Structural comparisons of the species of recognition of Ni$^{2+}$, Cu$^{2+}$, Zn$^{2+}$, Cd$^{2+}$, Hg$^{2+}$ and Ag$^{+}$ with 1,3-di-conjugates of calix[4]arene by DFT computations

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Calixarenes are versatile macromolecular systems, which would exhibit better host-guest chemistry upon appropriate derivatization. Our group has been involved in the synthesis of a variety of calix[4]arene conjugates, which were found to selectively recognize metal ions. Though such recognitions have been demonstrated, based on different experimental studies, the qualitative and semi-quantitative understanding of the coordination aspects are still lacking owing to the non-availability of the crystal structures of the species of recognition. Hence, in order to understand the coordination chemistry of the interactions between the host calix[4]arene receptor and the guest metal ion species, computational modelling studies have been carried out. In this regard, five conjugates of calix[4]arene, which are 1,3-di-O-derivised at the lower-rim and appended over an amide as well as Schiff base linker have been studied. The computational modelling studies have been carried out at the level of density functional theory to yield coordination geometries, which are in tune with the experimental observations and comparable to those reported in the literature in the case of similar receptor systems. The species of recognition obtained with the metal ions are found to have some structural resemblances with the metal sites present in metalloproteins. The modelling studies and the coordination geometries are discussed in this article.

Keywords: Density functional calculations, Calix[4]arene conjugates, Macromolecules, Species of recognition, Coordination core, Computational modelling

Calix[4]arenes are macrocyclic molecules made of ‘n’ number of phenol units connected by ortho methylene bridge centers. Calix[4]arenes are the simplest, and most common examples of this family, with four phenolic residues being present in the macrocyclic ring that provides a platform for organic derivatization. Reactions of simple calix[4]arene with transition metal complexes can produce metal phenolates with substitution of one to four hydrogen atoms. Derivatives of calix[4]arenes have been extensively studied in the recent times, for their conformational flexibility, derivatizable and complexing ability. Thus, calix[4]arene conjugates have been exploited in a variety of chemical applications, including catalysis, host-guest chemistry, separation chemistry, selective ion transport, and sensors. Calix[4]arene can exist in four conformations designated as cone, partial cone, 1,2- alternate, and 1,3-alternate. It is well documented that the introduction of bulky substituents at the lower rim will impart conformational immobilization to these derivatives. Any of these conformations represents a special three-dimensional arrangement with different complexation behaviours and hence with potential applications as a molecular scaffold and advantageously provide useful building blocks in supramolecular chemistry.

It is known in the literature that the calixarene-conjugates are expected to form host-guest complexes by exhibiting a variety of ensembles. Apart from the experimental determination of these conjugates for their applicability, the structural features of the complexes formed are also important for understanding the details of binding. In the absence of crystal structures, there exists a dire need to model the structure of the species of recognition or the complexation with the experimental clues available to improve the understanding. In this regard, computational modelling is one of the useful and affordable means of predicting the structure. In recent years, computational investigations of these compounds have revealed that calix[4]arenes provide a congenial platform for the design of new systems with improved properties, and also to interpret their chemical and physical properties. During the past few years, our group has been actively involved in the design and development of the conjugates of calix[4]arene to sense ions of biological and
Environmental importance by selectively bringing about synthetic modifications such that the resultant conjugates possess suitable binding core(s). The focus of the present article is the computational modeling of the species of recognition of these calixarene conjugates towards different metal ions of biological importance to provide structural features for such selective recognition.

Methodology

Computation and visualisation tools

All geometry optimization calculations of the complexes of the conjugates of calixarenes (species of recognition) were carried out using Gaussian 98 or 03. The visualization and input preparations were done using chemcraft or GaussView softwares.

The quantum computational calculations were carried out in a cascading fashion, in case of geometry optimization, beginning from a primitive semi-empirical method followed by an ab-initio HF method and then to a density functional theory based on B3LYP method. This sequence can be easily understood from the representation, i.e., AM1 or PM3 → HF/STO-3G → HF/3-21G → HF/6-31G → B3LYP/3-21G → B3LYP/6-31G. Such a cascade methodology has been used throughout the present paper, unless otherwise mentioned. This is expected to minimize errors that may crop in if the crude model is directly taken to DFT methods, without going through this sequence of computations.

Semi-empirical methods

The Austin Model 1 (AM1) method was considered as the initial step for carrying out the computational modelling studies. The coordinates of the model generated or the one which is taken from the single crystal XRD structure are the first geometry minimised using this method. A full optimisation was carried out by using ‘FOPT’ keyword available without altering the default convergence criteria.

