HOMOGENEOUS MODELS OF THIOPHENE HDS REACTIONS. SELECTIVITY IN THIOPHENE C-S CLEAVAGE AND THIOPHENE REACTIONS WITH DINUCLEAR METAL COMPLEXES.


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Received August 1, 1996

Abstract: The reactive 16 e\textsuperscript{-} metal fragment [(C\textsubscript{5}Me\textsubscript{5})Rh(PMe\textsubscript{3})] inserts into a wide variety of thiophene C-S bonds. The structures of the thiophene, benzothiophene, and dibenzothiophene insertion complexes have been determined. While the thiophene complex adopts a planar 6-membered ring structure the other metallacycles are bent, and all molecules possess localized diene structures. The mechanism of C-S cleavage was found to proceed by way of initial sulfur coordination. 2-Methylbenzothiophene gives a kinetic product resulting from cleavage of the sulfur-vinyl bond, but then rearranges to cleave the sulfur-aryl bond. A number of substituted dibenzothiophenes were examined, showing little electronic effect of substituents, but showing a large steric effect of substituents at the 4 and 6 positions. 4,6-Dimethyldibenzothiophene does not undergo cleavage, but instead forms an S-bound complex. Reactions of a cobalt analog, (C\textsubscript{5}Me\textsubscript{5})Co(C\textsubscript{2}H\textsubscript{4})\textsubscript{2} with thiophenes also lead to C-S cleaved products, and the use of a dinuclear iridium system produces a butadiene complex in which both C-S bonds have been cleaved.

Introduction

The hydrodesulfurization of petroleum is one of several steps in the hydrotreating of oil in which sulfur is removed from thiols and thiophenes as H\textsubscript{2}S.\textsuperscript{1} The process uses a supported molybdenum sulfide catalyst containing cobalt, and operates under pressures of 150-600 psi hydrogen at 300-450 °C. The sulfur content in oil of 1-5% is reduced to 0.1% in gasoline, and future sulfur limits may be reduced to as little as 0.003-0.04%.

Important studies performed by Kabe, Ishihara, and Tajima using a mixture of polyaromatic sulfur-containing compounds in light oil with a commercial HDS catalyst under typical operating conditions show the limitations of the current generation of catalysts.\textsuperscript{2} Figure 1 shows a GC trace of these sulfur containing compounds prior to treatment. Figure 2 shows how this original mixture of compounds is changed upon HDS treatment at temperatures of 350 - 390 °C. While all of the benzothiophenes are removed, it can be observed that specific dibenzothiophenes are difficult to desulfurize. These include 4-methyl dibenzothiophene, 4,6-dimethyl dibenzothiophene, and other polyalkylated dibenzothiophenes.

Work in our laboratory has focussed upon homogeneous models for the desulfurization of thiophenes.\textsuperscript{3-12} Mechanistic studies might provide


insights into the workings of the heterogeneous catalyst systems that could lead to new catalyst developments with improved desulfurization characteristics. In this manuscript, results with homogeneous systems that cleave the C-S bonds of thiophenes are summarized.

Figure 1. GC trace of polyaromatic sulfur containing compounds in light oil. (Reproduced with permission from ref. 2)

Results and Discussion

The reactive fragment [(C₅Me₅)Rh(PMe₃)] produced by heating solutions of (C₅Me₅)Rh(PMe₃)(Ph)H has been found to insert into a wide variety of thiophene C-S bonds (Scheme I).³ One of the first products to be structurally characterized was the adduct formed with 2,5-dimethylthiophene. This C-S insertion product shows a bent 6-membered ring in which the sulfur and butadiene portion of the ring form a plane that is oriented at an angle of 26° to the rhodium-sulfur-Cᵦ plane. The distances around the ring are consistent with a localized bonding structure (Scheme II). In contrast, the structure of the parent thiophene insertion product displays a planar metallathiabenzen ring, although once again bond length alternation is seen in the diene portion of the ring.¹³

Scheme I:

Figure 2. GC trace of desulfurized oil at each temperature. (Reproduced with permission from ref. 2)

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A number of other mononuclear transition metal complexes have been found to be capable of inserting into the C-S bonds of thiophenes, benzothiophenes, and dibenzothiophenes.\textsuperscript{14-20} As shown in Scheme II, the products formed have been characterized as containing in some cases a delocalized metallathiabenzenne ring and in others as an isolated thiobutadiene fragment. In most examples, the 6-membered ring formed upon insertion is planar. Delocalization of the π-electrons into an electron deficient metal has been cited as a reason for the molecule adopting a planar geometry, although steric factors seem to be more important in determination of the planarity.\textsuperscript{14,16}

