Revealing the remarkable structural diversity of the alkali metal transfer agents of the trans-calix[2]benzene[2]pyrrolidine ligand†

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Excellent reagents for transferring their heterocalix[4]arene ligand to f-block organometallic complexes, lithium, sodium and potassium trans-calix[2]benzene[2]pyrrolidines have been found to adopt a fascinating series of structures in their own right.

Whilst following the recent seminal breakthroughs in f-block elements literature, the contributions of trans-calix[2]benzene[2]-pyrrolyl, H₂L₄⁺, in its doubly deprotonated ligand form (L₄⁻)²⁻, caught our eye. Introduced by Sessler et al. as one of a series of heterocalix[4]arene hybrids, (L₄⁻)²⁻ is made up of four aromatic rings of alternate pyrrole and arene units that connect via dimethylmethane linkers. Possessing interrupted conjugation, the ligand is blessed with conformational and coordinative flexibility that renders it effective at supporting a range of organometallic lanthanide and actinide complexes. These include uranium[II] and samarium[III] and thorium[V] complexes and most recently organoneptunium[II] complexes. Characterisation of these complexes especially via single crystal X-ray crystallography, a particularly noteworthy feat for the technologically challenging radiotoxic organoneptunium(III) complexes, have uncovered an assortment of trans-calix[2]benzene[2]pyrrolidine σ and π bonding modes, typically mono or bis(κ⁵-arene) coordination, bis(κ²-pyrrolyl) coordination, bis(κ⁵-pyrrolyl) sandwiching and bis(κ⁶-arene) sandwiching.

Alkali metal reagents play an important if somewhat understated role in this emerging organometallic chemistry of f-block elements. For example, potassium intermediate K₂(L₄⁻) reacted with SmCl₃(thf)₃ to generate the corresponding Sm₃Cl(L₄⁻), where L adopts a σ, π bonding mode with the pyrrolide N atoms σ-bonded to Sm, which occupies a bis(κ⁶-arene) π-pocket. Lithium intermediate, Li₂(L₄⁺), made by reaction of H₂L₄⁻ with LiHMDS† [HMDS = 1,1,1,3,3,3-hexamethyldisilazide; also known as bis(trimethylsilyl)amide] in toluene solution, was used to convert U₁ to dinuclear U₂⁴¹¹(L₄⁻), with one U in a bis(κ⁵-pyrrolyl) pocket and the second U in a bis(κ⁶-arene) pocket. Moreover, alkali metals can be incorporated into actinide products as seen in K[Th[N(SiMe₃)₂](L₄⁻)]† synthesised by reaction of Th⁴Cl₂(L₄⁻) with excess KHMDS. Here, aside from substituting Cl ligands by the silylamide, the K occupies the bis(arene) pocket of the macrocycle but interestingly also facilitates double C–H metallation of the arene by the Th⁴⁺ centre.

Recently, theoretical investigations by Schrekenbach, Pan et al. have put the alkali metal (Li, Na and K) derivatives of the trans-calix[2]benzene[2]pyrrolidine ligand in the spotlight for accessing low-valent uranium and transuranium complexes. Surprisingly, these important alkali metal intermediates AM₂(L₄⁻) have not been studied in their own right. Knowing that the structures of alkali metal compounds can have a profound influence on the outcome of reactions, we set out this study to isolate and characterize these intermediates and where possible to resolve their structures. As outlined here, the results obtained for the congeneric series of AM = Li, Na, K are remarkable.

In previous work potassium and lithium derivatives of trans-calix[2]benzene[2]pyrrolide were generated in situ by treating the free ligand with potassium hydride or lithium HMDS respectively. Adapting this procedure, a set of the alkali metal macrocycles was accessed by reaction of the free ligand with an appropriate metal-lating agent, namely either nBuLi, NaHMDS or KHMDS (Scheme 1). Crystallisation of the solvated potassium complex [K₂(L₄⁻)(thf)₃], was accomplished in neat thf solution. Initially obtained as a white solid, crystals of [Li₂(L₄⁻)(thf)₂] 2, suitable for X-ray analysis, were secured via slow diffusion of n-hexane into a tetrahydrofuran solution of 2. The distinction between thf-rich [Na₂(L₄⁻)(thf)₃], 3 and its thf-poor variant [[Na₂(L₄⁻)(thf)₃][Na₂(L₄⁺)]]ₐₕ, 4 came about by diffusing n-hexane into a tetrahydrofuran solution of the latter but growing the former from an all thf solution, while thf-free [Na₂(L₄⁻)ₚ], 5 was obtained by performing the sodiation reaction in neat methylcyclohexane.

