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# Pushing the limits of electron donation for cis-chelating ligands via an alliance of phosphonium ylide and anionic abnormal NHC†

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The grafting of a -(CH<sub>2</sub>)<sub>2</sub>PR<sub>3</sub><sup>+</sup> moiety on an NHC ligand backbone in the Mn(I) complex [Cp(CO)2Mn(IMes)] followed by double deprotonation opens a route to bidentate ligands with extreme electrondonating character. Such remarkable electronic properties can even allow intramolecular sp<sup>2</sup> C-H functionalization in typically inert square-planar Rh(ı) dicarbonyl complexes.

Strongly electron-donating ligands have been widely investigated over the last three decades, leading to decisive advances in many fields such as homogeneous catalysis, photophysics<sup>2</sup> and materials science.3 The isolation of the first stable 2Himidazol-2-ylidene (A, Chart 1) by Arduengo in 1991<sup>4</sup> was a key milestone in this continuing quest, from which N-heterocyclic carbenes (NHCs) became essential for modern chemistry. The successive preparations of imidazol-4-ylidene (B)<sup>5</sup> and anionic imidazol-2,4-diylidene (C),6 so-called "abnormal"  $(aNHC)^7$  and "ditopic" (dNHC)8 carbenes, were integral parts of the progress made to access increasingly electron-donating carbon ligands based on the imidazolyl core (Chart 1, top). Indeed, aNHC ligands **B** exhibit stronger  $\sigma$ -donor and weaker  $\pi$ -acceptor character than their "normal" NHC counterparts A<sup>10</sup> and these electronic effects are even more pronounced in dNHCs metallated at the C2 position, 11 due to the presence of an additional delocalized negative charge in the heterocyclic moiety. Beyond different classes of NHCs, phosphonium ylides derived from triarylphosphines (D) and trialkylphosphines (E) (Chart 1, down) constitute an alternative class of electron-rich neutral C-ligands for transition metals<sup>12</sup> and main group elements.<sup>13</sup> Moreover, a systematic IR spectroscopy investigation of a series of C,C-chelating Rh(1) dicarbonyl complexes with three different combinations of NHC (A) and phosphonium ylide (D) donor extremities revealed a decrease in the average  $\nu_{CO}$  stretching frequencies in the order A-A > A-D > D-D, thus demonstrating the superior electron donation of phosphonium ylide vs. conventional NHC (Chart 1, bottom).14 In addition, it was recently shown for Pd(II) carbonyl complexes that a PCy3-derived phosphonium ylide fragment (E) is an even stronger donor than is its PPh3-containing analogue (D).15 In a logical continuation of our research program focused on the association of classical NHCs with phosphonium ylides in bidentate, 14-16 pincer 15,17 and tetradentate<sup>18</sup> scaffolds, we envisioned to combine for the first time the strongest donors of each family with the aim of approaching the upper limits of the electronic scale in chelating systems. 19 In this contribution, we report the preparation, coordination chemistry with rhodium and evaluation of electronic properties of anionic abnormal NHC/phosphonium ylide ligands

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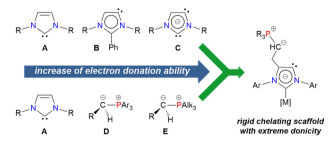


Chart 1 General trends in electron donation for different types of imidazole-based N-heterocyclic carbenes (top) and phosphonium ylides (bottom) as well as the structure of hybrid bidentate ligands proposed in this work (right)

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(Chart 1, right) as well as unusual C-H functionalization in Rh(I) dicarbonyl species.

For the assembly of the anionic abnormal carbene platform, we chose a readily available, on a multi-gram scale, Mn(1) complex 1<sup>20</sup> (Scheme 1) bearing the common 1,3-bis(2,4, 6-trimethylphenyl)-2H-imidazol-2-ylidene (IMes) ligand, in which the organometallic fragment in addition to protecting the C2 carbene position can also serve as an excellent IR<sup>11</sup> and electrochemical<sup>21</sup> probe. The -CH<sub>2</sub>CH<sub>2</sub>OH arm required for grafting the phosphonium moiety was installed on the IMes ligand backbone via a direct functionalization approach<sup>22</sup> consisting of the deprotonation of 1 with *n*BuLi and the trapping of the resulting anionic abnormal carbene [Cp(CO)<sub>2</sub>Mn(dIMesLi)] with ethylene oxide. The subsequent transformation of complex 2 to the corresponding triflate derivative followed by onepot quaternization with PPh3 and PCy3 afforded phosphonium pre-ligands [3a](OTf) and [3b](OTf) in good yields (Scheme 1). Both of them were fully characterized and their exact structures were obtained using X-ray diffraction (Fig. 1). Their deprotonation with 2.5 eq. of nBuLi in THF at room temperature led to a nearly quantitative generation of extremely moisture-sensitive aNHC-ylides 4a and 4b as confirmed by IR spectroscopy having revealed the presence of two low-frequency  $\nu_{\rm CO}$  bands (Table 1) characteristic for Mn(1) complexes bearing anionic aNHC ligands. 20,22 The coordination of these species with a slight excess of [Rh(CO)<sub>2</sub>Cl]<sub>2</sub> dimer was performed in toluene at low temperature similarly as described earlier for the related MnFe dIMes derivative, 11 leading to the zwitterionic complexes 5a and 5b isolated in excellent yields (Scheme 1).

