Tuning the Excited-State Properties of Luminescent Rhenium(V) Benzylidyne Complexes Containing Phosphorus and Nitrogen Donor Ligands

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Received December 2, 1997

Efficient methods are developed for the synthesis of rhenium(V)-substituted benzylidyne complexes with various auxiliary ligands to facilitate the tuning of their excited-state properties. The electronic structures and spectroscopic and photophysical properties of [Re- $(\equiv CAr')$ (pdpp)₂Cl]⁺ (Ar' = C₆H₂Me₃-2,4,6, pdpp = *o*-phenylenebis(diphenylphosphine), **2**), $[Re(\equiv CAr')L_2(CO)(H_2O)Cl]^+$ (L = PPh₃, **3**; P(C₆H₄OMe-*p*)₃, **4**; PPh₂Me, **5**), $[Re(\equiv CAr')(dppe)$ - $(CO)_2Cl$ ⁺ (dppe = 1,2-bis(diphenylphosphino)ethane, **6**), $[Re(\equiv CAr')(L-L)(CO)_2Cl]$ ⁺ (L-L)) 2,2′-bipyridine, **⁷**; 4,4′-dichloro-2,2′-bipyridine, **⁸**; 4,4′-dimethoxycarbonyl-2,2′-bipyridine, **9**), $[Re(\equiv CAr') (Tp') (CO)_2]^+$ (Tp' $=$ tris(3,5-dimethyl-1-pyrazolyl)borohydride, **10**), and [Re- $(\equiv CC_6H_4-R)(pdpp)(CO)_2(O_3SCF_3)]^+$ (**13**, R = OMe, **a**; Me, **b**; H, **c**; Cl, **d**; Br, **e**; CN, **f**) are studied and compared. The molecular structures of **⁷**'CHCl3**, 10**, **12f**, **13a**'CH3OH'H2O, and **13d**'2CH₂Cl₂ are determined by X-ray crystallography and reveal $Re \equiv C$ distances in the 1.766(8)-1.786(7) Å range. HF-SCF calculations on the model compounds $[Re(\equiv CC_6H_5)(H_2-V_6H_6)]$ $PCH=CHPH_2)_2Cl$ ⁺ (**2m**), $[Re(\equiv CC_6H_5)(PH_3)_2(H_2O)(CO)Cl$ ⁺ (**3m**), and $[Re(\equiv CC_6H_5)(H_2-O)$ $PCH=CHPH₂)(CO₂(OH)⁺$ (13m) suggest that the HOMO is π (Re \equiv C-Ph) and the LUMO is $\pi^*(\text{Re} \equiv \text{C}-\text{Ph})$. CI-singles calculations on the excited state of optimized **2m** indicate that the lowest energy UV-vis absorption of **²**-**6**, **¹⁰**, and **¹³** originates from a HOMO to LUMO spin-forbidden transition. This is identified as ${}^3[\pi(\text{Re}\equiv\text{CAr}) \rightarrow \pi^*(\text{Re}\equiv\text{CAr})]$, where d(Re) \rightarrow p(\equiv C) MLCT character is apparent and the p(\equiv C) orbital and phenyl π system are conjugated. The UV-vis absorption spectra of **⁷**-**⁹** are significantly different, and their lowest energy absorption is assigned d(Re) $\rightarrow \pi^*(X_2$ -bpy). The rhenium(V) benzylidyne complexes are highly emissive at room temperature and 77 K. The combination of spectroscopic studies and theoretical calculations suggest that the emitting state of **²**-**6**, **10**, and **13** is ${}^{3}[\pi(\text{Re}=\text{CAT})\rightarrow \pi^{*}(\text{Re}=\text{CAT})]$ but that of **7**-**9** is d(Re) $\rightarrow \pi^{*}(X_{2}$ -bpy). The emission energies in dichloromethane can be adjusted from 520 to 610 nm by variation of the benzylidyne and ancillary ligands. Their electrochemical behaviors are examined and provide further evidence to support the excited-state assignment.

Introduction

Fluid luminescence from metal alkylidyne complexes was first observed in $1985¹$ However, emissive complexes containing a metal-carbon triple bond remain rare. To our knowledge, the only reported examples are the d² species $[W(\equiv CPh)(L_2)X(CO)_2]$ (L_2 = tmen (*N,N*,- N , N -tetramethylethylenediimine), $(py)_2$, dppe $(1,2$ -bis-(diphenylphosphino)ethane); $X = Cl$, Br, I),¹ Cp(CO)-
[P(OMe)₃]M=CR (M = Mo, W; R = Ph, o -tolyl),² and $[P(\overline{OMe})_3]M \equiv \hat{C}R$ (M = Mo, W; R = Ph, o -tolyl),² and $[Os(\equiv CPh)(NH_3)_5](O_3SCF_3)_3^3$ and the d¹ derivatives Re- $(\equiv$ CAr')(PPh₃)(H₂O)X₃ (Ar' = 2,4,6-C₆H₂Me₃; X = Cl, Br).⁴ Hopkins has studied the molecular structures of the redox compounds $W(\equiv CPh)(dmpe)_2Br$ and $[W(\equiv CPh)-$

 $(dmpe)_2Br][PF_6]$, and examination of their electronic absorption spectra at high resolution provided a direct assignment of the frontier molecular orbitals.⁵ McElwee-White reported the observation of the lowest excited states of tungsten and molybdenum arylcarbyne complexes by laser flash photolysis.6 A number of accounts on alkylidyne/carbyne-based photochemical reactions have appeared,⁷ although the nature of the excited states remains elusive.

Following the work by Schrock and Williams on d2 rhenium benzylidyne complexes,8 our studies have revealed that congeners bearing phosphine ligands exhibit a long-lived emissive excited state in fluid solution.⁹ We have continued to develop this theme with the purpose of (1) defining the nature of the emissive excited state in order to gain further insight into the photochemistry of metal benzylidyne complexes

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and (2) exploring the relationship between molecular structure and excited-state properties, such as excitedstate energy, nonradiative and radiative decay rate constants, and excited-state redox potentials. A key component of this work is the convenient synthesis of a series of rhenium(V)-substituted benzylidyne complexes with different ancillary ligands. Rhenium alkylidyne complexes are relatively rare compared to the group 6 analogues, 10 and previous routes do not allow for the modification of both the alkylidyne and auxiliary groups. We now present synthetic strategies to two series of luminescent rhenium(V) benzylidyne complexes: (1) 2,4,6-trimethylbenzylidyne species containing various phosphorus or nitrogen donor ligands (**2**-**10**) and (2) para-substituted benzylidyne derivatives supported by *^o*-phenylenebis(diphenylphosphine) (pdpp) (**13a**-**f**). Detailed spectroscopic investigations demonstrate that the anticipated tuning of the excited-state properties in these complexes is possible.

Experimental Section

Spectroscopic Procedures. Infrared spectra were recorded with KBr disks on a BIO-RAD FTS165 FT-IR spectrophotometer. Mass spectra were obtained on a Finnigan Mat 95 mass spectrometer. Elemental analyses were performed by Butterworth Laboratory, U.K. ¹H, ¹³C, and ³¹P NMR spectra (ppm, in CDCl₃ unless otherwise stated) were obtained on a JEOL 270 or a Bruker DRX 300 or 500 multinuclear FT-NMR spectrometer with tetramethylsilane (¹H and ¹³C NMR) and H_3PO_4 (³¹P NMR) as internal references. UV-vis absorption spectra were obtained on a Milton Roy Spectronic 3000 diode-array spectrophotometer.

Emission Spectra and Lifetimes. Steady-state emission spectra were recorded on a SPEX 1681 FLOUROLOG-2 series F111AI spectrometer and corrected for monochromator and photomultiplier efficiency and xenon lamp stability. The absolute emission quantum yield was measured by the method of Demas and Crosby¹¹ using quinine sulfate in 0.1 N sulfuric acid as the standard, where the emission quantum yield is 0.546 with 355 nm excitation.

