Calculation of a Methane C–H Oxidative Addition Trajectory: Comparison to Experiment and Methane Activation by High-Valent Complexes

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Abstract: An effective core potential (ECP), parallel supercomputing study of methane activation by 14-electron Ir(PH3)2(X) complexes (X = H, Cl) is presented. Considerable weakening of the coordinated methane C–H bond occurs upon formation of an α2-CH coordinated (X)(Ir=H)(2H) adduct. A more strongly bound adduct (with greater weakening of the coordinated C–H bond) occurs when X = Cl versus X = H. The calculated Ir(PH3)2(H) + CH4 = Ir(PH3)2(H)2(Me) reaction enthalpy is -12.8 kcal mol-1, and -41.6 kcal mol-1 for the chloro analogue. The intrinsic reaction coordinate is calculated and compared to an experimental trajectory. Analysis of the wave function along the intrinsic reaction coordinate (IRC) suggests that although donation of electron density from methane to metal is essential for adduct formation, it is not until backdonation to σ*IrH increases that the C–H bond is activated and cleaved. The electronic and molecular structure of the reacting system along the IRC suggest a two-stage mechanism: substrate to complex donation is important in the early part of the reaction (electrophilic stage) while complex to substrate backdonation is necessary later on (nucleophilic stage) for C–H scission. Finally, comparison of IRCs for low- and high-valent methane-activating complexes shows similar topology in the early portion of the activation event; differentiation between oxidative addition and σ-bond metathesis occurs at the point at which there is a shift from the electrophilic to nucleophilic stage of the reaction.

Introduction

Selective C–H activation of methane by transition metal and lanthanide complexes has been the subject of considerable study as a result of its importance in catalytic methanation. Two main strategies have been pursued to effect concerted (as opposed to nonconcerted radical or electrophilic activation) C–H activation in homogeneous systems: oxidative addition5 and σ-bond metathesis. An 1,2-addition of C–H across an appropriate metal–ligand bond (single or multiple) is effected by high-valent (d9 and d10) complexes. Representative examples of high-valent, methane-activated complexes are Cp2,2LiCH37 (NH5)2Zr==NSi2 (Si' = Si(t-Bu)3)8 and thoracyclobutanes.9 Oxidative addition entails 1,1-addition of C–H to a low-valent transition metal and results in an increase in the formal oxidation state of the metal by two units.9 The 16-electron Cp*M(L) (M = Rh, Ir; L = PR3, CO) intermediates5,10 are perhaps the best known examples of complexes for which oxidative addition is the putative C–H activation mechanism.

Another family of low-valent, C–H-activating complexes are 14-electron species of the general form M(L)(PR3)2, where L is a univalent ligand such as hydrido, chloro, or an n-carboxylate group.11,12 As Crabtree has pointed out,13 14-electron complexes have advantages over 16-electron species from the point of view of building a catalyst system. The hydrido(alkyl) complex, eq 1a, is still electron-deficient (i.e., 16-electron), typically with

\[ \text{L}_n\text{M} + \text{R} = \text{CH}_2 = \text{CH}_2 \rightarrow \text{L}_n\text{M}(\text{H})(\text{CH}_2 = \text{CH}_2 = \text{R}) \] (1a)

\[ \text{L}_n\text{M}(\text{H})(\text{CH}_2 = \text{CH}_2 = \text{R}) \rightarrow \text{L}_n\text{M}(\text{R}_2\text{C} == \text{C}(\text{H})\text{R}) \] (1b)

\[ \text{L}_n\text{M}(\text{H})(\text{CH}_2 = \text{CH}_2 = \text{R}) \rightarrow \text{L}_n\text{M}(\text{H}) + \text{H}_2\text{C} == \text{C}(\text{H})\text{R} \] (1c)

\[ \text{L}_n\text{M}(\text{H})_2 \rightarrow \text{L}_n\text{M} + \text{H}_2 \] (1d)

vacant coordination sites, allowing for β-H transfer to form an 18-electron bis(hydrido)olefin, eq 1b, which can dissociate olefin to yield a bis(hydrido) complex, eq 1c. The bis(hydrido) complex can regenerate the reactive intermediate either by photolytic dissociation of H2 or by addition of an H2 scavenger (e.g., tert-butylethylened), eq 1d.11 Crabtree11,12 has developed an alkane dehydrogenation catalyst starting from Ir(H2)(CO)2(η2-carboxylate)(PR3): the reactive intermediate is thought to be the 14-electron Ir(PR3)2(η2-carboxylate). Sakakura and Tanaka13 have studied catalytic alkane carbylation by a Ru catalyst; the C–H activating species is thought to be Rh(C)(PR3)2. Recently, Schulz and Werner12a have studied the rearrangement between vinyl(hydrido) complexes and olefin isomers of Ir(PR3)2(η2-CI)(C,H2).


