Self-assembly of tetrareduced corannulene with mixed Li–Rb clusters: dynamic transformations, unique structures and record $^7$Li NMR shifts†

Alexander S. Filatov, Sarah N. Spisak, Alexander V. Zabula, James McNeely, Andrey Yu. Rogachev and Marina A. Petrukhina

Self-assembly processes of the highly reduced bowl-shaped corannulene generated by the chemical reduction with a binary combination of alkali metals, namely Li–Rb, have been investigated by variable-temperature $^1$H and $^7$Li NMR spectroscopy. The formation of several unique mixed metal sandwich products based on tetrareduced corannulene, C$_{20}$H$_{10}$$^{4–}$ ($^{1}$–$^{4}$), has been revealed followed by investigation of their dynamic transformations in solutions. Analysis of NMR data allowed to propose the mechanism of stepwise alkali metal substitution as well as to identify experimental conditions for the isolation of intermediate and final supramolecular products. As a result, two new triple-decker aggregates with a mixed Li–Rb core, [Rb(THF)$_2$]$^2$/[Li$_3$Rb$_2$(C$_{20}$H$_{10}$)$_2$(Li$^+$THF)]$_2$ and [Rb(diglyme)$_3$]$^2$/[Li$_3$Rb$_2$(C$_{20}$H$_{10}$)$_2$(diglyme)$_2$] 0.5THF (3·0.5THF), have been crystallized and structurally characterized. The Li$_3$Rb$_2$-product has an open coordination site at the sandwich periphery and thus is considered transient on the way to the Li$_3$Rb$_5$-sandwich having the maximized intercalated alkali metal content. Next, the formation of the LiRb$_5$ self-assembly with $^{1}$–$^{4}$ has been identified by $^7$Li NMR as the final step in a series of dynamic transformations in this system. This product was also isolated and crystallographically characterized to confirm the LiRb$_5$ core. Notably, all sandwiches have their central cavities, located in between the hub-sites of two C$_{20}$H$_{10}$$^{4–}$ decks, occupied by an internal Li$^+$ ion which exhibits the record high negative shift (ranging from $–21$ to $–25$ ppm) in $^7$Li NMR spectra. The isolation of three novel aggregates having different Li–Rb core compositions allowed us to look into the origin of the unusual $^7$Li NMR shifts at the molecular level. The discussion of formation mechanisms, dynamic transformations as well as unique electronic structures of these remarkable mixed alkali metal organometallic self-assemblies is provided and supported by DFT calculations.

Introduction

Non-planar carbonaceous aromatic compounds, such as fullerences and carbon nanotubes, have recently emerged as promising light-weight materials for electronics and energy-related applications. Their potential application in lithium-ion battery technology has also attracted special attention. Over, the unique thermoelectric and superconducting properties of the reduced forms of curved carbon frameworks have triggered a renewed interest in these materials. All these recent advances stimulated a rapid expansion of the family of curved carbon-rich compounds which now includes a great variety of carbon bowls, warped nanographene sheets, as well as nanobelts and nanoribbons of different sizes. Bowl-shaped polyaromatic hydrocarbons (also referred to as carbon or π-bowls) have been broadly investigated over the last two decades to reveal their unique coordination and redox properties. For example, they can readily accept multiple electrons in stepwise reduction reactions to form the sets of the consequently reduced non-planar polyaromatic carbonanes. The latter have been the subjects of numerous investigations due to their unique interplay of strain and conjugation. It was demonstrated that the smallest bowl-shaped polyarene, corannulene (C$_{20}$H$_{10}$, Scheme 1a) is able to undergo four reduction steps to form a set of corannulene anions, C$_{20}$H$_{10}$$^{n–}$ ($n = 1–4$). The final fourfold reduction is accompanied by a significant core rearrangement of the corannulene tetraanion which can be considered to have an annulene-within-an-annulene electronic structure (Scheme 1b). Notably, C$_{20}$H$_{10}$$^{4–}$ bears one electron per five carbon atoms and is more electron rich than the fullerene-related applications.

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hexaion (one electron per ten carbon atoms in C_{60}^{6−}). This makes it a very interesting and unique carbanion for investigation of metal coordination and self-assembly reactions.