Parametric method 3 (PM3) was used when the AM1 method did not give appreciable results. In most of the cases the PM3 method was used as the starting level of computational calculations and was used for full optimization procedure too.

Ab initio method

Hartree-Fock (HF) level of theory was used for all the geometry optimization calculations. This level of calculations was used for those models which were obtained from the semi-empirical calculations. The basis sets used in this are, STO-3G, 3-21G and 6-31G.

Density functional theory method

B3LYP method was used along with a set of basis sets, 3-21G, 6-31G with diffusion and polarisation functions involved wherever required. Also, ECP basis sets like CEP-121G, SDD, LanL2DZ were used for the metal ions wherever required.

Computational calculations

The initial guess structures used for computational calculations of L₁, L₂, L₃, and L₄ were taken from the single crystal XRD data, whereas in the case of L₅, the initial model was prepared from the published data of copper complex with the following modifications: (a) removing the copper center, (b) replacing tert-butyl moiety by hydrogen, (c) protonating the phenolate moieties, and, (d) introducing a -CH₂OH moiety ortho to the phenolic OH of the Schiff base part of each of the arm. In order to reduce the computational times, cut short models of L₁, L₂, L₃ and L₄, viz., L₁’, L₂’, L₃’ and L₄’, were made by simply replacing each ‘t-butyl’ group by ‘H’ and the benzimidazole by a simple imidazole moiety in the case of L₄. Such modifications are not expected to affect either the conformation or the platform or the binding core to any significant extent and hence used in the computational calculations. Subsequently, the computations were carried out with the unmodified models also and no change was found in the conformation or in the binding core as demonstrated herein. The optimization of the metal complexes were carried out with the structures obtained at B3LYP/6-31G and deprotonated at both the phenolic centers of the Schiff base of the arms wherever required. The metal ion was placed well away from the binding core and allowed to minimize.

Results and Discussion

During the past several years, our group has synthesized different 1,3-di-conjugates of calix[4]arene (Scheme 1) and demonstrated their selective recognition towards ions, viz., Ni²⁺, Cu²⁺, Zn²⁺, Ag⁺, Cd²⁺ and Hg²⁺ by using a variety of experimental methods. Since the formation of single crystals of the species of recognition has been the limiting step, structures of such complexes could not be established. However, the presence of crystal structure of either the receptor conjugate or its precursor was a boon to start the computational studies. Therefore, the structural features of the species of recognition of the complexes of the
conjugates of calix[4]arene have been presented based on computational methods carried out at the levels of DFT.

**Reducing computational times**

As can be seen from the Scheme 1, the conjugates of calix[4]arenes studied were all large in size with the total number of atoms ranging from ~150 to 200. All of these possess t-butyl moieties at their upper rim, while the recognition or binding is through the binding groups present on the arms at the lower rim. Therefore, chopping of these four t-butyl groups will bring down the size of the conjugates to two-thirds of its original size. This indeed saves a lot of computational time without changing the conformational features of the arms which in turn provides the binding core. In order to keep the valency satisfied for the carbon center from which the corresponding t-butyl group was chopped off, the same was attached with hydrogen. Such a process yields not only a conformational invariant but also a molecule which is chemically intact. Hence, a number of computational calculations were carried out using such truncated conjugates of calix[4]arene as reported in this paper. All the receptor systems, viz., L1, L2, L3, L4 and L5 were subjected to this truncation and the resultant conjugates, viz., L1’, L2’, L3’, L4’ and L5’ were used for the computations. However, in the case of L1, an additional truncation was done by converting the benzimidazole to a simple imidazole moiety. Thus, all the truncations carried out on the original conjugate were balanced for their valencies by attaching the requisite number of H atoms. Some of the calculations were carried out with both the truncated as well as the original conjugate, and no significant change was found in the conformation of the arms as well as in the binding features. The cone conformation has been maintained throughout.

**Cascade approach used in the computations**

All the computations carried out and reported in this paper have been done through a cascade methodology. In this approach, a well-optimized receptor geometry, which in turn is obtained from the crystal structure of the receptor, is taken initially in the semi-empirical methodology, and the outcome of this has been taken to the next higher level of calculations. Such computations are continued in a cascading manner by climbing to higher levels of calculations smoothly by going through AM1 or PM3 → HF/3-21G → HF/6-31G → B3LYP/3-21G → B3LYP/6-31G, etc. The metal ion is introduced in this cascade process at an appropriate stage and then the cascade taken further.

