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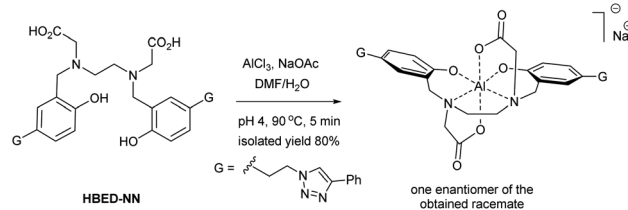
The HBED chelator is used to stabilize small and hard metal ions such as Fe^{3+} , Ti^{4+} , Ga^{3+} and Al^{3+} in both medicine and industry. While the coordination of hexadentate HBED^{4-} is known in the case of Fe^{3+} , Ti^{4+} and Ga^{3+} , it is unknown in the case of the small Al^{3+} ion since its corresponding complex has never been fully characterized. Thus, in this work the coordination pattern in a newly synthesized aluminum HBED-based complex ($[\text{Al}-\text{HBED}-\text{NN}]^-\text{Na}^+$) was determined using 2D NMR in conjunction with DFT calculations.

While X-ray crystallography remains the preferred method for determining the coordination of metal complexes, the requirement to obtain X-ray quality crystals can sometimes not be met.¹ In such instances NMR can be an alternative means to determine the coordination pattern and geometry of complexes.^{2–5} HMBC in particular is an especially useful NMR technique for structural analyses as it reveals multiple-bond connectivity between protons and X nuclei.² *N,N'*-Bis(2-hydroxybenzyl)ethylenediamine-*N,N'*-diacetic acid (HBED) is a popular polyaminocarboxylate-type chelator which can tightly bind to metal ions *via* its 6 coordinating atoms (two phenolate groups, two amine groups and two carboxylate groups) resulting in three possible octahedral isomers,^{6–12} each as a racemate.¹³ From an application point of view, it is important to distinguish these isomers from each other since isomeric structures can exhibit distinct (bio)chemical profiles.^{8,14} However, the content of the predominant complex species in the HBED chelation depends on a number of factors such as the reaction conditions, particularly the temperature, as well

as the nature of the metal in terms of its size, valence, hardness/softness, *etc.*^{3,7}

Although the synthesis and application of the Al-HBED hexacoordinate complex ($[\text{Al}-\text{HBED}]^-$) have already been described in basic and applied research, it has not yet been investigated from a structural point of view; and even characterization data to support its existence have not been reported.^{15–18} Only in one case was the occurrence of $[\text{Al}-\text{HBED}]^-$ reported in the research literature where it was detected by mass spectroscopy as a byproduct, but the compound was not isolated and characterized as a single substance.¹⁹ However, in contrast to $[\text{Al}-\text{HBED}]^-$, other HBED hexacoordinate scaffolds such as $[\text{Fe}-\text{HBED}]^-$, $[\text{Ti}-\text{HBED}]$ and $[\text{Ga}-\text{HBED}]^-$ have had their structures determined.^{3,11,12} Herein, we employed NMR in conjunction with DFT calculations to determine the coordination of the Al-HBED cage using the newly synthesized $[\text{Al}-\text{HBED}-\text{NN}]^-\text{Na}^+$ complex (Scheme 1).

For the preparation of the $[\text{Al}-\text{HBED}-\text{NN}]^-\text{Na}^+$ complex, the HBED-NN chelator (Scheme 1) bearing 1,2,3-triazolyl terminal groups was first prepared according to literature.³ As mentioned, metal chelation in general is highly dependant on the reaction conditions, such as the temperature, pH, *etc.*, often resulting in a mixture of complex species, isomers, side products and conformers with similar physical properties and spectra. Consequently, the ensuing analysis of the chromatographic and spectroscopic data can be challenging in order to



Scheme 1 Synthesis of the $[\text{Al}-\text{HBED}-\text{NN}]^-$ complex at elevated temperature as a single, racemic isomer.

^aGerman Cancer Research Center (DKFZ), Molecular Structure Analysis, INF 280, 69120 Heidelberg, Germany

^bUniversity of Hamburg, Institute of Pharmacy, Bundesstraße 45, 20146 Hamburg, Germany. E-mail: ata.makarem@chemie.uni-hamburg.de

^cUniversity of Neyshabur, Department of Chemistry, Neyshabur, Iran

† Electronic supplementary information (ESI) available: Methodologies and experimental procedures, characterization data for the synthesized compounds, and copies of spectra. See DOI: <https://doi.org/10.1039/d2dt01702d>

‡ Project leader.