As has been shown in papers, intuitively one would be inclined to (chem)draw the structure of the potassium intermediate with...
symmetry equivalent K cations attached to N atoms (i.e., replacing the two NH in H$_2$LA$^-$ by two NK in Scheme 1). Surprisingly, this representation of thf-rich I is found to be inappropriate. In actuality, the K1 cation bridges the two N atoms with one thf in a π-bonded N$_2$O pseudo-equatorial plane, while occupying a sandwich position between the two axially disposed κ$^2$-π-bonded aryl rings (Fig. 1a). On the other hand, K2 occupies the bis(η$^5$-pyrrolyl) pocket with its coordination completed by two thf ligands. An assessment of bond lengths emphasizes this bonding distinction (mean K1–N, 2.88 Å; mean K2–N, 3.20 Å). In fact, reflecting the distinct coordination mode for K1 vs. K2, I could be seen as a rare example of a potassium potassiate$^7$ where K1 represents the ate moiety in strongly binding one disordered component of two thf ligands for 1, whereas Na2 and Na4 occupy the bridging π-system of the η$^5$-pyrrolyl) if disregarding the long, weak contacts with the pyrrolyl N atoms. This is the first time that K has been found in the bis(η$^5$-pyrrolyl) pocket, drawing comparisons with the coordination of Np$^{III}$ in K(DME)$_n$[(L$^{Ar^+}$)Np$^{III}$OCH$_3$]]$_2^+$.9

where the two thf ligations in 1 match the actinide ion’s engagement with two MeO bridges though Np$^{III}$ also displays an additional interaction with a deprotonated aryl ring.$^1$d

Though molecular like 1, the structure of the smaller lithium congeners [Li$_2$(L$^{Ar^+}$)(thf)$_2$]$_2$ (Fig. 1b), differs significantly. Within its centrosymmetric arrangement, the bis(π-benzene) pocket lies empty. The Li cations cling to the outside of the macrocycle through a σ-bonded N atom [Li1–N1, 1.940(3) Å]. This N atom displays a distorted planar geometry (sum of bond angles, 358.49°) with distortion pronounced inside the pyrrolyl ring [C4NC1 bond angle, 105.68(12)°]. Structures of this type in which one or more pyrrole units has a N atom at the macrocycle exterior have been labelled as N-confused$^8$ isomers as in the samarium complex [[(L)Sm(HL$^+$)],$^1d$ where the protonated macrocycle HL$^+$ has one exterior N atom bound to Sm. Completed by two thf ligands, the primary Li (N × 1; O × 2) coordination sphere in 2 is essentially planar (sum of bond angles, 359.62°), though there are two exceedingly long interactions with C(Me) atoms of the dimethylmethylene linkers [Li1–C7, 2.727(3); Li1–C16, 2.815(3) Å].

Turning to the intermediate-sized alkali metal sodium, we crystallised the tris(thf) solvate [Na$_2$(L$^{Ar^+}$)(thf)$_2$]$_2$, 3, which bears a resemblance to the tris(thf) solvated potassium congener 1. The main distinction is that the smaller size of sodium leads to reduced hapticities of the bis(benzene) and bis(pyrrrole) rings with its two distinct Na centres, the former being η$^5$ bound [Na1–C9 2.8254(17); Na1–C25 2.8509(16)] and the latter being η$^5$ bound [Na2–C2 2.756(2); Na2–C18 2.8443(18); Na2–C19 2.8653(18); Na2–C3 2.8954(19)] (see ESI† for full details). Interestingly, as alluded to earlier [[Na$_2$(L$^{Ar^+}$)(thf)$_2$][Na$_2$(L$^{Ar^+}$)]$_2$]$_{0.5}$, 4, a thf-poor variant was also crystallised. Its zigzag polymeric structural arrangement exhibits four distinct sodium centres (Fig. 2a). Unfortunately, disorder affecting the Na1 and Na4 metal centres hampers the discussion of its structural parameters in detail. It can be said that the Na1 and Na3 centres occupy pockets of the intramolecular bis(benzene) type whilst the Na2 and Na4 centres make intermolecular Na bis(pyrrrole) units that bridge between ligands. A similar K coordination environment to that of the Na metal centres in 4 is also found in the potassium calix[4]pyrrolyl complex [[K$_2$[calix[4]pyrrolyl-3$^3$-thf(toluenes)$_2$]]$_2$]$_{0.5}$.9

