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Theoretical study of the activity in Rh-catalysed hydroformylation. The origin of enhanced activity of \( \pi \)-acceptor phosphinidine ligand

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The factors governing the activity in Rh-catalyzed hydroformylation were investigated using a set of computational tools. We performed DFT calculations on the phosphinidine-modified Rh catalyst [HRh(CO)\(_3\)(PC\(_5\)H\(_5\)R\(_3\))] and compared it to the phosphine-modified HRh(CO)\(_3\)(PR\(_3\)) and HRh(CO)\(_2\)(PR\(_3\))\(_2\) complexes. The \( \pi \)-acceptor phosphinidine ligand coordinates preferentially at the equatorial site of pentacoordinated Rh complex with the heterocycle perpendicular to the equatorial plane, although the ligand freely rotates around the Rh-P bond. The overall energy barrier can be divided into the following contributions: alkene complex formation, alkene rotation and alkene insertion. In the absence of steric effects (model systems), the overall barrier correlates with the computed barrier for alkene rotation. This proves that \( \pi \)-acceptor ligands reduce backdonation to the alkene, leading to a lower rotational barrier, and consequently, to a higher activity. The Rh-P donor-acceptor interactions were quantified using a modified version of energy decomposition analysis (EDA). In Rh-phosphinidine systems, the efficient directionality of the \( \pi \)-backdonation, rather than the overall acceptor ability, is responsible for the high catalytic activity. Introducing steric effects increases the energy required to coordinate the alkene, increasing the overall barrier. The factors governing the activity in Rh-monophosphine catalysts seem to be related to those derived for Rh-diphosphine during the development of a QSAR model (Catal. Sci. Technol. 2012, 2, 1694). To investigate whether the findings for mono- can be extrapolated to diphosphine ligands, we re-examine our previous QSAR model using the Topological Maximum Cross Correlation (TMACC) method based on easy-to-interpret 2D-descriptors. The TMACC descriptors highlight heteroatoms close to phosphorus as activity-increasing atoms, whereas highly substituted carbon atoms groups are highlighted as activity-decreasing groups.

Introduction

The hydroformylation of alkenes is one of the largest applications of homogeneous catalysis in industry.\(^1\) It consists of the addition of carbon monoxide and hydrogen to alkenes, catalyzed by cobalt, rhodium or platinum catalysts, leading to the formation of aldehydes. For industrial production, the phosphane-modified rhodium catalysts are widely used because they can show high activity and regioselectivity towards the usually desired linear aldehyde.\(^1,2\)

Scheme 1 shows the generally accepted mechanism for the hydroformylation of alkenes catalyzed by phosphane-modified rhodium catalysts.\(^3\) Different reaction kinetics have been observed experimentally depending on ligand properties that, in general, fit into one of two extreme cases. For electron-poor ligands such as bulky monophosphites and the unmodified rhodium-carbonyl catalysts, the hydrogenolysis of acyl species \(^6\) controls the overall rate of hydroformylation.\(^3,5\) On the contrary, the kinetics for electron-rich ligands are consistent with a rate-determining step early in the catalytic cycle. Recently, a combination of isotope effects study and computational analysis have demonstrated that the overall process from the resting state species 1 to hydride migration (from complex 3 to complex 4) governs the overall activity in 1-octene hydroformylation catalyzed by the rhodium-xantphos complexes.\(^6\) Independently, a theoretical study by Jensen and co-workers led to the same conclusion for phosphane and moderately electron-withdrawing phosphite ligands; whereas for strongly electron-withdrawing ligands, calculations supported hydrogenolysis as the rate-determining step.\(^7\)

Recent clarification of the rate determining step\(^6,7\) indicates that the ligands that promote CO dissociation, alkene coordination, or hydride migration might yield higher catalytic turnovers. Besides this, some systematic studies have attempted to establish correlations between ligand structure and catalytic activity.\(^8,9,10,11\) Early experimental studies on monodentate P-donor ligands showed that a relationship exists between ligand basicity and catalyst activity; thus the least basic phosphanes enhance the activity.\(^8\) In addition, van Leeuwen and coworkers showed that phenoxophosphane (phosphacyclic) moieties are less basic than diphenylphosphino moieties and exhibit an increase in
the rate.\textsuperscript{9} In general, for the ligands enclosing a phosphorus atom inside a cycle, an increase of the activity was observed and was attributed to a lower basicity.\textsuperscript{12}