The highly reduced corannulene anions exhibit remarkable ability to form unique supramolecular aggregates in solutions, as shown by extensive ¹H and ⁷Li NMR spectroscopy studies. However, structural investigations of the charged π-bowls were lagging behind their solution spectroscopic studies until we have accomplished the first X-ray diffraction characterization of the product formed by the tetrareduced corannulene with lithium counterions. The formation of a remarkable aggregate with a Li₃-core sandwiched between two C_{20}H_{10}^{4−} decks has been established (Scheme 2a). This supramolecular aggregation with lithium ions allows to achieving a tetrareduced state of corannulene, as the electrochemical generation of C_{20}H_{10}^{4−} cannot be accomplished due to a very large negative standard potential located outside of the current experimental window. We have also demonstrated that the triple-decker supramolecular aggregate, [Li₅(C_{20}H_{10}^{4−})]₃³⁻, can be crystallized in different external coordination environments, showing no significant effect on the geometry of the sandwich core. In addition, the formation of the [Li₅(C_{20}H_{10}^{4−})]₃³⁻ product (abbreviated as Li₅ below) is confirmed in solution based on the observed shifts for sandwiched Li⁺ ions (−11.70 ppm) and a proper 3 : 5 integration of external vs. internal ions in the ⁷Li NMR spectra. Importantly, these results illustrated the ability of tetrareduced corannulene to engage all of its adjacent six-membered rings in alkali metal binding and thus to encapsulate a large amount of Li⁺ ions, which can be related to the high charge capacity of the corannulene-based electrodes in Li-ion batteries.

Remarkably, the coordination limit of the highly electron rich corannulene tetrabenion can be further extended through the synergistic use of two alkali metals as the reducing agents. The concomitant reduction of C_{20}H_{10} using a Li–K mixture resulted in the recent discovery of a novel class of mixed metal supramolecular products, in which tetrareduced corannulene exhibited its new coordination record. The C_{20}H_{10}^{4−} anion is able to engage all its sites, five benzene rings along with a central five-membered ring, for binding of six alkali metal ions in the resulting mixed metal core, Li₃K₃ (Scheme 2b) or LiK₅ triple-decker sandwiches. Notably, the previously unseen involvement of the hub-site of corannulene tetraanion in lithium ion binding is accompanied by unprecedented negative shifts in ⁷Li NMR spectra (up to ca. −25 ppm). These recent results established a new paradigm for curved polyaromatic ligands in alkali metal binding and opened new opportunities for design and synthesis of novel mixed metal organometallic supramolecular products. However, the origin of the observed high negative shifts in ⁷Li NMR spectra was not understood and required additional attention.

Herein, we set to investigate the reduction reactions of corannulene using a binary alkali metal combination comprised of Li and Rb, the heavier congener of K. For the Li–Rb combination, we expect that a greater size mismatch of two alkali metals could not only facilitate the transformations observed in the Li–K–C_{20}H_{10} system but also afford novel supramolecular products and open new reaction pathways that have not been seen before. We also expect that isolation and analysis of novel mixed metal organometallic products formed by the highly charged corannulene could help to shed light on the record negative shifts observed in ⁷Li NMR spectra. We therefore used a combination of NMR spectroscopy, X-ray crystallography and DFT theoretical methods for thorough investigation of these complex systems. As a result, we have been able to reveal a direct relation between the structures of supramolecular assemblies and coupling effects of the highly charged polyaromatic bowls at the molecular level and to correlate those with the observed ⁷Li NMR shifts.

Results and discussion

Corannulene reacts with alkali metals, Li through Cs, to initially produce intense green solutions characteristic for the corannulene monoanion. The excess of alkali metal quickly reduces the resulting C_{20}H_{10}^{−} monoanion to the purple C_{20}H_{10}^{2−} dianion. In the case of Li metal, dianions can then be further reduced to tetraanions that form supramolecular [Li₅(C_{20}H_{10}^{4−})]₃³⁻ aggregates having characteristic ⁷Li NMR shifts for sandwiched Li⁺ ions (−11.70 ppm). While mono- and dianions of corannulene do not form such supramolecular species, they exhibit a variety of coordination modes that depend on the size of alkali metal ions and other experimental variables. The corannulene bowl is not flattened upon acquisition of the first electron but more pronounced changes can be seen upon addition of the second electron. In contrast, a significant bowl depth decrease and C–C bond length alteration pattern are observed for tetrareduced corannulene.

