the **data** were **corrected** for Lorentz and polarization effects. The **structures** were solved by direct methods (MVLTAN)' or Patterson methods, and successive cycles of difference Fourier maps followed by least-squares refinement. Scattering factors were taken from ref **10.**

1: Monoclinic, space group $P2_1/c$ (No. 14), $Z = 4$; $a = 15.910$
(4), $b = 10.294$ (6), $c = 20.817$ (7) A; $\beta = 98.63$ (2)°; $V = 3371$ A³; $d(\text{calcd}) = 1.282 \text{ g cm}^{-3}$; $3^{\circ} < 2\theta < 55^{\circ}$ (Mo K_a, $\lambda = 0.71069 \text{ Å}$; $\mu = 25.2$ cm⁻¹). 2: Monoclinic, space group $P2_1/c$ (No. 14), Z = 4; a = 16.118 (7), b = 10.398 (2), c = 20.996 (4) Å; β = 98.87 (3)°; $V = 3477$ Å³; $d(\text{calcd}) = 1.495$ g cm⁻³; $3^{\circ} < 2\theta < 50^{\circ}$ (Mo K_a, λ) = **0.71069 A;** *p* = **51.4** cm-'). Totals of **5919** and **6108** unique

(10) *Intemtional Tables for X-ray Crystallography;* **Kynoch Press: Birmingham, UK, 1974; Vol. 4.**

reflections were collected at 24 °C for 1 and 3, respectively. Of these, 3035 and 3414 reflections with $I > 3.0\alpha(I)$ were used to solve (direct methods) and refine (full-matrix, least squares) the structures of 1 and 3, respectively. The final R and R_w values were **0.053** and **0.061** for **1** and **0.042** and **0.050** for 3. **C13** in the structure of **1** is disordered. There is a **5050** occupancy of pos- itions **C13A** and **C13B.**

Acknowledgment. We are grateful to the National Science Foundation, the Robert A. Welch Foundation and the ARO (Grant No. **DAAL** 03-89-K-0164) for generous financial support.

Supplementary Material Available: Tables of bond distances, bond angles, atomic coordinates, and thermal parameters for **1** and 3 **(13** pages); listings of structure factor amplitudes for **1** and 3 **(33** pages). Ordering information is given on any current masthead page.

Reactlvlty of Early-Transition-Metal Fulvene Complexes. Transformation of a 2,3,4,5-Tetramethylfulvene Ligand into a Bidentate Diaikoxide with Ti[(OCHPh)₂C₅Me₄(CH₂)]CI₂ **Four Asymmetric Carbon Atoms. Molecular Structure of**

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Summary: Cp*FvTiCl (1; Cp* = η^5 -C₅H₅, Fv = η^6 - $C_5Me_4CH_2$) reacts with benzaldehyde, yielding Cp^*T i- $[$ (OCHPh)₂C₅Me₄(CH₂)]CI (2) as the result of insertion in two Ti-C bonds at the **2-** and 4-positions of the fulvene ligand in 1. Complex 2 reacts with HCI to give Cp*TiCl₃ **(3)** and **the** corresponding dialcohol and with TCI, to give **3 and TI (OCHPh)₂C₅Me₄(CH₂) CI₂ (5). X-ray diffraction** analysis of 5 shows **It** to be a dlalkoxide-dichloride monomeric titanium complex with a tetrahedral arrangement of the ligands around the metal center. Crystal data **for** C,H&I,O2Ti: monoclinic, **~2,/n,** a = **10.602 (5) A,** *^b* $=$ **14.157 (4) Å,** $c = 15.274$ (5) Å, $\beta = 103.76$ (3)^o, *V* $= 2227 (1)$ \hat{A}^3 , $Z = 4$, $R = 0.037$. 3 and TI((OCHPh)₂C₅Me₄(CH₂)]Cl₂ (5). X-ray diffractionally sis of 5 shows it to be a dialkoxide-dichloride momenci thanium complex with a tetrahedral arrangement of the ligands around the metal center. Crystal d

The bonding of a tetramethylfulvene ligand to early transition metals has been described **as** intermediate be-

