Ligand-Promoted Solvent-Dependent Ionization and Conformational Equilibria of
Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] (tim = 1-methylthioimidazolyl). Crystal Structures of
Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] and
\{Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]\}(PF$_6$)

Rosalice M. Silva
Marquette University

Brendan J. Liddle
Marquette University

Sergey V. Lindeman
Marquette University, sergey.lindeman@marquette.edu

Mark D. Smith
University of South Carolina - Columbia

James R. Gardinier
Marquette University, james.gardinier@marquette.edu
Ligand-Promoted Solvent-Dependent Ionization and Conformational Equilibria of Re(CO)₃Br[CH₂(S-tim)₂] (tim = 1-methylthioimidazolyl). Crystal Structures of Re(CO)₃Br[CH₂(S-tim)₂] and {Re(CO)₃(CH₃CN)[CH₂(S-tim)₂]}(PF₆)

Rosalice M. Silva
Department of Chemistry, Marquette University, Milwaukee, Wisconsin

Brendan J. Liddle
Department of Chemistry, Marquette University, Milwaukee, Wisconsin

Sergey J. Lindeman
Department of Chemistry, Marquette University, Milwaukee, Wisconsin

Mark D. Smith
Synopsis

The synthesis, structure, and solution chemistry of two tricarbonylrhenium(I) complexes of bis(1-methylthioimidazolyl)methane have been studied. The bromotricarbonylrhenium(I) complex was found by NMR and voltammetry to partially ionize on dissolution in CH$_3$CN; this behavior is not observed in acetone or CH$_2$Cl$_2$. The stereochemical nonrigidity of the complexes involving conformational changes in the eight-member chelate rings was also studied.

Abstract

The compounds Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] (1) and {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}(PF$_6$) (2), where tim is 1-methylthioimidazolyl, were prepared in high yields and characterized both in the solid state and in solution. The solid-state structures show that the ligand acts in a chelating binding mode where the eight-member chelate ring adopts twist-boat conformations in both compounds. A comparison of both solid-state IR data for CO stretching frequencies and the solution-phase voltammetric measurements for the Re$^{1+}/2+$ couples between 1, 2, and related N,N-chelates of the rhenium tricarbonyl moiety indicate that the CH$_2$(S-tim)$_2$ ligand is a stronger donor than even the ubiquitous dipyridyl ligands. A combination of NMR spectroscopic studies and voltammetric studies revealed that compound 1 undergoes spontaneous ionization to form {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}(Br$^-$) in acetonitrile. Ionization does not occur in solvents such as CH$_2$Cl$_2$ or acetone that are less polar and Lewis basic (less coordinating). The equilibrium constant at 293 K for the ionization of 1 in CH$_3$CN is 4.3 × 10$^{-3}$. The eight-member chelate rings in each 1 and 2 were found to be conformationally flexible in all solvents, and boat-chair conformers could be identified. Variable-temperature NMR spectroscopic studies were used to elucidate the various kinetic and thermodynamic parameters associated with the energetically accessible twist-boat to twist-boat and twist-boat to boat-chair interconversions.

Introduction

There is a great deal of interest in the coordination chemistry of the tricarbonylrhenium(I) moiety because of its applications in both materials and biomedical research. One of the attractive features of tricarbonylrhenium complexes in such applications is the generally inert
nature of the metal–ligand bonds that usually gives rise to reliable, simple synthetic protocols and straightforward characterizations. We recently reported\(^3\) on the synthesis and electrochemical properties of CH\(_2\)(S-tim)\(_2\) (tim = 1-methylthioimidazolyl), a compound that contains electroactive, heterocyclic thioimidazoline ring systems but for which little is known about its coordination chemistry.\(^4,5\) The free ligand exhibits two one-electron oxidations (as determined by a combination of voltammetry and redox titrations), but the voltammograms were reminiscent of ECE-type behavior and, unfortunately, the nature of the oxidized products remains unclear. We therefore embarked on studies in the coordination chemistry of this ligand with the intention of finding systems that will allow for the unequivocal elucidation of the unusual electrochemical behavior of the ligand. The tricarbonylrhenium(I) complex appeared well-suited for such a study because of its anticipated inert nature and the metal’s well-known electrochemical behavior. However, during the course of this study we discovered that the rhenium complexes of this ligand displayed unexpected reactivity that may prove useful to others in the future design of ligands for the tricarbonylrhenium moiety.

**Experimental Section**

Solvents were dried and distilled prior to use except where indicated. The syntheses of the rhenium complexes were carried out under an argon atmosphere using standard Schlenk techniques. Literature procedures were used to prepare Re(CO)\(_5\)Br,\(^6\) Re(CO)\(_3\)Br(X\(_2\)-bpy) (X = H, Me),\(^7\) CH\(_2\)(S-tim)\(_2\),\(^3\) and {Ag\(_2\)[CH\(_2\)(S-tim)\(_2\)]\(_2\)}(PF\(_6\))\(_2\).\(^5\) Elemental analysis was performed by Midwest Microlab Inc., Indianapolis, IN. Melting point determinations were made on samples contained in glass capillaries using an Electrothermal 9100 apparatus and are uncorrected. Infrared spectra were recorded on a Nicolet Magna-IR 560 spectrometer. \(^1\)H and \(^13\)C NMR spectra were recorded on a Varian 300 MHz spectrometer. Chemical shifts were referenced to solvent resonances at \(\delta H = 1.94\) and \(\delta C = 118.9\) for CD\(_3\)CN. Absorption measurements were recorded on an Agilent 8453 spectrometer. Electrochemical measurements were collected with a BAS CV-50V instrument for ca. 0.2 mM CH\(_3\)CN solutions of the complexes, with 0.25 M NBu\(_4\)PF\(_6\) as the supporting electrolyte, in a three-electrode cell composed of a Ag/AgCl electrode, a platinum working electrode, and a glassy carbon counter electrode. Mass spectrometric measurements recorded in ESI(+) mode were obtained on a Micromass Q-Tof spectrometer.

