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Photoreactions of Group 6 metal carbonyls with ethene: syntheses of trans-(.eta.2ethene)2M(CO)4 (M = chromium, molybdenum, tungsten

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troscopically and chromatographically identical with a sample of 20 produced in the same way from bilobalide-derived 19. Synthetic diol 20 was converted to bilobalide 6-acetate (22) by a two-step process: (1) reaction with methyl oxalyl chloride (1.3 equiv) and diisopropylethylamine (2 equiv) in acetonitrile at 0 °C for 2 h to form 21 in 100% yield and (2) reaction in dilute toluene solution at reflux with a low concentration of tri-n-butyltin hydride (3 equiv of a 0.05 M solution in toluene containing 0.05 equiv of azobisisobutyronitrile added by syringe drive over 4 h) to give the (\pm)-bilobalide acetate 22, mp 270–274 °C (55%), along with recovered diol 20 (30%).¹⁴ Synthetic (\pm)-22 was identical with an authentic sample of bilobalide acetate1 by 500-MHz 1H NMR, infrared, mass spectral, and TLC comparison. Hydrolysis of synthetic (\pm) -22 with 3 N hydrochloric acid at reflux for 36 h afforded (±)-bilobalide 1 (70% yield), spectroscopically and chromatographically identical with an authentic sample.¹⁵ The synthesis of (±)-bilobalide described above solves one of the classical problems of organic synthesis by the use of novel chemistry and highly selective reactions. The one-step synthesis of 2 is both unprecedented and practical. In addition the conversions $7 \rightarrow 9$, $9 \rightarrow 12$, $12 \rightarrow 13$, $18 \rightarrow 19$, and $19 \rightarrow 1$ are noteworthy. The unusual tert-butylcarbinol unit of 1 greatly complicates the task of synthesis because the conventional methods of synthesis are inapplicable.16

Supplementary Material Available: Spectral and analytical data for 1-3, 5-7, 9a, 9b, 10-16, and 17-22 (3 pages). Ordering information is given on any current masthead page.

(14) See: Dolan, S. C.; MacMillan, J. J. Chem. Soc., Chem. Commun. 1985, 1588-1589. It is essential that the tin hydride be added slowly, otherwise the production of diol 20 increases relative to 22

(15) We are grateful to Dr. Pierre Braquet, Institute Henri Beaufour, Le Plessis Robinson, France, for a generous sample of naturally derived bilobalide. (16) This research was assisted financially by a grant from the National

Science Foundation.

Photoreactions of Group 6 Metal Carbonyls with Ethene: Syntheses of trans- $(\eta^2$ -Ethene)₂M(CO)₄ (M = Cr, Mo, W)

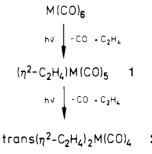
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Photolysis of Cr(CO)₆ in ethene-doped liquid xenon at -78 °C has been reported¹ to yield (η^2 -ethene)Cr(CO)₅ (1a), trans- $(\eta^2$ -ethene)₂Cr(CO)₄ (2a), and cis- $(\eta^2$ -ethene)₂Cr(CO)₄ (3a). All three compounds were said to be thermally unstable under these experimental conditions, their stability decreasing in the order 1a > 2a > 3a. More recently 2 1a was photochemically generated from Cr(CO)₆ and ethene in the gas phase. Subsequent pulsed-laser photolysis produced a transient species, detected by means of time-resolved IR spectroscopy, which decayed on the ms time scale and was assigned as $cis-(\eta^2-ethene)_2Cr(CO)_4$ (3a). The lability and cis structure of 3a was set in contrast with the stability and trans configuration of the known $(\eta^2$ -olefin)₂M(CO)₄ complexes of molybdenum and tungsten.^{3,4}

(2) Weiller, B. H.; Grant, E. R. J. Am. Chem. Soc. 1987, 109, 1252-1253.

Scheme Ia



 $^{a}M = Cr (1a, 2a), Mo (1b, 2b), W (1c, 2c).$

Table I. Spectroscopic and Analytical Data of 2a-c

	2a	2b	2c
$\overline{\nu}(CO)$, a cm ⁻¹	≃1986 (vw)	1997.5 (vw)	1991 (vw)
	1952.5 (vst)	1965 (vst)	1965 (vst)
$\delta (^{1}H)^{b}$	2.27	2.33	2.11
$\delta (^{13}C[^{1}H])^{b} C_{2}H_{4}$	48.25	41.89	31.26
CO	220.44	208.74	198.08
C found (calcd)	43.42 (43.65)	36.45 (36.38)	27.86 (27.30)
Н	3.62 (3.66)	3.16 (3.05)	2.09 (2.29)
metal	23.72 (23.62)	36.21 (36.33)	52.32 (52.23)

^a In *n*-hexane. ^b In toluene- d_8 .