The mechanism of insertion of [(C\textsubscript{5}Me\textsubscript{3})Rh(PMe\textsubscript{3})] into the C-S bond of thiophene has been investigated using a variety of labelling techniques.\textsuperscript{4} Attempts to directly observe an intermediate in the reaction at low temperature failed. Irradiation of (C\textsubscript{5}Me\textsubscript{3})Rh(PMe\textsubscript{3})(η\textsuperscript{2}-C,S-C\textsubscript{6}H\textsubscript{4}S) and (C\textsubscript{5}Me\textsubscript{3})Rh(PMe\textsubscript{3})(2-thienyl)H. Upon warming the sample, the 2-thienyl hydride complex converts to the C-S insertion complex. If this rearrangement is carried out in the presence of an excess of thiophene-d\textsubscript{4}, no incorporation of the deuterated thiophene into the product is observed, indicating that the rearrangement takes place without dissociation of the thiophene from the metal center.

The rearrangement of the 2-thienyl deuteride complex was examined by the reaction of (C\textsubscript{5}Me\textsubscript{3})Rh(PMe\textsubscript{3})Cl\textsubscript{2} with 2-thienyllithium followed by sec-butylborodeuteride. Rearrangement to the C-S insertion complex takes place in a non-regiospecific fashion, producing a 1:1 mixture of the two deuterated isomers. This observation could be interpreted in terms of an η\textsuperscript{2}-thiophene complex that rapidly equilibrates between the double bonds of the thiophene (Scheme III).

\textbf{Scheme III:}

A second labelling experiment proceeded similarly using 3-thienyllithium followed by sec-butylborodeuteride to generate the 3-thienyl deuteride complex. Rearrangement of the 3-thienyl deuteride complex to the 2-thienyl hydride complex proceeds regiospecifically, \textit{i.e.}, only the 3-deutero-2-thienyl product is formed (Scheme IV). A second, slower rearrangement to the C-S insertion product then proceeds non-regiospecifically, as observed in the previous experiment. The η\textsuperscript{2}-thiophene intermediate shown in Scheme IV explains the

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regiospecificity, since the metal remains on a single double bond in the thiophene molecule. Only by proceeding on to the S-bound complex does the molecule become symmetric, and then the C-S bond is broken.

Scheme IV:

Figure 3. Free energy diagram for the interaction of the fragment (C₅Me₅)Rh(PMe₃) with thiophene.

These reactions are summarized in the free energy diagram shown in Figure 3. Note that the figure shows the preferred kinetic selectivity for C-S bond cleavage over C-H bond activation, and that the C-S insertion product is more stable than the C-H insertion product. The presence of the η²-thiophene and S-thiophene intermediates, and their connections with the observed species are also made clear. Support for this reaction sequence comes from ab initio calculations done by Sargent, which indicates the presence of a low energy pathway from the S-thiophene complex to the C-S insertion complex.

Only one thiophene examined did not undergo C-S cleavage, tetramethylthiophene. Reaction with (C₅Me₅)Rh(PMe₃)PhH leads to the formation of the S-bound thiophene complex (C₅Me₅)Rh(PMe₃)(S-C₄Me₄S). The reason for this lack of reactivity has not been determined, but possibilities include a ‘buttressing effect’ that increases the steric interactions of the methyl groups in the 6-membered ring, or a combined electronic effect of the four methyl groups that changes the energetics of the interaction. A simple steric effect of a methyl group on the carbon attached to sulfur can be ruled out, since 2,5-dimethylthiophene is readily cleaved.

Benzothiophene was also found to undergo cleavage upon reaction with (C₅Me₅)Rh(PMe₃)PhH. The product formed arises from insertion of rhodium into the sulfur-vinyl bond only. An X-ray structure of this product shows puckering of the rhodathiabenzenene ring and bond length alternation consistent with localized bonding.

Reaction of (C₅Me₅)Rh(PMe₃)PhH with 2-methylbenzothiophene results in a similar product with cleavage of the sulfur-vinyl bond. In this case, however, continued heating of the sample leads to the rearrangement to a more stable product in which the metal has inserted into the sulfur-aryl bond (Scheme V). Apparently, the methyl group destabilizes the kinetic product sufficiently to allow it to rearrange to the thermodynamically preferred product. The origin of this preference can be attributed to the greater bond strength of a rhodium-aryl bond compared to a rhodium-vinyl bond. X-ray examination of both of these adducts show puckered, localized bonding structures. The rearrangement of the kinetic product to the thermodynamic product is intramolecular, as addition of 10 equivalents of thiophene during the rearrangement does not result in the formation of any of the thiophene C-S insertion product. The selectivities and interconversion can be accounted for in terms of the choices available to the S-bound

