has to be attributed to the quickness of the monoanion destruction rather than to the irreversibility of the heterogeneous charge transfer; in addition it **seems** likely that the electrode formation of the dianion takes place at potentials more negative than the solvent discharge itself. Also in this case a reoxidation peak, indicative of the decomposition of the cluster anion, is present in the reverse scan at about -0.3 **V,** well anodic to the cathodic peak.

Finally compounds VII and VIII reduce through similar cathodic processes involving one two-electron step. In both *casea* the relevant cathodic **peak** shows the typical features due to a charge transfer totally irreversible in character. *As* an example Figure 10 shows the response from VII.

Thus the electrochemical reduction process of $\mu_3 - \eta^2$ -||alkyne clusters **results** to be completely irreversible, **givmg** rise to a rapid declusterification. This is consistent with the EHMO picture of Schilling and Hoffman:⁴ for a 48electron species just a highly destabilized asymmetric MO (having metal-metal antibonding character due to ita parentage with the 2e_a orbital) is available, so the added electrons cause strong destabilization of the entire polyhedral frame.

Experimental Section

The $Fe_3(CO)_9(RC_2R)$ (I–IV) clusters were synthesized from Fe3(C0)1z and the appropriate alkyne in **1:l** molecular ratio in n-hexane at reflux under N_2 for 4 h (MeC₂Me and EtC₂Et) or 6 h (PhCzMe and PhCPh). The separations of **the** reaction **mixture** were carried out on TLC plates $(SiO₂,$ eluant petroleum etherdiethyl ether, 9:1 v/v). The preparation of ¹³CO-enriched samples of $Fe₃(CO)₉(RC₂R)$ complexes was performed by a similar procedure using Fes(CO)12, **-20%** enriched in I3CO, **as** the starting material.

Compounds V-VIII were synthesized according to the published procedures.^{9,30,31} After crystallization in n-heptane at 0^o C, purity of all tested complexes was checked by **IR** and 'H and 13C NMR spectroscopy.

The IR spectra were recorded on a Perkin-Elmer **580** B instrument and the lH and 13C NMR on a JEOL **GX-270-89** machine (for variable-temperature **'H** NMR experiments a JEOL C-60-HL was also employed).

The electrochemical apparatus **has** been described elsewhere.% Potential values refer to a saturated aqueous calomel electrode (SCE). The temperature was controlled at 20 ± 0.1 °C. The exhaustive macroelectrolysis of $Fe_3(CO)_9(EtC_2Et)$ (I) was brought about at the controlled potential of **-1.0** V by using a platinum gauze working macroelectrode. **As** the two-electron reduction proceeds, the solution turns from brown into dark-red. All the manipulations of the electrogenerated $[Fe_3(CO)_9(EtC_2Et)]^{2-}$ dianion were carried out in a glovebox purged with dry dinitrogen. The solventa used for electrochemistry, dichloromethane and acetone, were from Burdick and Jackson, were "distilled in glass", and were used without purification. Tetrabutylammonium perchlorate *(TBAP,* **Fluka)** and tetraethylammonium perchlorate (TEAP, Carlo Erba) supporting electrolytes were dried in a vacuum oven and used without further purification. Lithium perchlorate and sodium perchlorate supporting electrolytes were prepared by neutralizing perchloric acid with the corresponding alkali carbonate, twice crystallizing from methanol, and finally drying at 110 °C. Bis(η^5 -cyclopentadienyl)iron(II) (ferrocene, Alfa Products) was used **as** purchased. Extra pure dinitrogen was employed to remove oxygen from tested solutions.

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Registry No. I, **69402-19-3;** 11, **12171-93-6;** 111, **101315-98-4;** IVa, **101315-99-5;** IVb, **101316-00-1;** V, **78109-40-7;** VI, **101316-01-2;** VII, 80303-37-3; VIII, 101316-02-3; $[Fe_3(CO)_9(EtC_2Et)][NBu_4]_2$, **101316-04-5; [Fe,(CO)g(EtCzEt)][NEt4]z, 101316-05-6;** [Fe3- $(CO)_{9}(EtC_{2}Et)$]Li₂, 101400-35-5; [Fe₃(CO)₉(EtC₂Et)]Na₂, **101316-06-7;** Fe, **7439-89-6; Os, 7440-04-2.**

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Methyl-Ethylene and Methyl-Carbonyl Complexes of Platinum(I I) and Plathum(IV)

