Fluorocarbene, fluoroolefin, and fluorocarbyne complexes of Rh†

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The manuscript reports the synthesis, characterization, and analysis of electronic structure in a series of complexes of small perfluorocarbon ligands with the (PNP)Rh fragment (where PNP is a diarylamido/bis(phosphine) pincer ligand). Reactions of (PNP)Rh(TBE) as the source of (PNP)Rh with CHF3 and C2HF5 produced perfluoroalkylidene complexes (PNP)Rh=CF2 and (PNP)Rh=CF3. (PNP)Rh=CF2 could also be obtained via the reaction of (PNP)Rh(TBE) with Me2SiCF2/CsF, with an admixture of (PNP)Rh(C2F4), where TBE = tert-butylethylene. Abstraction of fluoride from these neutral (PNP)RhCFx complexes was successful, although only abstraction from (PNP)Rh=CF2 allowed unambiguous identification of the Rh product, [(PNP)Rh=CF]+. DFT computational studies allowed comparison of relative energies of (PNP)Rh(C2F4) and [(PNP)Rh(C2F3)]+ isomers as well as comparisons between the electronic structure of the =CF2, C2F4, and ≡CF complexes and their hydrocarbon analogues.

Introduction

Organofluorine chemistry’s major impact on the world of industrial chemistry has inspired many investigations into the unique properties that are inherent to molecules and materials containing C–F bonds. Transition metal complexes containing perfluorocarbon ligands are an important subset of these studies since they exhibit distinctive bonding properties1 and can mediate perfluoroalkyl–carbon bond forming processes.2

Group 9 perfluoroalkylidenes have garnered interest in the past decade after Hughes developed a simple reductive method for making Ir=CFR complexes (Fig. 1, top) from iridium-fluoroalkyl precursors.3 These complexes have been analyzed in the context of their potential intermediacy in perfluoroeflin metathesis,4 and more recently the Baker group has shown that analogous cobalt perfluoroalkenens (Fig. 1, top)5 are capable of undergoing a [2 + 1] cycloaddition with CF2 (ref. 6) and [2 + 2] cycloaddition with C2F4.7 Analogous chemistry was also reported for a difluorocarbene complex of Ni(0).8 Baker has also shown that cationic cobalt(III) difluorocarbenes could undergo migratory insertion into perfluoroalkyl ligands, possibly providing a blueprint for transition metal catalyzed perfluoroolefin polymerization.9

The only family of isolable terminal fluoromethylidyne complexes known to date are the Cp*M(CO)2(CF) compounds (M = Cr, Mo, W) reported by Hughes and co-workers (Fig. 1, bottom).10 The Andrews group has reported a number of fluoromethylidyne complexes of the general formula X,M(CF) (Fig. 1, bottom; X = halogen) via trapping laser ablated metal atoms in argon/halocarbon matrices at ca. 10 K.11 Most of the isolable terminal carbene complexes are complexes of metals of groups 6,12,13 7,14,15 and 8.15,16 A few examples are known for group 5.17 In group 9, one 18-electron complex has been fully characterized for Ir by Bergman et al. (Fig. 1, bottom),18 and one square planar 16-electron complex was mentioned in passing for Rh by Werner et al.,19 as a component of a reaction mixture.

Fig. 1 Perfluoroalkylidenes from Hughes and Baker. Isolated fluorocarbenes by Hughes, the Ir carbine by Bergman, and matrix-trapped fluorocarbenes by Andrews.

Hughes’ Reductive Route to Perfluoroalkylidenes

Baker’s Co Analogue

| Hughes | Bergman | Andrews |
|--------|---------|---------|
| ![Image](image1.png) | ![Image](image2.png) | ![Image](image3.png) |

**Hughes**

**Baker**

**Andrews**

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The “concentration” of metal carbyne complexes in the middle of the transition metal series can be compared with similar trends for other metal-element multiple bonds.\textsuperscript{20,21} In this report, we describe the synthesis, characterization, and analysis of electronic structure of a rare cationic fluoromethylidyldyne complex of Rh, as well as related Rh perfluoroalkylidene complexes.

Results and discussion

Synthesis of CF₂, C₂F₄, and CFCF₃ complexes

We recently reported reactions of the (PNP)Rh fragment with aryl carboxylates, including aryl-oxygen oxidative addition.\textsuperscript{22} The (PNP)Rh aryl-oxygen oxidative addition product of phenyl trifluoroacetate, (PNP)Rh(C==OCF₃)(OPh), could be thermolysed to produce (PNP)Rh(CO) and (PNP)Rh(CF₃)(CO)(OPh) as major products. In that report, we noted that some other unidentified products were evident in trace amounts. We continued to be intrigued by one apparent trace product in particular that was consistently observed in 2–5% yield. For it, we observed a doublet of triplets both in the $^{31}$P{1H} NMR and $^{19}$F NMR spectra (coupling constants: $J_{\text{Rh-P}} = 146$ Hz, $J_{\text{F-P}} = 49$ Hz, $J_{\text{F-F}} = 30$ Hz). These multiplicities implied a P$_2$RhF$_2$ NMR spin system – rather unexpected given the three fluorines in the CF$_3$ group of the starting material. We noted that the $^{19}$F NMR chemical shift was itself uncommon (95.6 ppm) and in the range reported for various difluorocarbene complexes (i.e., $M$=CF$_3$).\textsuperscript{5,10w,23,24} The observed $J_{\text{F-P}}$ was also quite similar to that of Grushin’s trans-(PPh$_3$)$_2$RhCF$_2$ which possessed a $J_{\text{F-P}} = 33$ Hz.\textsuperscript{23}

