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Pd-$\eta^3$-C$_6$H$_9$ Complexes of the Trost Modular Ligand: High Nuclearity Columnar Aggregation Controlled by Concentration, Solvent and Counterion.

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The Trost modular ligand (TML) series induces high levels of asymmetric induction in an extraordinarily wide range of reactions involving palladium $\pi$-allyl intermediates. Prior mechanistic investigations into reactions involving simple Pd-$\pi$-cyclohexenyl intermediates has focussed on the monomeric 13-membered ring formed via P,P-chelation of the ligand to Pd. However, ring-opening oligomerisation was recognised as providing a major pool of polynuclear palladium species that, through low or even opposite asymmetric induction to the mononuclear species, are responsible for a reduction in selectivity under non-optimised conditions. Herein we describe an investigation by NMR, molecular mechanics, molecular dynamics, and small-angle neutron scattering, of a Pd-$\pi$-cyclohexenyl cation bearing the 1,2-diaminocyclohexane TML ligand (2). Using both nondeuterated and perdeuterated (D$_2$) isotopologues of the resulting complexes ([I]$^+$), we show that a two-stage oligomerisation-aggregation process forms large cylindrical particles of high nuclearity (up to 56 Pd centres).

**Introduction**

The Trost modular ligand (TML) series has been applied to an extraordinarily wide range of Tsuji-Trost (allylic alkylation) reactions. Under carefully optimised conditions, these ligands frequently provide high enantioselectivity in reactions that have proven challenging with other chiral ligands, particularly those involving cyclic allylic substrates. These features have led to broad use of the TML in the synthesis of natural products, as well as industrial application for the construction of high enantiopurity chiral building blocks. However, reactions involving the TML frequently exhibit, memory effects and a high sensitivity of the enantioselectivity to reaction temperature, catalyst concentration, solvent and nucleophile counter-ion.

Our previous mechanistic studies of this system focussed on the monomeric cationic complex [I]$^+$, in which the 1,2-diaminocyclohexane derived TML ligand (2) chelates the Pd($\eta^3$-C$_6$H$_9$) moiety. This was identified as an intermediate capable of leading to high asymmetric induction on attack of, for example, a malonate anion nucleophile, Scheme 1. Detailed NMR studies, facilitated by isotopic labelling, in conjunction with MM-DFT simulations, led to a model in which the amide units in the catalyst facilitate enantioselective ligand-accelerated catalysis.

![Scheme 1. Asymmetric allylic alkylation of racemic 2-cyclohexenyl acetate; 2 = 1,2-diaminocyclohexane TML ligand, THAB = tetraxethylammonium bromide.](image-url)

For the ligand-accelerated catalysis to function efficiently, cation [I]$^+$ requires a degree of flexibility. This flexibility is provided by the 13-membered chelate ring, but at a cost: complex [I]$^+$ can readily undergo ring-opening oligomerisation to generate polynuclear species ([I]$^+$)$_n$, Scheme 2. Competing nucleophilic attack of the oligomer, rather than the monomer...
[1] is, in part, responsible for a reduction in overall enantioselectivity under non-optimised conditions. \[1,14\]

\[
\begin{align*}
\text{MONOMER [1]}^+ & \quad \leftrightarrow \quad \text{OLIGOMER ([1])}_n \\
\end{align*}
\]

Scheme 2. Oligomerisation of [1]' erodes enantioselectivity during asymmetric Pd-catalysed allylic alkylation mediated by 2 (as Scheme 1). Nu = nucleophile.

To date, the structure and driving forces for formation of the polynuclear oligomeric species has not been studied in detail. Herein we describe an investigation of the oligomerisation of D₈ and D₄₇ isotopologues of [1]', employing NMR spectroscopy, molecular mechanics (MM), molecular dynamics (MD), and small-angle neutron scattering (SANS). The data obtained indicate that the impact of a first-stage depletion of the monomeric species [1]' from the catalyst pool, via cyclic oligimerisation, is amplified by a second-stage process involving columnar aggregation of the oligomers, leading to species with very high nuclearity (up to 56 Pd centres). The effects of solvent, ligand enantiopurity and counter-ion on the degree of aggregation are explored in detail, and it is concluded that a relatively small and restricted set of conditions facilitate dissolution of the complexes in a low aggregation state, consistent with the extensive optimisation frequently required for these catalyst systems.