Scheme 1 Corannulene (a) along with the electronic structure of C_{20}H_{10}^{4−} (b).

Scheme 2 Supramolecular aggregates with (a) Li₅− and (b) Li₃K₃−cores sandwiched between two C_{20}H_{10}^{4−} decks.
These experimental observations are in line with multiple theoretical predictions.\textsuperscript{11,18,20}

In Li\textsubscript{5}-sandwiches all five benzene rings of C\textsubscript{20}H\textsubscript{10}\textsuperscript{4−} are engaged in lithium ion binding leaving an internal space between two corannulene hub-sites empty. Notably, the distance between the centroids of 5-membered rings in the triple-decker Li\textsubscript{5}-products is 3.5 Å\textsuperscript{11} and this may not be sufficient to accommodate an additional lithium ion in that space. For comparison, the separation between 5-membered rings in lithium cyclopentadienide, [(η\textsuperscript{5}-Cp)\textsubscript{2}Li], is ca. 4.0 Å,\textsuperscript{2} and the Li-Cp distances range from 3.8 to 4.1 Å in some other cyclopentadiene Li-organometallic compounds.\textsuperscript{22,23} We have recently demonstrated\textsuperscript{15} that in order to get access to the central cavity larger alkali metals, such as K, should be introduced into the reaction along with Li. The initially formed Li\textsubscript{5}-sandwich having both corannulene decks parallel to each other starts to open up upon stepwise substitution of small Li\textsuperscript{+} ions by larger K\textsuperscript{+} ions, leading to the products having angled corannulene decks (Scheme 3). This results in the opening of a channel which allows the insertion of Li\textsuperscript{+} ion from periphery into the previously inaccessible inner cavity of the sandwich (Scheme 3, pathways A and B). It is necessary to have two large alkali metal ions at the rim of the sandwich for the lithium insertion process to take place, as the first substitution step (formation of Li\textsubscript{4}K) does not provide a sufficient opening for such insertion. For the second step, there are two possible pathways, A and B (through the formation of α and β isomers), where rings 2 and 5 or 2 and 3 are occupied by larger alkali metals (Scheme 3). Our DFT calculations showed that in the Li-K systems the insertion occurs through the β-isomer only (pathway A).\textsuperscript{15}

Monitoring the reduction reaction of 1 with a mixture of Li and Rb metals in THF-d\textsubscript{8} by NMR spectroscopy confirms the formation of supramolecular products formed by C\textsubscript{20}H\textsubscript{10}\textsuperscript{4−}. The reaction seems to proceed faster than in the case of Li-K and substantial amounts of Li-Rb-sandwiches are quickly observed within a few hours (Fig. 1a).

At the initial stage of reaction, which is determined by its characteristic brown-red color, two major broad peaks at −23.92 and −24.46 ppm and one very small peak at −24.86 ppm are observed in \textsuperscript{7}Li NMR spectrum (Fig. 1a). The significant upfield-shift of these peaks is associated with the Li\textsuperscript{+} ion squeezed between two 5-membered rings of tetrareduced corannulene, revealing the formation of several mixed metal Li-Rb sandwich-type assemblies. The latter peaks are assigned to the Li\textsubscript{4}Rb\textsubscript{2} and Li\textsubscript{3}Rb\textsubscript{2} products (α- and β-isomers, see Fig. 2 for schematic representation of the sandwich cores) based on integration of these peaks and the possibility to crystallize these major products in the single-crystalline form, as reported below. It can be mentioned here that all NMR peaks coalesce into a single broad signal at temperatures ranging from +20 to −20 °C, showing the rapid lithium ion exchange between all molecular structures existing in solution (ESI, Fig. S1†).