Having crystallised variations of the sodium macrocycle containing different amounts of coordinated thf, we pondered whether crystals of the thf-free version could be obtained. This was achieved by performing the dideprotonative metallation reaction in a methylcyclohexane medium. An X-ray crystallographic determination revealed a remarkably eye-catching tetrameric, octanuclear structure [Na$_5$(L$^{Ar^+}$)], 5 (Fig. 2b). As highlighted in Fig. 2b, a 16-atom [Na$_{16}$] ring runs through the structure. Measuring from H nuclei to H nuclei the “hole” in the ring is approximately 125 Å$^3$, but given the van der Waals radii of an H approaches 1 Å that reduces the hole size to about 27 Å$^3$, approximately the size of a water molecule. There are four crystallographically distinct Na centres, which fall into two distinct types. Na1 and Na3 occupy the bis(η$^5$-benzene) pocket and form bonds of predominate σ character to the pyrrolyl N atoms as reflected by their short lengths (mean Na–N, 2.40 Å); whereas Na2 and Na4 occupy the bridging bis(η$^3$-pyrrolyl) units with a corresponding mean Na–N length.

**Fig. 1** Molecular structures of (a) [K$_2$(L$^{Ar^+}$)(thf)$_2$]$_2$, 1, and (b) [Li$_2$(L$^{Ar^+}$)(thf)$_2$]$_2$, 2. Thermal ellipsoids are displayed at 35% probability. Hydrogen atoms and one disordered component of two thf ligands for 1 have been omitted for clarity. The dashed lines illustrate the K–C and Li–Me contacts for 1 and 2, respectively. For 2, the symmetry operation used to generate equivalent atoms denoted with ′ is −x + 2, −y + 1, −z + 1.
of 2.62 Å. The contacts between Na1 and Na3 with their respective bis[η⁵-arene] cavities are in the range 2.707(2)–2.968(2) Å. The chameleonic character of sodium to switch from primarily a σ-bonded stance to a π-bonded stance with a NH-deprotonated pyrrole ligand has previously been observed in a series of sodium pyrrolylzincate structures. 10

In addition, compounds 1–5 were successfully characterised by ¹H and ¹³C NMR spectroscopy either in [D₈]thf or C₆D₆/[D₈]thf solutions (see ESI† for full details). All of them are distinguished by the absence of the pyrrolyl NH resonance and the presence of corresponding signals for the macrocyclic framework. 2 exhibits a characteristic broad singlet at 7.94 ppm in the ¹H NMR spectrum for the aromatic H flanked by the two CMe₂ groups; whereas this H resonates as a triplet in the range 6.69–6.93 ppm for 1 and 3–5. This situation reflects the distinction between the transoid conformation of the aryl units of the macrocycle in its Li salt form 2 and the alternative cisoid arrangement in the heavier K and Na congeners 1 and 3–5. For all of them, a singlet in the range 5.73–6.08 ppm is found for the pyrrolyl moiety in the ¹H NMR spectra.

In conclusion, prior to this work the structures of alkali metal derivatives of the dideprotonated trans-calix[2]benzene[2]pyrrole ligand were concealed since these compounds had only been studied as in situ transfer agents in organometallic lanthanide and actinide chemistry. Here, their isolation and structural characterisation have uncovered a surprisingly diverse range of novel structures from dinuclear monomers, to octanuclear tetramers through to chain polymers.

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