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# Monodentate coordination of the normally chelating chiral diamine (*R*,*R*)-TMCDA<sup>+</sup>

Ana I. Ojeda-Amador, Antonio J. Martínez-Martínez, Alan. R. Kennedy, David R. Armstrong and Charles T. O'Hara\*

After isolating an unusual binuclear, but monosolvated NaHMDS complex  $[\{(R,R)-TMCDA\}\cdot(NaHMDS)_2]_{\infty}$  which polymerises *via* intermolecular electrostatic Na $\cdots$ Me<sub>HMDS</sub> interactions, further (*R,R*)-TMCDA was added to produce the discrete binuclear amide  $[\{\kappa^2-(R,R)-TMCDA\}\cdot(NaHMDS)_2\{\kappa^1-(R,R)-TMCDA\}]$ , whose salient feature is the unique monodentate coordination of one of the chiral diamine ligands.

Chiral diamine ligands, for example (-)-sparteine, its (+)-sparteine surrogate and N,N,N',N'-(1R,2R)-tetramethylcyclohexane-1,2-diamine [(R,R)-TMCDA] have attracted considerable attention in asymmetric synthesis in a whole host of transition metal catalysed methodologies.<sup>1</sup> From an s-block perspective, when paired with an organolithium reagent it can be envisaged that 'chiral carbanions' are created, which can be used in subsequent enantioselective syntheses.<sup>2</sup> Focusing particularly on the  $C_2$ -symmetric ligand (R,R)-TMCDA, it has come to prominence recently as the availability of the historically more widely utilised diamine (-)-sparteine, has been unreliable over the past few years.<sup>3</sup> In terms of its coordination chemistry, (R,R)-TMCDA has worldwide interest and has been well studied. Over 50 metal complexes containing its ligated form have been reported, spanning both the s- (Li,<sup>4</sup> Na,<sup>4e</sup> K,<sup>4e</sup> and Mg,<sup>5</sup>) and d-block metals (Cu,<sup>6</sup> Zn,<sup>7</sup> Ru,<sup>8</sup> Pd,<sup>9</sup> Pt<sup>10</sup> and Hg<sup>11</sup>). Within s-block chemistry and germane to this work, Strohmann has comprehensively studied (R,R)-TMCDA complexes of synthetically important organolithium reagents (such as <sup>t</sup>BuLi,<sup>4a</sup> MeLi,<sup>4b</sup> <sup>i</sup>PrLi,<sup>4b</sup> <sup>s</sup>BuLi,<sup>4b</sup> <sup>n</sup>BuLi,<sup>4c</sup> BH<sub>3</sub>P(Ph)(Me)CH<sub>2</sub>Li,<sup>4d</sup> MeLi,<sup>4g</sup> PhLi,<sup>4h</sup> (allyl)Li<sup>4h</sup> and (benzyl)Li<sup>4i</sup> derivatives). An all-encompassing feature of all known structures is that the chiral diamine ligand adopts exclusively a  $\kappa^2$ -bidentate chelating mode. Due to the less flexible, fixed bite angle in (R,R)-TMCDA, with respect to that of N,N,N',N'tetramethylethylenediamine (TMEDA),12 it is a stronger chelating ligand than the latter,<sup>13</sup> with a recent study noting that it 'displays