We hypothesized that this minor side product might be (PNP)Rh(CF$_2$) and attempted an independent synthesis of it based on the procedure of Grushin et al. that yielded trans-(PPh$_3$)$_2$RhCF$_2$.\textsuperscript{23} Indeed, treatment of (PNP)Rh(TBE) (TBE = tert-butyl-ethylene) with CsF/Me$_3$SiCF$_3$ (Ruppert’s reagent) resulted in complete consumption of (PNP)Rh(TBE) and the formation of (PNP)Rh(CF$_2$) and (PNP)Rh(CF$_3$) in about 85 : 15 ratio (NMR evidence).\textsuperscript{22} We were able to isolate (PNP)Rh(CF$_2$) in 52% yield and of >98% purity by recrystallization. The $^{31}$P{1H} NMR and $^{19}$F NMR spectra of (PNP)Rh(CF$_2$) obtained in this fashion were identical to that of the impurity we observed in the reaction of Scheme 1.

(PNP)Rh(TBE) showed no reaction with Me$_3$SiCF$_3$ alone. Similar to the other cases of use of CsF/Me$_3$SiCF$_3$, we propose that these reagents generate a CF$_3$ anion equivalent that displaces TBE and then loses fluorine, resulting in the formal transfer of CF$_2$ to Rh. Alternatively, CsF/Me$_3$SiCF$_3$ could be generating free CF$_3$ which then binds to Rh. Using the CsF/Me$_3$SiCF$_3$ protocol, we could not avoid the formation of (PNP)Rh(C$_2$F$_4$) due to the generation of free CF$_3$ from the CsF/Me$_3$SiCF$_3$ mixture. CF$_3$ was observed by $^{19}$F NMR spectroscopy in control experiments where CsF and Me$_3$SiCF$_3$ were mixed in C$_6$D$_6$ and heated at 80 °C. No reaction was observed when (PNP)Rh(CF$_2$) was treated with another equivalent of CsF/Me$_3$SiCF$_3$. This contrasts the reactivity of Baker’s difluorocarbenes cobalt(i) complexes\textsuperscript{2} which undergo a $[2 + 1]$ cycloaddition with free CF$_2$ to form cobalt tetrafluoroethylene complexes.

To date, we have not been able to formulate a reasonable proposal for how (PNP)Rh(CF$_2$) could be formed from (PNP)Rh(COCF$_3$)(OPh) (Scheme 1). The formation of $M$=CF$_2$ by fluoride migration from M–CF$_3$ is well preceded\textsuperscript{26} and is likely the key step in forming (PNP)Rh(CF$_2$); the difficulty is with conceiving of a plausible fate of the other atoms of the original phenyl trifluoroacetate molecule.

Goldman et al. documented formation of (PCP)Ir(CF$_2$) in a reaction of a (PCP)Ir source with HCF$_3$.\textsuperscript{24} This reaction proceeded via C–H oxidative addition of HCF$_3$ to Ir followed by loss of HF. In a similar vein, we found that (PNP)Rh(TBE) reacted with HCF$_3$ at 80 °C to provide a mixture of compounds containing (PNP)Rh(CF$_2$) as a major product (>80%) with (PNP)Rh(CO) and [(PNP)Rh(CF$_3$)$_2$] as minor products (Scheme 2). Commercial HCF$_3$ contains dinitrogen as an impurity. Hydrolysis of a difluorocarbene complex to a carbonyl complex has precedent,\textsuperscript{27} but attempts to purposefully hydrolyze (PNP)Rh(CF$_2$) proved to be unsuccessful, reminiscent of Baker’s cobalt fluorocarbene complexes.\textsuperscript{5} It is possible that hydrolysis of (PNP)Rh(CF$_2$) only takes place in the presence of HF (a by-product of (PNP)Rh(CF$_3$) generation). We observed no intermediates\textsuperscript{28} in the reaction of (PNP)Rh(TBE) with HCF$_3$, which may indicate that dissociation of TBE\textsuperscript{29} is the rate-limiting step.

An analogous reaction of (PNP)Rh(TBE) with C$_2$HF$_3$ was attempted as a potential means to access (PNP)Rh(CF$_3$). However, the major product of this reaction turned out to be a tetrafluoroethylene complex (PNP)Rh(CF$_3$) (Scheme 2). Dinitrogen impurity in C$_2$HF$_3$ led to the known\textsuperscript{29} [(PNP)Rh]$_2$(μ-N$_2$) as a major side product, whose content could be reduced by degassing C$_2$HF$_3$ using the “freezepump–thaw” technique. (PNP)Rh(CF$_2$) was also observed as a side product composing 13% of the reaction mixture when (PNP)Rh(TBE) was treated with 2 atm of C$_2$HF$_3$ and heated overnight at 80 °C. (PNP)Rh(CF$_3$) could be isolated in

![Scheme 1](https://example.com/scheme1.png)

**Scheme 1** Initial observation of (PNP)Rh==CF$_2$.

![Scheme 2](https://example.com/scheme2.png)

**Scheme 2** Synthesis of rhodium fluorocarbenes and tetrafluoroethylene complexes.
they are probably unavoidable in these molecules, these distances are short enough to be considered weak F⋯H interactions.89 C–F⋯H interactions have been observed in pincer-ligated zirconium complexes bearing a trifluoromethyl as a pendant group, which have also exhibited through-space H–F coupling visible in their 1H NMR spectra.89 However (PNP)Rh(C2F4) and (PNP)Rh=CF2 showed no through-space 13F⋯1H coupling to the isopropyl arms.

