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Synthesis, Structural Characterization, and Ligand Replacement Reactions of *gem*-Dithiolato-Bridged Rhodium and Iridium Complexes

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The reaction of *gem*-dithiol compounds $R_2C(SH)_2$ (R = Bn (benzyl), Pr; $R_2 = -(CH_2)_4 - 0$ with dinuclear rhodium or iridium complexes containing basic ligands such as $[M(\mu-OH)(cod)]_2$ and $[M(\mu-OMe)(cod)]_2$, or the mononuclear [M(acac)(cod)] (M = Rh, Ir, cod = 1,5-cyclooctadiene) in the presence of a external base, afforded the dinuclear complexes $[M_2(\mu-S_2CR_2)(cod)_2]$ (1-4). The monodeprotonation of 1,1-dimercaptocyclopentane gave the mononuclear complex $[Rh(HS_2Cptn)(cod)]$ (5) that is a precursor for the dinuclear compound $[Rh_2(\mu-S_2CP_1)(cod)_2]$ (6). Carbonylation of the diolefin compounds gave the complexes $[Rh_2(\mu-S_2CR_2)(CO)_4]$ (7-9), which reacted with P-donor ligands to stereoselectively produce the trans isomer of the disubstituted complexes $[Rh_2(\mu-S_2CR_2)(CO)_2(PR'_3)_2]$ (R' = Ph, Cy (cyclohexyl)) (10-13) and $[Rh_2(\mu-S_2CBn_2)(CO)_2[P(OR')_3]_2]$ (R' = Me, Ph) (14-15). The substitution process in $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) by P(OMe)_3 has been studied by spectroscopic means and the full series of substituted complexes $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) by P(OMe)_3 $_n$] (n = 1, 4) has been identified in solution. The cis complex $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-n_1^P(OR)_3)_n]$ (n = 1, 4) has been identified in solution. The cis complex $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-n_1^P(OR)_3)_n]$ (n = 1, 4) has been identified in solution. The cis complex $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (R = Bn (1), Pr (2); $R_2 = -(CH_2)_4 - (6)$) and that of the cis complex 16 have been studied by X-ray diffraction.

Introduction

Transition metal–sulfur complexes have attracted considerable attention due to their significant relevance to biological and industrial processes.¹ In particular, thiolato derivatives of transition metals are of great importance in coordination chemistry and a large number of mono- and polynuclear compounds containing both simple and functionalized thiolato and dithiolato ligands have been synthesized and structurally characterized.² The long-standing interest in transition metal–thiolate chemistry has been promoted from their relevance as model compounds of metal

6090 Inorganic Chemistry, Vol. 47, No. 13, 2008

sites in metalloenzymes,³ catalytic intermediates in the hydrodesulfurization process (HDS) in homogeneous phase,⁴ and in the study of sulfur poisoning of solid catalysts.⁵

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However, the use of thiolate derivatives as building blocks for heteropolymetallic transition-metal complexes⁶ and supramolecular entities,⁷ their potential application in transitionmetal catalysis for the synthesis of organosulfur compounds,⁸ and their possible use as precursors for metal sulfides with important technological applications⁹ are additional aspects that have stimulated further developments in this field.

In contrast with the plentiful transition-metal complexes containing dithiolato ligands, the number of *gem*-dithiolato complexes, unlike those 1,1-ethylenedithiolato complexes and related unsaturated dithiolene ligands that exhibit planar geometry,¹⁰ is very scarce. On the other hand, trialkylphosphonium–dithioformate complexes also have a four-membered dithiametallacyclobutane structural unit that results from the coordination of the two thiolato groups on the sp³ carbon atom; however, these ligands are electronically very different because of the zwitterionic character of the [S₂C(H)PR₃] ligand.¹¹ Surprisingly, all of the known *gem*-dithiolato complexes have been obtained by indirect meth-

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ods^{12–19} which do not involve *gem*-dithiol compounds as reactants despite being already described in the early 1960s.²⁰ Rational synthetic routes to mono- and dinuclear complexes containing the simple methanedithiolato ligand (S₂CH₂)^{2–} involve the alkylation of several bis-hydrosulfido, M₂(μ -SH)₂ (M = Mo, Re, Fe),¹³ or bis-sulfido Pt₂(μ -S)₂ dinuclear complexes¹⁴ by dihalomethanes, generally in the presence of a base or the insertion of carbon disulfide into M–H bonds (M = Cr, Mo, Re, Ru, Rh).¹⁵ Other complexes with (S₂CR₂)²⁻ ligands have been obtained, for example, by alkylation of dinuclear methanedithiolato complexes,¹⁶ by the baseinduced double Michael addition of activated alkynes to

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dinuclear bis-hydrosulfido iron complexes,¹⁷ or by the addition of acetone to a mononuclear bis-hydrosulfido iridium compound in the presence of catalytic acid.¹⁸ Interestingly, a variety of dirhenium *gem*-dithiolato complexes have been prepared by the reaction of *cis*-[Re₂(μ -O₂CR)₂Cl₂(μ -dppm)₂] with dihydrogen sulfide in the presence of aldehydes or ketones in acid media.¹⁹

Dinuclear d⁸ transition-metal complexes with doubly bridged thiolates, [M2(µ-SR)2L4], have recently attracted widespread interest regarding their electronic, structural, and conformational properties.²¹ However, dinuclear rhodium thiolato bridged complexes were already discovered to be effective catalysts in the hydroformylation of olefins under mild conditions in the early 1980s.²² Subsequent advances in rhodium thiolate chemistry were focused on the utilization of modified thiolato and dithiolato ligands, looking for the influence of a more rigid structure on the catalytic activity.²³ Furthermore, as chiral information can be introduced on the backbone of the dithiolato ligand, these dinuclear systems have found application in asymmetric hydroformylation, especially in combination with chiral diphosphines.²⁴ Interestingly, complexes with dithiolato ligands have also been used as synthons for the preparation of heterobimetallic complexes with catalytic activity.²⁵

We have recently reported the straighforward synthesis of the complex $[Rh_2(\mu-S_2Chxn)(cod)_2]$ (ChxnS₂²⁻ = 1,1-cyclohexanedithiolato) which is, to our knowledge, the first example of direct synthesis of a *gem*-dithiolato-bridged

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complex from a *gem*-dithiol compound.²⁶ This compound, as the bis-thiolato and dithiolato dinuclear counterparts, is an active catalyst precursor for the hydroformylation of olefins under mild conditions. We report herein the scope and limitations of this new synthetic methodology for the direct synthesis of diolefin dinuclear *gem*-dithiolato complexes $[M_2(\mu$ -S₂CR₂)(cod)₂] (M = Rh, Ir) from several *gem*-dithiols (R₂C(SH)₂; R = Bn, ^{*i*}Pr; and R₂ = -(CH₂)₄-). In addition, the reactivity of the rhodium complexes with carbon monoxide, the replacement reactions with several P-donor ligands, and the comparison with the related thiolato dinuclear systems are also described.

Experimental Section

General. All manipulations were performed under a dry argon atmosphere using Schlenk-tube techniques. Solvents were dried by standard methods and distilled under argon immediately prior to use. Standard literature procedures were used to prepare the complexes [Rh(μ -OH)(cod)]₂, [M(μ -OMe)(cod)]₂ (M = Rh, Ir),²⁷ [M(acac)(cod)] (M = Rh, Ir),²⁸ [Rh(acac)(CO)(PPh₃)],^{28a} and [Rh(acac)(CO)(PCy₃)].²⁹

Physical Measurements. ${}^{1}H$, ${}^{31}P{}^{1}H$, and ${}^{13}C{}^{1}H$ NMR spectra were recorded on a Varian Gemini 300 spectrometer operating at 300.08, 121.47, and 75.46 MHz respectively. Chemical shifts are reported in parts per million and referenced to SiMe₄ using the residual resonances of the deuterated solvents (1H and ¹³C) and 85% H₃PO₄ (³¹P) as an external reference, respectively. Assignments in complex NMR spectra were done by simulation with the program gNMR ver. 3.6 (Cherwell Scientific Publishing Limited) for Macintosh. The initial choice of chemical shifts and coupling constants were optimized by successive iterations following a standard least-squares procedure, a numerical assignment of the experimental frequencies was used. IR spectra were recorded on a Nicolet-IR 550 spectrometer. Elemental carbon, hydrogen, and nitrogen analyses were performed with a PerkinElmer 2400 microanalyzer. Molecular weights were determined with a Knauer osmometer using chloroform solutions of the complexes. Mass spectra were recorded in a VG Autospec double-focusing mass spectrometer operating in the FAB⁺ mode. Ions were produced with the standard Cs⁺ gun at ca. 30 Kv, and 3-nitrobenzylic alcohol (NBA) was used as a matrix.

Synthesis and Characterization of *gem*-Dithiol Compounds. 1,1-Dimercaptocyclopentane, $Cptn(SH)_2$,^{20a,b} 2,4-dimethyl-3,3-dimercaptopentane, ${}^{1}Pr_2C(SH)_2$,^{20c} and 1,3-diphenyl-2,2-dimercaptopropane, $Bn_2C(SH)_2$,^{20d} were prepared according to reported

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methods. Analytical and NMR data: **Cptn(SH)**₂. Anal. Calcd for C₅H₁₀S₂: C, 44.73; H, 7.51; S, 47.76. Found: C, 44.70; H, 7.48; S, 47.70. ¹H NMR (CDCl₃, 293 K) & 2.69 (s, 2H, SH), 2.06 (m, 4H, >CH₂), 1,81 (m, 4H, >CH₂). ¹³C{¹H} NMR (CDCl₃, 293 K) & 55.6 (C¹), 48.1 (C² and C⁵), 23.6 (C³ and C⁴). **Bn₂C(SH)**₂. Anal. Calcd for C₁₅H₁₆S₂: C, 69.18; H, 6.19; S, 24.62. Found: C, 69.14; H, 6.10; S, 24.50. ¹H NMR (CDCl₃, 293 K) & 7.33-7.28 (m, 10H, Ph), 3.22 (s, 4H, >CH₂), 2.36 (s, 2H, SH). ¹³C{¹H} NMR (CDCl₃, 293 K) & 136.0, 131.5, 127.9, 127.4 (Ph), 56.8 (C(SH)₂), 52.8 (>CH₂). ⁱ**Pr₂C(SH)**₂. Anal. Calcd for C₇H₁₆S₂: C, 51.16; H, 9.81; S, 39.02. Found: C, 51.10; H, 9,70; S, 38.91. ¹H NMR (CDCl₃, 293 K) & 2.08 (sept, 2H, $J_{H-H} = 7$ Hz, CH, ⁱPr), 2.05 (s, 2H, SH), 1.07 (d, 12H, $J_{H-H} = 7$ Hz, $-CH_3$, ⁱPr). ¹³C{¹H} NMR (CDCl₃, 293 K) & 68.0 (C(SH)₂), 37.7 (CH), 18.2 (CH₃) (ⁱPr).

