Regioselective Simmons–Smith-type cyclopropanations of polyalkenes enabled by transition metal catalysis†

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A \([\text{PDI}][\text{CoBr}_2] \) complex (PDI = pyridine-diimine) catalyzes Simmons–Smith-type reductive cyclopropanation reactions using \(\text{CH}_2\text{Br}_2\) in combination with \(\text{Zn}\). In contrast to its non-catalytic variant, the cobalt-catalyzed cyclopropanation is capable of discriminating between alkenes of similar electronic properties based on their substitution patterns: monosubstituted > 1,1-disubstituted > (Z)-1,2-disubstituted > (E)-1,2-disubstituted > trisubstituted. This property enables synthetically useful yields to be achieved for the monocyclopropanation of polyalkene substrates, including terpene derivatives and conjugated 1,3-dienes. Mechanistic studies implicate a carbenoid species containing both Co and Zn as the catalytically relevant methylene transfer agent.

Introduction

Cyclopropanes are common structural elements in synthetic and natural biologically active compounds.\(^1\) The Simmons–Smith cyclopropanation reaction was first reported over half a century ago but remains today one of the most useful methods for converting an alkene into a cyclopropane.\(^2\) As compared to diazomethane, which is shock sensitive and must be prepared from complex precursors, \(\text{CH}_2\text{I}_2\) is both stable and readily available, making it an attractive methylene source. Additionally, the stereospecificity of the Simmons–Smith reaction allows diastereomeric relationships in cyclopropanes to be established with a high degree of predictability. Several advances have addressed many of the limitations of the initial Simmons–Smith protocol. For example, \(\text{Et}_2\text{Zn}\) can be used in the place of \(\text{Zn}\) to more reliably and quantitatively generate the active carbenoid reagent.\(^3\) Acidic additives, such as \(\text{CF}_3\text{CO}_2\text{H}\)\(^4\) and substituted phenols,\(^5\) have been found to accelerate the cyclopropanation of challenging substrates. Finally, \(\text{Zn}\) carbenoids bearing dialkylphosphate anions\(^6\) or bipyridine ligands\(^7\) are sufficiently stable to be stored in solution at low temperatures (Fig. 1).

Despite the many notable contributions in \(\text{Zn}\) carbenoid chemistry, a persistent limitation of Simmons–Smith-type cyclopropanations is their poor selectivity when attempting to discriminate between multiple alkenes of similar electronic properties. For example, the terpene natural product limonene possesses a 1,1-disubstituted and a trisubstituted alkene. Friedel reported that, under a variety of \(\text{Zn}\) carbenoid conditions, the two alkenes are cyclopropanated with similar rates, resulting in mixtures of monocyclopropanated (up to a 5 : 1 ratio of regioisomers) and dicyclopropanated products.\(^8\) This issue is exacerbated by the challenge associated with separating the two monocyclopropane regioisomers, which only differ in the position of a non-polar \(\text{CH}_2\) group. In general, synthetically useful regioselectivities in Simmons–Smith reactions are only observed for substrates containing directing groups.\(^9\) In principle, catalysis may provide an avenue to address selectivity challenges in Simmons–Smith-type cyclopropanations; however, unlike diazoalkane transfer reactions,
which are catalyzed by a broad range of transition metal complexes,
there has been comparatively little progress toward the
development of robust catalytic strategies for reductive
cyclopropanations. Lewis acids in substoichiometric loadings
have been observed to accelerate the Simmons-Smith reaction,
but in many cases, this rate effect is restricted to allylic alcohol
substrates.

Recently, our group described an alternative approach to
catalyzing reductive cyclopropanation reactions using a transition
metal complex that is capable of activating the dihaloalkane reagent by C-X oxidative addition. A dinickel catalyst
was shown to promote methylene and vinylidene transfer
using CH2I2 and 1,1-dichloroalkenes in combination with Zn
as a stoichiometric reductant. Here, we describe a mononuclear
[PD1]Co (PD1 = pyridine-diimine) catalyst that imparts a high
degree of steric selectivity in the cyclopropanation of polyalkene
substrates. Mechanistic studies suggest that the key intermediate
responsible for methylene transfer is a heterobimetallic conjugate of Co and Zn.