Synthesis of cationic fluorenyldiylene

With compounds (PNP)Rh=CF2, (PNP)Rh(=CFCF3), and (PNP)Rh(C2F4) in hand, we contemplating whether one of the fluorides could be removed to yield cationic C2F4 complexes. Hughes et al. previously demonstrated proton-induced loss of fluoride from α-positions of Ir perfluoroalkyls,84 and Baker recently demonstrated a Lewis-acid abstraction of a fluoride from N-heterocyclic fluoroalkenes to yield polyfluorooalkenyl imidazolium salts.84 There is significant precedent for electrophilic abstraction of an anionic heteroatom substituent from late-metal carbene complexes by a Lewis acid.64,85 Trialkylsilylium cations, in the form of their salts with halogenated carborane anions, are powerful Lewis acids with high affinity for fluoride.84 We and others have exploited them in catalytic C–F activation reactions86 and thus a [R3Si]+ reagent appeared perfect for fluoride abstraction.

Reactions of (PNP)Rh=CF2, (PNP)Rh(C2F4), and (PNP)Rh=C(F)CF3 with [Et3SiH–SiEt3][HCB11Cl11] or [(Et3Si)2OTf][HCB11Cl11]77 all generated the Et3SiF by-product, indicating that fluoride abstraction took place in all three cases. However, reactions of (PNP)Rh(C2F4) and (PNP)Rh=C(F)CF3 resulted in mixtures of several products as seen by 19F NMR spectroscopy and typically broad or no signals were observed by 31P[1H] NMR spectroscopy. The reaction mixtures produced from the reaction of (PNP)Rh(C2F4) or of (PNP)Rh=C(F)CF3 with [(Et3Si)2OTf][HCB11Cl11] did regenerate the corresponding starting material when treated with CsF. This indicates that fluoride abstraction from these two isomeric complexes generates isomers of [(PNP)Rh(C2F4)]+ that do not interconvert on the experimental time scale. Although we were not able to identify these compounds experimentally, DFT computational studies were used to investigated possible structures of the [(PNP)Rh(C2F4)]+ isomers (vide infra).

On the other hand, reaction of (PNP)Rh=CF2 with [Et3SiH–SiEt3][HCB11Cl11] cleanly and reproducibly generated a new Rh complex that displayed a P2RhF NMR spin system (Scheme 3).
The key NMR spectroscopic features of this compound were the unusual $^{19}$F NMR chemical shift (66.2 ppm), the very high $J_{C-F}$ coupling constant of 470 Hz, and the rather substantial $J_{Rh-F}$ = 136 Hz.

These spectroscopic data are similar to those exhibited by $\text{Cp}^*\text{(CO)}_2\text{Mo} = \text{CF}$, whose $^{19}$F NMR spectrum contained a singlet at 78.15 ppm, with a large $J_{C-F}$ coupling constant of 556 Hz evident by $^{13}$C NMR spectroscopy. Hughes’s other $\text{Cp}(\text{CO})_2\text{M} = \text{CF}$ (M = Cr, W) complexes also exhibited $^{19}$F NMR chemical shifts in this region with high $J_{C-F}$ coupling constants. Due to limited solubility in non-interactive solvents and the extensive coupling inherent to the fluoromethyldyne $^{13}$C NMR resonance in [(PNP)Rh≡CF]$, it was not observed by $^{12}$C($^{13}$P), $^{12}$C($^{1}$H), nor $^{12}$C($^{1}$F) NMR spectroscopy.

X-ray quality crystals of [(PNP)Rh≡CF][CHB$_11$Cl$_{11}$] were studied using X-ray diffraction to yield a structure fully supportive of a fluorocarbene formulation (Fig. 3). The structural and NMR spectroscopic features of [(PNP)Rh≡CF]$^-$ are best reviewed in comparison with Cl$_3$Rh$^-$ and a few other relevant compounds. Andrews et al. observed IR spectroscopic evidence for Cl$_3$Rh≡CF in reactions of laser-ablated rhodium atoms with CFCl$_3$. A DFT calculation of this product predicted a Rh–C bond length of 1.740 Å and a Rh–C–F bond angle of 143.4°. This compares with our observed Rh–C bond length of 1.702(7) Å and a Rh–C–F bond angle of 173.46°. Although both [(PNP)Rh≡CF]$^-$ and Cl$_3$Rh≡CF are four-coordinate, they contain different numbers of valence electrons: from a hypothetical point of view of a [CF]$^+$ ligand, it is attached to a d$^8$ Rh center in [(PNP)Rh≡CF]$^-$, but to a d$^7$ [Cl$_3$Rh]$^-$ fragment in Cl$_3$Rh≡CF. The geometry of the RhCF unit in [(PNP)Rh≡CF]$^-$ is similar to Bergman’s iridium carbene complex (Scheme 1), which possesses an Ir–C bond length of 1.734(6) Å and an Ir–C–C bond angle of 175.7(4)°.

The Rh–C distance in [(PNP)Rh≡CF]$^-$ is ca. 0.12 Å shorter than that in (PNP)Rh=CF$_2$, consistent with the increase in the Rh–C bond order. The Rh–C bond distance in [(PNP)Rh≡CF]$^-$ is also ca. 0.07 Å shorter than that in Werner’s trans-(P(Pr)$_3$)$_2$ClRh=CC(C(Me)(H)) square planar vinylidene complex.