Synthesis of the Complexes. $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1). Method A. To a solution of $[Rh(\mu-OH)(cod)]_2$ (0.500 g, 1.096 mmol) in CH₂Cl₂ (5 mL) was added 1,3-diphenyl-2,2-dimercaptopropane, Bn₂C(SH)₂, (0.312 g, 1.200 mmol) to give immediately an orangered solution that was stirred for 15 min. The addition of EtOH (5 mL) afforded an orange suspension that was concentrated under vacuum to ca. 5 mL and then filtered to give an orange microcrystalline solid, which was washed with EtOH (2 \times 3 mL) and dried under vacuum. Yield: 0.627 g (84%). Method B. [Rh(µ-OMe)(cod)]₂ (0.100 g, 0.206 mmol) and Bn₂C(SH)₂ (0.054 g, 0.206 mmol) were reacted in CH₂Cl₂ (5 mL) to give immediately a red solution that was stirred for 30 min. The compound was isolated as an orange-brown microcrystalline solid following the procedure described above. Yield: 0.122 g (87%). Method C. Triethylamine $(120 \ \mu L, 0.652 \ mmol, \rho = 0.728 \ g \ mL^{-1})$ and $Bn_2C(SH)_2 \ (0.091)$ g, 0.350 mmol) were successively added to a solution of [Rh(acac)-(cod)] (0.202 g, 0.650 mmol) in CH₂Cl₂ (5 mL) to give a red solution that was stirred for 15 min. Work-up as described above gave the compound as a red microcrystalline solid. Yield: 0.188 g (85%). Anal. Calcd for C₃₁H₃₈S₂Rh₂: C, 54.71; H, 5.63; S, 9.42. Found: C, 54.71; H, 5.90; S, 9.57. ¹H NMR (CDCl₃, 293 K) δ: 7.21 (m, 10H, Ph), 4.3 (m, 4H, =CH), 4.16 (m, 4H, =CH) (cod), 3.56 (s, 4H, >CH₂, Bn), 2.36 (m, 4H, >CH₂), 2.27 (m, 4H, >CH₂), 1.80 (m, 4H, >CH₂), 1.74 (m, 4H, >CH₂) (cod). ¹³C{¹H} NMR (CDCl₃, 293 K) δ: 137.8, 131.2, 127.6, 126.4 (Bn), 86.1 (CS₂), 79.9 (d, $J_{Rh-C} = 12$ Hz, =CH), 78.5 (d, $J_{Rh-C} = 12$ Hz, =CH) (cod), 60.9 (>CH₂, Bn), 30.6 (>CH₂, cod). MS (FAB⁺, CH₂Cl₂, m/z): 680 (M⁺, 25%), 572 (M⁺ - cod, 15%), 464 (M⁺ - 2cod, 22%).

 $[Rh_2(\mu - S_2C^iPr_2)(cod)_2]$ (2). $[Rh(\mu - OH)(cod)]_2$ (0.151 g, 0.331 mmol) and 2,4-dimethyl-3,3-dimercaptopentane, $Pr_2C(SH)_2$, (52 μ L, $\rho \simeq 1 \text{ g mL}^{-1}$, 0.317 mmol) were reacted in CH₂Cl₂ (10 mL) at -78 °C for 10 min. The solution was allowed to warm up to room temperature and stirred for 20 min. The resulting red-orange solution was concentrated under vacuum to about one-half the volume and then EtOH (5 mL) was added to give a red-orange solid that was filtered, washed with EtOH (2×2 mL), and dried under vacuum. The solid was dissolved in an n-hexane/CH₂Cl₂ (2:1) mixture and then eluted through an alumina column (14 \times 1.5 cm) to give an orange solution. Concentration of the solution afforded an orange microcrystalline solid that was filtered, washed with *n*-pentane (2 \times 3 mL), and dried under vacuum. Yield: 0.151 g (78%). Anal. Calcd for C₂₃H₃₈S₂Rh₂: C, 47.26; H, 6.55; S, 10.97. Found: C, 47.21; H, 6.40; S, 10.77. ¹H NMR (CDCl₃, 293 K) δ: 4.50 (m, 4H, =CH), 4.25 (m, 4H, =CH) (cod), 2.65 (sept, 2H, $J_{H-H} = 6.9$ Hz, CH, Pr), 2.45 (m, 8H, >CH₂), 2.0 (m, 4H, >CH₂), 1.85 (m, 4H, >CH₂) (cod), 1.20 (d, 12H, $J_{H-H} = 6.9$ Hz, $-CH_3$, ^{*i*}Pr). ¹³C{¹H} NMR (CDCl₃, 293 K) δ : 101.2 (CS₂), 79.5 (d, $J_{Rh-C} = 11.5$ Hz, =CH), 78.6 (d, J_{Rh-C} = 12.4 Hz, =CH) (cod), 46.5 (CH, ^{*i*}Pr), 31.6 and 31.1 (>CH₂) (cod), 20.1 (-CH₃, ⁱPr). MS (FAB⁺, CH₂Cl₂, *m*/*z*): 584 (M⁺, 100%).

[**Ir**₂(*μ*-**S**₂CBn₂)(cod)₂] (3). [Ir(acac)(cod)] (0.104 g, 0.260 mmol) and Bn₂C(SH)₂ (0.040 g, 0.153 mmol) were reacted in CH₂Cl₂ (5 mL) for 15 min to give a dark-red suspension. The suspension was concentrated under vacuum to about one-half the volume, and then MeOH (3 mL) was added to complete the precipitation. The compound was isolated by filtration, washed with MeOH (2 × 2 mL), and dried under vacuum. Yield: 0.099 g (88%). Anal. Calcd for C₃₁H₃₈S₂Ir₂: C, 43.34; H, 4.46; S, 7.46. Found: C, 43.29; H, 4.36; S, 6.61. ¹H NMR (CDCl₃, 293 K) δ: 7.23 (m, 10H, Ph), 4.04 (m, 2H, =CH), 3.81 (m, 2H, =CH) (cod), 3.51 (s, 4H, >CH₂, Bn), 2.09 (m, 8H, >CH₂), 1.66 (m, 4H, >CH₂), 1.27 (m, 4H, >CH₂) (cod). ¹³C{¹H} NMR (CDCl₃, 293 K) δ: 137.7, 131.2, 127.8, 126.6 (Bn), 95.1 (CS₂), 66.2 (=CH, cod), 63.0 (>CH₂, Bn), 62.9 (=CH), 31.8 and 31.6 (>CH₂, cod). MS (FAB⁺, CH₂Cl₂, *m/z*): 860 (M⁺, 40%), 661 (M⁺ - cod - Bn, 30%).

 $[Ir_2(\mu - S_2C^iPr_2)(cod)_2]$ (4). $[Ir(\mu - OMe)(cod)]_2$ (0.126 g, 0.190 mmol) and ${}^{i}Pr_{2}C(SH)_{2}$ (29 μ L, $\rho \simeq 1$ g mL⁻¹, 0.177 mmol) were reacted in CH₂Cl₂ (10 mL) at -78 °C for 10 min. The compound was isolated as a dark-red solid following the procedure described for 2. The crude compound was dissolved in a n-hexane/CH₂Cl₂ (3:1) mixture and then eluted through an alumina column (14 \times 1.5 cm) to give a violet solution. Concentration of the solution afforded a purple microcrystalline solid that was filtered, washed with *n*-pentane $(2 \times 3 \text{ mL})$, and dried under vacuum. Yield: 0.110 g (76%). Anal. Calcd for C₂₃H₃₈S₂Ir₂: C, 36.20; H, 5.02; S, 8.40. Found: C, 36.00; H, 4.73; S, 8.83. ¹H NMR (CDCl₃, 293 K) δ : 4.26 (m, 4H, =CH), 3.92 (m, 4H, =CH) (cod), 2.48 (sept, 2H, CH, $J_{H-H} = 6.9$ Hz, ^{*i*}Pr), 2.28 (m, 4H, >CH₂), 2.16 (m, 4H, >CH₂), $1.87 \text{ (m, 4H, >CH_2), } 1.41 \text{ (m, 4H, >CH_2) (cod), } 0.98 \text{ (d, 12H, }$ $-CH_3$, $J_{H-H} = 6.9$ Hz, ^{*i*}Pr). ¹³C{¹H} NMR (CDCl₃, 293 K) δ : 110.4 (CS₂), 65.7 and 63.2 (=CH, cod), 50.6 (CH, ⁱPr), 32.6 and 31.9 (>CH₂, cod), 19.9 (-CH₃, *i*Pr). MS (FAB⁺, CH₂Cl₂, *m/z*): 762 (M⁺, 100%), 613 (M⁺ - cod - ^{*I*}Pr, 24%).

[Rh(HS₂Cptn)(cod)] (5). To a solution of [Rh(acac)(cod)] (0.171 g, 0.550 mmol) in CH₂Cl₂ (10 mL) was added 1,1-dimercaptocyclopentane, Cptn(SH)₂, (77 μ L, 0.550 mmol, $\rho = 0.95$ g mL⁻¹) to give immediately a dark-red solution that was stirred for 15 min. The solution was concentrated under vacuum to ca. 5 mL, and then MeOH (5 mL) was added to give a dark-red suspension. The suspension was further concentrated, and then diethyl ether (5 mL) was added to complete the precipitation. The suspension was filtered and then recrystallized from CH2Cl2/diethyl ether to give the compound as a dark-red solid. Yield: 0.128 g (68%). Anal. Calcd for C₁₃H₂₁S₂Rh: C, 45.34; H, 6.15; S, 18.62. Found: C, 45.30; H, 5.95; S, 18.13. ¹H NMR (CDCl₃, 293 K) δ : 4.18 (m, 4H, =CH) (cod), 2.39 (m, 5H, >CH₂, cod and SH), 1.93 (br, 4H, >CH₂) (cod), 1.82 (m, 4H, >CH₂), 1.64 (m, 4H, >CH₂) (Cptn). ¹³C{¹H} NMR $(CDCl_3, 293 \text{ K}) \delta$: 81.7 (d, $J_{Rh-C} = 11 \text{ Hz}$, =CH, cod), 66.8 (C¹), 48.5 (C² and C⁵) (Cptn), 31.6 (>CH₂, cod), 23.8 (C³ and C⁴, Cptn). MS (FAB⁺, CH₂Cl₂, m/z): 554 (M⁺ + Rh(cod) - H, 89%), 343 $(M^+ - H, 67\%).$

[**Rh**₂(μ -S₂**Cptn**)(**cod**)₂] (6). [Rh(μ -OH)(**cod**)]₂ (0.350 g, 0.767 mmol) and Cptn(SH)₂, (108 μ L, 0.767 mmol, $\rho = 0.95$ g mL⁻¹) were reacted in CH₂Cl₂ (10 mL) at room temperature for 10 min. The compound was isolated as a dark-red solid following the procedure described for **1**. The crude compound was dissolved in CH₂Cl₂ and then eluted through an alumina column (12 × 1.5 cm) with *n*-hexane/dichloromehane (2:1) to give an orange solution. Concentration of the solution afforded an orange microcrystalline solid that was filtered, washed with *n*-pentane (2 × 3 mL), and dried under vacuum. Yield: 0.263 g (62%). Anal. Calcd for

Rivas et al.

 $\begin{array}{l} C_{21}H_{32}S_2Rh_2:\ C,\ 45.49;\ H,\ 5.82;\ S,\ 11.56.\ Found:\ C,\ 45.61;\ H,\ 5.69;\\ S,\ 11.23.\ ^{1}H\ NMR\ (C_6D_6,\ 293\ K)\ \delta:\ 4.85\ (m,\ 4H,\ =CH),\ 4.44\ (m,\ 4H,\ =CH)\ (cod),\ 2.54\ (m,\ 4H,\ >CH_2,\ Cptn),\ 2.46\ (m,\ 8H,\ >CH_2),\ 2.00\ (m,\ 4H,\ >CH_2),\ 1.85\ (m,\ 4H,\ >CH_2)\ (cod),\ 1.70\ (m,\ 4H,\ >CH_2),\ 1.85\ (m,\ 4H,\ >CH_2)\ (cod),\ 1.70\ (m,\ 4H,\ >CH_2,\ Cptn).\ ^{13}C\{^{1}H\}\ NMR\ (CDCl_3,\ 293\ K)\ d:\ 79.9\ (d,\ J_{Rh-C}=11\ Hz,\ =CH),\ 78.9\ (d,\ J_{Rh-C}=12\ Hz,\ =CH)\ (cod),\ 81.9\ (C^{1}),\ 60.0\ (C^2\ and\ C^5)\ (Cptn),\ 31.5\ and\ 31.1\ (>CH_2,\ cod),\ 25.0\ (C^3\ and\ C^4,\ Cptn).\ MS\ (FAB^+,\ CH_2Cl_2,\ m/z):\ 554\ (M^+,\ 100\%),\ 446\ (M^+-\ -cod,\ 45\%),\ 338\ (M^+-\ 2cod,\ 32\%). \end{array}$

 $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7). Carbon monoxide was bubbled through an orange solution of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1) (0.100 g, 0.147 mmol) in CH₂Cl₂ (5 mL) for 10 min to give a pale-red solution. n-Hexane (10 mL) was added, and the volume of the solution reduced by continuous bubbling of carbon monoxide until a red solid precipitated. Addition of n-hexane (3 mL) and concentration by bubbling led to further precipitation. The compound was isolated by filtration, washed with *n*-hexane $(2 \times 3 \text{ mL})$, and dried under vacuum. Yield: 0.078 g (92%). Anal. Calcd for C₁₉H₁₄O₄S₂Rh₂: C, 39.60; H, 2.45; S, 11.13. Found. C, 39.90; H, 2.63; S, 11.09. ¹H NMR (CDCl₃, 293 K) δ: 7.45 (m, 8H), 7.28 (m, 2H), 3.31 (s, 4H, >CH₂) (Bn). ${}^{13}C{}^{1}H{}$ NMR (CDCl₃, 293 K) δ : 184.4 (d, $J_{\text{Rh}-\text{C}} = 72$ Hz, CO), 136.9, 132.1, 128.3, 127.8, (Bn), 95.4 (CS₂), 60.6 (>CH₂, Bn). MS (FAB⁺, CH₂Cl₂, *m/z*): 576 (M⁺, 70%); 548 (M⁺ - CO, 95%), 520 (M⁺ - 2CO, 70%), 492 (M⁺ -3CO, 85%), $464 (M^+ - 4CO, 100\%)$, $430 (M^+ - 2CO - Bn)$, 80%). IR (CH₂Cl₂, cm⁻¹): ν (CO), 2078 (m), 2054 (s), 2010 (s).