Results and discussion

4-Vinyl-1-cyclohexene contains a terminal and an internal
alkene of minimal electronic differentiation and thus provided
a suitable model substrate to initiate our studies (Table 1). Under standard CH2I2/Et2Zn conditions (entry 1), there is a modest preference for cyclopropanation of the more electron-rich disubstituted alkene (rr = 1 : 6.7) with increasing amounts of competing dicyclopropanation being observed at higher conversions (entries 2 and 3). Other modifications to the conditions, including the use of a Brønsted acid (entry 4) or a Lewis acid additive (entry 5), did not yield any improvements in selectivity. Likewise, an Al carbeneid generated using CH2I2 and ALEt3 afforded a similar preference for cyclopropanation of the endocyclic alkene (entry 6).

In a survey of transition metal catalysts, the [PDI]CoBr2 complex 1 was identified as a highly regioselective catalyst for
the cyclopropanation of 4-vinyl-1-cyclohexene, targeting the less hindered terminal alkene (Table 2). CH2Br2 and Zn alone do not afford any background levels of cyclopropanation (entry 1); however, the addition of 6 mol% [PDI]CoBr2 (1) provided monocyclopropane 3 (81% yield) with a >50 : 1 rr and <1% of the dicyclopropane product (entry 5). The steric profile of the catalyst appears to be critically important for yield. For example, the mesityl (entry 6) and phenyl-substituted variants (entry 7) of the ligand provided only 58% and 4% yield respectively under the same reaction conditions. Related N-donor ligands similarly afforded low levels of conversion (entries 8–12) as did the use of other first-row transition metals, including Fe (entry 14) and Ni (entry 15), in the place of Co.

In order to define the selectivity properties of catalyst 1, we next conducted competition experiments using alkenes bearing different patterns of substitution (Fig. 2). Reactions were carried out using an equimolar amount of each alkene and run to full conversion of the limiting CH2Br2 reagent (1.0 equiv.). Mono-substituted alkenes are the most reactive class of substrates using 1 but are not adequately differentiated from 1,1-disubstituted alkenes (3 : 1). By contrast, terminal alkenes are significantly more reactive than internal alkenes, providing synthetically useful selectivities (≥31 : 1). Furthermore, a model Z-alkene was cyclopropanated in preference to its E-alkene congener in a 33 : 1 ratio. Using catalyst 1, trisubstituted alkenes are poorly reactive, and no conversion is observed for tetrasubstituted alkenes.

The synthetic applications of the catalytic regioselective cyclopropanation were examined using the terpene natural products and derivatives shown in Fig. 3. In all cases, the selectivity properties follow the trends established in the competition experiments. Substrates containing ether or free alcohol functionalities (e.g., 7, 10, and 11) exhibit a strong directing group effect under classical Simmons-Smith conditions; however, catalyst 1 overrides this preference and targets the less hindered alkene. Additionally, the presence of electron-deficient α,β-unsaturated carbonyl systems (e.g., 9, 13, and 14) do not perturb the expected steric selectivity.