The identity of heterobimetallic MnRh complexes 5a and 5b was confirmed from spectroscopic methods and HRMS data. IR spectra of both products showed two pairs of  $\nu_{CO}$  bands in an approximately 1:1 intensity ratio (Fig. S23, S24, ESI† and Table 1) in agreement with a presence of [Mn(CO)<sub>2</sub>] and [Rh(CO)<sub>2</sub>] fragments exhibiting a cis-arrangement of carbonyl groups. The presence of a P-ylide moiety coordinated to the Rh(I) center was confirmed by the observation of <sup>31</sup>P NMR

нó [3a](OTf) 75% R = Cy [3b](OTf) 80% [Rh(CO)2CI]2 2.5 eq. *n*BuLi toluene, -40°C THF, r.t. (THF)<sub>n</sub> OC R = Ph 5a 88% R = Ph 4a R = Cv 90% R = Cv 4b

Scheme 1 Synthesis of anionic aNHC-phosphonium ylide ligands 4a-b and their Rh(ı) complexes 5a-b. Reaction conditions: (i) 1.2 eq. nBuLi, THF, r.t., 2 h; (ii) 2 eq. ethylene oxide, -40 °C to r.t., 2 h; (iii) 1.1 eq. nBuLi, THF, r.t., 20 min; (iv) 1.1 eq. Tf<sub>2</sub>O, PhCl, -40 °C to r.t., 1 h; (v) 3.0 eq. PPh<sub>3</sub> or 1.5 eq. PCy<sub>3</sub>, PhCl, 60-80 °C, 16 h.

signals at  $\delta_P$  41.2 and 43.6 ppm for 5a and 5b, respectively, being deshielded compared to those of their phosphonium precursors ([3a](OTf):  $\delta_P$  22.7 ppm; [3b](OTf):  $\delta_P$  31.3 ppm) as well as characteristic high-field <sup>13</sup>C resonances with appropriate multiplicity (5a:  $\delta_{\rm C}$  13.9 ppm (dd,  ${}^{1}J_{\rm CRh}$  = 27.5 Hz,  ${}^{1}J_{\rm CP}$  = 25.5 Hz); **5b**:  $\delta_{\rm C}$  6.4 ppm (dd,  ${}^{1}J_{\rm CRh}$  = 25.7 Hz,  ${}^{1}J_{\rm CP}$  = 19.6 Hz)). The characteristic 5a and 5b13C NMR signals resulting from the carbenic atoms of anionic imidazol-2,4-diylidenes were observed as singlets at  $\delta_{\rm C}$  199.9–200.5 ppm (C2) and doublets at  $\delta_{\rm C}$ 162.2-162.7 ppm with a  ${}^{1}J_{CRh}$  coupling constant of ca. 42 Hz (C4). Finally, each of the bimetallic complexes yielded four distinct <sup>13</sup>C NMR spectral CO signals and a full set of inequivalent resonances of mesityl groups, probably due to blocked rotation across their respective C-N bonds and the existence of a chiral center in the NHC backbone.

Inspection of the X-ray diffraction structure acquired of complex 5a (Fig. 1) showed the Rh(1) atom in a square-planar environment with the dIMes-ylide behaving as a cis-chelating ligand. While the Rh-carbene bond distance (2.045(2) Å) was measured to be very close to the values found in other Rh(1) complexes bearing IMes-derived aNHCs (2.038(3)-2.061(4) Å), <sup>23</sup> the Rh-ylide bond (2.189(2) Å) was found to be slightly longer than those of similar bidentate ylide structures (2.1250(14)-2.155(8)). 14,24 The Rh-CO bond in the position trans to the aNHC moiety was also observed to be slightly longer than that opposite to the phosphonium ylide, presumably reflecting higher trans effect of the former ligand. Inspection of the structure also showed, notably, the 5-membered rhodacycle to be virtually coplanar with the NHC fragment. And structural comparisons showed the NHC-Rh-ylide angle in 5a (78.98(8)°) to be significantly more acute that those observed in Rh(1)  $(94.0(3)^{\circ})^{14}$  or Pd(II)  $(91.9(2)^{\circ})^{16a}$  NHC-ylide complexes featuring more flexible 7- and 6-membered chelating cycles, respectively, and to be comparable only with those of related species based on more rigid NHC-bis(ylide) ligands (82.61(6)-85.85(5)°)<sup>25</sup> with a similar size of the metallacycle.

