Molecular dynamics simulation of protein-mediated biomineralization of amorphous calcium carbonate

R. Sandya Rani and Moumita Saharay*

The protein-mediated biomineralization of calcium carbonate (CaCO$_3$) in living organisms is primarily governed by critical interactions between the charged amino acids of the protein, solvent, calcium (Ca$^{2+}$) and carbonate (CO$_3^{2-}$) ions. The present article investigates the molecular mechanism of lysozyme-mediated nucleation of amorphous calcium carbonate (ACC) using molecular dynamics and metadynamics simulations. The results reveal that, by acting as nucleation sites, the positively charged side chains of surface-exposed arginine residues form hydrogen bonds with carbonates and promote aggregation of ions around them leading to the formation and growth of ACC on the protein surface. The newly formed ACC patches were found to be less hydrated due to ion aggregation-induced expulsion of water from the nucleation sites. Despite favorable electrostatic interactions of the negatively charged side chains of aspartate and glutamate with calcium ions, these residues contribute minimally to the growth of ACC on protein surface. The activation barrier for the growth of partially hydrated ACC patches on lysozymes was determined from the free energy profiles obtained from metadynamics simulations.

Introduction

Calcium carbonate (CaCO$_3$) is a commonly found biomineral in the environment. The diversity in the morphology, origin, and composition exhibited by naturally occurring CaCO$_3$ makes it an intriguing system to investigate the mechanism, energetics and thermodynamics of biomineralization. Apart from the three anhydrous (calcite, vaterite, and aragonite) and two hydrous (monohydrocalcite, CaCO$_3$·H$_2$O; and ikaite, CaCO$_3$·6H$_2$O) crystalline polymorphs, calcium carbonate can also exist in metastable amorphous phases. Amorphous calcium carbonate (ACC) serves as a precursor in the biomineralization of almost all types of biogenic CaCO$_3$.

In living organisms, the process of mineral formation is primarily governed by the interactions between the charged biopolymers, ionic constituents (Ca$^{2+}$ and CO$_3^{2-}$), and water. For instance, the negatively charged proteins in corals and sea urchin spicules are shown to stabilize the amorphous phase of CaCO$_3$ even in *in vitro* experiments in the absence of any biological cells. A recent study on the formation of coral skeleton demonstrated that the ACC phase is indeed the metastable precursor to the crystalline CaCO$_3$, and that the growth rate of aragonite crystals through the non-classical crystallization process involving aggregation of ACC particles is ~100 times faster than the classical route of ion-by-ion formation of crystals. Earlier studies have also shown that the binding affinity of Ovoceidin-17 (OC-17), a hen eggshell protein, for ACC nanoparticles is much higher than that for the calcite surface suggesting that favorable binding of the protein to ACC is critical for biomineralization and eggshell formation. The lower calcite-protein binding affinity is leveraged for the detachment of the protein from the crystal after the ACC to crystal transition. Thus, OC-17 or a class of functionally similar proteins/biopolymers can be regarded as catalysts to facilitate the crystallization of CaCO$_3$.

Although previous computer simulation studies have investigated the mechanism of the attachment of the protein to the ACC nanoparticles and CaCO$_3$ crystal surfaces, little is known about the role of protein in promoting the formation of ACC from the ionic solution. Another key feature that is yet to be understood is the atomistic details of the mechanism and energetics of protein-ACC association/dissociation process. Some proteins/biopolymers hinder the crystallization of CaCO$_3$ by stabilizing the amorphous phase, and the biomineralization is activated as and when the living organism requires it. The nucleation of CaCO$_3$ in the presence of negatively charged biopolymers is thought to occur in two steps: (i) the binding of Ca$^{2+}$ with the biopolymer to form Ca$^{2+}$-bound globules, (ii) the diffusion and association of CO$_3^{2-}$ ions with the globules to form polymer-mediated ACC phase that then grows in size. These polymers, referred to as ‘ion sponge’, critically determine where and how the CaCO$_3$ nucleation will take place.

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place and proceed further. However, the precise molecular details associated with these steps are still missing.