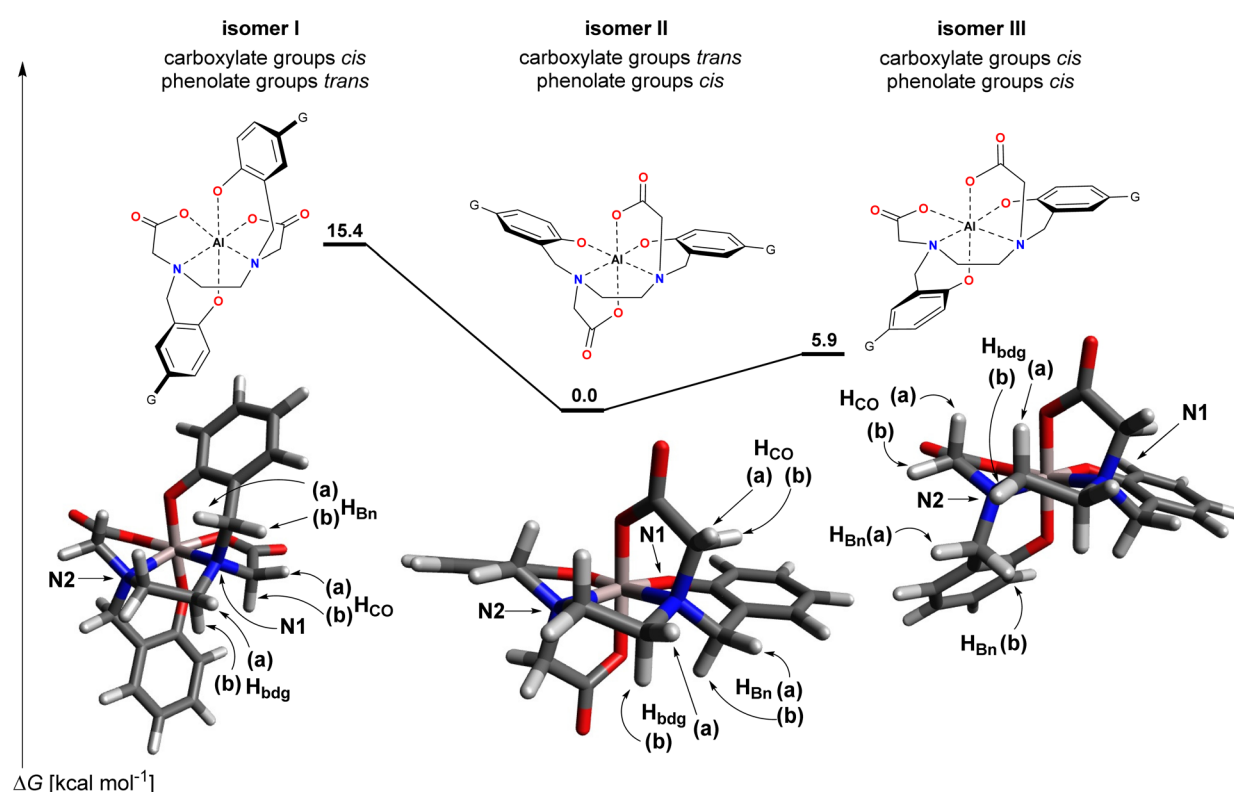
elucidate the reaction progress and optimize the process to a single product.^{7,8,20-22} For example, the upfield aliphatic region of ¹H NMR spectra in chelate systems especially is usually crowded with various multiplets and broad peaks.^{3,22-24} This challenge in the case of underderivatized HBED has been detailed in the literature.⁷ In contrast, the HBED-NN ligand facilitates convenient reaction monitoring by ¹H NMR since the sharp triazolyl singlet at 8.4 ppm (in DMSO) occurs in an uncrowded downfield region.³ Generally, triazolyl residue protons are very sensitive to structural modification, even to quite distant changes elsewhere in the molecule to which their δ_H 's respond accordingly.³⁻⁵ Thus, by utilizing the proton triazolyl singlet for reaction monitoring, the conversion and chemoselectivity of chelation is able to be followed and duly optimized.

