Introduction

The photooxidation of coordination compounds may occur by quite different mechanisms. Bimolecular electron transfer from electronically excited transition-metal complexes to reducing agents has been extensively investigated recently.\(^2\) However, the majority of photooxidations of a great variety of transition-metal complexes has been observed in halocarbon solvents; where solvent molecules coordinate ligands due to their importance in biological systems.\(^12\) Another example is the photooxidation of certain cyanide complexes in aqueous solution accompanied by the production of solvated electrons.\(^3\) Although, the majority of photooxidations of a great variety of transition-metal complexes has been observed in halocarbon solvents; where solvent molecules act as electron acceptors.\(^4\)\(^-\)\(^10\) Charge-transfer-to-solvent (CTTS) states may be important in some cases;\(^10\) the nature of the photoactive excited states does not seem to be very clear. The present study may contribute to a better understanding of these photooxidations.

For our investigation we selected 1,2-dithiolene complexes \(^1\) of Ni, Pd, and Pt. This choice was guided by the following considerations. By a variation of the metal, the dithiolene ligand, and the charge of the complex, its electronic structure can be altered to a great extent. It was expected that such changes would influence the photochemical behavior considerably and lead to a correlation between electronic structure and photoreactivity.

Another interesting aspect of the present study is the growing interest in coordination compounds with sulfur-containing ligands\(^12\) due to their importance in biological systems. So far only a few investigations of the photochemistry of sulfur-containing complexes have been carried out\(^8\)\(^-\)\(^12\) although such complexes are known to participate in photochemistry.\(^12\)

Experimental Section

Materials. The complexes \([\text{Ni(C}_2\text{H}_4\text{)}_2\text{][S}_2\text{C}_2\text{(CN)}_2\text{]}_2\), with \(M = \text{Ni}, \text{Pd}, \text{Pt}\), \(\text{Ni}, \text{Pt}, \text{Ni}, \text{Pd}, \text{Pt}\), and \(\text{Ni}, \text{Pt}, \text{Ni}, \text{Pd}, \text{Pt}\) were prepared according to published procedures. Their electronic absorption spectra agreed well with those reported previously.\(^16\)\(^-\)\(^19\) All solvents used in the photochemical
1,2-Dithiolene Complexes of Ni, Pd, and Pt

Figure 1. Spectral changes during the photooxidation of 1.39 × 10⁻⁴ M \([\text{Ni(C(H₂)₃)}₂][\text{PtS₂C₂(CN)₂}]\) in CHCl₃ at (a) 0 and (d) 20-min irradiation time, \(\lambda_{irr} = 333\) nm, and 1-cm cell.

Figure 2. Spectral changes during the photooxidation of 1.84 × 10⁻⁴ M \([\text{Ni(C(H₂)₃)}₂][\text{NiS₂C₂(CH₃)₂}]\) in CHCl₃ at (a) 0 and (e) 18-min irradiation time, \(\lambda_{irr} > 340\) nm, and 1-cm cell.

experiments were Spectro grade. Purification of the solvents by chromatography (Al₂O₃) and distillation from molecular sieves (5 Å) did not affect the results.

Photolyses. The light source was a high-pressure mercury lamp Osram HBO 200 W/2. The following Schott interference filters of the type PIL were used for the irradiation at selected wavelengths: 313, 333, 366, 406, and 436 nm. The photolyses were carried out at room temperature in 1- and 10-cm spectrophotometer cells. For quantum yield determinations the complex concentrations were such that room temperature in 1- and 10-cm spectrophotometer cells. For quantum yield determinations the complex concentrations were such as to have essentially complete light absorption. The total amount of photolysis was limited to less than 5% to avoid light absorption by the photoproduct. Absorbed light intensities were determined by ferrioxalate actinometry.⁶

Progress of the photooxidation was monitored by UV-visible and near-IR spectral measurements with a Varian-Techtron Super Scan 3 recording spectrophotometer and a Zeiss PMQ II spectrophotometer for measurements at selected wavelengths.