 $[Rh_2(\mu-S_2CBn_2)(CO)_2(PPh_3)_2]$ (10). Method A. To a solution of [Rh(acac)(CO)(PPh₃)] (0.103 g, 0.209 mmol) in CH₂Cl₂ (10 mL) was added Bn₂C(SH)₂ (0.027 g, 0.105 mmol) to give a red solution that was stirred for 15 min. The solution was concentrated under vacuum to about one-half the volume, and then MeOH (6 mL) was added to afford an orange solid that was filtered, washed with MeOH (2×3 mL), and dried under vacuum. Yield: 0.101 g (92%). Method B. To a solution of compound $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7), prepared in situ by carbonylation of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1) (0.100 g, 0.147 mmol) in CH₂Cl₂ (10 mL) for 15 min was added solid PPh₃ (0.077 g, 0.295 mmol) to give an orange solution with evolution of carbon monoxide. The solution was stirred for 10 min, and then MeOH (10 mL) was added to give a red suspension. Concentration of the suspension to about one-half the volume led to further precipitation. Filtration and washing with MeOH (3 \times 3 mL) followed by drying under vacuum gave the product as an orange-red solid. Yield: 0.138 g (90%). Anal. Calcd for C₅₃H₄₄O₂P₂S₂Rh₂: C, 60.93; H, 4.24; S, 6.14. Found: C, 60.43; H, 4.44; S: 6.04. ¹H NMR (C₆D₆, 293 K) δ: 7.6 (m, 12H), 7.3 (m, 18H) (PPh₃), 7.0 (m, 10H, Ph), 3.50 (AB q: $\delta_A = 3.98$, $\delta_B = 3.3$, $J_{AB} = 14.1 \text{ Hz}, 4\text{H}, > \text{CH}_2, \text{Bn}$). ¹³C{¹H} NMR (C₆D₆; 293 K) δ : 192.5 (dd, $J_{Rh-C} = 76$ Hz, ${}^{2}J_{P-C} = 18$ Hz, CO), 138.7 (Ph), 135.8 (d, $J_{P-C} = 72$ Hz), 134.8 (d, $J_{P-C} = 12$ Hz) (PPh₃), 131.8 (Ph), 130.5 (PPh₃), 128.9 (d, $J_{P-C} = 12$ Hz) (PPh₃), 127.1 (Ph), 89.1 (CS₂), 61.9 (>CH₂, Bn). MS (FAB⁺, CH₂Cl₂, *m/z*): 1044 (M⁺, 100%), 1016 (M⁺ - CO, 35%), 988 (M⁺ - 2CO, 11%), 754 (M⁺ - CO - PPh₃, 36%). IR (toluene, cm⁻¹): ν (CO), 1967 (s).

[**Rh**₂(μ -S₂C^{*i*}Pr₂)(**CO**)₂(**PPh**₃)₂] (11). Carbon monoxide was bubbled through a solution of [Rh₂(μ -S₂C^{*i*}Pr₂)(cod)₂] (2) (0.086 g, 0.147 mmol) in CH₂Cl₂ (10 mL) to give a yellow-brown solution of the compound [Rh₂(μ -S₂C^{*i*}Pr₂)(CO)₄] (8) in 15 min. Further reaction with PPh₃ (0.077 g, 0.294 mmol) gave a deep-red solution with evolution of carbon monoxide. Work-up as described above for **10** (method B) gave the compound as a red solid. Yield: 0.120 g (93%). Anal. Calcd for C₄₅H₄₄O₂P₂S₂Rh₂: C, 56.97; H, 4.67; S, 6.76. Found: C, 56.56; H, 4.73; S, 6.75. ¹H NMR (C₆D₆, 293 K) δ: 8.11 (m, 12H), 7.32–7.18 (m, 18H) (PPh₃), 3.07 (m, 2H, CH), 1.19 (d, $J_{H-H} = 6.9$ Hz, -CH₃), 1.13 (d, $J_{H-H} = 6.9$ Hz, -CH₃) ('Pr). ¹³C{¹H} NMR (C₆D₆, 293 K) δ : 191.7 (dd, $J_{Rh-C} = 75$ Hz, ² $J_{P-C} = 17$ Hz, CO), 135.2 (d, $J_{P-C} = 44$ Hz), 133.5 (d, $J_{P-C} = 12$ Hz), 129.6, 128.2 (d, $J_{P-C} = 5$ Hz) (PPh₃), 102.3 (CS₂), 47.6 (CH), 19.2 and 19.1(-CH₃) ('Pr). MS (FAB⁺, CH₂Cl₂, *m*/*z*): 948 (M⁺, 100%); 920 (M⁺ - CO, 50%); 890(M⁺ - 2CO, 40%); 686 (M⁺ -PPh₃, 35%). IR (CH₂Cl₂, cm⁻¹): ν (CO), 1956 (s).

[**Rh**₂(*μ*-S₂**Cptn**)(**CO**)₂(**PPh**₃)₂] (12). [Rh(acac)(CO)(PPh₃)] (0.065 g, 0.132 mmol) and Cptn(SH)₂ (10.7 μ L, 0.075 mmol, $\rho = 0.95$ g mL⁻¹) were reacted in CH₂Cl₂ (10 mL) for 15 min to give an orange solution. The compound was isolated as an orange solid following the procedure described for 10 (method A). Yield: 0.049 g (81%). Anal. Calcd for C₄₃H₃₈O₂P₂S₂Rh₂: C, 56.22; H, 4.17; S, 6.98. Found: C, 56.00; H, 4.16; S: 7.03. ¹H NMR (CDCl₃, 293 K) δ: 7.77–7.69 (m, 12H), 7.42–7.33 (m, 18H) (PPh₃), 2.19 (m, 4H, >CH₂), 1.57 (m, 4H, >CH₂) (Cptn). ¹³C{¹H} NMR (CDCl₃, 293 K) δ: 0.5191.4 (dd, *J*_{Rh-C} = 76 Hz, ²*J*_{P-C} = 17 Hz, CO), 134.8 (d, *J*_{P-C} = 46 Hz), 133.9 (d, *J*_{P-C} = 12 Hz), 129.8 (s), 128.1 (d, *J*_{P-C} = 12 Hz) (PPh₃), 84.3 (C¹), 60.0 (C² and C⁵), 24.5 (C³ and C⁵) (Cptn). MS (FAB⁺, CH₂Cl₂, *m/z*): 918 (M⁺, 82%), 890 (M⁺ – CO, 55%). IR (toluene, cm⁻¹): ν(CO), 1972 (s).

[Rh₂(µ-S₂CBn₂)(CO)₂(PCy₃)₂] (13). [Rh(acac)(CO)(PCy₃)] (0.103 g, 0.202 mmol) and Bn₂C(SH)₂ (0.026 g, 0.101 mmol) were reacted in CH₂Cl₂ (10 mL) for 15 min to give a pale-orange solution that was stirred for 15 min. The compound was isolated as a yellow solid following the procedure described for 10 (method A). Yield: 0.089 g (82%). Anal. Calcd for C₅₃H₈₀O₂P₂S₂Rh₂: C, 58.88; H, 7.46; S, 5.93. Found: C, 58.35; H, 7.22; S, 6.02. ¹H NMR (CDCl₃, 293 K) δ : 7.20 (m, 10H, Bn), 3.96 (AB q, $\delta_A = 4.02$, $\delta_B = 3.88$, $J_{AB} = 14.85 \text{ Hz}, 4\text{H}, > \text{CH}_2, \text{Bn}$, 2.04 (m, 18 H), 1.75 (m, 30 H), 1.23 (m, 18 H) (PCy₃). ${}^{13}C{}^{1}H{}$ NMR (CDCl₃, 293 K) δ : 192.1 (dd, $J_{\text{Rh-C}} = 76.9 \text{ Hz}$, ${}^{2}J_{\text{P-C}} = 17.5 \text{ Hz}$, CO), 138.6, 130.8, 128.2, 126.7 (Bn), 86.2 (CS₂), 62.0 (>CH₂, Bn), 37.0 (d, $J_{P-C} = 22$ Hz, -CHP), 30.7, 28.1 (d, $J_{P-C} = 12.4$ Hz), 26.9 (>CH₂, PCy₃). MS $(FAB^+, CH_2Cl_2, m/z)$: 1080 (M⁺, 17%), 1052 (M⁺ - CO, 20%), 1024 (M⁺ - 2CO, 15%), 744 (M⁺ - 2CO - PCy₃, 30%). IR $(CH_2Cl_2, cm^{-1}): \nu(CO), 1942$ (s).

[Rh₂(µ-S₂CBn₂)(CO)₂{P(OMe)₃}₂] (14). To a solution of [Rh₂(µ-S₂CBn₂)(CO)₄] (7) (0.024 g, 0.042 mmol) in CH₂Cl₂ (5 mL) at room temperature was added P(OMe)₃ (10 μ L, 0.084 mmol, ρ = 1.05 g mL $^{-1}\!\!,\,98\%)$ to give a yellow solution with evolution of carbon monoxide. The solution was stirred for 10 min and then n-hexane (5 mL) was added. The solvent was slowly eliminated under vacuum, and the resulting thick solid was washed with *n*-hexane $(3 \times 3 \text{ mL})$ to give a light orange solid that was filtered and dried under vacuum. Yield: 0.029 g (91%). Anal. Calcd for C₂₃H₃₂O₈P₂S₂Rh₂: C, 35.95; H, 4.20; S, 8.34. Found. C, 36.15; H, 4.18; S, 8.12. ¹H NMR (C₆D₆, 293 K) δ: 7.54 (d, 4H), 7.27 (m, 4H), 7.19 (d, 2H) (Bn), 3.79 (AB q: $\delta_A = 4.08$, $\delta_B = 3.50$, $J_{AB} =$ 13.35 Hz, 4H, >CH₂, Bn), 3.58 (d, ${}^{2}J_{C-P} = 12$ Hz, -OMe). ¹³C{¹H} NMR (C₆D₆, 293 K) δ : 190.24 (dd, $J_{Rh-C} = 74$ Hz, ${}^{2}J_{P-C}$ = 24 Hz, CO), 137.98, 131.59, 128.59, 127.20 (Bn), 90.42 (CS₂), 62.56 (>CH₂, Bn), 51.70 (OMe). MS (FAB⁺, CH₂Cl₂, m/z): 768 $(M^+, 32\%), 712 (M^+ - 2CO, 100\%), 616 (M^+ - CO - P(OMe)_3),$ 33%), 588 (M⁺ - 2CO - P(OMe)₃, 45%), 464 (M⁺ - 2CO - $2P(OMe)_3$, 25%). IR (toluene, cm⁻¹): $\nu(CO)$, 1986 (s).