**Re(CO)\(_3\)Br[CH\(_2\)(S-tim)\(_2\)] (1).** A mixture of 0.340 g (0.836 mmol) of Re(CO)\(_5\)Br and 0.201 g (1.30 mmol) of CH\(_2\)(S-tim)\(_2\) in 20 mL of toluene was heated at reflux 12 h. After the mixture was cooled to room temperature, the precipitate was collected by filtration washed with two 5 mL portions of Et\(_2\)O and vacuum dried to give 0.435 g (99% yield) of analytically pure 1 as a
colorless powder. mp: 233 °C (dec) gray-brown solid. Anal. Calcd (obsd): C, 24.41 (24.48); H, 2.05 (1.93); N, 9.49 (9.26). IR (CH3CN, vCO, cm⁻¹): 2018(s), 1902, 1887. IR (KBr, vCO, cm⁻¹): 2016, 1902, 1871. ¹H NMR (CD3CN, 293 K): δH [twist boat, 86% of signals, the following integrations are relative for only this species, see Figure 4 for labeling] 7.72 (br s, 1H, tim-Ha), 7.66 (br s, 1H, tim-Ha), 7.39 (br s, 1H, tim-Hb), 7.37 (br s, 1H, tim-Hb), 3.88 (AB multiplet, JAB = 13.2 Hz, Δν = 6.0 Hz, 2H, CH2), 3.81 (br, s, 6H, NCH3); [boat-chair, 6% of signal, (the following integrations are relative to this species, see Figure 4 for labeling)] 8.48 (d, J = 1.6 Hz, 2H, tim-Ha), 7.11 (d, J = 1.6 Hz, 2H, tim-Hb), 4.29 (AB multiplet, JAB = 14.8 Hz, Δν = 47.8 Hz, 2H, CH2), 3.70 (br, s, 6H, NCH3); [cation, 8% of signals, see 2, below, for assignments] 7.58, 7.50, 7.48 (d, J = 1.7 Hz), 7.46, 7.25 (d, J = 1.7 Hz), 3.89, 3.83, 3.72; signals for one tim-Ha (twist-boat) and onset of methylene hydrogens (AB, multiplet, boat-chair) not observed (either in baseline or overlapping more intense resonances). ESI(+MS: 552 (13) [M − Br + CH3CN], 511 (100) [M − Br], 411 (21) [Re(CO)₃(CH₃CN)³(H₂O)⁺], 388 (18) [Re(CO)₃(CH₃CN)₂(H₂O)₂⁺], 370 (3) [Re(CO)₃(CH₃CN)₂(H₂O)⁺], 329 (50) [Re(CO)₃(CH₃CN)(H₂O)⁺], 270 (7) [Re(CO)₃⁺]. HRMS Calcd (obsd) for C₁₂H₁₂N₄O₃ReS₂[Re(CO)₃(CH₃CN)(H₂O)⁺]: 510.9908 (510.9913). Single crystals suitable for X-ray diffraction were grown by slow evaporation of an acetone solution.

{Re(CO)₃(CH₃CN)[CH₂(S-tim)²]}(PF₆) (2). Method A. A mixture of 0.105 g of ReCO₅Br (0.259 mmol) and 0.127 g (0.259 mmol) of {Ag₂[CH₂(S-tim)²]}(PF₆) in 20 mL of CH₃CN was heated at reflux for 4 h; then, the solution was filtered through Celite, and the solvent was removed to leave crude 2 as a glassy solid. After it was washed with two 10 mL portions of Et₂O and vacuum dried, the resulting material was recrystallized by layering Et₂O onto concentrated acetonitrile solutions and allowing the solvents to slowly diffuse over 2 days. The mother liquor was decanted and the residue was vacuum dried to produce 0.170 g (94%) of analytically pure 2 as colorless plates. Method B. A mixture of 0.236 g (0.400 mmol) of 1 and 0.101 g (0.400 mmol) of AgPF₆ in 15 mL of CH₃CN was heated at reflux for 12 h. After the mixture was filtered and the solvent was removed by vacuum distillation, the residue was washed with two 5 mL portions of Et₂O and vacuum dried under vacuum to leave 0.235 g (84%) of 2 as a colorless powder whose characterization data were identical to that of the crystals obtained above. mp: 220 °C (dec) gray-brown solid. Anal. Calcd (obsd): C, 24.03 (23.88); H, 2.59 (2.94); N, 10.01 (10.27). IR (KBr, vCO, cm⁻¹): 2034, 1922, 1899. ¹H NMR (CD₃CN, 293 K): δH [twist-boat, 90% of signal (integrations relative to this species), see Figure 4 for labeling] 7.66 (s, br, 1H, tim-Ha), 7.58 (s, br, 1H, tim-Ha), 7.50 (s, br, 1H, tim-Hb), 7.46 (s, br, 1H, tim-Hb), 3.89 (AB multiplet, JAB = 14.9 Hz, Δν = 1.0 Hz, 2H, CH₂), 3.83 (br, s, 6H, NCH₃), 2.27 (br, s, 3H, CCH₃); [boat-chair, 10% of signal (integrations relative to this species)] 7.48 (d, J = 1.7 Hz, 2H, tim-Ha), 7.25 (d, J = 1.7 Hz, 2H, tim-Hb), 4.30 (AB multiplet, JAB = 14.8 Hz, Δν = 40.2 Hz, 2H, CH₂).
3.72 (br, s, 6H, NCH₃), 2.14 (br, s, 3H, CCH₃). ¹³C NMR (CD₃CN, 233 K): δH 194.0, 194.7, 142.32, 142.27, 136.2, 127.5, 127.1, 123.6, 44.5, 35.60, 35.62, 4.3. ESI(+) MS: 552 (100) [M − Br + CH₃CN], 511 (40) [M − Br], 411 (14) [Re(CO)₃(CH₃CN)₃(H₂O)+], 270 (27) [Re(CO)₃⁺]. HRMS Calcd (obsd) for C₁₄H₁₅N₅O₃ReS₂ [M⁺]: 550.0146 (550.0139). Single crystals suitable for X-ray diffraction were grown by layering a concentrated acetonitrile solution of 2 with Et₂O and allowing solvents to slowly diffuse over 2 days.

**X-ray Crystallography.** X-ray intensity data from a colorless hexagonal rod of Re(CO)₃Br[CH₂(S-tim)₂] (1) were measured at 150(1) K on a Bruker SMART APEX CCD-based diffractometer, while those for a colorless plate of {Re(CO)₃(CH₃CN)[CH₂(S-tim)₂]}(PF₆) (2) were measured at 100(2) K on a Bruker SMART diffractometer equipped with an APEX2 CCD-based detector for the data collection of 2. Both experiments used monochromatic Mo Kα radiation, λ = 0.71073 Å. Raw data frame integration and Lp corrections were performed with SAINT+. Final unit cell parameters were determined by least-squares refinement of 6976 reflections from the data set of 1 and of the strongest 7600 reflections from the data set of 2, each with I > 5σ(I). The analysis of the data showed negligible crystal decay during collection. The data were corrected for absorption effects with SADABS. Direct methods structure solution, difference Fourier calculations, and full-matrix least-squares refinement against F² were performed with SHELXTL.

