REACTIVE INTERMEDIATES IN THE PHOTOLYSIS OF CYCLOPENTADIENYLCOBALT(I) DICARBONYL AND IN THE REDUCTION OF CYCLOPENTADIENYLCOBALT(III) CARBONYL DIIODIDE

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(Received July 19th, 1976)

Summary

Photolysis of \((C_5H_5)Co(CO)_2\) (I) in toluene or petroleum ether solution at \(-78^\circ C\) generates the unsaturated monocarbonyl species \((C_5H_5)Co(CO)\), which was identified in solution by its IR spectrum. At room temperature, this monocarbonyl can associate with excess I to give \((C_5H_5)_2Co_2(CO)_3\), or dimerize to \((C_5H_5)_2Co_2(CO)_4\). The latter is stable as a solid, but in solution it is slowly converted to the insoluble trimer \((C_5H_5)_3Co_3(CO)_3\). \((C_5H_5)_2Co_2(CO)_2\) is symmetrically cleaved by phosphines to \((C_5H_5)Co(CO)(PR_3)\), while diolefins bring about unsymmetrical cleavage to give \((C_5H_5)Co(\text{diolefin})\) and \((C_5H_5)Co(CO)_2\).

Introduction

Among organometallic reactions involving coordinatively unsaturated intermediates, those of sandwich compounds with two and only two available coordination sites offer a unique opportunity to study reaction patterns and reactivity parameters. The availability in coordination compounds with a 14-electron core structure of more than one coordination site for substitutions, oxidative additions, and other basic reaction steps appears to be associated with a variety of reactions relevant to homogeneous catalysis. At the same time, however, structural degrees of freedom are limited by the fairly rigid sandwich skeleton, so that reasonably accurate descriptions of the transition states may be possible. A striking example of such a 14-electron coordination core structure is that of bis(cyclopentadienyl)titanium(II), for which a wide variety of modes

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of reaction and their synthetic and catalytic applications have been reported [1]. The later transition metals, particularly those of Group VIII could yield comparable 14-electron core structures with two available coordination sites only in half-sandwich arrangements. One such 14-electron half-sandwich core which contains its metal in a low, yet reasonably stable, oxidation state is that of cyclopentadienylcobalt(I) and its homologues. In accord with this unique position, the coordination chemistry of cyclopentadienylcobalt(I) is comparable to that of bis(cyclopentadienyl)titanium(II) [2]. The dicarbonyl complex I, \((C_5H_5)Co(CO)_2\), for example, is susceptible to photolysis; upon irradiation a trinuclear complex II, \((C_5H_5)Co_3(CO)_3\) has been obtained [3]. A diversity of products arise from thermal and photochemical reactions of the dicarbonyl I with substituted acetylenes [4–6]; this indicates the generation from I of coordinatively unsaturated intermediates e.g. in the photolytic reaction I \(\rightarrow\) II. Vollhardt et al. [7] recently investigated the nature of these intermediates. We report further observations concerning the identity and mutual conversion pathways of reactive intermediates in these systems.

Results

1. Low temperature photolysis of \((C_5H_5)Co(CO)_2\)

When toluene solutions of \((C_5H_5)Co(CO)_2\) are irradiated at \(-78^\circ\text{C}\) with UV light, while being continuously swept by a slow stream of \(N_2\), the IR spectra of the solutions change as follows. The initial dicarbonyl complex I exhibits two absorptions of nearly identical intensity in its IR spectrum at 2020 and 1955 \(\text{cm}^{-1}\). The band at 2020 \(\text{cm}^{-1}\) decreases in intensity upon irradiation. In dilute solutions, after an irradiation time of about 10 h, a species with only one carbonyl absorption at 1955 \(\text{cm}^{-1}\) is observed. We assign this absorption to the unsaturated monocarbonyl complex \((C_5H_5)Co(CO)\) (III). Addition of triphenylphosphine to these solutions converts this species quantitatively to the known carbonylphosphine complex \((C_5H_5)Co(CO)[P(C_6H_5)_3]\) with \(\nu(\text{CO})\) 1905 \(\text{cm}^{-1}\) [8]. Due to its instability, the monocarbonyl III cannot be isolated. At room temperature, its IR spectrum disappears, and, a new IR band in the \(\mu\)-CO region appears at 1785 \(\text{cm}^{-1}\). In more concentrated solutions, this conversion to some bridged carbonyl complex takes place even at \(-78^\circ\text{C}\). A dinuclear dicarbonyl compound can be obtained in low yield by irradiation of I at \(-78^\circ\text{C}\), removal of solvent at room temperature and subsequent sublimation at 40 \(^\circ\text{C}\) and 10\(^{-3}\) Torr. This green material, which shows a single CO absorption at 1798 \(\text{cm}^{-1}\) in petroleum ether, is identified, on the basis of its IR, NMR and mass spectra, as the dinuclear dicarbonyl IV, \((C_5H_5)_2Co_2(\mu-Co)\) *.

