up the unit cell contents. These two hexamers are not identical but nearly so.

A less cluttered view of the hexagonal prism formed by the lithium atoms and the oxygen atoms is given in Figure 2. The six atoms forming either hexagonal face deviate by less than 0.1 A from the best plane through the face.

Since all six enolate anions (i.e., C₆H₁₁OLi units) in the crystallographic asymmetric unit of the pinacolone enolate are similar, a labeled plot of only one of these is given in Figure 3. Selected bond distances and angles for the specific enolate depicted are given with standard deviations of the least significant digits in parentheses. One salient feature is the variation in bond angles for the three C(2)-O(1)-Li bonds. Particularly outstanding are the rather large C(2)-O(1)-Li(1) bond angle (140.0 (4)°) and the small C(2)-O(1)-Li(2) bond angle (88.0 (9)°).

The average bond lengths of the six crystallographically independent enols corresponding to the bonds represented by C-(1)—C(2) and C(2)—O(1) in Figure 3 are 1.33 and 1.34 Å, respectively. These values match the values of the four C=C (1.34) Å) and the four C-O (1.35 Å) bonds in the symmetrical THF-solvated pinacolone enolate.3a However, a significant difference between the hexameric pinacolone enolate and the tetrameric enolate exits. This difference is exemplfied by comparing average distances corresponding to C(1)-Li(2) and C(1)-Li(3a) in Figure 3. In the hexamer, the average C(1)-Li(2) distance is only 2.53 Å and the average C(1)-Li(3a) distance is 3.36 Å. For comparison, in the tetramer the average analogous distances are 3.25 and 3.28 Å, respectively. The terminal methylene carbon is $\simeq 0.7$ Å closer to one specific lithium atom in the hexameric aggregate. Hence, for the hexameric pinacolone enolate each π bond is paired with a unique lithium atom by virtue of a relatively short distance; whereas in the THF-solvated tetramer, the C-(1)-C(2) σ bond closely bisects a Li-O-Li bond angle and C(1) is more symmetrically disposed with respect to the two staggered lithium atoms. 10

Calculation of the structure of the "gas-phase" lithium enolate of acetaldehyde at the restricted Hartree-Fock level resulted in three different structures of almost identical minimum energy.¹¹ A fragment of Figure 3 represented by Li(2)-O(1)-C(2)-C(1)provides experimental realization of one of the calculated enolate geometries with a lithium atom roughly equidistant from both C(1) and C(2). The O-Li bonds are, however, ≈ 0.23 Å longer in the crystal than in the calculated structure.

The ramifications of this crystal structure for the mechanism of enolate reactions in solution remain to be determined. Evidence for the correlation of X-ray crystal structures of lithiated species with solution species exits. 12 Further investigation to establish this correlation for the hexameric pinacolone enolate as well as crystal structure analyses of additional alkali metal ketone enolates are under way.

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Registry No. tert-Butyl methyl ketone (lithium enolate), 70367-67-8.

Supplementary Material Available: Full crystallographic details including unit cell parameters, atomic coordinates, thermal parameters, bond lengths, bond angles, and structure factors (8 pages). Ordering information is given on any current masthead page.

Synthesis, Conformation, and Reactivity of the Rhenium Phosphide Complex $(\eta^5 - C_5H_5)Re(NO)(PPh_3)(PPh_2)$. The "Gauche Effect" in Transition-Metal Chemistry

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Factors that influence the conformations of metal-bound ligands are fundamentally important and can provide practical insight into reactivity, especially in metal-mediated asymmetric synthesis. Since Corey and Bailar applied concepts from organic conformational analysis to the structures of metal chelate rings over 25 years ago, many steric and electronic contributions to ligand conformations have been identified.² We now wish to draw a new, stereoelectronic analogy from organic and main-group element chemistry to transition-metal chemistry—namely, that the "lone pairs" of ligating atoms should prefer to be orthogonal to the metal fragment HOMO.3 In organic and main-group inorganic compounds, this is commonly called the "gauche effect" 4 and has a profound influence on physical^{4,5} and chemical⁶ properties.