Re(CO)₃Br[CH₂(S-tim)₂] (1) crystallizes in the hexagonal system. Systematic absences in the intensity data indicated a space group from either of the enantiomorphous pairs P₆/P₆ or P₆22/P₆22. The space group P₆22 was eventually confirmed by the successful solution and refinement of the data and supported by the final absolute structure (Flack) parameter of 0.04(3). The asymmetric unit contains half a molecule located on a 2-fold rotational axis. Re1 and C1 are on the 2-fold. The molecule is disordered about the 2-fold axis such that Br1 and the carbonyl group C22–O22 are interchanged. Occupancies for the Br1 and C22–O22 groups were initially refined and yielded values near the expected 0.5. Subsequently, these occupancies were fixed at 0.5 for the final refinement cycles. Refinement in P₆ with the 2-fold axis removed resulted in the same Br/CO disorder with one complete molecule per asymmetric unit. Since the disorder is still observed in the lower space group P₆, it is not imposed by the 2-fold axes present in the higher space group P₆22, and therefore, P₆22 was retained as the better space group choice. Careful examination of the intensity data showed no evidence of a larger unit cell which might resolve the disorder. All non-hydrogen atoms were refined with anisotropic displacement parameters except C22 and
O22 (isotropic). Hydrogen atoms were placed in geometrically idealized positions and included as riding atoms.

\{{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]}\}(\text{PF}_6) \ (2) \text{ crystallizes in the orthorhombic system. Space group } Pbca \text{ was unequivocally assigned from systematic absences. The molecule is positioned in a general symmetry position and was refined in anisotropic approximation for all non-hydrogen atoms. The positions of the hydrogen atoms were determined from a series of difference Fourier syntheses and refined with isotropic approximation.}

Table 1. Crystallographic Data for Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}] (1) and \{{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]}\}(\text{PF}_6) (2)

<table>
<thead>
<tr>
<th></th>
<th>Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}]</th>
<th>{{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]}}(\text{PF}_6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>formula</td>
<td>C\textsubscript{12}H\textsubscript{12}BrN\textsubscript{4}O\textsubscript{3}ReS\textsubscript{2}</td>
<td>C\textsubscript{14}H\textsubscript{15}F\textsubscript{6}N\textsubscript{5}O\textsubscript{3}PReS\textsubscript{2}</td>
</tr>
<tr>
<td>fw</td>
<td>590.49</td>
<td>696.60</td>
</tr>
<tr>
<td>cryst syst</td>
<td>hexagonal</td>
<td>orthorhombic</td>
</tr>
<tr>
<td>space group</td>
<td>\textit{P}_\text{6}2\text{2}</td>
<td>\textit{Pbca}</td>
</tr>
<tr>
<td>(a)</td>
<td>8.6926(3) Å</td>
<td>13.635(2) Å</td>
</tr>
<tr>
<td>(b)</td>
<td>8.6926(3) Å</td>
<td>13.746(2) Å</td>
</tr>
<tr>
<td>(c)</td>
<td>40.452(2) Å</td>
<td>23.788(4) Å</td>
</tr>
<tr>
<td>(\alpha)</td>
<td>90°</td>
<td>90°</td>
</tr>
<tr>
<td>(\beta)</td>
<td>90°</td>
<td>90°</td>
</tr>
<tr>
<td>(\gamma)</td>
<td>120°</td>
<td>90°</td>
</tr>
<tr>
<td>(V)</td>
<td>2647.11(19) Å\textsuperscript{3}</td>
<td>4458.6(12) Å\textsuperscript{3}</td>
</tr>
<tr>
<td>(Z)</td>
<td>6</td>
<td>8</td>
</tr>
<tr>
<td>(T) (K)</td>
<td>150(1)</td>
<td>100(2)</td>
</tr>
<tr>
<td>(\rho)\textsubscript{calcld}</td>
<td>2.222 Mg m\textsuperscript{-3}</td>
<td>2.076 Mg m\textsuperscript{-3}</td>
</tr>
<tr>
<td>(\mu) (Mo K\textsubscript{\alpha})</td>
<td>9.403 mm\textsuperscript{-1}</td>
<td>5.786 mm\textsuperscript{-1}</td>
</tr>
<tr>
<td>(R) 1 [(I &gt; 2\sigma(I))]\textsuperscript{a} (all data)</td>
<td>0.0327 (0.0328)</td>
<td>0.0184 (0.0225)</td>
</tr>
<tr>
<td>wR\textsuperscript{2} (all data)</td>
<td>0.0777 (0.0777)</td>
<td>0.0440 (0.0453)</td>
</tr>
</tbody>
</table>

Results and Discussion

**Synthesis.** Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}] (1) and \{{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]}\}(\text{PF}_6) (2) were prepared in high yields (>80%) according to Scheme 1. Compound 2 (prepared by two different routes) was originally synthesized with the intention of comparing its spectroscopic and electrochemical properties with its charge-neutral counterpart and was later found to be critical for understanding the complex solution behavior of 1. For this purpose, the silver complex in the bottom left of Scheme 1, which has been described previously,\textsuperscript{5} was found to be a convenient reagent for a one-step high yielding (94%) synthesis of 2 from ReCO\textsubscript{9}Br. Both 1 and 2 are air- and light-stable colorless solids that are soluble in DMSO, DMF, and
acetonitrile, slightly soluble in acetone, THF, or CH₂Cl₂, and insoluble in Et₂O or hydrocarbon solvents.