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complex, that is formed reversibly. At equilibrium, there is a 21:1 ratio of thermodynamic to kinetic product.\(^{12}\)

**Scheme V:**

Dibenzothiophene also reacts with \((C_5Me_5)Rh(PMe_3)PH\) to give a C-S insertion product. Examination of the structure shows a strongly puckered metallathiabenzene ring, and a twisting along the biphenyl axis.\(^{13}\) Similar insertion reactions are seen with a variety of substituted dibenzothiophenes. In an effort to determine the effects of methyl substitution, reactions of 2-methyldibenzothiophene, 3-methyldibenzothiophene, and 4-methyldibenzothiophene with \((C_5Me_5)Rh(PMe_3)PhH\) were examined. While the former two substrates showed little preference for cleavage of one C-S bond vs the other, the latter substrate showed exclusive cleavage of the C-S bond away from the adjacent methyl substituent. A similar observation was made with disubstituted 2,6-dimethyl dibenzothiophene (Scheme VI).\(^{22}\)

Heteroatom substituents on dibenzothiophene were found to have only modest effects on the insertion selectivity. Reactions of \((C_5Me_5)Rh(PMe_3)PH\) with 2-methoxy-, 2-fluoro-, 2-bromo-, 2-cyano, and 2-trifluoromethyl-dibenzothiophene were found to give mixtures of both insertion products, indicative of little electronic directing effect in the cleavage reaction (Scheme VII). Two benzonaphthothiophenes were examined. Reaction with one isomer gave only a single C-S insertion product that was characterized by X-ray crystallography as resulting from insertion away from the bulky naphthalene ring (Scheme VIII). Reaction with a second isomer gave a 2:1 ratio of products, since steric interference of the naphthalene ring was not problematic.\(^{22}\)

**Scheme VI:**

Most of the above dibenzothiophene products were characterized by X-ray crystallography, since this proved to be the only reliable method for determining the direction of insertion. In all cases

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except thiophene, puckering of the 6-membered ring was observed. For the dibenzo thiophenes, a twisting along the biphenyl linkage was also observed. Table I summarizes these structural features for the complexes.

Only one dibenzo thiophene complex did not undergo C-S cleavage upon reaction with \((\text{C}_5\text{Me}_5)\text{Rh(}\text{PMe}_3\text{)}\text{PhH}\). 4,6-dimethyl dibenzo thiophene reacts to give an S-bound complex that is labile, reacting with \(\text{PMe}_3\) to give \((\text{C}_5\text{Me}_5)\text{Rh(}\text{PMe}_3\text{)}_2\) plus free thiophene \((\Delta G^\ddagger = 24.5\text{ kcal/mmol}, 25^\circ \text{C})\). A similar S-bound complex is formed with 4,6-diethyl dibenzo thiophene.

The above reactivity patterns of substituted dibenzothiophenes are remarkably similar to that observed with the commercial Mo/Co industrial catalysts.\(^1\)\(^2\) There is difficulty associated with cleavage of a dibenzo thiophene C-S linkage when the adjacent carbon bears an alkyl substituent. No such steric barriers exist for substituted thiophenes or benzothiophenes, which are readily desulfurized and also react readily with \((\text{C}_5\text{Me}_5)\text{Rh(}\text{PMe}_3\text{)}\text{PhH}\). The similarity of reactivity suggests that a similar mechanism may be operative in the homogeneous and heterogeneous cases, i.e., it is the S-bound complex that is the immediate precursor of C-S bond cleavage. The similarity ends here, however, in that the \((\text{C}_5\text{Me}_5)\text{Rh(}\text{PMe}_3\text{)}\) fragment can only cleave one C-S bond and then becomes unreactive towards further cleavages or reaction with hydrogen.

The lack of further reactivity of the rhodium C-S insertion products can be attributed to the fact that the products are non-labile, coordinatively saturated Rh(III) complexes. In an attempt to produce a product with enhanced reactivity, the reactions of \((\text{C}_5\text{Me}_5)\text{Co(}C_2\text{H}_4\text{)}_2\) with thiophene,\(^6\) benzothiophene,\(^23\) and dibenzo thiophene were examined.\(^10\) In all three cases, a similar product was observed upon heating the cobalt complex (70°C) in the presence of the substrate. The product can be described as a C-S cleavage adduct similar to those found with \((\text{C}_5\text{Me}_5)\text{Rh(}\text{PMe}_3\text{)}\), except that the coordination site occupied by \(\text{PMe}_3\) is replaced by a bridging \((\text{C}_5\text{Me}_5)\text{Co}\) fragment (Scheme IX). Similar structures were observed in the reaction of benzothiophene with \(\text{Fe}_3(\text{CO})_{12}\)\(^{24}\) and of \((\text{C}_5\text{Me}_5)\text{Rh(}\eta^4\text{-}C_5\text{Me}_5\text{S)}\).\(^{20}\)