no tendency to bind as a monodentate ligand.<sup>14</sup> This has been attributed to the  $\kappa^1$  (or by implication  $\eta^1$ ) form of (R,R)-TMCDA inducing severe steric strain due to the juxtaposition of the metal-NMe<sub>2</sub> with the uncoordinated NMe<sub>2</sub> group. The structural chemistry of alkali metal amide complexes continues to be an important topic of research.<sup>15</sup> We have recently discovered that lithium and sodium 1,1,1,3,3,3-hexamethyldisilazide (LiHMDS and NaHMDS) can capture alkali metal halide salts in the presence of donor ligands to form ion pair metal anionic crown (MAC) complexes, for example  $[Li\{(R,R)-TMCDA\}_2]^+[Li_5HMDS_5Cl]^-$ . 4f,16 A key starting material which remained hitherto elusive in our studies involving sodium is the (R.R)-TMCDA-solvated NaHMDS complex. Crystallisation of other donor ligated [e.g., Me<sub>6</sub>TREN<sup>17</sup> and (-)-sparteine<sup>18</sup>] NaHMDS complexes has proven difficult, although the polymeric TMEDA  $[(\mu\text{-TMEDA})\cdot(\text{NaHMDS})_2]_{\infty}^{19}$ and N,N,N',N'-tetramethylpropanediamine (TMPDA) [(µ-TMPDA)·  $(NaHMDS)_2]_{\infty}^{20}$  complexes, which propagate *via* the nonchelating diamine ligand, are known (Fig. 1). These have similar structural motifs to Williard's lithium diisopropylamide (LDA) complex  $[(\mu\text{-TMEDA})\cdot(\text{LDA})_2]_{\infty}$ .<sup>19</sup>

In an effort to prepare the (*R*,*R*)-TMCDA complex of NaHMDS, an equimolar mixture of NaHMDS and (*R*,*R*)-TMCDA was combined in *n*-hexane medium and left to stir at ambient temperature for 1 hour (Scheme 1). The reaction mixture was then cooled to -33 °C and crystals suitable for X-ray crystallographic analysis deposited after 48 hours (27% non-optimised, crystalline yield; maximum yield 50% based on (*R*,*R*)-TMCDA consumption). X-ray data reveal the mono-(*R*,*R*)-TMCDA, binuclear [{(*R*,*R*)-TMCDA}-(NaHMDS)<sub>2</sub>]<sub>∞</sub> **1** (Fig. 2a). There are six crystallographically distinct but essentially chemically



Fig. 1 Structures of previously known polymeric  $[(\mu-TMEDA)\cdot(NaHMDS)_2]_{\infty}$  and  $[(\mu-TMPDA)\cdot(NaHMDS)_2]_{\infty}$ .

WestCHEM, Department of Pure and Applied Chemistry, University of Strathclyde, 295 Cathedral Street, Glasgow, G1 1XL, UK. E-mail: charlie.ohara@strath.ac.uk † Electronic supplementary information (ESI) available: General synthetic procedures, crystal structure determinations and NMR spectroscopic data. CCDC 1501992 and 1501993. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c6cc07190b



 $\label{eq:scheme1} \begin{array}{l} \mbox{Scheme1} & \mbox{Syntheses of } [\{(R,R)\mbox{-}TMCDA\}\mbox{-}(NaHMDS)_2]_{\infty} \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA\}\mbox{-}(NaHMDS)_2]_{\infty}$ \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA\}\mbox{-}(NaHMDS)_2]_{\infty}$ \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA\}\mbox{-}(NaHMDS)_2]_{\infty}$ \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA\}\mbox{-}(NaHMDS)_2]_{\infty}$ \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA]\mbox{-}(NaHMDS)_2]_{\infty}$ \mbox{1} and $[\{\kappa^2\mbox{-}(R,R)\mbox{-}TMCDA]\mbox{-}(R,R)$ 

equivalent molecules of  $[\{(R,R)\text{-TMCDA}\}$ ·(NaHMDS)<sub>2</sub>] in the structure of 1, thus for brevity only one is discussed here. Interestingly, the empirical formula of 1, *i.e.*,  $[(\text{donor})\cdot(\text{NaHMDS})_2]$  is identical to that for the aforementioned TMEDA and TMPDA derivatives; however, in keeping with previously known (R,R)-TMCDA complexes, the diamine adopts a chelating bonding mode, and with respect to the N donor atoms, renders one Na metal centre (Na1) four-coordinate in a distorted tetrahedral arrangement (bond angles range from 68.70(9) to 151.55(10)°, see ESI† for full details). Additionally, Na1 has two long Na…Me interactions with a methyl group from each HMDS ligand [Na1…C12 2.987(4) and Na1…C22 2.987(4) Å]. The second Na metal centre (Na2) remains only two-coordinate with respect to the bridging amido N atoms. To satisfy this electron deficiency, Na2



