

## Highly Z-Selective and Enantioselective Ring-Opening/Cross-Metathesis Catalyzed by a Resolved Stereogenic-at-Ru Complex

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## Supporting Information

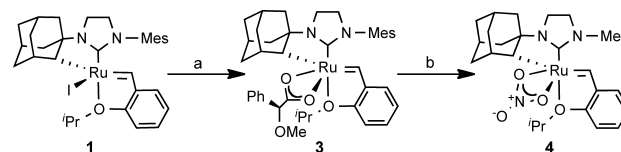
**ABSTRACT:** The synthesis of a ruthenium complex that catalyzes Z-selective (up to 98% Z) asymmetric ring-opening/cross-metathesis with high enantioselectivity (up to 95% ee) is reported. The synthesis of the catalyst features the resolution of a chelating N-heterocyclic carbene complex by ligand substitution with a chiral carboxylate.

Olefin metathesis is a powerful carbon–carbon bond-forming reaction that is widely used in organic synthesis,<sup>1</sup> polymer chemistry,<sup>2</sup> materials science,<sup>3</sup> and biochemistry.<sup>4</sup> Asymmetric olefin metathesis methodologies have proven useful for the synthesis of enantiopure natural products and other biologically relevant compounds.<sup>5</sup> Consequently, the development of chiral catalysts for methods such as asymmetric ring-opening/cross-metathesis (AROCM) is a field of ongoing interest.<sup>6</sup>

The earliest catalysts, which contained molybdenum, were capable of generating AROCM products in high ee (80–99%) but suffered from limited substrate scope and functional group compatibility.<sup>7</sup> Ruthenium-based catalysts wherein the chirality is built into the N-heterocyclic carbene (NHC) ligand have been developed.<sup>8</sup> Most of these molybdenum and ruthenium catalysts are capable of performing AROCM with high levels of *E* selectivity (up to >98% *E*).<sup>9</sup>

Recently, Z-selective AROCM of oxabicycles has been achieved with molybdenum catalysts.<sup>10</sup> While Z-selective AROCM has been accomplished with ruthenium catalysts, to date it has been limited to reactions involving heteroatom-substituted terminal olefin cross-partners.<sup>11</sup> Recent advances have produced ruthenium catalysts with chelating NHC ligands possessing exquisite Z selectivity in cross-metathesis.<sup>12</sup> We anticipated that enantiopure versions of the newly developed catalysts would exhibit high Z selectivity and enantioselectivity in AROCM because of the rigidity imparted by the Ru–C chelate. Herein we report a new homochiral stereogenic-at-ruthenium complex that exhibits high enantioselectivity in the AROCM of norbornene derivatives.

Enantioenriched **4** was synthesized by resolution as shown in Scheme 1. Treatment of racemic iodide **1**<sup>12c</sup> with silver carboxylate **2** cleanly formed a 1:1 mixture of diastereomers in 97% yield. Chromatographic separation of the mixture afforded a 45% yield of **3** (90% of the theoretical maximum) with >95:5 dr. The absolute stereochemistry of complex **3** was confirmed by X-ray crystallography (Figure 1). Sequential treatment of

Scheme 1. Synthesis of Homochiral Complex **4**<sup>a</sup>

<sup>a</sup>(a) (1) (S)-AgO<sub>2</sub>CCH(Ph)(OMe) (2) (2 equiv), THF, 23 °C, 1.5 h, 97%; (2) chromatographic separation, 45%. (b) (1) *p*TsOH·H<sub>2</sub>O, THF, 5 min; (2) NaNO<sub>3</sub>, THF/MeOH, 15 min, 43%.

carboxylate **3** with *p*-toluenesulfonic acid and sodium nitrate produced the enantioenriched nitrate complex **4** in 43% yield.

