Anagostic Interactions under Pressure: Attractive or Repulsive?

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The term “anagostic interactions” was coined in 1990 by Lippard and coworkers to distinguish sterically enforced M•••H-C contacts (M = Pd, Pt) in square-planar transition metal d8 complexes from attractive, agostic interactions.[1a] This classification raised the fundamental question whether axial M•••H-C interaction in planar d8-ML4 complexes represent (i) repulsive anagostic 3c-4e M•••H-C interactions[1] (Scheme 1a) or (ii) attractive 3c-4e M•••H-C hydrogen bonds[2] (Scheme 1b) in which the transition metal plays the role of a hydrogen-bond acceptor (Scheme 1b). The latter bonding description is related to another bonding concept which describes these M•••H-C contacts in terms of (iii) preagostic or preagostic interactions[3] (Scheme 1c) which are considered as being “on the way to becoming agostic, or agostic of the weak type”.[4]

preagostic interactions are considered to lack any “involvement of dz2 orbitals in M•••H-C interactions” and rely mainly on M(dxz,dyz) → σ*(H-C) π-back donation.[1b]

The first observation of unusual axial M•••H-C interaction in planar d8-ML4 complexes was made by S. Trofimenko, who pioneered the chemistry of transition metal pyrazolylborato complexes.[5,6] Trofimenko also realized in 1968, on the basis of NMR studies, that the shift of the pseudo axial methylene protons in the agostic species [Mo{Et2B(pz)2}((η2-allyl)(CO)3}] (1) (pz = pyrazolyl; allyl = H2CCHCH3) “is comparable in magnitude but different in direction from that observed in Ni{Et2B(pz)2}2” (2) (Scheme 2).[8]

Indeed, the protons of the methylene group which form a close M•••C contact of [2.954(2) Å][8] in 1 resonate at -2.41 ppm at RT,[9] while the corresponding signal of the methylene protons of 2 occurs at 3.64 ppm (q, CH2, 2JH-H 8 Hz) and does not show any coalescence upon cooling to -90 °C despite large differences in the calculated chemical shifts[10a] of both methylene protons (Scheme 2). In contrast, the 1H NMR signal of the agostic Mo•••H-C in 1 splits into two features (-0.83 and -4.3) below -53 °C[9] in agreement with the computed NMR properties of our static DFT model (1.4 and -5.88 ppm) for the agostic Mo•••H-C and non-coordinating methylene proton, respectively (Scheme 2). Trofimenko concluded that the agostic proton in 1 displays a “hydridic” character and that the agostic proton of 1 is “intruding into a suitable empty metal orbital.”[8a] He, therefore, suggested that the M•••H-C interactions in 1 is a bonding one causing an activation of the C-H bonds as evident by the “presence of CH stretch bands at remarkably low frequency” of 2704 cm-1[10b]. This concept was later developed and refined by Brookhart and Green (BG); who coined the expression “agostic” for these kind of interactions.[11]

This work was supported by the DFG (SPP1178) project number SCHE478/12-1 and the SNSF project Nr. 144534.

Supporting information for this article is available on the WWW under http://www.dc.doi.org/10.1002/anie.2014XXXX

Scheme 1.

In contrast to the first two types of interactions which require the presence of a fully occupied and axially oriented M(dz2) orbital, 

Scheme 2.

Supporting information for this article is available on the WWW under http://www.dc.doi.org/10.1002/anie.2014XXXX

DOI: 10.1002/anie.200((will be filled in by the editorial staff))
Echols and Dennis in 1974 who concluded that “steric restrictions dictate the conformation of the molecule, not an interaction of Ni with an apical H atom” of one of the methylene groups of the pyrazoylborato ligand.\textsuperscript{(18,12)} A subsequent Extended Hückel Theory (EHT) study by Saillard and Hoffmann (SH)\textsuperscript{(14c)} gave further evidence that the interaction between an axially approaching H-C ligand (e.g. CH₄) and the d⁸-ML₄ metal fragment might be actually even repulsive due to the presence of a fully occupied and axially oriented (M)dl²-orbital (Scheme 1a).\textsuperscript{(14c)} However, the nature of these d⁸-ML₄-H•••C interactions and their description in terms of either (i) anagostic interactions, (ii) hydrogen-bonds, or (iii) pregostic interactions remains an ongoing matter of debate.\textsuperscript{(13)} We will therefore outline in the following that all three bonding concepts (i–iii; Scheme 1) basically rely upon an oversimplified description of the electronic situation in d⁸-ML₄ complexes, which causes a misinterpretation of the fundamental C-H bond activation processes in these types of compounds.\textsuperscript{(13a)}

