Ruthenium-Catalyzed O- to S-Alkyl Migration: A Pseudoreversible Barton-McCombie Pathway

William Mahy,[a] Pawel Plucinski,[b] Jesús Jover[c] and Christopher G. Frost*[a]

Abstract: A practical ruthenium-catalyzed O- to S-alkyl migration affords structurally diverse thiocarbamates in excellent yields. Our studies suggest this catalytic transformation proceeds through a pseudoreversible radical pathway drawing mechanistic parallels to the classic Barton-McCombie reaction.

The prevalence of sulfur-containing functionality in chemical biology, pharmaceutical drugs, agrochemicals and material science dictates the development and application of new synthetic methods in organosulfur chemistry is of fundamental importance. Historically, O-thiocarbamates have proved valuable as directing groups for the chemoselective reduction of alcohol functional groups to the corresponding alkane (Barton-McCombie Reaction)[1] and also in the generation of Ar–S compounds from phenols via an OMe to SMe migration (Newman-Kwart rearrangement).[2] The emergence of efficient metal-mediated techniques for the construction and transformation of carbon-sulfur bonds provide valuable tools for contemporary organic synthesis.[3] In this context, Lloyd-Jones et al have reported an elegant palladium-catalyzed variant of the Newman-Kwart rearrangement facilitating rearrangement at lower temperatures via an oxidative addition-tautomerisation-reductive elimination sequence.[4] Herein, we report a selective ruthenium-catalyzed Oalkylation to Salkylation migration that occurs when the O-thiocarbamate functionality is constrained in a readily-available ring structure (Scheme 1).[5] Our studies suggest this catalytic transformation proceeds through a pseudoreversible radical pathway drawing mechanistic parallels to the classic Barton-McCombie reaction.[6]

At the onset of our investigation, we explored the O- to S-alkyl migration of N-phenylthiazolidine-2-thione (1a) under traditional Barton-McCombie deoxygenation conditions using a variety of organic-based radical initiators.[7] The use of a number of transition metal catalysts (10 mol% equivalence of metal) known for their catalytic activity in single electron transfer processes were also investigated.[8] [RuCl₃(p-cymene)]₂ showed exceptional reactivity and selectivity towards O- to S-alkyl migration, giving rise to 82% 2a after 1 hour, where no observable desulfonated N-phenyl oxazolidinone from the competitive radical pathway was observed. Given ligation of ruthenium species is well known to improve both physical and chemical properties in ruthenium mediated transformations we tested a range of mono- and bidentate ligands in combination with 1 mol% Ru dimer in toluene at 100 °C.[9] Monodentate biaryl phosphines proved most effective, where SPhos gave the highest improved reactivity after 3 hours, giving rise to 2a in 81% conversion. The effect of concentration was then explored and higher concentrations were found to be by far the most effective, allowing quantitative conversion of 1a at 100 °C in 3 h with just 1 mol% Ru dimer. In the absence of catalyst there was no detectable rearrangement.

<table>
<thead>
<tr>
<th>Entry</th>
<th>Cat. (mol%)</th>
<th>Solvent</th>
<th>Time (h)</th>
<th>Conv. (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-</td>
<td>PhMe</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>DTBP (10)</td>
<td>PhMe</td>
<td>3</td>
<td>6 (8)</td>
</tr>
<tr>
<td>3</td>
<td>Pd(OAc)₂ (10)</td>
<td>PhMe</td>
<td>1</td>
<td>7 (11)</td>
</tr>
<tr>
<td>4</td>
<td>NiBr₂(PPh₃)₂ (10)</td>
<td>PhMe</td>
<td>1</td>
<td>5 (4)</td>
</tr>
<tr>
<td>5</td>
<td>[RuCl₃(p-cymene)]₂ (5)</td>
<td>PhMe</td>
<td>1</td>
<td>82 (10)</td>
</tr>
<tr>
<td>6</td>
<td>[RuCl₃(p-cymene)]₂ (1)</td>
<td>THF</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>7</td>
<td>[RuCl₃(p-cymene)]₂ (1)</td>
<td>MeCN</td>
<td>3</td>
<td>6</td>
</tr>
<tr>
<td>8</td>
<td>[RuCl₃(p-cymene)]₂ (1)</td>
<td>PhMe</td>
<td>3</td>
<td>17</td>
</tr>
<tr>
<td>9</td>
<td>[RuCl₃(p-cymene)]₂ (1)</td>
<td>PhMe</td>
<td>3</td>
<td>79</td>
</tr>
<tr>
<td>10</td>
<td>[RuCl₃(p-cymene)]₂ (1)</td>
<td>PhMe</td>
<td>3</td>
<td>100</td>
</tr>
</tbody>
</table>