**Hg$^{2+}$ recognition by L1**

Based on our recent studies, L1 has been found to be sensitive and selective in the recognition of Hg$^{2+}$ in aqueous acetonitrile (1:1) based on fluorescence spectroscopy.$^{5h}$ The stoichiometry of the complex
species formed during the titration was found to be 1:1 based on both the ESI MS and the absorption spectral studies. The selectivity of \(L_1\) towards \(\text{Hg}^{2+}\) has been demonstrated in comparison with eleven different and biologically relevant \(\text{M}^{2+}\) ions, viz., \(\text{Mn}^{2+}, \text{Fe}^{2+}, \text{Co}^{2+}, \text{Ni}^{2+}, \text{Cu}^{2+}, \text{Zn}^{2+}, \text{Cd}^{2+}, \text{Hg}^{2+}, \text{Pb}^{2+}, \text{Ca}^{2+}\) and \(\text{Mg}^{2+}\), including those of the mercury group. None of these ions negatively influences the recognition of \(\text{Hg}^{2+}\) by \(L_1\). The influence of solvent polarity on the recognition of \(\text{Hg}^{2+}\) has also been spectrally demonstrated. The species of recognition has been addressed by DFT computations as reported in this paper.

\(L_1\) and \(L_1'\)

Both \(L_1\) and \(L_1'\) were optimized as explained in the methodology section and the corresponding structures obtained from B3LYP/6-31G are shown in Fig. 1. The optimized structures of \(L_1\) and \(L_1'\) exhibit almost similar changes in the dihedral angles (as shown in Fig. 1) of the arms when compared to that in \(L_1\) obtained from the single crystal XRD structure. The dihedral angles indeed confirmed the retainment of the arm-conformation on going from \(L_1\) to \(L_1'\). The hydrogen bond present between the two imidazoles of the \(L_1\) was found to be intact even after the geometry optimization though minimal changes were observed in the metric data. The predicted hydrogen bond stabilization energy was obtained by mapping the energy from the H...A distance correlation plot reported in the literature. It has been found that \(L_1\) (XRD), \(L_1\) (OPT) and \(L_1'\) (OPT) form a hydrogen bond with stabilization energies of -5.6, -4.6 and -5.3 kcal/mol respectively (where OPT refers to the optimized). The OPT geometries of both \(L_1\) and \(L_1'\) showed comparable changes in the dihedral angles, revealing minimal effect on crafting the system. The ground state existence of the OPT geometries were checked by performing a vibrational state analysis and found that both the geometries had no imaginary frequencies.

\(\text{Hg}^{2+}\) complex

The B3LYP/6-31G optimized geometries of \(L_1'\) and \(L_1\) were taken as the starting structures for modelling their \(\text{Hg}^{2+}\) complexes. The starting geometry was modelled by placing the \(\text{Hg}^{2+}\) ion away from the binding core of \(L_1/L_1'\) at a non-interacting distance, without disturbing the existing hydrogen bond. No attempt was made to deprotonate the imidazole proton of \(L_1\) or \(L_1'\) as the NMR spectra showed this proton intact. The geometry optimization of the modelled complex was carried out using B3LYP. The basis set used for \(\text{Hg}^{2+}\) was a compact effective potential basis set build by Stevens/Basch/Krauss, having a double and triple zeta, viz., CEP-31G followed by CEP-121G. The trail computations with CEP-31G or CEP-4G prior to the usage of CEP-121G did not yield any favourable structures as mentioned in the literature. The OPT structure of the \(\text{Hg}^{2+}\) complexes of \(L_1\) and \(L_1'\) are shown in Fig. 2. The ground state existence of the optimized geometry has been shown by performing vibrational state analysis.

The OPT \(\text{HgL}L_1'/\text{HgL}_{1}\) complex showed the existence of a linear coordination around Hg with two of the imidazole nitrogens being bound, forming a species of the type N2...Hg...N5 with an average Hg...N distance of ~2.09 Å and N...Hg...N angle of 178.6° (Fig. 3(a&b)). Such linear coordination about \(\text{Hg}^{2+}\) is indeed common in the literature. In addition to the interactions found with the imidazole N, a stabilizing interaction was observed between the \(\text{Hg}^{2+}\) and the two amide oxygens, wherein the Hg...O distance is ~2.7 Å. Such stabilizing interactions
were reported in the literature with different types of nitrogens in the range of 2.04 - 2.85 Å and oxygens in the range 2.57 - 2.95 Å.