The differences in catalytic activity have also been related to the steric features of the ligands. For a series of phosphinine ligands, the variation in catalytic performance was attributed to the steric properties of the ligand.\textsuperscript{13} According to the authors, the bulky phosphinine ligands favour the formation of a monoligand rhodium species, which should have a larger accessible space compared with diligand rhodium species. Similarly, in industrial conditions excess of phosphine ligand is used because the selectivity towards linear aldehyde is improved; however the activity is reduced because the phosphine dissociation equilibrium shifts from the monophosphine [HRh(CO)\textsubscript{2}(PR\textsubscript{3})] towards the less active HRh(CO)(PPh\textsubscript{3})\textsubscript{2} species. On the other hand for a series of diphenyl phosphine-type ligands,\textsuperscript{14} the rate increases with increasing the bite angle,\textsuperscript{15} while we showed that increasing the bite angle increases the steric hindrance around the metal center.\textsuperscript{16} During the development of a quantitative structure-activity relationship (QSAR) model for the hydroformylation catalyzed by Rh-diphosphine complexes, we discovered that complex relationships underlie the origin of the activity; and that both the shape and the electronic properties of the catalyst need to be considered.\textsuperscript{17}

Here, we focus on the study of phosphinine-modified rhodium catalyst \([HRh(CO)\textsubscript{2}(PC\textsubscript{2}H\textsubscript{2}R\textsubscript{3})_2]\), for which computational investigations are still lacking,\textsuperscript{18} and compared it with the phosphine-modified catalysts [HRh(CO)\textsubscript{2}(PR\textsubscript{3})] and [HRh(CO)\textsubscript{2}(PC\textsubscript{2}H\textsubscript{2}R\textsubscript{3})\textsubscript{2}] (Scheme 2). The Rh-phosphine system showed a much higher activity than classical PPh\textsubscript{3}-based catalysts,\textsuperscript{13,19} and followed an analogous kinetics.\textsuperscript{13} Our aim is to understand at molecular level the factors governing the activity, evaluating them and their interplay. In addition, we investigate whether the findings for monophosphine ligands can be extrapolated to diphosphine, re-examining a previous QSAR model\textsuperscript{17} by using the easy-to-interpret TMACC descriptors.\textsuperscript{20}

\begin{align*}
\text{Scheme 2}
\end{align*}

\textbf{Results and discussion}

\textbf{Coordination preferences of phosphinine ligand}

The coordination and interaction with rhodium of phosphine ligands have been extensively studied by computational methods\textsuperscript{21,22} but to the best of our knowledge there are no such studies on phosphinine ligands. Thus, initially, we analyzed the coordination mode on [HRh(CO)\textsubscript{2}(2,4,6-PC\textsubscript{2}H\textsubscript{2}R\textsubscript{3})] (R = H, Ph) complexes, which correspond to the resting states in Rh-catalyzed hydroformylation. The phosphinine ligand can coordinate as equatorial (e) and apical (a). It is also possible to generate two additional isomers for e coordination depending on whether the heterocycle is perpendicular to the equatorial plane of the complex or if it is in-plane (1e1 and 1e2, respectively in Scheme 3 and Figure 1).

Previous high pressure NMR studies on the analogous and more stable iridium(I) system had indicated that only one phosphinine ligand is coordinated to the transition metal center under hydroformylation conditions.\textsuperscript{13} These experiments also suggested an equatorial position for the phosphine, presumably as two rotamers. The DFT calculated relative energies support the experimental proposal. For [HRh(CO)\textsubscript{2}(PC\textsubscript{2}H\textsubscript{3})] model complex, the most stable isomer is 1e1\textsuperscript{H} followed by 1e2\textsuperscript{H} and 1a\textsuperscript{H} (+1.1 and +1.4 kcal.mol\textsuperscript{-1} higher in energy, respectively). The preference for equatorial coordination with the phosphinine cycle perpendicular can be explained from electronic arguments based on frontier molecular orbitals (FMO). The in-plane d\textsubscript{xy} orbital of the metal fragment is high in energy and hybridized away from the other equatorial ligands favouring the \pi-back donation to the phosphorus p orbital perpendicular to the ligand plane.\textsuperscript{23} This is reflected in the shortest Rh-P distance for 1e1\textsuperscript{H}: 2.37 Å vs. 2.39 and 2.41 Å for 1e2\textsuperscript{H} and 1a\textsuperscript{H}, respectively.

