Transition-Metal-Substituted Diphosphenes. 29.1 Cycloaddition Reactions of the Diphosphenyl Complex (η⁵-C₅Me₅)(CO)₂Fe-P=P-Mes* (Mes* = 2.4.6-tBu₃C₆H₂) with Hexafluoroacetone. X-ray Structure Analyses of $(\eta^5-C_5Me_5)(CO)Fe-P(=PMes^*)OC(CF_3)_2C(O)$ and $(\eta^5-C_5Me_5)(CO)_2Fe-P-P(Mes^*)OC(CF_3)_2$

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Summary: The diphosphenyl complex $(\eta^5-C_5Me_5)$ - $(CO)_{2}Fe^{-P}-Mes^{*}$ $(Mes^{*} = 2,4,6-tBu_{3}C_{6}H_{2})$ undergoes a [3 + 2] dipolar cycloaddition with hexafluoroacetone to give the metalla heterocycle (η^5 -C₅Me₅)(CO)-Fe-P(=PMes*)OC(CF₃)₂C(O) with a remarkably short Fe-P bond (2.084 (4) Å) and an exocyclic P-P bond. When stored in solution at -40 °C, this complex partly rearranges to the metalated 1-oxa-2,3-diphosphetane $(\eta^5\text{-C}_5\text{Me}_5)(\text{CO})_2\text{Fe}$ —P—P(Mes*)OC(CF₃)₂. The molecular structures of both Isomers were elucidated by singlecrystal X-ray analyses.

Electron-releasing organometallic complex fragments as substituents markedly enhance the nucleophilicity of a diphosphene. In molecules such as $(\eta^5-C_5Me_5)(CO)(L)$ -M-P-P-R (M = Fe, L = CO, R = 2,4,6- $tBu_3C_6H_2$ -(Mes*), 2,4,6-(CF₃)₃C₆H₂, 2,6-(CF₃)₂C₆H₃, C(SiMe₃)₃; M =Ru, Os, L = CO, $R = Mes^*$; M = Mn, Re, L = NO, R =Mes*) the HOMO is mainly represented by the lone pair at the metalated phosphorus atom. This is documented by a number of reactions of (η⁵-C₅Me₅)(CO)₂Fe—P=P— Mes* (1) with organic and organometallic electrophiles. Compound 1 is converted into oxaphospholenes I by treatment with acrolein, methacrolein, and methyl vinyl ether.² Azodicarboxylates and azodiamides give rise to the formation of oxadiazaphospholenes II.3 whereas the reaction of 1 with 1,2,4-triazoline-3,5-diones furnished the first 1,2-diaza-3,4-diphosphetidines III.⁴ [2 + 2] cycloadditions to 1,2-diphosphetanes IV are observed with fumarodinitrile, maleodinitrile dimethyl fumarate, dimethyl maleate,⁵ and maleimides.⁶ In all these processes we assume that the ring formation is initiated by the attack of the lone pair of the metalated P atom at the LUMO of the electrophile.

The step from electron-poor alkenes and azo compounds to other electrophiles with heteroatomic double bonds is obvious. Here we report on the chemical behavior of 1 toward anhydrous hexafluoroacetone, which is known to undergo oxidative additions to trivalent phosphorus compounds. Low-coordinated phosphorus systems such as

 R^1 , R^2 = H, Me; R^3 = OEt, OtBu, OCH₂Ph, NC₅H₁₀; E = CN, CO₂Me Aryl = Ph, 4-EtOC₆H₄; [Fe] = $(\eta^5 - C_5Me_5)(CO)_2$ Fe

iminophosphanes are converted by hexafluoroacetone to λ^5 -oxaphosphiranes V (for R = Me₃Si)⁸ or to λ^3 -1-oxa-3aza-2-phosphetanes VI (for R = alkyl).9 A different mode of reaction, however, is encountered with 1 and the ketone.

 $R^1 = Me_3Si$ or tBu $R^2 = Me_3Si$

Condensing an excess of gaseous hexafluoroacetone into

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Figure 1. Molecular structure of 3.