The coordination complexes formed by \( \text{L}_1 \) as well as \( \text{L}_1' \) with Hg\(^{2+} \) were almost the same, except for marginal variations in the metric data (shown in Fig. 3(a&b)). It has been reported in the literature that Hg\(^{2+} \) can exhibit interactions with groups or moieties other than N, O and S, viz., with aromatic rings, aliphatic groups, etc.\(^{34} \) The overall coordination around Hg, viz., the two nitrogens and the two amide oxygens, resemble an octahedral geometry with two vacant sites as depicted in the Fig. 3(a&b).

### Recognition of Zn\(^{2+} \) and Ni\(^{2+} \) by L\(_2\)

L\(_2\) was found to selectively recognize Zn\(^{2+} \) (switch-on) and Ni\(^{2+} \) (switch-off) by showing changes in the fluorescence intensity by forming 1:1 complex.\(^5e \) The binding studies of ions with L\(_2\) and the composition of the complex formed have been addressed by various experimental studies, viz., steady state and time resolved fluorescence spectroscopy, absorption spectroscopy and ESI MS. It was also found that L\(_2\) can detect Zn\(^{2+} \) and Ni\(^{2+} \) to a minimum concentration of 142 and 203 ppb respectively, making L\(_2\) an efficient molecular probe for these ions. L\(_2\) has also been experimentally found to recognize both these metal ions simultaneously in a cooperative manner, viz., Ni\(^{2+} \) and Zn\(^{2+} \). The experimentally observed fluorescence intensity of the complex [ZnL\(_2\)] decreases when titrated against Ni\(^{2+} \) and increases when [NiL\(_2\)] was titrated against Zn\(^{2+} \). These results clearly suggest the cooperative binding of Zn\(^{2+} \) and Ni\(^{2+} \) to L\(_2\), immaterial of whether the L\(_2\) was initially loaded with Zn\(^{2+} \) or with Ni\(^{2+} \).

Thus, the mononuclear (viz., Ni\(^{2+} \) or Zn\(^{2+} \)) and the dinuclear (viz., both Ni\(^{2+} \) and Zn\(^{2+} \)) species of recognition were modelled by DFT calculations.

### L\(_2\) and L\(_2'\)

Geometry optimization on both L\(_2\) and L\(_2'\) were carried out as mentioned under methodology and the structures of DFT (B3LYP with 6-31G basis set) optimized L\(_2\) and L\(_2'\) are shown in Fig. 4. The optimization of L\(_2\) and L\(_2'\) resulted in similar structures and are different from that of the crystal structure. The ground state existence of the OPT geometries of both L\(_2\) and L\(_2'\) were checked by performing a vibrational state analysis. When arm-to-arm dihedral angle comparison was made (Fig. 4), it was found that the dihedral angles were very similar among both the optimized structures (L\(_2\) and L\(_2'\)), though these were considerably different from that observed in the crystal structure. The orientation of the pyridyl groups present in the single crystal structure undergoes appreciable changes upon
optimisation to bring the nitrogens in a position that is favourable for the metal ion binding. The formation of favourable metal ion binding core can be quantified by the decrease observed in the inter arm N…N distances, viz., from 6.45–11.48 Å (XRD) to 3.46–8.96 Å (optimized structure).

Mononuclear Ni$^{2+}$ or Zn$^{2+}$ complex of L$_2'$$^\text{'$}$$^\text{'}$

The L$_2'$$^\text{'$}$$^\text{'}$ that was obtained from DFT calculations was taken as the initial structure for the computational modelling of the complexes. The model for the complex was generated by placing either Ni$^{2+}$ or Zn$^{2+}$ at a non-interacting position well above the pyridyl core of the L$_2'$$^\text{'$}$$^\text{'}$ and maintaining the overall charge on the complex to be +2. The computational calculations were carried out for both the [NiL$_2'$$^\text{'$}$$^\text{'}$]$^{2+}$ and [ZnL$_2'$$^\text{'$}$$^\text{'}$]$^{2+}$ complexes at B3LYP/6-31G(d) level. The optimized structures of these complexes are shown in Fig. 5(b&c) respectively. The complexation in the case of L$_2'$$^\text{'$}$$^\text{'}$ is a result of the metal ion induced conformational changes brought about in the pendant arms of L$_2'$$^\text{'$}$$^\text{'}$ so that the core possessing the ligating atoms is well poised for binding.