Scheme 2 Selected screening results. *Reaction conditions 1a (0.25 mmol), Catalyst, Toluene (2.5 mL), 100 °C, air. †Yields of isolated products are given. ‡Conversion of 1b was determined by 'H NMR. Additional 1.0 eq Bu₂SnH used. §Conversion to N-phenyl oxazolidinone. ‖Reactions performed at 70 °C. ‡‡2 mol% SPhos. §§Toluene (1.25 mL). Full screening results reported in the supporting information.

[a] W. Mahy, Prof. Dr. C. G. Frost
Department of Chemistry
University of Bath, Bath, BA2 7AY (UK)
E-mail: c.g.frost@bath.ac.uk
[b] Dr. P. Plucinski
Department of Chemical Engineering
University of Bath, Bath, BA2 7AY (UK)
[c] Dr. J. Jover
Departament de Química Inorgànica
Universitat de Barcelona
C/Marti i Franquès 1-11, 08028
Barcelona (Spain)

Supporting information for this article is available on the WWW under http://dx.doi.org/10.1002/anie.2014XXXXX.
With the optimized reaction conditions in hand we turned our attention to the scope and limitations of this ruthenium promoted rearrangement. To our delight a wide range of N-aryl oxazolidine-2-thiones\(^\text{[10]}\) were tolerated under the reaction conditions, affording the corresponding N-aryl thiazolidin-2-ones in excellent yields. (Scheme 3)

Notably, the reaction conditions are tolerant of a number of valuable and reactive functional groups, including Br, CN, CF\(_3\), OMe, and F (products 2e-h, 2l), in the case of the electron rich and strongly electron poor aromatics showed decreased rates with respect to the unsubstituted aromatic, this effect was most pronounced with electron poor systems, where an increase in catalytic loading to 2 \text{mol\%} of Ru dimer was required to afford improved yields (products 2f, 2g and 2m). Enantiomerically pure 4-substituted oxazolidine-2-thiones also showed excellent reactivity under the reaction conditions, affording the corresponding enantiopure 4-substituted thiazolidin-2-ones with complete enantiomeric retention (2n-s).

5-substituted oxazolidine-2-thiones were significantly less reactive, mono-substitution at this position required elevated temperatures (130°C), increased catalytic loading (5 \text{mol\%}) and extended reaction times (24 h) to afford modest conversion (2t-v), a slight erosion of the enantiom purity (91% ee) was also observed. Geminal substituents showed an even further pronounced reduction in reactivity (2w-x). Phenol derivatives (2y), more closely mimicking substrates used in Newman-Kwart rearrangements showed no conversion even at more forcing conditions.\(^\text{[2]}\)

![Scheme 3](image)

\[ \text{Scheme 3. Synthesis of N-aryl thiazolidin-2-ones from a variety of N-aryl oxazolidine-2-thiones. Reaction conditions: N-aryl oxazolidine-2-thiones (0.25 mmol), [RuCl}_2(p-cymene)]_2 (1 \text{mol\%}), \text{SPhos (2 mol\%)}, \text{Toluene (1.25 mL), 100 °C, 3 h, air. Yields of isolated products are given.} \]

