Comparison of [Pd2L4][BF4]4 cages for binding of n-octyl glycosides and nitrate (L = isophthalamide or dipicolinamide linked dipyrnyl ligand)

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Two dipyridyl ligands were synthesized, where the pyridyl donor fragments were separated by an isophthalamide (1) or a dipicolinamide moiety (2). Both ligands formed [Pd2(Ligand)4][BF4]4 complexes in CD2Cl2 containing 5% dmso-d6. It was found that while [Pd2(1)4][BF4]4 readily binds to n-octyl glycosides and to nitrate anions, [Pd2(2)4][BF4]4 did not. The difference in binding properties could be rationalized based on the reduced flexibility and size of the [Pd2(2)4]4+ cage and/or stronger interior binding of a BF4- counter anion.

Introduction

It is well-known that cage complexes of the type [M2L4]4+ are readily prepared when the metal (M) is a divalent Pd2+ ion and the ligand (L) is a dipyridyl ligand.1 Such complexes typically exhibit a hollow interior that is suitable to host smaller molecules.1a,b Recently, it was shown that such [M4L4]4+ complexes could be used to bind carbohydrates.2 In particular, it was shown that a dipyridyl ligand such as 1 in Fig. 1 formed a [Pd4L4]4+ cage4 that could bind to n-octyl-β-D-glucoside.3c As is highlighted in red in Fig. 1, the amides in a cage derived from 1 experience a steric clash with the isophthalamide’s central C–H fragment. We wondered what the effect would be of C–H → N adjustment of 1 to the dipicolinamide analogue 2, where the amides should be preorganized by intramolecular N–H⋯N hydrogen bonds (also highlighted in red).5 Herein, we report on the synthesis of 1 and 2 and their [Pd2(Ligand)4][BF4]4 complexes and shown that the C–H → N adjustment is detrimental to the binding properties of the cage.

Results and discussion

As is detailed in the ESI (section S2†), ligands 1 and 2 could easily be prepared according to known chemistry adapted from literature procedures.3c,6 Instead of the –C(O)NHCH2C(CH2)2–Bu group6 we opted for the –CH2O-p-Ph-C(p-tBu-Ph)3 group8 due to the more facile synthesis, particularly to obtain the dipicolinamide derived ligand. Stepwise addition of 0.5 equivalents of [Pd(OSMe)3][BArF]2 to a solution of 1 in CD2Cl2 containing 5% dmso-d6 led to the formation of single species on NMR, most likely [Pd2L4][BArF]4 (Fig. S32, † BArF = tetrakis[3,5-bis(trifluoromethyl)phenyl]borate). Following the same procedure with ligand 2 however, gave complicated spectra that did not resolve to a neat spectrum, not even after standing for a week (Fig. S33†). Repeating the procedure but with [Pd(NCMe)4][BF4]2 gave clearly resolved spectra for both 1 and 2 after standing for 7 days and these final spectra are shown in Fig. 2. The difference in complex formation is likely due to a templating effect of BF4-.‡ For the dipyridyl ligand 1 (Fig. 2a), the inwards facing s3NH, p2 and s4 displayed significant downfield shifts that are characteristic of [Pd2(Ligand)4]4+ formation.3c,d With ligand 2 (Fig. 2b) the inwards facing s3NH and p2 underwent an upfield shift upon Pd-coordination. With both ligands, the resonances belonging to the CH’s of the solubilizing groups remained unperturbed. Moreover, DOSY-NMR showed that the complexes had a larger diffusion constant than their parent ligand, which is consistent with formation of [Pd2(Ligand)4]4+ complexes. A thorough NMR spectroscopic evaluation of the complexes was also consistent with the formation of [Pd2L4]4+ and [Pd2L4]4+ and high resolution mass spectroscopy of both complexes was congruent with the 2:4 molar ratio of Pd to ligand (see Fig. S23 and S32†).

The binding affinities of both complexes for carbohydrates 3-6 listed in Table 1 was probed by 1H-NMR titration experiments in CD2Cl2 containing 5% dmso-d6. As is illustrated in Fig. 3 for the titrations with n-octyl-β-D-mannoside 3, significant peak shifting was observed for...
[Pd2L14]4+ (Fig. 3a) in the concentration range to 25 mM of 3. Shifting of peaks cannot result from the dilution of the complex, as dilution studies in the concentration range used during titrations (0.64–0.27 mM) revealed that all resonances remained stationary (see Fig. S36†). The resonances that shift most are the inwards pointing $s_3$NH, $p_2$ and $s_4$. The peak shifts could be analyzed using HypNMR, as is shown in the left-hand side of Fig. 3a. From this plot it is evident that all resonances display clear saturation behavior around 10 mM of 3. Fitting these shifts with HypNMR8 to a 1 : 1 binding model gave $K_a = 541 \pm 2.9 \text{ M}^{-1}$ with a reasonable accuracy ($r^2 = 0.9862$). Similar spectra and fits were obtained by titrations of [Pd2L14]4+ with the other carbohydrates. Moreover, NOESY spectroscopy of solutions of [Pd2L14]4+ containing galactoside 3 or glucoside 5 were consistent with carbohydrate binding to the interior of the cage (see Fig. S43 and S47†). Mass spectroscopic analysis of a solution of [Pd2L14]4+ with glucoside 5 supports a 1 : 1 binding stoichiometry (see Fig. S44†).

