent fragment coupling using a *trans*-decalin tertiary radical generated from a tertiary alcohol precursor. J. Org. Chem. 2016; 81:7029–7035. [PubMed: 27254137]

- 49. Shi Y, Jung B, Torker S, Hoveyda AH. N-Heterocyclic carbene–copper-catalyzed group-, site-, and enantioselective allylic substitution with a readily available propargyl(pinacolato)boron reagent: Utility in stereoselective synthesis and mechanistic attributes. J. Am. Chem. Soc. 2015; 137:8948–8964. [PubMed: 26172476]
- 50. Lee Y, Hoveyda AH. Efficient boron–copper additions to aryl-substituted alkenes promoted by NHC-based catalysts. Enantioselective Cu-catalyzed hydroboration reaction. J. Am. Chem. Soc. 2009; 131:3160–3161. [PubMed: 19256564]
- Jang H, Jung B, Hoveyda AH. Catalytic enantioselective protoboration of disubstituted allenes. Access to alkenylboron compounds in high enantiomeric purity. Org. Lett. 2014; 16:4658–4661. [PubMed: 25153792]
- 52. Lee J, Torker S, Hoveyda AH. Versatile homoallylic boronates by chemo-, S_N2^2 -, diastereo- and enantioselective catalytic sequence of Cu–H addition to vinyl-B(pin)/allylic substitution. Angew. Chem. Int. Ed. 2017; 56:821–826.

Shibasaki & Kanai, 2006

NBn
G Me
Dis-phosphine-Cu
catalyst

Frost, 2010

NTs
G Me
catalyst

Phosphoramidite-Pd
catalyst

Me₃Si
OAc

OAc

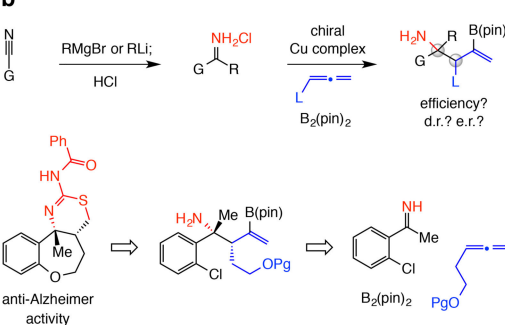


Figure 1. State-of-the-art in allyl additions to ketimines and goals of this study

There are significant exisiting limitations and a number of compelling issues remain unaddressed. **a,** There are only a small number of methods for catalytic enantioselective addition of an allyl group to a ketimine. The substrate is typically equipped with an activating/protecting group, which might prove difficult to remove in the presence of similar functional groups within a product structure (e.g., another *N*-benzylamine). **b,** A direct approach to synthesis of α -tertiary amines may entail preparation of the requisite unprotected N-H ketimine through alkylation of readily available nitriles followed by catalytic site-, diastereo- and enantioselective multicomponent addition of 2-boryl-

substituted allyl groups. One application relates to synthesis of the core tricyclic structure of a set of heterocyclic molecules that exhibit strong anti-Alzheimer activity. Bn, benzyl; Ts, tosyl; Ac, acyl; pin, pinacolato; G, R, L, functional groups; Pg, protecting group.

