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# Charge effect on the formation of polyoxometalate-based supramolecular polygons driven by metal coordination

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Abstract: The metal driven self-assembly of a Keggin-based hybrid bearing two remote pyridine units was investigated. The resulting supramolecular species were identified by combination of 2D diffusion NMR spectroscopy (DOSY) and electrospray ionization mass spectrometry (ESI-MS) as a mixture of molecular triangles and squares. This behaviour is different to that of the structural analogue Dawson-based hybrid displaying a higher charge, which only led to the formation of molecular triangles. This study highlights the decisive effect of the charge of the POMs in their self-assembly processes that disfavours the formation of large assemblies. An isothermal titration calorimetry (ITC) experiment confirmed the stronger binding in the case of the Keggin hybrids. A correlation between the diffusion coefficient D and the molecular mass M of the POM-based building block and its coordination oligomers was also observed. We show that the diffusion coefficient of these compounds is mainly determined by their occupied volume rather than by their shape.

### Introduction

Self-assembly is an attractive approach to elaborate complex nanostructured molecular architectures. Starting from molecular-sized building blocks of known topologies, 0D to 3D supramolecular systems can be built in a straightforward manner. Among the various synthetic approaches, metal-driven self-assembly is a powerful route for controlling the topology of the desired molecular architectures by using suitable metallic ions and ligands.<sup>1-6</sup> Polyoxometalates (POMs) are anionic oxo-clusters of high oxidation-state transition metal

ions whose remarkable electronic properties are suitable for various technological applications.<sup>7,8</sup> Owing to their nanoscale size, their structural complexity and their sensitivity to various types of interactions,<sup>9</sup> POMs are attractive molecular building-blocks for the elaboration of supramolecular architectures with emergent<sup>10-13</sup> or synergistic properties such as molecular separation,<sup>14</sup> self-healing,<sup>15</sup> luminescence,<sup>16, 17</sup> or crystal morphogenesis.<sup>18</sup> Their functionalization into covalent organic-inorganic hybrids is a first step toward their integration into more complex assemblies. In this context, we recently described the selfassembly of covalent POM-based hybrids through metal coordination.<sup>19, 20</sup> The hybrids developed in these studies were organosilyl functionalized Dawson-type POMs bearing remote binding sites (pyridine or terpyridine units). These hybrids display two organic arms approximately perpendicularly oriented,<sup>21-23</sup> a topology that is well suited for elaborating molecular polygons by coordination to a metal linker. Furthermore, when using a cationic linker, the discrete supramolecular assembly could further self-assemble through intermolecular electrostatic interactions into an unprecedented multiscale organized system according to the solvent composition.<sup>19</sup> The discrete molecular polygons were identified as supramolecular triangles, while molecular squares were expected owing to the ca. 90° angle between the pendant arms of the starting hybrid. This suggests that the charge of the POM plays a major role in the formation of the cyclic assembly. Indeed, the formation of highly charged supramolecular species has potentially an important entropic cost due to the confinement of the associated counter-ions.

Organo-silyl derivatives of monovacant POMs can be also prepared in the Keggin series. Interestingly these Keggin (*i.e.*  $[PW_{11}O_{39}{O(SiR)_2}]^{3-}$ ) and Dawson (*i.e.*  $[P_2W_{17}O_{61}{O(SiR)_2}]^{6-}$ )-based hybrids display a very similar topology with approximately orthogonal organic arms but distinct charge.<sup>24</sup> This drove us to evaluate the effect of the charge of the molecular building block on the structure of the resulting self-assembly. We herein report the metal-directed self-assembly of a Keggin-based molecular building block. We show that the resulting supramolecular structures are a mixture of molecular triangles and squares. This study outlines the major role of the charge of the POMs in their self-assembly.