Results and Discussion

Preliminary NMR Studies and Synthesis of [D₂₋₁][B(C₆F₅)₃].

Despite extensive efforts, we have been unable to crystallise any Pd(η⁵-C₅H₅) complexes of 2, in either oligomeric or monomeric forms. Indeed, to date, the only X-ray crystal structures of Pd-allyl complexes of Trost ligand 2 are η²-C₅H₅ complexes with triflate counter anions: one a racemic tetranuclear cyclo-oligomer, the other an acyclic dinuclear bis-P,O-chelate.

The extent of solution-phase oligomerisation of cationic complexes of type [1]' can be conveniently determined \[³¹P \{¹H\} NMR spectroscopy. \[15\] Analysis of \([\{R,R\}]-[1][BAr₅]\) complexes in CH₂Cl₂, where Ar = C₆Cl₅, 3,5-(CF₃)₂C₆H₃, or C₆F₅ (‘BAr₅’, Figure 1), indicates a maximum monomer concentration ([1]') of about 4 mM. In THF the monomer maximum is lower (approx. 1.6 mM) and decreases as [Pd]₀ is raised above 10 mM. With smaller, less charge-diffuse, counter-ions such as chloride or triflate, the monomer maxima are lower still. We were unable to fit simple analytical solutions for monomer-oligomer distributions to any of the \[³¹P NMR data, indicative that physicochemical effects dominate over simple solution-phase equilibria, even at low [Pd]₀.

\[\text{Figure 1. Speciation of} \quad \text{[}(R,R)\text{-}[BAr₅]) \quad \text{([B(C₆F₅)₃])} \quad \text{determined by} \quad \text{³¹P \{¹H\} NMR (CDCl₃, 25 °C). Solid lines through data are solely a guide to the eye.}\]

We thus elected to study the oligomeric species by SANS – a technique that can be used for characterising the shape and dimensions of large scale molecular aggregates and colloids in solution. \[16\] We began with \([\{R,R\}]-[1][BAr₅]\) and, to aid the studies, also synthesised the perdeuterated enantiomeric complex \([\{S,S\}-[D₂₋₁]-[1][BAr₅]\). Not only does this facilitate SANS in a non-deuterated solvent, thus providing greater neutron scattering contrast, it also allows pseudo racemic and pseudo scalemic mixtures to be prepared by mixing \([\{S,S\}-[D₂₋₁]-[1][BAr₅]\) with \([\{R,R\}]-[1][BAr₅]\). The perdeuterated complex was synthesised from benzoic acid ([D₉]-3), chlorobenzene ([D₄]-4) and cyclohexene ([D₆]-5), Scheme 3. A major hurdle was the ortho-metallation of ester \([D₃]-6\) with (TMP)₂Mg:LiCl \[38\] which proceeded with an unexpectedly large net kinetic isotope effect (k₂/k₁ ≈ 30). \[16\] This required an excess base to be employed, and interfered with a planned direct phosphination of the metallated intermediate. Instead, the intermediate was trapped with I₂. The iodide \([D₃]-7\) was then converted to a more conventional Grignard reagent \[39\] before reaction with chlorophosphine \([D_{10}]-8\) \[40\] to give phosphine \([D_{14}]-9\), and thus acid \([D_{14}]-10\).