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\textbf{Scheme IX:}
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These molecules display an interesting fluxional process that equilibrates the two inequivalent \(\text{C}_5\text{Me}_5\) rings. These two rings differ only due to the bonding of the \(\text{C}_\beta\) carbon, since the sulfur and \(\text{C}_\alpha\) both bridge the two cobalt centers. The migration of \(\text{C}_\beta\) from one metal to the other equilibrates the two \(\text{C}_5\text{Me}_5\) rings. This process occurs most readily for the dibenzo thiophene complex, and less easily for the thiophene complex, since \(\eta^2\) coordination of the olefinic portion of the molecule disrupts the aromaticity in the dibenzo thiophene.

Since the rhodium complex was only capable of breaking one C-S bond, and the cobalt complex showed a tendency to form a dinuclear complex, a binuclear iridium complex was next examined for reactivity with thiophene. The complex \(\{(\text{C}_5\text{Me}_5)\text{IrH}_3\}_2\) was found to react with thiophene in the presence of a hydrogen acceptor to give a product in which both C-S bonds had been cleaved to give a butadiene ligand, and the sulfur remained as a \(\mu\)-sulfido ligand.\(^5\) Two intermediates are seen in the course this reaction, each of which grows in and then goes away. Scheme X shows the proposed mechanism for the formation of the butadiene containing product.\(^7\) The role of the \(t\)-butylethylene is believed to be needed to produce the

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\(^{23}\) W. D. Jones and J. H. Roache, unpublished results.

coordinatively unsaturated and reactive iridium complex, [(C₅Me₅)IrH]₂. Heating [(C₅Me₅)IrH₂]₂ alone under vacuum produces [(C₅Me₅)IrH]₂, which can also be used to desulfurize thiophene, but the reaction is still slow without added t-butylethylene. Coordination of sulfur followed by insertion of Ir(I) into the C-S bond gives intermediate A, a structure similar to that observed with cobalt but now with hydride ligands present. Reductive elimination of the vinyl hydride gives a butadienethiolate ligand, which can then coordinate to the coordinatively unsaturated iridium center to give intermediate B, also seen by ¹H NMR spectroscopy. From here, insertion of the internal olefin into the remaining Ir-H bond followed by β-elimination of the sulfur would lead directly to the butadiene µ-sulfido product with the correct regiochemistry for the butadiene, i.e. with the endo hydrogens above the sulfido ligand.

**Scheme X:**

![Scheme X](image)

This model for thiophene C-S cleavage would require two adjacent metal centers, a situation that could readily be accommodated at the edge of a MoS₂ crystallite in the commercial catalyst, either on the 1010 or 1010 edge. Figure 4 illustrates some of the different types of coordinatively unsaturated sites that are possible at the edge of a MoS₂ crystallite.¹ One can imagine a thiophene molecule undergoing C-S cleavage at one of these exposed metal centers and forming a µ-sulfido bridge as in the dinuclear cobalt complexes. Further reaction as seen with the diiridium system would then lead to desulfurization and the generation of the organic product plus a sulfur filled coordination site. Regeneration of the vacant site would occur under the reducing conditions of high temperature and high hydrogen pressure. Future studies and future catalysts will have to be able to also cleave C-S bonds of the hindered dibenzothiophenes, and it is hoped that studies such as these will lead to the next generation of catalysts.

**Figure 4.** Illustration of some of the different types of coordinatively unsaturated sites which are possible at the 1010 (a-e) and 1010 (f-j) edges of MoS₂. (Reproduced with permission from ref. 1)

**Conclusions**

A variety of C-S insertion reactions into thiophenes, benzothiophenes, and dibenzothiophenes are observed with rhodium, cobalt, and iridium fragments in the +1 oxidation state. The rhodium system provides evidence for the coordination of thiophene through sulfur as the immediate precursor to C-S bond cleavage. The inability to cleave certain substituted dibenzothiophenes is attributable to a steric effect on the accessibility to the C-S bond, rather than to an electronic deactivation of the C-S bond towards cleavage. Two metal centers in a bimetallic complex are found to be capable of activating both C-S bonds in thiophene, and the presence of the µ-sulfido bridge may be important in this second bond cleavage.

**Acknowledgement** is made to the National Science Foundation (CHE-9421727) for their support of this work. WDJ also thanks NATO for a travel grant.
Table I. Summary of X-ray Structural Data for Thiophene Insertion Compounds.

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