This conflicts with our experience with olefin-substituted group 6 metal carbonyls^{3,5-7} and led us to suspect that the apparent disconformity should not be associated with the particular metal but rather originates from differences in the reaction conditions, i.e., gas-phase photochemistry in the chromium case² vs photoreactions of the molybdenum and tungsten compounds in solution.3,4,6,7 Recent experiments allow us to expand on this point and prompt us to report on the photoreaction of Cr(CO)6 with ethene in alkane solution, which provides convenient preparative access to the stable trans- $(\eta^2$ -ethene)₂Cr(CO)₄ (2a).

Irradiation of Cr(CO)₆ (10 mM) in ethene-saturated pentane solution (solidex immersion well apparatus, $\lambda \ge 280$ nm; Philips HPK 125-W mercury lamp) leads to sequential photosubstitution of CO for ethene; Scheme I, M = Cr. As monitored by infrared spectroscopy,8 initially formed 1a [$\bar{\nu}(CO)$ = 2078 (w), 1967 (st), 1961.5 (vst) cm⁻¹]⁹ is largely converted into trans- $(\eta^2$ ethene)₂Cr(CO)₄ (2a)⁹ upon extended irradiation. The first step is most conveniently performed at -20 °C, whereas the second step requires cooling to -50 °C. After filtration of the concentrated solution over silica gel and recrystallization from pentane, pure 2a is obtained as pale yellow crystals, mp 67-68 °C, 30% yield.

Relevant spectroscopic and analytical data on 2a are summarized in Table I. The observation of only one strong CO stretching vibrational band in the IR spectrum immediately rules out a $cis-(\eta^2-\text{ethene})_2\text{Cr}(\text{CO})_4$ structure [3a; four IR active $\nu(\text{CO})$ vibrations] in favor of the trans- $(\eta^2$ -ethene)₂Cr(CO)₄ geometry, most probably with orthogonal orientation of the two ethene ligands $[D_{2d}]$ symmetry; $\nu(CO)$: a_1 (IR inactive), b_2 (formally IR active, but extremely weak), e (IR active, very strong)]. This has been confirmed by the X-ray structure analysis of 2a.¹⁰ The ¹H and ¹³C{¹H} NMR spectra (Table I) exhibit only one and two signals, respectively, in accordance with structure 2a.

[†]Permanent address: Middle East Technical University, Ankara, Turkey. (1) Gregory, M. F.; Jackson, S. A.; Poliakoff, M.; Turner, J. J. J. Chem. Soc., Chem. Commun. 1986, 1175-1177.

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⁽⁹⁾ The IR data of 1a and 2a are in good agreement with those obtained in liquid xenon.

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 $trans-(n^2-Ethene)_2Cr(CO)_4$ (2a) is thermally stable not only in the solid state but also in the gas phase,11 in liquid xenon,12 and in alkane solution. By contrast, $(\eta^2$ -ethene)Cr(CO)₅ (1a) is only moderately stable. Monitoring a 1a/2a mixture in ethenesaturated alkane solution at ambient temperature by infrared spectroscopy we observe that 1a decomposes gradually with formation of Cr(CO)₆, whereas 2a survives for an extended period of time. Under CO atmosphere 1a has vanished within a few minutes, but 2a is still present after several hours, as long as oxygen is strictly excluded.

The photoreactions of $Mo(CO)_6$ and $W(CO)_6$ take essentially the same route as illustrated for $Cr(CO)_6$; Scheme I, M = Mo, W. The only difference is that the formation of 2c via 1c [M = W; $\bar{\nu}(CO) = 2085.5$ (w), 1974 (st), 1955.5 (vst) cm⁻¹] occurs at ambient temperature, and the synthesis of 2b via 1b M = Mo; $\bar{\nu}(CO) = 2086 \text{ (w)}, 1976 \text{ (st)}, \approx 1960 \text{ (vst) cm}^{-1}$ is most conveniently performed at -20 °C. The photochemical generation of these compounds has been reported previously together with their infrared data,4 but they were considered to be too unstable for isolation. However, we have obtained both of the two trans- $(\eta^2$ -ethene)₂M(CO)₄ complexes as colorless crystals: **2b**, mp 70-73 °C, 48% yield; 2c, mp 130-135 °C, 52% yield. The spectroscopic data (Table I) closely resemble those of 2a, and consequently the same structure can be assigned.