[**Rh**₂(μ -S₂CBn₂)(**CO**)₂{**P**(**OPh**)₃}₂] (**15**). [Rh₂(μ -S₂CBn₂)(CO)₄] (7) (0.020 g, 0.035 mmol) and P(OPh)₃ (19 μ L, 0.070 mmol, ρ = 1.184 g mL⁻¹, 97%) were reacted in CH₂Cl₂ (5 mL) at room temperature to give a yellow solution with the evolution of carbon monoxide. Work-up as described above gave the compound as a yellow solid. Yield: 0.035 g (88%). Anal. Calcd for C₅₃H₄₄O₈P₂S₂Rh₂: C, 55.80; H, 3.89; S, 5.62. Found: C, 55.68; H,

Table 6. Selected Bond Distances (Angstroms) and Angles (Degrees) for ${\bf 16}$

Rh(1)-S(1)	2.3995(5)	Rh(2)-S(1)	2.4001(5)
Rh(1) - S(2)	2.3926(5)	Rh(2)-S(2)	2.3801(5)
Rh(1)-P(1)	2.2655(5)	Rh(2) - P(2)	2.2618(5)
Rh(1) - C(1)	1.840(2)	Rh(2) - C(2)	1.837(2)
S(1) - C(3)	1.869(2)	S(2) - C(3)	1.867(2)
P(1)-C(21)	1.851(2)	P(2) - C(18)	1.834(2)
P(1)-C(22)	1.830(2)	P(2) - C(34)	1.819(2)
P(1)-C(28)	1.823(2)	P(2) - C(40)	1.826(2)
C(1) - O(1)	1.142(2)	C(2) - O(2)	1.144(3)
C(3)-C(4)	1.540(3)	C(3) - C(11)	1.551(3)
C(18)-C(19)	1.536(3)	C(19)-C(20)	1.531(3)
C(20)-C(21)	1.528(3)		
S(1) - Rh(1) - S(2)	71.047(16)	S(1) - Rh(2) - S(2)	71.248(16)
S(1) - Rh(1) - P(1)	96.810(17)	S(1) - Rh(2) - P(2)	95.385(18)
S(1) - Rh(1) - C(1)	172.34(6)	S(1) - Rh(2) - C(2)	168.50(6)
S(2) - Rh(1) - P(1)	166.816(18)	S(2) - Rh(2) - P(2)	166.080(18)
S(2) - Rh(1) - C(1)	101.30(6)	S(2) - Rh(2) - C(2)	98.92(6)
P(1) - Rh(1) - C(1)	90.82(6)	P(2) - Rh(2) - C(2)	94.77(6)
Rh(1) - S(1) - Rh(2)	75.894(15)	Rh(1) - S(2) - Rh(2)	76.399(14)
Rh(1) - S(1) - C(3)	85.21(6)	Rh(1) - S(2) - C(3)	85.46(6)
Rh(2) - S(1) - C(3)	85.19(6)	Rh(2) - S(2) - C(3)	85.82(6)
S(1)-C(3)-S(2)	96.36(9)	C(4) - C(3) - C(11)	112.35(16)
S(1) - C(3) - C(4)	110.01(13)	S(2) - C(3) - C(4)	113.87(13)
S(1)-C(3)-C(11)	112.95(14)	S(2) - C(3) - C(11)	110.36(13)
Rh(1) - C(1) - O(1)	179.6(2)	Rh(2) - C(2) - O(2)	176.16(19)

4.02; S, 5.55. ¹H NMR (C₆D₆, 293 K) δ : 7.71 (d, 12H, OPh), 7.51–7.33 (m, 10H, Bn), 7.27 (m, 12H), 7.06 (m, 6H) (OPh), 3.68 (AB q: $\delta_{A} = 3.68$, $\delta_{B} = 3.48$, $J_{AB} = 14.4$ Hz, 4H, >CH₂, Bn). ¹³C{¹H} NMR (C₆D₆, 293 K) δ : 189.2 (d, $J_{Rh-C} = 74$ Hz, ² $J_{P-C} = 24$ Hz, CO), 152.4 (d, J = 6 Hz, OPh), 137.6, 131.7 (Bn), 130.2 (OPh), 128.5, 127.3 (Bn), 125.5, 121.9 (OPh), 89.1 (CS₂), 61.8 (>CH₂, Bn). MS (FAB⁺, CH₂Cl₂, m/z): 1140 (M⁺, 20%), 1112 (M⁺ – CO, 30%), 1084 (M⁺ – 2CO, 45%), 802 (M⁺ – P(OPh)₃ – CO, 35%), 774 (M⁺ – P(OPh)₃ – 2CO, 85%). IR (toluene, cm⁻¹): ν (CO), 1994 (s).

 $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-dppb)]$ (16). To a solution of 7 prepared in situ by carbonylation of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1) (0.059 g, 0.087 mmol) in CH₂Cl₂ (10 mL) was added solid dppb (0.036 g, 0.087 mmol) to give a yellow solution with evolution of carbon monoxide. The solution was stirred for 10 min and then concentrated under vacuum to about one-half the volume. The addition of MeOH (5 mL) gave the compound as a yellow solid that was filtered, washed with cold MeOH (3 \times 3 mL), and dried under vacuum. Yield: 0.072 g (87%). Anal. Calcd for C45H42O2P2S2Rh2: C, 57.09; H, 4.47; S, 6.77. Found: C, 56.94; H, 4,35; S, 6.65. ¹H NMR (CDCl₃, 293 K) δ: 7.80 (m, 4H), 7.67 (m, 4H), 7.64-7.00 (m, 14H), 7.04 (m, 8H) (Bn and dppb), 3.56 (AB q, $\delta_A = 3.59$, δ_B = 3.54, J_{AB} = 14.10 Hz, 4H, >CH₂, Bn), 2.46 (m, 4H), 2.20 (m, 2H), 1.40 (m, 2H) (>CH₂, dppb). ³¹P{¹H} NMR (CDCl₃, 293 K) δ: 32.25 (d, $J_{Rh-P} = 157$ Hz). MS (FAB⁺, CH₂Cl₂, *m/z*): 946 (M⁺, 95%), 918 (M⁺ - CO, 26%), 890 (M⁺ - 2CO, 72%). IR (toluene, cm⁻¹): ν (CO), 1980 (s).

Crystal Structure Determination of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1), $[Rh_2(\mu-S_2CPr_2)(cod)_2]$ (2), $[Rh_2(\mu-S_2Cptn)(cod)_2]$ (6), and $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-dppb)]$ (16). Suitable crystals for X-ray diffraction of 1, 2, 6, and 16 were obtained by slow diffusion of *n*-hexane into CH₂Cl₂ solutions of the complexes at 258 K. A summary of crystal data and refinement parameters for the structural analyses is given in Table 6. The crystals used in the analyses were glued to a glass fiber and mounted on a Bruker SMART APEX diffractometer. The instrument was equipped with a CCD area detector, and data were collected using graphite-monochromated Mo K α radiation ($\lambda = 0.71073$ Å) at low temperature (173 K (1) or 100(1) K (2, 6, and 16)). Cell constants were obtained from the least-squares refinement of 3D centroids (3185 reflns, $4.5 \le 2\theta \le 29.3^{\circ}$ for **1**; 6234, 9863, or 19 904 reflns, $4.5 \le 2\theta \le 56.6^{\circ}$ for **2**, **6**, or **16**, respectively). Data were measured through the use of CCD recording of narrow ω rotation frames (0.3° each). All data were integrated with the Bruker *SAINT* program,³⁰ which includes Lorentz and polarization corrections. Absorption correction was applied by using the *SADABS* routine.³¹

The structures were solved by Patterson methods, completed by subsequent difference Fourier techniques, and refined by full-matrix least-squares on F^2 (*SHELXL-97*)³² with initial isotropic but subsequent anisotropic thermal parameters for all non-hydrogen atoms. Hydrogen atoms in 1 were included in the model from calculated or observed (olefinic hydrogens) positions but were refined with riding positional and displacement parameters. In the case of **2**, **6**, and **16**, hydrogen atoms were obtained from difference Fourier maps and refined as free isotropic atoms. Atomic scattering factors were used as implemented in the program.³²

Results

Synthesis and Characterization of Diolefin gem-Dithiolato-Bridged Rhodium and Iridium Dinuclear **Compounds.** The synthesis of *gem*-dithiolato-bridged dinuclear complexes can be accomplished directly from gemdithiol compounds using standard mono- or dinuclear compounds containing basic ligands. Thus, reaction of $[Rh(\mu OH(cod)]_2$ with 1,3-diphenyl-2,2-dimercaptopropane, $Bn_2C(SH)_2$, in dichloromethane at room temperature gave a red-orange solution of the compound $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1), which was isolated as an orange-red microcrystalline solid in good yield. The synthesis of 1 can be also carried out in similar yields starting from $[Rh(\mu-OMe)(cod)]_2$ or [Rh(acac)(cod)] (1:2 molar ratio), although addition of an external base (NEt₃) is convenient for the later compound to drive the reaction to completion. The related iridium dinuclear compound $[Ir_2(\mu-S_2CBn_2)(cod)_2]$ (3) was obtained as a purple microcrystalline solid in good yield from $Bn_2C(SH)_2$ and [Ir(acac)(cod)] (1:2 molar ratio), although without the need of an external base, probably because of the low solubility of the complex in the reaction media (Scheme 1).

The synthetic flexibility observed in the case of Bn₂C(SH)₂ is not completely applicable to the preparation of dinuclear complexes with 2,4-dimethyl-3,3-dimercaptopentane, ^{*i*}Pr₂C-(SH)₂, as the synthesis is generally problematic due to side reactions. Thus, following a similar synthetic protocol, reaction of ^{*i*}Pr₂C(SH)₂ with [Rh(acac)(cod)] (1:2 molar ratio) in dichloromethane gave a deep-red solution from which the trinuclear hydride cluster [Rh₃(μ_3 -S)₂(μ -H)(cod)₃]^{33a} was isolated in 55% yield (Scheme 2). It is known that some *gem*-dithiol compounds are transformed into the corresponding thione by SH₂(g) elimination.³⁴ Thus, the formation of the cluster is probably driven by the SH₂(g) formed by the

⁽³⁰⁾ SAINTPLUS, ver. 6.28; Bruker AXS: Madison, WI; 2001.

⁽³¹⁾ Blessing, R. H. Acta Crystallogr. 1995, A51, 33; implemented in SADABS: Area-Detector Absorption Correction, ver. 2.03, 2002.

⁽³²⁾ SHELXTL Package, ver. 6.10; Bruker-AXS: Madison, WI, 2000; Sheldrick G.M. SHELXS-86 and SHELXL-97, 1997.

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Rivas et al.

Scheme 1. Synthesis of Rhodium and Iridium *gem*-Dithiolato-Bridged Dinuclear Diolefin Complexes



Scheme 2. Formation of a Trirhodium Hydride Cluster by Deprotonation of ${}^{\prime}Pr_{2}C(SH)_{2}$



decomposition of ${}^{i}Pr_{2}C(SH)_{2}$ mediated by [Rh(acac)(cod)]. This interpretation is sustained together by the detection in the reaction media of the thione ${}^{i}Pr_{2}C=S$ (GC/MS evidence) and by the clean formation of [Rh₃(μ_{3} -S)₂(μ -H)(cod)₃] by reaction of [Rh(acac)(cod)] with SH₂(g).