Cyclohexene \([D₆]-5\) was epoxidised, to give \([D_{10}]-11\), and this ring-opened \[12\] to give aminoalcohol \([D_{12}]-12\). Aziridine \([D₁₀]-13\), obtained via Mitsunobu conditions, was converted to azide \([D_{10}]-14\) \[41\]. After diastereoisomer separation, hydrogenolysis \[41\] gave (S,S)-diamine \([D_{10}]-15\) which was coupled with acid \([D_{14}]-10\) to afford Trost ligand \([D₉]-2\). The chloro-dimer \([D₉]-16\), prepared \[42\] from \([D_{10}]-5\), was converted to cationic complex \([D₄]-17\), and then reacted with \([D_{10}]-2\) to generate \([\{S,S\}-[D₂₋₁]-[1][BAr₅]\) in good yield.
**SANS Analysis of Aggregation of [1][BAr₄] in THF**

We began SANS analysis of enantipure [(R,R)-1][BAr₄] in THF-D₈ at 25 °C, with [Pd]₀ concentrations at and above the oligomerisation threshold (1.6 to 64 mM). The combined data sets, presented here as plots of scattered intensity (I(Q), y-axis) versus neutron momentum transfer (Q, x-axis) were analysed with standard mathematical models corresponding to various simple shapes. Figure 2 This clearly identified the particles as cylindrical or rod-like, and Guinier analyses (see supporting information, Figure S69) established the radii (8-9 Å) and length (150-200 Å) as invariant across the range of [Pd]₀ explored. In other words, the number of particles changes in response to [Pd]₀ but not their average dimensions, clearly indicative of a set of factors that tightly control the particle scale.

![Figure 2](image)

We have previously used ³¹P NMR to analyse the constitution of the solution-phase (i.e. lower-order) oligomers generated from various complexes of type [1][BAr₄] in CD₂Cl₂. Using PPOCS in combination with pairs of isotopically-differentiated ligands ([D₆]-2), we were able to determine that the oligomers are: i) non-chelated species, i.e. each of the ligands (2) in the oligomer are coordinated to two different Pd centres; ii) present in predominantly homochiral form, i.e. [(R,R)-1] and [(S,S)-1] oligomerise independently, and iii) there no free (i.e. not Pd-coordinated) P-centres in the ligand (2). Although a cyclic oligomer structure (Scheme 2) is fully consistent with these features, we were unable to determine the number (n) of ring-opened monomer units incorporated within the cyclo-oligomer ([1][BAr₄]).

As [Pd]₀ in THF or CH₂Cl₂ solutions of complexes of type [1][BAr₄] is increased, the ³¹P NMR bandshape of the signals...
arising from the cyclo-oligomer do not change in appearance, but the samples do become increasingly turbid. This behaviour suggests that in response to an increase in [Pd]_{iso}, cyclo-oligomers ([I][BARf]), do not incorporate more monomer (n), but instead aggregate to form large particles, ([I][BARf]), containing ‘m’ cyclo-oligomers. It is these high nucleation particles that are detected by SANS.

In racemic or scalemic samples of ([I][BARf]), the homochiral cyclo-oligomers, ([I][BARf]), could aggregate in three general forms: discrete homochiral, ordered heterochiral (e.g. alternating or co-block), or statistically distributed. SANS data of mixtures of [(R,R)-I][BARf] and [(S,S)-I][BARf] representing enantiopure, scalemic, and racemic samples, was uniform across the series, within experimental error, Figure 3.

The absence of a change in particle number, shape or size, suggests a statistical distribution.

SANS data of the pseudoracemate [(S,S)-[D$_{25}$]-I][BARf] + [(R,R)-[D$_{25}$]-I][BARf]) in D$_o$-THF, and in D$_{25}$-THF, Figure 4, and enantioselectively pure [(S,S)-[D$_{25}$]-I][BARf] in D$_{25}$-THF, show that the fully and partially deuterated systems retain the concentration-independent, cylindrical shape in the aggregates. The main difference however is in the dimensions: [D$_{25}$]-[I][BARf] forms shorter (130 Å), slightly wider (10 Å radius) cylinders than [D$_o$]-[I][BARf]. The pseudo racemic mixture measures as an average of its precursors (9-10 Å radius, 150 Å length), again consistent with a statistical distribution of cyclo-oligomers in the aggregate.