In case of [NiL$_2'$$^\text{'$}$$^\text{'}$]$^{2+}$ complex, the Ni$^{2+}$ exhibited a distorted octahedral geometry with one of the coordination sites being vacant by forming covalent bonds with all the four pyridyl nitrogens along with one weak bond being formed with one of the lower rim phenolic-OH (Fig. 5b). The distortion in the Ni$^{2+}$ geometry can be clearly gauged from the angles observed in the coordination sphere, viz., 83–115° (as compared to 90°) and 155–160° (as compared to 180°) (Fig. 6a). A literature report$^\text{35}$ shows the Ni–N distance to be 1.84–2.05 Å or more, which is similar to that found in the present case, viz., 1.91–2.42 Å.

In the case of [ZnL$_2'$$^\text{'$}$$^\text{'}$]$^{2+}$, the Zn$^{2+}$ exhibits a distorted tetrahedral geometry by binding through all the four pyridyl nitrogens (Fig. 5c), similar to that found in the case of Ni$^{2+}$ complex but in a tetrahedral geometry. The angles observed in the coordination sphere, viz., 95–134°, clearly suggests a considerable distortion in the tetrahedral geometry (Fig. 6b). It has been reported in the literature that the Zn…N distances$^\text{36}$ are in the range of 2.03–2.48 Å.
Hetero-binuclear complex of [Zn-L₂''-Ni]⁴⁺

Based on the experimental results of simultaneous recognition of Ni²⁺ and Zn²⁺ by L₂ in a cooperative fashion, an attempt to model a hetero-bimetallic complex of this conjugate was initiated. Geometry optimization studies in the presence of both the cations was carried out with the optimized Zn²⁺ complex of L₂' followed by placing Ni²⁺ at a non-interacting position above Zn²⁺. To reduce the computational time required for these calculations further, L₂'' was derived from L₂', by replacing the phenyl rings of the calixarene by hydrogens, taking care of the hybridization at each center appropriately. The optimizations resulted in the formation of a new complex. In this new complex, while the Ni²⁺ occupies the N₄ core of the pyridyls, that of the Zn²⁺ has been pushed to the O-rich core of the lower rim by forming a distorted trigonal bipyramidal geometry (Fig. 7). The Ni…N and Zn…O bond distances were found to be in the range of 1.90 – 2.11 Å, which is similar to that reported in the literature.³⁵,³⁶

Cu²⁺ and Ag⁺ recognition by L₃

L₃ was studied for its metal ion binding properties towards various biologically relevant M⁺ transition/alkali/alkaline earth ions by fluorescence and absorption spectroscopy in two different solvent systems. L₃ was found to detect Cu²⁺ (switch-on fluorescence at 315 nm) and Ag⁺ (ratiometrically switch-on fluorescence at 445 nm) selectively in pure and aqueous methanol media.⁵⁵,⁵⁶ The stoichiometry of the copper or the silver complex has been found to be 1:1, based on the Job’s plot and was further confirmed by ESI-MS. The corresponding Ag⁺ and Cu²⁺ complexes were modelled by computational studies as reported here.

L₃ and L₃'

Both L₃ and L₃' were geometry optimized as given under Methodology and the corresponding structures are shown in Fig. 8. The geometry optimization of the L₃ and L₃' showed similar orientation of the arms, but both showed appreciable changes in the dihedral angles at the arms when compared to the angles present in the crystal structure. The distance between the pyridyl nitrogen in the intra arms as well as the inter arm of L₃ were found to change from 4.88 – 5.39 Å (with an average of 3.73 Å) in the case of intra arm, and 5.09 – 8.87 Å to 4.66 – 5.50 Å in the case of inter arms, upon optimization. The data clearly suggest the formation of a favourable binding core by the pyridyl moieties for metal ion by bringing
the nitrogen centers of these into a coordinating distance. The variations occurred in the conformation of the arms of $L_3/L_3'$ as compared to the single crystal structure as can be seen from the Fig. 8.

**Cu$^{2+}$ or Ag$^+$ complex of $L_3'$**

The DFT optimized geometry of $L_3'$ was taken as the initial structure for generating models for the copper and silver complexes. The model was generated by placing the Cu$^{2+}$ or Ag$^+$ at a non-interacting distance, well above the pyridyl-nitrogen core of $L_3'$. The models were then geometry optimized to the highest level of B3LYP with 6-31G basis set in the case of copper complex and SDD basis set in the case of silver complex. The overall charge of the complexes were maintained as $[\text{Cu}L_3']^{2+}$ and $[\text{Ag}L_3']^+$ respectively. The OPT structures of these complexes are shown in Fig. 9(b&c) respectively for $[\text{Cu}L_3']^{2+}$ and $[\text{Ag}L_3']^+$.