Fig. 2 (a) Molecular structure of  $[\{(R,R)-TMCDA\}\cdot(NaHMDS)_2]_{\infty}$  1 showing one molecule from the asymmetric unit. Hydrogen atoms omitted for simplicity and thermal ellipsoids are displayed at 35% probability. (b) Section of the zigzag polymeric chain of 1. The dashed lines illustrate  $Na\cdots Me(SiMe_2)$  interactions. The symmetry operation used to generate the atoms labelled with ' is -x + 1, y + 1/2, -z + 1.

engages a solitary intermolecular Na···Me(SiMe2) [Na2···C65 distance, 2.818(4) Å] electrostatic interaction (Fig. 2b), which is short in comparison to known literature examples [range Na···Me(SiMe<sub>2</sub>) 2.947–3.138 Å].<sup>21</sup> This sole intermolecular Na...Me interaction induces propagation of binuclear units in a zigzag polymer chain. This change in the coordination chemistry of (R,R)-TMCDA in 1 with respect to the bridging TMEDA and TMPDA ligands in the aforementioned polymeric sodium amides emphasises the propensity for the chiral 1,2-diamine to remain as a chelating ligand rather than binding in a monodentate fashion. As a consequence of this coordination mismatch, significantly shorter Na2-N<sub>HMDS</sub> bonds (mean distance, 2.356 Å) are observed when compared with Na1-N<sub>HMDS</sub> bonds (mean distance, 2.530 Å). Despite utilising a 1:1 ratio of NaHMDS: (R,R)-TMCDA in this synthesis, it is clearly evident that the ultimate ratio in 1 is 2:1. When this optimised ratio is used in the synthesis, 1 was again the sole product isolated (36% crystalline yield).

Complex **1** is a rare example of a solvated sodium amide which contains an unsolvated Na site. Bochmann revealed the mono(tetrahydrofuran), mono(THF), complex  $[(THF) \cdot (NaHMDS)_2]$ where one Na atom is two coordinate whilst the other binds to the ether to render it three coordinate.<sup>22</sup> Interestingly, seven years prior to this report Dehnicke published the bis(THF) analogue  $[(THF)_2 \cdot (NaHMDS)_2]$  where both Na atoms are three coordinate.<sup>23</sup> This begged the question: '*could the coordinatively unsaturated (Lewis acidic) Na atom in 1, act as a host for another Lewis base*?'

A logical route to address this question would be to utilise monodentate donors such as THF and diethylether, in an attempt to saturate the deficient metal centre; but, it is highly likely that these strong  $\sigma$ -donors would also displace the chelating (R,R)-TMCDA ligand. Therefore to maintain synthetic simplicity, we repeated the preparation of 1 but employing an excess (two molar equivalents) of (R,R)-TMCDA with respect to NaHMDS in an attempt to coordinate a second molecule of the Lewis base ligand to the donor-free metal centre. High quality crystals (39% crystalline yield) were obtained by storing the resultant solution at -33 °C for 24 h, which were analysed by X-ray crystallography and were pleasingly found to be the target bis(solvated) derivative  $[\{\kappa^2 - (R,R) - TMCDA\} \cdot (NaHMDS)_2 \{\kappa^1 - (R,R) - K(R,R) - K(R) - K(R) - K(R,R) - K(R) -$ TMCDA}] 2 (Fig. 3). The distorted tetrahedral coordination sphere of Na1 in 2 (bond angles around Na1 range from 66.90(6) to 151.05(8), see ESI<sup>+</sup>) is essentially identical to that found in 1, exhibiting additional long contacts with a methyl group from each HMDS amido ligand [Na1···C27 2.968(3) and Na···C24 2.976(3) Å]. However, the second sodium metal centre, Na2, is additionally coordinated to an extra molecule of (R,R)-TMCDA, giving rise to a distorted trigonal planar geometry. As such there are two distinct coordinated diamine ligands within the structure of 2. Undoubtedly, the most eye-catching feature is that one (R,R)-TMCDA ligand adopts a previously unseen  $\kappa^1$ -coordination mode. To change from a  $\kappa^2$ - to a  $\kappa^1$ -coordination mode, it appears that inversion of the N1 atom of the (R,R)-TMCDA has occurred, no longer allowing the ligand to chelate to Na2 (Fig. 3).