The \textsuperscript{7}Li signal at −11.70 ppm in Fig. 1a belongs to the monometallic Li\textsubscript{3}-product in which five Li\textsuperscript{+} ions are occupying the sites in between the benzene rings of C\textsubscript{20}H\textsubscript{10}\textsuperscript{4−} (Scheme 2a). Additional peaks between −6.2 and −8.3 ppm are the signals of lithium ions located between the same 6-membered ring sites but belonging to the mixed metal Li-Rb sandwiches (similar to the reported Li-K aggregates).\textsuperscript{15}

In contrast, the \textsuperscript{7}Li NMR signals of Li\textsuperscript{+} ions coordinating to the exterior of triple-decker sandwiches (filling the open concave cavities of C\textsubscript{20}H\textsubscript{10}\textsuperscript{4−} bowls) appear as broad peaks in the range between −4 to −6 ppm. The small intensity of these
peaks observed in this work suggests that the larger Rb\(^+\) ions are preferred over smaller Li\(^+\) ions as the external charge-compensating cations. The \(^7\)Li NMR peaks around 0 ppm belong to the solvent-separated Li\((\text{THF})_2\)^\(^+\) species. No other peaks are observed in the +20 to −30 ppm range (Fig. S2†). Notably, the ratio of [Li\(_2\)(C\(_{20}\)H\(_{10}\)\(^4\^-\))]\(^b\) to the mixture of all Li-Rb sandwiches is 0.38 to 1. This is consistent with the \(^1\)H NMR spectra which also show a 0.38 ratio of the representative peaks at 6.86 ppm (the signal of C\(_{20}\)H\(_{10}\)\(^4\^-\) in Li\(_3\)) and 6.69 ppm (a broader peak associated with all mixed Li-Rb products). It should be mentioned here that the precise time assignment to a particular point in the series of dynamic NMR measurements is problematic due to very strong dependence of the system reactivity on very minor and hard-to-control experimental variations. For example, we observed that diminutive changes in purity of the Li\(_5\)-sandwich, appearing as a very broad peak between −4 and −6 ppm, and the corresponding signals of external Li\(^+\) ions bound to other sandwich products existing in the mixed metal system all contribute to the elevated background.

Since the \(^7\)Li signal at −11.70 ppm belongs to the Li\(_5\)-sandwich, containing lithium ions only,\(^{11}\) the close small peak at −9.62 ppm is assigned to the product resulting from the first substitution of one sandwiched Li\(^+\) ion by a Rb\(^+\) ion (to form a Li\(_3\)Rb\(_2\) core). No peak correlates with this one in the region of −20 to −30 ppm in the \(^7\)Li NMR spectrum and, as expected, there is no internal lithium insertion at this stage. This transient species undergoes further substitution and thus does not accumulate in any significant amount in solution (Scheme 3).

The second substitution of Li by Rb leads to two Li\(_3\)Rb\(_2\) isomers that open a path for the internal lithium insertion (pathways A and B, Scheme 3). After insertion, the resulting \(\alpha\)- and \(\beta\)-isomers of Li\(_3\)Rb\(_2\) have an open coordination site between the two benzene rings of coronene and can be abbreviated for clarity as \(\alpha\)-Li\(_3\)Rb\(_2\) and \(\beta\)-Li\(_3\)Rb\(_2\). Subsequent addition of Li\(^+\) to this open site provides \(\alpha\)-Li\(_3\)Rb\(_3\) and \(\beta\)-Li\(_3\)Rb\(_3\) isomers, respectively. Final substitution of this loose Li\(^+\) ion by Rb\(^+\) affords the Li\(_3\)Rb\(_3\)-product (Fig. 2). Since the Li\(_3\)Rb\(_3\)-sandwich is a single product of both insertion pathways, the corresponding \(^7\)Li NMR signal at −23.92 ppm is notably growing with time (Fig. 1a–c). Three peaks at −6.21, −7.61, and −8.32 ppm can be identified in the \(^7\)Li NMR spectrum with a related triad of peaks having the same relative intensities being observed at the highly-negative region at −23.92, −24.44, and −24.86 ppm. Notably, the signals from these two triads correlate really well and therefore should belong to the same type of supramolecular sandwiches, as assigned in Fig. 2 with arrows. The triad on the right is associated with the Li\(^+\) ion internally located between the central 5-membered rings of C\(_{20}\)H\(_{10}\)\(^4\^-\), while the signals on the left stem from the Li\(^+\) ions sandwiched between the peripheral 6-membered rings of tetra-reduced coronene. The relative integration of peaks is ca. 2.6, 3.5, and 3.0 to 1 in pairs of −6.21/−23.92, −7.61/−24.44, and −8.32/−24.86 ppm, respectively. Slight overestimation of the integrated intensity may suggest that Li\(^+\) ions located at the sandwich periphery exchange with external environment. A small shoulder of the peak at −7.61 ppm may be indicative of differentiation between two peripheral Li\(^+\) ions sites observed in \(\beta\)-Li\(_3\)Rb\(_2\).