**Molecular Mechanics (MM) and Dynamics (MD)**

Computational modelling was employed to probe the factors that control the aggregation phenomena, and to estimate the size (n) and number of cyclo-oligomers in the particle (m). The number of atoms in the aggregates (>6,000, *vide infra*) means that density functional theory (DFT) calculations of their structures would demand currently unattainable computational resources and inordinate simulation times. On the other hand, molecular mechanics (MM) can provide a good approximation, in just a small fraction of the computational time required by DFT. We began by confirming that structure of the 84-atom monomeric cationic P,P-chelate [I]$^+$, optimised at MM3 theory level and dielectric constant ε = 9.0, was almost identical to that obtained using DFT (B3LYP-D3) in a polarisable continuum model for dichloromethane. Further calculations involving ion-pairs, oligomers and aggregates were then performed using MM3 to provide analysis of the energies involved, albeit at a course-grained level of detail.

Comparison of the MM3 optimised energies for monomeric P,P-chelate [I][BARF] with a series of homochiral cyclo-oligomers, ([I][BARf]), normalised by the number of Pd atoms (n, the x-axis in Figure 5) confirmed cyclo-oligomerisation to be exergonic. The cyclic dimer (n = 2) still suffers from ring strain, and a more substantial stability is afforded by trimerisation (n = 3) then tetramerisation (n = 4, Figure 6). Further increase in oligomer ring size (n = 5, 6, 8) yields a modest reduction in the system energy and generates species with significantly higher radii than the 8-9 Å cylinder radius detected by SANS.

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**Figure 3.** Fitted SANS profiles of 32 mM enantio-pure, racemic and scalemic samples of [I][BARf] in THF-D$_o$, 25 °C.

**Figure 4.** SANS data for pseudo racemate ([S,S]-[D$_{25}$]-I)[BARf] + [(R,R)-[D$_{25}$]-I][BARf]) in THF-D$_o$ at 25 °C, at [Pd]$_{iso}$ ranging from 13 to 51 mM.

**Figure 5.** MM3 energies of cyclo-oligomeric complexes ([I][BARf]), $\varepsilon$ = 9.0.
Aggregation of cyclic tetramers ([1][BAR$_F$])$_4$ was then probed by MM3 as a mechanism for generation of cylindrical particles. Positively charged rod-like structures with dissociated or removed anions were estimated by MM3 to be very much higher in energy.

Although BAR$_F$ is considered a weakly coordinating anion, the charge delocalisation over its surface reduces repulsive interactions with other BAR$_F$ anions as well as making it significantly lipophilic. Indeed, the calculations indicated a favourable interleaving of the anions in sandwich layers between cationic cyclo-oligomers. The estimated formation energies, (ΔE, Figure 7) of such species {([1][BAR$_F$])$_4$}$_m$ as a function of ‘m’ indicated that columnar aggregates are readily attainable, with the growing entropic cost (7ΔS) placing limits on the aggregate length.

These conclusions were further probed by molecular-dynamics (MD) simulations in which the MM3-minimised structures {([1][BAR$_F$])$_4$}$_m$ were computationally excited over a short period (300 ps) to test the structural integrity of the aggregate as a function of ‘m’. In the low dielectric constant medium used for the model, most systems (m = 4 to 16) did not undergo any significant changes in their tertiary structure at 300 K, over the full 300 ps simulation time. As the energy input was increased the aggregate models exhibited varying degrees of structural deformation and at 600-700 K rapidly fragmented. The most significant observations were made in the intermediate region (500-550 K) where aggregates with m = 10-14 (e.g. Figure 8, m = 12) retained a cylinder shape, albeit mildly distorted, whereas higher or lower order aggregates significantly deformed, in some cases losing one or more BAR$_F$ anions. The average dimensions of the MM3 aggregates with m = 10-14 (radius 8-9 Å and length 150-200 Å) are fully consistent with the dimensions determined by SANS.