Nevertheless, the formation of the hydride clusters $[M_3(\mu_3-S)_2(\mu-H)(cod)_3]$ (M = Rh, Ir)³³ is minimized by using rhodium and iridium dinuclear compounds as starting materials. Thus, reaction of ${}^{i}Pr_2C(SH)_2$ with $[Rh(\mu-OH)(cod)]_2$ or $[Ir(\mu-OMe)(cod)]_2$ afforded the compounds $[Rh_2(\mu-S_2C^iPr_2)(cod)_2]$ (2) and $[Ir_2(\mu-S_2C^iPr_2)(cod)_2]$ (4), which were isolated as orange and purple microcrystalline solids in 76% yield after chromatography purification to remove the hydride clusters (Scheme 1).

1–4 have been fully characterized by elemental analysis, FAB mass spectra, and multinuclear NMR spectroscopy. In addition, the structures of 1 and 2 have been determined by X-ray diffraction methods (Figures 1 and 2), showing the bridging and chelating coordination mode $(1:2\kappa^2 S, 1:2\kappa^2 S')$ of the *gem*-dithiolato ligands. The ¹H NMR of the compounds showed no resonances attributable to SH protons, indicating the complete deprotonation of the *gem*-dithiol compounds upon the formation of the complexes. The ¹H and ¹³C{¹H} NMR showed rigid structures that are in agreement with the dinuclear structures of $C_{2\nu}$ symmetry found in the solid. In particular, the ¹H NMR spectra showed



Figure 1. Molecular structure of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1).



Figure 2. Molecular structure of $[Rh_2(\mu-S_2C'Pr_2)(cod)_2]$ (2).

the expected two sharp resonances for the =CH protons and four sharp multiplets for the >CH₂ protons as a consequence of the differentiation between the two groups of exo and endo protons of the equivalent cod ligands. In accordance with the chemical equivalence of both benzyl and isopropyl fragments of the gem-dithiolato ligands in the structures, the benzylic and isopropylic protons were observed as a sharp singlet in 1 and 3 and as the expected septuplet and doublet (1:6 ratio, ${}^{3}J_{\rm H-H} \approx 6.9$ Hz) in 2 and 4, respectively. The formation of the dimetallic core in 1-4 strongly influences the chemical shift of the resonance of the geminal carbon atoms of the bridging *gem*-dithiolato ligands in the ${}^{13}C{}^{1}H$ NMR spectra. This resonance is shifted to higher frequencies, compared to the free gem-dithiols, by about 30 ppm in the rhodium (1 and 2) and 40 ppm in the iridium ones (3 and 4) and reflects the effect of the ligand coordination to the metal centers.

Stepwise Formation of Dinuclear gem-Dithiolato-Bridged Compounds. The monitoring of the reaction of [Rh(acac)(cod)] with Bn₂C(SH)₂ in CDCl₃ by ¹H NMR spectroscopy showed the exclusive formation of the dinuclear compound [Rh₂(μ -S₂CBn₂)(cod)₂] (1) independently of the utilized molar ratio (1:1 or 2:1) as a consequence of the simultaneous deprotonation of both –SH groups of

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Scheme 3. Stepwise Formation of the *gem*-Dithiolato-Bridged Dinuclear $\mathbf{6}$





The reaction of 1,1-dimercaptopentane, Cptn(SH)₂, with [Rh(acac)(cod)] (1:2 molar ratio) and NEt₃ in dichlorometane gave a brown solution from which a brown solid was isolated. The ¹H NMR analysis of this solid in CDCl₃ evidenced the formation of the dinuclear compound $[Rh_2(\mu S_2Cptn(cod)_2$ (6) together with the mononuclear [Rh(HS₂Cptn)(cod)] (5), which was present in, roughly, 20%. However, when the reaction was conducted in a 1:1 molar ratio without NEt₃ under the same conditions, the composition of the mixture was reversed, and the resulting dark-red solution showed almost 85% of the mononuclear species (Scheme 3). In fact, the mononuclear compound [Rh(HS₂Cptn)(cod)] (5) was isolated as a dark-red solid in 68% yield after recrystallization of the crude obtained following this procedure. The dinuclear compound $[Rh_2(\mu S_2Cptn(cod)_2$ (6) has been obtained as an orange solid in 62% yield from $[Rh(\mu-OH)(cod)]_2$ and $Cptn(SH)_2$ after purification by chromatography to remove 5 that is formed in trace amounts. Interestingly, the deprotonation of $[Rh(HS_2Cptn)(cod)]$ (5) can be accomplished by $[Rh(\mu OH)(cod)]_2$ in dichloromethane to give the dinuclear compound $[Rh_2(\mu-S_2Cptn)(cod)_2]$ (6) in good yield (Scheme 3).

The dinuclear formulation of **6** has been corroborated by an X-ray diffraction study (Figure 3). In addition, the NMR spectroscopic data of **6** closely resemble those of the previously described dinuclear compounds and do not deserve further comments. **5** has been characterized by elemental analysis, mass spectrum, and NMR spectroscopy as a mononuclear species containing a monodeprotonated Cptn(SH)₂ ligand with an unusual $\kappa^2 S, S'$ coordination mode. The FAB mass spectrum showed the molecular ion at m/z343 together with the ion at m/z 554 corresponding to the dinuclear species. This ion most likely results from the recombination of the produced ions because no mononuclear fragments were observed in the FAB mass spectrum of **6**. In contrast with **6**, which is rigid at room temperature, **5** is fluxional and the ¹H NMR showed average resonances for



Figure 3. Molecular structure of $[Rh_2(\mu-S_2Cptn)(cod)_2]$ (6).

the cod protons. Thus, the =CH and >CH₂ protons were observed as broad and featureless resonances at 4.18 ppm and at 2.39 and 1.93 ppm, respectively. The resonance of the SH proton probably is also broad and is hidden under the resonances of the cod ligand. The ¹³C{¹H} NMR spectrum showed the full equivalence of the =CH and >CH₂ carbons, which were observed at 81.7 ppm (d, J_{Rh-C} = 11 Hz) and at 31.6 ppm. The resonance of the geminal carbon atom in **5** was observed at 66.8 ppm and, as expected, is shifted to higher frequency compared to the resonance of Cptn(SH)₂ (23.6 ppm). As could be anticipated, this effect is more pronounced in **6** where this resonance was observed at 81.9 ppm and is the result of the (1:2 $\kappa^2 S$, 1:2 $\kappa^2 S'$) coordination mode of the doubly deprotonated Cptn(SH)₂ ligand.

The spectroscopic information obtained for [Rh(HS₂-Cptn)(cod)] (**5**) strongly suggests the existence of a dynamic behavior responsible for the chemical equivalence of the =CH protons and carbons of the cod ligand. Assuming a planar or a dynamic puckering four-membered metallacycle,³⁵ the intramolecular shift of the SH proton between both S-donor atoms together with the inversion of the sulfur atom bearing the mercapto group would account for the spectroscopic observations. A comparable dynamic processs involving the Au(PPh₃)⁺ fragment, that is isolobal with H⁺, has been observed in the early late heterotrimetallic compound [Cp^{tt}₂Zr(μ -S)₂{Ir(CO)₂}{Au(PPh₃)}] that contains a [Zr(μ -S)₂Ir] core.³⁶

Molecular Structures of $[Rh_2(\mu-S_2CBn_2)(cod)_2]$ (1), $[Rh_2(\mu-S_2CPr_2)(cod)_2]$ (2), and $[Rh_2(\mu-S_2Cptn)(cod)_2]$ (6). The molecular structures of 1, 2, and 6 are shown in Figures 1, 2, and 3, respectively, and their selected bond lengths and angles are all deliberately arranged in Table 1. In all of the three complexes, two Rh(cod) moieties are connected through a *gem*-dithiolato μ -S₂CR₂ double bridge,

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⁽³⁶⁾ Hernandez-Gruel, M. A. F.; Lahoz, F. J.; Oro, L. A.; Pérez-Torrente, J. J. Organometallics 2007, 26, 6437.

 Table 1. Crystal Data, Data Collection, and Refinement Parameters for 1, 2, 6, and 16

	1	2	6	16
empirical formula	C31H38Rh2S2	C23H38Rh2S2	$C_{21}H_{32}Rh_2S_2$	$C_{45}H_{42}O_2P_2Rh_2S_2 \cdot CH_2Cl_2$
cryst size, mm	$0.17 \times 0.09 \times 0.06$	$0.38 \times 0.11 \times 0.02$	$0.34 \times 0.34 \times 0.34$	$0.37 \times 0.23 \times 0.22$
fw	680.55	584.47	554.41	1031.59
space group	$P2_1/n$ (No. 14)	<i>P</i> 1̄ (No. 2)	<i>P</i> 1 (No. 2)	$P2_1/n$ (No. 14)
<i>a</i> , Å	6.3798(5)	6.4559(4)	7.5580(5)	17.6034(12)
<i>b</i> , Å	20.1991(16)	10.8020(6)	9.9802(6)	10.6870(7)
<i>c</i> , Å	21.1670(17)	16.8466(10)	15.1423(9)	24.2161(17)
α, deg	90.0	99.427(1)	108.7250(10)	90.0
β , deg	97.660(2)	91.151(1)	92.1170(10)	107.7410(10)
γ, deg	90.0	104.990(1)	108.5870(10)	90.0
$V, Å^3/Z$	2703.4(4)/4	1117.11(11)/2	1012.90(11)/2	4339.1(5)/4
$D_{\text{calcd}}, \text{g} \cdot \text{cm}^{-3}$	1.672	1.738	1.818	1.579
μ , mm ⁻¹	1.394	1.671	1.837	1.092
min., max. transm. factors	0.787, 0.919	0.566, 0.964	0.537, 0.542	0.685, 0.792
no. of measd reflns	$17795\ (1.40 \le \theta \le 28.61)$	$13859\ (1.98 \le \theta \le 28.34)$	$12\ 332\ (2.24 \le \theta \le 28.29)$	$51934\ (1.71 \le \theta \le 28.31)$
no. of unique reflns	$6347 \ (R_{\rm int} = 0.1029)$	5274 ($R_{\rm int} = 0.0252$)	4718 ($R_{\rm int} = 0.0132$)	$10466\ (R_{\rm int}=0.0286)$
no. of data/restraints/params	6347/0/324	5274/0/396	4718/0/354	10 466/0/681
GOF (all data) ^{a}	0.927	1.026	1.103	1.056
R1(F) (only for $F^2 > 2\sigma(F^2))^b$	0.0487 (4572 refl.)	0.0231 (4699 refl.)	0.0160 (4539 refl.)	0.0257 (9397 refl.)
$wR2(F^2)$ (all data) ^c	0.1005	0.0511	0.0398	0.0603

^{*a*} GOF = $(\sum [w(F_o^2 - F_c^2)^2]/(n - p))^{1/2}$, where *n* and *p* are the number of data and parameters. ^{*b*} R1(*F*) = $\sum ||F_o| - |F_o|/\sum |F_o|$ only for observed reflections (in parentheses). ^{*c*} wR2(*F*²) = $(\sum [w(F_o^2 - F_c^2)^2]/\sum [w(F_o^2)^2])^{1/2}$ where $w = 1/[\sigma^2(F_o^2) + (aP)^2]$ and $P = [\max(0, F_o^2) + 2F_c^2]/3$.

with the sulfur atoms showing a $(1:2\kappa^2 S, 1:2\kappa^2 S')$ coordination mode. The major differences among the three molecular structures merely concern the distinct *R* substituents of the dithiolato ligands: two benzyl groups in 1, two *iso*-propyl substituents in 2, and a cyclic four-membered saturated ring of carbon atoms in **6**.