In complexes of the type Cp*FvM and Cp*FvMCl (M = Ti, **Zr)** the fulvene ligand reads with ketones or nitriles, through an adduct intermediate, to give bidentate alkoxide- or amide-functionalized tetramethylcyclopentadienyl groups.2 This reactivity can be explained in terms of the cyclopentadienyl alkyl structure A. Nevertheless, spectroscopic and structural data for these compounds point to a significant contribution of structure **B.3** A different bonding of the fulvene group is observed in late-transition-metal fulvene complexes, where in some cases the ligand appears to be bonded as a η^4 -diene⁴ or η^2 -olefin⁵ and extrusion of the fulvene ligand is observed.⁶

We wish to report here an unprecedented reactivity of this type of ligand that can be well understood in terms of another bonding form, C, a special type of 1,3-diene complex, in which the diene coordinates in a σ^2 , π mode as has been observed for a number of Zr and Hf complexes.⁷ of the full vente group is observed in
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on of the full vene ligand is observed.⁶
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Table I. Selected Bond Distances (A) end Angles (deg) for

| Bond Distances | | | |
|-----------------------|------------|----------------------|----------|
| $Ti-Cl(1)$ | 2.236(1) | $O(1) - C(1)$ | 1.427(4) |
| $Ti-Cl(2)$ | 2.240(1) | $O(2) - C(12)$ | 1.429(4) |
| $Ti-O(1)$ | 1.738(2) | $C(5)-C(6)$ | 1.333(4) |
| $Ti-O(2)$ | 1.739(2) | $C(3)-C(7)$ | 1.326(5) |
| Bond Angles | | | |
| $Cl(1) - Ti - Cl(2)$ | 110.34 (4) | $Ti-O(1)-C(1)$ | 154.6(2) |
| $Cl(1) - Ti - O(1)$ | 111.33 (8) | $Ti-O(2)-C(12)$ | 153.0(2) |
| $Cl(1) - Ti - O(2)$ | 110.71 (8) | $C(3)-C(4)-C(5)$ | 102.6(3) |
| $C(2) - C(3) - C(4)$ | 109.8(3) | $C(4) - C(5) - C(6)$ | 112.4(3) |

Results and Discussion

Cp*FvTiCl* **(1)** reacts with benzaldehyde in a 1:2 ratio,

giving the yellow complex 2 in 47% yield (eq 1).
\n
$$
Cp*FvTiCl + 2C(O)HPh \rightarrow 1
$$
\n
$$
Cp*[(OCHPh)2C5Me4(CH2)]TiCl (1)
$$

When the reaction is carried out in a 1:l ratio, half of the starting material remains unchanged, indicating that reaction with a second molecule of aldehyde is much faster than with the first.

Complex **2** was obtained **as** a yellow microcrystalline material. It is very air-sensitive and poorly soluble in toluene or pentane. It was characterized by IR and 'H NMR spectroscopy and elemental analysis and identified as the dialkoxide $\text{Cp*Ti}[(\text{OCHPh})_2\text{C}_5\text{Me}_4(\text{CH}_2)]$ Cl. The exact geometry and bonding conformation of the modified fulvene ligand was confirmed by an X-ray crystal structure determination of a derivative (vide infra).

The dialkoxide ligand can be removed from the metal by treatment with HCl, which generates Cp^*Ticl_3 and the free dialcohol **4** (eq 2).

$$
\text{Cp*[(OCHPh)2C5Me4(CH2)]TiCl + 2HCl \rightarrow P \atop 2
$$

\n
$$
\text{Cp*TiCl}_3 + \text{C}_5Me_4(CH_2)(CHPhOH)2 (2)
$$

Treatment of **2** with TiCl, leads to formation of $Cp*TiCl₃$ and the titanium dialkoxide-dichloride complex **5** (eq 3).

5 (eq 3).
\n2 + TiCl₄
$$
\rightarrow
$$
 Cp*TiCl₃ + TiCl₂[(OCHPh)₂C₅Me₄(CH₂)]
\n8 (3)

Complex **5** was isolated **as** white crystals. It is very poorly soluble in pentane but dissolves reasonably well in toluene. **A** singlet at 61.21 ppm in the 13C *NMR* **spectrum** of 5 indicates the presence of one or more sp³ carbon atoms
in the ring, while a triplet at low field [118.3 ppm $(^1J_{\text{C-H}})$ = 157.69 Hz)] unequivocally proves the existence of a $=CH₂$ group. All this indicates that the insertion has taken place somewhere in the ring instead of in the expected position, i.e., into the methylene-titanium bond.² Two singlets at 136.71 and 139.52 ppm are assigned to the remaining sp2 carbon atoms in the ring. The *NMR* spectra of **2,4,** and **5** are relatively simple, and the small number of absorptions observed indicate a high symmetry of the new ligand. **'H** NMR spectra of **all** three compounds **also**

Figure 1. Molecular structure of Ti[(OCHPh)₂C₆Me₄(CH₂)]Cl₂ **(5).**

show similar characteristics, **as** can be expected for compounds having the same ligand, and confirm the interpretation of the ¹³C NMR spectrum given above.