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The methyl(ethylene)platinum(II) complex $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)_2]$ and methylcarbonylplatinum(II) complex $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2)]$ have been prepared and characterized. Reactions of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)]$ with alkenes or alkynes (un) generally give the products of substitution $[Pt_2(\mu\text{-Cl})_2\text{Me}_2(\text{un})_2]$, but $CF_3\text{C}=\text{CCF}_3$ gives insertion into the PtMe bond. Reaction of $[Pt_2Me_4(\mu\text{-}SMe_2)_2]$ with ethylene yields the dimethyl-(ethylene)platinum(II) complex cis-[PtMe₂(SMe₂)(C₂H₄)] and with carbon monoxide cis-[PtMe₂(SMe₂)(CO] is formed. Oxidative addition of iodomethane to cis-[PtMe₂(CO)(SMe₂)] produces a rare example of a platinum(IV) carbonyl complex fac -[PtIMe₃(SMe₂)(CO)].

Introduction

This paper reports the synthesis, characterization, and properties of several compounds containing cis-methyl- (ethylene)platinum(II) or **cis-methylcarbonylplatinum(I1)** units. Such compounds are unusual, although some interesting examples such **as cis-[PtMe(CzH4)(diars)]+,** diars $= 1,2-(Me₂As)₂C₆H₄$, and cis-[PtMe₂(CO)₂] are known¹⁻³

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of cis-[PtMe₂(SMe₂)₂] with Zeise's dimer $[Pt_2(\mu\text{-}Cl)_2Cl_2$ - $(C_2H_4)_2$ gave the complex $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)_2]$ by methyl for chloro exchange between platinum centers.^{7,8} **A** preliminary account has been published.8

Results and Discussion

Synthesis and Characterization of $[Pt_2(\mu\text{-}Cl)_2\text{-Me}_2]$ $(C_2H_4)_2$. The new methyl(ethylene)platinum complex was prepared by methylation of Zeise's dimer $[Pt_2(\mu\text{-Cl})_2Cl_2$ - $(C_2H_4)_2$ (1) with $[Pt_2Me_4(\mu-SMe_2)_2]^7$ in the presence of excess ethylene (Scheme I). The desired product **2** was separated from the second product $[Pt_2(\mu\text{-Cl})_2\text{Me}_2(\text{SMe}_2)_2]$ by chromatography on a Florisil column, when only **2** passed through the column. We have shown previously that $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(S\text{Me}_2)_2]$ is thermally unstable,⁷ and it is decomposed on the column. However, this complex is sufficiently stable to be positively characterized by its 'H NMR spectrum in the reaction mixture. Yields of **2** of 90% could be obtained from this reaction.

Complex **2** could **also** be prepared by reaction of Zeise's dimer with cis -[PtMe₂(SMe₂)₂], when the byproduct was *trans*-[PtClMe(SMe_2)₂], and with SnMe_4 , when the byproduct was SnClMe₃. However, isolated yields were much lower due partly to decomposition reactions (e.g., with SnMe₄) or to problems with purification of 2 (e.g., with $[PHMe₂(SMe₂)₂]$). Therefore the methylating agent Therefore the methylating agent $[Pt₂Me₄(\mu-SMe₂)₂]$, although expensive, was preferred in the synthesis **of 2.** More powerful methylating agents such **as** MeLi or MeMgCl led to decomposition of **Zeise's** dimer, and this is probably why the complex **2** has not been successfully prepared earlier.

Complex **2** was characterized by elemental analysis and by ita 'H and '3c NMR **spectra."** Of particular significance is the observation of satellites around the C_2H_4 resonance in the 'H or 13C NMR spectra at room temperature in $CD₂Cl₂$ solution showing that rapid reversible dissociation **of C2H4** does not occur. **Above** *-65* **"C,** the ethylene resonance in the lH NMR spectrum of **2** appears **as** a singlet with ¹⁹⁵Pt coupling, but below -65 °C this resonance splits into an [AA'BB'] pattern with ¹⁹⁵Pt coupling. This effect has been observed previously in $[PtCl(aceac)(C₂H₄)]$ and related complexes⁹ and is due to the freezing out of rotation about the Pt- C_2H_4 bond.⁴

In contrast Zeise's dimer gives a broad singlet without $^{195}\text{Pt satellites at temperatures as low as -90 °C in $\text{CD}_2\text{Cl}_2$$ solution, indicating a very rapid exchange process. We suggest that the strong σ -donor methyl group leads to stronger $Pt-C_2H_4$ back-bonding in 2 and hence the ethylene is less prone to dissociation or rotation compared to Zeise's dimer.¹⁰