The coordination geometry around the rhodium centers is in all cases distorted square-planar; the distortions fundamentally arise from the chelating behavior of the two ligands, cod, and μ -S₂CR₂, with the S(1)-Rh-S(2) bond angles in the range of $70.40(4) - 72.017(13)^{\circ}$ and the interolefinic angles (between midpoints of coordinated double bonds) in the range $87.44(16) - 88.31(5)^{\circ}$ (Table 1). The dihedral angle between the two metal coordination planes are rather similar $(91.32(6) (1), 92.75(3) (2) \text{ and } 93.59(2)^{\circ} (6))$ and the metal-metal separations are almost unaffected by the change in the thiolate substituent (2.8544(5) in 1, 2.8674(3) in 2, 3674(3) in 2)and 2.8608(2) Å in 6). As previously observed for other thiolate bridged Rh(I) complexes, both rhodium atoms are slightly displaced out of the coordination mean-plane toward the external part of the molecule (range 0.0197(4) - 0.1263(4)Å), proving the existence of a weak repulsion between the metal atoms, most likely due to the ligand-forced short metal-metal nonbonding distance;^{23d,25f,26} this fact could be also substantiated by the values of the torsion angle $Rh(1)-S(1)\cdots$ S(2)-Rh(2) (94.51(4) in 1, 96.50(2) in 2, and 96.22(1)° in **6**)) that, in all cases, are slightly larger (around 3°) than the dihedral angle between coordination planes (see above).

If compared with other related $(\mu$ -SR)₂ or $(\mu$ -SRS) dithiolato complexes, the presence of these *gem*-dithiolato ligands originates a feeble augment of the Rh–S–Rh bond angle (mean 74.12(1)°), but a significant decrease in the S–Rh–S bond angles (71.30(1)°); thus, values reported for closely related dinuclear dithiolato complexes [Rh₂(μ -SR)₂(cod)₂] are (Rh–S–Rh and S–Rh–S mean values, respectively): 77.75 and 75.90° (R = Me),³⁷ 75.68 and 88.83°

 $(R = C_6F_5)$,³⁸ 76.32 and 87.37° $(R = C_6F_4CF_3)$,³⁹ or 74.99 and 79.01° $(R_2 = -CH_2CH_2-)$,^{23d} respectively. These structural modifications—fundamentally those affecting the S-Rh-S bond angles—are also coupled with changes in the dihedral angles between the metal coordination planes (as evidenced by the RhS₂Rh torsion angles: 105.49° (R =Me),³⁷ 118.37° $(R = C_6F_5)$,³⁸ 117.39 $(R = C_6F_4CF_3)$,³⁹ or 104.16° $(R_2 = -CH_2CH_2-)^{23d}$) and by the existence of shorter intermetallic distances.

All the cycloocta-1,5-diene molecules in **1**, **2**, and **6** exhibit the classical tub conformation, showing analogous Rh–C and olefinic C–C bond distances to those observed in the above referred dithiolato complexes, reflecting the electronic similarities between *gem* and general dithiolato ligands. Also, the Rh–S bond distances observed (Table 1) are in the central region of the separations detected in related dinuclear dithiolato complexes [Rh₂(μ -SR)₂(cod)₂] (range 2.289–2.487, mean 2.38(3) Å).⁴⁰

Synthesis and Characterization of Carbonyl gem-Dithiolato-Bridged Rhodium Dinuclear Compounds. The dinuclear framework of the gem-dithiolato-bridged rhodium compounds is sustained in carbonylation reactions. The carbonylation of diolefin rhodium **1**, **2**, and **6** at room temperature in dichloromethane afforded colored solutions of the corresponding tetracarbonyl complexes $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (**7**), $[Rh_2(\mu-S_2C'Pr_2)(CO)_4]$ (**8**), and $[Rh_2(\mu-S_2Cptn)(CO)_4]$ (**9**). However, the presence of cod in the reaction media, the partial reversibility of the carbonylation processes, and the relative instability of these complexes make their isolation difficult.

Attempts to prepare the carbonyl complexes directly from *gem*-dithiol compounds using carbonyl starting materials, for example [Rh(acac)(CO)₂], were unsuccessful. Fortunately,

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Table 2. S	Selected Bond	Distances	(Å)	and	Angles	(deg)	for	1, 2,	and	6
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	1	2	6		1	2	6
Rh(1)•••Rh(2)	2.8544(5)	2.8674(3)	2.8608(2)	Rh(2) - S(1)	2.3748(12)	2.3633(6)	2.3627(4)
Rh(1) - S(1)	2.3979(12)	2.3514(5)	2.3690(4)	Rh(2) - S(2)	2.3650(12)	2.3581(5)	2.3618(4)
Rh(1) - S(2)	2.4078(11)	2.3819(6)	2.3896(4)	Rh(2)-C(9)	2.129(4)	2.126(2)	2.1367(16)
Rh(1) - C(1)	2.137(5)	2.142(2)	2.1406(15)	Rh(2) - C(10)	2.155(4)	2.157(2)	2.1419(16)
Rh(1) - C(2)	2.142(4)	2.143(2)	2.1449(16)	Rh(2)-C(13)	2.129(4)	2.138(2)	2.1411(15)
Rh(1) - C(5)	2.125(4)	2.119(2)	2.1358(16)	Rh(2) - C(14)	2.122(5)	2.135(2)	2.1334(16)
Rh(1) - C(6)	2.135(5)	2.141(2)	2.1444(16)	S(2)-C(17)	1.867(4)	1.875(2)	1.8640(16)
S(1)-C(17)	1.847(4)	1.870(2)	1.8613(16)	C(9)-C(10)	1.387(6)	1.389(3)	1.398(2)
C(1) - C(2)	1.378(7)	1.392(3)	1.397(2)	C(13)-C(14)	1.386(7)	1.394(3)	1.398(2)
C(5)-C(6)	1.373(7)	1.390(3)	1.400(2)	$C(17) - C(19)^{b}$	1.544(6)	1.559(3)	1.549(2)
C(17)-C(18)	1.541(6)	1.550(3)	1.543(2)				
S(1) - Rh(1) - S(2)	70.40(4)	71.122(19)	71.420(13)	S(1) - Rh(2) - S(2)	71.53(4)	71.331(19)	72.017(13)
S(1) - Rh(1) - M(1)	172.17(12)	172.95(5)	172.39(4)	S(1) - Rh(2) - M(3)	170.81(12)	170.87(5)	172.33(4)
S(1) - Rh(1) - M(2)	100.14(12)	98.73(5)	99.10(4)	S(1) - Rh(2) - M(4)	99.51(12)	99.91(5)	99.70(4)
S(2) - Rh(1) - M(1)	101.83(11)	102.07(5)	101.11(4)	S(2) - Rh(2) - M(3)	100.75(11)	100.67(5)	100.46(3)
S(2) - Rh(1) - M(2)	170.49(11)	169.21(5)	170.33(4)	S(2) - Rh(2) - M(4)	168.36(11)	170.89(5)	169.28(3)
M(1) - Rh(1) - M(2)	87.61(15)	87.94(7)	88.31(5)	M(3) - Rh(2) - M(4)	87.44(16)	87.85(7)	87.95(5)
Rh(1) - S(1) - Rh(2)	73.46(3)	74.916(17)	74.399(12)	Rh(1) - S(2) - Rh(2)	73.45(3)	74.445(16)	74.038(12)
Rh(1) - S(1) - C(17)	84.55(13)	86.65(7)	84.66(5)	Rh(1)-S(2)-C(17)	83.85(13)	85.66(7)	84.02(5)
Rh(2) - S(1) - C(17)	88.23(13)	86.80(7)	86.86(5)	Rh(2) - S(2) - C(17)	88.06(14)	86.84(7)	86.83(5)
S(1) - C(17) - S(2)	96.47(19)	94.65(10)	96.42(7)	$C(18) - C(17) - C(19)^{b}$	108.5(3)	111.23(17)	105.48(13)
S(1) - C(17) - C(18)	113.4(3)	110.12(14)	112.84(11)	S(2)-C(17)-C(18)	109.4(3)	110.03(14)	113.49(11)
$S(1)-C(17)-C(19)^{b}$	115.4(3)	114.61(14)	115.42(11)	$S(2) - C(17) - C(19)^{b}$	113.2(3)	115.12(15)	113.41(11)

^{*a*} M(1), M(2), M(3), and M(4) represent the midpoints of the olefinic C(1)–C(2), C(5)–C(6), C(9)–C(10), and C(13)–C(14) bonds. ^{*b*} In **6**, the atom label of carbon atom bonded to C(17), analogous to C(19) in **1** and **2**, corresponds to C(21).

the low solubility of $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) in *n*-hexane allowed their isolation under a carbon monoxide atmosphere as a red microcrystalline solid in excellent yield. The dinuclear formulation of **7** relies on the FAB+ spectra that showed the molecular ion at m/z 576 and the ions resulting from the sequential losses of four carbonyl ligands. The ¹H and ¹³C{¹H} NMR spectra in CDCl₃ were in agreement with a C_{2v} symmetry structure resulting from the replacement of a cod ligand by two carbonyls on each rhodium center. Thus, the equivalent carbonyl ligands were observed as a doublet at 184.4 ppm ($J_{Rh-C} = 72$ Hz) in the ¹³C{¹H} NMR spectrum and the benzylic protons of the *gem*-dithiolato ligand as a sharp singlet at 3.31 ppm in the ¹H NMR spectrum.

The IR spectrum of **7** in solution (CH₂Cl₂) showed three ν (CO) bands for the terminal carbonyl groups at 2078 (m), 2054 (s), and 2010 (s) cm⁻¹. The same pattern has also been observed in the IR of the in situ generated solutions of the complexes [Rh₂(μ -S₂CⁱPr₂)(CO)₄] (**8**) and [Rh₂(μ -S₂Cptn)(CO)₄] (**9**) and [Rh₂(μ -S₂Chxn)(CO)₄]²⁶ (Table 2), indicating that the formation of dinuclear tetracarbonyl complexes from the parent olefin complexes is clean and quantitative. The observed pattern of intensities (m, s, s) is typical for tetracarbonyl dinuclear complexes having both thiolato and dithiolato bridging ligands.^{21a,24d,41}

Replacement Reactions on [Rh_2(\mu-S_2CR_2)(CO)_4] by P-Donor Ligands. The carbonyl replacement reactions by P-donor ligands on the complexes $[Rh_2(\mu-S_2CR_2)(CO)_4]$ takes place quickly at room temperature and stops at the disub-

Scheme 4. Replacement Reactions on *gem*-Dithiolato-Bridged Dinuclear Complexes



stituted complexes $[Rh_2(\mu-S_2CR_2)(CO)_2(PR'_3)_2]$ in the case of phosphine ligands $(PR'_3 = PPh_3 \text{ or } PCy_3)$. Thus, the addition of 2 mol equiv of PPh₃ to solutions of tetracarbonyl 7 and 8, generated in situ by carbonylation of the corresponding diolefin complexes, afforded orange solutions of the complexes $[Rh_2(\mu-S_2CBn_2)(CO)_2(PPh_3)_2]$ (10) and $[Rh_2(\mu-S_2C^2Pr_2)(CO)_2(PPh_3)_2]$ (11), which were isolated as orangered solids in excellent yields. 10 can be alternatively prepared in similar yield by reaction of $Bn_2C(SH)_2$ with $[Rh(acac)-(CO)(PPh_3)]$ (1:2 molar ratio) (Scheme 4). This synthetic approach is also the more convenient for the synthesis of the complexes $[Rh_2(\mu-S_2Cptn)(CO)_2(PPh_3)_2]$ (12) and $[Rh_2(\mu-S_2CBn_2)(CO)_2(PCy_3)_2]$ (13), which were obtained as orange

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Figure 4. (a) ${}^{31}P{}^{1}H$ NMR (CDCl₃, 293 K) spectrum of [Rh₂(μ -S₂Cptn)(CO)₂(PPh₃)₂] (**12**). (b) Calculated ${}^{31}P{}^{1}H$ NMR spectrum for the AA'XX' spin system observed for the trans isomer.