The molecular structure of **5** is shown in Figure 1. The complex is monomeric and shows a tetrahedral arrangement of the ligands around the titanium atom. Table I provides a selected list of pertinent bond distances and angles.

 $\text{Ti}-\text{O}(1)$ and $\text{Ti}-\text{O}(2)$ distances (1.738 (2) and 1.739 (2) Å) together with the very obtuse $Ti-O(1)-C(1)$ and $Ti-O (2)-C(12)$ angles $(154.6 (2)$ and $153.0 (2)$ °, respectively), indicate extensive π electron donation from oxygen to titanium.⁹ In the same way the Ti-Cl(1) and Ti-Cl(2) distances (2.236 (1) and 2.240 (1) **A,** respectively) **also** show some double-bond character, as expected for an electronically unsaturated complex.

The C_5 ring is practically planar (mean deviation from a plane is 0.008 **A)** and has normal C-C distances for four single bonds (average 1.526 (4) **A)** and one double bond (1.333 (4) **A).** The exo double bond is also normal (1.326 *(5)* A). Both features indicate that the five-membered ring has no interaction with the metal. The very long Ti-C distances $Ti-C(7) = 4.548$ (4) Å and $Ti-C(3) = 3.664$ (3) **A** underline this conclusion. The two phenyl groups are both oriented in the same direction.

It is easier to understand the reaction if the starting material, Cp*FVr'iCl, is formulated **as** a cis diene (structure C). Structure C can be seen **as** a close analogue of the well-known Cp*M(1,3-diene)X complexes reported before.' If structure C is a good description of the bonding in **1,** then we can see the process as an insertion in both Ti-C σ -bonds.¹⁰ Insertion after initial complexation of the aldehyde carbonyl oxygen atom to titanium will lead to attack on the fulvene ring plane from the metal side, thus explaining the observed stereoselectivity with respect to the configuration of the sp3 ring carbon atoms.

The stereoselectivity of the aldehyde carbon atoms can be understood on the basis of steric arguments. It is reasonable to expect that insertion will take place in a manner in which the phenyl groups will be pointing away from the Cp* ligand. Once the dialkoxide ligand is formed, rotation around the $C(C_5\text{-ring})-C(alkyl)$ bond should be strongly obstructed.

The regioselectivity of attack in the fulvene ligand is not yet understood. Some mechanistic studies in order to clarify this point are in progress.

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Conclusions

We have shown that it is possible to **transform** a fulvene ligand **into** a dialkoxide **with** a specific geometry. Transfer of a Cp* ligand and replacement by a chloride ligand is quite possible, and this opens an interesting perspective for further derivatization and synthesis of new classes of compounds with challenging chemistry.

Experimental Section

General Remarks. *All* reactions were performed under nitrogen with **we** of Schlenk, glovebox, and vacuum-line techniques. Solvents were distilled from Na/K alloy prior to use; $CHCl₃$ and C(O)HPh were dried over molecular sieves (4 Å) . IR spectra were recorded on a Pye-Unicam **SP3-300** spectrophotometer using Nujol mulls between KBr disks. *NMR* spectra were recorded on a Bruker WH-90 or a Varian **VXR-300** spectrometer. Chemical *shifts* **are** reported in **parta** per million and referenced to residual proton signals of the solvent (¹H: benzene- d_6 , δ 7.15; chloroform-d, δ 7.24) or the solvent itself (¹³C: benzene- d_6 , δ 128.00; chloroform, **6 70.00).** Elemental analyses were carried out at the Micro-Analytical Department of the University of Groningen. All data given are the average of two independent determinations.