Reactions of $[Pt_2(\mu\text{-Cl})_2\text{Me}_2(C_2H_4)_2]$. (a) With Pyr**idine and Dimethyl Sulfide. A** common method for preparing monomeric olefin complexes of platinum(I1) involves reaction of a halogen-bridged dimer $[Pt_2(\mu-X)_2X_2(\text{olefin})_2]$ with a neutral ligand.¹¹ When 2 was X_2X_2 (olefin)₂] with a neutral ligand.¹¹ treated with a stoichiometric quantity of pyridine, two isomers of $[PtClMe(C₂H₄)(py)]$ (4a and 5a) were easily isolated, thus confirming the dimeric structure of **2.** $[PtCl₂(C₂H₄)(py)]$ exists only as the trans isomer,¹² but the presence of two of the three possible isomers of $[PtClMe(C₂H₄)(py)]$ was evident from two distinct sets of Pt-Me, Pt-C₂H₄, and pyridine H α resonances in the ¹H NMR **spectrum.** Isomers **4a** and **5a** were present in a ratio of 3:1, and the structures were deduced from the magnitudes of $J(PtH)$ coupling constants.^{1-4,12} Isomer **4a** gave signals in the ¹H NMR spectrum due to the C_2H_4 and pyridine *Ha* protons which had satellites due to coupling to ¹⁹⁵Pt, but no such satellites were observed for the corresponding resonances of isomer **5a.** This is presumably due to labilization of pyridine by the high trans influence of the methyl group which leads to ligand exchange^{4,12} but which does not lead to equilibration with **4a.**

A similar reaction of **2** with Me2S gave **4b** and **5b** as major products, but a disproportionation occurred on evaporation of the reaction mixture at room temperature to give **2** and trans-[PtClMe(SMe,),]. The mixture of **4b** and **5b** could be isolated by evaporation of the solvent from the reaction mixture at **-78** "C, but the solid product was thermally unstable at room temperature.

(b) With Alkenes and Alkynes. In $CD₂Cl₂$ solution at 35 "C in the presence **of** a large excess of ethylene, complex **2** apparently remains dimeric and undergoes rapid exchange of free and coordinated ethylene, as is evident from the unchanged MePt resonance in the 'H NMR spectrum and a broad (C_2H_4) Pt resonance (shifted upfield from free ethylene) possessing no ¹⁹⁵Pt satellites. At -60 "C there are two distinct MePt resonances; one can be attributed to complex **2** while the other, having a lower coupling constant, $^{2}J(\text{PtH}) = 68$ Hz, is due to trans- $[PtClMe(C₂H₄)₂]$ (3). It is evident that the monomeric species exists in equilibrium with complex **2** at low temperatures. The rapid exchange of free and coordinated ethylene in **2** at room temperature presumably occurs by rapid reversible formation of **3,** though the NMR parameters show that very little **3** is present at room temperature. Chatt has prepared the analogous complex $[PtCl_2(C_2H_4)_2]$ by reaction of Zeise's dimer with ethylene and other dichlorobis(olefin)platinum(II) monomeric complexes have also been prepared.¹³

Reaction of **2** with trans-stilbene or dimethyl fumarate gave the corresponding alkene complexes **6a or 6b** by displacement of ethylene. These complexes were isolated **as** yellow-orange solids, but they decomposed in 1-2 days at room temperature. In solution the 'H NMR spectra contained broad singlets due to the alkene protons at room

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Figure 1. *H NMR spectra **(100** MHz) of a **1:l** mixture of complexes 1 and 2 in CD₂Cl₂, to give an equilibrium mixture with 8: **(a)** at **-20** OC, in the fast-exchange regime; **(b)** at -80 OC, in the slow-exchange regime and with slow rotation about the $Pt-C_2H_4$ bonds for **2** and **8.** The peak marked asterisk **is** due to CHDC12.

temperature, indicating a rapid exchange process, but at -90 °C coupling to ¹⁹⁵Pt was observed and the signals appeared as an [AB] quartet. Thus at low temperature alkene exchange and rotation about the Pt-alkene bond can be frozen out.

The diphenylacetylene complex **7** was easily prepared by reaction of **2** with diphenylacetylene, but similar reactions with but-2-yne or phenylacetylene gave polymeric materials and it was not possible to identify the platinum-containing products. A similar reaction of Zeise's dimer with diphenylacetylene led to rapid reduction to metallic platinum,¹⁴ and again it seems that the presence of the σ -donor methyl group enhances the stability of 7 by increasing the platinum-alkyne d_{τ} -p, back-bonding.