### **Results and Discussion**

#### Formation of mixtures of molecular triangles and squares

The Keggin-type pyridyl-terminated POM-based building block  $TBA_3[PW_{11}O_{39}{O(Si-C_{13}H_8N)_2}]$ , denoted as **K**<sub>si</sub>[**pyr**]<sup>25</sup> (Figure 1) was synthesized according to a previously reported synthetic procedure.<sup>26, 27</sup>



**Figure 1.** Schematic molecular representations of the hybrids  $K_{si}[pyr]$  and  $D_{si}[pyr]$  (adapted from X-Ray structures of related compounds)<sup>21-24</sup> according to different views. The red plane corresponds to a pseudo-symmetry plane of  $K_{si}[pyr]$  that is absent in  $D_{si}[pyr]$ . Color code: WO<sub>6</sub> octahedra, blue; PO<sub>4</sub> tetrahedra, green.

The incremental addition of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] to a solution of **K**<sub>si</sub>[**pyr**] (2 mM in CD<sub>3</sub>CN) up to *ca.* 0.5 equiv. mostly leads to the appearance of a single set of signals attributed to coordinated pyridyl units (Figure 2). Between 0.5 and 1.0 equiv. of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] this set of signals splits into two sets with very close chemical shifts. After the addition of *ca.* 1 equiv., the signals of the starting parent hybrid have almost disappeared suggesting that the formed species display a 1:1 stoichiometry between the POM and the Pd unit. At first sight, this behaviour is quite close to the one observed in the Dawson-analogue system, denoted as **D**<sub>si</sub>[**pyr**] (Figure 1).<sup>20</sup> In the previously reported study, two sets of <sup>1</sup>H NMR signals were also observed after the addition of 1.0 equivalents of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] to a solution of **D**<sub>si</sub>[**pyr**]. Diffusion-ordered NMR spectroscopy (DOSY) revealed that both sets of signals decayed similarly thus indicating that they correspond to species of similar shape and size. Furthermore in this previous study, high-resolution ESI-MS only showed peaks corresponding to the molecular triangle [(**D**<sub>si</sub>[**pyr**])<sub>3</sub>·(**PdCl**<sub>2</sub>)<sub>3</sub>] with

various charge states depending on the number of TBA counter-ions, whereas peaks corresponding to any other bigger oligomer, such as a tetramer, were not observed. The presence of the two sets of signals was attributed to the fact that in the molecular triangle **[(Dsi[pyr])3·(PdCl2)3]**, the Dawson unit can present two different orientations with respect to the plane defined by the Pd units. Hence in the molecular triangle displaying a pseudo Cs symmetry species, the aromatic rings of the organic linker are not chemically equivalent. In the present case, the hybrid **K**si**[pyr]** display an additional pseudo-symmetry plane, neglecting the inner phosphate, compared to **Dsi[pyr]** (Figure 1). It is thus very unlikely that the presence of the two sets of <sup>1</sup>H NMR signals would arise from the different orientations of the POM units with respect to the plane defined by the Pd units.



**Figure 2.** Evolution of the aromatic region of the <sup>1</sup>H NMR (600 MHz) spectra of a solution of **K**si**[pyr]** (2 mM) in CD<sub>3</sub>CN upon titration with *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>]: a) 0 equiv.; b) 0.25 equiv.; c) 0.5 equiv., d) 0.75 equiv., e) 1.05 equiv. The circles, rectangles, triangles and squares correspond to signals of **K**si**[pyr]**, **[(K**si**[pyr])**<sub>2</sub>·(PdCl<sub>2</sub>)], **[(K**si**[pyr])**<sub>3</sub>·(PdCl<sub>2</sub>)<sub>3</sub>] and **[(K**si**[pyr])**<sub>4</sub>·(PdCl<sub>2</sub>)<sub>4</sub>] respectively.<sup>28</sup>