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Emission lifetimes were measured with a Quanta Ray DCR-3 pulsed Nd:YAG laser system (pulse output 355 nm, 8 ns). The emission signals were detected by a Hamamatsu R928 photomultiplier tube and recorded on a Tektronix model 2430 digital oscilloscope.

Cyclic Voltammetry. Cyclic voltammetry was performed with a Princeton Applied Research model 175 universal programmer and a model 273 potentiostat. Glassy carbon was used as the working electrode, with $Ag-AgNO₃$ (0.1 mol dm⁻³ in acetonitrile) as the reference electrode and platinum wire as the counter electrode. The supporting electrolyte was *n*-tetrabutylammonium hexafluorophosphate (0.1 mol dm-3). Ferrocene was added as an internal standard.

Materials. 4,4′-Dimethoxycarbonyl-2,2′-bipyridine ((Me- O_2C ₂-bpy),¹² 4,4'-dichloro-2,2'-bipyridine (Cl₂-bpy),¹² and Re-[C(O)Ar′](CO)5 ⁸ were prepared by literature methods. All other reagents were used as received. The synthesis of [Re- $(\equiv$ CAr')(CO)₄Cl][O₃SCF₃] (**1**), [Re(\equiv CAr')(pdpp)₂Cl][ClO₄] (**2**), $[Re(\equiv CAr')L_2(CO)(H_2O)Cl]^+$ (L = PPh₃, **3**; P(C₆H₄OMe-*p*)₃, **4**; PMePh₂, **5**), and $[Re(\equiv CAr)(dppe)(CO)_2Cl][ClO_4]$ (6) have been reported previously.9 All manipulations were carried out under a nitrogen atmosphere using standard Schlenk techniques unless otherwise stated.

Syntheses. Satisfactory elemental analysis and infrared and mass spectral data were obtained for all complexes (see Supporting Information).

 $[Re(\equiv CAr') (bpy)(CO)_2 Cl] ClO_4$ (7). A mixture of 1 (0.40) g, 0.65 mmol) and bpy (0.12 g, 0.78 mmol) in toluene (20 cm3) was refluxed overnight. The resultant pale yellow precipitate was filtered and dissolved in methanol (5 cm^3) . LiClO₄ (0.5 g) was added to the solution, and precipitation with diethyl ether gave a yellow solid. Recrystallization from an acetonitrile/ chloroform mixture afforded golden yellow crystals. Yield: 0.11 g, 25%. ¹H NMR (CD₃CN) 2.31 (s, 3, *p*-Me), 2.35 (s, 6, *o*-Me), 6.92 (m, 2, aryl H), 7.85, 8.44, 8.63, 9.34 (4 × m, 8, py H); 13C NMR (CD3CN) 28.1 (*p*-Me), 30.2 (*o*-Me), 126.0-165.2 (py and aryl C), 207.4, 194.0 (CO).

 $[Re(\equiv CAr)(X_2 \cdot bpy)(CO)_2Cl]ClO_4 (X = Cl, 8; MeO_2C, 9).$ A mixture of **1** (0.40 g, 0.65 mmol) and Cl2-bpy (0.18 g, 0.80 mmol) or $(MeO_2C)_2$ -bpy (0.21 g, 0.77 mmol) in tetrahydrofuran (40 cm3) was refluxed for 16 h. The solvent was removed in vacuo, and the residue was dissolved in methanol (10 cm^3) . The crude product was precipitated with $LiClO₄$ (0.5 g) and then diethyl ether. Recrystallization by diffusion of diethyl ether into a chloroform solution gave yellow crystals. Yield: **8**, 0.16 g, 34%; **9**, 0.12 g, 24%.

For **8**: 1H NMR (CD3CN) 2.16 (s, 3, *p*-Me), 2.25 (s, 6, *o*-Me), 6.89 (s, 2, aryl H), 7.93-8.82 (m, 6, py H); ¹³C NMR (CD₃CN) 27.4 (*p*-Me), 30.9 (*o*-Me), 125.2-157.7 (m, py and aryl C), 194.4, 207.6 (CO).

For **9**: 1H NMR 2.37 (s, 3, *p*-Me), 2.44 (s, 6, *o*-Me), 4.11 (s, 6, CO2Me), 6.83 (s, 2, aryl H), 8.16-9.38 (m, 6, py H); 13C NMR 25.1 (*p*-Me), 30.8 (*o*-Me), 54.4 (CO2*Me*), 124.2-157.6 (m, py and aryl C), 164.5 (CO₂), 194.8, 207.9 (CO).

[Re(t**CAr**′**)(Tp**′**)(CO)2][O3SCF3] (10).** A solution of **1** (0.20 g, 0.36 mmol) and KTp′ (0.14 g, 0.42 mmol) in THF (20 cm3) was refluxed for 16 h to give a yellow precipitate. This was filtered, washed with THF, and recrystallized by diffusion of diethyl ether into an acetonitrile solution to yield yellow crystals. Yield: 0.064 g, 22%. 1H NMR 2.04 (s, 3, *^p*-Me), 2.40- 2.47 (m, 24, Tp′ Me and *o*-Me), 6.03 (s, 3, Tp′ H), 6.97 (s, 1, aryl H), 7.05 (s, 1, aryl H); 13C NMR 11.5, 12.6, 12.8, 15.4, 15.9, 18.9 (Tp′-Me), 20.6 (*p*-Me), 22.6 (*o*-Me), 107.8-153.5 (m, Tp′ and aryl C), 192.8 (CO).

 $Re[C(0)C_6H_4R-p](CO)_{5}$ (11, $R = OCH_3$ (a), CH_3 (b), H (c), **Cl (d), Br (e), CN (f)).** A modified version of Schrock's method was used.⁸ $\text{Re}_2(\text{CO})_{10}$ (1.0 g, 1.53 mmol) was added to freshly

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prepared sodium amalgam (0.1 g of Na, 4.4 mmol, 10 g of Hg) in tetrahydrofuran (40 cm3), and the mixture was stirred for 12 h at room temperature. The resultant bright orange solution was filtered, and p -R-C₆H₄COCl (3.06 mmol, R = OCH₃, 0.52 g; CH₃, 0.40 cm³; H, 0.36 cm³; Cl, 0.40 cm³; Br, 0.67 g; CN, 0.50 g) was added to the filtrate. After 1 h of stirring at room temperature, the mixture was filtered and the solvent was removed in vacuo. Recrystallization of the residue in dichloromethane/methanol afforded yellow crystals.

For **11a**: yield 0.68 g, 48%; 1H NMR 3.86 (s, 3, CH3), 6.94 (d, 2, $J_{HH} = 9.0$, *m*-H), 7.56 (d, 2, $J_{HH} = 9.0$, o -H); ¹³C NMR 55.5 (OMe), 113.3, 129.6, 146.9, 162.1 (aryl C), 181.2 (ax-CO), 183.1 (eq-CO), 242.2 (C=O).

For **11b**: yield 0.66 g, 48%; 1H NMR 2.38 (s, 3, CH3), 7.23 (d, 2, $J_{HH} = 8.0$, *m*-H), 7.43 (d, 2, $J_{HH} = 8.0$, σ -H); ¹³C NMR 21.3 (CH3), 127.0, 129.0, 141.5, 151.5 (aryl C), 181.2 (ax-CO), 183.0 (eq-CO), 244.6 (C=O).

For **11c**: yield 0.58 g, 44%; 1H NMR 7.41-7.50 (m, aryl H); 13C NMR 126.3, 128.4, 130.9, 154.3 (aryl C), 181.1 (ax-COax), 182.9 (eq-CO), 246.0 (C=O).

For **11d**: yield 0.69 g, 47%; 1H NMR 7.42 (m, aryl H); 13C NMR 127.8, 128.6, 137.2, 152.1 (aryl C), 180.8 (ax-CO), 182.7 $(eq$ -CO), 244.1 $(C=0)$.

For **11e**: yield 0.77 g, 49%; ¹H NMR 7.35 (d, 2, $J_{HH} = 8.5$, *m*-H), 7.58 (d, 2, $J_{HH} = 8.3$, ρ -H); ¹³C NMR 126.1, 128.3, 131.9, 132.8 (aryl C), 181.1 (ax-CO), 183.0 (eq-CO), 244.9 (C=O).