### Computational studies and discussion

#### DFT structural studies

Modern Density Functional Theory (DFT) is a powerful tool with which to examine electronic structures and bonding trends in organometallic compounds. In addition, application of Natural Bond Orbital (NBO) methods allows insight into some of the subtleties of metal–ligand bonding. The NBO analysis also generates Wiberg Bond Indices (WBI), determined within the natural atomic orbital basis, providing one means of estimating bond orders between atoms. Trends in WBI values are also useful in tracking variations in bond multiplicities.

The newly synthesized family of fluorocarbon complexes (PNP) Rh(C$_2$F$_4$), (PNP)Rh=CF$_2$, and (PNP)Rh≡CF$^-$ prompted a computational comparison with their hypothetical hydrocarbon analogues (PNP)Rh(C$_2$H$_4$), (PNP)Rh=CH$_2$ and [(PNP)Rh≡CH]$^-$ in order to assess the effects of fluorination on the metal–carbon bonding, and, for the carbene and carbyne complexes, to probe the nature and extent of the multiple bonding between the metal and carbon. In addition, since the CF$^+$ ligand is isoelectronic with the well-known ligands NO$^-$ and CO, it was of interest to establish trends in metal ligand and intraligand bonding between [(PNP)Rh≡CF]$^-$, [(PNP)Rh(NO)]$^-$, and (PNP) Rh(CO).

Full molecule DFT studies were performed using the M06 functional and the triple-$\zeta$ LACV3P basis set, which uses extended core potentials on heavy atoms and a 6-311G basis for other atoms, as implemented in the Jaguar suite of programs. Full details are available as ESI.

Selected bond lengths and computed WBI values for the calculated complexes are provided in Table 1, with metric comparisons to the available crystallographic structures reported here. The DFT calculated metrics are in good agreement with crystallographic numbers, giving confidence in the DFT metrics for the unknown complexes. One exception appears in (PNP) Rh(C$_2$F$_4$) in which the C–C distance for the coordinated alkene (1.354 Å) is only slightly longer than that in C$_2$F$_4$ itself (1.318 Å) and is much shorter than all other transition metal complexes of this perfluoroalkene in the Cambridge Structure Database. In contrast, the DFT calculated value (1.416 Å) is in good agreement with other crystallographically determined values and is much more sensible with respect to calculated WBI values (see below).

All the complexes examined here can be formally viewed as square planar d$^8$ compounds, i.e., as complexes of a d$^8$, three-coordinate fragment (PNP)Rh with neutral or cationic ligands. The NBO perspective of bonding interactions in such compounds requires 4-electron/3-center bonds between the pair of trans ligands such that the alkene, carbene, and carbyne ligands of interest are always involved in a shared bonding interaction with the N of the PNP pincer. Clearly contributions to this shared interaction may be weighted differently in each case, and the WBI values should reflect this.

#### Comparison of C$_2$H$_4$ and C$_2$F$_4$ ligands

The bonding between alkene and metal–ligand fragments is well understood, but a comparison between C$_2$H$_4$ and C$_2$F$_4$ coordinated to identical metal–ligand fragments is rare. A
classic intramolecular example involves CpRh(C₂H₄)(C₂F₄)⁴⁺ but the hydrocarbon and fluorocarbon alkenes are necessarily bound to different fragments in this molecule. Fig. 4 illustrates the key Natural Localized Molecular Orbitals (NLMOs)⁴⁵,⁴⁶ arising from NBO calculations of interactions between C₂H₄ and C₂F₄ and the truncated⁴⁵ (PNP)Rh fragment. The bonding orbitals (σ and π) are essentially localized on Rh and the alkene ligand, while the corresponding antibonding NLMOs show significant “tailing” involving the σ and π orbitals on the trans-N of the pincer ligand. This “tailing” is indicative of delocalization of these N electrons into the corresponding σ* and π* components of the Rh-alkene interaction; it is significantly greater for the σ* component and corresponds to the 3-center/4-electron bonding expected between trans-ligands in a d⁸ Rh(Ⅰ) complex. The WBI values indicate significantly greater reduction in C-C bonding and increase in Rh-C bonding in coordinated C₂F₄ than in C₂H₄. This is consistent with the shorter Rh-C distances and with the idea of a more metallacyclopropane structure and stronger Rh–C bonding for the fluorinated alkene complex. Not surprisingly, stronger bonding to the fluorinated alkene results in weaker bonding to the trans ligand, with correspondingly lower WBI values for the Rh-N bonds (Table 1). In the π*-perp NMLO for the C₂F₄ there is also evidence for delocalization from F-lone pairs (see Fig. 4).

There are two Rh → alkene backbonding options involving π*-perp or σ*-in-plane. Clearly the ground state conformation of the C₂F₄ complex utilizes the former, but the latter is available for an in-plane C₂F₄ conformation, leading to a lower barrier for C₂F₄ rotation. Similar arguments for facile rotation of C₂F₄ ligands in Ru(n) complexes have been put forth elsewhere.⁴⁶ The free energy profile for C₂F₄ rotation was calculated using a truncated⁴⁵ version of the PNP ligand (identical to that shown in Fig. 4), and is unusual. Relative to the perpendicular conformation observed in the ground state, two transition states were located. The first, lying 7.1 kcal mol⁻¹ above the ground state, corresponds to a 45 degree rotation about the Rh-alkene bond axis, and the second, lying 8.7 kcal mol⁻¹ above the ground state, is the conformation in which the fluoroalkene lies in the coordination plane. These barriers are low enough in energy that rotation should be fast on the NMR timescale, consistent with the observed NMR data. The barriers contrast with those for the corresponding C₂H₄ analogue, for which the in-plane conformation is a minimum, lying only 0.4 kcal mol⁻¹ above the ground state, and the 45 degree conformation is a transition state lying 2.5 kcal mol⁻¹ above the ground state.