5-substituted oxazolidine-2-thiones were significantly less reactive, mono-substitution at this position required elevated temperatures (130°C), increased catalytic loading (5 \text{mol\%}) and extended reaction times (24 h) to afford modest conversion (2t-v), a slight erosion of the enantiom purity (91% ee) was also observed. Geminal substituents showed an even further pronounced reduction in reactivity (2w-x). Phenol derivatives (2y), more closely mimicking substrates used in Newman-Kwart rearrangements showed no conversion even at more forcing conditions.\(^\text{[2]}\)

![Scheme 4](image)

\[ \text{Scheme 4. Synthesis of N-alkyl thiazolidin-2-ones from a variety of N-alkyl oxazolidine-2-thiones. Reaction conditions: N-alkyl oxazolidine-2-thiones (0.5 mmol), [RuCl}_2(p-cymene)]_2 (1 \text{mol\%}), \text{SPhos (2 mol\%)}, \text{Toluene (2.5 mL), 100 °C, 4 h, air. Yields of isolated products are given.} \]

Subsequently, we turned our attention to the rearrangement of N-alkyl substituted oxazolidine-2-thiones. In all cases the corresponding N-alkyl thiazolidin-2-ones was formed in good to excellent yields. However, the reaction using an anthracene derivative gave only moderate conversion, which could be ascribed to the poor solubility of the substrate in toluene.

We next examined the rearrangement methodology on a multi-gram scale in order to probe the scalability of the process. N-phenyl oxazolidine-2-thione was subjected to the ruthenium catalyzed O- to S-alkyl migration with 1 \text{mol\%} [RuCl_2(p-cymene)]_2, under using the standard conditions (Scheme 5).
Isolation of the target thiazolidin-2-one was achieved in a similarly high yield of 96% following column chromatography.

In order to determine the mechanism of the \([\text{RuCl}_2(p\text{-cymene})]_2\) catalyzed reaction a number of kinetic and control experiments were performed. Cross-over experiments showed no formation of cross-over products when reacted under the optimized conditions. Reaction rates displayed a first-order dependency in both [1a] and [Ru] which eliminates a possible polymerization-depolymerisation process reported previously in the thermally mediated rearrangement of N-phenyloxazolidine-2-thiones.\(^{[11]}\)

The isomerization of cyclic thiocarbonate esters in the presence of potassium iodide is known to selectively generate primary monothiolcarbonates through an ionic pathway.\(^{[13]}\) In order to investigate the reactivity of N-phenyloxazolidine-2-thiones under ionic rearrangement conditions, 1a in the presence of catalytic potassium iodide (50 mol%) was heated and monitored over time (Scheme 6). Analysis of the crude reaction mixtures by \(^1\)H NMR showed that even after extended reaction times no rearrangement of the oxazolidine-2-thione occurred. Therefore it was concluded that it is unlikely that the reaction is proceeding through a nucleophilic ring opening / recombination process.

It has been reported that the treatment of cyclic thiocarbonates under Barton-McCombie promoted conditions results in the formation of the O- to S-rearrangement product when catalytic amounts of the radical promoter is used.\(^{[13]}\) Tsuda et al. found that cyclic thiocarbonates derived from glycosides gave a distribution of rearrangement, deoxy and o xo products when reacted under catalytic radially promoted conditions. Whilst only organic-based radical initiators were investigated, the choice of radical initiator and promoter were shown to have a significant effect on reactivity as well as selectivity. In contrast to ionic based mechanisms, both secondary and primary rearrangement products were observed in addition to the o xo derivative produced by the action of atmospheric oxygen and an intermediate radical.

In order to probe consistencies with the ruthenium catalyzed O- to S-alkyl migration of oxazolidine-2-thiones and this reported radical mechanism, \([\text{RuCl}_2(p\text{-cymene})]_2\) and cyclic thiocarbonate 5a were reacted using the standard conditions (Scheme 7). As with previous reported works, a distribution of products were observed, including the radical desulfurization product (6c, 4%). Interestingly both secondary and primary rearrangement products (6a and 6b) were observed in a 3:1 ratio, well within the ratios observed for classical radical promoted rearrangements. The observed consistencies would strongly suggest that the presented ruthenium catalyzed system is capable of proceeding via a similar radical promoted reaction pathway.\(^{[14]}\)

Analysis of the ruthenium catalyzed O- to S-alkyl migration by DFT (See SI for details) supports the proposed reaction proceeding through radical adducts (Scheme 8).

