

Cp₂ZrMeCl: A reagent for asymmetric methyl addition.

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Supporting Information Placeholder

ABSTRACT: The use of Cp₂ZrMeCl is described as a source of nucleophilic methyl in asymmetric catalysis. This easily prepared reagent is bench stable, weighable in air, and generally useful in highly enantioselective copper-catalyzed addition reactions at room temperature. Methyl is successfully (generally > 90% *ee*) added in 1,4-additions to cyclic and acyclic α,β -unsaturated ketones, to provide tertiary and quaternary centers. Examples of catalyst controlled diastereoselective 1,6-addition and dynamic kinetic asymmetric allylic alkylation reactions are also reported. The reagent is used in the catalytic asymmetric synthesis of naturally occurring fragrance (*R*)-(-)-muscone (82% yield, 91% *ee*).

Many biologically active compounds feature stereocentres bearing methyl groups, making methods for the asymmetric addition of Me-nucleophiles extremely important. Both tertiary and quaternary centers containing methyl groups are widely represented in natural products¹ and clinically used medicines.² Asymmetric conjugate additions (ACAs) of alkyl groups to electron-deficient α,β -unsaturated systems have been widely developed.^{3–7} ACAs to add methyl rely on Cu-catalysis, and asymmetric methods developed for addition of other alkyl nucleophiles (for example, ethyl) are sometimes suitable, but very often problematic, with methyl nucleophiles.³

While Me₃Al can be used to effectively add a methyl group in ACAs,^{8,9} the reactivity profile of the reagent presents real safety concerns.¹⁰ Additionally, asymmetric additions with Me₃Al must be performed at cryogenic temperatures, and the reagents Lewis acidity renders it incompatible with many functional groups, limiting applications in complex molecule synthesis.

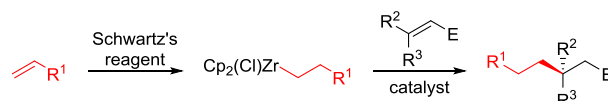
Other organometallic sources of nucleophilic Me also have limitations. Dimethyl zinc spontaneously combusts upon exposure to air and reacts violently in a number of common laboratory situations. Conversely, methyl Grignard reagents tend to have less aggressive reactivity profiles than other Grignard reagents, attributed to aggregation of MeMgX species.¹¹ However, the importance of asymmetric Me-addition has led to several useful procedures using Me-Grignard reagents to acyclic enones,¹² a variety of electrophilic thioester acceptors,^{13–15} and Loh's protocol which may be used with unsaturated esters.^{16,17} Despite high reactivity to air and water, Me₂Zn also shows sluggish reactivity in ACAs, requiring an excess of reagent and long reaction times – especially designed acceptors have recently been reported to address these issues.¹⁸

Alkylzirconium reagents can be used in ACAs (Scheme 1a) to acyclic¹⁹ and cyclic^{20–24} enones, lactones,²⁵ 1,4- and 1,6-additions to functionalized steroid derivatives,^{26,27} in the formation of quaternary centers, and remote asymmetric C-H activations sequences initiated by alkene isomerization.²⁸ These highly enantioselective reactions generally proceed at room temperature although in

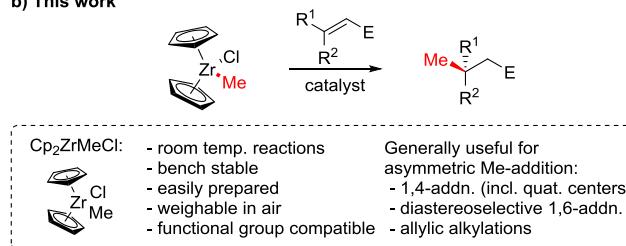
some cases better results are obtained at 0 °C. Recently, alkylzirconium species have been shown to undergo dynamic kinetic asymmetric kinetic transformations to allow highly enantioselective allylic alkylations.^{29,30} These methods generally work in a variety of solvents and have wide functional group tolerance. In all cases the nucleophilic zirconium species were prepared by hydrometallation of alkenes using Schwartz's reagent (Cp₂ZrHCl), and so the use of methyl nucleophile was not possible.

Scheme 1. Asymmetric addition of Me-groups

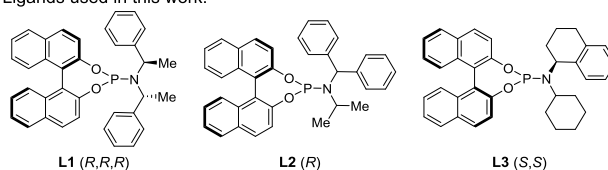
a) Previous work with alkyl zirconium reagents required hydrometallation



b) This work



Ligands used in this work:

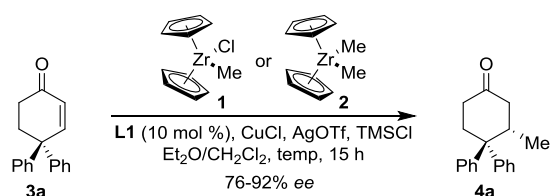


Recognizing the potential importance of a nucleophilic Me-reagent that could operate at room temperature in a variety of solvents, we hoped that the reactivity profile of Cp₂ZrMeCl **1** would be similar to the zirconocenes described above. We examined readily available zirconocenes **1** and **2** (Scheme 2) and began by applying conditions previously developed for hydrozirconated alkenes to cyclohexenones. Enone **3a** was used for testing because the product is non-volatile.