Table 1  Regioselectivity studies using Zn and Al carbeneid reagents a

| Entry | Reaction conditions | Yield (3 + 4) | rr (3 : 4) | Yield 5 |
|-------|-------------------|-------------|-----------|--------|
| 1     | CH2I2 (1.0 equiv.), Et2Zn (0.5 equiv.) | 28% | 1 : 6.7 | 3% |
| 2     | CH2I2 (1.0 equiv.), Et2Zn (1.0 equiv.) | 33% | 1 : 4.6 | 5% |
| 3     | CH2I2 (2.0 equiv.), Et2Zn (2.0 equiv.) | 53% | 1 : 6.5 | 16% |
| 4     | CH2I2 (1.0 equiv.), Et2Zn (1.0 equiv.), 3,5-difluorobenzolic acid (2.0 equiv.) | 28% | 1 : 3.5 | 19% |
| 5     | CH2I2 (2.0 equiv.), Et2Zn (2.0 equiv.), TiCl4 (0.2 equiv.) | 13% | 1 : 4.6 | 1% |
| 6     | CH2I2 (1.2 equiv.), ALEt3 (1.2 equiv.) | 38% | 1 : 3.1 | 9% |

 a Reaction conditions: 4-vinylcyclohexene (0.14 mmol), CH2I2 (1.0 mL), 24 h, 22 °C. Yields and ratios of regioisomers were determined by GC analysis against an internal standard.
Vinylcyclopropanes are a valuable class of synthetic intermediates that engage in catalytic strain-induced ring-opening reactions. The monocyclopropanation of a diene represents an attractive approach to their synthesis but would require a catalyst that is capable of imparting a high degree of regioselectivity and avoiding secondary additions to form dicyclopropane products. These challenges are addressed for a variety of diene classes using catalyst 1 (Fig. 4). Over the substrates that we have examined, the selectivities for cyclopropanation of the terminal over the internal double bond of the diene system are uniformly high. Additionally, the catalyst is tolerant of vinyl bromide (15) and vinyl boronate (23) functional groups, which are commonly used in cross-coupling reactions.

Like the non-catalytic Simmons–Smith reaction, the cyclopropanation using 1 is stereospecific within the limit of detection, implying a mechanism in which the two C–C σ-bonds are either formed in a concerted fashion or by a stepwise process that does not allow for single bond rotation. For example, cyclopropanation of the Z-alkene 24 affords the cis-disubstituted cyclopropane 25 in 95% yield as a single diastereomer (Fig. 5a). Furthermore, the vinylcyclopropane substrates 26 and 28, commonly used as tests for cyclopropylcarbinyl radical intermediates, react without ring-opening to afford products 27 and 29 (Fig. 5b).

### Table 2 Catalyst structure–activity relationship studies

| Entry | Metal source | Ligand | Yield (3 + 4) | rr (3 : 4) | Yield 5 |
|-------|--------------|--------|--------------|------------|--------|
| 1     | —            | —      | <1%          | —          | <1%    |
| 2     | CoBr₂       | —      | <1%          | —          | <1%    |
| 3     | Co(DME)Br₂  | —      | <1%          | —          | <1%    |
| 4     | —            | i-PPDI | <1%          | —          | <1%    |
| 5     | CoBr₂       | i-PPDI | 81%          | >50 : 1    | <1%    |
| 6     | CoBr₂       | MBPDI  | 58%          | >50 : 1    | <1%    |
| 7     | CoBr₂       | PDPDI  | 4%           | —          | <1%    |
| 8     | CoBr₂       | i-PDAD | <1%          | —          | <1%    |
| 9     | CoBr₂       | i-PIP  | 2%           | —          | <1%    |
| 10    | CoBr₂       | Terpy  | 4%           | —          | <1%    |
| 11    | CoBr₂       | i-RPyBOX| <1%          | —          | <1%    |
| 12    | CoBr₂       | PhPyBOX| <1%          | —          | <1%    |
| 13    | CoBr₂       | PPh₃ (12 mol%) | <1% | — | 0% |
| 14    | FeBr₂       | i-C0PrDAD | <1% | — | <1% |
| 15    | NiBr₂       | i-PDPI  | <1%          | —          | <1%    |

The reaction conditions: 4-vinylcyclohexene (0.14 mmol), THF (1.0 mL), 24 h, 22 °C. Yields and ratios of regioisomers were determined by GC analysis against an internal standard.

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**Fig. 2** Intermolecular competition experiments probing selectivity based on alkene substitution patterns. Reactions were conducted using a 1 : 1 : 1 molar ratio of the two alkene starting materials and CH₂Br₂.