In the case of $[\text{Cu}L_3']^{2+}$ complex, the Cu$^{2+}$ site exhibits a distorted tetrahedral geometry extending its four coordinations to four of the pyridyl nitrogens present. The distortion in the geometry can be measured from the coordination angles observed at the copper center, viz., 89.3 - 141.8° (as compared to 109°7′). The coordination bond lengths and bond angles are given in the caption of Fig. 10. Such a highly distorted tetrahedral center was observed for Cu$^{2+}$ in blue copper proteins, viz., plastocyanin.\(^\text{37}\) In case of the protein, the presence of such distorted copper center has been interpreted to its ease
to undergo redox process during the electron transfer activity. Comparable geometric parameters have also been observed in the literature about a copper center.38

In the case of [AgL₃]⁺ complex, Ag⁺ also exhibits a distorted tetrahedral geometry, extending its four coordination through the four pyridyl nitrogens. The extent of distortion present in the [AgL₃]⁺ complex is almost comparable to that of the copper complex [CuL₃]²⁺ (Fig. 11). Similar coordination geometry has been found in the literature.39

Recognition of Cu²⁺ by L₄

The receptor molecule, L₄ exhibited selectively the fluorescence quenching towards Cu²⁺ alone among the eleven divalent ions studied.5d The 1:1 stoichiometry of the complex formed between L₄ and Cu²⁺ has been demonstrated by electronic absorption and ESI-MS. The structural features of the Cu²⁺ complex of L₄ have been established by DFT.

Optimization of L₄'

L₄' was optimized as explained earlier and the corresponding structure obtained from HF/3-21G calculation is shown in Fig. 12. The optimized structure of L₄' exhibits no significant change in the dihedral angles of the arms as compared to the single crystal XRD structure of L₄. The hydrogen bond present between the lower rim phenolic core of the L₄' was found to be intact even after the geometry optimization. The optimized geometry of L₄' showed no comparable changes in the dihedral angles of the arms as compared to that of the single crystal X-ray structure of L₄, revealing minimal or no effect on replacing the t-butyl groups by hydrogen atoms.

Cu²⁺ complex of L₄'

The B3LYP/6-31G* optimized structure of L₄' was taken as the starting structure for modeling the Cu²⁺ complex. The optimization of the complex was carried out by simply placing the Cu²⁺ far above the binding core so that initially there are no interactions between L₄' and Cu²⁺. The minimization of the modeled complex was carried out using B3LYP/6-31G*. The optimization resulted in a four coordinated distorted trigonal bipyramidal Cu(II) complex with one vacant site. The resultant structure was further allowed to fill the vacant coordination by an acetonitrile. The optimization carried out in the
presence of acetonitrile in B3LYP/6-31G* resulted in a complex where Cu$^{2+}$ exhibits a distorted trigonal bipyramidal geometry wherein each arm of L$_4'$ acts as bidentate in providing a total of four coordinations, with the fifth coordination coming from the acetonitrile, resulting in NO$_2$S$_2$ binding core (Fig. 13(a&b)). Similar coordination geometry and bond distances were observed for a Cu(II) complex of 1,6-bis(2-benzimidazolyl)-2,5-dithiahexane$^{40}$.

**Differential recognition of Zn$^{2+}$ over Cd$^{2+}$ by L$_5$**

L$_5'$ showed selectivity toward Zn$^{2+}$ by switch-on fluorescence among the 12 metal ions studied.$^{3c}$ The interaction of Zn$^{2+}$ with L$_5$ has been further supported by absorption studies, and the stoichiometry of the complex formed (1:1) has been established on the basis of the absorption and ESI MS. The receptor also showed fluorescence enhancement with Cd$^{2+}$ but this is much less than that exhibited by Zn$^{2+}$. The differential recognition of L$_5$ towards Zn$^{2+}$ over Cd$^{2+}$ has been computationally established.

L$_5'$ was optimized as before and the structure obtained from B3LYP/6-31G is shown in Fig. 14. The OPT structure observed at B3LYP/6-31G level of calculations showed well ordered arms in an extended fashion wherein each arm is stabilized through phenolic O-H···N hydrogen bond interaction (intra-arm). In addition, one of the arms extends a weak hydrogen bond, viz., alcoholic O-H···O (phenolic). Further, the two arms are connected through a hydrogen bond formed between their alcoholic-OH groups (inter arm), as can be seen from Table 1. The hydrogen-bond interactions observed at the phenolic-OH groups indicates cone conformation; the inter arm interactions also supports the conformation.

