α-Cationic Phospholes: Synthesis and Applications as Ancillary Ligands

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Abstract: A series of structurally differentiated α-cationic phospholes containing cyclopropenium, imidazolium, and iminium substituents has been synthesized by reaction of chlorophosphole 1 with the corresponding stable carbenes. Evaluation of the donor properties of these compounds reveals that their strong π-acceptor character is heavily influenced by the nature of the cationic group. The coordination chemistry of these newly prepared ligands towards Au(I) centers is also described and their unique electronic properties exploited in catalysis. Interestingly, α-cationic phosphole containing catalysts were not only able to accelerate model cycloisomerization reactions, but also to efficiently discriminate between concurrent reaction pathways, avoiding the formation of undesired product mixtures.

Introduction

In sharp contrast to pyrrole, which is a planar and strongly aromatic heterocycle, its immediately heavier analogue, phosphole, is characterized by a highly pyramidal arrangement of the substituents around the phosphorus and, as consequence, a severely diminished degree of aromaticity.[1,2] This geometry is adopted due to the high energetic cost required to planarize the phosphorus atom, which is not compensated by the aromatic stabilization won in a hypothetically planar conformation, even though theoretical calculations predict that in such planar environment phosphole would be more aromatic than pyrrole.[3] Hence, in phospholes the P-lone pair is bent away from the heterocycle and its overlap with the butadienyl fragment is drastically reduced.[3] This fact primarily defines the different reactivities of phospholes and pyrroles; in particular, the possibility of the former to form P-oxides, quaternary salts, or metal complexes by reaction at phosphorus.[4]

With regard to the use of phospholes as ancillary ligands, both experimental and theoretical analyses indicate that while they depict very similar α-electron-releasing properties to these characteristic of classical phosphines of analogue structure; their low lying LUMO, which largely corresponds to the mixture of the exocyclic α*(P-R) with a π*-orbital, confers them stronger π-acceptor character.[5] This results in enhanced back-donation from orbitals of the metal and, subsequently, in lower net electronic transfer to the metal centres they coordinate (Figure 1).[6]

During the last few years, our group has established a research program focused on the design and synthesis of strong acceptor ancillary ligands. These have been utilized to promote transformations in which the energetic span (ΔE) of the complete catalytic cycle is minimized by the enhancement of the electronegativity of the metal.[7] Specifically, we have prepared a complete series of α-cationic ligands by formal exchange of at least one -R substituent on typical phosphines by positively charged groups, such as cyclopropenium, imidazolium or pyridinium rests. Using this design, a series of Au(I), Pt(I) and Rh(I) catalysts depicting unmatched reactivity in hydroarylation[8] and cycloisomerization reactions[9] have been developed, together with enantioselective versions of the already mentioned processes.[10]

Given the intrinsic π-acceptor character that the phosphole architecture imposes to ligands derived from that heterocycle, we envisaged that such platform might be an ideal starting point to develop even stronger π-acceptor ligands by incorporation of positively charged substituents as exocyclic rests. In the resulting α-cationic phospholes the electron withdrawing effect of the cationic group should lead to a larger localization at phosphorus of the σ*(P-R) orbital, favouring the σ*(P-R)-π*(ring) mixture and, as consequence, lowering the energy of the LUMO (Figure 1). Modulation of this interaction by employing cationic groups of different nature should additionally allow the fine tuning of the π-accepting properties of the final α-cationic phosphole.

Figure 1. Frontier orbitals in phospholes and impact of the cationic rest on the LUMO energy.
Herein, the synthesis of such cationic phospholes is reported for the first time, their structure elucidated by means of NMR spectroscopy and X-ray crystallography, and their donor properties evaluated. Finally, their coordination chemistry towards AuI centres is studied, and the first practical applications in homogeneous catalysis of the obtained complexes are described.