Reaction of Cp*FvTiCl with C(O)HPh. To a solution of Cp*FvTiCl (0.4 g, 1.13 mmol) in 10 mL of toluene at 50 °C was added **0.23** mL **(2.25** mmol) of benzaldehyde. The mixture was stirred for **2** h, and the resultant yellow precipitate, **2,** was separated and washed first with toluene/pentane and then with pentane; yield 0.30 g $(0.53 \text{ mmol}, 47\%)$. **IR** (cm^{-1}) : 3075 (w), 3020 (w), **1635** (w), **1600** (w), **1485** (w), **1240** (w), **1230** (w), **1200** (m), **¹¹⁰³(w), 1089 (w), ¹⁰⁶⁵(e), ¹⁰⁵⁰(w), ¹⁰²⁰(81,940** (w), **908** (m), **880** (w), **795** (m), **768 (a), 738 (a), 728** (m), **700** (vs), **620** (w), **630** (w), **530** (w), **500** (w), **445** (m), **412 (s), 402** *(8).* Anal. Calcd for TiC102CMH11: C, **72.27;** H, **7.31;** Ti, **8.47;** C1, **6.27.** Found: C, **20** OC): **0.83 (e, 6** H, Me), **1.88 (a, 6** H, Me), **1.98 (s,15** H, Cp*), **4.70** *(8,* **2** H, =CHz), **5.47 (s, 2** H, H-C-O), **7.267.48** (m, **10** H, Ph). **72.22; H, 7.10;** Ti, **8.40; C1,6.26.** 'H *NMR* **(6,** benzenede,90 **MHz,**

Reaction of **2 with HCl.** To a suspension of **2 (0.1** g, **0.17** mmol) in 7 mL of CHCl₃ was added 0.1 mL of a solution of HCl (3.6 M) in Et₂O. The mixture turned orange immediately, and after the solvent was removed under vacuum, the product was analyzed by IR and ¹H and ¹³C NMR measurements, which showed that it was a 1:1 mixture of compounds 3 and 4. IR (cm⁻¹): **3225** (br), **1639** (w), **1600** (w), **1205** (w), **1114** (w), **1090** (w), **1050 (e), 1030 (e), 917** (m), **770** (m), **706 (w), 621** (m), **466** *(8).* 'H **NMFt** (δ , benzene- d_6 , 300 MHz): dialcohol $[C_5Me_4(CH_2)(CHPhOH)_2]$ 0.85 *(8,* **6** H, Me), **1.57 (e, 6** H, Me), **4.07 (e, 2** H, =CH2), **4.48 (s, 2** H, H-C-O), **5.75** (8, **2** H, O-H), **7.15-7.35** (m, **10** H, Ph); Cp*TiCls **1.89 (s,15** H, Cp*). **'BC NMR** (6, benzene-d,, **75.4** *MHz):* dialcohol $[C_5Me_4(CH_2)(\text{CHPhOH})_2]$ 10.06 $(q, {}^1J_{C-H} = 125.46 \text{ Hz},$ $C-Me$), 26.79 $(q, {}^{1}J_{C-H} = 127.76$ Hz, $C-Me$), 57.58 $(s, C-C)$, **77.71** (d, $^{1}J_{\text{C-H}} = 142.73$ Hz, H-C-O), 115.90 (t, $^{1}J_{\text{C-H}} = 156.54$ Hz, +HZ), **127.01** (d, 'JC-H = **158.84** Hz, Ph), **127.11** (d, *'Jc-H* = **161.14** Hz, Ph), **128.89** (d, *'JGH* = **151.85** Hz, Ph), **137.36** *(8,* C-Me), **142.14** (8, C-Me), **154.18 (e,** a-Ph); Cp*TiC13 **14.05** (q, *'JGH* = **128.91** Hz, Me), **135.85** *(8,* C-Me).

Reaction of **2 with TIC4.** To a suspension of **2 (0.29** g, **0.51** mmol) in 15 mL of CHCl₃ was added 54 μ L (0.49 mmol) of TiCl₄, yielding an orange solution. The solvent was pumped off, and after several recrystallizations from toluene/pentane **0.12** g **(0.25** mmol,50%) of **5** was obtained **as** white *Cryetals.* **IR** (cm-'1: *3090* (w), **3030** (w), **1495** (w), **1305** (w), **1240** (w), **1200** (w), **1110** (sh), **¹¹⁰⁰(e), 1090 (a), ¹⁰⁵⁰(w), ¹⁰³⁰(a), 916** (m), **870** (w), **795** (w), **778** (m), **765 (a), 740 (a), 730** (sh), **708** (vs), **638** (w), **620** (w), **540** (w), 468 (vs), 409 (s). Anal. Calcd for TiCl₂O₂C₂₄H₂₆: C, 61.95;