We first sought evidence for a "gauche effect" in metal phosphide complexes (L_nMPR₂) containing pyramidal phosphorus and hence an unambiguously defined "lone pair".7 We noted that in Paine's crystal structure of (η⁵-C₅Me₅)Fe(CO)₂(PN(CH₃)-CH₂CH₂NCH₃)^{7b} the phosphorus ligand did adopt a conformation in which the lone pair was orthogonal to the iron HOMO. However, the three HOMOs of the $(\eta^5-C_5H_5)$ Fe(CO)₂+ fragment are close in energy. 2b,8 Hence, we synthesized rhenium phosphide complexes $(\eta^5-C_5H_5)Re(NO)(PPh_3)(PR_2)$, where the metal fragment HOMO (d orbital shown in 1) would be energetically

⁽¹⁰⁾ A similarity in both the hexameric and tetrameric enolates is the rather large bond angle of C(2)-O-Li for the one lithium that is approximately anti to C(1) and syn to C(3). Perhaps this is a consequence of the steric bulk of the tert-butyl group

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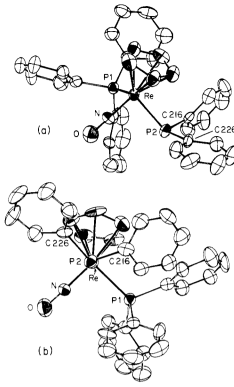


Figure 1. Two views of the molecular structure of $(\eta^5 - C_5 H_5) Re(NO)$ -(PPh₃)(PPh₂) (4); (b) approximates a Newman projection down the Ph₂P-Re bond. Important bond lengths (Å): Re-P(1), 2.358 (3); Re-P(2), 2.461 (3); Re-N, 1.738 (10); P(2)-C(216), 1.813 (14); P(2)-C-(226), 1.811 (15); N-O, 1.208 (15). Important bond angles (deg): P(1)-Re-P(2), 92.5 (1); P(1)-Re-N, 91.5 (4); P(2)-Re-N, 92.5 (4); Re-P(2)-C(216), 111.7 (5); Re-P(2)-C(226), 107.5 (4); C(226)-P-(2)-C(216), 103.8 (6); Re-N-O, 177.9 (10).

well separated from other occupied orbitals.9

Reaction of tosylate complex $(\eta^5 - C_5H_5)Re(NO)(PPh_3)(OTs)$ (2, OTs = OSO₂-p-C₆H₄CH₃)¹⁰ with PPh₂H (CH₂Cl₂, 25 °C, 2 days) gave $[(\eta^5-C_5H_5)Re(NO)(PPh_3)(PPh_2H)]^{+-}OTs$ (3, 94%, eq i).11 Reaction of 3 with t-BuO- K+ yielded the desired

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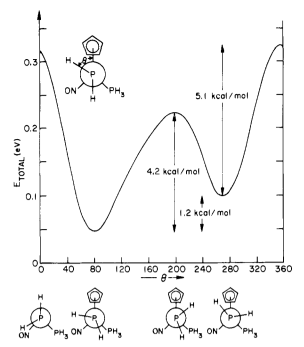


Figure 2. E_{total} for $(\eta^5 - C_5H_5)\text{Re}(\text{NO})(\text{PH}_3)(\text{PH}_2)$ as the PH₂ ligand is rotated, calculated (≥1 data point/10°) by the extended Hückel method with weighted H_{ij} formula.

phosphide complex $(\eta^5 - C_5 H_5) \text{Re}(NO) (PPh_3) (PPh_2) (4, 97\%).^{11}$ The X-ray crystal structure of 4 (red crystals from benzene/ hexane) was determined as described in the supplementary material.

Two views of the molecular structure of 4, and important bond lengths and angles, are given in Figure 1. The Re-PPh2 bond (2.461 (3) Å) is significantly longer than the Re-PPh₃ bond (2.358 (3) Å). The PPh₂ phosphorus is distinctly pyramidal, but the bond angles (107.5°, 111.7°, 103.8°) are markedly greater than usually found in trivalent phosphorus compounds such as PMe₃ (99°) and PPh₃ (103°). This suggests greater than normal p character in the PPh₂ lone pair. The torsion angle between the PPh₂ lone pair¹³ and the rhenium fragment HOMO (taken as Re-PPh₃ bond) is 59.7°, as illustrated by projection b in Figure 1. Hence, the expected near orthogonality of orbitals is observed.