It should be mentioned here that the insertion mechanism through the \(\alpha\)-isomer of Li\(_3\)Rb\(_2\) (pathway B, Scheme 3) is much less energetically favourable. The internal lithium migration for \(\beta\)-isomer was found to be barrierless, whereas for \(\alpha\)-isomer the barrier was calculated to be +15.48 kcal mol\(^{-1}\).\(^{14}\) As a result, the corresponding NMR signals of \(\alpha\)-isomer are always much less intense, being at about 10–20% relative to the \(\beta\)-isomer. In the case of Li–K aggregates, the formation of \(\alpha\)-isomer of the Li\(_3\)K\(_2\) composition was not observed at all, thus showing that a larger Rb\(^+\) ion opens a new reaction pathway in comparison with K\(^+\).

After the Li\(_5\)-sandwich is fully consumed in the stepwise substitution reactions (as seen by the disappearance of the 6.85 ppm peak in \(^1\)H and −11.70 ppm peak in \(^7\)Li NMR spectra, Fig. 1c) the major product in solution is the Li\(_3\)Rb\(_3\) sandwich, with the \(\beta\)-isomer of Li\(_3\)Rb\(_3\) being a minor component of the mixture. Interestingly, there is a new peak appearing at −21.46 ppm which is assigned to the fully substituted product having five rubidium ions at the rim (Li\(_3\)Rb\(_3\)). Similar to the Li–K system
in solution, namely Li3Rb3 and LiRb5 constitutes approximately 20% of the reaction mixture in solution. At this point, the amount of solvent-separated Li(THF)+ species (broad peak at ca. 0 ppm) and non-solvated lithium ions bound to the external surface of C20H10-4-4(very broad, 0 to −4 ppm) is quite significant. This is also accompanied by the appearance of some precipitation-decomposition products in the NMR ampoule.

Notably, NMR monitoring shows that the α-isomer of Li4Rb2 can no longer be seen and the amount of the corresponding β-isomer is very small. At the final stage, LiRb3 is revealed as the major component in the mixture (Fig. 1d), and it finally becomes the only product observed in solution (Fig. 1e). The formation of LiRb3 is accompanied by the appearance of a single peak at −21.46 ppm in 7Li NMR and at 6.49 ppm in 1H NMR spectra. It should be mentioned here that a substantial amount of unidentified precipitate is observed in the NMR ampoule at this stage.

X-ray crystallographic studies

As revealed by NMR studies (Fig. 1a–c), the mixed metal Li–Rb sandwiches are represented by two major contributors existing in solution, namely Li1Rb3 and β-Li4Rb2. From these systems, by applying suitable crystallization techniques (see ESI†), we were able to crystallize two products, [(Rb(THF))2Li/[(C20H10)]Li(THF)]2 (2) and [(Rb(diglyme))2Li/Rb(C20H10)]2 (diglyme)2·0.5THF (3·0.5THF). Importantly, while the Li1Rb3 sandwich constitutes a close analogue of the previously reported Li1K3 product,12 the Li1Rb2-sandwich having an empty coordination site between coronulene decks is observed for the first time. To clearly indicate this open site (abbreviated as □) and the β-disposition of two adjacent rubidium ions we designate this novel sandwich as β-Li4Rb2□. The single crystal X-ray diffraction analyses revealed that geometric parameters of C20H10-4-4 anions in the Li1Rb3 and β-Li4Rb2 products (Fig. 3) are similar to those in the Li1K3 sandwiches (see Table S2†).

It should be emphasized here that the fact that the transient β-Li4Rb2□ sandwich can be isolated (Fig. 3a and c) provides compelling experimental evidence for the proposed Li-insertion mechanism. The sandwich opening (measured as an angle between the mean planes passing through the rim of two α-bowls) in the case of β-Li4Rb2□ is greater than in the Li1Rb3 or Li1K3 sandwiches (27° vs. 21° and 20°, respectively). The distance between the rim C-atoms at the sandwich opening side is about 6.6 Å, which is substantially longer than the corresponding separations in Li1Rb3 (Fig. 3b) or Li1K3 cases (6.2 and 6.1 Å, respectively).