Introducing the ligand steric effects via calculations on [HRh(CO)\textsubscript{2}(2,4,6-PC\textsubscript{2}H\textsubscript{2}Ph\textsubscript{3})\textsubscript{2}] complexes did not change the order in relative energies (0.0, +0.5 and +3.3 kcal.mol\textsuperscript{-1} for 1e1, 1e2 and 1a). Nevertheless, the energy difference between the two equatorial rotamers diminishes, while between equatorial and
apical coordination it increases. Both trends can be attributed to the steric effects. The ideal P-Rh-CO angle in 1e1 and 1a is 90°, whereas in 1e2 it is 120° (see Figure 1). Thus, the two former, 1e1 and 1a, with a smaller angle are slightly destabilized respect to 1e2 by the steric interactions between phosphinine substituents and the auxiliary ligands.

Fig. 1. Calculated 3D molecular structures for the different geometric and conformational isomers of [HRh(CO)(PC₅H₅)] complex 1e1, 1e2 and 1a. Relative energies in kcal.mol⁻¹.

In order to analyze the rotation around the Rh-P bond in e compounds, we performed a relaxed energy scan of the H-Rh-P-Cortho dihedral angle. The calculations did not show any barrier connecting the rotational isomers 1e1H and 1e2H. This along with the small energy difference between the rotamers, even for the 2,4,6-triphenylphosphinine system, strongly indicates that the Rh-P bond rotates freely. This might hamper the application of axially chiral monodentate phosphinines in the asymmetric hydroformylation of prochiral substrates. As matter of fact, the preferred equatorial coordination would place the chiral centers far away from the apical region, in which the key ligand-substrate interactions should take place. For example, ligands based on axial chirality such as Binaphos, which shows excellent performance in Rh-catalyzed asymmetric hydroformylation, induce enantioselectivity via interactions between the substrate and the axially chiral groups of the apical ligand moiety. Moreover, the very low barrier computed for ligand rotation would allow easy ligand reorganization upon interaction with the incoming substrate, leading to low enantiodiscrimination.

Table 1. Overall (ΔE(rot)alkene insertion (ΔE(inkr) and rotation (ΔE(inkr) energy barriers, and alkene coordination energy (ΔE(alkene)).

| phosphinine | mono-phosphane | bi-phosphane |
|-------------|----------------|-------------|
| ΔE(rot)alkene | 19.0 (19.3) | 22.4 (21.9) | 25.9 (22.5) |
| ΔE(inkr) (3→TS) | 7.7 (10.8) | 10.9 (13.5) | 10.8 (13.3) |
| ΔE(inkr) (3→TS) | 4.5 (5.9) | 7.1 (7.3) | 5.6 (8.6) |
| ΔE(alkene) (1→3) | 11.3 (8.5) | 11.5 (8.5) | 15.1 (10.2) |

*Energies in kcal.mol⁻¹ [HRh(CO)(PC₅H₅)] systems (R = Ph, and values in parenthesis for R = H).

Next, we analyzed the coordination preferences of the phosphinine ligand in the rate-determining step, in which the transition state for alkene insertion into the Rh-H bond is involved (Figure 2). Using ethene as a model substrate, the computed equatorial path for the model system, TSe1H, is lower in energy than the apical, TSe1H, by 2.0 kcal.mol⁻¹, increasing somewhat the preference found in the resting-state (1.4 kcal.mol⁻¹). For equatorial coordination, the attempts to locate a transition state with the phosphinine parallel to the equatorial plane ended in the corresponding TS for path e1. The perpendicular disposition of the ligand favours back-donation from the Rh, and in turn, reduces back-donation to the alkene, which then can easily rotate to reach the TS for insertion. If we recall the low rotational barrier for the Rh-P bond, it reasonable to think that in case that the alkene complex 3e2 is formed, the system would tend to switch to the lower-energy easily-accessible e1 reaction channel. Thus, calculations indicate that most of the reaction will occur through a channel with the phosphinine in an equatorial position and perpendicular to the equatorial plane.