To support our hypothesis that a gauche effect contributes to the conformation of the PPh2 ligand in 4, extended Hückel MO calculations were conducted on the model compound (η^5 -C₅H₅)Re(NO)(PH₃)(PH₂). Key results are shown in Figure 2.14 In contrast to the ~3 kcal, degenerate 3-fold Re-CH₃ rotational barrier calculated for $(\eta^5-C_5H_5)Re(NO)(PH_3)(CH_3)$, 9a $(\eta^5-$ C₅H₅)Re(NO)(PH₃)(PH₂) exhibits a higher, nondegenerate 2-fold Re-PH₂ rotational barrier. Energy minima correspond to near-orthogonal orientations of the PH2 lone pair and 1. One staggered Re-PH₂ conformation ($\theta = 180^{\circ}$), which should be a steric local minimum, is in fact calculated to be almost a local maximum due to eclipsing of the PH2 lone pair and 1.

Finally, compounds that exhibit the gauche effect often show enhanced solution-phase nucleophilicity ("\alpha effect"). 6a-d Accordingly, 4 and weakly electrophilic CH₂Cl₂¹⁵ readily react (2 h, 25 °C) to give $[(\eta^5-C_5H_5)Re(NO)(PPh_3)(PPh_2CH_2C1)]^+$ Cl⁻

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(5, 79%, eq i).¹¹ Also, 4 is readily oxidized by static air (THF, 21 h) to $(\eta^5 - C_5 H_5) \text{Re}(\text{NO}) (\text{PPh}_3) (\text{PPh}_2 O) (6, 71\%).^{11}$

In view of the numerous common transition-metal ligands with lone pairs on the ligating atoms (OR, SR, SR2, NR2, etc.), we believe that the ideas set forth above will prove useful in interpreting a large body of structural and reactivity data. Our results also suggest several reasons for the ease of formation and stability of bridging phosphide 16 ligands and may bear on the extremely low phosphorus inversion barriers observed with 4 and related complexes.17

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Supplementary Material Available: Tables of analytical (3-6) and crystallographic (4) data (31 pages). Ordering information is given on any current masthead page.

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An Extremely Short Way to Prostaglandins¹

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Among various strategies for prostaglandin (PG) synthesis, the three-component coupling process¹ is one of the ideal approaches in view of the directness and synthetic flexibility. Obviously, the ultimate goal along this line is, as illustrated by eq 1 (M = metal,X = halogen), the single-pot construction of the whole frameworks

via organometallic-aided conjugate addition of the ω side-chain unit to 4-oxygenated 2-cyclopentenones followed by trapping of the regiochemically defined enolate species by organic halides having α side-chain structures. However, Syntex groups² among others, after pioneering, extensive study on this possibility, noted extreme difficulty in achieving the direct alkylation.³ Here we wish to announce the realization of this earnestly desired convergent synthesis. The success relies simply on the lithium (or copper) to tin transmetalation in the enolate stage, a technique elaborated earlier by Tardella (simple alkylation)⁴ and Itoh et

al. (vicinal carba-condensation).5

The requisite optically active cyclopentenone and ω side-chain blocks are now accessible in various ways. 1,6 An organocopper reagent was prepared under our standard conditions⁷ by mixing the vinyllithium derived from 2a^{6,8} in ether and a THF solution containing copper(I) iodide (1 equiv) and tributylphosphine (2.6 equiv). Sequential treatments of the enone 1 with this copper

 $SiR_3 = Si(CH_3)_2 - t - C_4H_9$

reagent (1:1 molar ratio, -78 °C, 1 h), hexamethylphosphoramide (11 equiv, -78 °C, 30 min), triphenyltin chloride (1 equiv, -78 °C, 10 min), and the allylic iodide 3¹⁰ (5 equiv, -30 to -20 °C, 17 h) afforded stereoselectively the PGE_2 derivative $\boldsymbol{5a}$ in 78%yield, $^{11-13}$ [α] 19 _D -49.9° (c 1.02, CH₃OH). No PGA derivatives were detected. Natural PGE₂ can be obtained from 5a by removal of the silyl protective group with HF-pyridine^{3b} followed by enzymatic ester hydrolysis.¹⁴ In a like manner, **5b** (a versatile precursor of D series of PGs), $[\alpha]^{16}_D$ -60.0° (c 1.02, CH₃OH), was prepared in 77% yield by the one-pot condensation of 1, 2b, and 3.13 Use of methyl 7-iodoheptanoate, a saturated alkylating agent (-20 °C, 16 h), gave the corresponding PGE₁ derivative in only 20% yield.13

Utilization of the propargylic iodide¹⁵ as the α side-chain unit allowed the synthesis of 6 in 82% yield, $[\alpha]^{17}$ _D -13.2° (c 0.59,

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