Dimethyl zirconocene **2** gave encouraging preliminary results (*ee*'s over 75%) for the room temperature ACA shown in Scheme 2, but we found that **2** slowly degrades when stored at room temperature, even under an inert atmosphere. In contrast, chloro(methyl)zirconocene **1** is stable for at least 6 months on the bench without any noticeable changes. Control experiments showed that Cp₂ZrMeCl is unreactive to enone **3a** in the absence of a catalyst and no traces of 1,2- nor 1,4-addition products were observed after 4 days at room temperature in Et₂O/CH₂Cl₂.

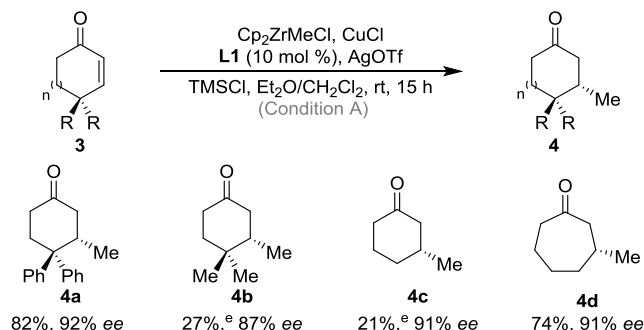
Cp_2ZrMeCl ³¹ is readily prepared on a 10-gram scale from Cp_2ZrCl_2 (see SI for protocol), although this current procedure is straightforward it does use Me_3Al as the source of methyl.

Scheme 2. Examining readily available Me-zirconocenes



Reported conditions²⁴ for the asymmetric addition of alkyl zirconocenes to **3a** were highly effective with Cp_2ZrMeCl , and screening temperatures and solvents only gave moderate improvement. Optimized conditions use 1.6 equiv of Cp_2ZrMeCl with 10 mol % copper(I)triflate and **L1** (10 mol %), in a mixture of Et₂O and CH₂Cl₂ at room temperature (**4a** obtained in 82%, 92% ee, Scheme 3). The role of silver salts in these reactions appears to be limited to counterion exchange with CuCl. These conditions (condition A) were applied to other cyclic enones **3** (Scheme 3). 6-Membered ring substrates **3b** and **3c** (R = Me or H) gave ACA products **4b** and **4c** with high ee's (87% and 91%, respectively). Low isolated yields (27% and 21%, respectively) are due to product volatility but complete conversion (TLC control) and clean crude NMR spectra were obtained. Cyclopentenone gave a complex mixture of products as expected,^{6,32} and the 7-membered ring enone **3d** gave **4d** (74% yield, 91% ee).

Scheme 3. ACAs to form tertiary centers.^{a-d}



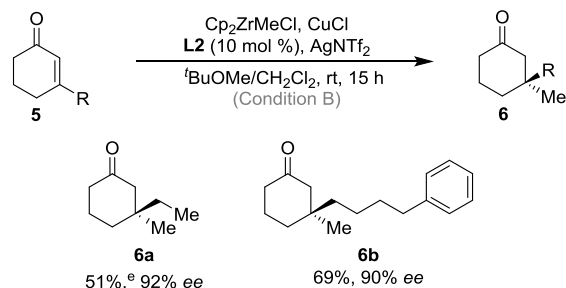
^areactions performed on 0.4 mmol scale; ^bisolated yields; ^cee determined by chiral GC or chiral HPLC; ^dcondition A: Cp_2ZrMeCl (1.6 equiv), **L1** (10 mol %), CuCl (0.10 equiv), AgOTf (0.11 equiv), TMSCl (5.0 equiv), 5:1 Et₂O/CH₂Cl₂, rt, 15 h; ^evolatile product.

On more challenging all-carbon quaternary centers, applying conditions B (Scheme 4) to 3-substituted enones **5**, gave excellent results.^{20,21} Cyclohexanones **6a** and **6b** were obtained in good yield (51% and 69%, respectively) with high ee (92% and 90%).

Other substrates were then assessed (Table 1). ACA of methylzirconocene **1** to 5-membered protected-acetal **7**²² using related conditions (condition C) and ligand **L3** at 0 °C cleanly afforded methyl adduct **8** in 57% yield with excellent (94%) ee.

Linear enones such as **9** and **10**¹⁹ can also be used, with far shorter reaction times (~45 min) than cyclic enones. TLC monitoring is essential to avoid by-products, but quenching at the appropriate time gave very clean crude reaction mixtures and β -methyl ketones **10** and **12** with high ee's (92% and 94%). The yield of **12** was excellent (84%), while **10** suffers from volatility (29%).