Table I. Selected Bond Lengths (Å) and Angles (deg) for 3

Fe-P(1)	2.084 (4)	P(2)-C(16)	1.860 (13)
Fe-C(11)	1.795 (16)	O(1)-C(11)	1.128 (19)
Fe-C(13)	1.937 (14)	O(2)-C(12)	1.427 (15)
P(1)-P(2)	2.014 (5)	O(3)-C(13)	1.191 (16)
P(1)-O(2)	1.647 (9)	C(12)-C(13)	1.640 (19)
Fe-P(1)-P(2)	140.3 (2)	P(1)-O(2)-C(12)	115.6 (7)
Fe-P(1)-O(2)	109.5 (3)	O(2)-C(12)-C(13)	111.8 (10)
P(2)-P(1)-O(2)	109.9 (3)	Fe-C(13)-C(12)	116.3 (9)
P(1)-P(2)-C(16	99.0 (4)	P(1)-Fe-C(13)	86.4 (4)

a pentane solution of 1 at -196 °C and warming to 20 °C afforded the cycloadduct 3 as an orange crystalline solid. No other product could be observed in the reaction mixture by means of ³¹P NMR spectroscopy, and no intermediate was detected during the course of the reaction.

[Fe]
$$+ \frac{(cF_3)_2cO}{D}$$

1

 F_3c
 F_3c

The structure of 3 was assigned on the basis of spectral evidence¹⁰ and confirmed by the single-crystal X-ray diffraction study.¹¹ The ³¹P NMR spectrum exhibits two doublets at δ 483.49 and 180.68 with the large coupling

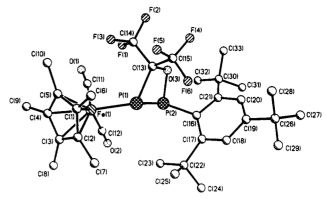


Figure 2. Molecular structure of 4.

Table II. Selected Bond Lengths (Å) and Angles (deg) for 4

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Fe-P(1)	2.327 (4)	P(1)-C(13)	1.927 (13)
Fe-C(11)	1.779 (15)	P(2)-O(3)	1.722 (8)
Fe-C(12)	1.744 (15)	O(3)-C(13)	1.409 (13)
P(1)-P(2)	2.249 (5)	O(1)-C(11)	1.139 (18)
P(2)-C(16)	1.865 (11)	O(2)-C(12)	1.144 (18)
Fe-P(1)-P(2)	110.7 (2)	O(3)-P(2)-C(1	6) 96.6 (5)
Fe-P(1)-C(13)	118.5 (4)	P(1)-Fe-C(11)	103.0 (4)
P(1)-P(2)-C(16) 105.9 (4)	P(1)-Fe-C(12)	92.0 (5)
P(1)-C(13)-O(3) 102.2 (8)	C(11)-Fe-C(12	90.4 (7)
P(2)-O(3)-C(13) 104.2 (7)		

constant $^1J_{\rm PP}$ = 633 Hz, suggesting the presence of a P—P double bond in 3. In the $^{19}{\rm F}$ NMR spectrum two quartets at δ -70.55 and -71.35 ($^4J_{\rm FF}$ = 7.3 Hz) agree with two magnetically nonequivalent CF₃ groups with no PF coupling and infer the absence of any direct PC(CF₃)₂ linkage. The appearance of two discrete singlets for the *o-tert*-butyl substituents of the Mes* ring in the $^1{\rm H}$ NMR spectrum at δ 1.65 and 1.68 is due to the chiral Fe center in 3. The IR spectrum (Nujol mull) displays only one intense band at ν = 1960 cm $^{-1}$ for the stretching mode of one terminal CO ligand. The acylic carbonyl group gives rise to a band at 1650 cm $^{-1}$ of medium intensity.