H, 5.63; Ti, **10.29;** C1, **15.23.** Found C, **61.59;** H, **5.66,** Ti, **10.30;** C1, **15.20.** 'H NMR **(6,** chloroform-d, **300** MHz): **1.05** *(8,* **6** H, C-Me), **1.81** *(8,* 6 H, C-Me), **3.97** *(8,* **2** H, H-C-O), **5.53 (e, ²**H, =CH2), **7.30** (m, **10** H, Ph). **'q** NMR (6, chloroform-d, **75.4 11. 10.36 (g, ¹J_{C-H} = 125.46 Hz, C-Me)**, **25.61 (g, ¹J_{C-H} = 127.76 Hz, C-Me)**, **61.21 (s, C-C-C-O)**, **101.28 (d, ¹J_{C-H} = 149.64** *HZ,* H-C-O), **118.23** (t, *'J~H* = **157.69,** +Ha, **127.40** (d, *'JGH* = **159.99** Hz, Ph), **127.46** (d, *'Jc-H* = **159.99** Hz, Ph), **128.12** (d, *'Jc-H* = **160.00** Hz, Ph), **136.71** *(8,* C-Me), **139.52** *(8,* C-Me), **151.44 (e,** a-Ph).

Crystal Structure Analysis. Crystal Data: C₂₄H₂₈Cl₂O₂Ti, $M_r = 465.3$, monoclinic, space group $P2_1/n$ $(P2_1/c$, nonstandard setting), $a = 10.602$ (5) Å, $b = 14.157$ (4) Å, $c = 15.274$ (5) Å, β = 103.76 (3)°, $V = 2227$ (1) Å³, $Z = 4$, $D_{\text{caled}} = 1.39$ g cm⁻³, μ (Mo $K\alpha$) = 6.38 cm⁻¹.

Suitable colorless, irregular-shaped prisms were obtained by slow diffusion of pentane into a saturated solution of **5** in toluene. A crystal with the approximate dimensions $0.22 \times 0.15 \times 0.10$ mm was mounted in epoxy resin on a glass fiber and transferred to a Rigaku AFC6R diffractometer. Diffracted intensities (+ $h, +k, \pm \tilde{l}$ were measured at -150 ± 1 °C, with use of graphitemonochromated Mo Ka radiation from an RU200 rotating anode operated at 9 kW (50 kV; 180 mA). The $\omega/2\theta$ scan mode was employed for $3.5 < 2\theta < 50$ ° with an ω scan rate of 8° min⁻¹ and a scan width of $(1.31 + 0.30 \tan \theta)$ ^o. Weak reflections $(F \leq 10\sigma(F))$ were rescanned up to three times and counts accumulated to improve counting statistics. Stationary background counts were recorded on *each* side of the reflection. The ratio of **peak** counting time vs background counting time was **2:l.** Of the **3914** independent reflections measured, excluding those systematically absent, 2686 had $I > 3\sigma(I)$ and were regarded as being observed. That the crystal was not subject to decay during measurement of intensities was checked by monitoring three reflections at regular intervals (after measurements of **150** reflections). Intensity data were corrected for Lorentz and polarization effects, but no correction was made for the effects of absorption. Unit-cell parameters were obtained from diffractometer setting angles for **25** reflections in the range **44.6'** < **28** < **48.7'.**

Structure Determination and Refinement. The structure was solved by direct methods (MITHRIL)¹¹ and subsequent electron density calculations. Full-matrix least-squares refinement of positional and anisotropic thermal parameters for the non-hydrogen atoms, with the hydrogen atoms **as** a fixed contribution in positions as located from a difference map, gave a final $R =$ 0.037 , $R_w = 0.042$, for 262 parameters and 2686 reflections. Atomic scattering factors and anomalous dispersion **corrections** were taken from ref 12, and the F_0 values were weighted according to $w =$ $[\sigma^2(F_{\alpha})]^{-1}$. A final difference map showed maximum and minimum residual densities of 0.47 and -0.29 e Å⁻³, respectively. All calculations were carried out with the TEXSAN program system.18

Acknowledgment. We thank the Ministerio Español de Educacion y Ciencia for a postdoctoral fellowship for **R.F.**

Supplementary Material Available: Tables of bond lengths and angles, positional and thermal parameters, and least-squares planes for **5 (12** pages); a listing of observed and calculated structure factors for **5 (27** pages). Ordering information is given on any current masthead page.

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