In addition, other species are observed in irradiated solutions, especially at higher concentrations of starting material I in particular, the trinuclear carbonyl II, identified by its IR absorptions at 1830, 1770 and 1670 \(\text{cm}^{-1}\). This compound is isolated, following earlier accounts [3], when the residue obtained after irradiation, removal of solvent at room temperature and sublimation, is extracted with diethyl ether.

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* The anion \((C_5H_5)Co_2(\mu-CO)_2^\text{-}\) has recently been found to form when \((C_5H_5)Co(CO)_2\) is reduced e.g. with sodium amalgam [9]; IR and NMR absorptions assignable to IV have also been observed by Vollhardt et al. [7].
Recently, Vollhardt et al. have investigated related reaction systems and observed the intermediate formation of a dinuclear tricarbonyl V, \((\text{C}_5\text{H}_5)_2\text{Co}_2(\text{CO})_3\)[7]. This compound is characterized by IR absorptions at 1965 and 1814 cm\(^{-1}\) in Nujol. The intermediacy of this species V in the consecutive photo- and dark reaction steps leading to compounds IV and II is clearly observable in high-boiling petroleum ether as a solvent. Here, irradiation at \(-78^\circ\text{C}\) produces this species V with an absorption at 1810 cm\(^{-1}\), observable at room temperature immediately after removal from the low-temperature reaction mixture. In addition, the terminal CO band at 2025 cm\(^{-1}\) decreases relative to that at 1970 cm\(^{-1}\). The residual absorption at 1970 cm\(^{-1}\) must be due, at least in part, to the terminal CO-band of the tricarbonyl V; we cannot say whether some monocarbonyl III contributes to this absorption.

A subsequent dark reaction then occurs at room temperature: the band at 1810 cm\(^{-1}\) associated with the dinuclear tricarbonyl V, decreases to about half its initial intensity with an initial half-life of about 1 h. At the same rate, a new band grows in at 1798 cm\(^{-1}\), indicating the formation of dinuclear dicarbonyl IV. This change is accompanied by an increase of the terminal CO absorption at 2025 cm\(^{-1}\), i.e. by regeneration of starting material I. When the two bands at 1810 and 1798 cm\(^{-1}\), associated with the dinuclear tri- and di-carbonyl complexes, respectively, have reached comparable intensities, both remain fairly stationary, decaying synchronously with a half-life of several hours, while a band at 1762 cm\(^{-1}\), probably due to trimer II, continues to grow. Essentially in accord with Vollhardt et al. [7], we interpret these observations according to Scheme 1. The primary photoproduct III, \((\text{C}_5\text{H}_5)_2\text{Co}(\text{CO})\), associates with excess

\[ \text{SCHEME 1} \]

Generation and interconversion pathways for coordinatively unsaturated cyclopentadienylcobalt(0) carbonyl intermediates. Photochemical reaction steps are indicated by wavy arrows, thermal reactions by straight arrows: numbers in parentheses are carbonyl frequencies in cm\(^{-1}\).

I in a concentration-dependent equilibrium to form the dinuclear tricarbonyl V. This species can then be converted, presumably via the monocarbonyl III in equilibrium with it and I, to the dinuclear dicarbonyl IV with regeneration of starting material I. Probably V is also convertible to IV by direct photolysis under loss of CO. The coordinatively unsaturated dicarbonyl IV can then associate with an additional monocarbonyl unit to form the trinuclear particle II, which appears
to be the final product under our conditions.

