Photochemistry of Pentaamminethiocyanato-cobalt(III) Ion

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Irradiation of the first ligand field band of Co(NH₃)₅SCN²⁺ in aqueous solution did not lead to any efficient photochemical reaction. Irradiation of the first CT band led to a redox decomposition with formation of Co²⁺ and to a simultaneous linkage isomerization with formation of Co(NH₃)₅NCS²⁺. At 333 nm irradiating wavelength the quantum yields were 0.48 for redox reaction and 0.24 for isomerization. In the solid state CT excitation led to the linkage isomerization without redox decomposition. These results are discussed with regard to the photochemistry of the complex ions Co(NH₃)₅NO₂²⁺ and Co(NH₃)₅NCS²⁺.

Introduction

The photochemistry of aqueous Co(III) complex ammines and acidoammines has been extensively investigated. The majority of these photochemical reactions is certainly initiated by CT excited states. However, the mechanism of the reactions which follow CT excitation is not yet completely understood. The main reactions are redox decompositions and substitutions. A remarkable exception is Co(NH₃)₅NO₂²⁺ which also undergoes nitro-nitrito linkage isomerization.

The complex Co(NH₃)₅SCN²⁺ offered the opportunity to study a second example of a light induced linkage isomerization. The choice of Co(NH₃)₅SCN²⁺ was initiated by an accidental observation of Buckingham, Creaser, and Sargeson, who investigated the thermal thiocyanato isothiocyanato linkage isomerization of Co(NH₃)₅SCN²⁺. They found that this isomerization occurred also under the influence of light in the infrared spectrometer.

The comparison of the photochemistry of Co(NH₃)₅SCN²⁺ and of its linkage isomer Co(NH₃)₅NCS²⁺, which has been investigated earlier, should be another interesting aspect of the present study. The homolytic bond splitting which may follow CT excitation of both complexes should generate the same radical pair Co(NH₃)₅⁺ and NCS. If the reactivity of this radical pair is independent of its origin CT excitation of both linkage isomers should lead to the same product distribution.

Experimental

Materials

[Co(NH₃)₅NCS][ClO₄]₂ and [Co(NH₃)₅SCN]Cl₂ were prepared according to known procedures. The compounds were of good spectroscopic purity; their visible and ultraviolet absorption spectra, shown in Figure 1, and their i.r. spectra agreed well with those reported for analyzed samples.

Photolysis Procedures

The light source used was a 100 W Osram high-pressure mercury lamp. A Schott interference filter 333 limited the ultraviolet radiation emitted to the wavelength of the mercury line at 333 nm. Alternatively, the Schott glass filter GG 495 transmitted light longer than 480 nm.

The photolyses were carried out at room temperature in a 5 cm cylindrical spectrophotometer cell, using concentrations of complex such as to have essentially complete light absorption. The solutions were 0.01M in perchloric acid. An aliquot of the solution to be analyzed was kept in the dark, at the same temperature, so that any thermal reaction could be noted and allowed for. However, correction for the thermal reaction was negligible. The total amount of photolysis was limited to about 10% to avoid secondary photochemical reactions. Absorbed light intensities were determined by Ferrioxalate actinometry or by Reineckate actionmetry in the visible region.

Analytical Procedures

Cobalt (II)

Cobaltous ion was determined by forming the blue complex ion, Co(NCS)₂⁻ in a water-acetone solution and measuring the absorbance (ε = 1.7 x 10³ at 625 nm), according to a literature procedure.

Thiocyanate

Free thiocyanate ion was determined by adding an aliquot of the irradiated (or dark) solution to a four-fold volume of a reagent consisting of 0.1M ferric nitrate in 0.5M perchloric acid, and measuring the absorbance due to the resulting ferric thiocyanate complex (ε = 4.3 x 10³ at 450 nm).
Measurements

Infrared spectra were measured with a Perkin Elmer 325 IR spectrophotometer. Samples were KBr pellets. Visible and ultraviolet spectra were measured in 1 cm cells with a Cary 14 recording spectrophotometer and a Zeiss PMQ II spectrophotometer for measurements at selected wavelengths.