Complex 2 is a discrete dimeric entity, despite the potential availability for N2 to coordinate further. In theory, this could be



**Fig. 3** Molecular structure of  $[{\kappa^2-(R,R)-\text{TMCDA}} \cdot (\text{NaHMDS})_2{\kappa^{1-(R,R)-\text{TMCDA}}]$  **2**. Hydrogen atoms and one disordered component of the mono-dentate (*R*,*R*)-TMCDA ligand are omitted for simplicity. Thermal ellipsoids are displayed at 35% probability.

achieved if this N atom could also invert thus allowing an additional exo-coordination site; however, it is unlikely that this would occur due to high steric strain (buttressing).<sup>14</sup> The  $\kappa^{1}$ -coordinated (*R*,*R*)-TMCDA is disordered over two domains, but its atomic connectivity and geometry are unequivocal. The  $\kappa^2$ - and the hitherto unseen  $\kappa^1$ -coordination mode (*R*,*R*)-TMCDA observed in 2 can be compared with DFT calculations (at the B3P86/6-311+G\* level) performed for its diamine relative (-)-sparteine (Fig. 4).<sup>24</sup> It has been shown that when (-)-sparteine binds to a metal complex, it always adopts a chelating 'cis' configuration. However, in the absence of a metal complex, it is actually slightly more stable (by 3.4 kcal  $mol^{-1}$ ) in a ring-flipped '*trans*' configuration [akin to our  $\kappa^1$ -coordinated (*R*,*R*)-TMCDA] where the lone pairs of electron present on the N atoms are not adjacent to each other. We have performed similar DFT studies (ESI<sup> $\dagger$ </sup>) on (*R*,*R*)-TMCDA and have shown that there is negligible difference (less than 1 kcal mol<sup>-1</sup>) between the potentially  $\kappa^{1}$ - and  $\kappa^{2}$ -coordination modes.

As **1** and **2** are both highly soluble in non-polar hydrocarbon and arene solutions, solutions of these compounds were studied by NMR spectroscopy. Using <sup>1</sup>H NMR spectroscopy, it was evident that the expected **1**:2 and **2**:2 (*R*,*R*)-TMCDA:HMDS ratios were observed respectively. For **1**, a single amido resonance (at  $\delta$  0.25) was observed and the (*R*,*R*)-TMCDA resonances (at  $\delta$  2.01, 1.90, 1.47 and 0.74) in C<sub>6</sub>D<sub>6</sub> solution appeared to correspond to a metallocoordinated ligand (see ESI† for full details). For **2**, the amido resonance appears at  $\delta$  0.31 in the same solvent. If the solid state structure of **2** was to be retained in solution, two unique sets of (*R*,*R*)-TMCDA resonances would be expected. In reality a single set of resonances (at  $\delta$  2.06, 1.99, 1.51 and 0.80 in C<sub>6</sub>D<sub>6</sub> solution) is observed. This indicates that a single (*R*,*R*)-TMCDA environment exists at 300 K in arene solution, indeed, a variable temperature



Fig. 4 Relative stabilities of *cis* and *trans* isomers of uncoordinated (–)-sparteine.<sup>24</sup>

NMR spectroscopic study of 2 in  $[D_8]$ -toluene solution unveiled that this situation was maintained even at low temperature (down to 206 K, see ESI<sup>†</sup>). In addition, <sup>1</sup>H and <sup>13</sup>C NMR spectra obtained in non-polar  $[D_{12}]$ -cyclohexane also reveal this situation (see ESI<sup>†</sup>). Therefore due to the steric bulk of the HMDS ligands within the molecule [thus precluding a dual  $\kappa^2$ -situation for the (*R*,*R*)-TMCDA ligands], it is likely that the spectra show a time-averaged situation between dynamic  $\kappa^1$ - and  $\kappa^2$ -coordinated (*R*,*R*)-TMCDA ligands.

In closing, we have shown that counter to previous studies, (R,R)-TMCDA can indeed bind to an alkali metal in a nonchelating  $\kappa^1$ -manner.

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