Analysis of the overall energy barrier for Rh-phosphinine and phosphane systems.

The overall energy barrier can be computed as the energy difference between the transition state for alkene insertion (TS) and the rhodium hydride-carbonyl resting-state of the catalyst 1.67 To identify the individual factors governing the activity, we decomposed the overall barrier into the energetic cost of several steps: the formation of the alkene complex (ΔE(alkene)) and the energy barriers for alkene rotation (ΔE(inkr) and insertion (ΔE(inkr)). Note that the alkene insertion proceeds through rotation of the alkene moiety out of the equatorial plane of the trigonal-bipyramidal (bpt) complex 3, followed by the transfer of the hydride moiety to generate the alkyl species. Table 1 collects the values of the most favourable path for each system, and Figure 3 shows some of the key structures. For mono- and bicoordinated PPh₃ systems the lowest energy paths are the equatorial and the equatorial-equatorial (see Supporting Information). Interestingly, at the ONIOM level, in which the electronic effects of Ph substituents are neglected, the equatorial-apical path is more favoured. This can be rationalized as follows: assuming similar steric interactions at both levels, the electronic effects of the Ph groups reduce the donation ability of the phosphane; and consequently, its tendency to be placed in apical position. Thus, although the results need to be viewed cautiously, it is more straightforward to compare equatorial paths.
The overall energy barriers reproduce the experimental observations, providing a clear picture about the activity differences. The phosphinine system shows a lower value (19.0 kcal.mol\(^{-1}\)) than the monophosphane complex (22.4 kcal.mol\(^{-1}\)), while the bis(phosphane)-based complex shows a higher value (25.9 kcal.mol\(^{-1}\)); see Table 1. For monocoordinated phosphinine and phosphane complexes, the overall barrier follows the same trend as the alkene insertion and rotation barrier, the energy increasing by ~3 kcal.mol\(^{-1}\) on going from phosphinine to phosphane. This indicates that the alkene rotation process governs the overall barrier, and it is responsible for the observed higher activity in Rh-phosphinine system. Furthermore, it proves previous statements suggesting that for electron-withdrawing ligands the amount of back-donation is small, leading to facile rotation of the alkene moiety and therefore a low barrier for alkene insertion.\(^{7,21}\)

Following the previous arguments, for electron-donor bis(phosphane) systems one would expect higher rotational barriers, and consequently higher overall barriers. This is observed for the model system, in which both barriers increase by ~1 kcal.mol\(^{-1}\) with respect to the monophosphane system (Table 1, values in parenthesis). On the other hand, for bulky PPh\(_3\) phosphanes the alkene rotation barrier decreases upon bi-coordination, whereas the overall barrier is still higher.\(^{13}\)

Introducing the bulky groups increases the steric repulsion between the alkene substrate and the equatorial PR\(_3\) ligands in 3Pee (see Figure 3). This was reflected in longer Rh-alkene carbon distances for 3Pee than for 3Pe (2.287 vs. 2.272 Å on average); and in smaller alkene interaction energies (-24 and -30 kcal.mol\(^{-1}\)); and therefore, the more loosely bound alkene in 3Pee can rotate more easily. On the other hand, the formation of the alkene complex 3Pee is energetically more costly than the formation of 3Pe (15.1 vs. 11.5 kcal.mol\(^{-1}\)). The latter effect dominates, explaining the higher overall barrier. Thus, the increase of the overall barrier upon coordination of a second phosphane is not a direct consequence of the electronic properties of the ligands but of their steric properties. This is in line with our previous findings which indicated that the activity of Rh-catalyzed hydroformylation depends on both the basicity and the shape of the ligand;\(^{17}\) and with the proposed rate-controlling factors by Jensen and co-workers.\(^7\) Thus, we can propose for the design of new active ligands that reducing their steric hindrance and increasing their π-acidity will favour alkene coordination and rotation to reach the insertion TS, causing a reduction of the overall energy barrier and higher catalytic activities.

**Fig. 3** Molecular structures of the alkene complexes [HRh(CO)(L)(H,CCH\(_3\))] 3e1, 3Pe and 3Pee; transition states for alkene rotation (TS\(_{rot}\)) and insertion into the Rh-H bond (TS). Ligand hydrogen atoms are omitted for clarity. Distances in Å.