The photoproducts \(\text{M}[\text{S}_2\text{C}_2\text{C} = \text{N}]\) and \(\text{M} = \text{Ni}, \text{Pd}, \text{Pt}\) were identified by their absorption spectra. The extinction at the maximum of the intense near-IR absorptions was used to determine the concentrations of the photoproducts: \(\text{Ni}[\text{S}_2\text{C}_2\text{C} = \text{N}]\), \(\lambda = 848\) nm (\(\epsilon = 8000\)); \(\text{Pd}[\text{S}_2\text{C}_2\text{C} = \text{N}]\), \(\lambda = 1111\) nm (\(\epsilon = 13800\)); \(\text{Pt}[\text{S}_2\text{C}_2\text{C} = \text{N}]\), \(\lambda = 855\) nm (\(\epsilon = 11700\)); \(\text{Ni}[\text{S}_2\text{C}_2\text{C} = \text{N}]\), \(\lambda = 866\) nm (\(\epsilon = 30903\)).¹⁶ As indicated by the spectral changes, the photolysis of \(\text{M}[\text{S}_2\text{C}_2\text{C} = \text{N}]\) (\(\text{M} = \text{Ni, Pd, Pt}\)) took place without any side reaction. In the case of \(\text{Ni}[\text{S}_2\text{C}_2\text{C} = \text{N}]\), isosbestic points were observed but were less pronounced, indicating some side reactions. But these side reactions were not important since the quantum yield for product formation and disappearance of the starting complex were equal within 5%.

Results and Discussion

The most interesting aspect of the chemistry of 1,2-dithiolene complexes is the redox behavior because the ligands themselves can undergo consecutive one-electron-transfer processes while the metal may retain its oxidation state.¹¹ The limiting electronic structures of the ligand are the oxidized dithioketone \(\text{S} = \text{C} = \text{O} - \text{C} = \text{S}\) and the reduced ethylenedithiolate dianion \(\text{S} = \text{C} - \text{O} - \text{C} - \text{S}\), which has two electrons more. For square-planar complexes of Ni, Pd, and Pt in the oxidation state +II, five members of a complete electron-transfer series \([\text{M(S}_2\text{C}_2\text{R}_2\text{)}_2]\) with the charges \(z = 2⁻\) to \(2⁺\) are feasible. But dependent on the substituent \(R\), only neutral and anionic complexes are well characterized.¹¹

The electronic spectra of 1,2-dithiolene complexes of Ni, Pd, and Pt are very rich.¹¹,¹⁶-¹⁹ The most important orbitals are 2b₂ and 3b₂, which are nondegenerate and essentially ligand-localized \(\pi\) orbitals.¹⁶,¹⁹,²¹ In the neutral complexes with \(z = 0\), \(2b₂\) is empty while the lower \(2b₁\) is occupied. The allowed transition \(2b₁ \rightarrow 2b₂\) gives rise to an intense absorption band in the near-infrared region. This transition is possible when the charges are \(z = 1⁻\) (half-filled \(2b₂\)) and \(z = 1⁺\) (half-filled \(2b₁\)). The corresponding absorption is observed for \(z = 1⁻\) (Experimental Section and Figure 1). Although cationic 1,2-dithiolene complexes are not stable, this absorption band does occur in the spectra of the similar diimine complexes with \(z = 1⁺\) where the sulfur of the dithiolate is replaced by imino groups NH.¹⁰ʰ These long-wavelength bands are missing in the spectra of the dianionic complexes.¹⁶,¹⁸,¹⁹ (Figure 1) with \(z = 2⁺\) (filled \(3b₂\)). In addition, many other absorptions occur down to at least 600 nm irrespective of the charge. The assignment of these bands is controversial. They may be of the ligand field (LF), charge-transfer metal to ligand (CTML), and intraligand (IL) type.¹⁴,¹⁶,¹⁸,¹⁹,²¹,²² They should occur at widely differing energies depending on the metal, the substituent \(R\) at the dithiolene ligand, and the charge \(z\).