**Scheme 1. Preparation of Rhenium Derivatives**

IR Spectra. The solid-state IR spectra of 1 and 2 each showed three CO stretching bands expected for fac-rhenium complexes with local C₅ symmetry. A summary of the CO stretching frequencies for derivatives with Re(CO)₃(Br or CH₃CN)(N,N-chelate) cores is given in Table 2. As expected, the electron-poor cationic derivative 2 is less capable of participating in back-bonding interactions and, thus, has higher carbonyl stretching frequencies than its charge-neutral counterpart 1. This is also observed when the carbonyl stretching frequencies of [Re(CO)₃(CH₃CN)(bpy)](B[3,5-(CF₃)₂C₆H₃]₄) were compared with the charge-neutral halo-derivatives Re(CO)₃(Cl, Br, or I)(bpy). Also, by comparison of the CO stretching frequencies of 1 with those of the ReCO₃Br complexes of other N,N-chelating ligands (for samples prepared as KBr pellets), the electron-donating ability of CH₂(S-tim)₂ (νbio = 2016, 1902, 1871 cm⁻¹) was found to be better than that of the dipyridyls (only two of the expected three bands are observed in many of the dipyridyl cases because of a fortuitous overlap of the lower energy stretches; the two observed bands typically occur near νbio = 2019, 1900 cm⁻¹) which are, in turn, better donors than the diazabutadienes (νbio = 2025, 1925, 1910 cm⁻¹), followed by the bis(pyrazolyl)methanes (νbio = ca. 2030, 1920, 1880 cm⁻¹).
Table 2. IR Spectroscopic Data for \textit{fac-Re(CO)}\textsubscript{3}(ligand) N,N-Chelate Complexes

<table>
<thead>
<tr>
<th>Compounds</th>
<th>(v_{CO}) (KBr, cm\textsuperscript{-1})</th>
<th>ref</th>
</tr>
</thead>
<tbody>
<tr>
<td>Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}] (1)</td>
<td>2016, 1902, 1871</td>
<td>this work</td>
</tr>
<tr>
<td>Re(CO)\textsubscript{3}Br(bpy)</td>
<td>2019, 1919, 1895</td>
<td>10a</td>
</tr>
<tr>
<td>Re(CO)\textsubscript{3}Br(4,4'-Me\textsubscript{2}bpy)</td>
<td>2019, 1889</td>
<td>10b</td>
</tr>
<tr>
<td>Re(CO)\textsubscript{3}Br(1,4-diazabutadiene)</td>
<td>2025, 1925, 1910</td>
<td>10c</td>
</tr>
<tr>
<td>Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(pz)\textsubscript{2}]</td>
<td>2029, 1925, 1876</td>
<td>11</td>
</tr>
<tr>
<td>{Re(CO)\textsubscript{3}(CH\textsubscript{3}CN)[CH\textsubscript{2}(S-tim)\textsubscript{2}]}(PF\textsubscript{6}) (2)</td>
<td>2034, 1922, 1899</td>
<td>this work</td>
</tr>
<tr>
<td>[Re(CO)\textsubscript{3}(CH\textsubscript{3}CN)(bpy)] (B[3,5-(CF\textsubscript{3})\textsubscript{2}C\textsubscript{6}H\textsubscript{3}]\textsubscript{4})</td>
<td>2044, 1943</td>
<td>7</td>
</tr>
</tbody>
</table>

**Solid-State Structures.** The structures of both Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}] (1) and {Re(CO)\textsubscript{3}(CH\textsubscript{3}CN)[CH\textsubscript{2}(S-tim)\textsubscript{2}]}(PF\textsubscript{6}) (2) are given in Figure 1. In each, the ReN\textsubscript{2}C\textsubscript{2}S\textsubscript{2}C eight-member chelate ring has a twist-boat geometry, as emphasized in the bottom of Figure 1. While the structure of 2 is well-behaved, that of Re(CO)\textsubscript{3}Br[CH\textsubscript{2}(S-tim)\textsubscript{2}] (1) is afflicted with a 50:50 disorder of the axial Br and CO groups that produces a pseudo-2-fold rotation axis along the C(1)–Re vector that can be visualized in Figure S1 of the Supporting Information. The common metal–ligand bond distances in 1 and 2 are statistically identical and all are in agreement with values found in related systems.\textsuperscript{10-13} The similarity in the metal–ligand bond lengths in 1 and 2 signifies that the structural impact resulting from any difference in the back-bonding capabilities of the rhenium centers in each complex is, in fact, negligible. Also of potential structural significance is the intramolecular S⋯S distance, which may be important in determining the electrochemical activity of the ligands because of the possible formation of intramolecular disulfide linkages (3.056(8) Å in complex 1 and 3.069(2) Å in 2). Both distances are shorter than the corresponding distances (av 3.11 Å) in the silver complexes [with bridging CH\textsubscript{2}(S-tim)\textsubscript{2} ligands], which were, in turn, shorter than twice the van der Waals radii of sulfur.\textsuperscript{5}
Figure 1 (left) Structure of one disordered component of Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] (1). Selected bond distances (Å) and angles (deg): Re(1)−C(21) = 1.902(8), Re(1)−C(22) = 1.92(2), Re(1)−N(11) = 2.210(7), Re(1)−Br(1) = 2.577(3), C(21)−O(21) = 1.153(10), C(22)−O(22) = 1.17(3); C(21)−Re(1)−Br(1) = 90.1(3), C(21)−Re(1)−Br(1) = 94.7(3), C(21)−Re(1)−C(21) = 86.6(5), C(21)−Re(1)−C(22) = 91.8(6), C(21)−Re(1)−C(22) = 86.3(6), C(22)−Re(1)−C(22) = 177.4(11), C(21)−Re(1)−N(11) = 174.5(3), C(22)−Re(1)−Br(1) = 177.9(6).

(right) Structure of {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}(PF$_6$) (2). Selected bond distances (Å) and angles (deg): Re(1)−C(10) = 1.907(2), Re(1)−C(11) = 1.9157(19), Re(1)−C(12) = 1.9172(19), Re(1)−N(5) = 2.1404(16), Re(1)−N(1) = 2.2119(14), Re(1)−N(3) = 2.2333(14), C(10)−O(10) = 1.158(2), C(11)−O(11) = 1.148(2), C(12)−O(12) = 1.150(2); C(10)−Re(1)−C(11) = 88.43(8), C(10)−Re(1)−C(12) = 88.19(8), C(11)−Re(1)−C(12) = 88.77(8), C(10)−Re(1)−N(5) = 94.91(7), C(11)−Re(1)−N(5) = 94.49(7), C(12)−Re(1)−N(5) = 175.56(7), C(10)−Re(1)−N(1) = 174.53(7), C(11)−Re(1)−N(1) = 86.13(7), C(12)−Re(1)−N(1) = 92.23(7), N(1)−Re(1)−N(5) = 84.98(6), C(10)−Re(1)−N(3) = 89.62(7), C(11)−Re(1)−N(3) = 176.57(7), C(12)−Re(1)−N(3) = 93.99(6), N(3)−Re(1)−N(5) = 82.87(6), N(1)−Re(1)−N(3) = 95.79(5). The views at the bottom emphasize the twist-boat conformation of the eight-member chelate ring.

