

Preliminary communication

LIGAND SCAVENGING AND CATALYTIC UTILIZATION OF THE PHOTOTRANSIENT $\text{ReH}_5(\text{PMe}_2\text{Ph})_2$

MARK A. GREEN, JOHN C. HUFFMAN, KENNETH G. CAULTON,
Department of Chemistry and Molecular Structure Center, Indiana University,
 Bloomington, Indiana 47405 (U.S.A.)*

WITOLD K. RYBAK and JÓZEF J. ZIÓŁKÓWSKI
Institute of Chemistry, University of Wrocław, Wrocław (Poland)
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Summary

Photogenerated ReH_5P_2 ($\text{P} = \text{P}(\text{CH}_3)_2\text{C}_6\text{H}_5$) effects hydrogen exchange selectively at the β -carbons of naphthalene. Under one atm H_2 , hex-1-ene is hydrogenated catalytically. Cyclopentene, however, forms the complex $\text{ReH}_3\text{P}_3(\eta^2\text{-C}_5\text{H}_8)$, whose crystal structure shows hydride coordinated *cis* to the olefin. Photolysis of this complex results in disproportionation of the carbocycle to yield cyclopentane, $(\eta^5\text{-C}_5\text{H}_5)\text{ReH}_2\text{P}_2$ and $(\eta^5\text{-C}_5\text{H}_5)\text{ReH}_4\text{P}$. In the presence of *t*-butylethylene as a hydrogen acceptor, ReH_5P_2 is transformed into a species which dehydrogenates cyclopentane to $(\eta^5\text{-C}_5\text{H}_5)\text{ReH}_2\text{P}_2$. Reaction of ReH_5P_2 and *t*-butylethylene in benzene solvent under N_2 yield the arene complex $(\text{C}_6\text{H}_6)\text{ReP}_2(\text{CH}_2\text{CH}_2\text{CMe}_3)$ and *fac*- $\text{Re}(\text{PMe}_2\text{C}_6\text{H}_5)_3(\text{PMe}_2\text{C}_6\text{H}_4)\text{N}_2$, whose crystal structure is reported.

Following our observation that 366 nm irradiation of ReH_5P_3 ($\text{P} = \text{PMe}_2\text{Ph}$) results in phosphine dissociation [1], we report here a broader study of the reactivity of photogenerated ReH_5P_2 . Photogeneration (25°C) of ReD_5P_2 in a C_6D_6 solution in the presence of naphthalene leads to deuteration of the naphthalene. Under these circumstances, the C_6D_6 serves as a reservoir of deuterium and extensive deuteration of the naphthalene occurs in 3 h (550 W Hanovia lamp). Analysis of deuterium incorporation into recovered (sublimed) naphthalene by a combination of ^1H and ^2D NMR shows a substantial** prefer-

* Contribution No. 3773.

** A sample enriched to 46% deuterium overall is 74% enriched at the 2,3,6,7-positions and 19% enriched at the 1,4,5,8-positions. The mass spectrum of this sample shows it to be ~12% d_0 (totally unenriched).

ence for H/D exchange at the four equivalent β -carbons of naphthalene.

Photolysis of $\text{ReH}_5(\text{PMe}_2\text{Ph})_3$ and 1-hexene (mole ratio 1/8) in cyclohexane- d_{12} or hexafluorobenzene under 1 atm of H_2 results in complete hydrogenation of the olefin in 3 h (25°C). The reaction is catalytic in rhenium, but requires continuous photolysis: shuttering the light after an initial irradiation period of 10 min gives no significant additional hydrogenation over a dark period of 12 h.

The situation differs with internal olefins. Irradiation of ReH_5P_3 and cyclopentene (mole ratio 1/10) for 1 h in C_6H_6 results in nearly quantitative (by NMR) conversion to a stable hydridoolefin complex, $\text{ReH}_3(\text{PMe}_2\text{Ph})_3(\text{cyclopentene})$ (I)*. No olefin hydrogenation is observed. The X-ray structure** of I has been determined using crystals grown from hexane (Fig. 1). Although the