**In vitro** experiments have shown that the positively charged lysozyme increases the nucleation rate of CaCO$_3$ crystals. The large negative $\zeta$-potential (14 mV) around this protein favors the formation of negatively charged double layers of CO$_3^{2-}$ around its surface. To understand the microscopic details of the onset of protein-mediated aggregation of Ca$^{2+}$ and CO$_3^{2-}$ and to identify the vital molecular attributes and the key residues that govern the mineralization process, we have performed a series of atomistic molecular dynamics simulations of lysozyme, a hen eggshell protein. Our results reveal that the positively charged side chains of solvent-exposed arginine residues act as nucleation sites for the formation and growth of ACC on protein surface, while the negatively charged side chains of glutamate and aspartate residues contribute minimally to biomineralization.

### Simulation details

We have investigated the following four model systems: (i) lysozyme in water (hereafter referred to as LW), (ii) hydrated ACC (hACC) having 1 : 0.67 composition of CaCO$_3$ : H$_2$O, (iii) lysozyme in CaCO$_3$ solution with two different CaCO$_3$ : H$_2$O concentrations (S$_1$LC and S$_2$LC). The dimensions and the number of constituents of the simulated model systems are provided in Table 1.

The LW model allows us to characterize the lysozyme–water interactions in the absence of Ca$^{2+}$ and CO$_3^{2-}$ ions, whereas S$_1$$_2$LC models are used to probe the influence of these ions on lysozyme. The initial coordinates of the hen egg-white lysozyme (PDB Id: 2cds) was taken from the X-ray crystal structure. To neutralize the net positive charge on lysozyme, 8 Cl$^-$ were added in LW system, while 4 extra CO$_3^{2-}$ were added in S$_1$LC models. The initial configurations of S$_1$LC were generated by randomly placing desired numbers of Ca$^{2+}$ and CO$_3^{2-}$ ions in cubic water boxes of sizes specified in Table 1. The lysozyme was then solvated in this ionic solution by removing the ions and water molecules that were in hard contact with the protein using CHARMM-GUI$^{27-29}$ package. CHARMM36 (ref. 30) force field (FF) parameters were used for the protein and ions (Ca$^{2+}$, CO$_3^{2-}$) together with the standard TIP3P water model.

For hACC, H$_2$O : CaCO$_3$ ratio was maintained at 0.67 to be consistent with our earlier investigation. The MD simulations were performed using NAMD$^{31}$ package. The systems were equilibrated for 1 ns in NPT ensemble followed by 15 ns of MD simulations in NVT ensemble with the time step of 2 fs. The temperature and pressure were maintained at 303 K and 1 atm, respectively. The temperature was controlled by the Langevin thermostat using a damping coefficient of 1 ps$^{-1}$. The electrostatic interactions were evaluated using particle mesh Ewald method with a grid spacing of 1 Å. Periodic boundary conditions were used and a distance cutoff of 12 Å was chosen to calculate the non-bonded interaction energies between different species.

To understand the flexibility of the near neighbor environment around the positively charged side chains, we studied the evolution of hydrogen bonds formed by the side chains with the nearest CO$_3^{2-}$ and water molecules. An arginine side chain was considered to be hydrogen bonded to CO$_3^{2-}$ if the CZ-O$_r$ (refer Fig. 1 for atom labels) distance was less than 4.5 Å or to a water molecule if the CZ-O$_w$ distance was less than 4.0 Å$^{33-35}$ For each type of h-bond, the time correlation function, $C_{h_{bs}}(t)$, which is defined as follows,

$$C(t) = \langle h(0)h(t)\rangle/h$$

calculated with a time resolution of 2 ps. Here, $h(t) = 1$ when two groups are h-bonded, and $h(t) = 0$ otherwise. The ‘$\langle \rangle$’ represents the ensemble average.

We performed well-tempered metadynamics$^{36-38}$ simulations, using NAMD package, to estimate the free energy required for the association of ions and dehydration of hydrated Ca$^{2+}$–CO$_3^{2-}$ clusters around selected side chains of lysozyme. The separation ($r_{CV}$) between the tagged ions and the residue of interest was chosen as the collective variable (CV). The Gaussian hills were added at a regular interval of 100 fs, and the width of each Gaussian hill was 1 Å. The total run length for the metadynamics simulation was 100 ns.