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(c) A good discussion of the requirements for a successful methane conversion catalyst from an industrial point of view can be found in: Parkyns, N. D. Chem. Br. 1990, 9, 841.

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As part of a continuing focus on transition metal bonding and reactivity, a study of the important C–H activation reaction, eq 2, was undertaken. For the present, the focus is on C–H activation by 14-electron complexes of the form Ir(PH₃)ₓ(X)ₒ₉₈₉₆, where X = H and Cl. In previous work, the concentration of high-valent C–H-activating systems operating through π-electron meta-stability. In order to come to a more global understanding of the C–H activation problem, we instituted research on oxidative addition by 14-electron systems. This work also constitutes further testing of the coupling of promising new technologies (like parallel computing) and methods (such as effective core potentials) as answers to the challenges of computational transition metal (TM) and lanthanide (Ln) chemistry.

Computational Methods

Calculations employ the quantum chemistry program GAMESS. One important difference between this and previous work is that the GAMESS code has now been made to run in parallel on a variety of platforms. This research is part of a continuing focus on oxidative addition. In the standard implementation, TM valence basis sets and core orbitals for main-group (MG) elements. Thus, the ns, np, nd, (n + 1)s, and (n + 1)p are treated explicitly for the d-block; for the main-group, ns and np are treated explicitly. In the standard implementation, TM valence basis sets are quadruple- and triple-f for the sp and d shells, respectively, while main-group elements have a double-f valence basis. Basis sets for heavy, main-group elements are augmented with a d polarization function in our work. Transition metal ECPS are generated from all-electron, Dirac–Fock calculations and thus include Darwin and mass velocity effects, while spin–orbit coupling is averaged out in potential generation. No degradation in accuracy has been found upon descending a transition metal triad toward heavier members (for which relativistic effects will be most important).

Geometries are optimized at the restricted Hartree Fock (RHF) level for closed-shell singlets. Previous work has shown that bond lengths and angles for TM complexes (involving complexes in a variety of geometries and oxidation states and of metals from the entire transition series) are predicted to within 1–3% of experiment using the present computational scheme. Vibrational frequencies are calculated at stationary points to identify them as minima (zero imaginary frequencies) or transition states (one imaginary frequency). Plotting the imaginary frequencies is used to assess which TS connects which reactants and products. In some cases the intrinsic reaction coordinate (IRC) is followed using the steepest descent algorithm of Ishida et al., with the stabilization method described by Schmidt et al.

Although geometries are accurately predicted at the RHF level, energetics will typically be poor if correlation is ignored. For species described well at the RHF level, the correlation contribution can be treated as a perturbation of the RHF energy and calculated using Moller–Plesset second-order perturbation theory (MP2). Koga and Morokuma used a similar scheme in their work, as have Krogh-Jespersen et al. in studies of oxidative addition. A simple RHF geometry/MP2 energy scheme yielded good agreement with experimental data for high-valent C–H-activating systems, both in terms of absolute numbers and trends.

Results and Discussion

1. Reactants. The geometry of methane is well-known. Reactant complexes are three-coordinate Ir(X)(PH₃)₂₂ complexes (X = H, Cl), models of the putative 14-electron, C–H-activating species in dehydrogenation catalysts. As in previous computational work, we have focused on the lowest energy, singlet surface. Geometries for bis(phosphino)Ir chloride and bis(phosphino)Ir hydride minima (C₂ symmetry) are shown in Figure 1. As expected for a d⁸M₉ complex, the geometry is T-shaped.
to a vinyl; hydrocarbons are known to have large trans influences.²⁹
Ir–Cl bond lengths in Ir(H)(Cl)(PF₃)₂(C₂H₅)(C₆H₅) are 2.32(6) Å for Ir–H bonds with a range 1.57 (7)–2.82 (2) Å. The sum of the single-bond metallic radius of Ir (1.27 Å) and the covalent radius of H (0.77 Å) is 2.04 Å. Three additional X–H bonds have been characterized by neutron diffraction: Ir–H = 1.603 (9) Å (average Ir–H in Ir(H)(Cl)(PF₃)₂), 1.59 Å ([H₃Ir(P(Et₂)₃)] +), and 1.594 (3) Å ([H₃Ir(Si(Et₂)₂)Cl] +). All in good agreement with the calculated value of Ir–H in 1a. Thus, calculated Ir–X bonds in model reactants (1) are in accord with experimental data. Good agreement between theory and experiment has been found in previous work using ECPs and valence basis sets; it is satisfying to find this trend continued for a new family of low-valent TM complexes.