Results

The absorption spectra of Co(NH$_3$)$_2$SCN$^{2+}$ and Co(NH$_3$)$_2$NCS$^{2+}$ (Figure 1) exhibit their first LF maxima at 512 nm ($\epsilon = 74$) and at 498 nm ($\epsilon = 179$). The intense absorption maxima at 288 nm ($\epsilon = 15,600$) and at 306 nm ($\epsilon = 1490$) are assigned to SCN$^-$ and NCS$^-$ to Co(III) CT transitions of both complexes. Corresponding assignments have been made for the absorption bands of the two linkage isomers Co(CN)$_2$SCN$^-$ and Co(CN)$_2$NCS$^-$.

Photolysis in Aqueous Solution

The complex ion Co(NH$_3$)$_2$SCN$^{2+}$ was sufficiently stable under our experimental conditions. Hence any interference by thermal reactions$^6$ was excluded. Co(NH$_3$)$_2$SCN$^{2+}$ did not undergo any efficient photothermal reaction if the irradiating wavelength was restricted to the region of the LF band ($\lambda > 480$ nm). The quantum yield of any of the possible reactions was estimated to be smaller than 10$^{-3}$.

Upon irradiation of the CT band (333 nm irradiating wavelength) Co(NH$_3$)$_2$SCN$^{2+}$ underwent a redox decomposition. Co$^{2+}$ was formed with the quantum yield $\Phi = 0.48$. This redox reaction was accompanied by a change of the absorption spectrum (Figure 2). It was assumed that the products of the redox decomposition have a negligible absorption down to at least 300 nm. The extinction at longer wavelengths should then decrease according to the amount of decomposition. However, the decrease of the extinction was dependent on the wavelength and at 475 nm there was no change at all (Figure 2). It follows that in addition to the redox products another complex must have been formed, which had a larger extinction than the starting complex at least at 475 nm. The most probable candidate for another photoprodut was Co(NH$_3$)$_2$NCS$^{2+}$ which has a higher extinction than Co(NH$_3$)$_2$SCN$^{2+}$ almost over the entire wavelength region down to 320 nm (Figure 1). A comparison showed that the ratio of the extinction coefficients of the isothiocyanato and thiocyanato isomers is indeed highest at 475 nm. If the redox decomposition and linkage isomerization were the only photochemical reactions the ratio of both reaction modes can be calculated from the extinction coefficients of the two linkage isomers at 475 nm. The calculation showed that two thirds of the photolyzed ions underwent a redox decomposition and the remaining third was transformed into its linkage isomer. The change of the extinction at any other wavelength should then satisfy the equation:

$$A_0 - A = [\text{Co-NCS}] (3\epsilon_{\text{Co-SCN}} - \epsilon_{\text{Co-NCS}})$$

The agreement was better than 5% for all wavelengths above 400 nm. Hence the assumption that only the redox decomposition and linkage isomerization took
Photochemistry of Co(NH₃)₅NCS⁺⁺

Figure 2. Spectral changes during the photolysis of 3.8 × 10⁻³ M Co(NH₃)₅SCN⁺⁺ in 10⁻² M HClO₄; 333 nm irradiating wavelength. Reading downward: 0 min (a), 15 min, 30 min, 60 min (d) irradiation time.

place is certainly justified. However, the accuracy of the measurements is not great enough to exclude the formation of very small amounts of other photoproducts as for example Co(NH₃)₅H₂O⁺⁺. If secondary photolysis of Co(NH₃)₅NCS⁺⁺ was avoided the change of the optical density at any wavelength was not large due to the high extinction of Co(NH₃)₅NCS⁺⁺. The photolysis followed a zeroth order kinetics up to 7% of photolyzed starting complex.

Since the efficiency of the photoisomerization is independent of the concentration of Co(NH₃)₅SCN⁺⁺ the rearrangement should occur by an intramolecular mechanism. The photolysis produced also free NCS⁻ with the quantum yield Φ = 0.33. The formation of NCS⁻ must then result from side reactions of the redox decomposition as it has been observed in similar cases.²⁻²⁻¹⁻¹

Photolysis in the Solid State

Since the extinction of the CT bands of Co(NH₃)₅SCN⁺⁺ and its linkage isomer is rather high any photoisomerization of pure [Co(NH₃)₅SCN]Cl₂ would be restricted to the surface of the crystals and hence difficult to detect. A dilution was achieved by incorporation of [Co(NH₃)₅SCN]Cl₂ in KBr pellets. These pellets are also useful for the detection of linkage isomerization by i.r. spectroscopy.