**Zn$^{2+}$ complex of L$_5'$**

To obtain the Zn$^{2+}$ complex, the structure of L$_5'$ obtained at B3LYP/6-31G level was deprotonated at both the phenolic centers of the Schiff base of the arms to form deprotonated L$_5'$. The metal ion was placed well above the arms of the deprotonated L$_5'$ and allowed to minimize. The minimization in the presence of metal ion was carried out in a cascade fashion starting from PM3 → HF/STO-3G → HF/3-21G → HF/6-31G → B3LYP/3-21G → B3LYP/6-31G → B3LYP/LanL2DZ. Such minimizations resulted in formation of the Zn$^{2+}$ complex by breaking the hydrogen-bond interactions present between the two arms. The Zn$^{2+}$ interacts primarily at the Schiff base core (Fig. 15) and showed distorted four-coordination geometry bonded through two imine nitrogens and two Schiff base phenolate.

### Table 1 – Hydrogen bond data for L$_5'$

<table>
<thead>
<tr>
<th>Hydrogen bond</th>
<th>H...A (Å)</th>
<th>D...A (Å)</th>
<th>D-H...A (°)</th>
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<tr>
<td>O6-H30...N2</td>
<td>1.479</td>
<td>2.467</td>
<td>150.0</td>
</tr>
<tr>
<td>O8-H46...O6</td>
<td>1.828</td>
<td>2.641</td>
<td>137.3</td>
</tr>
<tr>
<td>O7-H41...O8</td>
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<tr>
<td>O4-H13...N1</td>
<td>1.618</td>
<td>2.545</td>
<td>147.7</td>
</tr>
</tbody>
</table>

Fig. 13 – (a) B3LYP/6-31G* optimized structure of [CuL$_4'$]$^{2+}$, and (b) coordination geometry around Cu$^{2+}$. [The bond distances (Å) and bond angles (°) around Cu$^{2+}$ in the coordination sphere are: Cu-N5 = 1.922, Cu-S1= 2.493, Cu-S2 = 2.522, Cu-O1 = 2.139 and Cu-O5 = 2.095 Å; and S1-Cu-S2 = 168.2, N5-Cu-O1 = 124.1, N5-Cu-O5 = 128.5, O1-Cu-O5 = 107.4, S1-Cu-N5 = 96.2, S1-Cu-O1 = 77.3, S1-Cu-O5 = 95.2, S2-Cu-N5 = 95.5, S2-Cu-O1= 95.5, S2-Cu-O5 = 77.9 ].

Fig. 14 – B3LYP/6-31G-optimized structure of L$_5'$ stabilized through both the intra- and inter-arm hydrogen bonds.
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Fig. 15 – Space filling models of (a) B3LYP/6-31G optimized structure of \( L_5' \), (b) B3LYP/LanL2DZ optimized structure of \([\text{ZnL}_5']\), and, (c) coordination core of \([\text{ZnL}_5']\). [The unfilled circle ‘X’ represents vacant site. The metric data for \([\text{ZnL}_5']\): N1-Zn = 2.071, N2-Zn = 2.072, O6-Zn = 1.979 and O4-Zn = 1.979 Å; N2-Zn-N1 = 161.7, N2-Zn-O6 = 91.3, N2-Zn-O4 = 94.8, N1-Zn-O6 = 94.8, N1-Zn-O4 = 91.3 and O6-Zn-O4 = 141.5].

oxygens. The Zn-O and Zn-N distances were found to be 1.979 and 2.072 Å, respectively, which are found commonly in the literature. The coordination angles observed with the Zn\(^{2+}\) core provide a better fit if it were to be considered as a trigonal bipyramidal with one trigonal center being vacant, where the trans-angle\(^{41}\), viz., N1-Zn-N2, is found to be 162°.