The activated alkyne $CF_3C=CCF_3$ reacted rapidly with **2,** and, after treatment with pyridine, complex **9** was obtained. Considerable polymer was also formed. The cis insertion is proved by the magnitude of the coupling *6J-* $(FF) = 15$ Hz, which is typical of this stereochemistry and much larger than for the trans isomer.¹⁵

In general, substitution of ethylene in **2** for other alkenes or alkynes is very easy and presumably occurs by way of intermediates analogous to 3. This ease of substitution is also partly responsible for the rapid insertion of CF_{3} - $C=CCF₃$ into the Pt-Me bond¹⁶ and for the easy polymerization of some alkynes by **2.** Reaction of **2** with excess ethylene at 50 °C leads to formation of some low molecular weight polyethylene, but, since some decomposition of **2** occurs under these conditions, it is not clear that **2** is the active catalyst for this process.

(c) With Carbon **Monoxide.** Carbon monoxide displaced ethylene from 2 to give the methylcarbonylplatinum derivative **10,** which could be isolated as a colorless solid but which decomposed at room temperature in about 0.5 day. Complex **10** was characterized by a carbonyl stretch, $\nu(\text{CO}) = 2040 \text{ cm}^{-1}$ (CH₂Cl₂ solution), in the IR, by a

methylplatinum resonance in the 'H NMR at **6** 1.24 **(2J-** $(PtH) = 76 Hz$ in CD_2Cl_2 , and by the mass spectrum, which gave a parent ion at m/e 546 with the expected isotope pattern for a Pt_2Cl_2 derivative. The analogous complex $[Pt_2(\mu\text{-Cl})_2Cl_2(CO)_2]^{17}$ has $\nu(CO) = 2146$ cm⁻¹, indicating weaker back-bonding than in **10 as** expected from the earlier discussion. There are very few methylcarbonylplatinum(II) complexes, and **10** appears to be the first binuclear derivative. $5,6$

(d) With Zeise's Dimer. Reaction of equimolar **amounts** of complex **2** and Zeise's dimer **1** leads to rapid equilibration with the unsymmetrical dimer 8. At temperatures above -50 "C this equilibration was rapid on the NMR time scale, as illustrated by Figure la. The spectrum shows methylplatinum, (ethy1ene)platinum cis to Me, and (ethy1ene)platinum cis to C1 signals with the expected intensity ratios. This spectrum is consistent either with **the** pure complex 8 or with a rapidly equilibrating mixture of 8, **1,** and **2,** occurring by rapid exchange of PtClMe- (C_2H_4) and $PtCl_2(C_2H_4)$ fragments. That the latter interpretation is correct is shown by the spectrum at *-80* "C shown in Figure lb. Separate resonances are clearly seen for the methylplatinum groups of **2** and 8. The ethylene resonances are less clearly resolved because rotation about Pt-C2H4 bonds is **also** frozen out for complexes **2** and 8. However, the singlet due to 1 is resolved from the $[A_2B_2]$ multiplet of the $Pt(C_2H_4)Cl$ resonance of 8 under these conditions. Integration indicates that **1,** 8, and **2** are present in approximately the expected statistical ratio of 1:2:1 at -80 "C.

Synthesis of cis -[PtMe₂(C₂H₄)(SMe₂)] and *cis-* $[PtMe₂(CO)(SMe₂)].$ Reaction of cis- $[Pt₂Me₄(\mu-SMe₂)₂]$ (11) with ethylene in CD_2Cl_2 solution gave an equilibrium mixture of the reagents and cis- $[PtMe₂(C₂H₄)(SMe₂)]$ (12; Scheme II), but only 11 was recovered on evaporation of such solutions.

The cis stereochemistry is shown by the observation of two MePt resonance in the 'H NMR spectrum of **12, as** well as by the very low coupling constants, $\frac{3J(PtSMe_2)}{2}$ = 21 Hz and ² $J(Pt\ddot{C}_2H_4) = 37$ Hz for the methylsulfur and ethylene resonances. The equilibrium constant changes with temperature **as** is clearly shown by Figure 2. Although exchange between free and coordinated ethylene was slow on the NMR time scale at **35** "C, rotation about the Pt- C_2H_4 bond appeared to be fast even at -90 $^{\circ}$ C since a singlet for the coordinated ethylene of *12* was observed at all temperatures.

No products analogous to **12** could be detected from reactions of diphenylacetylene or trans-stilbene with **11** even at low temperatures. Clearly, there is a fine balance as to whether the dimer **11** or the monomeric complexes analogous to **12** will be preferred in this system.