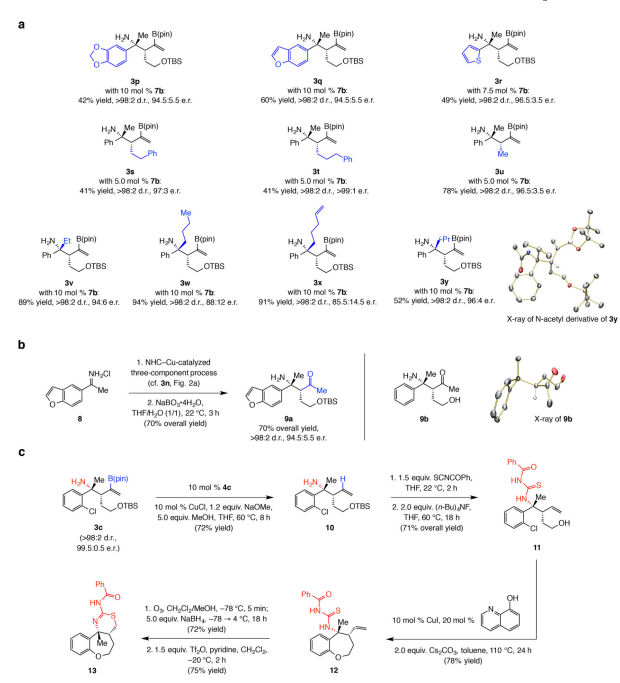


Figure 2. Further exploration of scope and illustration of utility

A variety of desirable products can be synthesised. **a,** The method is applicable to a variety of heterocyclic substrates and allenes. Products derived from ketimines containing *n*-alkyl or *iso*-alkyl substituents (vs. methyl) can be obtained efficiently, in >98:2 d.r. and 85.5:14.5–96:4 e.r., depending on the substituent identity. For results with achiral imidazolinium salt 4c, see the Supplementary Information Table 1. b, Oxidation of the alkenylboronate moiety within the products derived from the NHC–Cu-catalyzed multicomponent reactions proceed efficiently to deliver the corresponding β -amino ketones (e.g., 9a), which represent the products of diastereo- and enantioselective Mannich-type additions. c, The method may be

applied to the synthesis of the polycyclic core of compounds recently implicated in the fight against Alzheimer's disease. Conversion of the C–B(pin) to a C–H bond promoted by a readily accessible NHC–Cu complex afforded 10. Formation of the derived thiourea and another NHC–Cu-catalyzed reaction generated the oxepane ring of 12. A two-step procedure involving oxidative cleavage/reduction and activation of the resulting primary alcohol delivered the desired aminothiazine ring and the strained tricyclic 13. Reactions were performed under N_2 ; there was >98% disappearance of ketimine in all cases (might include decompositiopn products). Yields correspond to isolated and purified products and represent an average of at least three runs ($\pm 5\%$). Diastereomeric ratios were determined by analysis of the 400 MHz 1 H NMR spectra of unpurified product mixtures ($\pm 2\%$). Enantiomeric ratios were determined by HPLC analysis ($\pm 1\%$). See the Supplementary Information for experimental details and spectroscopic analyses.

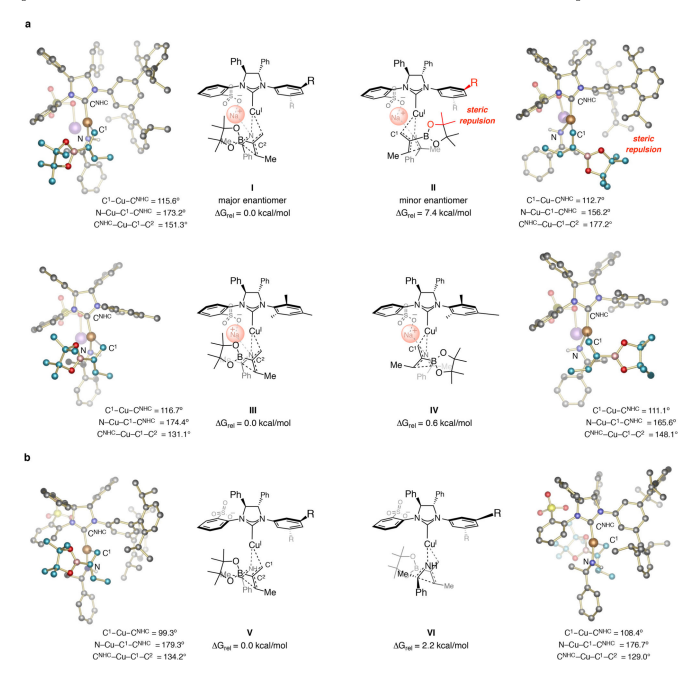
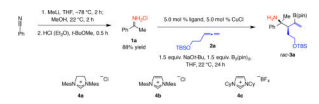


Figure 3. Stereochemical models

DFT calculations shed light on the origins of enantioselectivity. **a**, Transition states with a $N \rightarrow Na$ interaction account for high e.r.; **I** represents the preferred mode. **b**, The model suggests that disruption of the $N \rightarrow Na$ coordination by the long, flexible alkyl ketimine chain ($3\mathbf{w}$, \mathbf{x} Figure 2) in **V** might render mode **VI** competitive, leading to lower e.r. Free energy values relative to the major pathway refer to the MN12SX/Def2TZVPP_{THF(PCM)} level after geometry optimization performed with either MN12SX/Def2SVP_{THF(PCM)} (for **a**) or M06L/Def2SVP_{THF(PCM)} (for **b** and **c**). For details, see Sections 4 and 5 of the the

Table 1

Probing efficiency and diastereoselectivity with different (achiral) catalyst types.