The Effect of Counter-ion and Solvent on Shape and Extent of Aggregation

The effect of solvent on aggregation was probed by MM, $^{31}$P[1H] NMR (Figure 9) and SANS, also comparing [1][BAR$_F$] with [1][OTf] to explore the impact of counter-ion. The aggregation mode for [1][BAR$_F$] determined by MM3, e.g. Figure 8, involves multiple close-range electrostatic interactions that reduce the overall system energy. It is thus not surprising that the solvent dielectric constant, $\varepsilon$, was found to modulate aggregation, and thus also solubility: precipitation being the consequence of complete aggregation. As indicated in Figure 9, both [1][OTf] and [1][BAR$_F$] readily oligomerise and in all of the solvents that were explored, becoming essentially insoluble at the extremes of $\varepsilon$, e.g. in alkanes, most ethers, chloroform, aromatic hydrocarbons, and at the opposite end of the scale, in water. The lipophilicity and charge density of the anion also affects the solubility: [1][BAR$_F$] readily dissolves in...
THF, while [I][OTf] does not, and at the opposite end of the ε scale, [I][OTf] is soluble in aqueous-organic mixtures, whereas [I][BAR$_F$] is not.

SANS was employed to explore how the macromolecular composition of aggregates {[(I)[X]]$_m$} is affected by solvation. Although [I][BAR$_F$] is not soluble in organic-aqueous mixtures, SANS data was attainable in polar aprotic solvents (e.g. MeCN, $\varepsilon_r = 37.5$; and DMSO, $\varepsilon_r = 47$). This confirmed that cylindrical aggregates were still formed, but were significantly shorter than those in THF, Figure 10. Medium length cylinders were detected in a 50:50 mixture of THF and acetonitrile, consistent with the intermediate solvent polarity ($\varepsilon_r \approx 23$). In all cases, the cylinders were of radius 8-10 Å, strongly suggesting the prevalence of the tetranuclear cyclo-oligomer building blocks, with the solvent modulating the aggregation number ‘m’: {[(I)[BAR$_F$]]$_m$}.

The [I][OTf] aggregates behaved differently. Although, cylinders of radius 8-10 Å were again detected in all cases, indicative of {[(I)[OTf]]$_m$}, the flexibility, lengths and charge distribution in the particles were very different to those formed from [I][BAR$_F$]. In CD$_2$Cl$_2$, [I][OTf] forms long cylinders (up to 160 Å; Figure 11) that are semi-flexible, a phenomenon that can be attributed to the small and interactive triflate anion being less able to rigidify the structures than the larger and more lipophilic BAR$_F$ anion.

Figure 9. Oligomerisation of [I][BAR$_F$] and [I][OTf] at [Pd]$_{tot}$ = 15 mM in various solvents, as determined by $^3$P NMR; overlaid areas in green (OTf) and blue (BAR$_F$) indicate regions in which the complexes are insoluble.

Figure 10. SANS data for [I][BAR$_F$] in various solvents at 25 °C.

Figure 11. SANS for [I][OTf] in CD$_2$Cl$_2$ at 25 °C.

The anion effect became even more pronounced in media of higher dielectric constant. The SANS data indicated the presence of charged particles in acetonitrile-based solvent mixtures ($\varepsilon_r = 47-58$), Figure 12, indicating solvation-induced ion-pair separation of the triflate from the cationic Pd(II) oligomeric core. The increased cationic repulsion between the cyclo-oligomers thus results in much shorter cylinders, just 30 Å in length, with misleadingly simple $^3$P NMR spectra.

Similar conclusions were drawn from MD simulations with the medium set at $\varepsilon_r = 35$: only the shortest aggregates {[(I)[OTf]]$_{2-4}$} were structurally stable at elevated energies (500 K; 300 ps).

Finally, to probe the relevance of the higher aggregates to asymmetric alkylation (Scheme 1) SANS data was acquired on reaction mixtures in which [I][BAR$_F$] was employed as a pre-
catalyst (10 mol%) for addition of tetrabutylammonium dimethylmalonate to cyclohexenylacetate in THF. While the effects of substrate background scattering, varying acquisition times and shorter Q-range slightly affected the accuracy of data, it remained clear that the dominant structures in solution were large cylinders, for the whole duration of the catalytic process. This result is consistent with previous conclusions that in THF the catalytic turnover proceeds via a small pool of highly-active monomeric catalyst species, in competition with cyclo-
oligomers and aggregates.