Carbon monoxide is a stronger **ligand,** and reaction with **¹¹**gave both **13 and ¹⁴**(Scheme 11). Complex **14** has been

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Figure **2.** 'H NMR spectra (100 MHz) of a mixture of $[Pt_2Me_4(\mu\text{-}SMe_2)_2]$ and C_2H_4 in equilibrium with cis- $[PtMe_2 (\text{SMe}_2)(\text{C}_2\text{H}_4):$ (a) at 35 °C, mostly $[\text{Pt}_2\text{Me}_4(\mu\text{-SMe}_2)_2]$; (b) at -50 °C, mostly *cis*-[PtMe₂(SMe₂)(C₂H₄)].

prepared previously,^{2,3} and we find that 13 can be prepared in pure form by reaction of cis- $[PtMe₂(SMe₂)₂]$ with CO. Apparently the extra SMe_2 is sufficient to prevent further reaction of **13** with CO to give **14.**

A **Methylcarbonylplatinum(1V)** Complex. Oxidative addition of iodomethane to **13** gave the complex fuc- $[PtIME₃(CO)(SMe₂)]$ (15) along with byproducts fac- $[PtIME₃(SMe₂)₂]$ and $[(PtIME₃)₄].$ Complex 15 could also be prepared in solution by reaction of fac -[PtIMe₃(SMe₂)₂] with excess CO. It appears to be the first example of a **methylcarbonylplatinum(1V)** complex. Complex **15** is characterized by a carbonyl stretching frequency, $\nu(CO)$ $= 2090$ cm⁻¹ in benzene, and by the observation of three MePt and one $Me₂S$ resonances in the ratio 1:1:1:2, respectively. The methylplatinum group trans to CO had a lower value for ${}^{2}J(\text{PtCH}_3) = 64 \text{ Hz}$ than those trans to iodide or SMe₂, each of which gave ² $J(PtCH_3)$ = 70 Hz, due to the higher trans influence of CO. The methylsulfur resonance of **15** split **into** two resonances, due to freezing out of inversion at **sulfur,l*** at temperatures below -20 "C.

Conclusions

A number of new but very simple complexes containing $cis-PtMe(C₂H₄)$ and $cis-PtMe(CO)$ groups have been obtained. The ethylene complexes were only found with **platinum(II),** but carbonyl derivatives of both platinum(I1) and platinum(IV) were observed. Many of the complexes were thermally unstable, but they could be characterized unambiguously by low-temperature NMR studies. **Only** with the activated alkyne $CF_3C=CCF_3$ did insertion of the unsaturated group into the Pt-Me bond occur.

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Experimental Section

The ¹H, ¹⁹F, and ¹³C(H) NMR spectra were recorded on Varian XL100 and xL200 spectrometers. 'H and **'9c** chemical shifts were measured relative to Me4Si, and the 19F chemical shifts were measured relative to CFCl₃.

Infrared spectra were recorded by using NaCl solution cells on a Beckman 4250 spectrophotometer.

Gas chromatographic analyses were performed on a **Varian** 1400 gas chromatograph equipped with a porapak Q column.

Elemental analyses were performed by Guelph Chemical Laboratories, Ltd., Guelph, Ontario.

Complexes $[Pt_2Me_4(\mu\text{-sMe}_2)_2]$ and $[Pt_2(\mu\text{-Cl})_2Cl_2(C_2H_4)_2]$ were prepared by modifications of known procedures.^{7,1}

Synthesis of $[\mathbf{Pt}_2(\mu\text{-}\mathrm{Cl})_2\mathbf{Me}_2(\mathrm{C}_2\mathbf{H}_4)_2]$. $[\mathrm{Pt}_2(\mu\text{-}\mathrm{Cl})_2\mathrm{Cl}_2(\mathrm{C}_2\mathrm{H}_4)_2]$ $(0.3502 \text{ g}, 0.5955 \text{ mmol})$ was suspended in CH_2Cl_2 (40 mL) saturated with C₂H₄ at 0 °C. [Pt₂Me₄(μ -SMe₂)₂] (0.3422 g, 0.5955 mmol) in CH_2Cl_2 (20 mL) was added to the stirred solution. The solution was warmed slowly to room temperature and allowed to react for 0.5 h. The solvent volume was reduced to \sim 5 mL, and the mixture was added to a 12-in. column packed with Florisil and eluted with $CH₂Cl₂$ (50 mL). The solvent was removed, yielding a white solid identified as $[Pt_2(\mu\text{-Cl})_2\text{Me}_2(C_2H_4)_2]$ (0.2837 g, 91%); mp 77-90 °C dec. Anal. Calcd for $[Pt_2(\mu$ -Cl)₂Me₂- $(C_2H_4)_2$: C, 13.17; H, 2.58. Found: C, 13.30; H, 2.57. NMR in CD_2Cl_2 , 35 °C: ¹H, δ 0.52 [s, ²J(PtH) = 81 Hz, MePt], 3.73 [s, MePt], 63.38 [s, 1 J(PtC) = 254 Hz, $(C_{2}H_{4})$ Pt]. At low-temperatures (below -65 °C) the (C_2H_4) Pt resonances in the ¹H NMR resolved into an $[AB]_2$ multiplet (see ref 8). ${}^{2}J(\dot{P}tH) = 79$ Hz, $(C_{2}H_{4})Pt$]; ¹³C, δ -8.63 [s, ¹ $J(PtC) = 682$ Hz,