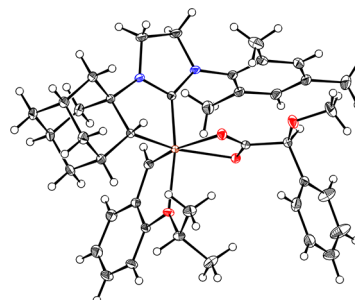
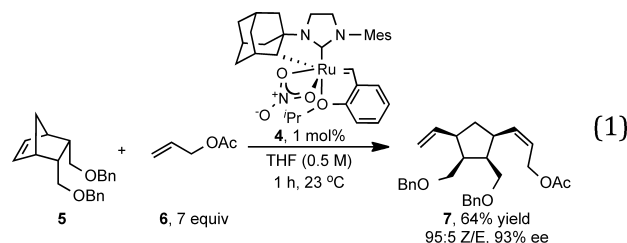


Figure 1. ORTEP drawing of **3**.

While complex **3** exhibited low enantioselectivity in AROCM, a 1 mol % loading of complex **4** catalyzed the reaction of norbornene derivative **5** with excess allyl acetate (**6**) to produce a 64% yield of the diene (1*S*,2*R*,3*S*,4*R*)-**7** with 95% Z selectivity and 93% ee (eq 1).<sup>13</sup> The highly selective reaction



produces four contiguous stereocenters in a tetrasubstituted cyclopentane ring. Optimization of the process revealed that **7**

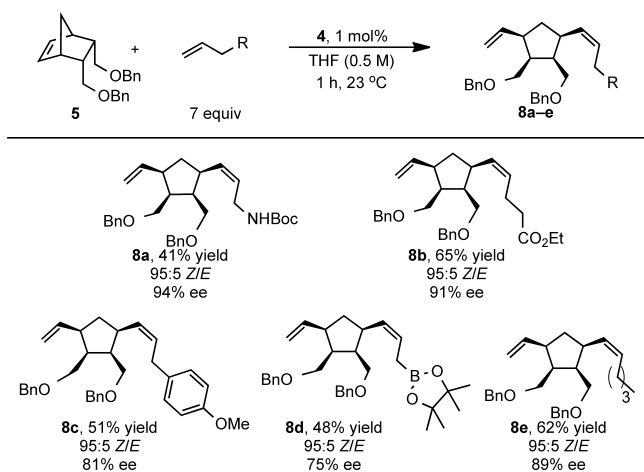
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equiv of terminal olefin, a catalyst loading of 1 mol % at 23 °C and a concentration of 0.5 M in tetrahydrofuran (THF) afforded the highest yield and selectivity. Ethereal solvents were optimal, with the catalyst solubility improved in THF over diethyl ether.

To demonstrate the scope of Z-selective catalyst **4**, a variety of terminal olefins bearing diverse functionality were employed in order to determine their effect on the efficiency and enantioselectivity of the reaction. As illustrated in Table 1,

**Table 1. AROCM with Different Terminal Olefin Partners<sup>a</sup>**



<sup>a</sup>Yields correspond to isolated products; Z/E ratios were determined by 500 MHz <sup>1</sup>H NMR analysis of the crude reaction mixtures; the ee's of the pure products were measured with chiral supercritical fluid chromatography (SFC).

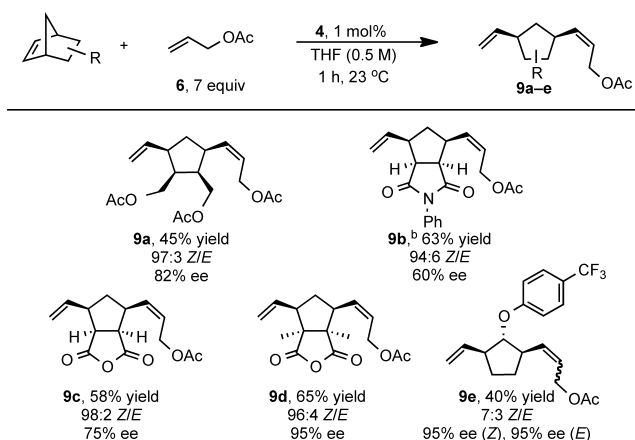
replacing allyl acetate with *N*-Boc-allylamine provided amine-containing product **8a** with equally high enantioselectivity (94% ee). Utilizing an olefin bearing a remote ester did not impact the Z selectivity and afforded **8b** with 91% ee.