In sharp contrast to the titrations with [Pd2L14]4+, the spectrum of [Pd2L24]4+ (Fig. 3b) remained unperturbed when adding 3 to a concentration of 25 mM. Particularly surprising was the complete absence of any shifting of the inwards facing $p_2$, which is very characteristic for binding to the interior of these type of M2L4 cages.3,6 Very similar titration data could be collected with carbohydrates 4–6 (see Table 1).

Another noticeable observation from Table 1 is the lack of binding of [Pd2L24]4+ for all four carbohydrates. This made us wonder if the interior of [Pd2L24]4+ was capable of binding at all. To this end, a titration was conducted with (n-Bu)4N+NO3−. As is shown in Fig. 4, very significant peak shifting was observed which appear to saturate around 13 mM of NO3−.

Interestingly, while $s_3$NH and $p_2$ only shifted downfield, the resonance of $p_3$ initially shifted upfield, but then downfield. Such behavior is evidence of a binding stoichiometry
that is more complex than simple 1 : 1 binding. As is detailed in the ESI (Fig. S52†), these shifts could be modelled to a 1 : 3 binding model with a $K_{a1:1} = 159$, $K_{a1:2} = 63 \text{ M}^{-1}$ and $K_{a1:3} = 31 \text{ M}^{-1}$ ($r^2 = 0.9968$). Such a stoichiometry is consistent with a nitrate anion binding to the interior of $[\text{Pd}_2\text{La}_4]^+$ (s3NH and p2 shifts) as well as with both exterior sites involving p3, in close proximity to both $[\text{Pd}($pyridyl$)_4]^+$ environments. It has indeed been noted that such a binding mode, involving four charge assisted $[\text{C}–\text{H}]^+\cdots\text{nitrate}$ interaction, is common in $[\text{Pd}($pyridyl$)_4]^+$ complexes.5

A very similar titration involving $[\text{Pd}_2\text{La}_4]^+$ could also be modelled with this 1 : 3 stoichiometry (Fig. S46†), but in this instance with a $K_{a1:1} = 1862 \text{ M}^{-1}$ (and smaller $K_{a1:2} = 537 \text{ M}^{-1}$ and $K_{a1:3} = 316 \text{ M}^{-1}$, with $r^2 = 0.957$).

It thus appears that $[\text{Pd}_2\text{La}_4]^+$ readily hosts other molecules, but its CH $\rightarrow$ N analogue does not. To gain insight into the possible origin for this loss in binding capability, molecular modeling was conducted using density functional theory (DFT). The resulting models are shown in space filling mode in the left-hand side of Fig. 5 and are similar to previously reported crystal structures (see also Fig. S54 and S55†).4b,5b Also given in the figure are the inner dimensions (in Å) of the

Table 1 Overview of binding studies performed using $n$-octyl-glycosides 3–6 (with axial groups highlighted in blue) and $\text{NO}_3^–$ in $\text{CD}_2\text{Cl}_2$ containing 5% dmso-$d_6$.

| Host $\rightarrow$ Guest | $K_a$ of 1 : 1 binding$^a$ (M$^{-1}$) | $[\text{Pd}_1\text{La}_1]^+$ | $[\text{Pd}_2\text{La}_2]^+$ |
|------------------------|----------------------------------|--------------------------|--------------------------|
| 3                      | 541 i.p.s.$^b$                   |                           |                           |
| 4                      | 262                             |                           |                           |
| 5                      | 447                             |                           |                           |
| 6                      | 262                             |                           |                           |
| $\text{NO}_3^–$        | 1862$^c$                        |                           |                           |

$^a$ Binding constants were obtained by fitting observed chemical shift differences with HypNMR$^8$ as is detailed in section S3 of the ESI. $^b$ i.p.s. stands for the relatively ‘insignificant peak-shifts’ that were observed in the concentration range of 0–25 mM glycoside.

$^c$ Incorporating the higher concentration ranges, the affinities could only be modelled with a more complicated stoichiometry than simple 1 : 1 binding, but the 1 : 1 stoichiometries were still dominant or representative of the binding strength of the cages for nitrate anions. Details are provided in the text and in Fig. S42† for $[\text{Pd}_1\text{La}_1]^+$ and Fig. S48† for $[\text{Pd}_2\text{La}_2]^+$.