Results and Discussion

Synthesis and Structure of α-Cationic Phospholes

The preparation of a prototype imidazolium-substituted phosphole was initially envisaged by reaction of 4,5-di-(Me)IMes with tetraethyl-substituted chlorophosphole 1, which was synthesized using the available routes.[11] After anion exchange with NaSbF$_6$, and washing of the crude reaction mixture with n-pentane, 2a was isolated as a white solid in 76% yield. In an analogue manner adducts 2b, 2c and 2d where obtained by reaction of 1 with 1,2-(diisopropylamino) cyclopropenyl-1-ylidene[12] or cAACs of different steric demand,[13] respectively (Scheme 1). The $^{31}$P [1H]NMR of these adducts display characteristic resonance signals at $\delta = -38.3$ (2a), $-41.0$ (2b), $-16.0$ (2e) and 0.8 ppm (2d). Compared with that of 1-phenyl-2,3,4,5-(tetraethyl)phosphate (s = 3.0 ppm), the significant up field shift of 2a, 2b and 2c can be attributed to the effect of the positively charged groups (Scheme 1).[7b,c] The shift of the $^{31}$P signal in 2d cannot be compared due to the tremendous steric hindrance of the Ad(cAAC) fragment, which significantly alters the geometry of the ligand.

The connectivity of salts 2a-d was subsequently confirmed by X-ray crystal analysis (See Figure 2 for the ORTEP plots, and the Supporting Information). In these structures the C1-P1 distances do not significantly differentiate along the series (1.80–1.86 Å), which are in the range of typical values for P–C(sp$^3$) bonds.[14] In addition, the sum of angles around P1: 302.64(12)$^\circ$, 2a; 304.47(18)$^\circ$, 2b; 295.00(52)$^\circ$, 2c; and 311.11(18)$^\circ$ for 2d, indicate that the central P atom adopts a pyramidal environment; albeit the pyramidalization slightly diminishes when bulkier cationic substituents need to be accommodated. Finally, in the solid state the P-atoms of 2a-d, are displaced out of the plane defined by the butadiene moiety (0.46 Å in 2a; 0.39 Å in 2b; 0.47 Å in 2c; and 0.78 Å in 2d). These parameters, together with the alternation of the C–C bond distances in the butadiene moieties corroborate the lack of aromatic character in the phosphole ring, something already known for the non-charged members of the family as well (see the Supporting Information).

Figure 2. X-ray structures of 2a–d. Hydrogen atoms, SbF$_6$ counterions and solvent molecules are removed for clarity; ellipsoids are set at 50% probability[31].

Steroelectronic Properties of α-Cationic Phospholes

IR stretching frequencies in trans-[RhCl(CO)L$_2$] complexes are generally used to evaluate the donor endowment of new ligands and establish their Tolman electronic parameter (TEP). Hence, we initially prepared complex 3b by reaction of [RhCl(CO)$_3$], with 2b. Comparison of the IR spectrum of 3b ($\nu_{CO} = 1984$ cm$^{-1}$) with that of 4 ($\nu_{CO} = 1971$ cm$^{-1}$), which contains a phosphine bearing a cyclopropenium substituent as well but no phosphole unit reveals, within experimental error, that 2b is a weaker net donor ligand.[16] This is attributed to the stronger acceptor properties imparted by the phosphole architecture (Scheme 2). Probably due to their excessive steric demand, ligands 2a and 2c,d do not form trans-[RhCl(CO)L$_2$] complexes by reaction with [RhCl(CO)$_3$], making this method inappropriate to compare their donor properties.[31] The X-ray structure of 3b has been resolved and is shown in the Supplementary Information.