For 11f: yield 0.61 g, 44%; ¹H NMR 7.46 (d, 2, $J_{HH} = 8.5$, *m*-H), 7.75 (d, 2, *J*_{HH} = 8.5, *o*-H); ¹³C NMR 113.8 (CN), 118.3, 125.6, 132.6, 157.0 (aryl C), 180.5 (ax-CO), 182.2 (eq-CO), 245.9 $(C=O)$.

 $Re[C(0)C_6H_4R$ - $pl(pdpp)(CO)_3$ (12, $R = OCH_3$ (a), CH_3 **(b), H (c), Cl (d), Br e), CN (f)).** A mixture of $Re[C(O)C_6H_4R$ *^p*](CO)5 (**11a**-**f**, 0.87 mmol) and pdpp (0.39 g, 0.87 mmol) in tetrahydrofuran (40 cm^3) was refluxed for 12 h. The solvent was removed in vacuo, and the residue was recrystallized in dichloromethane to give yellow to orange crystalline solids.

For **12a**: yield 0.51 g, 69%; 1H NMR 3.79 (s, 3, CH3), 6.72- 7.78 (m, 28, aryl H); 13C NMR 55.2 (CH3), 112.2 (*p*-C), 128.0- 148.9 (m, aryl C), 160.6 (*ipso*-C), 193.1 (d, ²J_{CP} = 4.6, ax-CO), 196.6 (d, ² J_{CP} = 10.7, eq-CO), 197.3 (d, ² J_{CP} = 10.7, eq-CO), 263.7 (C=O); ^{31}P NMR 44.2.

For **12b**: yield 0.54 g, 74%; 1H NMR 2.28 (s, 3, CH3), 7.00- 7.79 (m, 28, aryl H); ¹³C NMR 21.2 (CH₃), 127.2-141.9 (m, aryl C), 153.1 (*ipso*-C), 193.2 (t, ²J_{CP} = 5.4, ax-CO), 196.5 (d, ${}^{2}J_{CP}$ = 10.7, eq-CO), 197.2 (d, ² J_{CP} = 10.7, eq-CO), 265.8 (t,
 ${}^{2}J_{CP}$ = 10.7, C=O); ³¹P NMR 44.2.
For **12c**: yield 0.50 g, 70%; ¹H NMR 7.08–7.80 (m, aryl H);

¹³C NMR 126.6-141.8 (m, aryl C), 155.5 (*ipso-*C), 193.2 (t, ²*J*_{CP} $=$ 5.3, ax-CO), 196.2 (d, ² J_{CP} = 10.7, eq-CO), 196.9 (d, ² J_{CP} = 10.7, eq-CO), 268.1 (C=O); ³¹P NMR 44.1 .

For **12d**: yield 0.57 g, 77%; 1H NMR 7.07-7.79 (m, aryl H); 13C NMR 127.4-141.8 (m, aryl C), 153.6 (*ipso*-C), 193.0 (ax-CO), 196.1 (d, ² J_{CP} = 10.7, eq-CO), 196.9 (d, ² J_{CP} = 10.7, eq-CO), 265.1 (C=O); ^{31}P NMR 44.1.

For **12e**: yield 0.58 g, 74%; 1H NMR 7.04-7.80 (m, aryl H); 13C NMR 123.7-141.6 (m, aryl C), 154.2 (*ipso*-C), 193.0 (ax-CO), 196.3 (d, ² J_{CP} = 10.2, eq-CO), 196.7 (d, ² J_{CP} = 10.2, eq-CO), 265.1 (C=O); ^{31}P NMR 44.1.

For **12f**: yield 0.53 g, 72%; 1H NMR 6.95-7.82 (m, aryl-H); 13C NMR 111.6 (CN), 119.1, 125.6-141.6, 158.7 (aryl C), 192.9 (d, ² J_{CP} = 9.1, ax-CO), 195.7 (d, ² J_{CP} = 10.7, eq-CO), 196.5 (eq-CO), 266.3 (C=O); ^{31}P NMR 43.8.

[Re(t**CC6H4R-***p***)](pdpp)(CO)2(O3SCF3)][O3SCF3] (13, R**) **OCH3 (a), CH3 (b), H (c), Cl (d), Br (e), CN (f)).** Trifluoromethanesulfonic anhydride (40 mm3, 0.23 mmol) was added to complexes **12a**-**^f** (0.23 mmol) in dichloromethane (10 cm^3) at -40 °C. After 30 min, the volume of the solution was reduced in vacuo at $0 °C$ to 5 cm³ and diethyl ether (40 cm³) was added to precipitate the product. Recrystallization in dichloromethane/diethyl ether/methanol mixtures at -20 °C afforded yellow crystals.

For **13a**: yield 0.08 g, 32%; 1H NMR 3.89 (s, 3, CH3), 6.61 (d, 2, $J_{HH} = 9.0$, $m-H$), 6.71 (d, 2, $J_{HH} = 9.0$, $o-H$), 7.27-7.98 (m, 24, aryl H); ¹³C NMR 56.8 (CH₃), 114.9 (CF₃), 126.1-137.9 (m, aryl C), 166.6 (*ipso*-C), 185.6 (d, ²J_{CP} = 8.7, CO), 186.1 (d, $^{2}J_{\rm CP} = 8.4, \text{ CO}$, 301.5 (t, ²*J*_{CP} = 10.0, Re=C); ³¹P NMR 39.7.

For **13b**: yield 0.09 g, 35%; 1H NMR 2.35 (s, 3, CH3), 6.12 $(d, 2, J_{HH} = 8.2, m-H)$, 6.94 $(d, 2, J_{HH} = 8.2, o-H)$, 7.32-8.00 (m, 24, aryl H); 13C NMR 22.8 (CH3), 126.0-149.8 (m, aryl C), 185.2 (d, ²*J*_{CP} = 8.3, CO), 185.6 (d, ²*J*_{CP} = 8.4, CO), 300.4 (t, ²*J*_{CP} = 10.0, Re=C); ³¹P NMR 39.5.

For **13c**: yield 0.08 g, 32%; ¹H NMR 6.20 (d, 2, *J*_{HH} = 7.3, *m*-H), 6.94 (d, 2, *J*_{HH} = 7.3, *o*-H), 6.93–8.07 (m, 24, aryl H); *m*³C NMR 125.8-142.1 (m, aryl C), 184.9 (d, ²*J*_{CP} = 8.6, CO), 185.4 (d, ² J_{CP} = 8.6, CO), 299.7 (t, ² J_{CP} = 10.1, Re=C); ³¹P NMR 39.4.

For **13d**: yield 0.09 g, 36%; ¹H NMR 6.12 (d, 2, *J*_{HH} = 8.5, *m*-H), 7.05 (d, 2, *J*_{HH} = 8.5, *o*-H), 7.31–8.00 (m, 24H, aryl H); ¹³C NMR 125.9-143.7 (m, aryl C), 184.6 (d, ²*J*_{CP} = 8.2, CO), 185.1 (d, ² J_{CP} = 8.2, CO), 297.6 (t, ² J_{CP} = 10.3, Re=C); ³¹P NMR 39.4.

For **13e**: yield 0.09 g, 34%; ¹H NMR 6.06 (d, 2H, $J_{HH} = 8.5$, *m*-H), 7.24 (d, 2H, $J_{HH} = 8.5$, σ -H), 7.38-8.01 (m, 24H, aryl H); ¹³C NMR 126.0-140.7 (m, aryl C), 184.6 (d, ² $J_{CP} = 8.5$, CO), 185.1 (d, ²*J*_{CP} = 8.5, CO), 297.7 (t, ²*J*_{CP} = 10.2, Re=C); ³¹P NMR 39.4.

For **13f**: yield 0.08 g, 30%; ¹H NMR (CD₃CN) 6.74-8.06 (m, aryl H); ¹³C NMR (CD₃CN) 126.9-145.9 (m, aryl C), 183.4 (d, ${}^{2}J_{\rm CP} = 8.0, \text{CO}$, 183.8 (d, ${}^{2}J_{\rm CP} = 7.8, \text{CO}$), 302.5 (t, ${}^{2}J_{\rm CP} = 9.8, \text{CO}$ $Re \equiv C$); ³¹P NMR (CD₃CN) 39.6.