When solutions containing a mixture of the photolysis products III, IV, V and II are exposed to carbon monoxide, complete conversion to the initial dicarbonyl I is observed. Addition of triphenylphosphine to IV, V or II leads to formation of the known mixed carbonylphosphine (C₅H₅)Co(CO)[P(C₅H₅)₃] [8]. These reactions indicate the possibility of a symmetrical cleavage of the di- and tri-nuclear species by ligand attack.

The dinuclear dicarbonyl IV appears to undergo unsymmetrical cleavage as well. In its mass spectrum the ions (C₅H₅)Co(CO)₃⁺ and (C₅H₅)Co⁺ are observed with higher yields than the symmetrical cleavage product (C₅H₅)Co(CO)⁺. This tendency toward unsymmetrical cleavage is also manifested in substitution reactions in solution. Thus when a solution of IV in toluene is treated with a diolefin, the corresponding diolefin complex [10] (C₅H₅)Co(diene), is formed together with an approximately equal yield of dicarbonyl I. Apparently, the difunctionality of the attacking ligand directs this substitution toward unsymmetrical cleavage of the dimer, in contrast to the symmetrical reaction mode preferred for cleavage by the monofunctional phosphine ligand.

2. Reduction of (C₅H₅)Co(CO)₁₂

In order to further establish the intermediacy of the reactive species we have investigated additional modes of obtaining these intermediates. From related studies on reactive titanium and molybdenum compounds [1,11], it is known that reduction of dihalide compounds by a strongly reducing metal can lead to reactive intermediates. Reduction of the cobalt(III) diiodocarbonyl, (C₅H₅)Co-(CO)₁₂ [8], in a toluene solution at room temperature with sodium amalgam yielded a mixture of the dinuclear dicarbonyl IV, and the trinuclear carbonyl II. This establishes an independent, chemical entry into the series of intermediates represented in Scheme I.

Discussion

The generation of coordinatively unsaturated monocarbonyl III and its interconversion reactions involving the binuclear adducts IV and V are undoubtedly the primary steps in a number of photochemical and thermal reactions between the dicarbonyl compound I and other, reactive ligand molecules. Applications of this basic reaction pattern toward reactions systems involving various acetylene derivatives will be discussed in a subsequent paper [12].

Concerning the stability of the binuclear dicarbonyl IV, some comparisons with related compounds are warranted. Among the comparable cyclopentadienyl-rhodium complexes, the only known stable dinuclear carbonyl derivative has the composition (C₅H₅)₂Rh₂(CO)₃ [13], whereas the dinuclear cobalt carbonyls (C₅H₅)₂Co₂(CO)₂ (IV) and (C₅H₅)₂Co₂(CO)₃ (V) appear to exist in equilibrium with each other and with I. This difference might be related to increased steric crowding around the cobalt atom as compared to rhodium, which would favor smaller coordination numbers for the (C₅H₅)Co unit.

Formally, a cobalt–cobalt double bond has to be postulated for the dinuclear dicarbonyl IV in order to arrive at an 18-electron valence shell configuration. For the dinuclear iron–nitrosyl compound (C₅H₅)₂Fe₂(NO)₂ a comparatively
short Fe—Fe distance of 2.33 Å in a coplanar Fe₂(µ-NO)₂ ring has been observed by Calderon et al. [14]; this was interpreted as arising from a 4-electron bond between the two metal centers. The occurrence of only one stretching mode in the IR spectrum of the isoelectronic dicarbonyl (C₅H₅)₂Co₂(µ-CO)₂ indicates a similar, coplanar geometry of the central Co₂(µ-CO)₂ ring, and hence would be compatible with an analogous bonding situation in these two isoelectronic compounds.

Experimental

General techniques and chemicals

All manipulations of air-sensitive compounds were carried out under an inert gas using either Schlenk or high-vacuum manifold techniques. The preparations of (C₅H₅)Co(CO)₂, (C₅H₅)Co(CO)I₂ and (C₅H₅)₂Co₂I₄ were based on the procedures described in ref. 3 and 8. All solvents used for this work were doubly distilled and dried with butyllithium or methyllithium.