**Fig. 4** Representation of the interactions between the transition metal fragment and the phosphinine (a) and phosphane (b) ligands in terms of Dewar-Chatt-Duncanson model.

Rhodium-phosphorus and -alkene bond analysis: the origin of enhanced activity in π-acceptor ligands.

To evaluate the electronic properties of the P-ligands, we can use the IR stretching frequencies of the CO ligand (\(v_{CO}\)) in trans-L\(_2\)Rh(CO)Cl complex. They follow the order: P(OPh)_3 > P(OEt)_3 > 2,4,6-triphenylphosphinine > PPh\(_3\) > PEt\(_3\). A large \(v_{CO}\) value indicates π-acceptor properties due to a reduced π-backdonation from the metal center to the CO ligand, while a small value is indicative of strong σ-donation. Thus, these values show that phosphinines are poorer electron donors than phosphanes but richer than phosphites. Besides the overall electronic donating ability of the ligand, it is interesting to consider the decomposition into the individual contributions; σ-donation and π-backdonation, as done within the classical Dewar-Chatt-Duncanson model. To evaluate these contributions we used a modified version of energy decomposition analysis (EDA) based on an orbital deletion procedure, which allows the bonding
to be broken into physically meaningful components (see ESI and Figure S1 for details). Figure 4 schematically describes the orbital interactions in the analysis of the Rh-P bonding. We focused on the model systems because steric effects are put aside.

Table 2 Interaction energies (ΔEint) and σ-donation/π-backdonation evaluation for the Rh-phosphorus and Rh-alkene bonds.

| Rh-phosphorus | P(OH)3 | PC7H11 | PH3 | Rh-alkene | 3ε1H | 3PeH | 3PeH |
|---------------|--------|--------|-----|-----------|------|------|------|
| ΔEint         | -29.6  | -24.8  | -20.8 | -30.4     | -33.9 | -35.9 |
| ΔEint(L→Rh)   | -13.1  | -13.5  | -13.6 | -14.2     | -13.7 | -13.8 |
| ΔEint(Rh→L)   | -10.3  | -8.5   | -6.9  | -22.6     | -24.7 | -27.1 |
| ΔE1(a')       | -3.2   | -3.2   | -3.2  | -         | -     | -     |
| ΔE1(a'')      | -5.3   | -3.7   | -3.7  | -         | -     | -     |

Energies in kcal.mol⁻¹. For P(OH)₃, values taken from ref. 26 for the [HRh(CO)₂(PH₃)(P(OH)₃)] complex with the P(OH)₃ ligand in equatorial position. For 1ε₁H and 1PeH, calculations imposed C₂ symmetry to decompose the π-backdonation into in-plane (a') and out-of-plane (a'').

Table 2 collects the main results of the EDA bonding analysis for the Rh-P and –alkene ligands; and the full analysis is shown in Table S2 of ESI. We also show the values for the Rh-phosphite bond in [HRh(CO)₂(H₂P₃)](P(OH)₃) complex obtained at the same computational level in a previous work. For the P-ligands we observed that the interaction energies (ΔEint) follow the same trend of the CO stretching frequencies: P(OH)₃ > PC7H11 > PH₃ (29.6, -24.8 and -20.8 kcal.mol⁻¹, respectively). The values of σ donation [ΔEint(L→Rh)] are very similar for all the ligands (~13 kcal.mol⁻¹). On the other hand, the π backdonation decreases within the series (~10.3, -8.5, and -6.9 kcal.mol⁻¹ for P(OH)₃, PC7H11 and PH₃), explaining the higher overall charge donation from the ligand to the metal. The analysis of the Rh-alkene bond nicely correlates with rotational barriers and with the donor/acceptor properties on going from phosphinite to monophosphine and to bis(phosphine). Thus, this analysis quantifies the electronic factors and proves that π-acceptor property enhances the activity in Rh-catalyzed hydroformylation.