Table 3. ν (CO) Stretching Frequencies (cm⁻¹) Observed in the IR Spectra in Solution (CH₂Cl₂) of Carbonyl **7–9**

complex	ν(CO)	color
$[Rh_2(\mu-S_2Chxn)(CO)_4]^a$	2082 (m), 2058 (s), 2013 (s) cm ⁻¹	red-brown
$[Rh_2(\mu-S_2CBn_2)(CO)_4] (7)$	2078 (m), 2054 (s), 2010 (s) cm ⁻¹	red
$[Rh_2(\mu-S_2C^iPr_2)(CO)_4]$ (8)	2080 (m), 2055 (s), 2011 (s) cm ⁻¹	yellow-brown
$[Rh_2(\mu-S_2Cptn)(CO)_4]$ (9)	2082 (m), 2059 (s), 2014 (s) cm ⁻¹	yellow-brown
^a Ref 26.		

and yellow microcrystalline solids in good yield straightforwardly from the corresponding gem-dithiol and the appropriate [Rh(acac)(CO)(PR'₃)] complexes. The dinuclear formulation of 10-13 relies on the FAB + spectra, which showed the molecular ions and the sequential losses of two carbonyl ligands. The complexes exist as the trans isomer $(C_2 \text{ symmetry})$, although small amounts of the cis isomer (C_s symmetry) (<3%) were observed in the ³¹P{¹H} NMR spectra of those complexes containing the less-steric demanding PPh₃ ligands (10-12). The trans isomer showed a distinctive complex resonance in the ³¹P{¹H} NMR spectra corresponding to an AA'XX' spin system (Figure 4).⁴² The observed signals correlate well with the calculated spectra using the parameters reported in Table 3. Interestingly, each phosphorus atom is coupled to both rhodium atoms with a standard J_{Rh-P} and a negative small ${}^{2}J_{Rh-P}$ coupling constants. In addition, the observed ${}^{3}J_{P-P}$ (≈ 6 Hz) and J_{Rh-Rh} (3–4 Hz) coupling constants are in good agreement with a trans disposition of the PR'₃ ligands and the expected short rhodium-rhodium bond distances in the dinuclear complexes. Salient features in the ¹H NMR of **10** and **13** are the AB quartet ($J_{AB} \approx 14$ Hz), corresponding to the diastereotopic >CH₂ protons of the benzylic fragment in agreement with the C_2 symmetry of the compounds. Similarly, two doublet resonances were observed for the diastereotopic methyl groups of the isopropyl fragments in **11**. The equivalent carbonyl ligands in **10–13** were observed as a doublet of doublets (dd) resonance in the ¹³C{¹H} NMR spectra, with J_{Rh-C} and ² J_{P-C} couplings of \approx 75 and 18 Hz, respectively. Further evidence for the trans disposition of the CO ligands in the structures comes from the observation of a strong ν (CO) absorption in the IR of the complexes in solution.^{41b,43}

In contrast with 10-13, which do not react further with PR'_{3} (PPh₃ or PCy₃), the reaction of complexes $[Rh_{2}(\mu S_2CR_2$ (CO)₄ with phosphite ligands, P(OMe)₃ and P(OPh)₃, goes further. The reaction of $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) with $P(OMe)_3$ has been studied in detail by ${}^{31}P{}^{1}H{}$ NMR (C₆D₆) and four species $[Rh_2(\mu-S_2CR_2)(CO)_{4-n}{P(OR)_3}_n]$ (Pn, n =1, 4) resulting from the partial or total replacement of the CO ligands have been identified depending on the $P(OR)_3$ / [Rh₂] molar ratio (Table 4). The reaction of 7 with 1 mol equiv of P(OMe)₃ gave both the mono- and the disubstituted complexes, P1 and P2. Interestingly, the reaction with 2 mol equiv of P(OMe)₃ gave predominantly the disubstituted P2 (mainly the trans isomer), although a small amount of the monosubstituted P1 was still observed (Figure 5). Increasing the $P(OR)_3/7$ molar ratio to 3.5 resulted in the formation almost exclusively of P3 and P4 (Figure 6). The molecular ions of both species have been detected in the FAB+ spectrum of the resulting solution at m/z 836 (P3) and 960 (P4), respectively. Interestingly, the integrity of the dinuclear unit is preserved when the $P(OR)_3/7$ molar ratio was further increased because the formation of the cationic mononuclear species $[Rh{P(OMe)_3}_4]^+$ was not detected.⁴⁴

The attempt of isolation of P4 in the solid state was unsuccessful because the compound was obtained as a yellow-orange oil. However, the lack of formation of P3 when the $P(OR)_3/7$ molar ratio was adjusted to 2 suggests that the isolation of the P2 complexes could be possible. In fact, the complexes $[Rh_2(\mu-S_2CBn_2)(CO)_2\{P(OMe)_3\}_2]$ (14) and $[Rh_2(\mu-S_2CBn_2)(CO)_2\{P(OPh)_3\}_2]$ (15) have been obtained as light orange and yellow solids, respectively, in excellent yield. Both complexes have been fully characterized by elemental analysis, FAB+ mass spectra, and NMR. The ${}^{31}P{}^{1}H$ NMR spectra showed that the compounds exist mainly as the trans isomer, although the cis isomer is also present in less than 5%. The resonance of the trans isomer of both complexes closely resembles those of the related phosphine 10-13 and has been simulated with the same AA'XX' spin system using the parameters of Table 3. Noteworthy, similar ${}^{2}J_{Rh-P}$ and J_{Rh-Rh} but larger ${}^{3}J_{P-P}$ (≈ 15 Hz) coupling constants have been observed.

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Table 4. ³¹P{¹H} NMR (121.47 MHz, 293 K) Data for the Compounds $[Rh_2(\mu-S_2CR_2)(CO)_2(PR'_3)_2]^a$

compound	isomer	δ	$J_{ m Rh-P}$	${}^{2}J_{\mathrm{Rh-P}}$	${}^{3}J_{\mathrm{P-P}}$	$J_{ m Rh-Rh}$
$[Rh_2(\mu-S_2CBn_2)(CO)_2(PPh_3)_2]$ (10) ^b	trans	41.59	163.70	-1.46	6.22	3.12
	cis	39.60	163			
$[Rh_2(\mu-S_2C^iPr_2)(CO)_2(PPh_3)_2]$ (11) ^b	trans	41.76	163.10	-1.80	6.45	3.42
	cis	40.20	163			
$[Rh_2(\mu-S_2Cptn)(CO)_2(PPh_3)_2]$ (12) ^c	trans	41.37	163.50	-1.23	6.19	3.58
	cis	39.00	162			
$[Rh_2(\mu-S_2CBn_2)(CO)_2(PCy_3)_2]$ (13) ^c	trans	51.56	156.88	-0.76	4.29	3.33
$[Rh_2(\mu-S_2CBn_2)(CO)_2\{P(OMe)_3\}_2]$ (14) ^b	trans	139.96	258.07	-1.95	14.00	2.89
	cis	141.00	252			
$[Rh_2(\mu-S_2CBn_2)(CO)_2\{P(OPh)_3\}_2]$ (15) ^c	trans	122.08	276.82	-2.00	15.94	2.63
	cis	123.30	273			

^{*a*} The data of the trans isomers correspond to the calculated spectrum: AA'XX' spin system, $A = {}^{31}P$ and $X = {}^{103}Rh$ (δ in ppm and J in Hz). ^{*b*} In C₆D₆. ^{*c*} In CDCl₃.



Figure 5. ${}^{31}P{}^{1}H$ NMR (C₆D₆) of the reaction of [Rh₂(μ -S₂CBn₂)(CO)₄] (7) with 2 mol equiv of P(OMe)₃ showing exclusively the presence of [Rh₂(μ -S₂CBn₂)(CO)₂{P(OMe)₃}₂] (P2, cis and trans isomers, **14**) and [Rh₂(μ -S₂CBn₂)(CO)₃{P(OMe)₃}] (P1).



Figure 6. ³¹P{¹H} NMR (C₆D₆) of the reaction of $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) with 3.5 mol equiv of P(OMe)₃ showing the presence of $[Rh_2(\mu-S_2CBn_2)\{P(OMe)_3\}_4]$ (P4), $[Rh_2(\mu-S_2CBn_2)(CO)\{P(OMe)_3\}_3]$ (P3), and traces of P2.

In contrast with monodentate P-donor ligands, that mainly give rise to trans disubstituted complexes, the replacement reactions on $[Rh_2(\mu-S_2CBn_2)(CO)_4]$ (7) using diphosphines with a large bite angle, for example dppp (1,3-bis(diphenylphosphino)propane) or dppb (1,4-bis(diphenylphosphi-



Figure 7. Molecular structure of $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-dppb)]$ (16). no)butane), resulted in the formation of cis disubstituted complexes. The reaction of 7 with 1 mol equiv of dppb gave a yellow solution of the compound $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-S_2CBn_2)(\mu-S_2CBn_2)(CO)_2(\mu-S_2CBn_2)(\mu-S_2)(\mu-S_2)(\mu-S_2)(\mu-S_2)(\mu-S_2)(\mu-S_2)(\mu-S_2)$ dppb)] (16), which was isolated as a yellow solid in good yield. The structure of 16 has been determined by X-ray diffraction methods and is shown in Figure 7. The spectroscopic data obtained from the IR and NMR are in agreement with an approximate C_s molecular symmetry as a result of the coordination of the diphosphine in a cis fashion. The diastereotopic >CH₂ protons of the benzyl groups of the bridging gem-dithiolato ligand are isochronus in C₆D₆ and gave a single resonance at 3.94 ppm in the ¹H NMR spectrum. However, the spectrum in CDCl₃ showed the expected AB quarted centered at 3.56 ppm ($J_{AB} \approx 14$ Hz). The ³¹P{¹H} NMR spectrum features a slightly broad doublet with no fine structure centered at 32.25 ppm ($J_{\rm Rh-P} = 157$ Hz). Finally, the equivalent carbonyl ligands were observed as a broad absorption at 1980 cm⁻¹ in the IR spectrum in dichloromethane.

The related compound $[Rh_2(\mu-Bn_2CS_2)(CO)_2(\mu-dppp)]$ has also been identified in the reaction of **7** with dppp (27.87 ppm, d, $J_{Rh-P} = 170$ Hz) together with other minor unidentified species that could not be easily separated by recrystallization. However, the observed J_{Rh-P} coupling constants for these species (170–150 Hz) ruled out the formation of cationic compounds such as $[Rh(dppp)_2]^+$ or $[Rh(CO)(dppp)_2]^+$, potential products of the degradation of the dinuclear structure.⁴⁵

Table 5. ${}^{31}P{}^{1}H{}$ NMR Data (C₆D₆) for the Pn Compounds [Rh₂(μ -S₂CBn₂)(CO)_{4-n}{P(OMe)₃}_n] (n Denotes the Number of P(OMe)₃ Ligands in the Dinuclear Framework)^{*a*}

compound	δ	J (Hz)
P1	138.7	$J_{\rm Rh-P} = 256$
$P2^c$	139.96 (trans isomer) ^b	$J_{\rm Rh-P} = 258.07, {}^{2}J_{\rm Rh-P} = -1.95,$
		${}^{3}J_{\rm P-P} = 14.0, J_{\rm Rh-Rh} = 2.89$
	141.0 (cis isomer)	$J_{\rm Rh-P} = 252$
$P3^b$	146.1 (P _a), 145.3 (P _b),	$J_{\rm Rh1-Pa} = 275, J_{\rm Rh2-Pb} = 275,$
	142.2 (P _c)	$J_{\rm Rh2-Pc} = 258, J_{\rm Pb-Pc} = 10.4,$
		$J_{\rm Pa-Pc} = 3$
P4	146.6	$J_{\rm Rh-P} = 272$

 a For the labeling scheme used in P3, see Figure 6. b Calculated spectrum. c P2 is actually 14.