Instruments

IR spectra were taken on a Shimadzu 400 spectrophotometer, proton magnetic resonance measurements with a JEOL-MH-100 high resolution spectrometer, and mass spectra on a Varian CH 7 mass spectrometer.

1. Photochemical preparation of (C₅H₅)₂Co₂(µ-CO)₂ (IV) and (C₅H₅)₃Co₃(CO)₃ (II)

A solution of 0.7 ml (5.44 mmol) of (C₅H₅)Co(CO)₂ in 200 ml of toluene was irradiated at dry ice temperature with a 125 W mercury lamp. Nitrogen was slowly bubbled through the solution during photolysis and IR spectra of the irradiated solution were recorded occasionally. After 15 h, the irradiation was discontinued. After removal of solvent, a green solid was sublimed at 40°C under high vacuum from the residue. This material was identified as (C₅H₅)₂Co₂(µ-CO)₂ (IV): Its IR spectrum in Nujol exhibits a strong, broad peak at 1765 cm⁻¹. Its proton NMR spectrum in toluene-d₈ shows a sharp singlet at 7 5.61. Its molecular weight was determined as 304 by mass spectrometry (main fragment ions discussed above).

The crude black sublimation residue was extracted with diethyl ether in a Soxhlet extractor. This process was completed in half a day. The dark, partly crystalline extracts were then cooled in a -78°C bath, the solvent syringed off, and the black crystals dried in vacuum. The identity of the crystals as (C₅H₅)₃-Co₃(CO)₃ (II) was confirmed by comparison of the IR spectrum with that given by King [3]. Yield: (C₅H₅)₂Co₂(µ-CO)₂ (V), 0.16 mmol (6.0%); (C₅H₅)₃Co₃(CO)₃ (II), 1.01 mmol (55.6%).

2. Reduction of (C₅H₅)Co(CO)I₂, by sodium amalgam

A toluene solution (50 ml) of (C₅H₅)Co(CO)I₂, (1.60 g, 3.94 mmol) was stirred with 9 mmol of sodium amalgam. After 15 min, the purple solution changed to brown. Stirring was continued overnight. Filtration of the mixture gave a dark brown solution. The solvent and (C₅H₅)Co(CO)₂ produced from the reaction
were removed by vacuum distillation. A green solid was sublimed from the residue and identified as \((C_5H_5)_2Co_2(\mu-CO)_2\) (IV) by comparing its IR and mass spectra to that obtained in section 1.

The black residue was extracted with diethyl ether in a Soxhlet extractor. The black solid obtained was identified as \((C_5H_5)_3Co_2(CO)_3\) (II), by comparison of its IR spectrum with that given by King [3]. Yield: \((C_5H_5)_2Co(CO)_2\) (I) 0.67 mmol (17.0%); \((C_5H_5)_2Co_2(\mu-CO)_2\) (IV), 0.13 mmol (6.6%); \((C_5H_5)_3Co_3(CO)_3\) (II), 0.75 mmol (56.7%).

3. IR observations of intermediates in solution

Photolysis of the dicarbonyl I was carried out at \(-30^\circ C\) in toluene or petroleum ether solution in a low-temperature IR cell by irradiation with a focused UV beam derived from a 250 W high-pressure Xenon lamp (Hanovia). IR spectra of the irradiated solutions were recorded at various intervals at \(-30^\circ C\). Subsequently the temperature was raised to \(25^\circ C\), where the decay of the IR band at 1955 cm\(^{-1}\) assigned to the monocarbonyl III, and of the spectra of V and IV was followed over several hours. Similar observations were made when solutions irradiated at \(-78^\circ C\) in the synthetic irradiation vessel (section 1) were transferred to a normal solution IR cell, where their spectra were recorded at room temperature for 2–3 h.

Acknowledgements

This work has been supported by grants from the National Science Foundation (GP 33130X), from Deutsche Forschungsgemeinschaft and from fonds der Chemischen Industrie. We are grateful to Drs. Vollhardt, Bercaw and Bergman for making available their results on the formation of \((C_5H_5)_2Co_2(CO)_3\) in related reaction systems prior to publication.

References

12 W.S. Lee and H.H. Brintzinger, submitted for publication.