Fortunately, the interpretation of our results does not depend on the assignments of the absorption bands since the light sensitivity of those complexes which are photooxidized is es-

Table 1. Photooxidation Quantum Yields (\(\phi_{oxy} = 313\) nm) and Half-Wave Potentials of Oxidation (vs. SCE) of \([\text{M(S}_2\text{C}_2\text{R}_2\text{)}_2]\) in CHCl₃

<table>
<thead>
<tr>
<th>metal</th>
<th>ligand</th>
<th>charge (z)</th>
<th>(\phi_{oxy} (313) nm)</th>
<th>(E_{1/2}^{\beta} (V))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni</td>
<td>(\text{S}_2\text{C}_2\text{C} = \text{N})</td>
<td>2⁻</td>
<td>0.25</td>
<td>+0.23</td>
</tr>
<tr>
<td>Pd</td>
<td>(\text{S}_2\text{C}_2\text{C} = \text{N})</td>
<td>2⁻</td>
<td>0.12</td>
<td>+0.44</td>
</tr>
<tr>
<td>Ni</td>
<td>1⁻</td>
<td></td>
<td>0.23</td>
<td>+0.21</td>
</tr>
<tr>
<td>Pd</td>
<td>1⁻</td>
<td></td>
<td>0</td>
<td>+1.02</td>
</tr>
<tr>
<td>Pt</td>
<td>1⁻</td>
<td></td>
<td>0</td>
<td>c</td>
</tr>
<tr>
<td>Ni</td>
<td>(\text{S}_2\text{C}_2\text{C} = \text{N})</td>
<td>0</td>
<td>0</td>
<td>c</td>
</tr>
<tr>
<td>Pt</td>
<td>0</td>
<td></td>
<td>0</td>
<td>c</td>
</tr>
<tr>
<td>Ni</td>
<td>1⁻</td>
<td></td>
<td>0.1</td>
<td>+0.22</td>
</tr>
</tbody>
</table>

a Experimental error ± 5%.  b Reference 17.  c Not reported but expected to be above 1 V; see ref 11.
sentially restricted to wavelengths between 300 and 350 nm regardless of the metal, the substituent R, and the charge z.

While in most solvents, e.g., acetonitrile, all complexes investigated in the present study are not light sensitive, some of them are oxidized with high quantum yields upon irradiation with light of wavelengths 300–350 nm, when dissolved in CHCl₃ (Table I). By analogy with other photooxidations in halocarbon solvents, our results are consistent with eq 1.

\[ [\text{M(S₂C₂R₂)}]^{+} + \text{CHCl₃} \rightarrow [\text{M(S₂C₂R₂)}]^{+ \cdot 1} + \text{Cl}^{-} + \text{CHCl₃} \] (1)

While the one-electron oxidation product of the complex is stable, the CHCl₃ radical may undergo secondary reactions.

The electronic structures and hence the electronic transitions of all complexes which undergo the photooxidation are so different that any common explanation seems to be impossible. However, a careful survey of the published data shows that all photoactive complexes are oxidized at low potentials in the same region \( (E_{1/2} = 0.1–0.5 \text{ V vs. SCE; see Table I}) \). It follows that the photooxidation is controlled only thermodynamically and occurs regardless of the type and energy of the intramolecular absorption bands displayed by any particular complex. Those complexes which are oxidized at potentials below 0 V are subject to air oxidation and were not included in the present study. They are also expected to be photooxidized at irradiating wavelengths longer than 350 nm. Those complexes which are oxidized above 0.5 V, e.g., \( [\text{M(S₂C₂(CN)}]^{+} \) (M = Ni, Pd, Pt) and \( [\text{M(S₂C₂(C₆H₅)}]^{+} \) (M = Ni, Pt) were not observed to be photosensitive. They are expected to be photooxidized with light of wavelengths shorter than 300 nm. Unfortunately, CHCl₃ absorbs in this region.

It is interesting to note that a variety of other complexes which are photooxidized between 300 and 350 nm are also oxidized at potentials between 0.1 and 0.5 V vs. SCE. Some of them are related to the 1,2-dithiolene complexes, and others are completely different (Table II). In some cases these complexes displayed CTTS absorption bands when dissolved in halocarbon solvents. Photooxidation may then be due to light absorption into CTTS bands.