 $(0.1156 \text{ g}, 0.2112 \text{ mmol})$ was dissolved in CH_2Cl_2 (10 mL), and pyridine $(0.0334 \text{ g}, 0.4224 \text{ mmol}, 34.0 \mu\text{L})$ was added as a CH_2Cl_2 solution (5 **mL)** at 0 "C. The solution immediately turned clear yellow, and the product identified as $[PtClMe(C₂H₄)(py)]$ was isolated **as** a yellow solid (0.1402,98%) by concentration and then precipitation using n-pentane; mp 64-72 °C dec. Anal. Calcd for [PtClMe(C_2H_4)(py)]: C, 27.24; H, 3.43. Found: C, 27.73; H, 3.53. ¹H NMR in \widehat{CD}_2Cl_2 : **4a**, δ 0.66 [s, ²J(PtH) = 72 Hz, MePt], 3.53 [s, ²J(PtH) = 72 Hz, (C_2H_4) Pt]; 5a, δ 0.41 [s, ²J(PtH) = 76 Hz, MePt], 4.12 [s, ²J(PtH) not resolved, (C_2H_4) Pt]. **Preparation of [PtClMe(C₂H₄)(py)].** [Pt₂(μ -Cl)₂Me₂(C₂H₄)₂]

Preparation of $[PtCIME(SMe₂)(C₂H₄)]$. $[Pt₂(\mu$ -Cl)₂Me₂and 2 equiv of SMe₂ (0.02841 g, 0.4572 mmol, 33.6 μL) was added as a $CH₂Cl₂$ solution (5 mL) at 0 °C. The solution immediately turned clear yellow and was cooled rapidly to -78° C. Removal of the solvent at -78 "C yields the thermally unstable beige solid identified **as** [PtClMe(SMez)(CzH4)] (0.1443 g, 94%); mp 30-39 °C dec. Anal. Calcd for $[\text{PtC1Me(SMe₂)(C₂H₄)]:$ C, 17.88; H, 3.90. Found: C, 17.66; H, 4.17. ¹H NMR in CD_2Cl_2 , -80 °C: 4b, δ 0.53 [s, ²J(PtH) = 70 Hz, MePt], 3.62 [s, ²J(PtH) = 72 Hz, (C_2H_4) Pt], 2.43 [s, ³J(PtH) = 16 Hz, (Me₂S)Pt]; 5b, δ 0.74 [s, ${}^{2}J(\text{PtH}) = 74 \text{ Hz}, \text{MePt}, 3.73 \text{ [s, } {}^{2}J(\text{PtH}) = 72 \text{ Hz}, (\text{C}_{2}\text{H}_{4})\text{Pt},$ 2.53 [s, ${}^{3}J(\text{PtH}) = 22 \text{ Hz}$, $Me₂ \text{S}$)Pt]. $(C_2H_4)_2$] (0.1251 g, 0.2286 mmol) was dissolved in CH_2Cl_2 (10 mL),

Reaction of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2\text{H}_4)_2]$ with Ethylene. (A) A solution of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)_2]$ (0.0543 g, 0.099 21 mmol) in CD_2Cl_2 (1.5 mL) in an NMR tube was degassed on the vacuum line, a large excess of C_2H_4 was condensed into the tube, and the tube was sealed. Low-temperature 'H NMR showed conversion to trans-[PtClMe(C₂H₄)₂]. ¹H NMR in CD₂Cl₂, -60 °C: δ 0.32 [s, $^{2}J(\text{PtH}) = 68$ Hz, $MePt$], 3.62 [s, $^{2}J(\text{PtH})$ not resolved, $(C_{2}H_{4})Pt$].