The most interesting feature of the molecular structure of 3 (Figure 1 and Table I) is the geometry of the nearly planar five-membered metalla heterocycle (mean deviation from the plane 0.024 Å). The Fe-P bond of 2.084 (4) Å is remarkably short and strongly suggests multiple-bond contributions. Similarly short Fe-P bond distances were reported for complexes 5 (2.117 (2) Å)¹² and 6 (2.112 (1) and 2.202 (1) Å).¹³ In 1 the Fe-P contact was determined

 $(Aryl = 2,6-tBu_2-4-MeC_6H_2)$

to be 2.260 (1) Å.¹⁴ The exocyclic P-P bond length (2.014 (5) Å) is comparable to the P-P bond in 1 (2.027 (3) Å). The bond length P(1)-O(2) (1.647 (9) Å) is shorter than the sum of the covalent single-bond radii for P (1.10 Å) and O (0.66 Å).¹⁵ The atoms Fe, P(1), P(2), and O(2) are

^{(10) 3: &}lt;sup>1</sup>H NMR (100 MHz, C_6D_6) δ 1.33 (s, p-tBu), 1.49 [s, $C_5(CH_3)_5$], 1.65 (s, o-tBu), 1.68 (s, o-tBu), 7.56 (m, m aryl H); ¹⁹F{¹H} NMR (84.2 MHz, n- C_5H_{12} , $CFCl_3$ standard) δ -70.55 (q, $^4J_{FF}$ = 7.3 Hz, CF_3), ^{-71.35} (q, $^4J_{FF}$ = 7.3 Hz, CF_3); ³¹P[¹H} NMR (40 MHz, n- C_5H_{12} , 85% H₃PO₄ standard) δ 483.49 (d, $^1J_{PP}$ = 633 Hz, Fe-P), 180.68 (d, $^1J_{PP}$ = 633 Hz, P-Mes*).

⁽¹¹⁾ Crystal data for complex 3: space group $P2_1/c$, a=21.809 (4) Å, b=10.066 (2) Å, c=16.250 (3) Å, $\beta=95.87$ (1)°, V=3560 (2) ų, Z=4, $\rho_{\rm catcd}=1.344$ g cm⁻³, Mo K α (graphite monochromator, $\lambda=0.710.73$ Å), ω -scan data collection at 183 K (3.0 $\leq 2\theta \leq 50.0^{\circ}$), 6228 unique reflections, 2383 unique observed reflections ($F>4.0\sigma(F)$), Siemens P2₁ four-circle diffractometer, structure solved by direct methods and refinement by full-matrix least squares, with use of the Siemens SHELTXL PLUS software on a Micro VAXII computer. All non-hydrogen atoms were refined anisotropically with 232 parameters (hydrogen atoms in calculated positions riding on the corresponding C atoms), U(H)=0.08 Ų, R=0.106, $R_w=0.072$, $w^{-1}=\sigma^2(F)$, and maximum rest electron density 0.88 e/A^3

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located in the same plane (mean deviation from plane 0.023

Crystals of 3 were grown from pentane solutions at -40 °C in several experiments. In one sample after 4 weeks of crystallization compound 3 decomposed to some extent. The ³¹P NMR spectrum displayed a doublet at δ 194.68 $(^{1}J_{PP} = 94 \text{ Hz})$ and a doublet of quartets at δ 136.00 $(^{1}J_{PP})$ = 94, $^{3}J_{PF}$ = 26 Hz) in addition to the resonances of 3. Both compounds were present in the ratio 3:4 = 3:1. The PP coupling constant is consistent with a PP single bond, whereas the size of the PF coupling suggests the presence of a PC(CF₃)₂ group. Both complexes could not be separated on a preparative scale.