Solution Phase. Further characterization data shows that these two rhenium compounds, 1 and 2, exhibit rich solution-phase chemistry. First, the ESI(+) mass spectrum of each compound in CH$_3$CN, which is thought to give one indication of the solution behavior of compounds, are nearly identical showing signals at m/z = 552, 511, and 270 (among others) that correspond to {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}$^+$, {Re(CO)$_3$(CH$_2$(S-tim)$_2$)}$^+$, and [Re(CO)$_3$]$^+$, respectively. That is, the molecular ion is observed in 2 but not in 1; the main fragmentation peak of 1 is from the unipositive complex, sans bromide. While the signal for the {Re(CO)$_3$(CH$_2$(S-tim)$_2$)}$^+$ ion is a result of the fragmentation process, both solution NMR and electrochemical studies indicate that the bromide dissociation to give {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}$^+$ occurs to a certain extent in CH$_3$CN under typical laboratory conditions (vide infra).
To facilitate the discussion of the NMR spectra of \( \text{Re(CO)}_3\text{Br[CH}_2\text{(S-tim)}_2 \) (1) and \( \{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2\text{(S-tim)}_2]\}\text{(PF}_6 \) (2), it will be useful to examine the expected NMR spectra of the possible isomers of these species in more detail. After possible interchanges of bromine (or acetonitrile in the case of 2) and the axial carbonyl groups on rhenium, as well as what is known for other semirigid eight-member ring systems, such as \( \text{cis,cis-1,4-cyclo-octadiene,} \)\(^{15}\) its dibenzo analogue,\(^{16}\) or its heteroatomic dibenzo derivatives \( \text{(I–III, respectively, Figure 2),} \)\(^{17}\) are considered, it can be shown that there are seven possible conformers of the \( \text{ReN}_2\text{C}_2\text{S}_2\text{C} \) ring in either 1 or 2 leading to a total of 10 isomers (including optical isomers), as shown for 1 in Figure 3. The relative stabilities of the various possible conformers were probed via the evaluation of the AM1 geometry-optimized structures calculated using the SPARTAN\(^{24}\) molecular modeling program. While such models may be of limited value, certain qualitative features concerning factors responsible for the relative stabilities of the conformers become apparent, thereby allowing us to make more confident, yet still tentative, assignments for the two species responsible for the complicated NMR spectrum of 1 and 2 in acetone or \( \text{CD}_2\text{Cl}_2 \) (vide infra). Gratifyingly, similar to the NMR spectral data, the calculations yield two (nearly isoergonic) low-energy conformers that can be distinguished from the remaining five conformers. These two low-energy conformers were the twist-boat (with \( \text{C}_1 \) symmetry) as found in the solid-state structure and the \( \text{C}_s \)-symmetric boat-chair with the CO oriented toward sulfurs. The remaining conformers could be differentiated in energy by consideration of the steric interactions between the sulfur atoms of the chelate ring and rhenium–bromide moiety: the more sulfur–bromine interactions, the higher in energy is the conformer. For instance, the \( \text{C}_s \)-symmetric boat-chair with bromine oriented toward the sulfurs is about 7 kcal/mol higher in energy than the other boat-chair. Also, because of considerable ring strain, the \( \text{C}_s \)-symmetric boat-boat conformers are about 25 kcal/mol higher in energy than the \( \text{C}_s \)-symmetric boat-chairs. Similar observations are obtained in the case where acetonitrile replaces bromine, as in 2; however, the differences in energy between conformers were smaller presumably because of the lower steric demand of an acetonitrile versus a bromide. Thus, we interpret the complicated NMR spectra of 1 and 2 (in acetone or \( \text{CD}_2\text{Cl}_2 \)) to result from a mixture of the twist-boat (with \( \text{C}_1 \) symmetry) and the \( \text{C}_s \)-symmetric boat-chair conformers with the CO oriented toward sulfurs.
Figure 2: Semirigid organic-based eight-member ring systems. For III, X and Y can be carbon, any reasonable heteroatom, or heteroatom-containing groups such as S, SO₂, NH, etc.

Since the experimental NMR spectra can be satisfactorily interpreted to be the result of the presence of twist-boat and boat-chair (with CO oriented toward the sulfurs) conformers, the remainder of the discussion will be limited to these two cases. From Figure 4, it is expected that the NMR spectrum of the C₁-symmetric twist-boat conformer of 1 should consist of eight resonances: four thioimidazolyl hydrogen resonances (Hₐ, Hₐ', Hₖ, Hₖ'), two AB “doublet” resonances for the methylene hydrogens (Hₐ, Hₖ), and two N-methyl resonances. The expected spectrum of the Cₛ-symmetric boat-chair conformer should exhibit five resonances: two doublet resonances for thioimidazolyl hydrogens, two AB “doublet” methylene resonances,
and a single $N$-methyl resonance. The corresponding spectra of the cation in 2 should have an additional resonance for the acetonitrile methyls.

![Figure 4 Labeling diagram used for NMR assignments. Important short intramolecular contact distances are also shown.](image)