### Table 1: Calculated and crystallographic bond lengths (Å) and Wiberg Bond Indices (WBI)

| Compound | Rh–P̄ave | Rh–N | Rh–C̄ave | C–C | C–X̄ave (ligand) |
|----------|---------|------|---------|-----|-----------------|
| (PNP)Rh(C₂H₄) | 2.332 | 2.098 | 2.165 | 1.394 | 1.088 |
| (PNP)Rh(C₂F₄) | 2.378 | 2.100 | 2.048 | 1.416 | 1.346 |
| (PNP)Rh≡CH₂ | 2.338 | 2.216 | 1.850 | — | 1.104 |
| (PNP)Rh≡CF₂ | 2.331 | 2.156 | 1.864 | — | 1.325 |
| [(PNP)Rh≡CH]⁺ | 2.378 | 2.075 | 1.728 | — | 1.110 |
| [(PNP)Rh≡CF]⁺ | 2.384 | 2.061 | 1.740 | — | 1.247 |
| [(PNP)Rh≡C–CF₄]⁺ | 2.337 | 2.041 | 1.499 | 1.065 |
| (PNP)Rh(CO) | 2.336 | 2.117 | 1.855 | — | 1.152 |
| [(PNP)Rh(NO)]⁺ | 2.393 | 2.019 | 1.778 | 1.141 (NO) | 1.917 |

* DFT calculated (M06/LACV3P**/++*) values are in plain text; X-ray crystallographic values are in bold. WBI values are in italics. This crystallographic value is questionable. See discussion in the text.

### Comparison of CH₂ and CF₂ ligands

Fig. 5 presents the corresponding NLMOs for the CF₂ complex with a truncated⁴⁵ PNP ligand. Those for the CH₂ analogue are very similar, except for the fluorine delocalizations into the π*-NLMO, and are not illustrated here but can be found in the ESI (Fig. S17). The σ and π NLMOs are consistent with a formal double bond between Rh and the CF₂ (or CH₂) ligands, with the p-orbital on C and d-orbital on Rh providing the π-component. These NLMOs look essentially identical to those in Fig. 4, except for a more significant delocalization of the fluorine lone pairs in the π*-perp NLMO. But now there is competition between the rhodium d-orbital and the fluorine lone pair p-orbitals for π-bonding with the carbene carbon, as expected; in (PNP)Rh≡CH₂ only the metal can provide this π-bonding. Consequently π-bonding with fluorines diminishes π-bonding with Rh and, relative to the CH₂ complex, the Rh–C WBI decreases significantly and the Rh–C distance increases; notably the C–F WBI is greater.
than in the previously discussed C2F4 complex. In contrast to the alkene ligands discussed above there is overall weaker metal–carbon bonding to CF2 than CH2 and the corresponding trans-Rh–N WBI value is larger for (PNP)Rh[CF2].

Facile CF2 ligand rotation is expected due to the availability of the π*-perp and π*-in-plane interactions. Calculations on the truncated ligand analogue of the CF2 complex reveal the same trend in conformational energetics observed for the C2F4 complex (vide supra). The in-plane CF2 conformation, with the π*-perp interaction, is the ground state, with two transition states at 45 degree and 90 degree (perpendicular to the coordination plane) lying 2.0 and 4.8 kcal mol\(^{-1}\) higher in energy, respectively. The barriers are consistent with the experimental observation of fast rotation on the NMR timescale. In contrast, the rotation of the corresponding CH2 ligand is even more facile, with the perpendicular, 45 degree and in-plane conformations lying at essentially equal energies.

Comparison of CH+, CF+ and CCF3+ ligands

Fig. 6 presents the corresponding NLMOs for [(PNP)Rh≡CF]+. As with the carbene complexes, those for [(PNP)Rh≡CH]+ are similar except for enhanced “tailing” in the antibonding NLMOs for the CF+ complex. In contrast to the alkene and carbene complexes (vide supra) there is now a second fully engaged π-component for the Rh–C bond involving the in-plane d-orbital and a second p-orbital on the CF+ (or CH+) ligand. In the antibonding NLMOs we see the expected σ-donation from the trans-N in σ*, a small π-donation from trans-N in π*(perp), and a small donation from the Rh-P bonds in π*(in-plane). But once again the largest delocalizations in the π* NLMOs comes from the F lone pairs, interactions that cannot occur in [(PNP)Rh≡CH]+. Consequently the Rh–C WBI for [(PNP)Rh≡CH]+ is substantially larger than that for the CF+ analogue, consistent with the shorter Rh–C distance in the former; as before, a smaller Rh–C WBI in [(PNP)Rh≡CF]+ leads to a larger WBI for the trans-Rh–N bond. In the CCF3 analogue, in which F lone pair participation with the carbyne carbon is removed, a larger Rh–C WBI is calculated, with a correspondingly smaller Rh–N WBI.

Degree of Rh–C multiple bonding

The Rh–C bonding in these carbene and carbyne complexes is inextricably linked to interactions with the trans-N since there are necessarily shared bonding components between trans-ligands, with additional complexities introduced by the fluorine substituents on carbon. So we cannot expect the Rh≡CX2 interaction to be a true double bond, or that in the cationic Rh≡CX to be a triple bond, even though we may draw resonance structures that reflect these prejudices. However, the WBI values for both Rh≡CX2 bonds (CH2 1.250; CF2 1.168) are
significantly larger than unity, though not close to two, while those for the Rh\textsuperscript{CX} cations are significantly larger still (CH\textsubscript{1.714}; CF\textsubscript{1.587}; CCF\textsubscript{3} 1.770), though not close to the bond order of three. Clearly there is significant multiple bonding between Rh and these unsaturated carbon ligands with higher bond orders to these ligands being reflected in lower bond orders to the trans-N.