Interestingly, for phosphine π-backdonation occurs preferentially in the equatorial plane of the complex to the out-of-plane pₜ-type orbitals of phosphorus (ΔE₁(a'')) in Table 2). This directionality would lead to a more effective competition for metal electron density with the alkene, resulting in activities close to those of electron-poorer ligands such as phosphites. Accordingly, the computed overall barriers for HRh(CO)₂(PC₇H₁₁) and HRh(CO)₂(P(OH)₃) systems are similar, 19.3 and 20.6 kcal.mol⁻¹ respectively. Thus, although the overall π-acceptor property in phosphinines is somewhat reduced respect to phosphites, the efficient directionality of π-backdonation in phosphinines leads to a very active catalysts.

Factors governing the catalytic activity. Correlation with QSAR models for diphasphene ligands.

In this section, we analyzed whether the findings for monophosphines can be extrapolated to diphasphene ligands. As stated in the introduction, we had discussed the factors governing the activity for Rh-diphasphene catalysts during the development of a 3D-QSAR model. We had observed the correlation between high activity and low basicity for a given subset of structurally related complexes, while when comparing ligands of similar basicity the shape of the catalyst seems to determine the activity differences. Nevertheless, the use of alignment-independent 3D-descriptors made the chemical interpretation of the mathematical model difficult. Thus, here we re-examined the previous QSAR model using the Topological Maximum Cross Correlation (TMACC) method based on easy-to-interpret alignment-independent 2D-descriptors. During recent years, QSAR approaches have emerged as an alternative in the theoretical study of catalysis, including those based on alignment-independent descriptors.

The dataset defined from the experimental work in ref. 34 consist of 19 diphasphene ligands (set 1: L₁ – L₁₉, see Figure 5) and uses the percentage of conversion (%conv) as the response variable. Unfortunately, we could not expand the model including monophosphine ligands because there are no suitable data available that allows one to compare conversion values. We considered four atomic properties: Gasteiger partial charges represent electrostatics, Crippen-Wildman molar refractivity represent steric properties; and in addition, Crippen-Wildman logP parameters representing lipophilicity, and logS parameter representing solubility. Table 3 collects the statistical parameters of 10-fold cross-validation. All the defined individual descriptors yielded models that are close to the limit of prediction ability (0.55 < q² < 0.65), a model with q² < 0.5 being considered non-predictive. When we combined the four properties, an acceptable...
model was obtained, \( q^2 = 0.71 \). Closer inspection to the data revealed that ligand L13 has the largest difference between experimental and predicted conversion values, 61 \%, and consequently it could be classified as an outlier. The analysis of the chemical space showed that all ligands with medium or high activities have heteroatoms in their structures except L13. This means that the features of this type of structures are probably not well represented in the training data. When we set aside ligand L13 (set 2), the statistical parameters improved significantly \( (q^2 = 0.89 \) and \( r^2 = 0.92 \)). These findings indicate that the structure-activity relationship requires sophisticated descriptors that include electronic and steric factors.

**Table 3** Statistical parameters of 10-fold cross-validation for activity using different type of descriptors.

| Descriptor   | set 1 (19 ligands) | set 2 (18 ligands) |
|--------------|--------------------|--------------------|
|              | \( q^2 \) | \( r^2 \) | \( q^2 \) | \( r^2 \) |
| electrostatic| 0.59              | 0.64              | 0.72  | 0.78  |
| Steric       | 0.64              | 0.69              | 0.72  | 0.76  |
| lipophilicity| 0.58              | 0.78              | 0.76  | 0.91  |
| solubility   | 0.59              | 0.89              | 0.78  | 0.99  |
| combination of all | 0.71 | 0.76 | 0.89 | 0.92 |

*Pearson correlation coefficient \( (r^2) \) and cross-validated coefficient of determination \( (q^2) \).

Fig 6 Visualization of TMACC/PLS QSAR model for set 2 combining all the descriptors derived for ligands L1 (chiraphite), L5 (binaphos), L7 (diazophosphane) and L11 (kelliphite). In the interest of clarity, we do not show the lower contribution of other atoms. High positive contribution to activity in blue and high negative in red.