Scheme 4. ACAs to form quaternary centers.^{a-d}



^areactions performed on 0.4 mmol scale; ^bisolated yields; ^cee determined by chiral GC or chiral HPLC; ^dcondition B: Cp_2ZrMeCl (1.6 equiv), **L2** (10 mol %), CuCl (0.10 equiv), AgNTf₂ (0.05 equiv), 5:1 ^tBuOMe/CH₂Cl₂, rt, 15 h; ^evolatile product.

Table 1. Other substrates^a

substrate		conditions		product	
		up to 94% ee			
entry	substrate	product	conditions ^b	yield, % ^c	ee, %
1			C	57	94 ^d
2			D	29 ^e	92 ^f
3			D	84	94 ^f
4			E	61 ^g	5.1:1 crude dr
5			F	72 ^h	94 ⁱ

^areactions performed on 0.4 mmol scale; ^bcondition C: Cp_2ZrMeCl (2.5 equiv), **L2** (22 mol %), CuCl (0.20 equiv), AgOTf (0.20 equiv), TMSCl (5.0 equiv), 5:1 Et₂O/CH₂Cl₂, 0 °C, 15 h; ^ccondition D: Cp_2ZrMeCl (1.6 equiv), **L3** (10 mol %), CuCl (0.10 equiv), AgOTf (0.11 equiv), TMSCl (5.0 equiv), 5:1 Et₂O/CH₂Cl₂, 0 °C, 45 min; ^dcondition E: Cp_2ZrMeCl (1.6 equiv), *ent*-**L1** (10 mol %), CuCl (0.10 equiv), AgOTf (0.11 equiv), TMSCl (5.0 equiv), 1:1 Et₂O/CH₂Cl₂, rt, 15 h; ^econdition F: Cp_2ZrMeCl (1.6 equiv), **L1** (10 mol %), CuI (0.10 equiv), CDCl₃, rt, 15 h; ^fisolated yield unless stated otherwise; ^gdetermined by chiral HPLC; ^hvolatile product; ⁱdetermined by chiral GC; ^jpure major isomer only; ^kNMR yield against internal standard; ^ldetermined by chiral GC analysis of epoxidised crude mixture of **16**.

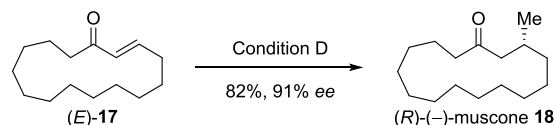
We next tested the compatibility of the reagent with complex molecules using commercially available steroid **13**.²⁶ **13** bears a number of stereogenic centers as well as an acetate and is capable of 1,2-, 1,4- and 1,6-additions. In the event, using conditions E

gave a separable mixture (5.1:1 crude *dr*) of 2 diastereomers where **14** could be isolated as a pure single isomer in 61% yield.

We also examined Cp₂ZrMeCl in an asymmetric allylic alkylation reaction with racemic allyl chloride **15**.²⁹ This Cu-catalyzed dynamic kinetic asymmetric transformation gave excellent results with 72% yield (based on an NMR standard) and 94% *ee* (GC analysis of epoxidized crude material).

Finally, we synthesized (*E*)-enone **17** from commercially available cyclopentadecanone³³ and applied conditions D due to the low rigidity of the 15-membered macrocycle.³⁴ These conditions, developed for linear substrates, gave a quick (45 min) and clean reaction to afford natural product (*R*)-(-)-muscone **18** (82% yield, 91% *ee*) which compares favorably with previous catalytic asymmetric approaches to this fragrance.^{34–39}

Scheme 5. Synthesis of natural muscone.^{a,b}



^aisolated yield; ^b*ee* determined by GC analysis of crude material reduced to the corresponding alcohols.

In conclusion, we have shown that Cp₂ZrMeCl can be used as a methylating agent in a wide variety of copper-catalyzed asymmetric reactions. Excellent levels of enantioselectivity (87–94% *ee*) were obtained in all cases. These procedures occur by a variety of mechanisms, tolerate acid-labile functional groups such as acetals and esters, and have been used in short synthesis of a natural product. Cp₂ZrMeCl was easily synthesized on a 10-gram scale in our laboratory and presumably this scale can be increased. The reagent is a readily manipulated crystalline powder that can be weighed out in air and is stable for at least 6 months when stored on the bench under inert gas.

Ongoing studies in our laboratory aim to expand the range of asymmetric reactions alkyl zirconocene species can be used in and will be reported in due course.

ASSOCIATED CONTENT

Supporting Information

All procedures, characterization data, NMR spectra, GC and HPLC traces are provided in the supporting information. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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- (lisdexamfetamine), Shire Pharmaceuticals Group; d) Symbicort® (budesonide), AstraZeneca; e) Xeloda® (capecitabine), Roche.
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