Access to the Rh(1) dicarbonyl derivatives 5a-b provided a perfect opportunity to quantify the electronic properties of the novel aNHC-ylide ligands using the frequencies of  $\nu_{\rm CO}$  bands in solution IR spectra, which is applicable for benchmarking both mono-9 and bidentate ligands. 19 While  $\nu_{CO}$  values for the [Cp(CO)<sub>2</sub>Mn] fragment in 5a and 5b were observed to be almost the same (Table 1) and to remain very close to that of the previously reported dIMes complex, 11 the average  $\langle \nu_{\rm CO} \rangle$  value for the  $[Rh(CO)_2]$  moiety of **5b** was found to be ca. 10 cm<sup>-1</sup> lower, consistent with the higher electron-donation ability of PCy3-derived ylide vs. its PPh3 analogue. The same trends were confirmed using cyclic voltammetry studies revealing a reversible Mn(I)/Mn(II) oxidation wave at  $-0.38 \text{ V} \text{ vs. Fc/Fc}^+$  followed by the occurrence of irreversible one-electron oxidation at +0.61 and +0.51 V vs. Fc/Fc<sup>+</sup> for the Rh(1) moiety in complexes 5a and **5b**, respectively. Very importantly, the values of the  $\langle \nu_{\rm CO} \rangle$ parameter for both 5a (2001 cm<sup>-1</sup>) and 5b (1991 cm<sup>-1</sup>) were observed to be far lower than those for known Rh(I) dicarbonyl complexes bearing bis(NHC) (2057 cm<sup>-1</sup>), 14 phosphinephosphonium ylide (2053 cm<sup>-1</sup>),<sup>24a</sup> bis(1,2,3,-triazol-4-ylidene)

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Fig. 1 Molecular geometries of complexes [3a](OTf), [3b](OTf) and 5a (25% probability ellipsoids, hydrogen atoms of aryl groups, co-crystallized solvent molecules and counter-anions are omitted for clarity). Selected bond lengths [Å], and angles (°). [3a](OTf): Mn1a-C3a 2.012(3), C4a-C5a 1.344(4), C7a-P1a 1.805(3); [3b](OTf): Mn1-C3 2.012(2), C4-C5 1.337(3), C7-P1 1.806(3); 5a: Mn1-C3 2.011(2), C4-C5 1.349(3), Rh1-C4 2.045(2), Rh1-C7 2.189(2), C7-P1 1.769(2) Rh1-C8 1.890(2), Rh1-C9 1.868(2), C4-C5 1.399(3), C4-Rh1-C7 78.98(8), C4-Rh1-C8 176.67(10), C7-Rh1-C9 174.16(8), C8-Rh1-C9 87.43(10), P1-C7-Rh1 115.69(10), C4-C5-C6-C7 3.9(3).

Table 1 Experimental IR data for complexes 4–6 in THF solution and the corresponding  $\nu_{\rm CO}$  frequency values computed at the wb97xd/tzvp level

	Experimental $\nu_{\rm CO}$ (THF, cm <sup>-1</sup> )		Calculated $\nu_{\rm CO}$ (non-scaled, cm <sup>-1</sup> )	
No.	$[Mn(CO)_2]$	$[Rh(CO)_2]$	$[Mn(CO)_2]$	$[Rh(CO)_2]$
4a 4b 5a 5b 6a	1899, 1829 1898, 1828 1906, 1838 1906, 1838 1912, 1845		2028, 1966 2028, 1965 2032, 1969 2031, 1968 2040, 1979	

 $(2047-2048~cm^{-1})$ ,  $^{26}$  NHC-iminophosphorane  $(2046~cm^{-1})$ ,  $^{27}$  NHC-phosphonium ylide  $(2039~cm^{-1})^{14}$  and bis(phosphonium ylide) ligands  $(2017~cm^{-1})$ ,  $^{14}$  thus highlighting the extreme donation ability of our developed systems.