We searched for the presence of CTTS bands by measuring the difference spectra in acetonitrile and CHCl₃. In the case of Pt[\( \text{S₂C₂(CN)}]^{+} \), careful measurements revealed two new maxima at 318 and 363 nm. For Ni[\( \text{S₂C₂(CN)}]^{+} \) there was an indication of a less well-pronounced new band around 335 nm. However, all our complexes have very intense intramolecular absorptions (~20 000) in the 300–350 nm region. Hence any other absorption of much lower intensity may be obscured. In addition, any weak new absorption detected by difference spectroscopy in two solvents may be artificial and due to any unspecified solvent shift. But irrespective of the presence of CTTS bands, most of the light causing photooxidation of the 1,2-dithiolene complexes is certainly absorbed by bands of a different origin.

The energy of the CTTS state should depend not only on the redox potential of the complex but also on that of the solvent. The photoactive wavelength region is expected to shift to the red with increasing oxidizing power of the solvent. The photoactive wavelength region is expected to shift to the red with increasing oxidizing power of the solvent. This assumption was confirmed. In addition to CHCl₃ \( (E_{1/2} = -1.67 \text{ V vs. SCE}) \) as solvent, the complex Ni[\( \text{S₂C₂(CN)}]^{+} \) was also photolyzed in CH₂Cl₂ \( (E_{1/2} = -2.33 \text{ V vs. SCE}) \) and CCl₄ \( (E_{1/2} = -0.78 \text{ V vs. SCE}) \). The quantum yield \( (\lambda_{\text{air}} = 366 \text{ nm}) \) of photooxidation increased from \( \phi = 0.002 \) in CH₂Cl₂ to \( \phi = 0.013 \) in CHCl₃ and \( \phi = 0.07 \) in CCl₄. While 366-nm light leads to a fairly efficient population of the reactive CTTS state in CCl₄, it is apparently not sufficient to reach this state at higher energies in CH₂Cl₂ as solvent. Such a solvent dependence was also observed for the photooxidation of \( [\text{S₄C₄H₄Fe(CO)}]^{+} \).

Shirom et al. investigated extensively the photooxidation of cyanide complexes in aqueous solution with concomitant formation of solvated electrons. They concluded that in the main pathway the electron is ejected to the solvent from CTTS excited states. These states can be reached by direct light absorption into a CTTS band or by internal conversion from other excited states. Even when a CTTS band was not detected, an efficient production of solvated electrons was observed. Consequently, the quantum yield of photooxidation does not depend on the fraction of light absorbed by a CTTS band. It was suggested that the reaction from the CTTS state is an extremely rapid process which competes successfully with other modes of deactivation. We assume that the same mechanism applies to the photooxidation of 1,2-dithiolene complexes of Ni, Pd, and Pt. The energy of the CTTS state depends only on the redox potentials of the complex but also on that of the solvent. It may be populated by an efficient internal conversion from any higher intramolecular excited state. The photooxidation takes place before a deactivation to lower excited states that are nonreactive occurs.

A further point of interest is the observation that the quantum yield of photooxidation increases with decreasing wavelength of irradiation (Table III). This behavior seems to be characteristic for the reactivity of CTTS excited states. Shirom et al. suggested that the electron ejection is rapid enough to occur from higher vibrational levels of the CTTS excited state. Such a hot excited-state mechanism may account for the observed wavelength dependence.

In addition to the photooxidation starting from CTTS states, it was concluded that other excited states, particularly of the CTML type, can lead to the release of an electron. These reactions seem to be much slower processes. The photooxidation of isocyanide complexes of Cr, Mo, and W may lead to the release of an electron. The excited state, can lead to the release of an electron.
wavelength CTML bands (436 nm) of these complexes dissolved in inert solvents is followed by an emission from CTML excited states. In CHCl₃ this emission was quenched with simultaneous oxidation of the complexes. A further example of this type is apparently the photooxidation of (2,2'-bipyridine)(3,4-toluenedithiolato)platinum(II) in CHCl₃ following charge-transfer ligand-to-ligand (CTTL) excitation at long wavelength (577 nm). It is interesting to note that the similar complex Ni⁴⁺(phen)[S₂C₂(C₆H₅)₂] (phen = 1,10-phenanthroline) is photooxidized in CHCl₃ only by short-wavelength irradiation (λ < 350 nm), while the CTLL band at longer wavelength is not photoactive. Compared to the Pt complex, the Ni compound should have LF excited states at much lower energies which can quench the CTLL state. In distinction, the CTTS excited states at higher energies react so rapidly that other lower excited states apparently do not interfere.