Comparison of CF\textsuperscript{+}, NO\textsuperscript{+}, and CO ligands
The NLMOs for the CO and NO\textsuperscript{+} complexes are similar to those of the CF\textsuperscript{+} compounds discussed previously and are not shown here. Considering this series of isoelectronic complexes as involving a linear Rh\textendash X\textendash Y array the three resonance forms (A, B, C) for the contiguous \(\pi\)-system are shown in Fig. 7, along with the WBI values for the appropriate bonds in the Rh\textendash C\textendash O, Rh\textendash N\textendash O, and Rh\textendash C\textendash F complexes.

![Fig. 6](image)

**Fig. 6** NLMOs for the bonding interactions between the CF\textsuperscript{+} ligand and the (PNP)Rh fragment. For clarity the PP\textsubscript{Pr} groups have been replaced by PM\textsubscript{2} groups and the aryl part of the pincer truncated to P\textendash CH\textendash CH\textendash N linkers.

![Fig. 7](image)

**Fig. 7** Resonance forms for the \(\pi\)-system in a linear Rh\textendash X\textendash Y ligand array, with WBI values for the bonds in Rh\textendash C\textendash O, Rh\textendash N\textendash O, and Rh\textendash C\textendash F complexes. All three complexes are isoelectronic and no formal charges are shown.

Relative energies of isomeric fluorocarbon ligands
It was of interest to compare the relative energies of the (PNP) Rh(C\textsubscript{2}F\textsubscript{4}) complex with its carbene isomer (PNP)Rh(CF\textsubscript{3}CF\textsubscript{3}). At the DFT/M06/LACV3P\textsuperscript{**} level the free energy of the carbene isomer is found to be 2.4 kcal mol\textsuperscript{-1} uphill from its alkene analogue. Interestingly the carbene CFC\textsubscript{3} ligand lies perpendicular to the (PNP)Rh plane, in contrast to the CF\textsubscript{2} analogue described above, presumably due to steric interactions between the CF\textsubscript{3} and the \(\text{cis}\)-PR\textsubscript{2} groups.

Potential products arising from fluoride abstraction from these isomeric complexes were also subjected to DFT evaluation. Abstraction of fluoride from (PNP)Rh(CF\textsubscript{3}CF\textsubscript{3}) could occur from the \(\alpha\)-position to yield a carbyne complex (PNP)Rh(CF\textsubscript{3})\textsuperscript{+}, analogous to the characterized CF\textsuperscript{+} complex described above, or from the \(\beta\)-position to afford the corresponding isomeric \(\eta\)-perfluorovinyl cation (PNP)Rh(CF\textsubscript{3}C\textsubscript{2}F\textsubscript{3})\textsuperscript{+}. These are found to have almost identical free energies, with the carbyne complex lying only 0.4 kcal mol\textsuperscript{-1} higher than its perfluorovinyl isomer. An \(\eta\)-perfluorovinyl isomer, the potential initial product of fluoride abstraction from the (PNP)Rh(C\textsubscript{2}F\textsubscript{4}) was found to lie 12.1 kcal mol\textsuperscript{-1} above its \(\eta\)-perfluorovinyl analogue. Structures of all these compounds and their relative energies are provided in Fig. 8.

Conclusions
In summary, we have shown that (PNP)Rh perfluorocarbene complexes can be synthesized by treating a Rh(\textit{i}) precursor with

![Fig. 8](image)
Ruppert’s reagent or a fluoroalkane containing a C–H bond. Using silylum reagents, a fluoride can be abstracted from (PNP)Rh–CF₂ to form a cationic fluoromethylidyne. Thus the (PNP)Rh system conveniently allows synthesis and comparison of perfluoroolefin, perfluorocarbene, and perfluorocarbyne complexes. Using DFT calculations we were able to compare the natural localized molecular orbitals of these fluoroorganic complexes to their hypothetical hydrocarbon analogues, as well as to the CO and NO⁺ complexes. We established that the fluoride atoms on the carbene and carbonye ligands participate in π donation to the acceptor orbitals on carbon to compete with back donation from the metal. This resulted in a longer Rh–C bond in the fluorinated complexes compared to their hydrocarbon analogues. However, C₃F₂ was calculated to form a shorter Rh–C bond than the C₂H₄ complex. Calculated Wiberg bond indices also showed that although the unsaturated fluoroorganic ligands have bond orders greater than one to rhodium, the nitrogen trans to these ligands interacts with their antibonding orbital and decreases the bond order to less than a true double and triple bond.