We first reanalyze the potential energy surface (PES) of the σ model complex [Co(CO)]⁺[CH₄] (3) formed by methane and the positively charged d⁸[ML₄]⁺ metal fragment. In line with SH’s findings\textsuperscript{(13)} (see above), the EHT-PES predicts a repulsive 3c-4e \( M \leftrightarrow H \cdot \cdot \cdot C \) interaction in the methane adduct 3, which was selected by BS as benchmark model. In these model calculations one C-H moiety of the methane molecule was geometrically enforced to approach the square-planar d⁸-ML₄ metal fragment in an axial coordination mode (Supporting Information S2). However, considering electronic correlation effects via DFT methods reveals in contrast to the EHT results the actually attractive nature of the \( M \leftrightarrow H \cdot \cdot \cdot C \) interaction in this d⁸-ML₄ benchmark system (Figure S2a,b). The electronic situation of 3 is therefore characteristic for a H-C-H•••M₄ sigma complex, which at an early stage of the oxidative addition reaction coordinate exhibits a modest C-H activation due to a covalent interaction between an occupied \( \sigma(C-H) \) orbital and a vacant metal orbital (Scheme 1d).

Motivated by this result, we continue to reanalyze the electronic situation in 2 – as a benchmark system of an uncharged d⁸-ML₄ complex displaying an axial \( M \leftrightarrow H \cdot \cdot \cdot C \) interaction. In the presence of a \( M(dz²) \leftrightarrow \sigma(C-H) \) σ-type donation, as displayed by model complex 3 (Scheme 1d), we would expect at least a subtle elongation of the bridging C-H bond. Indeed, 2 displays a \( r(C-H\text{-ago}) \) stretching frequency at 2805 cm⁻¹ \([2878\text{ cm}^{-1}]\) which is clearly shifted to lower frequencies with respect to the \( v(C-H\text{-ago}) \) stretching mode of 2864 cm⁻¹ \([2917\text{ cm}^{-1}]\) (Figure 2) of the non-coordinating methane hydrogen atom of the same methylene group (Scheme 2).

Note, that theoretical values will be specified in square brackets in the following. Utilizing McKeen’s empirical correlation, which links \( r(C-H) \) bond lengths to isolated \( \sigma(C-H) \) stretching frequencies, predicts a slight activation of the C-H\text{-ago} bond \( r(C-H\text{-ago}) = 1.111 \text{ [1.108 Å]} \) in comparison to its non-agostic reference moiety \( r(C-H\text{-ago}) = 1.105 \text{ [1.104 Å]} \).\textsuperscript{(14a,b)} Hence, the bridging M\leftrightarrowH•••C moiety of 2 displays a minute, noticeable C-H bond elongation which supports the presence of weak but attractive \( M(dz²) \leftrightarrow \sigma(C-H) \) interaction in 2. Also in case of our agostic benchmark system 1 the observed \(2704\text{ cm}^{-1}\) and calculated \(2709\text{ cm}^{-1}\) \( v(C-H) \) stretching modes of the M\leftrightarrowH•••C moieties are in agreement with a subtle C-H bond activation. Indeed, the theoretically predicted C-H\text{-ago} bond length of 1.128 Å is slightly longer than the non-coordinating C-H’’ reference moiety \( 1.104 \text{ Å} \). Hence, the observed \( v(C-H\text{-ago}) \) stretching modes in the bridging C-H\leftrightarrowM moieties of 1 and 2 fall into the typical range of complexes displaying weak agostic interactions; e.g. in the β-agostic complexes EtTiCl₃(dmpe) \( (4, \text{ where } \text{dmpe} = \text{Me}_2\text{PCH}_2\text{CH}_2\text{PMe}_2) \) \( v(C-H\text{-ago}) = 2585\text{ cm}^{-1}\); \( r(C-H\text{-ago}) = 1.131 \text{ Å} \)\textsuperscript{(14d)} and \( \text{CpTi(iPr)(N)Cl}_3 \) \( (5, v(C-H\text{-ago}) = 2716\text{ cm}^{-1}; r(C-H\text{-ago}) = 1.120 \text{ Å}) \).\textsuperscript{(15)}

**Figure 1.** Overlay of structural fragments of 2 obtained by experimental X-ray diffraction studies at 0.0 and 5.0 GPa pressure using a diamond anvil cell (DAC); 50% ellipsoids. For two of the nitrogen atoms (marked by stars) only the thermal ellipsoids at 5 GPa were drawn to illustrate the significant reduction of thermal motion in 2 at elevated pressures. Values of the Ni-C bond distances are given in Å.