The resonances for the thioimidazolyl and the methylene hydrogens of each conformer merit some further comment. First, the chemical shifts of the thioimidazolyl hydrogen resonances are anticipated to provide useful information regarding the local environment around rhenium. Changes in the environment around rhenium are expected to affect the chemical shifts of the hydrogen resonances in the ligand scaffold in a predictable way via dipole–dipole-type interactions, delineated by the Ramsey equation. That is, these interactions rapidly decrease with increasing distance ($E$ is proportional to $r^3$) giving the relative order of impact: proximal thioimidazolyls > distal thioimidazolyls > $N$-methyls ≥ CH$_2$'s. Thus, the downfield-most thioimidazolyl hydrogen resonances that correspond to those hydrogens closest to the metal center are expected to be most affected by changes in the environment surrounding rhenium. In the $C_1$-symmetric twist-boat conformer of 1, the proximal hydrogens are nearest in space to the equatorial carbonyls (Figure 4 left), but one hydrogen is oriented toward the axial bromide and the other hydrogen toward the axial carbonyl, thereby differentiating these two hydrogens. As was seen in related systems, weak CH···Br hydrogen bonding interactions tend to result in downfield shifts in hydrogen resonances. In the twist-boat conformer of 1, the H···Br distance is fairly long at 3.1 Å (from the X-ray structure), and because of the presumably weak nature of this CH···Br interaction, only a small splitting of proximal thioimidazolyl hydrogen resonances is expected. In the boat-chair conformer with the axial CO oriented toward the sulfurs, the thioimidazolyl hydrogens proximal to rhenium are equivalent but are now nearest to the axial bromide on rhenium (rather than the equatorial CO ligands as in the twist-boat). In this low-energy conformer, the thioimidazolyl hydrogens are now only ca. 2.5 Å from the
bromide (from the energy-minimized calculated structure); thus, a large downfield shift is expected for these resonances. Importantly, large chemical shift differences of proximal thioimidazolyl hydrogen resonances are expected in the spectrum of the boat-chair conformer (with CO oriented toward the sulfurs) of 1 compared to that of 2 because of the gross changes in the axial ligating sites in 1 versus 2, where the chemical shifts may be modified by weak \( \text{CH} \cdots \text{Br} \) and \( \text{CH} \cdots \pi(\text{CH}_3\text{CN}) \) interactions, respectively. By analogy, the chemical shifts of the proximal thioimidazolyl hydrogen resonances for the twist-boat conformers of 1 and 2 should have greater similarity than the corresponding resonances of the boat-chair conformers of these compounds because of the nearly identical \( \text{CH} \cdots \pi(\text{CO}_{\text{equatorial}}) \) interactions: the differences in the spectra would be caused by long-range \( \text{CH} \cdots \text{Br} \) and \( \text{CH} \cdots \pi(\text{CH}_3\text{CN}) \) interactions. As indicated by the solid-state structural studies, any differences as a result of the different back-bonding capabilities of the metal center or because of the trans effect modulating the strength of otherwise similar (thioimidazolyl)\( \text{CH} \cdots \pi(\text{CO}) \) interactions is expected to be minimal. Similarly, for the other possible (high-energy) boat-chair conformers of 1 and 2 with a carbonyl oriented between thioimidazolyl hydrogens, only minimal differences in the mean chemical shifts of thioimidazolyl resonances are expected.

The \(^1\text{H} \) NMR spectra of 1 and 2 in various solvents is more complicated than expected from the solid-state structures because of (i) the equilibrium ionization of \( \text{Re(CO)}_3\text{Br}[\text{CH}_2(\text{S-tim})_2] \) to give \( \{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]^+\}(\text{Br}^-) \), (ii) the presence of equilibrium mixtures of interconvertible conformers of the complexes that comes from the conformational flexibility of the eight-member \( \text{ReN}_2\text{C}_2\text{S}_2\text{C} \) chelate ring, or (iii) both. Thus, the \(^1\text{H} \) NMR spectra of 2 in CH\(_3\)CN contains 15 resonances, expected for a mixture of twist-boat and boat-chair isomers. However, the NMR spectrum of an analytically pure sample of 1 in CH\(_3\)CN has 28 resonances: 15 of these resonances match the chemical shifts of those in 2, while the remaining 13 resonances are for a mixture of twist-boat and boat-chair conformers of complex 1. For 1 in acetone or CH\(_2\)Cl\(_2\) (Supporting Information), the total number of resonances drops to thirteen, again, for a mixture of twist-boat and boat chair isomers of the intact complex. Thus, the ionization of \( \text{Re(CO)}_3\text{Br}[\text{CH}_2(\text{S-tim})_2] \) to give \( \{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]^+\}(\text{Br}^-) \) occurs in acetonitrile (and more polar solvents such as DMF and DMSO) but does not appear to occur to any appreciable extent in less polar (or possibly less nucleophilic) solvents. This is shown in Figure 5, which provides the downfield portion of the NMR spectrum (for thioimidazolyl resonances) of 1 (top) and of 2 (bottom) in acetonitrile at 293 K. The spectrum of analytically pure 1 in CD\(_3\)CN (top of Figure 5) clearly contains resonances in the baseline at ca. \( \delta = 7.6, 7.5, \) and 7.25 ppm that are for the \( \{\text{Re(CO)}_3(\text{CH}_3\text{CN})[\text{CH}_2(\text{S-tim})_2]^+ \) cation (bottom Figure 5, the nature of these resonances will
be described later). An NMR-scale reaction between 2 and NBu4Br in CD3CN gave a spectrum identical to that of an independently prepared mixture of 1 and NBu4PF6 in the same solvent, verifying the equilibrium. The apparent equilibrium constant for the ionization of pure 1 in CH3CN at 293 K is $4.3 \times 10^{-3}$ (assuming the density of the CH3CN in the solution is close to that for pure CH3CN at 293 K so that the equilibrium concentration of CH3CN is 19.1 M).

The broad nature of the resonances for the twist-boat conformer and the presence of small amounts of the boat-chair conformer compelled us to acquire variable-temperature NMR spectra to obtain thermodynamic and kinetic parameters associated with the variety of processes that occur in solutions of each compound. Some results of these studies are provided in Figure 6 and in Figures S4 and S5 in the Supporting Information. At 263 K and lower temperatures, the NMR spectra of 1 and 2 consist of sharp resonances indicative of slow exchange on the NMR time scale. Especially noteworthy in this temperature regime are the resonances for the thioimidazolyl hydrogens and those for the methylene moiety. First, the similarity in the mean chemical shifts of thioimidazolyl hydrogen resonances in the twist-boat conformers of 1 and 2 is indicative of the minimal structural impact of different metal–carbonyl back-bonding in each complex. By analogy, the very large difference in the chemical shifts of the proximal thioimidazolyl hydrogen resonances in the boat-chair conformers of 1 and 2 provides support that this conformer is indeed that with the carbonyl oriented toward sulfurs as from the earlier discussion. Also, on first inspection, the methylene resonance of the

Figure 5 Downfield portion of the 1H NMR spectrum of Re(CO)3Br[CH2(S-tim)2] (top) and {Re(CO)3(CH3CN)[CH2(S-tim)2]}(PF6) (bottom) in CD3CN at 293 K.

\[\delta, \text{ppm} \quad 8.5 \quad 8.0 \quad 7.5 \quad 7.0\]
twist-boat conformer of 2 at room temperature appears to be a singlet, but the variable-temperature studies confirm that this is actually an AB multiplet. The intensity of the outer lines of an AB spectrum, I, from theory can be calculated from $I = 1 - \frac{1}{2} \frac{J_{AB}}{\left(\frac{(\Delta \nu)^2 + J_{AB}^2}{2}\right)^{1/2}}$,\textsuperscript{18,19} where $J_{AB}$ is the geminal coupling constant and $\Delta \nu$ is the chemical shift difference between resonances. The chemical shift difference of the resonances is very small, on the order of 6 Hz for 1 and 1 Hz for 2, and the signals for the outer lines of the AB multiplet patterns are of very low intensity, almost indiscernible in the baseline.