2. Initial Interaction of Substrate and Reactant Complex.

Substantial interest has focused on loosely bound, alkane adducts of TM complexes and their relevance to alkane activation. For silanes, Si–H adducts are stable enough to permit structural analysis. Parkin and Bercaw have measured inverse kinetic and deuterated isotopomers, as have Gould and Heineke for isotope effects for methane elimination from CpIr(H)(CH₃)(H)₂Rh(CO). Burkey has measured ΔH° values as high as 17 kcal mol⁻¹ (Mo(CO),)₂(CO)₆). All three iridium hydride complexes have been characterized by neutron diffraction: Ir–H = 1.603 (9) Å (average Ir–H in Ir(H)(Cl)(PF₃)₂), 1.59 Å ([H₃Ir(P(Et₂)₃)] +), and 1.594 (3) Å ([H₃Ir(Si(Et₂)₂)Cl] +). Both findings have been interpreted in terms of adduct formation. Bergman et al. have used time-resolved infrared spectroscopy (TRIS) to probe alkane adducts of CpRh(CO). Rayner et al. have performed gas-phase TRIS studies of alkane interaction with W(CO),. Burke has used photoacoustic calorimetry to measure enthalpies of adduct formation between hydrocarbons and coordinatively unsaturated species such as M(CO), (M = Cr, Mo, W). Thus, there is a growing body of kinetic, spectroscopic, and thermochemical data supporting the presence of alkane adducts of TM complexes. The structure of these transients and their function, if any, in the crucial bond scission step are less clear.

On the basis of TRIS studies, an adduct-binding enthalpy (ΔH°) of 10 kcal mol⁻¹ has been estimated for alkane adducts of CpRh(CO). Burke has measured ΔH° values as high as 17 kcal mol⁻¹ (Mo(CO), + heptane) in solution. Gas-phase ΔH° values for Cr(CO), with alkanes and fluorinated alkanes range from <5 (methane) to 12.2 kcal mol⁻¹ (EtF). Ziegler et al. have measured a linear geometry for Cp(L)M–HCH₃ (M = Rh, Ir; L = Ph₃, CO); binding energies are ~13 kcal mol⁻¹ for CpIr(L) and 6–7 kcal mol⁻¹ for CpRh(L). Blomberg et al. have carried out calculations on a new family of low-valent TM complexes.

Level. Thus, as Koga and Morokuma have clearly shown, different adduct geometries can be very close in energy. Reducing optimization convergence criteria by an order of magnitude yields no significant change in geometry (<0.1% changes in bond lengths and bond angles) or energy (<0.1 kcal mol⁻¹). It seems likely that at ambient temperatures a methane adduct will be fluxional, sampling different coordination modes. Simple replacement of the methane by another alkane shows that calculations on the RHF level are not adequate to support the presence of alkane adducts of TM complexes. The growing body of kinetic, spectroscopic, and thermochemical data...
H with Cl yields an \( \eta^2 \)-CH minimum (C\(_2\) symmetry, 3b) for which Ir--C and Ir--\( \cdot \)H distances are much shorter. The differences in calculated geometries are reflected in a greater \( \Delta H_{\text{add}} \) for Ir(PH\(_3\))\(_2\)(Cl) versus Ir(PH\(_3\))\(_2\)(H). Calculated \( \Delta H_{\text{add}} \) values for 3a and 3b are \(-6.8 \) and \(-15.6 \) kcal mol\(^{-1}\), respectively. \(^{46}\) Calculated values for \( \Delta H_{\text{add}} \) are in line with the experimental and computational estimates given above; more quantitative assessments will require higher levels of theory.

Calculated intrinsic stretching frequencies\(^{47}\) show coordinated C--H bonds to be much weaker than spectator C--H bonds: 2873 cm\(^{-1}\) versus 3014 cm\(^{-1}\) (3a) and 2680 cm\(^{-1}\) versus 3032 cm\(^{-1}\) (3b). Note that the coordinated C--H is much weaker in 3b than in 3a, consistent with bond lengths of the coordinated C--H: 1.10 Å in 3a; 1.13 Å in 3b. The charge on the methane fragment (from Mulliken population analyses) in 3a is \(+0.14 \) versus \(+0.25 \) for 3b. Within the Saillard-Hoffmann\(^{13}\) model two explanations seem feasible for the difference in charges. Replacement of X with H with X = Cl makes 1b a better acceptor for \( \sigma^* \text{CH} \) than 1a; 1b is less effective at \( \pi \)-backbonding to \( \sigma^* \text{CH} \) than 1a. The Mulliken charge on Ir is \(-1.10 \) in 1a and \(-0.69 \) for 1b. A less negative charge on 1b should lower in energy metal-based MOs (as the calculations indicate); however, \( \pi \)-donation from the chloro destabilizes the \( \sigma^* \) orbital perpendicular to the Ir-P\(_2\)X plane (the one backbonding to \( \sigma^* \text{CH} \)), making it roughly the same energy in 1a (\(-0.314 \) hartree) and 1b (\(-0.324 \) hartree).