Entry	Ligand	Conv. (%)*	Yield (%) [†]	d.r.*
1	PPh ₃	60	49	>98:2
2	PCy_3	71	52	>98:2
3	rac-binap	75	72	>98:2
4	4a	80	66	>98:2
5	4b	55	53	>98:2
6	4c	>98	90	>98:2

Reactions were carried out under N2 atmosphere; 1.2:1 ratio of ketimine:allene was used.

Abbreviations: TBS, *tert*-butyldimethylsilyl; pin, pinacolato; Mes, 2,4,6-Me₃C₆H₂; Cy, cyclohexyl; *rac*, racemic; binap, 2,2'-bis(diphenylphosphino)-1,1'-binaphthalene.

^{*} Conversion (based on allene consumption; (includes desired and decomposition products) and diastereomeric ratio (d.r.) values were measured by analysis of 400 MHz 1 H NMR spectra of unpurified mixtures; variance of values is estimated to be <±2%.

Table 2

Studies to identify an effective chiral catalyst.

Entry	Ligand	Conv. (%)*	Yield (%) [†]	e.r. ^{††}
1	R-binap	70	65	44:56
2	5a	55	51	18:82
3	5b	65	51	19.5:80.5
4	5c	29	27	69:31
5	6	80	51	40:60
6	7a	>98	65	55:45
7	7b	54	51	95:5

Reactions were carried out under N2 atmosphere; 1.2:1 ratio of ketimine:allene was used.

Abbreviations: TBS, *tert*-butyldimethyl silyl; pin, pinacolato; Mes, 2,4,6-Me₃C₆H₂; Cy, cyclohexyl; binap, 2,2'-bis(diphenylphosphino)-1,1'-binaphthalene.

^{*}Conversion (consumption of allene; (includes desired and decomposition products) and d.r. was measured by analysis of 400 MHz 1 H NMR spectra of unpurified mixtures; variance of values is estimated to be <±2%.

 $^{^{\}dagger}$ Yield of isolated and purified products (<±5%).

 $[\]dot{7}\dot{7}$ Enantiomeric ratio (e.r.) values were determined by HPLC analysis ($<\pm1\%$) (see the Supplementary Information for details).

Table 3

Catalytic diastereo- and enantioselective additions to N-H ketimines.

Entry	G	Mol %; Yield (%) [†]	e.r. ^{††}
1	$C_6H_5(\mathbf{a})$	7.5; 76	95:5
2	$o ext{-MeOC}_6 ext{H}_4\left(\mathbf{b}\right)$	10; 95	98.5:1.5
3	o-ClC ₆ H ₄ (c)	7.5; 91	99.5:0.5
4	$o ext{-} ext{FC}_6 ext{H}_4\left(\mathbf{d} ight)$	7.5; 72	97.5:2.5
5	o,o - $F_2C_6H_3$ (e)	7.5; 91	98:2
6	1-naphthyl (f)	10; 66	98.5:1.5
7	$o ext{-} ext{MeC}_6 ext{H}_4\left(\mathbf{g} ight)$	7.5; 59	99.5:0.5
8	m-FC ₆ H ₄ (h)	5.0; 64	94:6
9	m -MeOC $_6$ H $_4$ (\mathbf{i})	10; 57	98.5:1.5
10	2-naphthyl (j)	7.5; 81	93:7
11	$p\text{-ClC}_6\text{H}_4\left(\mathbf{k}\right)$	7.5; 56	92.5:7.5
12	$p ext{-FC}_6 ext{H}_4$ (I)	5.0; 71	95:5
13	p-MeOC ₆ H ₄ (m)	7.5; 39	92.5:7.5
14	$\text{CyCH}_2\left(\mathbf{n}\right)$	10; 38	91:9
15	Cy (o)	10; 48	95:5

Reactions were carried out under N2 atmosphere; >98% disappearance of ketimine in all cases (includes desired and decomposition products).

Abbreviations: TBS, tert-butyldimethylsilyl; pin, pinacolato; Cy, cyclohexyl.

Experiments were performed at least in triplicate. See the Supplementary Information for details and the results with achiral imidazolinium salt 4c.

 $^{^{\}dagger}$ Yield of isolated and purified products (<±5%).

 $[\]dot{\tau}\dot{\tau}$ Enantiomeric ratios determined by HPLC analysis (<±1%; see the Supplementary Information for details).