Bulkier allylic substituents such as *p*-methoxyphenyl and pinacol boronic ester gave products **8c** and **8d** with moderate enantioselectivity (81 and 75% ee, respectively). A simple  $\alpha$ -olefin such as 1-hexene also gave good yield, Z selectivity, and enantioselectivity (**8e**, 89% ee), demonstrating that allylic functionality is not required to confer a selective reaction. The examples in Table 1 suggest that catalyst **4** is capable of producing a range of AROCM products.<sup>14</sup>

The norbornene component was then altered to investigate its impact on the Z selectivity and enantioselectivity. As a basis for comparison, the substrates were treated with 7 equiv of allyl acetate under the optimized catalytic conditions. Norbornenes bearing coordinating functionality such as acetate (to form **9a**) and *N*-phenylsuccinimide (to form **9b**) resulted in reduced yield and slower reaction, respectively. The dimethyl-substituted anhydride afforded a 65% yield of **9d**, which contains two vicinal all-carbon quaternary stereocenters, demonstrating the power of AROCM to afford otherwise synthetically challenging products with high ee (95%). Aryl ether **9e** was produced with 95% ee, although interestingly as a 7:3 Z/E mixture. The results in Table 2 support the observation that substrates bearing 2,3-endo substitution react with high Z selectivity, while substrates lacking this substitution pattern show reduced diastereoselectivity.

The fact that (Z)-**9e** and (E)-**9e** were formed with identical enantioenrichment has important mechanistic implications and

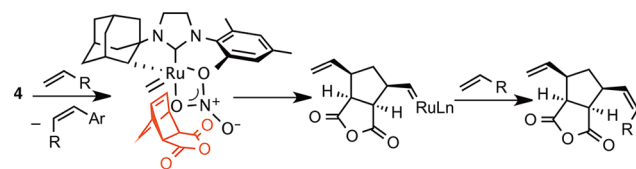
**Table 2. Influence of the Norbornene Reactant<sup>a</sup>**



<sup>a</sup>Yields correspond to isolated products; Z/E ratios were determined by GC; the ee's of the pure products were measured with chiral SFC. <sup>b</sup>Conducted at a catalyst loading of 3 mol % for 5 h.

offers indirect evidence of the active catalytic species. The result suggests that the enantiodetermining step most likely precedes the olefin-geometry-determining step.<sup>15</sup> This conclusion requires the initial enantiodetermining ring-opening event to occur with a ruthenium methylidene (Scheme 2). Subsequent cross-metathesis of the ring-opened product bearing a ruthenium alkylidene with 1 equiv of terminal olefin would then afford the observed product.

**Scheme 2. Proposed Model of Enantioselectivity**



On the basis of this indirect mechanistic evidence and the absolute configuration of the isolated product, we propose that the methylidene shown in Scheme 2 initially reacts with the norbornene component in an enantioselective ring-opening event. It is hypothesized that the enantioselectivity is governed by the approach of the methylidene to the less-hindered exo face, while the mesityl "cap" forces the bulk of the norbornene component to be oriented away from the NHC ligand.<sup>16</sup> The proposed methylidene is most likely produced by initial cross-metathesis of **4** with a molecule of terminal olefin, resulting in epimerization at the ruthenium center. Studies to provide a better understanding of the mechanism and origin of the enantioselectivity in the AROCM catalyzed by complex **4** are currently underway.

In summary, we have developed an enantioenriched ruthenium metathesis catalyst capable of highly Z-selective and enantioselective ROCM. An NHC ligand that chelates through a Ru–C bond is key to the design of the catalyst, which features a stereogenic Ru atom. The reaction is amenable to modification of both the terminal olefin and norbornene components, which significantly broadens the scope of this methodology.

## ■ ASSOCIATED CONTENT

### ■ Supporting Information

Experimental procedures, spectral data for the products, and crystallographic information files. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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### Notes

The authors declare no competing financial interest.

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(13) Absolute configurations were assigned by analogy to that of **9c**, which was determined by X-ray crystallography (see the Supporting Information for details).

(14) Attempts to employ a heteroatom-substituted olefin (butyl vinyl ether) resulted in no ROCM product, presumably because of catalyst deactivation.

(15) This assumes that secondary metathesis processes proceed at a negligible rate relative to the productive (ROCM) reaction. Measurements of the formation of **9e** (see the Supporting Information) showed that the Z/E ratio was constant during the course of the reaction and for several hours after complete conversion.

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