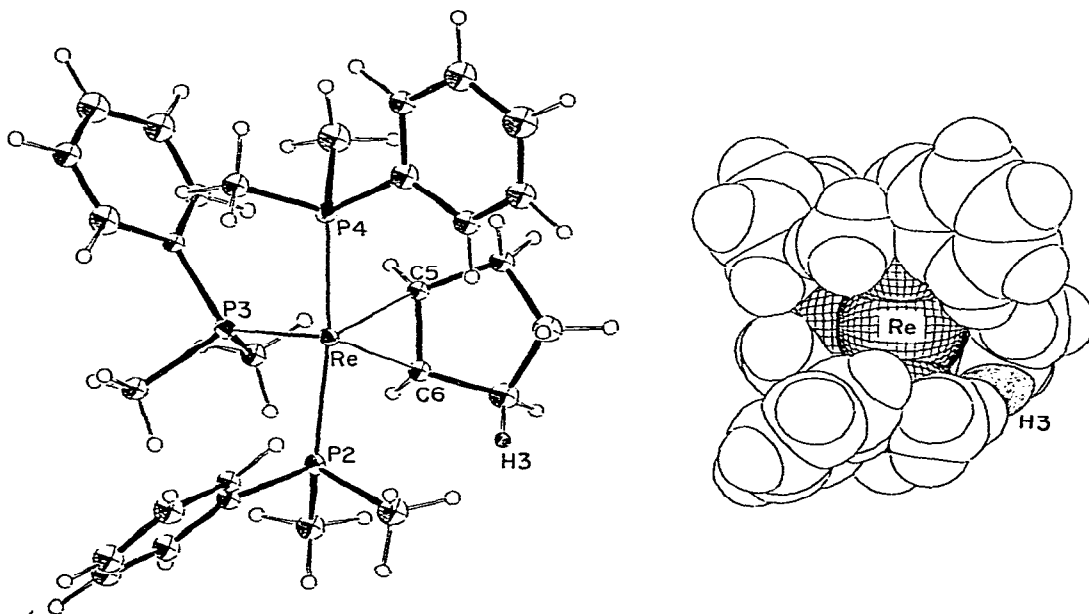


Fig. 1. Conventional ORTEP and space-filling drawings of $\text{ReH}_3(\text{PMe}_2\text{Ph})_3(\eta^2\text{-C}_5\text{H}_8)$. In this view, the edge of the equatorial plane of a proposed pentagonal bipyramid is horizontal, and the three hydrogens bound to rhenium are proposed to project out towards the reader. These hydrogens reside in the crevice shown in the space-filling view; other views of the molecule show no channel through which the metal is visible. One of the allylic hydrogens (H(3)) of the cyclopentene ligand is densely hatched.

Selected bond lengths: $\text{Re}-\text{P}(2) = 2.355(3)$ Å, $\text{Re}-\text{P}(4) = 2.374(3)$ Å, $\text{Re}-\text{P}(3) = 2.423(4)$ Å, $\text{Re}-\text{C}(5) = 2.275(14)$ Å, $\text{Re}-\text{C}(6) = 2.267(13)$ Å, $\text{C}(5)-\text{C}(6) = 1.423(18)$ Å.

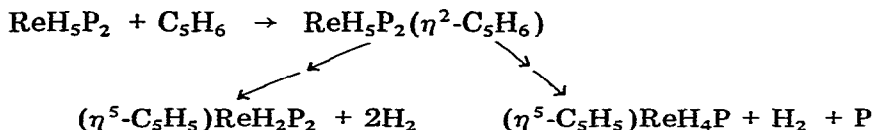
*NMR data for $\text{ReH}_3(\text{PMe}_2\text{Ph})_3(\text{C}_5\text{H}_8)$ in toluene- d_8 : ^1H NMR at 220 MHz and 50°C, δ (ppm) -6.90(3), quartet, J 17 Hz, $\text{Re}-\text{H}$; 1.45(18), doublet, J 5 Hz, $\text{P}-\text{CH}_3$; 1.28(1), 2.04(3), 2.39(1), 2.68(2), all multiplets, C_5H_8 ; 7.59(6), broad singlet, *ortho*- C_6H_5 . The *meta* and *para* protons of the phosphine overlap the protic impurity of the solvent. $^{31}\text{P}\{^1\text{H}\}$ NMR at 40.5 MHz and -70°C (chemical shifts referenced to external H_3PO_4 with downfield shifts reported as positive values) δ -14.5 ppm, doublet, J 10.3 Hz; δ -35.6 ppm, triplet, J 10.3 Hz.

**Crystallographic data (-162°C): a 16.334(5) Å, b 8.378(2) Å, c 10.739(2) Å, β 105.90(1)°, $Z = 2$ in polar space group $P2_1$; $R(F) = 0.033$, $R_w(F) = 0.033$ for 1832 observed ($F_o > 3\sigma(F_o)$) and absorption-corrected reflections using anisotropic temperature factors for Re and three phosphorus atoms; all hydrogen atoms except those bound to Re were refined with fixed β 's. A complete tabulation of crystallographic parameters is contained in Molecular Structure Center Report No. 81020, available from the Chemistry Library, Indiana University.

molecule is chiral and has no crystallographically-imposed symmetry, the coordination sphere has an approximate mirror plane of symmetry through Re, P(3), and bisecting the C=C bond of the cyclopentene. This is consistent with the AB₂ pattern seen in the ³¹P{¹H} NMR spectrum at -70°C. Space-filling models and nonbonded contact calculations show that the (crystallographically undetected) metal-bound hydrogens must lie almost in the plane of Re, P(3), and the olefin midpoint.