Conclusions

Since the initial report that [1]+ readily oligomerises and that this is a largely undesirable property of an otherwise highly efficient catalyst, e.g. Scheme 1, there has been limited understanding of the oligomer structure. We have now identified, through NMR, SANS, and MM/MD simulations of [D₄₋][1][BARF] (n = 0, 47), that monomeric cations [1]+ undergo chelate-opening to form a tetranuclear cyclo-oligomer; this being thermodynamically favoured on higher and lower nuclearity species. A second-stage process involving columnar type interactions between the cyclo-oligomers then forms polynucler aggregates. For [1][BARF] in THF, these comprise 10-14 alternating layers of cyclo-oligomer and interleaved BARF anions, containing up to 56 Pd(n-C₆H₄) centres, ligands (2) and anions, and generate cylindrical particles of radius 8-9 Å and length 150-200 Å.

Figure 13. Summary of upper-range aggregate cylinder lengths of [1][BARF] and [1][OTf] samples, as determined by SANS experiments, in selected solvents.

The identity of the counter-anion has a pronounced effect on the proportion of oligomer generated from the monomer. Bulky, weakly-coordinating anions reduce the extent of oligomerisation, particularly in low polarity solvents that cannot effectively stabilise charged particles. Here the role of the bulky and relatively lipophilic anions is to solvate the monomer [1]+. Smaller harder, less lipophilic anions are less able to solvate the monomer, and have the indirect effect of shifting the equilibrium towards the oligomer; an undesirable feature for catalysis. The diminutive size of the anion also results in greater flexibility of the resulting columnar aggregates, which are more ionic in nature, reducing their solubility in less polar solvents.

Overall, although the solvent polarity, counter-anion (X), and net concentration ([Pd₉]) all contribute to determining the degree of oligomerisation and aggregation (Figure 13) of monomer [1][X], the solvent perhaps offers the greatest degree of scope for optimisation under the conditions of catalysis. In this regard, CH₂Cl₂ is favourable: solutions can be virtually free of oligomer at ambient temperature, provided [Pd₉] ≤ 4 mM. Intriguingly, SANS studies of non-ionic surfactants have revealed specific solvent combinations that can lead to “dead zones” where aggregation is suppressed, even for concentrated monomer solutions. If such “dead zone” solvent combinations can be found for complexes of type [1][X], this may be highly advantageous for improving catalytic productivity whilst maintaining selectivity.

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Notes and references


14. Stoichiometric reaction of [(RR)-1][BAR], (Ar = 3,5-(CF3)2C6H3) with tetrabutyllammonium dimethylmalonate in THF at 21 °C: [Pd]ext = 1 mM affords allylation product in 98% ee, whereas product of 6% ee is obtained when [Pd]ext = 27 mM.

15. Around 100 attempts were made to crystallise the following complexes [(RR)- and rac [I][X]], X = BARn, Br(3,5-(CF3)2C6H3)2n, [Al(O(OC)(CF3))3] and OTf using diffusion, evaporation, cooling, seeding, co-crystallisation, and other methods, from a variety of binary and ternary solvent mixtures, including but not limited to chloroform, CH2Cl2 / pentane, 1,2-dichloroethane / hexane, THF / heptane, 1,2-difluorobenzene / 1,3-difluorobenzene, tetrachloroethane / hexafluorobenzene, MeCN / mesitylene, aqueous MeCN / MTBE, water, diethyl ether, etc. without success.


20. Perfluoro tetraphenylborate [Br(C6F5)4] (“BARn”) was chosen to avoid SANS data being complicated by neutron scattering from protons or deuterons in the counter-ion, as this could then be used without perturbation in the perdeuterated complex.


22. The large KIE, estimated as kH/kD = 30, by comparison of approximate rates of metallation of the non-deuterated substrate, may arise from multiple primary effects, involving the TMP-amide, and in situ generated [D]-TMP, or from tunnelling.