Upon irradiation of the LF band of Co(NH₃)₅SCN⁺⁺ (λ > 480 nm) the i.r. spectrum did not change. However, shorter-wavelength irradiation into the CT band initiated a linkage isomerization with formation of [Co(NH₃)₅NCS]Cl₂ as indicated by the change of the i.r. spectrum as has been observed previously.⁶ The first observation of this photoisomerization⁶ which stimulated the present investigation was possibly due to the exposure of the starting complex to the UV light of the light source in the i.r. spectrometer. The linkage isomerization was apparently the only photochemical reaction in the solid state. Co⁺⁺ was not detected in an aqueous solution of the irradiated KBr pellets.

Discussion

It was shown that aqueous Co(NH₃)₅SCN⁺⁺ underwent a photoredox decomposition which is accompanied by a linkage isomerization with formation of Co(NH₃)₅NCS⁺⁺. At 333 nm irradiating wavelength the quantum yields were Φ = 0.48 for redox reaction and Φ = 0.24 for isomerization. If solid [Co(NH₃)₅ SCN]Cl₂ was irradiated only the linkage isomerization took place.

These photochemical reactions of Co(NH₃)₅SCN⁺⁺ are quite similar to those of Co(NH₃)₅NO₃⁺⁺ which underwent a redox decomposition and a simultaneous linkage isomerization with formation of Co(NH₃)₅ ONO₂⁺⁺. The photoisomerization also occurred in the solid state.⁵ However, in contrast to Co(NH₃)₅ NO₃⁺⁺ which was photosensitive over its entire absorption spectrum the photoreactivity of Co(NH₃)₅ SCN⁺⁺ was restricted to the CT region. Upon irradiation of the LF band no photochemical change was observed. Hence we conclude that the photochemical reactions of Co(NH₃)₅SCN⁺⁺ were initiated by an excited CT state.
In analogy to Co(NH$_3$)$_2$NO$_2$$^{2+}$ the photochemistry of Co(NH$_3$)$_2$SCN$^{2+}$ may be explained on the basis of a radical pair model.$^{3-5,7,8,15}$ The separation of the radicals Co(NH$_3$)$_2^+$ and NCS which were produced in the primary step following CT excitation led to the redox reaction whereas the linkage isomerization should be the result of a primary or secondary geminate recombination of the radicals. In the solid state a separation of the radical pair is apparently not possible. Hence only the linkage isomerization was observed.

It is then interesting to compare the photochemistry of Co(NH$_3$)$_2$SCN$^{2+}$ and its linkage isomer since CT excitation of both complexes should produce identical radicals. Earlier investigations have shown that CT excitation of aqueous Co(NH$_3$)$_3$NCS$^{2+}$ led to a redox decomposition and an aquation with formation of Co(NH$_3$)$_2$H$_2$O$^{3+}$. The relative efficiencies of the two reaction modes were 70% redox reaction and 30% aquation.$^8$ Both reactions have been assumed to have also the radical pair Co(NH$_3$)$_3^+$ and NCS as a common precursor. The absence of any photoisomerization of Co(NH$_3$)$_3$NCS$^{2+}$ may be an indication that the radical recombination led always to the isothiocyanato isomer. However, the observation that only one of the linkage isomers, Co(NH$_3$)$_2$NCS$^{2+}$, undergoes an efficient photoaquation suggests that the reactivity of the radical pair generated in the first step is not independent of the way in which it is produced.

Any conclusive explanation of the different behavior of both isomers cannot yet be given. However, the occurrence of the photochemical linkage isomerization of the thiocyanato isomer may be related to the observation that the Co–SCN bond is bent at the sulfur atom with an angle of 105° while the Co–NCS bond is linear.$^{15}$ The radical pair formation following CT excitation should be associated with a motion of the coordinating ligand atom away from the metal center. In the case of the thiocyanato isomer this motion could lead to a rotation of the SCN radical. Its nitrogen atom may then move into a favorable position to reattach to the complex fragment. Alternatively, the different reactivity of both isomers may be used to support an excited state model.$^{17}$ However, it is our opinion that in comparison to the radical pair model the excited state mechanism is based on much more arbitrary assumptions which are difficult to verify.

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References

17. see Ref. 9 and references cited therein.