Photolysis of ReH₃P₃(C₅H₈) in C₆H₆ results in both hydrogenation and dehydrogenation of the C₅ ring*. (η⁵-C₅H₅)ReH₂P₂ and (η⁵-C₅H₅)ReH₄P (~1/1) are detected in the ¹H NMR spectrum along with cyclopentane. The identity of these cyclopentadienyl hydride complexes has been established by ¹H NMR chemical shifts and integrations and by the multiplet patterns observed in the ¹H and selectively decoupled ³¹P NMR spectra**.

CpReH₄(PMe₂Ph) can be isolated in 20% yield after photolysis of ReH₅P₃ and cyclopentadiene in hexane (1 h at 20°C), with CpReH₂P₂ also being formed as a minor product. This chemistry bears a striking similarity to that recently reported by Baudry and Ephritikhine [3], who found that refluxing ReH₇(PPh₃)₂ in THF in the presence of cyclopentadiene produces exclusively CpReH₂(PPh₃)₂. Although those authors postulate that ReH₅(PPh₃)₂ is an intermediate in their thermal reactions, the product distribution of η⁵-C₅H₅ complexes clearly differs from that obtained with our photogenerated ReH₅P₂***. Since we have shown that CpReH₄(PMe₂Ph) fails to react with 3.5 equivalents of PMe₂Ph in 10 h at 70°C, we are observing a kinetically-controlled product distribution. We conclude that ReH₅P₂, either thermally generated from ReH₇P₂ or photogenerated from ReH₅P₃, reacts with the diene C₅H₆ in a step-wise manner, whose later steps exhibit considerable thermal control:



Since Crabtree [4] and later Ephritikhine et al. [5] have shown that *t*-butylethylene can markedly influence hydrogen transfer reactivity, we explored the effect of this olefin on our photosystem. The most dramatic influence we have encountered occurs in cyclopentane. Irradiation of ReH₅P₃ in pure cyclopentane yields Re₂H₆P₅, the product of scavenging of ReH₅P₂ by ReH₅P_n in an otherwise unreactive environment [1]. If the same experiment is repeated

*Pt/Re catalysts are widely employed in hydrocarbon reforming (disproportionation) [2].

**NMR data for CpReH₄PMe₂Ph in C₆D₆: ¹H NMR at 60 MHz and 35°C, δ (ppm) -8.4 [4], doublet, *J* 20 Hz, Re-H; 1.72(6), doublet, *J* 10 Hz, P-CH₃; 4.24(5), broad singlet, C₅H₅; 6.9-7.6, multiplet, C₆H₅. The ³¹P{¹H} spectrum is a singlet at δ -17.5 ppm which becomes a quintet upon selective decoupling of the downfield protons.

NMR data for CpReH₂(PMe₂Ph)₂ in C₆D₆: ¹H NMR at 60 MHz and 35°C, δ (ppm) -11.5(2), triplet, *J* 42.5 Hz, Re-H; 1.72(12), doublet, *J* 10 Hz, P-CH₃; 4.39(5), singlet, C₅H₅; 7.1-7.7, multiplet, C₆H₅. The ³¹P{¹H} spectrum is a singlet at δ -17.5 ppm which becomes a triplet upon selective decoupling of the downfield protons.

***We have established that the identity of the phosphine does not alter this conclusion. A hexane solution of ReH₇(PMe₂Ph)₂ and cyclopentadiene heated to 70°C for twenty minutes produces only CpReH₂(PMe₂Ph)₂. Also, photolysis of ReH₂(PPh₃)₃ and cyclopentadiene in THF cleanly produces CpReH₄(PPh₃)₂; no CpReH₂(PPh₃)₂ is detected.

with 8 volume percent *t*-butylethylene added to the cyclopentane, (η^5 -C₅H₅)-ReH₂P₂ is produced (unoptimized yield: 5%). No other cyclopentadiene or hydride-containing products and no rhenium metal are detected in the resulting yellow-brown homogeneous solution. We conclude that ReH₅P₂ is incapable of reacting with cyclopentane, but that *t*-butylethylene dehydrogenates this phototransient to a species sufficiently devoid of ligands and sufficiently electron rich to react with the C—H bonds of cyclopentane.