DOSY <sup>1</sup>H NMR was performed at different steps during the addition of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] to a solution of **Ksi[pyr]** in order to characterize the different species formed upon metal coordination (Table 1). All sets of aromatic signals of **Ksi[pyr]** decay similarly according to the pulsed-field gradient strengths with a unique diffusion coefficient  $D_1 =$  $8.6 \times 10^{-10}$  m<sup>2</sup>.s<sup>-1</sup> (Figure S2). Interestingly, the diffusion coefficient of the TBA counter-ions ( $D_{TBA} = 1.50 \times 10^{-9}$  m<sup>2</sup>.s<sup>-1</sup>, Figure S2) is higher than that of the POM-based hybrid, suggesting that in acetonitrile, the TBA are not tightly associated to the POM. Upon the addition of 0.25 equiv. of trans-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>], the new doublets at 7.57, 7.75 and 8.80 ppm display a unique diffusion coefficient  $D_2 = 5.0 \times 10^{-10} \text{ m}^2 \text{ s}^{-1}$  (Table 1, Figure S4), a value that is slightly smaller than the diffusion coefficient of the starting building block. Furthermore the diffusion coefficient of the signals corresponding to the uncoordinated arms show an apparent weak decrease compared to Ksi[pvr] (Figure S5), while their decay vs the pulsed-field gradient strength cannot be satisfactorily modelled by using a single diffusion coefficient but by a twocomponent with D<sub>1</sub> and D<sub>2</sub> (Figures S5 & S6). In the previous study, the ratio of the diffusion coefficient of the free molecular building block to that of the molecular triangle  $[(D_{si}[pyr])_3 \cdot (PdCl_2)_3]$  was 2.3. The modest difference between D<sub>1</sub> and D<sub>2</sub> indicates that the first species formed at low concentration of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] is smaller than a trimer and more probably corresponds to the dimer [(Ksi[pyr])2·(PdCl2)] (Figure 3). This species contains uncoordinated ligands, which display <sup>1</sup>H NMR signals similar to those of K<sub>si</sub>[pyr]. This explains the apparent decrease of the diffusion coefficient of the signals at 7.50, 7.72 and 8.63 ppm in the presence of 0.25 equiv. of trans-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>]. The fact that the dimer [(Ksi[pyr])<sub>2</sub>·(PdCl<sub>2</sub>)] is the dominant species at intermediate concentration of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] suggests that the formation of the cyclic assembly is a non-cooperative process,<sup>29</sup> probably owing to a high entropic term that is not compensated by the chelate cooperativity.

**Table 1.** Diffusion constant of selected peaks during the addition of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] to a solution of **K**<sub>si</sub>[**pyr**].

[PdCl <sub>2</sub> (MeCN) <sub>2</sub> ]	$\delta / \text{ppm} (D \times 10^{-10} / m^2 . s^{-1}) \text{ assignment}$							
		8.63	7.94	7.72			7.50	
0 equiv.		(8.66)	(8.64)	(8.59)		(8.64)		
		М	М	М		М		
0.25 equiv.	8.80	8.63		7.75	7.72	7.57	7.50	
	(4.86)	$(8.6^a / 5.0^a)$		(5.01)	$(8.6^a / 5.0^a)$	(5.00)	(8.6 <sup>a</sup> / 5.0 <sup>a</sup> )	
	D	M+D		D	M+D	D	M+D	
0.50 equiv.	8.82	8.63	7.96	7.75	7.71		7.50	
	$(5.0^a / 3.0^a)$	$(8.6^a / 5.0^a)$	$(5.0^a / 3.0^a)$	$(5.0^a / 3.0^a)$	$(8.6^a / 5.0^a)$		(8.6 <sup>a</sup> / 5.0 <sup>a</sup> )	
	D+Te	M+D	D+Te	D+Te	M+D		M+D	
0.75 equiv.		8.63			7.71		7.50	
	$(8.6^a / 4.03)$			$(8.6^a / 4.00)$			$(8.6^a/4.32)$	
		M+D+bigger			M+D+bigger		M+D+bigger	
	acyclic species			acyclic species			acyclic species	

	8.82	8.80	7.96	7.93	7.75	7.72
1.05 equiv.	(2.93)	(3.58)	(3.01)	(3.74)	(3.08)	(3.64)
	Te	Tr	Te	Tr	Te	Tr

<sup>a</sup> Value obtained by imposing a constrain on the diffusion coefficient. M = monomer, D = dimer, Tr = trimer, Te = Tetramer