Structural Determination. Intensity data for **10**, **12f**, $13a$ ^{\cdot}CH₂OH \cdot H₂O, and $13d$ \cdot ²CH₂Cl₂ were collected at 301 K on a Rigaku AFC7R diffractometer with graphite-monochromated Mo Kα radiation ($λ = 0.71073$ Å) using $ω$ -2θ scans at a speed of 16.0 deg min-1. Intensity data were corrected for Lorentz and polarization effects. The structures were solved by Patterson methods, expanded by Fourier techniques (*PATTY*13), and refined by full-matrix least-squares using the software package *TeXsan*¹⁴ on a Silicon Graphics Indy computer.

Crystallographic data are summarized in Table 1. For complex **10**, a crystallographic asymmetric unit consists of one complex cation and one $\rm CF_3SO_3^-$ anion. $\rm All$ 45 non-H atoms were refined anisotropically, H(1) bonded to B(1) was located in the difference Fourier synthesis and its positional parameters were refined, and 32 H atoms at calculated positions with thermal parameters equal to 1.3 times that of the attached C atoms were not refined. For complex **12f**, all 49 non-H atoms were refined anisotropically, and 28 H atoms at calculated positions were not refined. For complex **13a**'CH3OH'H2O, all 65 non-H atoms were refined anisotropically and 31 H atoms at calculated positions were not refined. The H atoms of the methanol molecule and water molecule were not located. For complex $13d \cdot 2CH_2Cl_2$, all 53 non-H atoms of the complex cation, the S atom of the $CF_3SO_3^-$ anion, and the 4 Cl atoms of the dichloromethane molecules were refined anisotropically. The O, F, and C atoms of the anion and solvent molecules were refined isotropically. A total of 33 H atoms at calculated positions with thermal parameters equal to 1.3 times that of the attached C atoms were not refined.

The crystal structures of **⁷**'CHCl3, **12a**, and **12d** have also been determined and are presented in the Supporting Information. The quality of the crystal data for 7⁻CHCl₃ was relatively poor and resulted in large standard deviations; nevertheless, its structural parameters are discussed.

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Table 1. Crystallographic Data

 ${}^{a}R = \sum (||F_{0}| - |F_{c}||)\sum |F_{0}|$. ${}^{b}R_{w} = [\sum w(|F_{0}| - |F_{c}|)^{2}/\sum w|F_{0}|^{2}]^{1/2}$. ${}^{c}S = [\sum w(|F_{0}| - |F_{c}|)^{2}/(n-p)]^{1/2}$.

Figure 1. Geometry of the model molecule $[Re(\equiv CC_6H_5)(H_2-H_6)$ $PCH=CHPH_2)_2Cl$ ⁺ (**2m**).

Molecular-Orbital Calculations. Hartree-Fock selfconsistent-field (HF-SCF)15 and single-excitation configuration interaction (CI-singles)16 calculations were performed using the Gaussian 94/DEF program package¹⁷ on a Silicon Graphics Indigo 2 workstation.

The symmetry of the model compounds $[Re(\equiv CPh)(H_2-d_1)]$ $PCH=CHPH₂)₂Cl⁺$ (2m, Figure 1), $[Re(=CPh)(PH₃)₂Cl(CO) (OH_2)$ ⁺ (3m; see Supporting Information), and $[Re(\equiv CPh)(H_2-$ PCH=CHPH₂)(CO)₂(OH)]⁺ (13m; see Supporting Information) may only be as high as *C*2*v*, *Cs*, and *Cs* for **2m**, **3m**, and **13m**, respectively, depending on the orientation of ligands. These symmetries were chosen for the calculations because full optimizations for the models indicate that they are the most stable, plus the metrical parameters of the crystal data show that $[Re(\equiv CAr')(pdpp)_{2}Cl]^{+}$ (2), $[Re(\equiv CAr')(PPh_{3})_{2}(CO)(H_{2}O)$ Cl^+ (3),⁹ and $\text{[Re}(\equiv CC_6H_5Cl-p)(pdpp)(CO)_2(O_3SCF_3)^+$ (13d) have virtual $C_{2\nu}$, C_s , and C_s symmetry, respectively.

The main optimized bond distances (Å) and angles (deg) are as follows: in **2m** Re=C 1.76, C-Ph 1.44, Re-Cl 2.54, Re-P 2.52, P-Re-P 81.25; in **3m** Re=C 1.76, C-Ph 1.43, Re-Cl 2.52, Re-C 1.99, Re-O 2.28; in **13m** Re=C 1.79, C-Ph 1.43, Re-C 2.04, Re-O 2.01, Re-P 2.57, P-Re-P 80.38, C-Re-^C 93.37.

The ab initio calculations utilized the quasirelativistic effective core potential (ECP) developed by Hay and Wadt;¹⁸ only the outermost electrons of each atom were treated explicitly. For the Re atom, this includes electrons in the 5s, 5p, 5d, and 6s orbitals, and for P and Cl, this includes 3s and 3p electrons. All the electrons were included for C, O, and H atoms. The double-*ú* valence Gaussian basis set associated with the pseudopotential is adopted. The basis sets were taken as Re (8s6p3d)/[3s3p2d], P (3s3p)/[2s2p], Cl (3s3p)/[2s2p], C (10s5p)/[3s2p], O (10s5p)/[3s2p], and H (4s)/[2s].

Results and Discussion

Synthesis and Characterization. The synthetic strategy⁹ used to prepare the $2,4,6$ -trimethylbenzylidynerhenium(V) complexes $[Re(\equiv CAr')(pdp)p_zCl]^+$ (2), $[Re(\equiv CAr')L_2(CO)(H_2O)Cl]^+$ (L = PPh₃, **3**; P(C₆H₄OMe p)₃, **4**; PMePh₂, **5**), and $[Re(\equiv CAr)(dppe)(CO)_2Cl]^+$ (**6**) is also applicable for the synthesis of derivatives containing nitrogen donor ligands: $[Re(\equiv CAr')(X_2-bpy)(CO)_2$ -Cl^{$+$} (X = H, 7; Cl, **8**; CO₂CH₃, **9**) and [Re(\equiv CAr')(Tp')- $(CO)_2$ ⁺ (10) are afforded by the treatment of [Re(\equiv CAr['])- $(CO)_4Cl$ ⁺ (1) with the corresponding diimine or KTp'. They are the first examples of rhenium benzylidyne complexes supported by nitrogen donors. The IR spectra of **⁷**-**⁹** are similar, with two strong CO bands at *ca*. 2090 and 2030 cm-¹ while the CO bands of **10** appear at 2076 and 2010 cm^{-1} .

Efficient synthesis of the para-substituted benzylidynerhenium(V) compounds $[Re(\equiv CC_6H_4R-p)](pdpp)$ -

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 $(CO)_2(O_3SCF_3)[O_3SCF_3]$ (13, R = OMe, a; Me, b; H, c; Cl, **d**; Br, **e**; CN, **f**) has been developed by modification of Schrock's method.8 The para-substituted benzoylrhenium(I) complexes $Re[C(O)C_6H_4R$ -*p*](CO)₅ (11a-f) are readily formed from $\text{Re}_2(\text{CO})_{10}$, and substitution of CO with pdpp yield the air- and moisture-stable species Re[C(O)C6H4R-*p*](pdpp)(CO)3 (**12a**-**f**). Trifluoromethanesulfonic anhydride is subsequently used as an oxygenabstracting reagent to give the series of benzylidyne derivatives **13a**-**^f** (Scheme 1).