The calculations thus point to 1a and 1b being equivalent \( \sigma \)-donors with the latter a better \( \sigma \)-acceptor, leading to a more exothermic \( \Delta H_{\text{add}} \) and increased C--H weakening upon adduct formation for 3b. The weakening of coordinated C--H which occurs upon adduct formation may assist in C--H scission, as suggested for intermolecular\(^{48}\) and intramolecular\(^{39,49}\) M--H--C complexes. The calculations show the adduct to be sensitive to electronic effects induced by ancillary ligands, in addition to steric factors.\(^{15,17,38}\) Tunable properties are attractive from the standpoint of designing a catalyst precursor in which C--H bond weakening is maximized in an adduct, particularly if the interactions carry through to the transition state and lower the activation barrier.

3. Products. Products are 16-electron, five-coordinate Ir\(^{11}\) complexes--Ir(X)(H)(CH\(_3\))(PH\(_3\))\(_2\) (4). The phosphines remain trans to each other with angles close to 180°. The geometry of the complexes is closer to a square pyramid (SQP5) than a trigonal bipyramid (TBP5). Frontier orbital arguments suggest that a \( d^6 \) TBP5 complex will be Jahn-Teller unstable and distort to an SQP5 geometry;\(^{25}\) furthermore, a Walsh diagram shows the preferred \( L_{\text{spatial}} - M - L_{\text{axial}} \) angle to be near 90°.\(^{25}\) The optimized geometries of the products (4) bear out these predictions in all respects.

The Ir--X bonds X = H, Cl, P are in good agreement with experimental models described above. The most interesting changes in bond lengths highlight the trans influence of hydrido and methyl ligands. The Ir--Cl bond in 4b, trans to methyl, lengths by 0.12 Å (versus 1b); the Ir--H bond in 4a, trans to a methyl, lengths by 0.10 Å, still within reported limits.\(^{33}\) Rausch et al.\(^{49}\) quote Ir--C distances ranging from 1.982(26) to 2.202(4) Å depending on metal coordination environment.

The geometry of the products (4) bear out these predictions in all respects. The calculated IR spectra of 4a and 4b are found in Table 4. Differences in bond lengths reflect the trans influence of hydrido and methyl groups.

4. Transition State. It is generally agreed\(^{30,31,14,54,49}\) that low-valent complexes activate C--H bonds through a triangular M--H--C oxidative addition transition state (TS), 5a. The distance in the complex with a trans hydrido (Ir--C = 2.18 Å, 4a) is much longer than that with a trans chloro (Ir--C = 2.10 Å, 4b), in line with the trans influence order H > CH > Cl.\(^{29}\) Apical H atoms in 4a and 4b are trans to a vacant coordination site and show Ir--H bond lengths (1.55 Å) nearly identical to that of the Ir(PH\(_3\))\(_2\)(H) reactant (1.58 Å).

The Ir(PH\(_3\))\(_2\)(H) \( + \) CH\(_4 \) \( \rightarrow \) Ir(PH\(_3\))\(_2\)(H)(CH\(_3\)) reaction enthalpy is \(-12.8 \) kcal mol\(^{-1}\) versus separated reactants, while that for Ir(PH\(_3\))\(_2\)(Cl) is \(-41.6 \) kcal mol\(^{-1}\). Theoretical estimates for methane C--H oxidative addition are \(-15, -17, -33, -36 \) kcal mol\(^{-1}\) for CpRh(CO), CpRh(PH\(_3\)), Cplr(CO), and Cplr(PH\(_3\)), respectively.\(^{15}\) Koga and Morokuma\(^{15}\) calculate reaction energies for (C)(PH\(_3\))\(_2\) + CH\(_4\) \( \rightarrow \) Rh(C)(PH\(_3\))(CH\(_3\))(PH\(_3\)) from -17.3 to -24.4 kcal mol\(^{-1}\). Experimental Ir--H and Ir--R bond strengths for Cplr(R)\(_2\) are 74 and 56 kcal mol\(^{-1}\) (R = Me).\(^{50}\) Combining these values with a methane C--H bond strength of 105 kcal mol\(^{-1}\) yields an estimate of \(-25 \) kcal mol\(^{-1}\) for CpIr(PMe)(H) + CH\(_4\) \( \rightarrow \) CpIr(PMe)(H)(CH\(_3\)).\(^{51}\) Ir--C bond enthalpies in Ir(C\(_3\))\(_2\)(PR\(_3\))(CO)(C(O)Me) span a range of 12 kcal mol\(^{-1}\) for a variety of PR\(_3\) ligands;\(^{52}\) Ir--C bond energies in Ir(Cl\(_2\))\(_2\)(CO)(PPPh\(_3\))(R) are spread over a 17 kcal mol\(^{-1}\) range depending on the nature of R.\(^{53}\) Bond strengths of Ir--C are thus very sensitive to the chemical environment, a contention further supported by the large range of Ir--C bond lengths quoted by Rausch et al.\(^{49}\) The stronger trans influence of hydrido versus chloro leads one to predict that the Ir--C bond will be weaker in 4a than in 4b; thus, reaction 2 will be more exothermic for X = Cl than for X = H, as found. The calculations suggest that the relative trans influence of hydrido versus chloride is great enough to lead to a large difference in reaction exothermicities. Detailed studies of transition metal-carbon bond enthalpies as a function of trans ligand would be of great value in this regard.