\(\text{Cd}^{2+}\) complex of \( L_5' \)

The optimization of Cd\(^{2+}\) complex of \( L_5' \) was carried out with the structure of \( L_5' \) obtained at B3LYP/6-31G deprotonated at its both the phenolic centers of the Schiff base of the arms. The metal ion was placed well above the arms of the deprotonated \( L_5' \) and allowed to minimize. The minimization in the presence of the metal ion was carried out in a cascade fashion starting from HF/STO-3G → HF/3-21G → B3LYP/3-21G → B3LYP/LanL2DZ. Such minimizations resulted in the formation of Cd\(^{2+}\) complex by breaking the hydrogen-bond interactions present between the two arms. The Cd\(^{2+}\) primarily interacts at the lower rim phenolic core and shows a distorted six coordination geometry bonded through two imine nitrogens, two Schiff base phenolic oxygens, and two lower rim phenolic groups (Fig. 16). The Cd-O and Cd-N distances were found to be 2.229-2.445 and 2.323-2.377 Å, respectively and are in agreement with what is commonly found in the literature.\(^{42}\) The coordination angles observed with the Cd\(^{2+}\) core provide a better fit if it were to be considered as a seven-coordinated capped octahedral structure with one vacant site.

Conclusions and Correlations

The experimentally observed species of recognition has been computationally modelled in a cascade fashion from semi empirical to DFT level, via, several stages of calculations. These are done with six different ions using five different conjugates of calix[4]arene. Important features of the species of recognition are given in Fig. 17 for appropriate comparison for all these cases. Since the coordinating atoms and geometry of the metal center are important for the recognition process, all such features can be seen on a comparative basis from this figure. As can be seen from Fig. 17, the binding cores vary from N\(_2\) to N\(_2\)O\(_x\)X, N\(_2\)O\(_2\)X\(_2\), O\(_3\), N\(_2\)O\(_2\) to NO\(_2\)S\(_2\), N\(_2\)O\(_4\)X, N\(_2\)O\(_4\), NO\(_3\) (X = vacant coordination site), while the geometries vary from linear to distorted tetrahedron (T\(_d\)) to a situation that is in between distorted tetrahedron and distorted square planar (SP) to distorted trigonal bipyramid (TBP) to distorted octahedral (O\(_h\)) with vacant site(s), distorted square
Fig. 17 – Species of ion recognition of calix[4]arene conjugates. \([\text{Oh} = \text{octahedral}; \text{Td} = \text{tetrahedral}; \text{TBP} = \text{trigonal bipyramidal}; \text{SP} = \text{square planar}]\).
pyramidal to distorted octahedral (O₈) to capped octahedral with one vacant site. Thus, a large variation is found both in the binding core as well as in the geometry, and hence, the arms are able to sense ions with diverse coordination preferences.

Most of these geometries were found to be distorted owing to their wide range of angles in the coordination sphere. Indeed several metal ion cores present in metalloproteins also possesses highly distorted geometries, since such distortions favour the function and/or reactivity. A vivid example of this is the Cu(II) center present in plastocyanin; the geometry observed in the copper complex of L₃ is very similar to that observed in plastocyanin. Therefore, the distorted geometries observed based on the present computational studies (Fig. 17) may very well support the geometries observed at the metal centers of metalloproteins.

While the Hg²⁺ complex of L₁ exhibits a perfectly linear geometry as N-Hg-N, the inclusion of weak interactions exhibited by the carbonyl oxygens of the arms, viz., HgN₂O₂, leads to a geometry that best fits a distorted octahedron with two vacant sites. Even the recognition species of L₂ with Ni²⁺ having the N₄O core can best be fitted to a distorted octahedron, but with one vacant site. The same receptor, L₂ exhibits an N₄ core with distorted tetrahedron for Zn²⁺. However, when both Ni²⁺ and Zn²⁺ are present, Ni²⁺ goes predominantly to the nitrogen core (viz., N₄ and distorted tetrahedron) and Zn²⁺ goes predominantly to the oxygen core (viz., O₅ and distorted trigonal bipyramid). This seems to suggest that in hetero-bimetallo-proteins, selective preference of one metal ion over the other may depend on the nature of the binding core as well the conformational mobility of the arms attached.

The Cu²⁺ complex of L₄ exhibits distorted trigonal bipyramidal geometry wherein each arm of L₄ acts as bidentate in filling a total of four coordinations, with the fifth coming from the acetonitrile, resulting in a NO₃S₂ binding core. Even the species of recognition of L₅ with Zn²⁺ having N₂O₄ core provides a better fit to a trigonal bipyramidal with one trigonal center being vacant. The Cd²⁺ complex of the same receptor, L₅ exhibits N₂O₄ binding core to result in a capped octahedral geometry with one vacant site. While the Zn²⁺ binds at the Schiff base region, the Cd²⁺ binds primarily at the lower rim phenolic oxygen atoms. All these studies seem to suggest that selective preference of one metal ion over the other depends on the nature of the binding core as well the conformational mobility of the arms attached.

References


Gaussian View 3.0 (Gaussian Inc, Pittsburgh, PA) 2003.