### Results and discussion

#### Ionic structure around lysozyme

Fig. 2 compares the arrangement of ions around lysozyme in the initial configuration of S$_1$LC with that in a snapshot taken after 15 ns of MD simulation. The ions are distributed around lysozyme in the initial structure, while dense ionic patches are observed around specific surface residues of lysozyme in the equilibrated system. The aggregation of ions occurs primarily around arginine residues, but to some extent also around glutamate and aspartate. The clustering of Ca$^{2+}$ and CO$_3^{2-}$ at specific surface sites of the protein underscores the key role of protein-mediated aggregation of ions in the formation of biominerals. The average mass density of the dense ionic

<table>
<thead>
<tr>
<th>System</th>
<th>CaCO$_3$ pairs</th>
<th>Water molecules</th>
<th>Orthorhombic box dimension (Å$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LW</td>
<td>—</td>
<td>30 284</td>
<td>70.0 × 70.0 × 70.0</td>
</tr>
<tr>
<td>S$_1$LC</td>
<td>235</td>
<td>14 289</td>
<td>76.2 × 76.2 × 76.2</td>
</tr>
<tr>
<td>S$_2$LC</td>
<td>474</td>
<td>14 238</td>
<td>76.5 × 76.5 × 76.5</td>
</tr>
<tr>
<td>hACC</td>
<td>1620</td>
<td>1094</td>
<td>50.0 × 50.0 × 50.0</td>
</tr>
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</table>
patches around arginine is \( \sim 1.3 \text{ g cm}^{-3} \), which is comparable with that of hydrated ACC.\textsuperscript{26} However, the local environment around glutamate (0.7 g cm\(^{-3}\)) and aspartate (0.71 g cm\(^{-3}\)) are relatively less dense than the microenvironment of arginine residues.

The clusters of ions are observed to grow in size during the course of MD simulation. The largest ion cluster is formed around ARG14, ARG125, and ARG128. Due to their close proximity, these three surface residues form an ‘attractive triangle’, which favors the rapid growth of amorphous calcium carbonate on lysozyme. The positively charged guanidinium (Gdm\(^+\)) functional groups of arginine residues play a crucial role in the nucleation and growth\textsuperscript{25} of crystalline CaCO\(_3\) by favorably interacting with the negatively charged CO\(_3^{2-}\) ions. The Gdm\(^+\)–CO\(_3^{2-}\) and Ca\(^{2+}\)–CO\(_3^{2-}\) interactions facilitate the ion association and further growth of amorphous CaCO\(_3\)-like domains around these residues.

### Radial distribution function

To further investigate the local environment around the key residues of the protein, we examined the distribution of ions and water around them by calculating the radial distribution functions (RDFs) that describes how the density of neighbors vary around a reference atom/particle as a function of radial separation between the two.

**Local environment around ions.** Fig. 3 compares the RDFs involving Ca\(^{2+}\), CO\(_3^{2-}\), and water oxygen (O\(_w\)) of hACC with that of S\(_2\)LC. The water structure around Ca\(^{2+}\) and CO\(_3^{2-}\) are mostly similar in both the systems as evident from the peak positions and the peak heights of Ca–O\(_w\) and Co–O\(_w\) distributions. However, the population of water molecules (Table 2) around Ca\(^{2+}\) and CO\(_3^{2-}\) differs significantly between these two systems due to the difference in CaCO\(_3\) : H\(_2\)O ratio (1620 : 1094 in hACC and 474 : 14 238 in S\(_2\)LC).

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**Fig. 1** Atom names used for the descriptions of side chains in (a) arginine, (b) aspartate, and (c) glutamate.

**Fig. 2** Arrangement of Ca\(^{2+}\) (red spheres) and CO\(_3^{2-}\) (black) ions around arginine (blue), aspartate (green), and glutamate (green) residues in S\(_1\)LC for (a) initial configuration and (b) after 15 ns of MD simulation. Water molecules are not shown for clarity. VMD\textsuperscript{39} software package was used for visualization.
Although the peak intensities in ion–ion RDFs are relatively higher in S2LC than in hACC, peak positions remain mostly unchanged. The ion–ion RDFs indicate that the ions are held together strongly to form a semi-rigid and well-structured coordination shells in S2LC. The present structural analysis reveals that local environment around lysozyme favors the aggregation of ions and the formation of amorphous calcium carbonate, although, the concentration of ions is much lower than the hydrated ACC (hACC) studied here.

Distribution of ions around lysozyme. Here we investigated the near neighbor structures around arginine, glutamate, and aspartate side chain residues. The high intensity peaks in the protein-ion RDFs indicate well resolved first and second coordination shells of ions around these key residues, preferably for arginine (Fig. 4), in both S1LC and S2LC.