**Fig. 3** Catalytic regioselective monocyclopropanations of terpene natural products and derivatives. Isolated yields following purification are averaged over two runs.
Under standard catalytic conditions, the reaction mixtures using 1 adopt a deep violet color, which persists until complete consumption of the alkene. The UV-vis spectrum of the catalytic mixture at partial conversion is consistent with a Co(I) resting state (Fig. 6a). The authentic 

i/CoPrPDI]CoBr complex (30) can be prepared by stirring the 

i/CoPrPDI]CoBr2 complex 1 over excess Zn metal. Cyclic voltammetry data (Fig. 6b) indicates an $E_{1/2}$ for the Co(i)/Co(i) redox couple of $-1.00 \text{ V vs. } \text{Fc/Fc}^+$. The large peak-to-peak separation (0.96 V in 0.3 M [n-Bu4N][PF6]/THF) is characteristic of a slow bromide dissociation step following electron transfer. The second Co(i)/Co(0) reduction event is significantly more cathodic at $-1.93 \text{ V}$ and is inaccessible using Zn.

In order to decouple the cyclopropanation steps of the mechanism from catalyst turnover, we conducted stoichiometric reactions with the isolated 

i/CoPrPDI]CoBr complex in the absence of Zn (Fig. 6c). The reaction of 30 with 4-vinylcyclohexene and CH3Br2 generates the 

i/CoPrPDI]CoBr2 complex 1 within 24 h at room temperature but forms cyclopropanated products in a relatively low combined yield of 26%, which is not commensurate with the efficiency of the catalytic process. Furthermore, the regioselectivity is only 3 : 1, whereas the catalytic cyclopropanation achieves a >50 : 1 selectivity for this substrate. When the same stoichiometric reaction is conducted in the presence of ZnBr2, the yield and selectivity of the catalytic process is fully restored.

The Co-containing product (31) of the stoichiometric reaction in the presence of ZnBr2 is green, which is notably distinct
from the tan color of the $[^{125}\text{PrPDI}]\text{CoBr}_2$ complex 1. This green species is NMR silent but may be crystallized from saturated solutions in Et$_2$O to afford 31 (Fig. 6f). The solid-state structure reveals the expected $[^{125}\text{PrPDI}]\text{CoBr}_2$ fragment in a distorted square pyramidal geometry ($\tau_5 = 0.36$) with a Zn(THF/Et$_2$O)Br$_2$ Lewis acid coordinated to one of the Br ligands. This interaction induces an asymmetry in the structure, causing the Co–Br1 distance ($2.557(1)$ Å) to be elongated relative to the Co–Br2 distance ($2.358(2)$ Å).

Collectively, these studies suggest that both Co and Zn are present in the reactive carbenoid intermediate, and that ZnBr$_2$ may interact with the $[^{125}\text{PrPDI}]\text{Co}$ complex through Lewis acid–base interactions. There is a notable similarity between the observed Co/Zn effect and previous studies of Lewis acid acceleration in the Simmons–Smith cyclopropanation. For example, Zn carbenoid reactions are known to be accelerated by the presence of ZnX$_2$,

Conclusions

In summary, transition metal catalysis provides a pathway to accessing unique selectivity in reductive carbenoid transfer reactions. A $[^{125}\text{PrPDI}]\text{CoBr}_2$ complex functions as a robust catalyst for Simmons–Smith type cyclopropanation using a CH$_3$Br$_2$/Zn reagent mixture. This system exhibits the highest regioselectivities that have been observed in reductive cyclopropanations based solely on the steric properties of the alkene substrate. Accordingly, a range of terpenes and conjugated dienes may be converted to a single monocyclopropanated product. Ongoing studies are directed at exploring the applications of transition metal catalysts to other classes of carbenoid transfer reactions.

Conflicts of interest

There are no conflicts to declare.

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