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Notes and references

1 F. L. Taw, A. E. Clark, A. H. Mueller, M. T. Janicke, T. Cantat, B. L. Scott, P. J. Hay, R. P. Hughes and J. L. Kiplinger, Organometallics, 2012, 31, 1484.
2 (a) O. A. Tomashenko and V. V. Grushin, Chem. Rev., 2011, 111, 4475; (b) W. Zhu, J. Wang, S. Wang, Z. Gu, J. L. Aceña, K. Izawa, H. Liu and V. A. Soloshonok, J. Fluorine Chem., 2014, 167, 37; (c) M. Ohashi, H. Shirataki, K. Kikushima and S. Ogoshi, J. Am. Chem. Soc., 2015, 137, 6496.
3 (a) R. P. Hughes, R. B. Laritchev, J. Yuan, J. A. Golen, A. N. Rucker and A. L. Rheingold, J. Am. Chem. Soc., 2005, 127, 15020; (b) C. J. Bourgeois, R. P. Hughes, J. Yuan, A. G. DiPasquale and A. L. Rheingold, Organometallics, 2006, 25, 2908; (c) R. P. Hughes, J. Fluorine Chem., 2010, 131, 1059.
4 J. Yuan, R. P. Hughes, J. A. Golen and A. L. Rheingold, Organometallics, 2010, 29, 1942.
5 D. J. Harrison, S. I. Gorelsky, G. M. Lee, I. Korobkov and R. T. Baker, Organometallics, 2013, 32, 12.
6 G. M. Lee, D. J. Harrison, I. Korobkov and R. T. Baker, Chem. Commun., 2014, 50, 1128.
7 (a) D. J. Harrison, G. M. Lee, M. C. Leclerc, I. Korobkov and R. T. Baker, J. Am. Chem. Soc., 2013, 135, 18296; (b) J. T. Fuller, D. J. Harrison, M. C. Leclerc, R. T. Baker, D. H. Ess and R. P. Hughes, Organometallics, 2015, 34, 5210.
8 D. J. Harrison, A. L. Daniels, I. Korobkov and R. T. Baker, Organometallics, 2015, 34, 5683.
9 M. C. Leclerc, J. M. Bayne, G. M. Lee, S. I. Gorelsky, M. Vasiliiu, I. Korobkov, D. J. Harrison, D. A. Dixon and R. T. Baker, J. Am. Chem. Soc., 2015, 137, 16064.
10 (a) H. Huang, R. P. Hughes, C. R. Landis and A. L. Rheingold, J. Am. Chem. Soc., 2006, 128, 7454; (b) H. Huang, R. P. Hughes and A. L. Rheingold, Dalton Trans., 2011, 40, 47.
11 (a) J. T. Lyon and L. Andrews, Inorg. Chem., 2006, 45, 9858; (b) J. T. Lyon, H.-G. Cho and L. Andrews, Organometallics, 2007, 26, 6373; (c) J. T. Lyon and L. Andrews, Eur. J. Inorg. Chem., 2008, 1047; (d) H.-G. Cho and L. Andrews, Organometallics, 2010, 29, 2211.
12 R. R. Schrock, Acc. Chem. Res., 1986, 19, 342.
13 R. R. Schrock, Chem. Rev., 2002, 102, 145.
14 O. V. Ozerov, L. A. Watson, M. Pink and K. G. Caulton, J. Am. Chem. Soc., 2007, 129, 6003.
15 G. Jia, Coord. Chem. Rev., 2007, 251, 2167.
16 (a) S. R. Caskey, M. H. Stewart, Y. J. Ahn, M. J. A. Johnson, J. L. C. Roswell and J. W. Kampf, Organometallics, 2007, 26, 1912; (b) M. Shao, L. Zheng, W. Qiao, J. Wang and J. Wang, Adv. Synth. Catal., 2012, 354, 2743.
17 (a) R. Ramirez-Contreras, N. Bhuvanesh and O. V. Ozerov, Organometallics, 2015, 34, 1143 and references within; (b) T. Kurogi, P. J. Carroll and D. J. Mindiola, J. Am. Chem. Soc., 2016, 138, 4306.
18 H. F. Luecke and R. G. Bergman, J. Am. Chem. Soc., 1998, 120, 11008.
19 T. Rappert, O. Nürnberg, N. Mahr, J. Wolf and H. Werner, Organometallics, 1992, 11, 4156.
20 W. A. Nugent and J. M. Mayer, Metal-Ligand Multiple Bonds, John Wiley & Sons, New York, 1988.
21 (a) R. H. Holm, Chem. Rev., 1987, 87, 1401; (b) J. M. Mayer, Comments Inorg. Chem., 1988, 8, 125; (c) T. A. Betley, Q. Wu, T. V. Voorhis and D. G. Nocera, Inorg. Chem., 2008, 47, 1849.
22 Y. Zhu, D. A. Smith, D. E. Herbert, S. Gatard and O. V. Ozerov, Chem. Commun., 2012, 48, 218.
23 J. Goodman, V. V. Grushin, R. B. Larichev, S. A. Macgregor, W. J. Marshall and D. C. Roe, J. Am. Chem. Soc., 2009, 131, 4326.
24 J. Choi, D. Y. Wang, S. Kundu, Y. Choliy, T. J. Emge, K. Krogh-Jespersen and A. S. Goldman, Science, 2011, 332, 1545.
25 (PNP)Rh(CO) accounted for approximately 1% of the reaction mixture.
26 D. Huang, P. R. Koren, K. Folting, E. R. Davidson and K. G. Caulton, J. Am. Chem. Soc., 2000, 122, 8916.
27 (a) R. P. Hughes, Eur. J. Inorg. Chem., 2009, 4591; (b) P. J. Brothers and W. R. Roper, Chem. Rev., 1988, 88, 1293.
28 We do not observe the HF by-product spectroscopically, but it is possible it is consumed by a reaction with borosilicate glass.
29 S. Gatard, C. Guo, B. M. Foxman and O. V. Ozerov, Organometallics, 2007, 26, 6066.
30 K. Reichenbächer, H. I. Süß and J. Hulliger, Chem. Soc. Rev., 2005, 34, 22.
31 S. C. F. Kui, N. Zhu and M. C. W. Chan, Angew. Chem., Int. Ed., 2003, 42, 1628.
32 M. C. Leclerc, S. I. Gorelsky, B. M. Gabidullin, I. Korobkov and R. T. Baker, Chem.–Eur. J., 2016, 22, 8063.
33 (a) E. O. Fischer, G. Kreis, C. G. Kreiter, J. Müller, G. Huttner and H. Lorenz, Angew. Chem., 1973, 85, 618; (b) E. O. Fischer,
35 L. P. Press, B. J. McCulloch, W. Gu, C.-H. Chen, B. M. Foxman and O. V. Ozerov, Chem. Commun., 2015, 51, 14034.