In β-Li1Rb3□, the fourth Li+ ion (also having a coordinated THF molecule) is closely approaching the sandwich (to show this side binding, the sandwich is abbreviated as β-Li1Rb3□-Li from the Li3-triangular side (Fig. 3c)), thus forming a Li4 parallelogram with the Li⋯Li contacts of 2.86 Å inside the sandwich and of 2.92 Å outside the sandwich. This additional lithium cation brings both coronulene bowls to a very short distance of 3.8 Å (compared with 4.1 Å in Li1Rb3). In Li–Rb sandwiches in 2 and 3, both external concave cavities of C20H10-4-4 are filled with rubidium ions that participate in the formation of a 1D network through the shared external solvent molecules (Fig. S5†). In 2, these 1D chains are further packed into 2D sheets via additional C–H⋯π interactions (Fig. S6†).

Crystallization of the final product in this series of dynamic transformations, namely of LiRb3, has been very problematic. By the time this sandwich appears as the only product in solution, which is determined by its characteristic red color, there is a significant amount of precipitation-decomposition observed in the system. Removal of this unidentified solid affords rather dilute solutions, which do not allow efficient crystal growth of LiRb3. After numerous attempts we were able to isolate the single crystals of LiRb3 sandwich, but even the best crystals show diffraction only up to ca. 1 Å resolution. Nevertheless, these X-ray diffraction data allowed us to confirm the LiRb3 core structure of this product (Fig. 4), as predicted based on the stepwise alkali metal substitution mechanism and NMR data.

7Li NMR shifts and structural correlations

The record negative 7Li NMR shifts characteristic of supra-molecular Li–K and Li–Rb sandwich products formed by C20H10-4-4 anions are worthy of special discussion. Such unprecedented shifts of the central Li+ ion occupying the cavity between the two hub-sites of coronulene tetraenions (signals range from −21 ppm to −25 ppm) are due to strong shielding of this internal cation from the external magnetic field. This implies that the centrally located Li+ ion is surrounded by some electron density, which clearly separates and distinguishes it from other alkali metal ions (Li, K, or Rb) sandwiched in-between the peripheral 6-membered rings of.
C_{20}H_{10}^{4-} anions. Importantly, this provides the first indication that some electronic communication should exist between the negatively charged corannulene decks separated by an alkali metal layer in the triple-decker organometallic products. The most plausible way for such electronic exchange should be through the orbital interactions between the two tetrareduced π-bowls. This is clearly confirmed by the observation that the shift of $^7$Li NMR signal of the central Li$^+$ cation correlates with the distance between two corannulene decks in the isolated sandwich structures.

In Li$_3$K$_3$, the central Li$^+$ ion resonates at –24.48 ppm but the signal is shifted to ca. –22.40 ppm in Li$_3$K$_5$ (the separations between the hub-sites of two C$_{20}$H$_{10}^{4-}$ decks are 3.8 and 4.0 Å, respectively). The centrally located Li$^+$ ions in Li$_3$Rb$_3$ and LiRb$_5$ products appear to be deshielded (–23.92 and –21.46 ppm) in comparison with their Li$_3$K$_3$ and LiK$_5$ analogues, which is consistent with the size of Rb$^+$ ions requiring more space than K$^+$ ions. As a result, the corresponding separations between two corannulene bowls in Li$_3$Rb$_3$ and LiRb$_5$ are 4.1 and 4.2 Å, respectively. Clearly, Rb$^+$ ions move the charged decks farther apart and thus weaken communication between two π-bowls in the triple-decker aggregates. Consequently, the central Li$^+$ cation becomes less shielded and the corresponding $^7$Li NMR signals shift downward for sandwich structures having higher amount of heavier alkali metals.

The close analogy can be drawn with the strong shielding effect of the C$_{60}$ fullerene hexaanion. The $^3$He atom inside the C$_{60}$'6- cage was found to be significantly more strongly shielded (by ca. 20 ppm) than any other previously reported encapsulated $^3$He atom, suggesting the ability of electrons to move freely about the surface of a charged spheroidal π-system. In the case of mixed alkali metal sandwich-type assemblies formed by highly charged corannulene, electrons can also move freely about the π-bowl surface of $^1$-$^1$, but in addition to that, they are able to move from one surface to the other, as we found here for the first time.