(B) A solution of $[Pt_2(\mu\text{-Cl})_2\text{Me}_2(C_2H_4)_2]$ (0.1304 g, 0.2383 mmol) in dry benzene (100 mL) was stirred at 50 \degree C for 20 h under 10 atm of C_2H_4 in a pressure reactor. A clear colorless solution was obtained, but decomposition giving an insoluble black solid (possibly Pt and $PtCl₂$) occurred on partial evaporation of the solution to a volume of 25 **mL.** The solids were filtered off (0.0158 **g),** leaving a clear yellow solution which when evaporated to dryness yielded a yellow solid (0.1417 g). This product easily dissolved in CDCl₃ (2 mL), and the ¹H NMR showed the presence of both $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)_2]$ and low molecular weight poly-
ethylene: δ 1.25 [s, $(\text{CH}_2)_n$]; 0.84 [t, CH₃].

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Preparation of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(\text{PhC=CPh})_2]$ **.** $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(\text{PhC=CPh})_2]$. $\text{Cl}_2\text{Me}_2(\text{C}_2\text{H}_4)_{2}$] (0.0801 g, 0.1464 mmol) was dissolved in CH_2Cl_2 was added at 0 "C. The colorless solution immediately turned clear orange. The mixture was allowed to warm to room temperature, and evaporation of the solvent yielded an orange solid $\text{identity of } \frac{1}{2}$ **(Pt2)** μ -Cl)₂Me₂(PhC=CPh)₂] (0.1141 g, 92%); mp 81-90 °C dec. Anal. Calcd for $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(\text{PhC}=\text{CPh})_2]$: C, 42.51; H, 3.09. Found: C, 42.15; H, 2.95. ¹H NMR in CD₂Cl₂, 35 °C : δ 0.90 [s, ²J(PtH) = 86 Hz, MePt], 7.44 and 8.06 [m, Ph]. (5 mL) , and PhC=CPh $(0.0521 \text{ g}, 0.2928 \text{ mmol})$ in CH_2Cl_2 (5 mL)

Similarly were prepared complexes 6a, yield 94%, and 6b, yield **98%,** from trans-stilbene and dimethyl fumarate, respectively. ¹H NMR in CD₂Cl₂: **6a** at 35 °C, δ 0.42 [s, ²J(PtH) = 84 Hz, MePt], 5.96 [s, $\sqrt[2]{U(\text{PtH})}$ not resolved, $(=\text{CH})\text{Pt}$]; 6a at -90 °C, δ 0.30 [s, ²J(PtH) = 84 Hz, MePt], 5.69 [m, ²J(PtH) not resolved, $Pt(H_aC=)]$, 6.15 [m, ²J(PtH) not resolved, $Pt(H_bC=)]$; 6b at -90 $^{\circ}$ C, δ 0.45 [s, ²J(PtH) = 84 Hz, MePt], 4.91 [AB, ²J(PtH) not resolved, $Pt(HC=)$], 3.73 [s, OMe]. Anal. Calcd for $[Pt_2(\mu-$ **C1)zM~(Me02CCH=CHCOzMe)z] (6b):** C, **21.57;** H, **2.85. Found** C, **21.46;** H, **3.36. 6b** mp **100-103** "C.

 ${\bf trans\text{-}[PtCl(CCF₃)=CCCF₃)Me}(C₆H₅N)₂]$ (9). Excess $CF₃$ C $=$ CC $F₃$ was condensed into a tube containing a solution of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(C_2H_4)_2]$ (0.051 **g**) in CHCl₃ (5 mL), and the tube was sealed. The tube was allowed to warm to room temperature, and, after **0.5** h, the tube was opened and the insoluble polymer was removed by filtration to give a clear solution. A solution of pyridine (0.029 g) in CH_2Cl_2 (5 mL) was added, the solution was concentrated, and pentane was added to precipitate the product as a yellow solid (0.091 g) . NMR in CD₂Cl₂: ¹H, δ 1.24 [s, MeC]; 19 F, δ 40.5 [q, ³J(PtF) = 93 Hz, ⁵J(FF) = 15 Hz, (F₃CCPt)], 54.8 $[q, {}^{5}J(\text{FF}) = 15 \text{ Hz}, (F_3 \text{CCCPt})].$