The IR spectra of benzoylrhenium(I) complexes Re- [C(O)R](CO)5 (**11a**-**f**) are comparable and depict a sharp peak at *ca*. 2135 cm^{-1} and a broad band in the 2055- 1975 cm⁻¹ region for the CO stretching frequencies. In the 13C NMR spectra, no systematic effect by the parasubstituents is observed upon the benzoyl resonances, which are located at $242.2-246.0$ ppm. The ¹³C NMR signal for the CO ligand trans to the benzoyl group appears at *ca*. 181 ppm, while those for the four equatorial COs are found at *ca*. 183 ppm. The IR spectra of $Re[C(O)C_6H_4R-p](pdpp)(CO)_3$ (12a-f) contains three CO bands at *ca*. 2010, 1940, and 1910 cm-1, which are consistent with a *fac* configuration at the rhenium center. The 13C NMR shifts of the benzoyl carbons $(265-268$ ppm) and axial $(193$ ppm) and equatorial (196 and 197 ppm) CO groups are downfield from those of the pentacarbonyl derivatives **11a**-**f**. The IR spectra of $[Re(\equiv CC_6H_4R-p)](pdpp)(CO)_2(O_3SCF_3)][O_3-$ SCF3] (**13a**-**f**) show CO bands at *ca*. 2110 and 2070 cm^{-1} , while their ¹³C NMR spectra reveal resonances for the benzylidyne carbon in the 297.6-302.5 ppm range.

Complexes **⁷**-**⁹** with substituted bpy ligands are stable for *ca*. 1 month in air in the solid state but decompose after only a matter of days in acetonitrile solution. In contrast, **²**-**⁶** (containing phosphine ligands) and **10** (with Tp') can be stored in air for several months. Complexes **13a**-**^f** are slightly air-sensitive and decompose in aerobic dichloromethane after a few hours at room temperature.

Figure 2. Perspective view of $[Re(CAr)(Tp')(CO)_2]^+$ (10; 35% probability ellipsoids).

Figure 3. Perspective view of $Re[C(O)C_6H_4CN \text{-} p]$ (pdpp)-(CO)3 (**12f**; 40% probability ellipsoids).

Crystal Structures. Perspective views of complexes **10**, **12f**, **13a**[·]CH₃OH·H₂O, and **13d**·2CH₂Cl₂ are presented in Figures 2-5, respectively; those of complexes **⁷**'CHCl3, **12a**, and **12d** are given in the Supporting Information. Selected bond distances and angles are listed in Table 2.

The Re \equiv C distances in this work (\AA ; 1.772(7) for **7**,

Figure 4. Perspective view of $[Re(\equiv CC_6H_4OMe-p)(pdp)]$ $(CO)_{2}(O_{3}SCF_{3})$ ⁺ (13a; 40% probability ellipsoids).

Figure 5. Perspective view of $[Re(\equiv CC_6H_4Cl-p)(pdpp)$ - $(CO)_2(O_3SCF_3)]^+$ (**13d**; 40% probability ellipsoids).

1.786(7) for **10,** 1.769(10) for **13a,** 1.766(8) for **13d**) are typical for rhenium-carbon triple bonds¹⁰ and are comparable to those in the previously reported structures of **2** and **3** (1.802(5) and 1.784(8) Å, respectively).9 The distances from the benzylidyne carbon to the neighboring *ipso*-carbon (Å; 1.49(2) for **7**, 1.424(10) for **10**, 1.43(1) for **13a**, 1.43(1) for **13d**) are slightly shorter than normal C-C single bonds and suggest *^π*-electron delocalization throughout the $Re \equiv C - Ar$ moiety (Ar $=$ substituted phenyl). The $Re \equiv C$ distances also approach those in $[Re(\equiv CCH_2 tBu)(dppe)_2F][BF_4]$ (1.772(7) Å),^{10j} $[Re(\equiv CNHMe)(dppe)_2Cl][BF_4]$ (1.798(30) Å),^{10k} and

Table 2. Selected Bond Distances (Å) and Angles (deg) for Complexes 10, 12f, 13a'**CH3OH**'**H2O, and 13d**'**2CH2Cl2**

u NUILZUIZ					
Compound 10					
$Re(1) - N(1)$	2.158(6)	$Re(1) - C(2)$	1.972(9)		
$Re(1) - N(3)$	2.142(5)	$Re(1) - C(3)$	1.786(7)		
$Re(1) - N(5)$	2.226(6)	$C(3)-C(4)$	1.424(10)		
$Re(1) - C(1)$	1.962(9)				
$N(1) - Re(1) - N(3)$	85.6(2)	$N(1) - Re(1) - C(3)$	96.4(3)		
$N(1) - Re(1) - N(5)$	83.3(2)	$N(3)-Re(1)-C(3)$	96.9(2)		
$C(1) - Re(1) - C(3)$	90.3(3)	$N(5)-Re(1)-C(3)$	179.5(3)		
$C(2)-Re(1)-C(3)$	91.4(3)	$Re(1)-C(3)-C(4)$	178.8(6)		
Compound 12f					
$Re(1) - P(1)$	2.425(2)	$C(4)-C(5)$	1.54(1)		
$Re(1) - C(1)$	1.952(9)	$Re(1) - C(4)$	2.199(9)		
$C(1) - Re(1) - C(4)$	176.6(3)	$P(1) - Re(1) - C(4)$	90.5(2)		
$C(2)-Re(1)-C(4)$	89.5(3)	$P(2)-Re(1)-C(4)$	83.7(2)		
$C(3)-Re(1)-C(4)$	88.0(3)	$Re(1)-C(4)-C(5)$	116.7(6)		
Compound $13a \cdot CH_3OH \cdot H_2O$					
$Re(1) - P(1)$	2.461(2)	$Re(1) - C(3)$	1.769(10)		
$Re(1) - O(4)$	2.231(6)	$C(3)-C(4)$	1.43(1)		
$P(1) - Re(1) - C(3)$	93.5(3)	$C(1) - Re(1) - C(3)$	89.8(4)		
$P(2)-Re(1)-C(3)$	94.5(3)	$C(2)-Re(1)-C(3)$	88.5(4)		
$O(4)-Re(1)-C(3)$	175.6(3)	$Re(1) - C(3) - C(4)$	175.3(7)		
Compound $13d \cdot 2CH_2Cl_2$					
$Re(1) - P(1)$	2.472(2)	$Re(1) - C(3)$	1.766(8)		
$Re(1) - O(3)$	2.228(5)	$C(3)-C(4)$	1.43(1)		
$P(1) - Re(1) - C(3)$	94.8(2)	$C(1) - Re(1) - C(3)$	89.2(3)		
$P(2)-Re(1)-C(3)$	94.6(2)	$C(2)-Re(1)-C(3)$	89.3(3)		
$O(3) - Re(1) - C(3)$	177.3(3)	$Re(1)-C(3)-C(4)$	175.9(6)		

 $[Re(\equiv CNH_2)(dppe)_2Cl][BF_4]$ (1.802(4) Å).^{10k} The Re $\equiv C-$ Cipso angles are all close to linearity as expected.

The substantial trans influence of the benzylidyne ligand is illustrated in these structures. In complex **7**, a trans chloride group $(C(3)-Re(1)-Cl(1)$ 174.7(4)^o) results in a rather long $Re(1)-Cl(1)$ distance of 2.485(2) Å, similar to that in **2** (2.497(2) Å).9 For **10** with the facial Tp' ligand, the nitrogen atom N(5) trans to the benzylidyne moiety $(N(5)-Re(1)-C(3)$ 179.5(3)^o) is located further from the metal center $(A; Re(1) - N(5))$ 2.226(6), *cf*. Re(1)-N(1) 2.158(6), Re(1)-N(3) 2.142(5)). For **13a** and **13d**, a relatively long Re-O contact to the trans triflate group is revealed in each case (Å; 2.231(6) for **13a**, 2.228(5) for **13d**).

The structures of the benzoyl complexes **12a**, **d**, and **f** are similar, and depict a distorted octahedral configuration. The benzoyl ligand is trans to one of the three CO groups with virtually linear $C(1)-Re(1)-C(4)$ angles (176.5(2)° for **12a**, 177.2(2)° for **12d**, and 176.6(3)° for **12f**). The angles around the benzoyl carbon C(4) approach 120°, as expected for sp² hybridization (for example, in complex **12f** $Re(1) - C(4) - C(5)$ 116.7(6)°, $Re(1)-C(4)-O(4)$ 128.2(7)°, $C(5)-C(4)-O(4)$ 115.0(8)°).