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Registry No. [Ni(S₂C₂(CN)₂)₃]⁺, 14876-79-6; [Pd(S₂C₂(CN)₂)₃]⁺, 19555-33-0; [Pt(S₂C₂(CN)₂)₃]⁺, 15152-99-5; [Ni(S₂C₂(CN)₂)₂]⁺, 46761-25-5; [Pt(S₂C₂(CN)₂)₂]⁺, 19570-29-7; [Pt(S₂C₂(CN)₂)₂], 14977-45-8; Ni(S₂C₂(C₆H₅)₂), 28984-20-5; Pt(S₂C₂(C₆H₅)₂), 15007-55-3; [Ni(S₂C₂(C₆H₅)₂)₂]⁺, 14879-11-9.

High-Pressure Mechanistic Studies of the Photochemical Reactions of Transition-Metal Complexes. 3. Medium Effects on the Photoaquation of Some Co(II) and Cr(II) Complexes in Solution

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The viscosity dependence of the photoaquation reactions of Co(CN)₃⁻, Cr(CN)₆³⁻, Cr(CNS)₆³⁻, Cr(NH₃)₅NCS⁺, and Cr(NH₃)₅Cl⁺ was studied in glycerol/water mixtures (565 wt.%) at normal pressure. The results indicate that significant cage recombination occurs during the photosubstitution of NH₃ in Cr(NH₃)₅NCS⁺ and Cr(NH₃)₅Cl⁺ in pure water. However, no significant cage effect was observed in the photosubstitution of CN⁻ and NCS⁻ in the studied systems. In addition, the combined medium–pressure dependence of the photoaquation of NCS⁻ in Cr(NH₃)₅NCS⁺ was studied over the ranges 1 ≤ p ≤ 1500 bar and 1 < η < 30 ep. A rather unexpected pressure dependence of the quantum yield is reported for this reaction in highly viscous media. The results are discussed in reference to data reported in the literature.

Introduction

The earlier papers in this series studied high-pressure mechanistic studies of the ligand field photolysis of some cationic Cr(III) ammine and some anionic pseudohalogeno Co(III) and Cr(III) complexes in aqueous solution. From the pressure dependence of the observed quantum yields for the photoaquation processes, apparent volumes of activation were calculated. These data permitted a detailed discussion of the molecular nature of the intimate mechanism involved in such photoaquation reactions.

It was concluded that reactions of the type Cr(NH₃)₅X₃⁻ + H₂O → Cr(NH₃)₅OH₂⁻ + X⁻ (1)

proceed according to an 1ₐ mechanism, during which substantial Cr–X bond cleavage occurs in the transition state. Furthermore, reactions of the type Cr(NH₃)₅X₃⁻ + H₂O → cis-Cr(NH₃)₅O⁺(OH)X₂⁻ + NH₃ (2)

also follow such a mechanism but with significantly less Cr–NH₃ bond cleavage in the transition state. In contrast, it was found that reactions of the type

ML₅⁻ + H₂O → ML₅(OH)²⁻ + L⁻ (3)

M = Co(III) for L = CN

M = Cr(III) for L = NCS, CN

proceed according to an 1ₐ mechanism in which some M–OH₂⁻ bond formation occurs in the transition state.

Scandola et al. observed a different approach. They studied medium effects on the photoaquation of Co(CN)₆³⁻ and presented evidence for the occurrence of cage recombination in such photosubstitution reactions. The viscosity of the solvent had a meaningful influence on the quantum yield of the aquation reaction, and a good correlation between these variables was reported. The cage effect seemed to be quite efficient in more viscous solvents but relatively unimportant in pure aqueous solution for the studied system.

Scandola et al. concluded that these tendencies may not hold for other photosubstitution reactions and are expected to depend on the nature of the leaving group.

Wong and Kirk observed similar viscosity effects for the photoaquation of Co(CN)₆³⁻ in water and in water/ethylene glycol, and in water/ethanol mixtures. In contrast, however, they argued that cage recombination is not established by the data since they do not find such an effect in water/acetonitrile and water/poly(vinylpyrrolidone) mixtures. Alternative explanations such as changes in the extent of solvent structure.