Upon the addition of 0.5 equivalent of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>], the decay of the signals at 7.50, 7.71 and 8.63 ppm (corresponding to non-coordinated organic linkers) can still be fitted by a double component with  $D_1 = 8.6 \times 10^{-10} \text{ m}^2 \text{.s}^{-1}$  and  $D_2 = 5.0 \times 10^{-10} \text{ m}^2 \text{.s}^{-1}$  (Figure S8). Ksi[pyr] and [(Ksi[pyr])2·(PdCl<sub>2</sub>)] are thus the major species containing non coordinated arms in these conditions. When 0.75 equivalent of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] are added to Ksi[pyr], the decay of these signals at 7.50, 7.71 and 8.63 ppm cannot be satisfactorily fitted with  $D_1 = 8.6 \times 10^{-10}$  $m^2.s^{-1}$  and  $D_2 = 5.0 \times 10^{-10} m^2.s^{-1}$ , but with values of  $8.6 \times 10^{-10} m^2.s^{-1}$  and  $4.1 \times 10^{-10} m^2.s^{-1}$ (Figure S11). This suggests that bigger acyclic oligomers can be present at this stage of the titration. Finally, upon the addition of a very slight excess of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] (1.05 equiv.) the two sets of signals corresponding to species with a 1:1 stoichiometry do not decay similarly indicating thus that they correspond to species of different sizes and compositions. The peaks at 7.72, 7.93 and 8.80 ppm can be nicely fitted with a single component  $D_3 =$ 3.7×10<sup>-10</sup> m<sup>2</sup>.s<sup>-1</sup> (Figure S13) while those at 7.75, 7.96 and 8.82 ppm also give a unique diffusion coefficient  $D_4 = 3.0 \times 10^{-10} \text{ m}^2 \text{.s}^{-1}$  (Figure S14). Interestingly the  $D_1/D_3$  ratio is 2.3, *i.e.* the value found for the molecular triangle formed from  $D_{si}[pyr]$ <sup>20</sup> The D<sub>4</sub> value, significantly smaller than D<sub>3</sub>, is thus attributed to the presence of a bigger species, most probably a molecular square. Note that the decay of the signals of the coordinated arms at 7.75, 7.96 and 8.82 ppm in the presence of 0.5 equiv. of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] can nicely be fitted by a double component with  $D_2$  and  $D_4$ , attesting the presence of the tetramer at this stage of the titration (Figure S9). NMR integration of the aromatic signals allows estimating the species distribution during the addition of trans-[PdCl2(MeCN)2] to Ksi[pyr] (Figure 3, Table S1).<sup>30</sup> In practice, the ratio between the integration of the signals at 7.57 ppm and 8.80-8.82 ppm versus those at 7.50 ppm and 8.63 ppm give the proportion of coordinated pyridine versus uncoordinated pyridine, while the proportion of the peak at 8.80 ppm within the overall signal at 8.80-8.82 ppm allows estimating the ratio between the different coordination oligomers. The comparison of the integration of all aromatics signals versus those of the TBA cations shows a slight decrease of ca. 10% after the addition of 1.05 equiv. of trans-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>]. This loss in the NMR signals is attributed to the minor formation of oligomeric species displaying Pd / POM ratio higher than 1. Finally the two sets of signals at 7.93 ppm and 7.96 ppm (corresponding to the triangle and the square respectively) after the addition of 1.05 equiv. of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] have similar integration indicating an identical distribution of the hybrid within its triangle and square forms.



**Figure 3.** Left) Metal driven formation of the molecular triangle and square from the POM hybrid **K**si[**pyr**]. Right) Species distribution during the addition of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] on a solution of **K**si[**pyr**] in CD<sub>3</sub>CN; green: **K**si[**pyr**], red: [(**K**si[**pyr**])<sub>2</sub>·(**PdCl**<sub>2</sub>)], blue: [(**K**si[**pyr**])<sub>3</sub>·(**PdCl**<sub>2</sub>)<sub>3</sub>] and [(**K**si[**pyr**])<sub>4</sub>·(**PdCl**<sub>2</sub>)<sub>4</sub>], black: remaining non attributed NMR signals.<sup>19</sup>