Fig. 3 Partial $^1\text{H}$ NMR spectra of titration experiments with $n$-octyl-$\beta$-D-mannoside 3 added to a solution of $[\text{Pd}_1\text{La}_1][\text{BF}_4]_4$ (a) or $[\text{Pd}_2\text{La}_2][\text{BF}_4]_4$ (b). The peak shifting observed in the titration with $[\text{Pd}_1\text{La}_1][\text{BF}_4]_4$ were fitted with HypNMR$^8$ as shown in the left. The solvent is $\text{CD}_2\text{Cl}_2$ containing 5% dmso-$d_6$. Further details are provided in section S40 of the ESI.$^†$

Fig. 4 Partial $^1\text{H}$ NMR spectra of a binding study with $($n-Bu$)_4\text{N}^+$-$\text{NO}_3^–$ added to $[\text{Pd}_2\text{La}_2][\text{BF}_4]_4$. The solvent is $\text{CD}_2\text{Cl}_2$ containing 5% dmso-$d_6$. See Fig. S52† for details.
models that were obtained by measuring intramolecular distances and subtracting twice the van der Waals radius of Hydrogen (1.09 Å) or Nitrogen (1.55 Å). While the complex with ligand 2 is about 1.4 Å wider (N–N versus CH–HC distance), the complex is also 1.4 Å less high (2.5 Å versus 3.9 Å). Actually, the height of [Pd2(2)]4+ of 2.5 Å is just large enough for host a nitrate anion (3.0 Å in height) when assuming van der Waals overlap in the order of 0.5 Å. However, the 2.5 Å height of [Pd2(2)]4+ is much smaller than the height of a glucoside (4.9 Å) and as a result very unlikely to fit. The dimensions of [Pd1L4]1+ on the other hand are much more congruent with the dimensions of NO3− and a glucoside, thus rationalizing why this complex binds to both (and with much greater affinities). Moreover, due to the preorganization of the amides in ligand 2 (see Fig. 1) its complex is expected to be very rigid. As a result, [Pd2(2)]4+ might well lack the conformational flexibility that could enable it to encapsulate a glucoside. Such a rational was also proposed previously for the comparison of isophthalamide versus dipicolinamide covalent cages in carbohydrate binding. Actually, inspection of the NOESY spectrum of [Pd2(2)]4+ (Fig. S27†) reveals the complete absence of a close proximity of the amide hydrogens (s3NH) and the aromatic CH hydrogens (s2) of the dipicolinamide fragment. This indeed implies that rotation of the amides is locked into a position with the NH hydrogens pointing to the interior of the complex (bound by the dipicolinic N). In the NOESY spectrum of [Pd1L4]1+ (Fig. S20†) on the other hand there is a clear nuclear Overhauser effect cross peak between the amide NH’s and the inwards pointing s4, as well as the outwards pointing s2. This is consistent with the reported flexibility of these amides in the solid state structures of related cage complexes. Finally, modelling of interior bound BF4− (Fig. S51†) shows a much tighter and energetically more stable fit within the [Pd2(2)]4+ binding pocket, suggesting that interior bound BF4− might hamper further binding. Such an adverse effect of BF4− on the binding potential of an M2L4 cage has been reported before. The increased flexibility of [Pd1L4]1+ compared to [Pd2(2)]4+, the smaller size of [Pd2(2)]4+, and the increased stability of the BF4− complex with [Pd2(2)]4+ together rationalizes the stark contrast in binding properties observed for both complexes.

Summary and conclusion

In summary, ligands were synthesized that bear two diprydyl donor groups linked by an isophthalamide (1) or a dipicolinamide moiety (2). Both ligands formed [Pd4(Ligand)4][BF4]4 complexes in CD2Cl2 containing 5% dmso-δ6. Cage [Pd4(1)4][BF4]4 was shown to bind to n-octyl glycosides 3–6 with affinities of about 250–500 M−1 in the order 3 > 5 > 4 = 6, and to nitrate anions with a 1 : 1 affinity Kα = 1862 M−1. In sharp contrast, cage [Pd4(2)4][BF4]4 did not appear to bind to glycosides and bound to nitrate with a 1 : 1 affinity of merely 159 M−1. The difference in binding properties could be rationalized based on the reduced flexibility and size of the [Pd4(1)4][BF4]4 cage, and its stronger complexation to a BF4− anion. It is thus concluded that preorganization of the amides in 2 by intramolecular NH–N hydrogen bonding has an adverse effect on the binding properties of [Pd4(2)4][BF4]4 compared to its CH analogue [Pd4(1)4][BF4]4, at least for n-octyl glycosides 3–6 and nitrate.

Conflicts of interest

There are no conflicts to declare.

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