23. To avoid the formation of by-products, it was found that it was essential to slowly add D10-chlorodiphenylphosphine [D10-8] to the Grignard drop-wise at -78 °C.


30. In contrast to the report of Singh, the hydrogenolysis with Pd(OH)2/C under acidic conditions in methanol failed to cleave the α-methylbenzyl auxiliary. Instead, after reduction of the azide, the hydrogen source was changed to ammonium formate: a) WO 2005/092840 PCT/IB2005/000619; b) T. Bieg, W. Szeja, Synthesis 1985, 76-77.

31. The dinner was prepared using the literature procedure (see: a) B. M. Trost, P. E. Strege, L. Weber, T. J. Fullerton, T. J. Dietsche, J. Am. Chem. Soc. 1978, 100, 3407-3415; b) S. Imaiuzzi, T. Matsuhashi, Y. Senda, J. Organomet. Chem. 1985, 280, 441-448). However we identified degradation of the crude organometallic complex during the early work-up stages. Constant exposure of the crude material to air, avoiding vacuum operations until the complex has been purified, significantly increases the yield of 17.

32. SANS experiments were conducted at the Institut Laue-Langevin (ILL) in Grenoble, France. Data was acquired in two detector movements on separate instrument: high-Q data on D16, and low-Q data on D22.

33. Very similar results were obtained on SANS2D at ISIS, UK using a different batch of complex, indicative of the reproducibility of the data.

34. Measurements were also conducted at higher and lower temperatures. Consistent with analysis by 1H NMR spectroscopy, aggregation is reversible, and favoured by lower temperatures.

35. At higher concentrations the radius shrunk slightly, possibly due to the associated increase in the ionic strength of the medium.

36. It should be noted that most of the concentrations employed for the SANS experiments were, by necessity, well above the oligomerisation and aggregation threshold. Changes in particle number due to differential selectivity between homochiral and heterochiral aggregation of cyclo-oligomers may be small, and thus hard to distinguish.

37. The resolution of the SANS instrument was insufficient to completely rule out the possibility that the scattering profile is the composite of two curves arising from segregated (homochiral) cylinders.


41. Comparison of the MM3 energy of the tetramer with a pre-oriented assembly of four P,P-chelates {4[1][BAR])}, or the energy of two non-covalently associated tetramers {([1][BAR])2} with a single cyclic octamer (Figure 5) again indicates that oligomerisation to form the tetramer (Figure 6) is most favourable. The dimensions of the MM3-minimised tetramer, are similar to that of the Pd ligand core of a cyclic tetranuclear complex ([g(-7-C,H4-Pd(2)]2)[OTf]4 determined by single crystal X-ray diffraction.

43. A variety of other anion arrangements explored by MM3 modelling did not reveal any plausible alternative modes of aggregation, that would be compatible with the SANS data.
44. The impact of solvent dielectric constant on ion-pair separation can be predicted by the Bjerrum length equation: \( \lambda_B = \frac{e^2}{4\pi \varepsilon_0 \varepsilon_r k_B T} \) which indicates that the ion separation distance, \( \lambda_B \), is inversely proportional to the dielectric constant of the media in which the solute is dissolved. The equation presumes point charges, however the general principles may still apply to more complex systems such as [1][X].
45. These systems present an apparently simple pair of doublets (\( J = 38 \) Hz), almost identical in appearance, but not chemical shift, to the pair of doublets \( \left( \frac{3J_{pp}}{2} \right) \) in the monomeric chelate system. However, it is very evident from the SANS data that the solutions are dominated by oligomers \( \left( [1][OTf]_n \right) \), and from careful inspection of the \( ^{31}P \) NMR spectrum that the monomer is indeed also present, but only in very low proportions (\( \leq 4\% \)).
47. All of the BAr\( _4 \) anions tested, as well as Al(OR\( _4 \))\( _4 \) (“Krossing anions”: Krossing, I. Chem. Eur. J. 2001, 7, 490-502), were effective in this regard.