5. Stationary Points (6) (with the geometry shown in 5a are found. That 6a and 6b are transition states at this level of theory is confirmed by calculation of the

46 Modification of ancillary ligands can substantially affect the acidity/ basicity of a metal center. Replacement of hydride with chloride makes the Os center significantly more acidic/basic in OsP\(_3\)(PPh\(_3\))\(_2\)X. The enthalpy of protonation by CF\(_3\)SO\(_3\)H is 23.2 kcal mol\(^{-1}\) less exothermic for OsP\(_3\)(PPh\(_3\))\(_2\)Cl than for OsP\(_3\)(PPh\(_3\))\(_2\)H. Rotkin, M. K.; Angelici, R. J., J. Am. Chem. Soc. 1992, 114, 8206.

47 (a) Boatz, J. A.; Gordon, M. S. J. Phys. Chem. 1989, 93, 1819. (b) Calculated stretching frequencies are typically 10% too high and thus scaled by 0.9 to account for correlation and anharmonicity. Pope, J. A.; Schüegg, H. B.; Krishnan, R.; DeFrees, D. F.; Binkley, J. S.; Frisch, M. J.; Whiteside, R.; Hout, R. F.; Hehre, W. J. Int. J. Quantum Chem., Proc. Sanibel Symp. 1981, 15, 269. We have used a scaling factor of 0.925, since this best reproduces the experimental C--H stretches of methane.


Contribution

A single-determinant description of the TS is not sufficient. Calculation at the RHF transition state shows a very large increase in internal coordinate displacements shows the major component would expect for a true oxidative addition TS (Figure 2). Probing the complete analogue is identical. Decomposition of the imaginary mode into internal coordinate displacements shows the major component would be of interest, although computationally expensive. Experimental trajectory "active site* am

The most interesting point to emerge from calculation of TS geometries is the extent to which Ir-H and Ir-C bond lengths are formed in the transition state. Relative to their values in the product, the Ir-C and Ir-H bond lengths in TS 6a (2.23 and 1.60 Å, respectively) are only 2% and 3% longer than those in product 4a; for the chloro derivative, Ir-C and Ir-H bond lengths in TS 4b (2.21 and 1.60 Å, respectively) are 6% and 3%, respectively, longer than those in product 4b. The C-H bond being activated is further removed from equilibrium values than Ir-H and Ir-C: +55% (6a) and +31% (6b). Crabtree11 has deduced a late TS for oxidative addition in 14-electron complexes on the basis of selectivity patterns; kinetic products arise from activation at the least hindered C-H bonds in the substrate, suggesting a short M-C distance in the TS and steric hindrance between substituents on C and ancillary ligands. The calculations support a "late" transition state for oxidative addition; not only is the Ir-C interaction substantial but Ir-H is as well. The tightness of transition states 6a and 6b should make it profitable to control the selectivity of alkane functionalization catalysts built around these 14-electron X(M)(PR3)2 species, particularly through choice of R groups on the phosphines.36

5. Intrinsic Reaction Coordinate. The intrinsic reaction coordinate (IRC)21 yields dynamic information about chemical interactions which dictate the ease of passage of reactants through the TS and on to products. Crabtree et al.37 have analyzed the crystal structures of agostic complexes and thus constructed a trajectory for C-H oxidative addition. This experimental trajectory is then used to follow the evolution of the C-H distance and M-H--H-C angle for the C-H being oxidatively added. Since the IRC is mathematically defined as the steepest descent path (in mass-weighted Cartesian coordinates) from the TS to reactants and products, it can also be calculated from first-principles quantum mechanics,21 making it of interest to compare the well-known experimental trajectory27 with a computational trajectory.