In all our experiments we see no evidence for the formation of any of the cis- $(\eta^2$ -ethene)₂M(CO)₄ complexes 3, but we have to consider the possibility that such a species could be involved as a short-lived intermediate.8 In fact, ab initio calculations13 on L-Mo(CO)₅ and low-temperature matrix experiments with $(\eta^2$ -olefin)W(CO)₅ in alkane glass¹⁴ or solid argon⁷ indicate initial photodissociation of cis CO. Nevertheless, we find it remarkable that in the gas-phase experiments² with $(\eta^2$ -ethene)Cr(CO)₅ and ethene the cis product 3a is the only observable $(\eta^2$ -ethene)₂Cr-(CO)₄ species, and no trace of the stable trans isomer 2a was observed. A plausible explanation 15 is that the formation of the latter involves $3a \rightarrow 2a$ photoisomerization. ¹⁶ At ambient temperature in the gas phase (or in alkane solution) this photoisomerization may be outstripped by the thermal decay of 3a. Cooling will provide sufficient stabilization of 3a, such that its photochemical rearrangement to 2a becomes feasible, as it happens in liquid xenon¹ or in alkane solution at -50 °C. In going from Cr to Mo and W the stability of type 3 complexes is expected to increase, and, consequently, cooling would become less essential in order to render the $3 \rightarrow 2$ photorearrangement feasible.

In a control experiment we used (η^4 -dimethylbutadiene)Cr-(CO)₄ (4)¹⁷ as a source of the cis-Cr(CO)₄ unit. ¹⁸ Upon treatment with ethene at ambient temperature in the dark 4 disappears gradually with formation of $(\eta^2$ -ethene)Cr(CO)₅ (1a) and Cr(CO)₆ as the only soluble products. Most probably these compounds originate from decomposition of labile cis- $(\eta^2$ -ethene)₂Cr(CO)₄ (3a). However, no trace of trans- $(\eta^2$ -ethene)₂Cr(CO)₄ (2a) could be detected, thus providing indirect evidence for the involvement of a second photochemical step, $3a \xrightarrow{h\nu} 2a$, in the synthesis of the

In conclusion we note that the coordination of two ethene ligands to the Cr(CO)₄ moiety nicely parallels the molybdenum and tungsten analogues: all three metals form stable trans- $(\eta^2$ -ethene)₂M(CO)₄ complexes 2. This and the distinctly lower stability of $(\eta^2$ -ethene)M(CO)₅ 1 and cis- $(\eta^2$ -ethene)₂M(CO)₄ 3 species is easily understood in terms of optimum metal $\rightarrow \pi^*$ (olefin) back donation. 6,7,19-21 Two olefins in trans orthogonal orientation do not compete for metal (d_{π}) electron density, owing to their single-faced π -acceptor character. By contrast a CO ligand (with its two orthogonal π -acceptor orbitals) situated trans to an olefin will strongly rival, thus weakening the metal-olefin bond.

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Novel Catalytic Chemistry of Iridium Polyhydride Complexes

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ordinated hydrogen.

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Polyhydride complexes of transition metals ([MH_xL_y], where L is a ligand such as a tertiary phosphine), constitute a distinctive class of compounds which have been widely studied in other contexts1 but whose catalytic chemistry has not thus far been extensively explored. Earlier studies^{2,3} have demonstrated the effectiveness of polyhydride complexes, e.g., [RuH2(PPh3)2- $\overline{(PPh_2C_6H_4)}$, $[RuH_3(PPh_3)_3]$, $[RuH_3(PPh_3)_2]$, and $[RuH_4]$ (PPh₃)₃], as catalysts or catalyst precursors for the hydrogenation of a variety of substrates including ketones, esters, and arenes which are not readily hydrogenated by most other homogeneous catalysts. Distinctive features of these systems appear to include the ability of such polyhydride complexes to serve as highly loaded "hydrogen reservoirs" and the accessibility of vacant coordination

Extending these studies we now have uncovered significantly different reactivity patterns for the iridium polyhydride complex $[IrH_5P_2]$ (P = P-i-Pr₃)⁴ and derivatives thereof, some features of which are unprecedented. We report here preliminary results of these studies.

sites through the facile reductive elimination or transfer of co-

[IrH₅P₂] reacts with the activated ester CF₃COOR (R = CH_2CF_3) in cyclo- $C_6D_{12}^5$ at 60 °C to yield $[IrH_2P_2(OR)]^6$ in

⁽¹¹⁾ In the gas-phase IR spectrum of 2a the intensity of the strong $\nu(CO)$ band (1967 cm⁻¹) remains constant over several hours.

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