In Fig. 4, the RDFs between N_{Gdm} (guanidinium-nitrogen) and C_c (carbonate carbon) atom pair shows three maxima at 3.34, 5.18, and 7 Å. The coordination number at the first minimum is 1.24. The first and second maxima are separated by a plateau of width ~0.7 Å with negligible intensity, showing a crystal-like ordering of C_c atoms in this region. The number of O_c (carbonate oxygen) around N_{Gdm} is ~1.4 at the first minimum (2.9 Å) of g_{N_{Gdm}-O_c}(r) (Fig. 4). The underlying geometry could possibly indicate a configuration where one of the H atoms of a N_{Gdm} forms single h-bond with O_c. This preferential arrangement compares well with the geometry optimized isolated cluster of Gdm−CaCO_3−2H_2O, where the distance between N_{Gdm} and O_c is ~2.65 Å. The second and third maxima in g_{N_{Gdm}-O_c}(r) appear at 3.4 and 4.6 Å, respectively. The coordination numbers at the corresponding minima in S1LC are 2.3 (2.0 in S2LC) and 4.7 (4.5 in S2LC), respectively.

The first maximum in g_{Ca-Ca}(r) (Fig. 4) shows two sub peaks at 4.2 and 5.5 Å in both the systems. This characteristic arrangement of Ca^{2+} around N_{Gdm} is somewhat similar to that of Ca^{2+}-environment around C_c in hACC (Fig. 3), where the first peak of g_{Ca-Ca}(r) is also divided into two sub peaks. This indicates two preferred arrangements: for longer distance, Ca^{2+} is coordinated with only one N_{Gdm} or O_c making a ‘monodentate’ geometry whereas for shorter distance Ca^{2+} is equidistant from two N_{Gdm} or O_c making a ‘bidentate’ geometry. With increased ionic concentration in S2LC, the intensity of these peaks decreases which implies a less-structured coordination shell with same number of neighboring Ca^{2+}.

Table 2 The peak positions and coordination numbers (N(r)) in the ion–ion and ion–water radial distribution functions obtained from hACC and S2LC.

<table>
<thead>
<tr>
<th>Atom pair</th>
<th>g(r) S2LC</th>
<th>hACC</th>
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<tbody>
<tr>
<td>Peak position</td>
<td>N(r)</td>
<td>Peak position</td>
</tr>
<tr>
<td>C_c−C_c</td>
<td>4.8 3.0 4.6 9.2</td>
<td>2.6, 3.2 2.3 2.6, 3.2 4.6</td>
</tr>
<tr>
<td>Ca−C_c</td>
<td>2.6 3.2 2.3 4.6</td>
<td>2.2 3.0 2.2 0.8</td>
</tr>
<tr>
<td>Ca−O_w</td>
<td>2.2 3.0 2.2 0.8</td>
<td>3.3, 3.8 1.3 3.3, 3.7 4.4</td>
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<tr>
<td>Ca−O_c</td>
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<td>3.3, 3.8 1.3 3.3, 3.7 4.4</td>
</tr>
<tr>
<td>C_c−O_w</td>
<td>3.3 12.8 3.1 4.04</td>
<td>2.6, 3.2 2.3 2.6, 3.2 4.6</td>
</tr>
</tbody>
</table>
similar in both the systems while the ion density increases significantly for S2LC beyond second coordination shell.

However, the near neighbor environment around the negatively charged functional groups (Fig. 5) of ASP and GLU residues are mostly populated by Ca\(^{2+}\). The RDF between carbonyl-oxygen (OD) of ASP and Ca\(^{2+}\) shows two sharp maxima at 2.2 and 4.2 Å, separated by a zero-intensity plateau of 1 Å width which signifies a crystal-like near neighbor ordered structure up to ∼5 Å from ASP. In addition, the \(g_{CG-Ca}(r)\) (Fig. 5) also shows two high-intensity peaks at ∼2.2 and 3.2 Å signifying two preferred geometries of the cations with respect to the COO\(^-\) group of ASP/GLU, bidentate and monodentate respectively. The radial distributions of \(C_Z\) and \(O_w\) (Fig. 5) around COO\(^-\) group of ASP/GLU are poorly structured with coordination numbers less than 1 up to the first minima at ∼5.0 Å for both the distributions.