To gain an even deeper insight into the nature of these subtle d⁸-M\leftrightarrowH•••C interactions we propose to analyze directly the chemical response of these d⁸-M\leftrightarrowH•••C moieties upon shortening the respective M\leftrightarrowH···H distances. Such concept can be experimentally accomplished by combined X-ray diffraction (Figure 1) and IR spectroscopic (Figure 2) studies at various pressures. Figure 1 reveals that the M\leftrightarrowH···H contacts between the metal and the coordinating methylene groups in 2 becomes significantly shortened with increasing pressure: \( r(M\leftrightarrowH···H) = 3.18 \) \([3.27 \text{ Å}]\) and \(3.06 \) \([3.14 \text{ Å}]\) at 0 and 5 GPa, respectively. Periodic DFT calculations\textsuperscript{(4b)} also reveal that the respective M\leftrightarrowH···H distances are shortened upon increasing pressure (about 0.1 Å at 5 GPa; \( r(M\leftrightarrowH···H) = 2.531 \text{ Å} \) and 2.431 Å at 0 and 5 GPa, respectively). To study the pressure induced effect on the individual C-H bonds we also recorded IR spectra at approximately the same pressures as employed for the X-ray studies. Figure 2 clearly demonstrates that all \( v(C-H) \) stretching modes of the pyrazolyl and ethyl groups in 2 are shifted to higher frequencies (blue shifts) – except for the \( v(C-H\text{-ago}) \) modes of the agostic Ni\leftrightarrowH···C moieties – at increased pressure. This is in line with the theoretical predictions which suggest that all C-H bonds of 2 display shorter lengths at elevated pressures, except for the agostic bonds: e.g. \( r(C-H\text{-ago}) \) bond shrinks significantly by 0.09 Å while the bond lengths of the agostic C-H’’ moieties remain rather constant between 0.0 and 5.0 GPa. Accordingly, the corresponding \( v(C-H\text{-ago}) \) mode does not show any significant pressure-dependency while the corresponding \( v(C-H\text{-ago}) \) mode and all other \( v(C-H) \) modes of 2 shift to higher frequencies (blue shifted). The expected pressure-induced shortening of the C-H bonds in the agostic Ni\leftrightarrowH···C moieties of 2 appears therefore to be equally balanced and compensated for by the increasing bond activation/elongation under pressure. Hence, shortening of the Ni\leftrightarrowH···H bond distance in the agostic C-H groups activates the bridging C-H\text{-ago} in line with the attractive nature of these Ni\leftrightarrowH···C bonds and their classification as \( M(dz²) \leftrightarrow \sigma(C-H) \) σ-type agostic interactions. In the case where this type of interactions is more pronounced and/or further enhanced (e.g. by \( M(dz², yz) \leftrightarrow \sigma(C-H) \) back donation) one should expect even a red
shift of the C-H$_{\text{agostic}}$ frequencies. This is indeed the case for our agostic benchmark system 1, where the C-H$_{\text{agostic}}$ frequencies shift from 2704 to 2670 cm$^{-1}$ by an increase of pressure from 0.0 to 7.4 GPa.

![Figure 2](image_url)  
**Figure 2.** Pressure-dependency of the υ(C-H) stretching modes of the pyrazolyl and ethyl groups in 2. Note that all these stretching modes display a blue shift with increasing pressure but the agostic υ(C-H$_{\text{agostic}}$) mode of the metal-coordinated Ni•••H$_{\text{agostic}}$-C moieties. This clearly signals the bonding character of these Ni•••H$_{\text{agostic}}$-C interactions in 2.

However, to characterize the nature of these agostic Ni•••H$_{\text{agostic}}$-C interactions in 2 in terms of a M(d$^5$) ← σ(C-H) σ-type donation process in our model system we need to clarify whether the Ni(d$^2$) orbital can be considered as vacant or at least as a partially depleted acceptor orbital (Scheme 1d). Indeed, analysis of the negative Laplacian of the experimental charge density distribution, $L(r) = -\nabla^2 \rho(r)$, in the valence shell of the metal atom of 2 clearly reveals the presence of a local charge depletion zone (denoted “CD”) in Figure 3c) in axial (z) direction (Figure 4c). This $L(r)$ feature is in conformity with a partially vacant Ni(d$^2$) orbital since the Laplacian is a well-established and experimentally accessible indicator to map regions of local charge concentration ($L(r) > 0$ eÅ$^{-3}$).