High-resolution ESI-MS was performed on a 10 μM solution of Ksi[pyr] in the presence of 1.0 equivalent of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] in MeCN. This 10 μM solution was obtained from a 2 mM solution similar to that analysed by NMR which was diluted a few minutes prior to the analysis. The resulting mass spectrum confirmed the stoichiometry of the species formed upon metal coordination. The mass spectrum mostly shows peaks that correspond to the trimer [(Ksi[pyr])<sub>3</sub>·(PdCl<sub>2</sub>)<sub>3</sub>] (Tr) with charge states 5- to 10-, depending on the number of TBA counterions attached ((Figure 4 and Figure S15). The other major peaks observed correspond to the tetramer [(Ksi[pyr])<sub>4</sub>·(PdCl<sub>2</sub>)<sub>4</sub>] (Te, Figure S16), the dimer [(Ksi[pyr])<sub>2</sub>·(PdCl<sub>2</sub>)] (D, Figure S17), the monomer Ksi[pyr] (M, Figure S18). Additional minor signals corresponding to the dimer with an additional PdCl<sub>2</sub> [(Ksi[pyr])<sub>2</sub>·(PdCl<sub>2</sub>)] (D<sup>\*</sup>) and the monomer with PdCl<sub>2</sub> [(Ksi[pyr])·(PdCl<sub>2</sub>)] (M<sup>\*</sup>) were also observed. Relative quantification of the species present in the solution from this mass spectrum is delicate because each oligomer appears with several charge states that all have different response factors so that the relative intensities cannot be considered directly proportional to their

relative quantities in solution. In addition, some oligomers (Tr and M\* or Te and D\*) appear at identical m/z values (Figures S15a, S15b, S16a). The last minute dilution and the instrumental parameters (electrospray ionization and ion transmission), although chosen to minimize any dissociation may also be responsible for the minor formation of the additional oligomer M\* and D\* that were not considered with NMR. Nevertheless, this study confirmed the concomittant formation of molecular triangles and squares in the presence of 1 equiv. of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>].



**Figure 4.** ESI mass spectrum of a solution of **K**si**[pyr]** (10  $\mu$ M) and *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] (1 equiv.) in MeCN. The main ion distribution corresponds to the molecular triangles and squares with a varying number of TBA counterions. Inset: Comparison of experimental (lower trace) and calculated (upper trace) isotopic distributions for *m*/*z* 1065, 1216 and 1261.

High-resolution ESI-MS was also performed on a 10  $\mu$ M solution of Ksi[pyr], in the presence of 0.5 equivalent of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] in MeCN (Figure S19). Similarly, this 10  $\mu$ M solution was obtained from a 2 mM solution and was diluted a few minutes prior to the analysis. Besides the major peaks of the starting monomer K<sub>Si</sub>[pyr], the mass spectrum mostly shows peaks corresponding to the dimer [(Ksi[pyr])<sub>2</sub>·(PdCl<sub>2</sub>)] (D) with charge states 4- to 6- and the monomer with an additional  $PdCl_2 [(Ksi[pyr]) \cdot (PdCl_2)] (M^*)$ . Only traces of the trimer  $[(Ksi[pyr])_3 \cdot (PdCl_2)_3]$  (Tr) were detected (Figure S20). This further evidenced that the metallomacrocycle species are hardly present at this intermediate ratio of *trans*- $[PdCl_2(MeCN)_2]$ .

### Correlation between diffusion coefficient and molecular weight

A power law between the diffusion coefficient D and the molecular mass M for a series of similar molecules has often been observed.<sup>31</sup>

$$D \propto \mathrm{M}^{-1/d_{\mathrm{F}}}$$

(1)

