Formation of an Amine-Water Cyclic Pentamer: A New Type of Water Cluster in a Polyaza cryptand

Musabbir A. Saeed,‡ Bryan M. Wong,‡ Frank R. Ronczek, # Ramaiyer Venkatraman,‡ and Md. Alamgir Hossain*†

†Department of Chemistry and Biochemistry, Jackson State University, Jackson, Mississippi 39212, 
‡Materials Chemistry Department, Sandia National Laboratories, Livermore, California 94551, and 
#Department of Chemistry, Louisiana State University, Baton Rouge, Louisiana 70803

Received February 2, 2010; Revised Manuscript Received February 25, 2010

ABSTRACT: Structural analysis of an adduct of a thiophene-based cryptand with tosylic acid shows the formation of a hybrid amine-water cyclic pentamer composed of four water molecules and one protonated amine in the charged hydrophobic cavity. The bulky tosylate groups remain outside the cavity, making the ligand favorable for hosting water molecules. Ab initio calculations based on density functional theory (DFT) confirm that the hybrid amine-water pentamer is stabilized inside the hydrophobic cavity of the cryptand.

Despite significant progress in experimental and theoretical studies of discrete water clusters, a detailed understanding of molecular water within a confined hydrophobic cavity remains a challenge.1 Within the peptide walls of a protein, molecular or clustered water can form in the hydrophobic cavity through hydrogen bonding and hydrophobic interactions,2 playing a critical role in controlling the function, activity, and folding of the protein.3 Protein-bound water clusters have also been identified structurally, showing stable polygons composed of three to eight water molecules.4 At the molecular level, water clusters are thought to facilitate the transfer of a proton along the hydrogen-bonded chain, which is not only a fundamental biological process5 but also a key mechanism for the generation of electrical power in hydrogen fuel cells.6 Interest in this area continues to intrigue the scientific community both in terms of basic research and technological applications.7 In particular, the structural characterization of water clusters is critical in correlating and predicting the properties of bulk water at a molecular level. In the solid state, water clusters (H2O)n are known to exist as discrete rings, chains, tapes, and layers8 with different numbers of water molecules. Among these arrangements, cyclic water clusters have been shown to be stabilized as trimers,9 tetramers,9b hexamers,10c,d hexamers,10e octamers,10f decamers,10g undecamers,10h and dodecamers10i in various organic environments. However, in contrast to their presence within crystal lattices or between hosts, cyclic water clusters inside the cavity are rare,11 with the exception of tricyclic amide with hexamer,11a p-xyllyl cryptand for quasi-prismatic hexamer,11b and self-assembled palladium-based cages for a decamer.11c It is known that, in the solid state, most of the organic hosts (ca. 96%) forming water clusters are directly linked to water molecules via hydrogen bonding interactions.12 However, to the best of our knowledge, the participation of a heteroatom in forming a ring has not been previously reported within a synthetic molecule. Herein, we present structural evidence and ab initio calculations based on a new type of hybrid pentamer [(H2O)4(NH2)]þ (Chart 1). This unique structure is stabilized within the charged cavity of a hexaprotonated cryptand L, bearing a similar structural topology to the cyclic pentamer, (H2O)5,10c or cyclopentane.

We recently reported that the bicyclic receptor L effectively binds chloride13a and nitrate13b ions in aqueous solution through the protonated bridgehead amines. While one chloride was found to be encapsulated linearly within the cavity, three nitrates were bridged in L forming a [NH(NO3)3HN]þ motif in the solid state. The formation of trinitrate complex, however, was inconsistent with a 1:1 binding observed in aqueous solution, suggesting that water molecules occupied the cavity and prevented the competitive binding of more than one nitrate in solution. This observation inspired us to further explore the cryptand with bulky anions that are too large to enter the cavity, thus making an ideal environment in L for hosting water molecules.

Crystallographic analysis of the tosylate salt of L revealed that the ligand in its hexaprotonated form crystallizes in the monoclinic space group P21/n with 6 tosylate anions and 12 water molecules.14 As shown in Figure 1, a cyclic pentamer is formed with four water molecules (O3W, O1W, O2W, and O6W) and one protonated amine (N6) group inside the cavity. The hydrogen-bonding interactions facilitate the formation of a pentamer which adopts a puckered conformation, with one water (OW1) lying at 1.410 Å above the mean plane formed by the three oxygens and with 2.894(2) and 2.754(2) Å for N6—O3W and N6—O6W, respectively. The linking nitrogen (Figure 1b). The protonated amine involved in the ring acts as two hydrogen bond donors for two waters with 2.894(2) and 2.754(2) Å for N6···O3W and N6···O6W, respectively.