Preparation of $[Pt_2(\mu\text{-}Cl)_2\text{Me}_2(\text{CO})_2]$ **.** Carbon monoxide was bubbled through a solution of $[Pt_2(\mu\text{-}CI)_2\text{Me}_2(C_2H_4)_2]$ (0.1055 g, **0.1928** mol) in CHzClz **(10 mL)** for **5** min at **0** "C. The solution was filtered, and the solvent was removed from the filtrate to yield the product **as** a white solid **(0.0981** g, **93%).** The complex was thermally unstable both **as** a solid and in solution above 0 "C. *NMR* in CD_2Cl_2 : δ 1.24 [s, ²J(PtH) = 76 Hz, MePt]. IR in CH₂Cl₂:

v(C0) **2040 cm-'.**

Preparation of cis-[PtMe₂(SMe₂)(CO)]. [Pt₂Me₄(μ -SMe₂)₂] **(0.1078** g, **0.1876** mmol) was dissolved in acetone **(10** mL), and $SMe₂$ (0.0233 g, 0.3752 mmol, 27.6 μ L) was added to generate cis -[PtMe₂(SMe₂)₂]. Carbon monoxide was bubbled through the solution for **5** min. Evaporation of the solvent gave the product as a beige oil which could not be solidified. **IR** in acetone: ν (CO) 2030 cm⁻¹. ¹H NMR in acetone- d_R at 35 °C: δ 0.22 [s, ²J(PtH) **²⁰³⁰**cm-'. 'H NMR in acetone-d6 at **35** "C: 6 **0.22 [s,** zJ(PtH) = **78** *Hz,* MePt **bans** to CO], **0.76 [s,** %T(PtH) = *89 Hz,* MePt trans to SMe₂, 2.68 [s, ²J(PtH) = 26 Hz, Me_2 S)Pt].

Preparation of fac-[PtIMe₃(SMe₂)(CO)] (15). (a) cis **-**[PtMez(SMez)(CO)] **(0.3752** mmol) was prepared **as** described above in acetone- d_6 (5 mL), and iodomethane (23.5 μ L) was added to the solution. The reaction was complete after **1** day giving fac -[PtIMe₃(SMe₂)(CO)] and byproducts fac -[PtIMe₃(SMe₂)₂] and $[(PtIME₃)₄]$. (b) fac -[$PtIME₃(SMe₂)₂$] (0.3752 mmol) was generated in acetone- d_6 (5 mL) by addition of iodomethane (23.5 μ L) to ~is-[PtMe~(SMe~)~] **(0.3752** mmol). CO was bubbled through the solution for 1 min. ¹H NMR in acetone- d_6 at -10 °C showed that MezS had been displaced by CO to generate fac-[PtIMe3- $(\text{SMe}_2)(CO)$]. ¹H NMR in acetone-d₆ for **15** at -10 °C: δ 1.31 [s, $^{2}J(\text{PtH}) = 70$ Hz, MePt trans to I or SMe₂], 1.46 [s, ²J(PtH) = 70 Hz, MePt trans to I or SMez], **1.20 [s,** zJ(PtH) = *64* Hz, MePt trans to CO], 2.72 [s, ³J(PtH) = 14.5 Hz , (Me₂S)Pt]. The same reaction can be performed in benzene, but the addition of indomethane to cis -[PtMe₂(SMe₂)₂] is slow in this solvent. IR in benzene for **15:** v(C0) **2090** cm-'.

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Effects of Ultrasonic Waves on the Generation and Reactivities of Some Metal Powders

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Ultrasound substantially accelerates the lithium reductions of a variety of metal halides to metal powders. Zinc powder produced in approximately **40** min at room temperature with ultrasound demonstrates reactivity in the Reformatsky reaction which rivals zinc produced in **4** h with potassium in refluxing tetrahydrofuran. Similarly copper and nickel powders produced in approximately 40 min with ultrasound promote Ullman coupling **as** effectively **as** copper and nickel powders produced in **>10** h in the presence of stirring. Results of the effect of iodide **salts** in the Ullman coupling of benzyl bromide are also presented.

Introduction

The field of reactive metal powders, as developed by Rieke et al., **has** a wide variety of synthetic applications.' It was found that the reduction of a metal halide with a group **1** element (i.e., Li, Na, **K)** in an ethereal solvent produces a metal slurry of exceptional reactivity.

 $MX_n + nA \rightarrow M^* + nAX$

$$
X = \text{Cl}, \text{Br}, \text{I}; A = \text{Li}, \text{Na}, \text{K}
$$

Ultrasonic waves are known to accelerate a number of heterogeneous reactions, particularly reactions involving metals.2 Recently, Suslick et **aL3** reported a method for

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