![Diagram of chemical structures]

Figure 6 Identical portions of the $^1$H NMR spectrum of Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] (1, top) and {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}(PF$_6$) (2, bottom) in CD$_3$CN at various temperatures revealing the dynamic behavior of the {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]} cation in both 1 and 2. See Supporting Information for full spectra.

When solutions of either 1 or 2 are warmed to room temperature and above, the various resonances for the twist-boat conformer broaden and coalesce between 293 and 303 K
for 1 and between 303 and 308 K for 2. At higher temperatures, the resonances for the boat-chair conformer broaden into the baseline and are no longer observed above 323 K. These results are interpreted by consideration of the two independent processes that interconvert twist-boat isomers: a low-energy twist-boat to twist-boat conversion and a higher but energetically accessible twist-boat to boat-chair to twist-boat conversion, as shown in the bottom portion of Figure 3. It should be noted that, similar to that found in the organic ring systems described earlier, the symmetric boat-chair isomers are more conformationally rigid than the twist-boat conformers. An examination of the CPK models supports the plausibility of a dual-pathway dynamic process, as it is found that simple torsion about the sulfur–methylene carbon bonds is sufficient to interconvert between two mirror-image twist-boat conformations, whereas a ring-flipping, in addition to bond torsion, is necessary to obtain the boat-chair conformer. From the relative integrations of resonances, it was found that the twist-boat is favored over the boat-chair by about 16:1 and 9:1 for 1 and 2, respectively, at 293 K in all solvents tested (CD$_2$Cl$_2$ for 1; acetone and CH$_3$CN for 1 and 2). This equilibrium distribution corresponds to free-energy differences of about 1.7 and 1.2 kcal/mol between the two conformers of each respective compound, values close to those predicted by the molecular mechanics calculations. The measurement of $K_{eq}$ from the integrations of thioimidazolyl resonances at temperatures between 283 and 233 K (where reliable integrations could be obtained) allowed for the determination of $\Delta H = -5.0$ kJ/mol, $\Delta S = -6.6$ J/Kmol and $\Delta H = -2.9$ kJ.mol, $\Delta S = -8.0$ J/Kmol for the boat-chair ⇄ twist-boat equilibrium for 1 and 2, respectively, from the straight-line plots of $\ln K$ versus $(1/T)$. The coalescence temperature ($T_c$) of the thioimidazolyl resonances for the twist-boat to twist-boat conversion for each 1 and 2 was used to estimate statistically identical activation barriers of $E_a = 4.57 T_c[10.32 + \log(\sqrt{2}T_c/\pi\Delta\nu)] = 18.6 \pm 0.3$ kcal/mol for 1 and $E_a = 19.2 \pm 0.3$ kcal/mol for 2 (where $\Delta\nu$ is the chemical shift difference of the resonances in absence of exchange).$^{20}$ Similar calculations were used to provide crude estimations of $E_a = 22.0 \pm 0.5$ kcal/mol for the activation energy of the boat-chair to twist-boat conversion in 1 and 2. In these latter calculations, a greater degree of uncertainty is introduced because of the disparity in relative populations of the boat-chair and twist-boat conformers, since a coalescence temperature is not easily obtained. Thus, the temperatures just before the resonances for the boat-chair conformers broaden into the baseline (323 K for 1 and 333 K for 2) were used in the estimation. The calculated energies are of similar magnitude, but they are a little higher than observed for the ring-flipping mechanisms in cyclooctatetraene and in related eight-member organic ring systems.$^{15-17}$

**Electrochemical Studies.** The cyclic and square-wave voltammograms for compounds 1 and 2 are given in Figure 7. In support of the findings of the NMR spectroscopic
studies, the electrochemical data are also indicative that Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] undergoes ionization in acetonitrile to give {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}$^+$ (Br$^-$). The ionization of 1 in acetonitrile is most clearly evident from the small irreversible (anodic) oxidation wave at ca. +0.7 V versus Ag/AgCl (top of Figure 7) for the Br$^-$/Br$_2$ couple which is nearly identical to the potential found for pure NBu$_4$Br in CH$_3$CN measured under the same conditions. Moreover, there are corresponding waves of similar (weak) intensity for the {Re(CO)$_3$(CH$_3$CN)[CH$_2$(S-tim)$_2$]}$^+$ cation present in CH$_3$CN solutions of 1. Thus, the voltammogram for 1 in acetonitrile consists of four oxidation waves, two overlapping waves at +1.58 and +1.53 V (best detected by square-wave voltammetry, by varying scan rates because of the significantly different diffusion rates, or by both), a wave at +1.17 V, and one at +0.7 V, in addition to two reduction waves at −2.17 and −2.52 V. For comparison, the voltammogram of 2 consists of one reversible oxidation wave at +1.58 V and one irreversible reduction wave at −2.17 V. Thus, in the voltammogram of 1, only the three intense waves at +1.53, +1.17, and −2.52 V are from the redox processes involving the intact charge-neutral Re(CO)$_3$Br[CH$_2$(S-tim)$_2$] complex.

Figure 7 Cyclic and square-wave voltammograms of Re(CO)$_3$Br[N,N-CH$_2$(S-tim)$_2$] in CH$_3$CN (top) and {Re(CO)$_3$(CH$_3$CN)[N,N-CH$_2$(S-tim)$_2$]}(PF$_6$) (bottom) in CH$_3$CN (NBu$_4$PF$_6$) at 50 mV/s.