The large anions are located outside the cavity without interacting with the macrocycle. The secondary amines are protonated in the macrocycle exhibiting a C3 symmetry along the bridgehead nitrogens, N1 and N4. The absence of an anion inside the cavity makes L favorable for hosting the water molecules. The distance between the bridgehead nitrogen sites is longer (7.891 Å) than the corresponding distance in the chloride (6.096(4) Å)13a or nitrate (5.528(6) Å)13b complex, where the internally bound anion brings the two bridgehead amines closer through electrostatic interactions. The geometry of the hybrid pentamer is similar to the
endothermic peak at 112 °C corresponds to the formation of a second phase on TGA is about 3.8%, corresponding to four water molecules that are assigned to the internally bound water in the hybrid pentamer observed in the crystal structure. The first weight loss, which corresponds to eight water molecules, could be due to the removal of water from the complex. The second weight loss, which corresponds to the removal of additional water molecules, is observed at a temperature of 120 °C. After losing the weight of the crystalline water, the complex appears to be stable up to 275 °C. The exothermic and endothermic peaks at ∼275 °C and ∼350 °C on the DSC curve could be due to the melting and decomposition of the complex, respectively. Furthermore, the presence of water in the complex was identified by FT-IR spectra (Supporting Information). The complex showed a broad band at around 3420 cm⁻¹ that can be assigned to the O···H stretching vibration of water present in the sample, similar to the calculated frequency, 3443 cm⁻¹ obtained for the pentamer or to the experimental value, 3400 cm⁻¹ observed for other water clusters. This band almost disappeared after heating the sample at 120 °C for 12 h, confirming the exclusion of water molecules from the complex, an observation that is consistent with the TGA observations and theoretical calculations discussed below.

In order to quantitatively understand the unique bonding involved in the hybrid amine-water pentamer, ab initio calculations based on density functional theory (DFT) were carried out. All quantum chemical calculations were carried out with Truhlar’s M06-2X meta-GGA hybrid functional which has been shown to accurately predict the relative energies of water hexamers and other noncovalent bonding interactions in large molecular systems. Ground-state geometries were optimized at the M06-2X/6-31G(d,p) level of theory, and harmonic frequencies were computed to verify that all structures were true minima. The initial crystallographic model was used as a starting point to optimize the geometries of the protonated cluster-cryptand. At the optimized geometries, single-point energies with a larger 6-311G(d,p) basis set were carried out (a diffuse 6-311+G(d,p) basis set was used for the H, O, and N atoms involved in the amine-water pentamer within the complex). Figure 3a,b shows the optimized geometry and electrostatic potential of the cluster-cryptand, respectively. From the DFT-optimized geometry, we find that the bond distances compare excellently with the experimental crystal structure, confirming the puckered conformation of the amine-water pentamer (Supporting Information). The electrostatic potential also shows a localization of positive charge on the protonated amine groups within the cryptand structure. From the optimized geometry, the stabilization energy of complexation was calculated as \( E_s = E(\text{cluster-cryptand}) - E(\text{cryptand}) - 4E(\text{water}) \), and was found to be \(-68.5 \text{ kcal/mol} \), resulting from strong electrostatic interactions between the hydrogen-bonded water molecules and the highly charged macrocycle, in addition to strong hydrogen bonds which stabilize the entire complex. In order to compare this binding energy with other complexes, we also calculated stabilization energies for an isolated water pentamer (Chart 1A) and an isolated NH4⁺···water pentamer (Chart 1B). At the M06-2X/6-31+G(d,p) level of theory, we find that the NH4⁺···water complex has a binding energy of \(-71.2 \text{ kcal/mol} \), while the isolated water pentamer complex has a smaller binding energy of \(-40.4 \text{ kcal/mol} \). The cluster-cryptand stabilization energy is slightly less (more positive) than the NH4⁺···water complex due to additional steric interactions (cf. Figure 3b) within the cryptand.

In conclusion, we have structurally identified a novel hybrid amine-water cyclic pentamer formed inside the cavity of a cryptand. Ab initio calculations not only support the stabilization of the hybrid pentamer inside the hydrophobic cavity but also reveal that the hybrid amine water pentamer is energetically more stable than the conventional water pentamer. The tosylate group used as a counteranion is not complementary in size to the cryptand, allowing neutral water molecules to reside inside the cavity. This observation clearly indicates that an appropriate selection of counteranions makes the ligand useful for caging water in the charged hydrophobic environment. The hybrid pentamer, in which a protonated amine is an important component in the ring, adopts a puckered conformation similar to cyclopentane or cyclic water pentamer. In vivo water clusters formed within a protein cavity or cleft are known to be stabilized.
predominantly through hydrogen-bonding interactions with CO groups at the C termini and NH groups at the N termini as well as with the charged amino acid side chains. While there is an abundance of discrete water clusters reported in the literature, the formation of the hybrid pentamer described in this work has not been observed before in a synthetic host. Given the novel role of water clusters in chemical and biological systems, the present findings provide insight and serve as a prototype for stabilizing other pure and hybrid water clusters.

Acknowledgment. The project described was supported by Grant Number G12RR013459 from the National Center for Research Resources. This material is based upon work supported by the National Science Foundation under CHE-0821357. Purchase of the diffractometer was made possible by Grant Number G12RR013459 from the National Center for Research Resources.

Supporting Information Available: One crystallographic file in CIF format, synthetic procedures, hydrogen bonding interactions, ORTEP diagram with numbering scheme, FTIR spectra, and CIF format, synthetic procedures, hydrogen bonding interactions, gas-phase electronic spectrum, and electrostatic potential of [H₆L(4H₂O)]₆⁺.

References

(14) Crystal data for [H₆L(4H₂O)]₆⁺. (b) Electrostatic potential of [H₆L(4H₂O)]₆⁺ at the M06-2X/6-31G(d,p) level of theory. (c) Figure 3. (a) Optimized geometry of the amine-water cluster, [H₆L(4H₂O)]₆⁺ at the M06-2X/6-31G(d,p) level of theory. (b) Electrostatic potential of [H₆L(4H₂O)]₆⁺.