Hence, the oxidation potential $E_{p}(ox)$ was chosen as an alternative parameter to rank the electronic properties of the

Scheme 1. Synthesis of α-cationic phospholes. Reagents and conditions: a) 1,3-dimesityl-4,5-dimethylimidazol-1-ylidene (1.0 equiv), Et$_2$O, −78°C, 2 h, and then NaSbF$_6$, −78°C−r.t., 76%; b) 2,3-diisopropylaminocyclopropenyl-1-ylidene (1.0 equiv), otherwise as a), 76%; c) Di-(Me)cAAC (1.0 equiv), otherwise as a), 60%; d) AdcAAC (1.0 equiv), otherwise as a), 46%.
new ligands. Cyclic voltammetry measurements afford the following values for the oxidation of ligands (\(E_{\text{p}}^{\text{ox}}\) = 1.32 V, \(2a\); 1.18 V, \(2b\) and 1.46 V, \(2c\), calibrated versus \(\text{Fc}^+/\text{Fc}\)), which suggest that \(2a\) and \(2b\) are, respectively, slightly better and worse donors than (MeO)\(_3\)P (1.287 V). Ligand \(2c\), due to the acceptor properties of the cAAC moiety, stands as the less basic member of the series. Its \(E_{\text{p}}^{\text{ox}}\) is very similar to the ones measured for \(\alpha\)-dicationic phosphines. These values however, should be considered with caution because of the irreversible nature of the oxidations monitored. The percent buried volume (%V\(_{\text{Bur}}\)) and the topographic steric maps of \(2a\)–\(d\) are shown in Figure 3. Given the quite rigid nature of the core architecture of these ligands (in particular \(2a\), \(2c\), \(d\)), which strongly limits the impact of conformational changes in solution; it can be concluded that upon coordination \(2a\) and \(2c\), \(d\) impose a geometry around the metal center very similar to that created by Buchwald biaryl phosphines, while this privileged environment cannot be provided by the more flexible \(2b\).[14]

**Scheme 2.** Synthesis of \(3b\) and comparison of its \(\nu_{\text{CO}}\) with that of \(4\).

**Figure 3.** Topographic steric maps for \(2a\)–\(d\).

**Synthesis of \(\alpha\)-Radical Phospholes**

Notably, during the electrochemical analyses, reversible reduction peaks were observed at \(-1.512\) V and \(-1.306\) (vs. Fe/Fe\(^+\)) for \(2c\), and \(2d\), respectively, which provided evidence of the stability of the corresponding neutral \(\alpha\)-radical phos- pholes. Encouraged by this result, we embarked on the isolation of these species. By treatment of the cationic precursor \(2c\) with one equivalent of potassium in graphite (K\(_\text{C}\)) in tetrahydrofuran (THF) at \(-78^\circ\)C, a deep-green solution was obtained, from which radical \(5c\) was isolated as black solid in moderate but reproducible yield (Figure 4a). Crystals suitable for X-ray analyses of this compound were grown by cooling their respective saturated pentane solutions to \(-20^\circ\)C. The Cl–P1 distance in \(5c\) (1.7976(9) Å) is notably shorter than that in their cationic precursor \(2c\) (1.867(6) Å), while the elongation of the Cl–N1 bond in radical \(5c\) (1.309(8) Å in \(2c\) and 1.3960(11) Å in \(5c\)) is consistent with a lesser degree of \(\pi\)-bonding between these two atoms as expected from a of the partial population of the C-N \(\pi^*\)-orbital (Figures 4b and 5).

A deeper insight into the electronic structure of \(5c\) was gained by X-band electron paramagnetic resonance (EPR) measurements from samples dissolved in dry toluene. The isotropic spectrum (Figure 4c) shows a g factor of 2.0028, which is consistent with an essentially C-based radical. Additionally, the resolved hyperfine splitting of the spectra

**Figure 4.** a) Synthesis of \(5c\). Reaction conditions: KC\(_8\), THF, \(-78^\circ\)C, 30%; b) X-ray structure of \(5c\), hydrogen atoms were removed for clarity and ellipsoids are set at 50 % probability;[15] c) Experimental (black) and simulated (red) X-band EPR spectra of \(5c\) (\(g = 2.0028\), \(\alpha_P = 5.36\) G, \(\alpha_N = 5.62\) G) in toluene solution at r.t. (G = gauss); d) Cyclic Voltamogram of \(5c\) in \(\text{CH}_2\text{Cl}_2\) calibrated versus \(\text{Cp}^\text{+}\)Fe/\(\text{Cp}^\text{+}\)Fe, \(\text{Bu}_4\text{NPF}_6\) (0.1 M).