A red crystal of 4 was picked out of the crop and submitted to an X-ray diffraction study (Figure 2 and Table The analysis shows the presence of an essentially planar 1-oxa-2,3-diphosphetane (mean deviation from plane 0.01 Å) which is linked to the $(\eta^5-C_5Me_5)(CO)_2Fe$ fragment via an Fe-P single bond (2.327 (4) Å). Compound 4 is obviously the result of a formal [2 + 2] cycloaddition between 1 and hexafluoroacetone. The P-P bond of 2.249 (5) A reveals a bond order of unity. In the fourmembered ring the oxygen atom of the ketone is added to the arylated P atom (P(2)-O(3) = 1.722 (8) Å). In keeping with this, the (CF₃)₂C fragment in 4 is connected

to the metalated phosphorus via a long P-C single bond (P(1)-C(13) = 1.927 (13) Å), which is quite common for diphosphetane derivatives.5,6

Due to the longer distance P(1)-P(2) the bond angles at the phosphorus atoms are markedly more acute (C- $(13)-P(1)-P(2) = 72.3 (4)^{\circ}, P(1)-P(2)-O(3) = 81.3 (3)^{\circ}$ as compared to the angles at carbon and oxygen (102.2 (9) and $104.2 (7)^{\circ}$, respectively). The E configuration of 1 has been maintained throughout the cycloaddition.

At room temperature solutions of 3 and 4 in benzene decompose within 1 week to give 1 and unidentified species. We suggest that the formation of 3 and 4 is initiated by a [2 + 1] cycloaddition to the transient adduct 2. Subsequent scission of the P-C linkage in 2 and attack of the carbanion at a positively polarized carbon atom of a terminal carbonyl ligand affords metallaheterocycle 3 (path a), whereas P-O bond rupture in 2 and attack of the oxygen at the arylated P atom yields 4. The dissociation of 3 into 1 and free $(CF_3)_2CO$ and a subsequent [2 + 2]cycloaddition to compound 4 cannot be excluded at the moment. The nature of the transformation of 3 into 4 is presently under investigation.

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Supplementary Material Available: Listings giving details of the crystal data, data collection, and structure solution and refinement and tables of atomic coordinates, bond lengths and angles, and thermal parameters for 3 and 4 (17 pages). Ordering information is given on any current masthead page.

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Platinum-Complex-Catalyzed 1,4-Disliplation of 1,3-Dienes Using Organodisilanes: Remarkable Effect of a Phenyl Functionality on a Silicon Atom

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Summary: Organodisilanes having a phenyl functionality on the silicon atom smoothly react with 1,3-dienes at 130 °C to afford the corresponding 1,4-disilylation adducts in high yields in the presence of platinum catalyst. However, other substituents such as Me, n-Bu, t-Bu, CH₂C₆H₅, CH₂CH=CH₂, and CH=CH₂ on the silicon atom are totally ineffective.

Activation of a silicon-silicon bond by a transition-metal catalyst¹ and subsequent insertions of 1,3-dienes,² acetycurrent interest. 1,4-Disilylation of 1,3-dienes is especially important, since it affords adducts⁵ having two versatile allylic silane moieties. Transition-metal-catalyzed 1,4-

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⁽¹⁶⁾ Crystal data for complex 4: space group C2/c, a=46.97 (3) Å, b=9.316 (5) Å, c=16.611 (5) Å, $\beta=101.52$ (3)°, V=7121 (6) ų, Z=8, $\rho_{\rm calcd}$ = 1.344 g cm⁻³, Mo K α (graphite monochromator, λ = 0.71073 Å), ω -scan data collection at 179 K (3.0 \leq 2 θ \leq 45.0°), 4661 unique reflections, 2143 unique observed reflections ($F > 4.0\sigma(F)$), Siemens P2₁ four-circle diffractometer, structure solved by direct methods and refinement by full-matrix least squares, with use of the Siemens SHELXTL PLUS software on a Micro VAXII computer. All non-hydrogen atoms were refined anisotropically with 254 parameters (hydrogen atoms in calculated positions, riding on the corresponding C atoms), $U(H) = 0.08 \text{ Å}^2$, R = 0.080, $R_w = 0.058$, $w^{-1} = \sigma^2(F)$, maximum rest electron density 0.50 e/Å^3 .

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