Besides, the TMACC descriptors provide a method for interpreting the results when combined with a linear regression method such as PLS. The predicted activity of the ligand can be partitioned among its constituent atoms. Figure 6 shows the representation of the interpretation for the diphosphane ligands L1 and L11, the phosphane-phosphate L5, and the diazophospholane L7 that have the highest activities (\( \% \text{conv} > 80 \)) of the dataset. TMACC method displays the atoms that most contribute positively to activity in blue and those that decrease the activity the most in red (yellow and orange colors represent intermediate positive and negative contributions). The oxygen atoms of the phosphate moiety and the hydrazine groups of the diazophospholane ligand are coloured blue (Figure 6), and hence have been identified as activity-increasing groups. On the other hand, disubstituted \( SP^2 \) carbons involving terminal alkyl groups and some other highly substituted carbons are colored in red (Figure 6), indicating activity-decreasing groups. These substituted carbons may be related with ligand bulkiness, which in turns is related with the reduction of catalyst activity. Figure 7 displays the interpretation the electrostatic to the steric-property models for ligand L5. The change in coloring for these properties indicates that the electrostatics has a more dramatic effect on activity of the ligand; the heteroatoms change from blue in the electrostatic-based model to uncolored in the steric-based model. A similar pattern was observed for ligands L1, L7 and L11, indicating that for them the high activity is also dominated by the electrostatic properties induced by the heteroatoms. To sum up, the interpretation of the QSAR model for diphosphane ligands can be related to our findings for monophosphane, indicating that similar rules govern the activity for both types of ligands.

**Conclusions**

We analyzed the behaviour of the \( \pi \)-acceptor phosphinine ligand in rhodium catalyzed hydroformylation of alkenes and compared it to phosphane-modified catalysts \([HRh(CO)\_3(PR_3)]\) and \([HRh(CO)\_2(PR_3)_2]\). The 2,4,6-PC\( \text{H}_2\text{Ph}_3\) phosphinine ligand coordinates preferentially at the equatorial site of the pentacoordinated rhodium complex with the heterocycle perpendicular to the equatorial plane of the complex, although the ligand freely rotates around the Rh-P bond.

We divided the overall energy barrier into several steps and/or contributions (alkene complex formation, alkene rotation and alkene insertion) and evaluated them. In the absence of steric effects (model systems), the overall barrier correlates with the barrier for alkene rotation. This proves that for \( \pi \)-acceptor ligands the amount of backdonation to the alkene is small, leading to its facile rotation, and consequently, to a higher activity. We also quantified the donor/acceptor interactions of the Rh-P bonds using a modified version of EDA analysis. Although the overall \( \pi \)-acceptor ability of phosphinines is lower than that of other ligands such as phosphites, the efficient directionality of their in-plane \( \pi \)-backdonation leads to very active catalysts. Introducing the steric effects of the ligands causes an increase of the energy required to form the alkene complex, and consequently an increase of the overall barrier. The factors governing the activity in Rh-monophosphane catalysts are closely related to those of Rh-diphosphane catalysts. This was confirmed by re-examining a previous QSAR model using the easy-to-interpret TMACC descriptors.

Thus, the design of active ligands in rhodium-catalyzed alkene hydroformylation should increase their \( \pi \)-acidity and reduce their
steric hindrance. Nevertheless, one should be aware that bulky monodentate ligands could favor the formation of \([\text{HRh(CO)}_2L]\) complexes which are more active than the \([\text{HRh(CO)}_3L]\) ones.

**Experimental section**

**Computational Details. DFT calculations**

The DFT calculations were carried out using the Amsterdam density functional program (ADFv2008).\(^{38}\) The electronic configurations were described by a triple-\(\xi\) plus polarization Slater-type basis set, as included in ADF package. The 1s-3d electrons for Rh, the 1s electrons for C and O, and the 1s-2p electrons for P were treated as frozen cores. We applied scalar relativistic corrections to them via the zeroth-order regular approximation (ZORA) with the core potentials generated using the DIRAC program.\(^{36}\) We used the GGA functional BP86.\(^{37}\) Full geometry optimization without any symmetry constraints were performed, unless otherwise stated. The transition states were characterized by a single imaginary frequency and the normal mode, which corresponds to the expected reaction path. In some structures, we found a residual imaginary frequency related with the loose torsion angles of bulky substituents. We are aware that most of the popular DFT methods such as BP86, B3LYP or PBEn are unable to describe noncovalent interactions in their attractive regime.\(^{41}\) In a recent study of Rh-catalyzed hydroformylation,\(^{26}\) we tested the M06 class of functional recommended to study noncovalent interactions,\(^{42}\) and B97D functional including explicit dispersion corrections.\(^{43}\) Both functional types gave qualitatively the same results as BP86 one for these systems, in which, the ligand-substrate interactions were dominated by repulsive interactions.\(^{46}\) We expect the same results here. Indeed, the interaction between phenylphosphino moieties and aliphatic alkenes were proved to be repulsive in nature for alkene insertion into the Rh-H bond by means of QM/MM calculations.\(^{29}\)