36 C. Douvris and O. V. Ozerov, Angew. Chem., Int. Ed. Engl., 2016, 120, 215.

37 S. P. Ho and F. Weinhold, Acc. Chem. Res., 2008, 41, 157.

38 W. R. Wadt and P. J. Hay, J. Chem. Phys., 1985, 82, 284.

39 W. R. Wadt and P. J. Hay, Discussing Chemistry with Natural Bond Orbitals, John Wiley & Sons, Hoboken, 2012.

40 E. D. Glendenning, C. R. Landis and F. Weinhold, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2012, 2, 1.

41 E. D. Glendenning, C. R. Landis and F. Weinhold, Valency and Bonding: A Natural Bond Orbital Donor-Acceptor Perspective, Cambridge University Press, Cambridge, 2005.

42 C. R. Landis and F. Weinhold, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2012, 2, 1.

43 F. Weinhold and C. R. Landis, Valency and Bonding: A Natural Bond Orbital Donor-Acceptor Perspective, Cambridge University Press, Cambridge, 2005.

44 F. Weinhold and C. R. Landis, Discovering Chemistry with Natural Bond Orbitals, John Wiley & Sons, Hoboken, 2012.

45 E. D. Glendenning, C. R. Landis and F. Weinhold, Valency and Bonding: A Natural Bond Orbital Donor-Acceptor Perspective, Cambridge University Press, Cambridge, 2005.

46 F. Weinhold and C. R. Landis, Valency and Bonding: A Natural Bond Orbital Donor-Acceptor Perspective, Cambridge University Press, Cambridge, 2005.

47 A. E. Reed, L. A. Curtiss and F. Weinhold, Chem. Rev., 1988, 88, 899.

48 K. B. Wiberg, Tetrahedron, 1968, 24, 1083.

49 Y. Zhao and D. G. Truhlar, Theor. Chem. Acc., 2008, 120, 215.

50 Y. Zhao and D. G. Truhlar, Acc. Chem. Res., 2008, 41, 157.

51 W. R. Wadt and P. J. Hay, J. Chem. Phys., 1985, 82, 284.

52 P. J. Hay and W. R. Wadt, J. Chem. Phys., 1985, 82, 299.

53 P. J. Hay and W. R. Wadt, J. Chem. Phys., 1985, 82, 270.

54 T. H. Dunning and P. J. Hay, Modern Theoretical Chemistry, in Applications of Electronic Structure Theory, ed. H. F. Schaefer III, Plenum, 1977, vol. 4, p. 461.

55 M. J. Frisch, J. A. Pople and J. S. Binkley, J. Chem. Phys., 1984, 180, 3265.

56 T. Clark, J. Chandrasekhar, G. W. Spitznagel and P. V. R. Schleyer, J. Comput. Chem., 1983, 4, 294.

57 A. D. McLean and G. S. Chandler, J. Chem. Phys., 1980, 72, 5639.

58 R. Krishnan, J. S. Binkley, R. Seeger and J. A. Pople, J. Chem. Phys., 1980, 72, 650.

59 A. D. Bochevarov, E. Harder, T. F. Hughes, J. R. Greenwood, D. A. Braden, D. M. Philipp, D. Rinaldo, M. D. Halls, J. Zhang and R. A. Friesner, Int. J. Quantum Chem., 2013, 113, 2110.

60 Jaguar, versions 7.0-9.3, Schrödinger, LLC, New York, 2007–2016.

61 D. Lentz, A. Bach, J. Buschmann, P. Luger and M. Messerschmidt, Chem. – Eur. J., 2004, 10, 5059.

62 C. R. Groom, I. J. Bruno, M. P. Lightfoot and S. C. Ward, Acta Crystallogr., Sect. B: Struct. Sci., Cryst. Eng. Mater., 2016, 72, 171.

63 R. H. Crabtree and M. Mingos, Comprehensive Organometallic Chemistry III, Elsevier, New York, 2006.

64 L. J. Guggenberger and R. Cramer, J. Am. Chem. Soc., 1972, 94, 3779.

65 We refer to a PNP ligand in which the PPr2 groups have been replaced by PMe2 groups and the aryl part of the pincer replaced by P=CH=CH=N linkers as “truncated.” It was used in calculations from which the NLMO Figures were derived and for calculations of rotational barriers in the C2H4/C2F4/CH2/CF2 complexes. All other calculations used the full version of the ligand. The truncated PNP ligands has been previously used by Wu and Hall (H. Wu and M. B. Hall, Dalton Trans., 2009, 5933).

66 (a) O. J. Curnow, R. P. Hughes and A. L. Rheingold, J. Am. Chem. Soc., 1992, 114, 3153; (b) O. J. Curnow, R. P. Hughes, E. N. Mairs and A. L. Rheingold, Organometallics, 1993, 12, 3102.