Intrigued by these outstanding electronic properties, we decided to study the structure and bonding situation in bimetallic species by performing DFT calculations. To our delight, all the trends observed in the experimental IR spectra of 4a-b and their Rh(1) complexes 5a-b were essentially perfectly reproduced in the  $\nu_{\rm CO}$  frequency calculations (Table 1). The calculations yielded rather similar HOMOs localized at the [CpMn(CO)<sub>2</sub>] moiety in both species (Fig. 2, top), but markedly different LUMOs for 5a and 5b, revealing the main contribution of PPh<sub>3</sub><sup>+</sup> and [Rh(CO)<sub>2</sub>] fragments, respectively (Fig. 2, bottom). The global AIM charge of the [Rh(CO)<sub>2</sub>] fragment in complex 5b was found to be slightly lower than that in the case of 5a (0.171 vs. 0.177), consistent with a stronger electron donation of the PCy<sub>3</sub>-derived ylide moiety. Interestingly, the AIM charge distribution in carbonyl ligands coordinating the Rh(1) center in bimetallic complexes was found to be quite asymmetrical, accumulating more electron density on the CO in the position trans to the aNHC ligand (-0.207 and -0.225 for 5a and 5b,respectively) than on those having the cis arrangement (ca. -0.160). An ETS-NOCV analysis of the NHC-ylide bonding (Fig. S4, see the ESI† for details)—beyond showing the major primary channel responsible for σ-donation from both

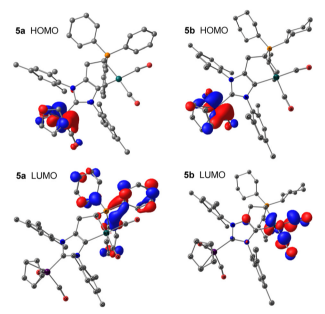


Fig. 2 Frontier molecular orbitals of complexes **5a** and **5b** (isosurface set at 0.05 a.u.).

*C*-ligands to the [Rh(CO)<sub>2</sub>] unit—also revealed an additional channel for the *a*NHC moiety having *ca.* 40% weaker associated energy, with the charge flowing only to the metal atom and opposite CO ligand (Fig. S4, ESI†). These results seemed to indicate the anionic *a*NHC ligand to be a stronger σ-donor than neutral phosphonium ylide regardless of the occurrence of some  $\pi$ -retrodonation for the metal–carbene bond (Fig. S5, ESI†).

Unlike complex  $5\mathbf{b}$ , which was observed to be stable in solution over long periods of time, its PPh<sub>3</sub><sup>+</sup> analogue  $5\mathbf{a}$  slowly transformed even at room temperature to another species, namely  $6\mathbf{a}$ , isolated in 95% yield after being heated at 50 °C (Scheme 2). While IR and cyclic voltammetry data for [Rh(CO)<sub>2</sub>] remained virtually unchanged (Table 1), both  $\nu_{\rm CO}$  frequencies (1912, 1845 cm<sup>-1</sup>)<sup>11</sup> and oxidation potential (-0.32 V  $\nu s$ .

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Scheme 2 Transformation of complex 5a into its isomer 6a via cyclometallation with the phenyl ring of the phosphonium moiety.

Fc/Fc<sup>+</sup>)<sup>21</sup> of the Mn(1) center clearly indicated its coordination with a regular imidazol-2-ylidene ligand, which was consistent with the re-appearance of the characteristic <sup>13</sup>C NMR resonance of the CH moiety belonging to the NHC backbone ( $\delta_{\rm C}$  122.9 ppm). Finally, the presence of the coordinated ylide moiety ( $\delta_{\rm C}$  22.5 ppm, dd,  ${}^1J_{\rm CP}$  = 32.2 Hz,  ${}^1J_{\rm CRh}$  = 24.1 Hz) and downfield signal at  $\delta_{\rm C}$  181.6 (pseudo t,  ${}^{1}J_{\rm CRh} = {}^{2}J_{\rm CP} = 33.7$  Hz) characteristic for cyclometallated phosphonium fragment, 17 unambiguously confirmed the structure of 6a as a zwitterionic heterobimetallic MnRh species bearing a bidentate aryl/phosphonium ylide ligand. A preliminary hypothesis concerning the formation of complex 6a might include the intramolecular activation of the ortho-C-H bond of a P+-Ph substituent by the electronrich metal site followed by a reductive elimination of the aNHC ligand in a putative Rh(III) hydride intermediate. Despite an extensive use of Rh(I) derivatives in catalytic C-H functionalization, <sup>28</sup> this transformation represented the first example, to the best of our knowledge, of intramolecular sp<sup>2</sup> C-H functionalization in typically inert square-planar Rh(I) dicarbonyl complexes.<sup>29</sup>

In summary, we have combined, for the first time, anionic abnormal carbene and phosphonium ylide to obtain structurally rigid cis-chelating ligands with extreme electron-donating properties. The incorporation of alkyl substituents at the phosphorus atom in such scaffolds improved both ligand donicity and stability of the resulting transition metal complexes. The application of these ligands in homogeneous catalysis as well as the studies on the comparison of electronic donation between aNHC and phosphonium ylide is currently ongoing in our group.

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# Data availability

The data supporting this article have been included as part of the ESI.†

## Conflicts of interest

There are no conflicts to declare.

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