To analyze the nature of Rh-phosphorus and -alkene bonds, we employed the Energy Decomposition Analysis (EDA) method;\(^{44}\) and a modified version based on orbital deletion that allows separating the \(\sigma\) and the \(\pi\) interactions in a physically meaningful manner.\(^{47}\) The details are provided in the Supporting Information.

**QSAR TMACC-based modelling**

These descriptors are generated using atomic properties (electrostatics, solubility, steric effect and lipophilicity) determined by molecular topology. The source code for computing TMACC descriptors is available for download from http://comp.chem.nottingham.ac.uk/download/TMACC. The electrostatic properties are represented by the Gasteiger partial charge,\(^{35}\) which is calculated using the method of partial equalization of orbital electronegativity. This procedure calculates atomic charges in \(\sigma\)-bonded and non-conjugated \(\pi\)-system using only the topology of the catalysts. The Crippen-Wildman molar refractivity (MR) is used as a measure of the steric effects that it is determined through the classification of atoms based on neighbour atoms.\(^{36}\) The Crippen-Wildman partition coefficients (logP) are assigned to each atom as a measure of atomic lipophilicity, determined in the same way as Crippen-Wildman molar refractivity (MR).\(^{46}\) The solubility properties are described by logS parameters, representing solubility and solvation phenomena.\(^{57}\) We scaled each contribution by the largest absolute value, so that the positive and negative values took maximum values of +1 and -1.

The TMACC autocorrelation descriptor \((x_{ac}(p,d))\) is

\[
x_{ac}(p,d) = \sum p_i p_j
d\]

where \(p\) is one of the properties and \(d\) is the topological distance between atoms \(i\) and \(j\), normally the shortest number of bonds between atoms. The sum is over all atom pairs that are separated by distance \(d\). We treat each atomic property that can take positive and negative values as separate properties. The molar refractivity is the exception that only takes positive values. Like for GRIND descriptor, we keep only the maximum value calculated for any given distance. For the purposes of interpretation (see above), for each descriptor, we recorded which atoms contribute to the maximum product. In the event of more than one pair having the same value, we record all pairs. The maximum distance was as large as the largest distance in the molecule. The minimum distance was zero, that is, we allowed \(i = j\).

We employed the Partial Least Squares (PLS) Regression\(^{45}\) as the multivariate regression technique. Ten-fold cross validation was used for model building and evaluation. Different statistical parameters facilitated evaluation of the predictive ability of models during the fitting and test stages, namely; the Pearson correlation coefficient \((r^2)\) and the cross-validated correlation coefficient \((q^2)\), see ESL.\(^{56}\) The TMACC descriptor string for each molecule comprises several hundred elements. Clearly, it is not practicable to present the resultant QSAR equation, in which the latent variables in the PLS regression are linear combinations of the descriptors. A more qualitative summary of the model is provided via the graphical interpretations in Figures 6 and 7, in which the PLS has effectively been reversed to ascribe contributions to activity to individual atoms.

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**Notes and references**

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\(^{d}\) Electronic Supplementary Information (ESI) available: Tables containing relative energy values for coordination isomers of mono- and bi-phosphate complexes, and the detailed results and description of EDA bond analysis. The \(xyz\) coordinates for the most relevant computed structures.

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This property requires 3D descriptors that are able to account for catalysts shape.

Note that for strongly electron-withdrawing phosphites, the rate-determining step can switch to hydrogenolysis (see Ref. 4 and 7). Nevertheless, the comparison could be useful to understand the effects of $\pi$-acceptor ligands. Note also that the simplified model ligand has been compared with the real phosphine-phosphate binaphos ligand in the previous work obtaining consistent results for electronic features (see ref. 26).

In the previous contribution we used 3D descriptors, which are more time-consuming and more difficult to interpret, because our main focus was the modeling of enantioselectivity. This property requires 3D descriptors that are able to account for catalysts shape.

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