Fortunately, due to the multiple permutations of the reaction conditions that we had conducted on the analogous Ga³⁺ complex,³ the optimal conditions found in that study were utilized for the [Al-HBED-NN]⁻ complex realizing its preparation in a straightforward manner (Scheme 1). Indeed, the chelation of the HBED-NN ligand to Al³⁺ was found to be high yielding (80% yield) when the reaction was conducted at an elevated temperature (90 °C) to conveniently produce a single geometric isomer after only 5 min. The isolated and purified [Al-HBED-NN]⁻Na⁺ product was fully characterized by NMR while

HR-ESI-MS provided an ion of m/z 753.2730 Da (calcd for C₄₀H₃₈AlN₈O₆, 753.2730 Da) to confirm its fabrication (see ESI†).

The three possible isomers for the isolated [Al-HBED-NN]⁻ complex – isomers **I** (carboxylate groups *cis*, phenolate groups *trans*), **II** (carboxylate groups *trans*, phenolate groups *cis*) and **III** (carboxylate groups *cis*, phenolate groups *cis*) – are depicted in Scheme 2. For both isomers **I** and **II**, the two halves of the ligand are equivalent due to the presence of a two-fold rotation axis while isomer **III** lacks any symmetry element. Thus, while isomer **III** should exhibit 36 unique carbon signals and 34 unique proton signals in the respective spectra, isomers **I** and **II** should only exhibit half these numbers due to the equivalent halves of the ligand. On the basis of the ¹H and ¹³C NMR spectra, isomer **III** can be immediately ruled out since in the ¹³C NMR spectrum only 18 signals are present and in the ¹H NMR spectrum only 17 signals are present (see ESI†).

The strategy applied here to distinguish between isomers **I** and **II** basically follows that applied previously³ where the distinction between the isomers relies on an examination of a comprehensive set of possible vicinal proton–carbon correlations in a HMBC spectrum, not only in terms of correlations observed, but also correlations not observed. Pertinent $^3J_{H,C}$'s, *viz.* those within the coordination sphere, are evaluated to be either large or small depending on the dihedral angle between



Scheme 2 Structures of the three possible isomers and their DFT-calculated models for the [Al-HBED-NN]⁻ complex. *N.b.* Each isomer depicted here is one of a pair of enantiomers and for the sake of convenience, the G groups were not included in the calculations and hence are missing from the depicted DFT-calculated structures. Methylenic protons that are oriented upwards in the depictions are designated as **a**, while methylenic protons oriented downwards are designated as **b**.

the nuclei concerned. If $^3J_{H,C}$ was anticipated to be large, *e.g.* >5 Hz, then a correlation should be observed in a HMBC spectrum optimized for a long-range coupling of 8 Hz; while values of $^3J_{H,C} < 5$ Hz would otherwise not be expected to give rise to an observable correlation.

The size evaluation of $^3J_{H,C}$ is dependent on vicinal proton–carbon nuclei following a Karplus-type function.²⁵ The well-known Karplus-type relationships for vicinal couplings between nuclei are functions relating 3J 's with the dihedral angles (α) between the nuclei.^{26,27} They since have been developed for various systems (*e.g.*, H–C–C–C, H–C–O–C, F–C–C–H, P–C–C–H, H–C–N⁺–H and H–C–C–N)^{25,28} however, they all exhibit quite similar trajectories, *viz.*, 3J is a maximum when $\alpha = 0^\circ$ or 180° due to maximum overlap of orbitals and a minimum when $\alpha = 90^\circ$ due to the orbitals being perpendicular to one another. The dihedral angle is not the only factor determining the size of 3J , other factors such as bond length, order and angle as well as the presence of electronegative/positive substituents *etc.* also have an influence; but for the comparison of systems containing the same atoms, Karplus-type relationships are very useful for evaluating vicinal coupling–dihedral angle relationships. The Karplus-type function for the $^3J_{H,C}$'s in propane is depicted in Fig. 1, used here as the function for the vicinal proton–carbon relationship.²⁹ For this analysis, a value of 5 Hz is defined as the cutoff between small and large couplings and from the plot in Fig. 1 it can be discerned that large $^3J_{H,C}$'s occur when $\alpha < 31.5^\circ$ or $\alpha > 135^\circ$.