Absorption Spectra and Electronic Structures. The UV-vis absorption data of the benzylidyne complexes are summarized in Table 3. The absorption spectra of $[Re(\equiv CAr')(pdpp)_{2}Cl]^{+}$ (2), $[Re(\equiv CAr')(PR_{3})_{2}$ - $(CO)(H_2O)Cl$ ⁺ (3-5), $[Re(\equiv CAr')(dppe)(CO)_2Cl$ ⁺ (6), $[Re(\equiv CAr)(Tp')(CO)_2]^+$ (**10**), and $[Re(\equiv CC_6H_4-R)-$ (pdpp)(CO)2(O3SCF3)]⁺ (**13a**-**f**) possess similar features: an intense band in the 310-354 nm region and weak absorption in the 370-430 nm range. The absorption spectrum of **13c** is shown in Figure 6. We note that complex **13a** displays an intense peak at 351 nm, but inexplicably, no other lower energy absorptions are

Table 3. UV-**Vis Spectral Data of Benzylidyne Complexes at Room Temperature***^a*

complex	λ_{max} , c nm (ϵ_{max} , d dm ³ mol ⁻¹ cm ⁻¹)
2 $[Re(\equiv CAr')(pdpp)2Cl]$ ⁺	318 (13 900), 410 (450)
3 [Re(\equiv CAr')(PPh ₃) ₂ (CO)(H ₂ O)Cl] ⁺	321 (13 900), 410 (1380)
4 [Re(=CAr'){P(C ₆ H ₄ OMe- p) ₃ } ₂ (CO)(H ₂ O)Cl] ⁺	320 (13 100), 420 (320)
5 $[Re(=CAr')(PPh2Me)2(CO)(H2O)Cl]+$	320 (12 100), 410 (470)
6 [Re(\equiv CAr')(dppe)(CO) ₂ Cl] ⁺	330 (7290), 430 (400)
7 [Re(\equiv CAr')(bpy)(CO) ₂ Cl] ^{+ b}	231 (31 500), 260 (20 300)
	313 (22 400), 320 (24 000)
	340 (16 400)
8 $[Re(\equiv CAr')(Cl_2-bpy)(CO)_2Cl]^+$	262 (22 200), 310 (17 700)
	318 (20 800), 346 (15 800)
9 [Re(\equiv CAr'){(CO ₂ Me) ₂ -bpy}(CO) ₂ Cl] ⁺	253 (20 800), 330 (23 600)
	337 (26 200), 390 (9560)
10 $[Re(\equiv CAr') (Tp')(CO)_2]^+$	354 (15 400), 425 (970)
13a [Re ($\equiv CC_6H_4OMe$)(pdpp)(CO) ₂ (O ₃ SCF ₃)] ⁺	351 (25 200)
13b $[Re(\equiv CC_6H_4Me)(pdpp)(CO)_2(O_3SCF_3)]^+$	326 (18 700), 425(300)
13c $[Re(\equiv CC_6H_5)(pdpp)(CO)_2(O_3SCF_3)]^+$	310 (14 800), 425 (580)
13d $[Re(\equiv CC_6H_4Cl)(pdpp)(CO)_2(O_3SCF_3)]^+$	325 (19 200), 370 (1440)
13e $[Re(\equiv CC_6H_4Br)(pdpp)(CO)_2(O_3SCF_3)]^+$	329 (23 200), 375 (3170)
13f [Re(\equiv CC ₆ H ₄ CN)(pdpp)(CO) ₂ (O ₃ SCF ₃)] ⁺	313 (18 900), 380 (860)

a In CH₂Cl₂ unless otherwise stated. *b* In MeCN. *c* Error ± 2 nm. *d* Error ± 1 in last significant figure.

Figure 6. UV-vis absorption spectrum of $[Re(\equiv CC_6H_5) (\text{pdpp})(CO)_2(O_3SCF_3]^+$ (13c) in CH_2Cl_2 at room temperature.

observed even at a concentration of 5×10^{-3} mol dm⁻³. Although it is difficult to precisely define the energy of the lowest energy absorption shoulder, it is apparent from Table 3 that the different electron-withdrawing para-substituents in complexes **13d**-**^f** cause a gradual blue shift (ca. 3500 cm^{-1}).

Complex **10** containing Tp′ displays a significant red shift to 354 nm for the high-energy band. Significantly, the absorption spectra of **⁷**-**⁹** containing para-substituted bipyridine are different from the other benzylidyne complexes in this work. A strong peak ($\epsilon > 2 \times$ 10^4 dm³ mol⁻¹ cm⁻¹) centered at 320, 318, and 337 nm for **7**, **8**, and **9**, respectively, is observed and is accompanied by a higher energy shoulder. The lowest energy absorption appears at *ca*. 340, 346, and 390 nm for **7**, **8**, and **9**, respectively, and is also intense ($\epsilon > 1.5$ \times 10⁴ dm³ mol⁻¹ cm⁻¹).

In solvent-dependent UV-vis studies on complexes **2** and **10**, no significant shifts were observed when the solvent is changed from chloroform to acetonitrile. The absorption spectra of **13e** in different solvents showed significant but irregular changes, and this is ascribed to the activity of the coordinated triflate group. A solvent effect on the electronic absorption bands of this class of Re(V) benzylidyne complexes is, therefore, discounted.

As an aid to interpreting the electronic absorption

Figure 7. HOMO (bottom) and LUMO (top) of **2m**. Coefficients are derived from the HF calculation.

spectra, HF-SCF calculations¹⁵ have been performed on the model molecules $[Re(\equiv CC_6H_5)(H_2PCH=CHPH_2)_2$ -Cl^{$+$} (**2m**, Figure 1; *x*, *y*, *z* axis defined), [Re $(\equiv CC_6H_5)$ -(PH3)2(CO)(H2O)Cl]⁺ (**3m**; see Supporting Information), and $[Re(\equiv CC_6H_5)(H_2PCH=CHPH_2)(CO)_2(OH)]^+$ (13m; see Supporting Information). The calculated energy and composition of the near frontier orbitals for **2m**, **3m**, and **13m** are summarized in the Supporting Information and reveal that the HOMO for each complex (for **2m**, Figure 7 bottom) is the π (Re=C-Ph) orbital with π bonding between d_{*xz*}(Re) and p_{*x*}(\equiv C) (2**m** d_{*xz*} 27.96%, p_{*x*}(\equiv C) 6.93%, p_x(Ph) 34.94%; **3m** d_{xz} 32.65%, p_x(\equiv C) 1.89%, p_x (Ph) 59.33%; **13m** d_{xz} 26.83%, p_x(\equiv C) 4.03%, p_x(Ph) 35.63%). This is different from previous assignments of the HOMO in related tungsten complexes to a metal d orbital that is nonbonding with respect to the benzylidyne ligand.4,5 The LUMO (for **2m**, Figure 7 (top)) is a $\pi^*(\text{Re} \equiv \text{C}-\text{Ph})$ orbital with antibonding between d_{xz}-(Re) and $p_x (\equiv C)$ (2m d_{xz} 20.49%, $p_x (\equiv C)$ 21.19%, $p_x (Ph)$