Molecular Structure of [Rh₂(µ-S₂CBn₂)(CO)₂(µ-dppb)] (16). A molecular diagram of this dinuclear complex is shown in Figure 7, and key bond lengths and angles are collected in Table 5. Like its diolefin analogues, 1, 2, and 6, complex 16 shows a folded Rh_2S_2 core, with both rhodium atoms in a distorted square-planar coordination. The greatest distortions from this square-planar geometry consist of the small S-Rh-S bond angles (71.047(16) and 71.248(16)°), although the sums of the four bond angles between pairs of cis ligands around each metal easily add up to 360.0°. The dihedral angle between the metal coordination planes is significantly larger (99.66(2)°) than those observed in the diolefinic analogues 1, 2, and 6 (range $91.32(6)-93.59(2)^{\circ}$); in accordance with this fact, the intermetallic separation elongates to 2.9514(3) Å. In this case, there are not structural features that could be indicative of an intermetallic repulsion, as the RhSSRh torsion angle detected is very similar $(98.60(2)^{\circ})$ to the interplanar angle between metal coordination planes.

The diphosphine ligand also bridges the two metals in **16** occupying relative cisoidal positions. Interestingly, within both metal coordination planes the larger cis angles are those between the dithiolato (S(2)) and the carbonyl groups (101.30(6) and 98.92(6)°), most probably as a subtle consequence of the bridging behavior of the diphosphine ligand in the other side of the metal coordination. The whole structure resembles quite well that of the dithiolato complex $[Rh_2(\mu-S'Bu)_2(CO)_2(\mu-dppb)]$, in which two monodentate *tert*-butyl thiolate groups bridge the two metals of an identical $Rh_2(CO)_2(\mu-dppb)$ moiety.⁴⁶

It could be consider that the cis disposition of carbonyl groups is a consequence of the bidentate nature of the dppb ligand. However, a detailed search on related structurally characterized dithiolato analogues, of general stoichiometry $[Rh_2(\mu-SR)_2(CO)_2(PR'_3)_2]$, reveals that these dithiolatobridged rhodium(I) systems exhibit a preference toward a cis disposition of the carbonyl groups, not only when the phosphine ligands could restrict the molecular geometry (bidentate phosphines)^{46,47} but also when the PR₃ is monodentate and no geometric restrictions could be envisaged, these are the cases of $[Rh_2(\mu-SPy'Bu)_2(CO)_2(PPh_3)_2]$,^{21a} $[Rh_{2}(\mu-S'Bu)_{2}(CO)_{2}(PPh_{3})_{2}],^{48} [Rh_{2}\{\mu-SC_{6}H_{2}(i^{2}Pr)_{3}\}_{2}(CO)_{2}-(PPh_{3})_{2}],^{49} [Rh_{2}(\mu-SPh)_{2}(CO)_{2}(PMe_{3})_{2}],^{50} \text{ or } [Rh_{2}(\mu-S'Bu)_{2}(CO)_{2}(PPh_{2}CH_{2}CH_{2}NMe_{3})_{2}],^{51}$

As for the Rh–S bond distances, those trans to the CO ligands (2.3995 and 2.4001(5) Å) are slightly longer than those trans to the phosphines (2.3926 and 2.3801(5) Å), a phenomenon previously described in other dithiolato complexes.^{21a,50} The rest of the metrical parameters, both in the metal environments and within the ligands, are very similar to those of literature complexes that have been structurally characterized.^{21a,48–51}

Discussion

The synthesis of dinuclear $[M_2(\mu-S_2CR_2)(cod)_2]$ (M = Rh, Ir) complexes can be accomplished by direct deprotonation of gem-dithiol compounds with rhodium or iridium complexes that contain basic ligands. The diolefin complexes $[M_2(\mu-S_2CR_2)(cod)_2]$ (M = Rh, Ir) are accessible from the dinuclear complexes $[M(\mu-OH)(cod)]_2$ and $[M(\mu-OMe)-$ (cod)]2, or the mononuclear [M(acac)(cod)], sometimes in the presence of an external base. The dinuclear complexes are generally obtained in high yield although, in the case of $^{i}Pr_{2}C(SH)_{2}$, the syntheses are complicated by the formation of the trinuclear hydride clusters $[M_3(\mu_3-S)_2(\mu-H)(cod)_3]^{33}$ which can be easily separated by chromatography. The formation of the sulfido clusters resulted from the $SH_2(g)$ produced in the decomposition of ⁱPr₂C(SH)₂ as has been evidenced by the detection of ${}^{i}Pr_{2}C=S$ by GC/MS. The stepwise deprotonation of the gem-dithiol compounds has only been observed in the case of Cptn(SH)₂, and both the mononuclear $[Rh(HS_2Cptn)(cod)]$ (5) and the dinuclear $[Rh_2(\mu-S_2Cptn)(cod)_2]$ (6) compounds have been isolated and fully characterized.

This synthetic methodology is also applicable to the preparation of the complexes $[Rh_2(\mu-S_2CR_2)(CO)_2(PR'_3)_2]$, which can be also prepared from $[Rh(acac)(CO)(PR'_3)]$ (R' = Ph, Cy) but not to the carbonyl complexes $[Rh_2(\mu-S_2CR_2)(CO)_4]$. However, these complexes have been obtained by carbonylation of the corresponding diolefin compounds and are precursors for the disubstituted complexes $[Rh_2(\mu-S_2CR_2)(CO)_2(PR'_3)_2]$ after carbonyl replacement by P-donor ligands (R' = Ph, Cy, OMe, OPh).

The above-described results on the synthesis of *gem*dithiolato-bridged complexes $[Rh_2(\mu-S_2CR_2)(L_2)_2]$ are, in some aspects, comparable to those previously described in bis-thiolato, $[Rh(\mu-SR)(L_2)]_2$, or dithiolato, $[Rh_2(\mu-S(CH_2)_nS)(L_2)_2]$ (n = 2-4), complexes as they all have an open-book structure. However, significant differences are observed in nuclearity, stereochemistry and reactivity that are probably a consequence of the special geometrical constrains imparted by the *gem*-dithiolato ligand in the M₂S₂ core. The bridging and chelating coordination mode (1:2 $\kappa^2 S$,

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 $1:2\kappa^2 S'$) of the *gem*-dithiolato ligands results in the formation of two four-membered metallocycles. In addition, the presence of a single bridgehead carbon atom between both sulfur atoms produces a compact and rigid M₂S₂ core with a small angle between the coordination planes of the rhodium centers and short metal-metal distances.

The bis-thiolato complexes can exist as different conformers (syn-exo, syn-endo, and anti) that arise from the spatial arrangement of the bridging thiolato ligands with a $1:2\kappa^2 S$ coordination mode.⁵² However, a large number of diolefin complexes $[M(\mu-SR)(cod)]_2$ (M = Rh, Ir) display a syn-endo conformation in the solid state and have a dynamic behavior by inversion of the nonplanar $Rh_2S_2\ ring.^{21a,41b,53,54}$ In contrast, the dinuclear complexes $[Rh_2(\mu-S_2CR_2)(cod)_2]$ are static as should be expected by the structure of the bridging ligand that blocks the ring-flipping process. On the other hand, dithiolato ligands afforded di- or tetranuclear complexes depending on the structure of the dithiolato ligand. For example, the nuclearity of the rhodium complexes $[Rh_2(\mu-S(CH_2)_nS)(cod)_2]_x$ is influenced by the number of methylenic units between the two sulfur atoms. Thus, dinuclear complexes (x = 1) were prepared for n = 2, 3,but a tetranuclear complex was obtained in the case of 1,4butanedithiolato ligand (n = 4).^{23d} More sophisticated ligands, for example, 1,1'-binaphthalene-2,2'-dithiolato, gave a mixture of both di- and tetranuclear complexes.^{24e} Interestingly, the tetranuclear complexes were easily converted to dinuclear complexes by carbonylation at atmospheric pressure, that is $[Rh_2(\mu-S(CH_2)_4S)(CO)_4]$, indicating the strong influence of the auxiliary ligands on the nuclearity.23d Similarly, dinuclear complexes [Rh(µ-SR)₂(CO)₂]₂ and [Rh₂(µ- S_2CR_2 (CO)₄ were obtained along the carbonyl series with bis-thiolato and gem-dithiolato ligands, respectively.

The replacement reactions by monodentate P-donor ligands on the tetracarbonyl complexes usually stop at the disubstituted complexes. The solid-state structure of the complexes $[Rh(\mu-SR)(CO)(PR'_3)]_2$ show a cis arrangement of the bulky phosphine or phosphite ligands and an anti conformation of the bridging thiolato ligands with the endo substituent at the side of the molecule with the small carbonyls ligands.^{21a,48–51,55} However, in some complexes a dynamic equilibrium between the cis and trans isomers has been observed in solution.^{41b,43b,50} In contrast, the complexes $[Rh_2(\mu-S_2CR_2)(CO)_2(PR_3)_2]$ that do not have any conformational restriction associated to the bridging ligand exist predominantly as the trans isomer. In the dithiolato series, the carbonyl replacement reactions in some dinuclear tetracarbonyl complexes encompass a change of nuclearity and the formation of tetranuclear complexes is quite common, depending on the structure of the dithiolato ligand.^{23c,24d,e} Nevertheless, some dinuclear complexes exist exclusively as the trans isomer,^{23d,56} although cis/trans mixtures have been observed in other cases.^{23c,57} Another striking difference between the three types of complexes concerns the formation of pentacoordinated species in the substitution process that evolve to the disubstituted complex by decarbonylation. Thus, the species [Rh(μ -SR)(CO)₂(PR₃)]₂ have been characterized in the bis-thiolate chemistry^{43b} but have not been observed in the *gem*-dithiolato complexes. Interestingly, related species have detected in dithiolate chemistry under CO pressure.⁵⁸

An additional remarkable difference between bis-thiolate and *gem*-dithiolate chemistry concerns the pattern of substitution by monodentate phosphite ligands. As described above, it is possible to replace the four carbonyl ligands in complex [Rh₂(μ -S₂CBn₂)(CO)₄] (7) as was evidenced by the characterization of the complex [Rh₂(μ -S₂CBn₂){P(OMe)₃}₄]. However, the disubstituted complex [Rh(μ -S'Bu)-(CO){P(OMe)₃}₂ does not react further with P(OMe)₃ despite that compound [Rh(μ -S'Bu){P(OMe)₃}₂]₂ can be prepared from [Rh(μ -Cl){P(OMe)₃}₂]₂ and LiS'Bu.⁵⁹

As far as the carbonyl replacement by diphosphine ligands is concerned, it is important to notice that a large number of bis-thiolato complexes $[Rh_2(\mu-SR)_2(CO)_2(\mu-diphos)]$ containing bridging diphosphine ligands have been characterized as the cis—anti isomer.^{46,60} In the same way, replacement reactions on the carbonyl *gem*-dithiolato complexes gives the expected cis isomer as has been exemplified by the synthesis of $[Rh_2(\mu-S_2CBn_2)(CO)_2(\mu-dppb)]$ (16). However, in the dithiolato series the dinuclear framework is not generally preserved as was evidenced by the formation of ion-pair compounds $[Rh(dppp)_2][Rh(dithiolato)(CO)_2]$ as the result of the sequestering of a rhodium atom by dppp.^{23b}

Concluding Remarks. We have described the high yield synthesis of dinuclear $[M_2(\mu-S_2CR_2)(cod)_2]$ (M = Rh, Ir) and $[Rh_2(\mu-S_2CR_2)(CO)_2(PR_3)_2]$ complexes by double deprotonation of diverse *gem*-dithiol compounds using mono- or dinuclear complexes containing basic ligands. The diolefin complexes are precursors for the carbonyl complexes $[Rh_2(\mu-S_2CR_2)(CO)_4]$, which undergo stereoselective replacement reactions by P-donor ligands to give the trans disubstituted complexes with monodentate ligands and the cis complexes with bridging diphosphines. The synthetic versatility of this kind of complexes, the singular structural features of the

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compact Rh₂S₂ core enforced by the *gem*-dithiolato bridging ligand, and the catalytic activity for the hydroformylation of alkenes open the possibility of study of the influence of the *gem*-dithiolato ligand on the catalytic activity. Further studies concerning the reactivity and catalytic activity of these complexes are currently in progress.

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Supporting Information Available: X-ray crystallographic file in CIF format for the structure determination of **1**, **2**, **6**, and **16**. This material is available free of charge via the Internet at http://pubs.acs.org.

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