**Figure 5.** a) Calculated spin densities of \(5c\) (left) and b) SOMO plot (right) at the UB3LYP-D3/def2-TZVP level. Total spin density drawn at an isosurface value of 0.0005 and SOMO drawn at 0.025.
reveals delocalization of spin density onto the adjacent N1 and P1 atoms; the quartet spectrum observed for 5c is the result of hyperfine coupling constants that are apparently identical (\( a_p = 5.36 \, \text{G} \); \( a_h = 5.62 \, \text{G} \)).

DFT calculations were performed at the UB3LYP-D3/def2-TZVP/UTPSS/def2-TZVP level[39] (see the Supporting Information for computational details). These calculations accurately reproduce the geometries observed experimentally in the solid state for 5c and provide a quantitative picture of the spin distribution (Figure 5). As can be seen from the plots of the localized Mulliken spin densities and that of the singly occupied molecular orbital (SOMO), the unpaired electron density is mainly located at the original carbene carbon atom derived from the cAAC fragment (ca. 68%). Moreover, there is considerable spin density on the N atom (ca. 21%), and only some residual one at the P center (less than 1%).

**Synthesis of Au\(^+\) Complexes**

Encouraged by this analysis and willing to exploit the stereoelectronic properties of the newly prepared cationic phospholes in catalysis, the corresponding Au\(^+\) derivatives 6a–d were prepared by reaction of (Me\(_2\)S)AuCl with phospholes 2a–d, respectively (Scheme 3). Very indicative of the formation of the desired Au\(^+\)-complexes is the downfield displacement of the \(^{31}\)P \([\text{H}]\)NMR chemical shifts upon coordination; \( \delta = -2.1 \) (6a); \(-2.2 \) (6b); 15.4 (6c); and 26.3 ppm (6d). The decision to employ Au-catalysis for the initial screening of cationic phospholes as ancillary ligands comes from our experience in Au-catalyzed processes, which often benefit from strong acceptor ligands, and other literature precedents[21].

The solid-state structures of complexes 6a–d were determined by X-ray diffraction analysis. The ORTEP diagrams of 6b and 6c are shown in Figure 6, and those of 6a and 6d can be found in the Supporting Information. The measured Au-1P1 bond lengths for these complexes, 2.2238(9) Å, 6a; 2.233(3) Å, 6b; 2.2151(6) Å, 6c and 2.2283(9) Å, 6d; are identical to the ones measured for neutral phosphole-Au\(^+\) complexes (P1-Au1, 2.20–2.24 Å).[17,22] Moreover, in structures 6a, 6c,d, a short contact between the Au atom and the C\(_{ipso}\) from the flanking aryl ring (3.0949(35) Å, 6a; 3.1691(22) Å, 6c; 2.9757(32) Å, 6d) can be observed.[23] In all cases the geometry of the P-Au-Cl fragment is slightly distorted from the expected linearity (P1-Au1-Cl1 angles: 176.5° in 6a; 175.1° in 6b; 173.2° in 6c and 174.4° in 6d).

**Catalysis**

As a preliminary comparison of the relative reactivity of complexes 6a–c with that of those derived from neutral or other cationic phosphines, we initially evaluated their performance in the intramolecular hydroarylation of sulfide 7 into thiepine 8 following a 7-exo-dig cyclisation.[24] The conversion versus time plot for the different precatalysts under otherwise identical conditions (2 mol% [Au], 2 mol% AgSbF\(_5\), r.t., CD\(_2\)Cl\(_2\)) is shown in Figure 7. To put our results into perspective, neutral phosphole 9, cationic phosphine 10, and phosphite Au complexes 11, were also prepared and used as precatalyst under otherwise identical reaction conditions (See the Supporting Information).