Table 4. Luminescence Data for Benzylidyne Complexes at Room Temperature*^a*

complex	λ_{max} , nm ^c	ϕ_{em} ^d	τ , μ s ^d
2 $[Re(\equiv CAr')(pdp)p_2Cl]^+$	573	0.042	2.08
3 $[Re(\equiv CAr') (PPh_3)_2 (CO)(H_2O)Cl]^+$	580	0.0020	2.25
4 [Re(=CAr'){P(C ₆ H ₄ OMe- p) ₃ } ₂ (CO)(H ₂ O)Cl] ⁺	588	0.0067	1.76
5 [Re(=CAr')(PPh ₂ Me) ₂ (CO)(H ₂ O)Cl] ⁺	611	0.0046	0.95
6 [$Re(\equiv CAr')$ (dppe)(CO) ₂ Cl] ⁺	567	0.0035	3.35
7 [Re(\equiv CAr')(bpy)(CO) ₂ Cl] ^{+ b}	555	0.00025	0.43
8 $[Re(\equiv CAr')(Cl_2-bpy)(CO)_2Cl]^+$	575	0.00012	0.07
9 $[Re(\equiv CAr')\{(CO_2Me)_2-bpy\}(CO)_2Cl]^+$	580	0.0028	0.61
10 $[Re(\equiv CAr')(Tp')(CO)_2]^+$	585	0.0027	1.48
13a $[Re(\equiv CC_6H_4OMe)(pdpp)(CO)_2(O_3SCF_3)]^+$	520	0.00015	0.02
13b $[Re(\equiv CC_6H_4Me)(pdpp)(CO)_2(O_3SCF_3)]^+$	527	0.0018	0.22
13c $[Re(\equiv CC_6H_5)(pdpp)(CO)_2(O_3SCF_3)]^+$	530	0.0026	0.53
13d $[Re(\equiv CC_6H_4CI)(\bar{p}dpp)(CO)_2(O_3SCF_3)]^+$	536	0.012	2.18
13e $[Re(\equiv CC_6H_4Br)(pdpp)(CO)_2(O_3SCF_3)]^+$	537	0.011	2.42
13f [Re(\equiv CC ₆ H ₄ CN)(pdpp)(CO) ₂ (O ₃ SCF ₃)] ⁺	559	0.016	4.84

a $\lambda_{\rm ex}$ = 300–350 nm; in CH₂Cl₂ unless otherwise stated. *b* In MeCN. *c* Error \pm 2 nm. *d* Error \pm 10%.

49.02%; **3m** d_{*xz*} 21.56%, p_{*x*}(≡C) 26.39%, p_{*x*}(Ph) 42.04%; **13m** d_{xz} 18.61%, $p_x (\equiv C)$ 26.30%, $p_x (Ph)$ 43.31%). By comparing the composition of these orbitals, we find that the percentage of d*xz* in the LUMO is smaller than that in the HOMO whereas the percentage of $p_x(\equiv C)$ increases tremendously in the LUMO. Hence, if the lowest energy transition involves these frontier orbitals, then this can be formulated as $\pi(\text{Re} \equiv \text{C}-\text{Ph}) \rightarrow \pi^*$ (Re=C-Ph) with $d_{xz} \rightarrow p_x (\equiv C)$ MLCT character, where the $p_x(\equiv C)$ orbital and phenyl π system are conjugated.

To aid in the assignment of the transitions in the absorption spectra of these complexes, a CI-singles calculation¹⁶ was performed on the excited states of the optimized model molecule **2m** (see Supporting Information), where all the orbitals and electrons of the ground state were included to take into account the electronic correlation for the excited states. The calculated lowest energy absorption at 461 nm (HOMO to LUMO triplet transition, see Supporting Information) can be visualized as a spin-forbidden transition of π (Re=C-Ph) \rightarrow *π*^{*}(Re≡C−Ph) (*vide supra*). From the absorption spectrum of **2**, a weak shoulder at *ca*. 410 nm is assigned to this transition. In addition, the calculated HOMO to LUMO singlet transition at 290 nm correlates with the 318 nm band in the absorption spectrum.

On the basis of these calculations, the lowest energy absorption for the rhenium(V) benzylidyne complexes **2–6**, **10**, and **13a–f** can be assigned to a $\frac{3}{\pi}$ (Re=CAr) $\rightarrow \pi^*(\text{Re} \equiv \text{CAT})$] transition and the higher energy band to $\frac{1}{\pi}$ (Re=CAr) $\rightarrow \pi^*$ (Re=CAr)]; both of these contain notable $d(Re) \rightarrow p(=C)$ MLCT character, and the $p_x(=C)$ orbital and phenyl π system are conjugated. Furthermore, if this is correct, the energies of these two absorption bands should be affected by substituents on the benzylidyne ligand but relatively insensitive to different auxiliary ligands. This is indeed consistent with our observation that while the variation of phosphine ligands only alters the absorption bands of complexes **²**-**⁶** to a small extent, the nature of the parasubstituents on the benzylidyne ligand in **13a**-**^f** significantly affects the absorption energies (Table 3). The fact that the absorption bands are not solvent-dependent is consistent with the proposal that the electronic transitions are not pure MLCT.

The UV-vis absorption spectra of complexes **⁷**-**⁹** containing 4,4′-substituted bipyridine ligands are different, and hence the above assignment of the electronic transitions is not applicable. The spectral features of

these complexes resemble those for a series of rhenium- (I) N-heterocyclic carbene complexes containing 4,4′-

substituted bipyridine ligands, [HNCH₂CH₂NHCRe(X₂bpy)(CO)₃]⁺, where the absorption peaks at $350-400$ nm are assigned to a $[d(Re) \rightarrow \pi^*(X_2-bpy)]$ ¹MCLT transition, and a correlation between the 1MLCT transition energy and the Hammett parameters (*σ*) of the substituents X was demonstrated.¹⁹ We therefore tentatively assign the lowest energy shoulders of complexes **⁷**-**⁹** to the $[d(Re) \rightarrow \pi^*(X_2-bpy)]$ ¹MLCT transition. Electronwithdrawing substituents such as $CO₂CH₃$ should lower the $\pi^*(X_2$ -bpy), and thus the absorption energy, and correspondingly, the lowest energy absorption of complex 9 containing (CH₃O₂C)₂-bpy appears at *ca*. 390 nm compared with that for **7** (340 nm) and **8** (346 nm).

Luminescence Studies and Modification of Excited-State Properties. All of the rhenium(V) benzylidyne complexes emit in fluid solution at room temperature upon excitation at 300-350 nm (Table 4). The profile of their emission spectra are very similar: a broad and structureless band which is indicative of a large degree of vibrational coupling like the solution emission of [Ru(bpy)₃]²⁺.²⁰ The emission energies λ_{max} range from 520 to 610 nm, and the emission lifetimes vary from 0.02 to 4.84 *µ*s. The relatively long lifetimes imply that the transitions involved are spin-forbidden. The emitting states of these complexes are assigned to the lowest energy transition. For **²**-**6**, **¹⁰**, and **13a**-**f**, the lowest energy excited states have been identified as $\frac{3[\pi(\text{Re} \equiv \text{CAr}) \rightarrow \pi^*(\text{Re} \equiv \text{CAr})]}{(vide \text{ supra})}$. This assignment predicts that the excited-state energy is affected by the electronic properties of the benzylidyne group, and changing the para-substituents would affect both the *π*(Re=CAr) and *π*^{*}(Re=CAr) energies. Electrondonating substituents such as $OCH₃$ and $CH₃$ should lower the energy of π (Re=CAr) and raise that of π ^{*}- $(Re \equiv CAr)$ and consequently increase the energy gap. Conversely, electron-withdrawing substituents such as Cl, Br, and CN should reduce this gap. Thus, a correlation between the emission energy of **13a**-**^f** and the Hammett parameters (*σ*) of the benzylidyne parasubstituents is anticipated and indeed observed (Figure 8). This demonstrates that the emission energy of these derivatives can be tuned and implies (1) association

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Figure 8. Correlation plot of room-temperature emission energy E_{em} vs Hammett parameters σ of the substituents for complexes $[Re(\equiv CC_6H_4R-p)(pdpp)(CO)_2(O_3SCF_3)]^+$ (**13af**), $R = -0.99$, slope $= -993 \pm 49$.

between the emitting state and the benzylidyne moiety and (2) delocalization of π (Re=C) and π ^{*}(Re=C) into the phenyl π system. The tunable range of the excited-state energy is 1300 cm^{-1} . This is the first example of systematic modification of the excited-state energy of metal benzylidyne complexes by variation of the benzylidyne moiety.