Calculated changes in bond lengths and angles along the IRC for IrCl(PR3)2(H)Ir-CH2CH3 (3a → 6a → 4a) are shown in Figure 3a using the reaction coordinate (r_eq, eqs 3 and 4) defined by Crabtree et al.27 (r = 0.28 Å; 1.27 Åbp)

\[ d_{bp} = \left[ d_{MH}^2 + r^2 d_{CH}^2 - r\left(d_{MH}^2 + d_{CH}^2 - d_{MC}^2\right)\right]^{1/2} \]

where \( r_{eq} \) is the covalent radius of Ir. \( r_{eq} \) is effectively the covalent radius of the C-H bonding electrons.21

In Figure 3b the same bond length and bond angles are plotted using \( S_{total} \) (eq 5), the \( S_{total} = \sum [(x_i(0) - x_i(s))^2 + (y_i(0) - y_i(s))^2 + (z_i(0) - z_i(s))^2]^{1/2} \)

mass along the reaction coordinate, on the abscissa. The values \( x_i(0), y_i(0), z_i(0) \) and \( x_i(s), y_i(s), z_i(s) \) in eq 5 are the mass-weighted Cartesian coordinates of atom \( i \) at the TS, while \( x_i(s), y_i(s), z_i(s) \) are the mass-weighted Cartesian coordinates of atom \( i \) at a subsequent point along the IRC. Proceeding from positive to negative values of \( S_{total} \) (Figure 3b) describes the oxidative addition process, the TS of which is at \( S_{total} = 0 \) (eq 6). Inspection of Figure 3b shows that using either \( r_{eq} \) or \( S_{total} \) as the reaction coordinate yields nearly identical results for describing the C-H oxidative addition trajectory. Thus, \( S_{total} \) will be used as the reaction coordinate in subsequent discussion.

The experimental trajectory27 derived by Crabtree and coworkers is reproduced in Figure 4. It seems reasonable to ascribe

(55) The two MOs located on the Ir=H=C "active site" are included in the M-CSCF active space as are their antibonding counterparts. This yields a four-orbital/four-electron active space. The FORS (full optimized reaction space)-M-CSCF wave function is generated by constructing all possible single


(56) Product selectivity for C-H activation by 16-electron CpML intermediates is kinetically controlled at low temperatures (where the reverse reaction, reductive elimination of hydrocarbons, is minimal) and thermodynamically controlled as one approaches room temperature. Jones, W. D. (Chemistry, Rochester). Personal communication.
that similar behavior for the C-H distance and M-H-C angle intermolecular Ma-H-C interaction). However, the fact remains tional geometry arises from a single set of reactants (with this stage in the reaction. Steric hindrance is expected to reaction coordinate not sampled by crystal structures, particularly in the region of the transition state.

Figure 3. (a, top) Calculated bond length and bond angle changes along the IRC for conversion of the Ir(PH3)2(H)(qZ-HCH3) adduct into Ir(PH3)2(H)2(CH3) using $\tau_{\text{dim}}$ as the reaction coordinate. See text for definition of $\tau_{\text{dim}}$. (b, bottom) Calculated bond length and bond angle changes along the IRC for conversion of the Ir(PH3)2(H)(qZ-HCH3) adduct into Ir(PH3)2(H)2(CH3) using $S_{\text{total}}$ as the reaction coordinate. See text for definition of $S_{\text{total}}$.

differences with computed ones to the fact that the experimental trajectory is derived from consideration of a variety of complexes (with intramolecular M--H--C interaction) while the computational geometry arises from a single set of reactants (with intermolecular M--H--C interaction). However, the fact remains that similar behavior for the C--H distance and M--H--C angle is exhibited in both experimental (Figure 3b) and computational trajectories (Figure 3), which is pleasing given the important contribution the Crabtree trajectory has made to the thinking on C--H activation. The computational trajectory does have an advantage in that it allows one to see important parts of the reaction coordinate not sampled by crystal structures, particularly in the region of the transition state.

Analysis of the changes in bond lengths and bond angles along the intrinsic reaction coordinate (Figure 3b) reveals three main points of interest.

1. The Ir--H--C angle is nearly constant at 105$^\circ$ for $S_{\text{total}} \geq 1$ bohr amu$^{-1/2}$. The value obtained from the experimental reaction coordinate is $\approx 130^\circ$. The experimental trajectory(57) was constructed from intramolecular, agostic complexes whose ability to sample the entire M--H--C conformational space is more restricted than in an intermolecular case. The near degeneracy of the edge-coordinated (2) and y =$\pi$-CH (3a) adducts for Ir(PH3)2(H)(CH3) suggests that distortion along the Ir--H--C mode is very soft at this stage in the reaction. Steric hindrance is expected to be less in a linear adduct than for a y =$\pi$-CH coordination geometry. However, an off-axis approach in the experimental and computational trajectories is consistent with a favorable orientation for donation from $\sigma_{\text{CH}}$ to a vacant metal orbital concomitant with backdonation from a filled metal orbital into $\sigma^*_{\text{CH}}$; a linear (Ir--H--C = 180$^\circ$) approach does not allow for any backdonation.