As expected, all cationic precatalysts tested performed better than 9. In addition, comparison of the plots obtained with 6b and 10 allows assessing the effect of the phosphole unit; this moiety makes 6b slightly more reactive than 10 and, interestingly, it seems to confer longer life to the catalytic species. Phosphite complex 11 depicts intermediate performance between 6b and 10. The most promising results in terms of conversion were provided by 6c containing the most strongly \( \pi \)-acceptor, cAAC-derived cationic moiety. We attribute the poor performance of 6a when compared to 6b to the higher steric demand imposed by the IMes moiety.

Finally, the catalytic performance of the Au complexes 6a–c was additionally evaluated in the Au-catalysed cyclisation of N-(3-iodoprop-2-ynyl)-N-rosylinelines 12a–c into the corresponding 1,2-dihydroquinolines. This particular transformation was chosen because two regioisomeric products are known to be produced; those derived from simple cyclisation 13a–c, and compounds 14a–c, which additionally involve a 1,2-iodo migration concomitant to the cyloisomerization. Moderate \( \pi \)-acceptor ligands attached to Au, as the phosphite in complex 11 have been reported to favor the first pathway.
Table 1: Effect of α-cationic phosphole ligands of different π-acceptor ability on the outcome of the Au-catalyzed cyclisation of tosylanylines 12 a–c.

| Entry | Substrate | Au cat. | Conv (%) | Ratio 13:14 |
|-------|-----------|---------|----------|-------------|
| 1     | 12a       | 11      | (64)     | 58:42       |
| 2     | 12a       | 15      | (85)     | 7:99        |
| 3     | 12a       | 6a      | >95      | 75:25       |
| 4     | 12a       | 6b      | >95      | 56:44       |
| 5     | 12a       | 6c      | >95 (80) | 97:3        |
| 6     | 12a       | 10      | >95      | 59:41       |
| 7     | 12b       | 11      | (79)     | 31:69       |
| 8     | 12b       | 15      | (80)     | <1:99       |
| 9     | 12b       | 6c      | >95 (79) | 97:3        |
| 10    | 12c       | 11      | 67       | 78:22       |
| 11    | 12c       | 6c      | >95 (95) | 99:1        |

[a] Results from ref. [24a]. [b] Conversions determined by 1H-NMR of the crude reaction mixtures; in parenthesis isolated yields; [c] Isomeric ratios determined by 1H-NMR.

While strong α-donating NHCS preferentially induce a 1,2-iodine shift prior to the cyclisation event (Table 1, Entry 2),[25] Hence, this transformation seems to be ideal to further test the unique donor properties of ligands 2a–c. Discrimination between competing reaction pathways and not only a kinetic effect is expected here.[26]

The product ratio obtained using precatalysts 6b is comparable to those afforded by 10 and 11, confirming the donor ability rank previously established. The selectivity towards 13a improves when precatalyst 6a is employed, and reaches a remarkable 97:3 ratio for cAAC-derived 6c (Table 1, Entries 3–5). Identical tendency is observed for substituted substrates 12b (R = Cl) and 12c (R = C(O)Me) (Table 1, Entries 7–11). Especially remarkable is the effect of precatalyst 6c in the cyclisation of 12b. In this case the phosphole ligand is even able to invert the product distribution of the reaction if compared to 11 and 15 (entries 7–9). Compound 13b is thus by far the major product of that reaction with a 13b:14b ratio of 97:3.

Conclusion

An efficient method for the preparation of α-cationic phospholes differing on the nature of the cationic group is reported, together with the synthesis of their Au-complex derivatives. Our experiments suggest exceptional electrophilicity at the Au center that can be used to accelerate otherwise difficult transformations and importantly, to discriminate between competing reaction pathways. Given the ease of handling of ligands 2a–d, we suspect that a potentially large number of Au-catalyzed transformations might benefit from their use. Ongoing research in our group is focused in this direction. We also target the application of cationic phosphole ligands in processes catalyzed by metals other than Au.

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Conflict of interest

The authors declare no conflict of interest.
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