In addition to effects on emission energy, the variation of benzylidyne and the auxiliary ligands also influences the emission lifetime and quantum yield. As shown in Table 4, lifetimes in the 20 ns to 4.8 *µ*s range and quantum yields from 1.2×10^{-4} up to 4.2×10^{-2} are observed, and these indicate that the nature of the ligands has considerable influence on the excited-state decay kinetics. For complexes $2-6$ with the $(\equiv CAr')$ moiety, the nonradiative decay rate constants increase as excited-state energies decrease, and this is consistent with the energy-gap law as described previously. $9,21$

We have assigned the lowest energy transition and, hence, the emitting state of complexes **⁷**-**⁹** containing 4,4'-substituted bipyridine ligands to $d(Re) \rightarrow \pi^*(X_2$ -bpy) MLCT. The red shift of the emission bands from **7** (555 nm) to **8** (575 nm) and **9** (580 nm), which bear more electron-withdrawing substituents, is consistent with this. Mayr reported that replacing tmen by bipyridine in the tungsten derivative $[W(\equiv CPh)(\text{tmen})(CO)_2Cl]$ totally quenched the fluid solution emission.¹

At 77 K, the Re(V) complexes $[Re(\equiv CCH_2C_6H_4R$ p)(dppe)₂Cl][BF₄] (R = H, Me)^{10j} with unconjugated alkylidyne groups were found to emit at 545 and 535 nm, respectively.²² These are comparable to the emission energies of the benzylidyne complexes (Table 4) and hence this implies that the emitting state is associated with the $[Re \equiv C]$ moiety. The effect of different solvents on the emissive behavior of complexes **2**, **10**, and **13e** were studied, but like the absorptions, no notable effects were seen upon the emission energies, quantum yields, and lifetimes for **2** and **10**. An irregular effect on the luminescence of **13e** was observed, but again the activity of the coordinated triflate is the likely cause.

The 77 K emission spectra of the benzylidyne complexes show well to poorly resolved vibrational structures. The corrected emission spectra of **13f** in the solid

Figure 9. Corrected emission spectra of $[Re(\equiv CC_6H_4CN)$ - $(\text{pdpp})(\text{CO}_2(\text{O}_3\text{SCF}_3))^+$ (13f) in CH₂Cl₂ at room temperature (-) and in solid state at 77K (- - -), $\lambda_{\rm ex}$ 310 nm, emission intensities are normalized.

Table 5. Electrochemical Data*^a*

complex	$E_{\rm ox}$, V	E_{red} , V
$2b$, c	1.48 d,h	
3 ^b	1.83 , e 1.39 d,h	$-1.31 f$
$\mathbf{4}^b$	1.50 , e 1.31 ^d	$-1.76 f$
5 ^b	$1.78,$ e 1.51 ^d	$-1.72 f$
6		-1.20 , β -1.54 β
13a		-0.89 , $g - 1.91$ f
13 b		-0.81 , β -1.84 β
13c		$-0.77, \frac{g}{r} - 1.81$ f
13d		-0.75 , β -1.80 β
13e		-0.68 , $\epsilon - 1.67$, $\epsilon - 1.81$
13f		$-0.53.5 - 1.51$ f,h

^a Measured in CH₂Cl₂ vs SCE; error \pm 0.02 V; irreversible (E_{pa} and *E*pc potentials for anodic oxidation and cathodic reduction waves, respectively) unless otherwise stated. *^b* In MeCN. *^c* Reference 9. ^{*d*} Re(VI)/(V). *e* Phosphine oxidation. *f* Benzylidyne reduction. *^g* Re(V)/(IV). *^h* Reversible.

state at 77 K and in dichloromethane solution at room temperature are shown in Figure 9. To our knowledge, this is the first observation of vibronic emission in metal alkylidyne complexes. Gaussian deconvolution of the vibronic structured 77 K emission spectra can be used to obtain the vibrational spacing $\hbar \omega_M$, excited-state distortion S_M , $v'_M = 0 \rightarrow v_M = 0$ emission energy E_{em} (0-0), and full-width at half-maximum of the individual vibrational components $\tilde{v}_{1/2}$.²³ The fitting results are available in the Supporting Information and show vibrational progression $\hbar \omega_M$ between 1080 and 1218 cm^{-1} and excited-state distortion S_M in the 1.21-1.72 range. The latter indicate that the emitting states are not pure MLCT, for which S_M values are typically *ca*. $1.0.²⁴$

Electrochemistry. Cyclic voltammetry was used to study the electrochemistry of the benzylidyne complexes (Table 5). The cyclic voltammograms of **³**-**⁵** are similar and reveal two oxidations and one reduction, all of which are irreversible. The first oxidation is tentatively assigned to Re(V)/Re(VI), and the second is assigned to oxidation of the phosphine ligands. The reduction is ascribed to occur at the benzylidyne ligand.

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Figure 10. Plot of room-temperature emission energy *E*em vs ground-state reductive potential E° [(CAr)^{0/-}] for [$\text{Re}(\equiv C$ - C_6H_4R -*p*)(pdpp)(CO)₂(O₃SCF₃)]⁺ (**13a–f**), $R = -0.96$, slope $= -3068 \pm 422.$

No oxidation wave is observed for complexes **6** and **13a**-**f**, which contain two coordinated CO groups, even up to 2.3 V vs SCE. Two irreversible reduction waves (the second for **13f** is reversible) are observed; they are tentatively assigned to metal-centered ($Re(V) \rightarrow Re(IV)$) and benzylidyne ligand reductions, respectively.

The electrochemical data of **13a**-**^f** provide further support for the assignment of the $\pi^*(\text{Re} \equiv \text{CAT})$ excited state. Reduction of the benzylidyne group results in addition of an electron into the $\pi^*(\equiv$ CAr) orbital. If the emitting state is associated with π^* (\equiv CAr), a linear relationship should exist between the emission energy (*E*em) and the reductive potential for the benzylidyne group (*E*°[(CAr)0/-], denoted with footnote *f* in Table 5). This is indeed observed for complexes **13a**-**^f** (Figure 10). Importantly, this correlation illustrates that the excited state is associated with the benzylidyne group and that emission in this system is occurring from states which have a common electronic origin.

Conclusion

Synthetic routes to rhenium(V) benzylidyne complexes have been developed so that both benzylidyne

and auxiliary ligands can be conveniently modified, and this has paved the way for exploring the excited-state properties of these luminescent compounds. Data from electronic absorption and emission spectra, electrochemistry, and molecular-orbital calculations provide compelling evidence to suggest that the excited state is $[\pi(\text{Re} \equiv \text{CAT}) \rightarrow \pi^*(\text{Re} \equiv \text{CAT})]$ with d(Re) \rightarrow p(\equiv C) MLCT character, where the $p(\equiv C)$ orbital and phenyl π system are conjugated. Significantly, the excited-state properties associated with this chromophore can be tuned by varying the benzylidyne and auxiliary ligands. Hence, the emission energy can be modified from 520 to 610 nm in CH_2Cl_2 , a range of 2840 cm⁻¹. The excited-state lifetimes can be adjusted from 20 ns to 4.84 *µ*s, many of which are significantly longer than those in related Mo, $W,$ ^{1,2} and Os³ alkylidyne systems, so that Stern-Volmer kinetic experiments are possible.

Acknowledgment. We are indebted to Prof. A. J. L. Pombeiro for providing samples of $[Re(\equiv CCH_2C_6H_4R$ p)(dppe)₂Cl][BF₄] (R = H, Me) and for helpful discussions. We thank The University of Hong Kong, the Croucher Foundation, and the Hong Kong Research Grants Council for financial support. M.C.-W.C. is grateful for a University Postdoctoral Fellowship from The University of Hong Kong.

Supporting Information Available: Tables of crystal data, atomic coordinates, calculated hydrogen coordinates, anisotropic displacement parameters, and bond distances and angles for **⁷**'CHCl3, **¹⁰**, **12a**, **12d**, **12f**, **13a**'CH3OH'H2O, and $13d·2CH₂Cl₂$, the HF-SCF calculated energy and composition of the near frontier orbitals for **2m**, **3m**, and **13m**, excited state assignment of **2m** by CI-singles calculation, analytical data for all complexes, emission parameters calculated from Gaussian deconvolution, and geometry of **2m**, **3m**, and **13m** (99 pages). Ordering information is given on any current masthead page.

OM971054T