Thus, the formation of the starting \([\text{RuCl}_2(p\text{-cymene})][\text{SPhos}]\) complex (S0) is followed by a thermoneutral loss of the p-cymene ligand, generating the catalytic species S1. The reaction proceeds by the coordination of the N-aryl oxazolidine-2-thione (1) through the sulfur atom to give rise to the complex S2. The reaction interchanges then from the singlet to the triplet energy surface through the Minimum Energy Crossing Point MECP_S-T, this transformation requires approximately 10 kcal mol\(^{-1}\). Subsequently, one electron is transferred from the metal to the substrate (SET1), automatically provoking the cleavage of the C–O bond and generating the Ru(III) diradical species T1. The single electron transfer process requires 12 kcal mol\(^{-1}\) but still remains at a quite reasonable height.

Once T1 is obtained the pendant radical rotates through the corresponding transition state (Rot_TS), which is less than 7 kcal mol\(^{-1}\) higher than the previous intermediate, to form the diradical complex T2. The second electron transfer process (SET2) generates the C–S bond, the reaction then returns to the singlet energy surface through the corresponding Minimum Energy Crossing Point MECP_T-S, giving rise to the Ru(II) intermediate S3. Finally, the product is liberated into the reaction mixture and the starting catalytic species is regenerated. The
computed Gibbs free energy of the catalytic cycle is -14.4 kcal mol⁻¹, indicating that the whole process is thermodynamically favoured. The computed overall barrier for the reaction, obtained as the free energy difference between S₂ and SET₁, is approximately 23 kcal mol⁻¹, which is highly plausible for a reaction occurring at 100 °C.

![Scheme 8. Proposed model for ruthenium catalyzed O- to S-alkyl migration. Numbers are DFT-derived free energies (in kcal mol⁻¹) at 100 °C.](image)

In addition, a classical, non-radical, oxidative addition/reductive elimination catalytic cycle through ruthenium C-O insertion has been also computed (See SI for details). DFT calculations showed the energy requirements for this pathway are much higher than those obtained for the radical mechanism; in this case, the massive barrier obtained for the concerted oxidative addition step (+52.2 kcal mol⁻¹) would completely shut down the reaction. In addition, the rotational transition state barrier is also very high (+42.7 kcal mol⁻¹ from the lowest species), likely due to the charge separation produced in the ligand replacement process on ruthenium.

In summary, we have developed a robust and efficient ruthenium catalyzed O- to S-alkyl migration process for the practical preparation of N-substituted thiazolidine-2-thiones from readily accessible N-substituted oxazolidine-2-thiones. Further studies are underway to explore new applications of catalytic O- to S-alkyl migrations in organic synthesis. Initial experimental and computational investigations into the mechanism of this transformation suggest a pseudoreversible Barton-McCombie-type pathway is plausible.

**Acknowledgements**

We are grateful to the University of Bath, EPSRC DTC in Sustainable Chemical Technologies for funding (W. M.). We are also grateful for the valuable assistance of Anneke Lubben (Mass Spectrometry) and John Lowe (NMR) and Dr. Dave Carbery and Christina Gulyace (Chiral HPLC, University of Bath).

**Keywords:** Catalysis • Barton-McCombie • O-thiocarbamate • Alkyl migration • Pseudoreversible reaction


A radical step in a new direction: A practical ruthenium-catalyzed O- to S-alkyl migration affords structurally diverse thiooxazolidinones in excellent yields. Experimental and computational studies suggest a pseudoreversible radical pathway drawing mechanistic parallels to the classic Barton-McCombie reaction.