Density map

**Distribution of Ca\(^{2+}\) and CO\(^3\(^{2-}\) ions around arginine.**

Among the three sp\(^3\)-hybridized nitrogen atoms (NH1, NH2, and NE) of the Gdm\(^+\) group of arginine side chain, NH1 and NH2 can form two h-bonds and NE can establish a single h-bond with the CO\(^3\(^{2-}\) ions. MD trajectory-based visual inspection of the local environment of arginine residues reveals that each arginine is coordinated to two near neighbor CO\(^3\(^{2-}\) ions to form the ‘twin N-twin O’ (TNTO) h-bond geometry (Fig. 6a). In this geometry, NH2 forms two h-bonds with these CO\(^3\(^{2-}\) ions, while NH1 and NE form single h-bonds with them. The CO\(^3\(^{2-}\) is mostly in the bidantate geometry with the nitrogen atom pairs forming in-plane h-bonds with the Gdm\(^+\) plane. The average length of arginine-CO\(^3\(^{2-}\) h-bonds is ∼1.65 Å.

The calculated atomic density distribution (Fig. 6), i.e. density map, of CO\(^3\(^{2-}\) around Gdm\(^+\) shows two density peaks of C\(_Z\) atoms within the CZ-C\(_Z\) distance cutoff of 4.3 Å, the first minimum in \(g_{CZ-C}(r)\) (Fig. 4). In contrast, \(O_w\) density map within the first minimum of \(g_{CZ-O_w}(r)\) at 3.5 Å shows four high-density regions along the N–H bond directions.

In general, the non-polar compounds approach the arginine side chain from the directions normal to Gdm\(^+\) plane, while the polar compounds, e.g. water, carboxylates/carbonyls, form in-plane h-bonds\(^{40,41}\) with Gdm\(^+\). Our results reveal that in the lysozyme–CaCO\(_3\) systems, the CO\(^3\(^{2-}\) ions replace the water molecules from the first coordination shell of arginine to form the TNTO coordination geometry. The second coordination shell is primarily formed by the Ca\(^{2+}\) ions with approximately three near-neighbor (NN) cations for each CO\(^3\(^{2-}\) (Fig. 6). The observed distribution of NN Ca\(^{2+}\) and CO\(^3\(^{2-}\) around arginine facilitates the formation of further layers of ions leading to their aggregation around the surface-exposed arginine side chains of lysozyme. In S1,2LC, the coordination number of water oxygen (O\(_w\)) around arginine is less than that of in LW. The aggregation of ions around Gdm\(^+\) in lysozyme–CaCO\(_3\) systems by expelling water molecules indicates that arginine residues act as nucleation sites for the formation of biominerals. The observed water expulsion at the protein-ACC interface is in-line with Freeman et al.\(^{16}\)’s work that showed that the water population at the ACC-protein interface is much less than that at the calcite-protein interface.

**Distribution of Ca\(^{2+}\) and CO\(^3\(^{2-}\) ions around aspartate/glutamate.**

The distribution of ions and water molecules around the negatively charged carbonyl groups of aspartate or glutamate side chains is entirely different from that of arginine side chain. In this case (Fig. 7), the Ca\(^{2+}\) ions in the first coordination shell are located mostly along the C=O bond and along the bisector vector of the angle OD1–CG–OD2. The density distribution along the bisector is similar to the most stable geometry...
of isolated cluster containing acetate (AcO⁻), Ca²⁺, and CO₃²⁻, where Ca²⁺ is coplanar to the COO⁻ plane and equidistant from the oxygen atoms. However, the cation population along the lone pairs of the oxygen was not observed in the H₂O containing clusters of acetate. The ensemble averaged coordination number of Ca²⁺ around CG is ~0.9 up to CG–Ca²⁺ distance of 7.0 Å. Unlike arginine side chain, the first layer of ions around COO⁻ is surrounded by water molecules (Fig. 7).

The binding energies of Ca²⁺–CO₃²⁻ and Ca²⁺–H₂O complexes are ~520.8 kcal mol⁻¹ (ref. 25) and 56.9 kcal mol⁻¹, respectively. Among these two attractive interactions, the hydrophilic nature of calcium dominates which results in the...
separation of the CO$_3^{2-}$ from its Ca$^{2+}$ counterpart via a water layer (Fig. 7) in between. These water molecules are ‘tightly bound’ by both types of ions. As the life time of Ca–H$_2$O h-bond is large, the possibility of breaking this h-bond within the simulation time limit is slim. Thus, the removal of water from the nearest hydration layer and aggregation of ions around aspartate/glutamate seems unlikely.