The weakness of the agostic interaction in 2 also becomes obvious by direct comparison with the d$^3$-Ni complex [(DCpH)Ni(dtbpe)]$^+$/[BF$_4$] (6 DCPH = dicyclopentenyl) which is our benchmark system for compounds displaying strong agostic interactions (Figure 4b): in that case the Ni•••H bonding is well pronounced ($\rho(r) = 0.553(4)$ e/Å$^3$), the Ni-H$_{\text{agostic}}$ bond distance (1.671(9) Å) is significantly shorter than in 2 (2.376 Å) and the C-H activation is large (C-H = 1.20(1) Å).

In the final step of our analysis we will focus on the NMR properties of 1 and 2 which display in $^1$H NMR experiments rather different chemical shifts of the agostic protons (Scheme 2). The agostic protons in 1 are clearly shifted upfield ($\delta (^1H) = -5.88$ ppm) but downfield in 2 ($\delta (^1H) = +6.1$ ppm). However, the calculations show, that the upfield/downfield shift of these agostic protons is not correlated with their “hydridic” character – as originally proposed by Trofimenko[36] and also postulated in the BG model[11] of agostic interactions. The agostic hydrogen atoms in 1 and 2 display both only small negative atomic charges and thus differ only marginally ($Q_{\text{agostic}} = -0.14$ and -0.08 e, respectively). As outlined in greater detail in a recent publication, the chemical shifts of agostic protons in transition metal alkyls rather depend on the polarization pattern of the valence shell density of the metal and/or the topology of the induced current density distributions in magnetic fields than on their “hydridic character”.[36]

In the case of all agostic complexes studied...
so far by topological analyses of the charge and/or current density, pronounced upfield shifts are only observed when agostic C-H moieties are pointing toward a pronounced local Lewis acidic site in the valence shell of the metal atom.\(^{[17b]}\) Indeed, inspection of the Laplacian maps of 1 and 2 (Figure 3) reveals, that only in case of 1 the agostic proton is “intruding” into a pronounced charge depletion zone, which represents a local Lewis acidic center in the charge density picture, or a vacant metal d-orbital in the MO model. A similar topological scenario is observed for the agostic protons in 2, however, the axial Lewis acidic sites in the valence shell of the nickel atom are less pronounced relative to the d\(^4\) complex 1. Furthermore, the chemical shifts in planar d\(^8\) complexes are also strongly influenced by the topology of the current density which is dominant in the equatorial plane of square-planar d\(^8\) complexes and strongly influenced by the topology of the current density which is provided a more direct insight in the nature of these interactions outlined above combined high pressure IR and diffraction studies.\(^{[13c]}\) Thus, the downfield chemical shift of the agostic proton in 2 does not rule out its agostic character. We therefore suggest, not to use the sign of \(^1\)H NMR shifts as major criterion to classify M····H–C interactions as attractive (agostic) or repulsive (anagostic). As outlined above combined high pressure IR and diffraction studies provide a more direct insight in the nature of these interactions – especially in combination with the analysis of the topology of the charge and current density.

To conclude: We have unequivocally revealed that square-planar d\(^8\)-ML\(_4\) complexes display subtle but noticeable local Lewis acidic sites in axial direction in the valence shell of the metal atom. These sites of local charge depletion provide the electronic prerequisites to establish weakly attractive 3c-2e M····H–C agostic interactions (Scheme 1d and Figure 4c) which are controlled by the extent of M(d\(^2\)) → σ(C–H) donation. We therefore suggest to use more carefully the expression “anagostic interactions” which describes M····H–C interactions in square-planar d\(^8\)-ML\(_4\) in terms of repulsive 3c-4e interactions. Indeed, the latter bonding concept is based on a purely electrostatic description of M····H–C interactions in d\(^8\)-ML\(_4\) complexes in terms of crystal field theory and ignores the consequences of strong covalent metal ligand interactions which might induce local Lewis acidic sites in the valence shell of the metal in axial direction.

Received: ((will be filled in by the editorial staff))
Published online on ((will be filled in by the editorial staff))

**Keywords:** charge density · bond activation · agostic interactions · high pressure


[7] a) Complex 2 was synthesized according to modified literature methods (see ref 5 and the Supporting Information). The high-resolution data set for the 100K data provided 98.9% completeness in a 2.48°<2\(θ\) range of 550–8000 cm\(^{-1}\) using a Bruker IFS 113v coupled to a Bruker IFS 66/S FT-IR spectrometer. A crystal among many possible cell was used with a pre-intended CuBe gasket and nitrogen as hydrostatic pressure-transmitting medium.


[10] (a) Molecular DFT calculations were performed with ADF using the BP86 or PBE0-functional, the ZORA for the description of scalar relativistic effects and the TZ2P or JCP basis set as implemented in ADF. (b) The periodic B3LYP/Def2-TZVP DFT calculations used the CRYSTAL09 code; see the Supporting Information for further details.