We can confidently assign the irreversible oxidation wave for 1 at +1.17 V to be from the Re$^I$/Re$^{II}$ couple by a comparison of this potential with those found in other well-known tricarbonyl Mungium(I) complexes of N,N-chelating ligands measured under similar conditions.$^{10-13}$ For instance, in the series of compounds [ReBr(CO)$_3$(pz)$_2$CHC$_5$H$_4$)$_2$Fe,$^{11}$ ReBr(CO)$_3$[CH$_2$(pz)$_2$],$^{11}$ReBr(CO)$_3$[CMe$_2$(pz)$_2$],$^{13}$ and ReBr(CO)$_3$[CH$_2$(3,5-Me$_2$pz)$_2$],$^{21}$ the rhenium oxidation waves were found to be +1.40, +1.37,
+1.35, and +1.29 V, respectively, versus Ag/AgCl measured under the same conditions as compound 1. This order shows that changing substituents on the bis(pyrazolyl)methane ligand system has a small but significant impact on the ReI/ReII couple: the more electron rich the ligand the easier it is to oxidize rhenium and the lower the oxidation potential. Similar comparisons can be made to gauge the relative electron-donating abilities of closely related ligand systems. Thus, in support of the findings of the IR spectroscopic data for CO stretches, the electrochemical data for compound 1 with an oxidation potential of +1.17 V shows that the N,N-chelating CH₂(S-tim)₂ ligand is a better electron-donor than the dipyridyl [Re(CO)₃Br(bpy), E₁/₂ = +1.29 V versus Ag/AgCl] or any of the bis(pyrazolyl)methane ligands (E₁/₂ = +1.4 to +1.3 V). In the cationic derivative, {Re(CO)₃(CH₃CN)[CH₂(S-tim)₂]}(PF₆), 2, the corresponding wave is shifted to more positive potentials (+1.58 V), as expected from Coulombic arguments. Interestingly, the metal-centered oxidation is now fully reversible in 2, whereas it is irreversible in 1.

The second most intense oxidation wave for intact Re(CO)₃Br[CH₂(S-tim)₂] (1) in CH₃CN at +1.53 V is assigned to a one-electron ligand oxidation. This assignment is made by comparison with the potentials found in the free ligand (+1.2 V) and its silver complexes (+1.5 V).³⁵ Complexation to the electron-withdrawing bromotricarbonylrhenium moiety makes the ligand oxidation more difficult by 0.3 V, similar to the effect caused by complexation to silver(I) cations. In the case of 2, the expected increased difficulty for the formation of a tricationic species (after the first metal oxidation of monocationic 2⁺ to give dicationic 2²⁺, followed by ligand oxidation to give 2³⁺) likely renders the first ligand oxidation so unfavorable as to be outside of the solvent potential window.

The nature of the reduction waves below −2.0 V versus Ag/AgCl for 1 and 2 is ambiguous as to whether these are metal- or ligand-based, however, it is clear from the right side of Figure 7 that the reduction wave for the cation in 2 is present in the voltammogram of pure 1 dissolved in CH₃CN. In terms of potential assignment of these waves, we favor ligand-centered reductions because the related purely organic dicationic derivative {CH₂(μ-tim)₂CH₂}}(PF₆)₂ showed an irreversible cathodic reduction wave at −1.2 V versus Ag/AgCl measured under similar conditions,³ therefore, a monocationic derivative is expected to have a reduction at more negative potential. As testament to the difficulty in this assignment, the voltammograms of Re(CO)₃(CH₃CN)₂Br²² gives an irreversible cathodic reduction wave at −2.41 V (in addition to a metal centered oxidation at +1.91 V versus Ag/AgCl in CH₃CN); however, as this potential approaches the solvent limit in our experiments, it is unclear as to whether this wave may be caused by reduction of the coordinated acetonitrile or reduction of
the Re+ center. It should be noted that, if the Re3+, Re2+, Re1+, and Re0 oxidation states for a given ligand system can be accessed, these redox couples (in nonaqueous environments) are usually each only separated by a approximately 1 V, much smaller than the potential separations in all of the above cases. Further experiments are underway to determine the nature of the reduction waves near −2.0 V in these and related systems. Regardless of their exact nature, the expected trend of increasing ease of reduction with increasing positive charge on going from the monocationic {Re(CO)3(CH3CN)[CH2(S-tim)2]}+ (2, −2.18 V) to the charge-neutral starting complex Re(CO)3Br[CH2(S-tim)2] (1, −2.52 V) is followed.

Conclusions
Two new tricarbonylrhenium complexes of the electroactive bis(1-methylthioimidazolyl)methane ligand Re(CO)3Br[CH2(S-tim)2] (1) and {Re(CO)3(CH3CN)[CH2(S-tim)2]}(PF6) (2) have been prepared. Although it was not possible to unequivocally establish the nature of all the redox processes in the rhenium complexes of this ligand, the multiple solid-state and solution characterization methods permitted further evaluation of the unique structural and electronic properties of the ligand, as well as the discovery of the unusual solution chemistry involving these rhenium complexes. From IR and electrochemical data, it is found that the bis(1-methylthioimidazolyl)methane ligand is more electron-rich than the dipyridyls or the bis(pyrazolyl)methanes, two common classes of N,N-chelating ligands. The solution NMR data indicate both 1 and 2 are stereochemically nonrigid in solution, as a result of the flexible eight-member chelate ring. Importantly, both NMR and electrochemical data indicate that 1 undergoes spontaneous ionization to {Re(CO)3(CH3CN)[CH2(S-tim)2]}(Br) in acetonitrile, where the equilibrium constant for ionization at 293 K was found by NMR to be 4.3 × 10⁻³. That this ionization does not occur in less polar solvents such as acetone implicates an associative mechanism for ionization, but further studies are needed to discern the mechanism of ionization. Studies are underway in our laboratory to determine the origin (whether electronic, steric, or both) and generality of the labilization of the axial ligands on these and other tricarbonylrhenium(I) N,N-chelate complexes.

Acknowledgment
We thank Bill Cotham and Mike Walla at the University of South Carolina for mass spectrometric measurements; J.R.G. also thanks Marquette University and the donors of the Petroleum Research Fund for support.
References


7 Hevia, E.; Perez, J.; Riera, V.; Miguel, D.; Kassel, S. Rheingold, A.


21 Gardinier, J. R. Unpublished observations.


A. Organometallics 1984, 3, 1515.