For the structural evaluation, all 16 vicinal proton–carbon relationships involving the methylene protons within the coordination sphere were taken into consideration. To evaluate the expected size of the $^3J_{H,C}$'s from the dihedral angles, the structures of isomers **I**–**III** were optimized by DFT calculations at the b3lyp/6-31 g(d,p)//b3lyp/6-31 g(d,p) level of theory with DMSO simulated as the solvent by a polarized continuum medium (Scheme 2). This level of theory was considered appropriate to attain geometries sufficiently reliable for the ensuing analysis. The starting geometry for isomer **II** was developed from the X-ray crystallographic structure of [Ti–HBED],¹² while isomers **I** and **III** were modified accordingly from the optimized geometry of isomer **II**. For the sake of con-

venience and to reduce computational cost, the terminal **G** groups were not included in the calculations as they were not considered to have any undue influence on the geometry of the coordination sphere.

The ΔG values for isomers **I**–**III** obtained from the DFT calculations overwhelmingly support isomer **II** as the thermodynamically favored structure (isomers **I** and **III** were computed to be higher in energy by 15.4 and 5.9 kcal mol⁻¹, respectively). In both isomers **I** and **III**, there appears to be a possible steric interaction between an acetyl segment and a phenyl ring, and as well, for isomer **I** there appears to be an additional possible steric interaction between the ethylenediamine bridge and the other phenyl ring. These possible steric interactions could account for some of the energy difference with respect to isomer **II**. Moreover, taking the preferred bond angles as those present in isomer **II** where the angles are effectively identical for the two halves of the ligand, for bond angles starting from carbon and ending in the Al³⁺ ion, isomer **I** had more distorted bond angles (taken as a deviation >4°, relative to the value in isomer **II**) than isomer **III** (6 *vs.* 4, respectively). Furthermore, for both isomers **I** and **III**, a majority of the distorted bond angles were in that half of the ligand experiencing possible steric interaction with the acetyl segment. By comparison for the X-ray crystallographic structure of [Ti–HBED],¹² only two degenerate angles were significantly different to those in isomer **II**. For bond angles with the Al³⁺ ion as the central atom and only comparing angles within the one ligand half, there was nothing of note between the four structures, and similarly for the bond lengths. A listing of the bond angles and bond lengths considered is presented in Table S5.†

Of note, calculated energies are highly sensitive to the environment and method of calculation and can change markedly. Moreover, it may be the kinetic product that is formed initially and if interconversion between the isomers is inexorably slow, the kinetic product only may be isolated. While it is noted that multiple isomers have been claimed to be observed in solution for the larger Ga³⁺ ion,^{7,8} perhaps only one isomer is overwhelmingly favored thermodynamically for Al³⁺. A possible reason for this is the smaller size of Al³⁺ which might be exacerbating the bond angle strain and steric interactions in isomers **I** and **III**. This would account for their large DFT-calculated energies. However, the geometries for the immediate atoms of the coordination sphere of these quite rigid complexes will not fundamentally alter due to the environment or higher levels of theory. The purpose of the DFT calculations was to use the optimized geometries for structural evaluation by NMR which now follows.

For isomer **II**, the corresponding dihedral angles calculated for the two halves are identical to one decimal place. Despite the differences calculated for isomer **I** between the two halves of the ligand, it is expected that rapid fluxional motion will interconvert the two halves of the ligand rendering them equivalent on the NMR timescale; thus the $^3J_{H,C}$ values were averaged for the structural evaluation. For isomer **III** – included for the sake of completeness though already eliminated from the

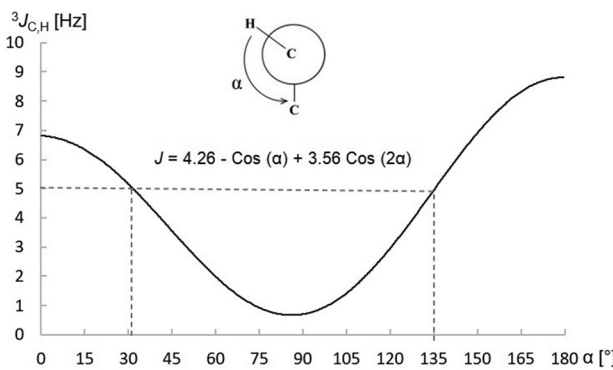


Fig. 1 Theoretical plot of $^3J_{H,C}$ vs. the dihedral angle between the proton and carbon nuclei for propane, used as the function here.