2. The Ir--H and Ir--C distances are nearly at the values they assume in the oxidative addition product even before the C--H bond has begun to be activated (at $S_{\text{total}} = 1$ bohr amu$^{-1/2}$, Figure 3b). Thus, the main motion, and energetic expenditure, in the region of the transition state ($S_{\text{total}} = 1$ bohr amu$^{-1/2}$ to $S_{\text{total}} = 1$ bohr amu$^{-1/2}$, Figure 3b) corresponds to cleavage of the C--H bond, as decomposition\(^{(47)}\) of the imaginary frequency (Figure 2) indicates.\(^{(57)}\) As mentioned above, a TS for C--H activation with substantial M--C bond formation has been inferred from selectivity patterns.\(^{(11)}\) Analysis of the IRC indicates that M--C (and M--H) bond formation is substantial even before the TS.

3. The sharp increase in the C--H bond length is correlated with a sharp decrease in the Ir--H--C angle at $S_{\text{total}} = 1$ bohr amu$^{-1/2}$ (Figure 3b). Calculation of Mulliken atomic charges shows that the "break point" at $S_{\text{total}} = 1$ bohr amu$^{-1/2}$ is roughly coincident with a decrease in the charge on the methane fragment. The calculated charge on the methane fragment is +0.14 in the adduct (3a) and increases to a value of +0.45 in the TS region, after which it decreases toward the value of +0.22 that is obtained in the product (4a). Gordon and Gano(58) have studied the IRC for a main-group analogue of this reaction (H2E + CH4 $\rightarrow$ H2E--CH3) and interpreted it in terms of a two-stage mechanism: an electrophilic, early stage (dominated by donation from $\sigma_{\text{CH}}$ to a vacant E(1)-based MO) followed by a nucleophilic, later stage (dominated by backdonation from an occupied EH2 orbital to $\sigma^*_{\text{CH}}$). In light of the Saillard--Hoffmann\(^{(13)}\) model of M--C--H bonding, the positive charge on the methane fragment at all stages between methane adduct and methyl(hydride) product indicates that forward donation from a C--H bonding MO to a vacant metal-based MO is greater than backdonation from an occupied

(57) Kubas et al. have concluded from analysis of neutron diffraction structures of $\sigma^*_E$-H2 complexes that the reaction coordinate for H--H bond breaking shows relatively little change in H--H distance until bond rupture is quite imminent, presumably when increased backbonding can no longer be tolerated.\(^{(23)}\) These deductions are consistent with this analysis of the IRC for the related C--H oxidative addition reaction. Kubas, G. J.; Burns, C. J.; Eckert, J.; Johnson, S. W.; Larson, A. C.; Vergamini, P. J.; Unkefer, C. J.; Khalsa, G. R. K.; Jackson, S. A.; Eisenstein, O. J. Am. Chem. Soc. 1993, 115, 569.
metal-based MO to a C–H antibonding MO all along the oxidative addition IRC. The increasing positive charge on the methane fragment as the adduct moves toward the TS indicates that donation is increasing relative to backdonation; the decreasing charge after the "break point" at $S_{\text{total}} = 1 \text{ bohr amu}^{-1/2}$ suggests that the reverse situation applies in the later stages of oxidative addition. Combining these threads leads to the conclusion that although donation of electron density from methane to metal is essential for formation of an adduct, it is not until significant population of $\sigma^*_{\text{CH}}$ occurs that C–H is cleaved.\(^{(19)}\)

One intriguing result which emerged from the experimental reaction coordinate\(^{(27)}\) is that both low- and high-valent, agostic complexes fit on the same trajectory, although occupying different positions along the reaction coordinate. To more closely probe the similarities between a typical $\sigma$-bond metathesis IRC and the present oxidative addition one, we have plotted both on the same reaction coordinate (Figure 5). The $\sigma$-bond metathesis IRC is that for the reaction $W(=\text{NH})_2 + CH_4 \rightarrow W(=\text{NH})_2(\text{CH}_3)$ (open points). The tungsten IRC shows the same features discussed at length in previous computational studies of methane activation by imido complexes.\(^{(16a-c,60)}\) A W example was chosen, since the metallic single-bond radii of W (1.30 Å) and Ir (1.27 Å) are roughly the same.\(^{(23)}\)