with  $d_{\rm F}$  being the fractal dimension of the molecular object. This correlation allows to predict the molecular mass from the diffusion coefficient without using complex optimized structures that are computationally challenging when dealing with large POM assemblies.<sup>19</sup> The fractal dimension can be seen as an indicator of the way an object fills up in the 3D space. This value ranges from 1 to 3 for purely one-dimensional objects and solid spheres respectively. This relation was applied to a wide range of cyclic POMs.<sup>32</sup> In this case a fractal dimension of 2.96 was observed, very close to the theoretical  $d_{\rm F} = 3$  value expected for perfect spheres. In the present case, owing to the high anisotropy of the POM-based building block Ksi[pyr], the fractal dimension is expected to be considerably lower. Furthermore, Ksi[pyr] and its coordination oligomers have different shapes, [(K<sub>Si</sub>[pyr])<sub>2</sub>·(PdCl<sub>2</sub>)] mostly featuring a linear shape, while the molecular triangle and square are cyclic assemblies, suggesting that the simple correlation of equation (1) may not be valid for this series of compounds. However, as shown in Figure 5, the relation of  $\log D vs \log M$  is linear with a slope of -0.735, giving a fractal dimension  $d_{\rm F} = 1.36$ . The fact that the power law between the diffusion coefficient D and the molecular mass M is applicable to this series of compounds, indicates that the shape of the molecular assembly has little effect on the diffusion coefficient and that the diffusion coefficient is probably mainly determined by the volume occupied by the compound. In other words for these discrete molecular assemblies, there is no need for complex models which would take into account the shape of the molecule. A similar observation has been previously done when comparing the diffusion coefficient of acyclic and macrocyclic drug-like molecules.33

![](_page_10_Figure_0.jpeg)

Figure 5. Relationship between diffusion coefficient and mass for K<sub>si</sub>[pyr] and its coordination oligomers.

## Comparison of complexation energy between Keggin type and Dawson type POM-based hybrids

In order to probe the thermodynamics of the formation of these complexes, an isothermal titration calorimetry (ITC) experiment was performed:<sup>34</sup> *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] was incrementally injected into a 1 mM solution of **Ksi[pyr]** (or **Dsi[pyr]**, respectively) in MeCN. The resulting exothermal heat effects are displayed on Figure 5 and qualitatively show an inflection point at a molar ratio of 1, which implies that (for both systems) the stoichiometry of the assembly is 1:1. Moreover, the steeper slope in the case of **Ksi[pyr]** indicates a stronger association for the Keggin-based system. Quantitative analysis of the data is tricky because (i) both sites on the Pd atom are probably dependent<sup>35</sup> and (ii) our NMR and ESI results unambiguously show that the self-assembly process involves several intermediates and products. Nevertheless, we tested whether the simplest association model considering a single set of identical binding sites could quantitatively describe the experiments. In other words, we analyse the data at the level of the elementary binding sites, without taking into account any possible interference between them:

### $L+n\;M \nleftrightarrow\; M_nL$

where L stands for one pyridine function of the ligand ( $[L] = 2[K_{si}[pyr]]$ ) and M stands for one binding site of the Pd(II) metal ( $[M] = 2[trans-[PdCl_2(MeCN)_2]]$ ). The results from the fits are displayed on Figure 5 and the values of fitted parameters are summarized in Table 2. The agreement between the data and this crude model is not perfect, but the slight discrepancy does not allow the use of a more elaborate model that would necessarily involve additional adjustable parameters. As expected, the stoichiometry of the interaction is 1:1 (*i.e.* one ditopic ligand to one ditopic metal) and the assembly is enthalpically-driven and entropically-opposed.

![](_page_11_Figure_1.jpeg)

**Figure 6.** ITC enthalpogram obtained by injecting a *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] solution in MeCN (10 mM) into a **K**si**[pyr]** (or **D**si**[pyr]**) solution in MeCN (1 mM) at 25°C. The lines correspond to fits according to a model considering a single set of identical sites.

The fit indicates a stronger binding in the case of the Keggin oxo-cluster that it is assigned to a more enthalpically favourable interaction. In **Ksi[pyr]** an identical distribution of the hybrid within its triangle and square forms are observed while only molecular triangles were detected with **Dsi[pyr]**. Consequently we assign the difference in binding enthalpy between **Ksi[pyr]** and **Dsi[pyr]** to reduced constrains in the molecular square that is only present with **Ksi[pyr]**. Weaker electrostatic repulsions between the POM building blocks forming the cyclic oligomers may also account for the stronger binding in the case of the Keggin oxo-cluster. The entropy formation in the self assembly of **Ksi[pyr]** and **Dsi[pyr]** is close while the formation of the tetramer *vs* trimer may have a significant difference of entropy.<sup>36</sup> The lower charge of **Ksi[pyr]** compared to **Dsi[pyr]** probably counter-balances the gain associated with the formation of the tetramer. Further studies on simpler systems (*e.g.* complexation of monotopic hybrids with a single metal ion) will be further investigated to evaluate more precisely the effect of the charge of the POM on the entropy reaction. **Table 2.** Thermodynamic parameters for the binding of POM ligands to *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>], deduced from Figure 5.