candidate structures – interconversion of the two halves of the ligand is not possible and was treated in two ways: firstly by averaging of the coupling values and; secondly, by consideration that if one dihedral angle of a methylene pair were to provide a large $^3J_{H,C}$ value, it is taken that a correlation would be expected for both that proton and its corresponding partner in the other half of the ligand. Of note, in the 1H NMR spectrum, the four protons of the ethylenediamine bridge segment are effectively isochronous; so, if one equivalent pair of protons in the ethylenediamine bridge were to provide a large $^3J_{H,C}$ value, it is taken that a correlation would be expected for the ethylenediamine bridge-methylene signal. The result of this is that one relationship for each isomer is rendered indeterminate (or in other words, a possible correlation expected to be present is masked by a possible correlation expected to not be present), leaving a total of 15 relationships that can be utilized for structural evaluation.

From a compilation of the evaluated dihedral angles, selected relationships and their expected *vs.* experimental results are presented in Table 1 for isomers I–III (a full listing

of the evaluated dihedral angles are presented in Tables S1–S4†). For isomer I, only in five cases was there a match between expected and experimental results and thus isomer I is eliminated as a candidate structure for the Al-HBED hexacoordinate cage (Table S2†). Interestingly, in the case of isomer III, both approaches described above only provide two contradictory predictions to experimental observations (Tables S3 and S4†). This is provided of course that corresponding protons from each half of the ligand were in fact isochronous; thus, isomer III can also be eliminated as a candidate structure on the basis of this analysis. On the other hand, for isomer II, all 15 expected results were in concert with experimental results consisting of seven observed correlations and eight possible correlations not observed (Table S1†). The complete match between observed and unobserved correlations undoubtedly confirms that the structure of $[Al\text{-HBED-NN}]^-$ isolated from the reaction has the carboxylate groups *trans* to each other and thus the coordination of HBED to Al^{3+} follows that of Fe^{3+} in $[Fe\text{-HBED}]^-$, Ti^{4+} in $[Ti\text{-HBED}]$ and Ga^{3+} in $[Ga\text{-HBED}]^-$.

Table 1 Selected proton–carbon vicinal relationships for isomers I–III and expected *vs.* experimental results

^a N1 and N2 refer to the ligand half. ^b ✓ = consistent; ✗ = inconsistent. ^c Despite the medium size of α , a correlation is expected for entry 3 due to the overlap of H_{bdg} (a) and (b).

It is worth noting that the structural determination analysis presented here can still be successfully performed without stereochemical assignment of the methylene protons, *i.e.* assigning the two protons of each methylene group. However, not stereochemically assigning the protons reduces the number of relationships that can be used for the analysis by eight (six relationships for the second evaluation approach in the case of isomer **III**). For isomer **I** this makes a sizeable difference as six contrarian predictions to isomer **II** are lost, though four contrarian predictions remain to enable a distinction between isomers **I** and **II**. Beguilingly, not stereochemically assigning the protons makes no difference in either of the two evaluation approaches for isomer **III** as no contrarian predictions to isomer **II** are lost.

Conclusions

A new aluminum HBED-type hexacoordinate complex has been prepared and fully characterized, and the coordination geometry of HBED⁴⁻ to Al³⁺ proven for the first time. Only for isomer **II** was there a complete match between observed correlations consistent with prediction and possible correlations that were not observed but also in line with prediction. The coordination of the hexadentate HBED chelator to Al³⁺ follows that of Fe³⁺, Ti⁴⁺ and Ga³⁺ in which the carboxylate ligands are *trans* to each other, and perhaps because of the small size of Al³⁺ only one isomer may be thermodynamically possible. The strategy of examining a comprehensive set of vicinal proton–carbon relationships in conjunction with DFT calculations – both for those correlations expected to be present as well as those expected to be absent in a 8 Hz-optimized HMBC spectrum – has once again proven successful.

Conflicts of interest

The authors declare no competing financial interest.

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