The resemblance of the oxidative addition IRC (Figure 3a) to the $\sigma$-bond metathesis IRC (Figure 5) is striking and suggests a good degree of similarity between C–H activation mechanisms, particularly early on in the reaction coordinate in the vicinity of the methane adduct ($S_{\text{total}} = 1 \text{ bohr amu}^{-1/2},$ Figure 5). In a study of methane adducts of high-valent complexes,\(^{(40)}\) $\pi^2$-CH coordination was also found to be preferred. For both $\sigma$-bond metathesis and oxidative addition, M–C bond lengths in the transition state ($S_{\text{total}} = 0 \text{ bohr amu}^{-1/2}$) are close to the values they assume in the product (Figure 5). Significant M–C bond formation has been inferred in the TS for C–H oxidative addition by 14-electron complexes\(^{(11)}\) from selectivity patterns. Extensive M–C interaction in the $\sigma$-bond metathesis TS (Figure 3a) should also allow one to engineer an alkane functionalization catalyst built on a high-valent imido active species which will permit control of selectivity through judicious choice of ancillary ligands. The M–H distances show distinctly different behavior along the IRC for the two mechanisms; this is not surprising, since in oxidative addition the H ends up coordinated to the metal and in $\sigma$-bond metathesis it is coordinated to a ligated atom. For $\sigma$-bond metathesis the importance of the M–H interaction in moderating the energetics of the transition state has been discussed.\(^{(18a-c)}\) However, in both cases the M–H distance in the TS is only slightly longer than a typical metal–terminal hydride bond length. The most interesting of the similarities between oxidative addition and $\sigma$-bond metathesis IRCs (Figure 5) is the same correlation between the abrupt increase in C–H distances and the sharp decrease in M–H–C angle. The same pattern of increasing charge on the methane fragment, followed by a decrease in the methane charge near this "break point" ($S_{\text{total}} = 1 \text{ bohr amu}^{-1/2}$) is seen for the high-valent system; however, it is not as dramatic as in oxidative addition, as expected since backdonation from the metal will be minimal for an electrophilic, $d^0$ complex. The frontier orbitals of this imido (and others we have studied)\(^{(18)}\) are metal–nitrone $\pi$ bonds, making them a potential source for electron density. Thus, while the metal acts as an electrophile in $\sigma$-bond metathesis and oxidative addition (which the calculations indicate is essential for adduct formation), the difference lies in the moity which acts as nucleophile (donating to $\sigma^*_{\text{CH}}$ which is crucial to cleavage of the C–H bond). The nucleophiles in oxidative addition and $\sigma$-bond metathesis are, of course, the metal and a polarized metal–ligand bond which directs the H in the C–H bond being activated to its eventual position. A two-stage C–H activation process provides an explanation for the similarity seen in the early part of the activation mechanism, since it is here that electron donation from substrate to metal is crucial for adduct formation for both types of methane-activating complexes. More work is needed to probe these conclusions for a wider variety of C–H-activating complexes; such research is currently underway in our lab.

Summary

An effective core potential, parallel supercomputing study of methane activation by oxidative addition was carried out for the 14-electron complexes Ir(Ph$_3$)$_2$(X), X = H, Cl. Several conclusions have been reached which are summarized below.

(1) Effective core potentials have again shown their ability to describe the bonding and structure of a transition metal system.\(^{(18)}\) The coupling of this new methodology with emerging technologies like parallel computing will continue to expand the ability to probe interesting chemistry in all regions of the Periodic Table. Given the importance of the transition metals (and lanthanides) in catalysis and advanced materials, such efforts are clearly warranted.

(2) Initial interaction of methane with the activating complexes is in line with previous computation and experiment. The chloro adduct is found to be more strongly bound (by $\sim 9 \text{ kcal mol}^{-1}$) due to electronegative Cl making the complex a better acceptor for $\sigma_{\text{CH}}$ density.

(3) Metal–carbon bond formation is substantial in the TS, as suggested by experiment.\(^{(11)}\) Additionally, Ir–H bonding in the transition state is also substantial. Thus, the main energetic expenditure in the TS region comes from scission of the C–H bond.

(4) Analysis of the intrinsic reaction coordinate shows good agreement with the experimental trajectory of Crabtree et al.\(^{(27)}\) The most interesting observation is the correlation between an abrupt increase in C–H bond distance and a decrease in M–H–C angle. Analysis of the wave functions shows that the methane fragment which had been growing more positively charged up to this point starts to accept electron density and become less positively charged. In view of the Saillard–Hoffmann\(^{(13)}\) model

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(59) (a) It is worth noting that all partitioning of the total electron density is arbitrary, although one expects trends in properties to be more reliable than absolute numbers. (b) Stretching the C–H bond and closing the M–H–C angle should greatly increase the effectiveness of metal to $\sigma^*_{\text{CH}}$ backdonation. See ref 13, in particular Table II.

(60) Full details of the reaction $W(=\text{NH})_2 + CH_4 \rightarrow W(=\text{NH})_2(\text{CH}_3)$ are to be published in a future contribution. C–H activation by high-valent, group IVB imido complexes is discussed in ref 18a.\(^{(61)}\)

of M/C–H bonding, we interpreted this in terms of a two-stage C–H activation mechanism: substrate to complex donation is important in the early stages of the reaction (electrophilic stage) while complex to substrate backbonding is necessary later in the reaction (nucleophilic stage) for C–H bond breaking.

Comparison of the intrinsic reaction coordinate for a low- and high-valent methane activation reaction showed surprisingly similar topology in the early portion of the C–H activation process. Differentiation between the two mechanisms occurs in the region in the TS at the point at which there is a shift from the electrophilic to nucleophilic stage.

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