DFT computational studies

To get further theoretical insights into the experimentally observed electronic communication between the highly charged corannulene anions, we carried out DFT calculations at the PBE0/def2-TZVPP(Li, Rb)//cc-pVDZ(C, H) level of theory. The choice of the model system was discussed elsewhere.

Indeed, theoretical calculations revealed that π-systems of two tetrareduced corannulene bowls are coupled with each other, forming shared electronic density that is mainly concentrated in the region between two 5-membered rings. The central lithium cation is thus wrapped into a negatively charged cocoon, which seems responsible for a record high shielding effect observed in $^7$Li NMR. The corresponding orthogonal molecular orbitals representing delocalization of electronic density between the C$_{20}$H$_{10}^{4-}$ bowls are shown in Fig. 5.

Theoretical analysis of components of shielding tensor revealed that the main difference between $^7$Li NMR signals of the lithium ions placed in-between six-membered rings and of the central Li$^+$ ion comes from the paramagnetic term of the Ramsey formula ($\sigma_{\text{tot}} = \sigma^p + \sigma^d$). The paramagnetic term involves mixing of ground and excited states of the molecule due to the magnetic field, and it is rather sensitive to the molecular electronic structure. Thus, the main reason for such a large shielding of the central Li$^+$ ion, in comparison with those located between six-membered rings of C$_{60}$H$_{10}^{4-}$, should arrive from the electronic structure of the sandwich and, more precisely, from the local electronic environment of the cation. MOs presented in Fig. 5 are found to provide the largest contributions to the $\sigma^p$ component of the shielding tensor. These MOs clearly show extended delocalization around the central Li$^+$ ion and no delocalization for the lithium(s) sitting between peripheral 6-membered rings. At the same time, the calculated charges of lithium cations of both types indicate essentially no charge transfer from the highly negatively charged corannulene tetraanions to Li$^+$ ions. The NBO charges are very similar (+0.88 and +0.91, respectively, in Li$_3$Rb$_3$ and +0.87 in LiRb$_5$). The larger distance between two bowls makes

Fig. 5 Canonical MOs, HOMO-10 (left) and HOMO-11 (right) in Li$_3$Rb$_3$ (a) and LiRb$_5$ (b) systems, showing coupling and sharing electron density between two tetrareduced corannulene bowls. MOs are presented with an isosurface of 0.03 a.u.
the coupling between them weaker, as suggested from experimental data. Consequently, this results in a downfield-shift of the $^7$Li NMR signal from the central Li$^+$ ion. The corresponding component of the shielding tensor was found to show the trend that is opposite to the distance between two C$_{24}$H$_{10}^{4-}$ anions. Thus, the signal from the central lithium cation could be used for qualitative evaluation of the strength of coupling between the charged bowls in sandwich-like aggregates.

Conclusions

Several remarkable triple-decker organometallic aggregates having a mixed metal core (Li$_x$Rb$_y$, Li$_x$Rb$_z$, and LiRb$_z$) sandwiched between two tetra-reduced corannulene decks have been isolated in this work, following the NMR studies of their dynamic transformations in solutions. These sandwiches have the central cavity located in between of the hub-sites of two C$_{24}$H$_{10}^{4-}$ decks occupied by an internal Li$^+$ ion that exhibits the record high negative shielding (up to $-25$ ppm) in $^7$Li NMR spectra. Theoretical investigation of these unique systems revealed that coupling of two highly-charged corannulene bowls results in a shared region of high negative electron density around the central lithium ion that is responsible for a record shielding effect observed in $^7$Li NMR spectra. Analysis of three new sandwich structures allowed us to identify the trend: the larger the negative chemical shift of the corresponding $^7$Li NMR signal from the internally encapsulated Li$^+$ ion. Consequently, these $^7$Li NMR signals can be used for qualitative evaluation of the strength of coupling between the charged carbon bowls in sandwich-like supramolecular aggregates. The higher the negative chemical shift is the stronger coupling should be expected.

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Notes and references


24 Energetic evaluation was performed with the help of xDH-PBE0 functional for PBE0-optimized geometries (see ESI† for more details). Correlation-consistent full-electron basis sets of double-ζ quality (cc-pVDZ) were applied for description of light elements (C and H), while def2-TZVP basis sets were applied for all metal atoms (combined with effective core potential for Rb). Importantly, for the α-isomer two different migration processes were observed. Here, we consider only one having the lowest barrier (Scheme 3).