Ligand	Ν	$K / M^{-1}$	$\Delta H$ / kJ/mol <sup>-1</sup>	$\Delta G \ / \ kJ.mol^{-1}$	$T\Delta S / kJ.mol^{-1}$
K <sub>Si</sub> [pyr]	0.90	4.62 10 <sup>4</sup>	-43.5	-26.5	-16.9
Dsi[pyr]	0.93	$2.46\ 10^4$	-39.5	-25.0	-14.4

#### Conclusions

We have synthesized new supramolecular polygons from a Keggin-based molecular building block through metal-directed self-assembly. These species consist of a mixture of molecular triangles and squares while in a previous study using a higher charged building block, only molecular triangles were formed. This studies outline the decisive effect of the charge of the POM disfavouring the formation of large assemblies. The formation of highly charged supramolecular species (e.g., a Dawson-based tetramer) is most likely hindered by an important entropic contribution. This finding will help us to develop larger POM-based supramolecular assemblies. While the Keggin-based molecular building block and its coordination oligomers display different shapes a power law between the diffusion coefficient D and the molecular mass M is applicable to this series of hybrids. This indicates that the diffusion coefficient of these compounds is mainly determined by their occupied volume rather than by their shape. This simple model will allow to efficiently estimating the stoichiometry of a POM-based assembly only thanks to its diffusion coefficient.

### **Experimental Section**

*Trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>], **K**si[**pyr**] and **D**si[**pyr**] were prepared according to published procedures.<sup>20, 27, 37</sup> All other reagents were used as supplied.

NMR spectra were recorded either on a Bruker Avance III 600 spectrometer equipped with a BBFO probe head. The 1H DOSY spectra were acquired by using the pulse sequence ledbpgp2s. The gradient strength G was varied in 16 linear steps from 2.49 to 47.3 G cm<sup>-1</sup> to ensure at least 90 % signal attenuation. The spectra were processed by using the fitting routine integrated in the TopSpin3.5 package.

Mass spectrometry experiments were carried out by using a Synapt G2 HDMS (Waters, Manchester, UK) instrument equipped with an electrospray ionization source, which was operated in the negative ion mode. All desolvation, transmission parameters were tuned to

limit gas-phase ion fragmentation. Mass spectra were processed with MassLynx 4.1 and the theoretical isotopic patterns were predicted using mMass 5.5.0.<sup>38</sup>

ITC data were recorded on a MicroCal VP-ITC apparatus at 25°C, injecting a  $10^{-2}$  M *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] solution in MeCN into a  $10^{-3}$  M **K**si**[pyr]** (or **D**si**[pyr]**) solution in MeCN. Injections of 4 µL over 20 seconds were performed every 900 seconds at a stirring rate of 300 rpm. The heat of dilution of *trans*-[PdCl<sub>2</sub>(MeCN)<sub>2</sub>] was measured in the same conditions and was confirmed to be negligible. The data were fitted with MicroCal software.

### Supporting Information available.

Fits of the DOSY NMR spectra, comparison of experimental and calculated isotopic distributions of the ESI mass spectrum.

### Acknowledgements

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### **Table of Contents Graphic**

### **Table of Contents Synopsis**

The metal driven self-assembly of a Keggin-based hybrid leads to the formation of a mixture of molecular triangles and squares. This behavior is different from that of the structural analogue Dawson-based hybrid displaying a higher charge, which only led to the formation of molecular triangles. This studies highlights the decisive effect of the charge of the POMs in their self-assembly processes.