We have exploited this "hydride-stripping" capacity of *t*-butylethylene to produce arene-rhenium complexes, something which our earlier work showed was not possible directly from ReH₅P₂. Irradiation of a benzene solution of ReH₅P₃ and *t*-butylethylene (mole ratio 1/16) in a stoppered NMR tube under 1 atm N₂ results in hydrogenation of the olefin and a mixture of rhenium products. The arene complex we sought is indeed produced; (η^6 -C₆H₆)ReP₂-(CH₂CH₂CMe₃) has appropriate ¹H NMR resonances*, as does the analogous π -toluene complex from toluene solvent. However, the identity of the other product was not established until the X-ray structure** (Fig. 2) was complete. The product is *fac*-Re(PMe₂Ph)₃(PMe₂C₆H₄)N₂***, resulting from the complete loss of Re—H ligands and the loss of one aryl *ortho* hydrogen upon *ortho*-

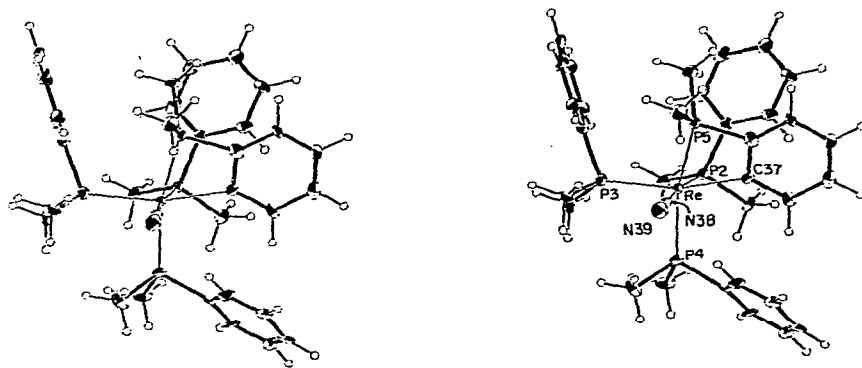


Fig. 2. ORTEP drawing of *fac*-ReN₂(PMe₂Ph)₃(PMe₂C₆H₄).

Selected structural data: Re—P(2) = 2.395(3) Å, Re—P(3) = 2.362(3) Å, Re—P(4) = 2.357(3) Å, Re—P(5) = 2.419(3) Å, Re—C(37) = 2.203(12) Å, Re—N(38) = 1.960(10) Å, N(38)—N(39) = 1.117(13) Å. Angle Re—N(38)—N(39) = 178.2(10)°, angle P(5)—Re—C(37) = 65.6(3)°.

metallation. This is a rare instance [6] where an arylphosphine with a cone angle as small as dimethylphenylphosphine has undergone *ortho*-metallation and contrasts with our observation that ReD₅P₂ undergoes no H/D exchange at the *ortho*-positions. This emphasizes the high reactivity of intermediates produced using *t*-butylethylene. Most remarkable is the fact that this reaction

*¹H NMR at 220 MHz and 16°C: δ (ppm) 4.07(6), singlet, C₆H₆; 1.18(9), singlet, C(CH₃)₃.

**Crystallographic data (–160°C): a 19.494(7) Å, b 10.403(2) Å, c 18.772(7) Å, β 120.64(2)°, Z = 4 in space group P2₁/a; $R(F)$ = 0.061, $R_w(F)$ = 0.054 for 4236 observed ($F_o > 2.33\sigma(F_o)$) reflections using anisotropic temperature factors for all nonhydrogen atoms. Placing hydrogen atoms in calculated positions reduced the residuals to 0.056 and 0.047, respectively. A complete tabulation of crystallographic parameters appears in Molecular Structure Center Report No. 81026, available from the Chemistry Library, Indiana University.

***³¹P {¹H} NMR at 40.5 MHz (referenced as above): δ –88.6 ppm, doublet (J 160 Hz) of pseudo-triplets (J 20 Hz), P(5); δ –25.7 ppm, complex multiplet, P(2) (or P(3)) and $\frac{1}{2}$ P(4); δ –30.5 ppm, complex multiplet, P(3) (for P(2)) and $\frac{1}{2}$ P(4). ν (NN) 2000 cm^{–1} (Nujol).

system (overall phosphine/Re ratio = 3/1) becomes so deficient of ligands that dinitrogen is scavenged to play this role. Our results also demonstrate that dinitrogen ($\sim 5 \times 10^{-3} M$) [7] is a superior ligand to the abundantly available *t*-butylethylene ($7 \times 10^{-1} M$).

It is clear that ligand photodissociation from $\text{ReH}_5(\text{PMe}_2\text{Ph})_3$ provides access to a transient of high and diverse reactivity, and one which readily shuttles between many oxidation states.

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