A convoluted picture of the distribution of ions and the ion density with reference to a given side chain residue are presented here by charge density plots (Fig. 8). The average charge density ($\rho$) around arginine demonstrates sharp oscillations around the neutral axis, $\rho = 0$, up to ~7 Å, indicating the formation of four alternating layers of cations and anions. However, the charge density around the COO$^-$ bearing groups (aspartate/glutamate) shows only one prominent positive peak with a much weaker next layer of anions before the distribution merges to the neutral axis. Thus, we can conclude that arginine may play the major role in the nucleation and growth of CaCO$_3$ crystal in lysozyme, whereas the other two residues may facilitate the protein to bind to the positively charged surface of the CaCO$_3$ crystal.

### Hydrogen bond dynamics

Fig. 9 shows the calculated hydrogen bond time correlation functions, $C_{hb}(t)$, for S1$_{2LC}$ and S2$_{2LC}$ systems. The time correlation for water-arginine h-bonds ($C_{hb}(t)$) decays much faster than that for CO$_3^{2-}$–arginine h-bonds ($C_{hb}(t)$) in both the systems. The slow decay of the $C_{hb}(t)$ implies that the CO$_3^{2-}$ ions remain h-bonded for an extended period of time, whereas the relatively faster decay of $C_{hb}(t)$ signifies frequent breaking and formation of h-bonds with water. The higher concentration of ions in S2$_{2LC}$ results in further slowing down of the decay of $C_{hb}(t)$ suggesting that the ions are held tightly to the protein surface. These tightly bound CO$_3^{2-}$ ions hinder water molecules to establish strong h-bonds with arginines and thereby promote faster decay of $C_{hb}(t)$ in S2$_{2LC}$.

Each time correlation function was fitted with a multi-exponential function to obtain average time constant or relaxation time ($\tau_{hb}$, Table 3). The values of $\tau_{hb}$ for CO$_3^{2-}$ are orders of magnitude higher than those for water, which is indicative of higher residence times of CO$_3^{2-}$ ions in the vicinity of the nucleation sites. The shorter relaxation times of ‘loosely bound’ water molecules may be necessary to facilitate the dehydration of the ACC patches on the protein surface.

### Free energies for the association of ions near arginine side chain

Free energy profiles were calculated to understand the energetics of ion association, growth of ion density, and the expulsion of water at the protein surface. As we have already studied the structure and energetics of ions in the first coordination shell of various side chain residues using radial distribution functions, density maps, and cluster analysis, our focus here is on the energetics of higher coordination shells. Since the binding affinity between arginine and CO$_3^{2-}$ depends on the thickness of the ionic-layer present on the protein surface, it is likely that the role of the ion–ion interactions would dominate over arginine–CO$_3^{2-}$ interactions in controlling the growth beyond some threshold thickness of ACC.

The calculated free energy profile estimates the energy cost to bring an anion or a Ca$^{2+}$–CO$_3^{2-}$ ion pair that is located away from the protein surface to the vicinity of arginine side chain. The separation ($r_{CV}$) between the C$_A$ atom of a distant CO$_3^{2-}$ and the CZ atom of ARG61 was chosen as a collective variable (CV) to describe arginine–Ca$^{2+}$–CO$_3^{2-}$ association. Fig. 10 shows the calculated free energy profile as a function of $r_{CV}$ in the range from 6.0 Å to 20.0 Å.

In the initial configuration, the chosen CO$_3^{2-}$ ion pairs with a neighboring Ca$^{2+}$ to form an isolated CaCO$_3$ unit surrounded by water. This particular Ca$^{2+}$ ion was observed to be bound to
the selected CO$_3^{2-}$ throughout the range of $r_{CV}$. The free energy profile shows a shallow and broad minimum at $r_{CV} = 11.4$ Å (Fig. 10a). The binding energy ($\Delta F$) between the dissociated state at $r_{CV} = 20$ Å and the global minimum is $\sim 8.4$ kcal mol$^{-1}$. This value compares well with the binding energy ($\sim 10$ kcal mol$^{-1}$) between OC-17 (a hen eggshell protein) and calcite surface, where arginine residues serve as the anchors to the crystal surface.$^{18}$ However, this barrier height ($\Delta F$) is 4 kcal mol$^{-1}$ higher than that estimated for a single CO$_3^{2-}$

**Fig. 8** (Up) Charge density ($\rho$) around (a) CZ of arginine and (b) CG of aspartate in S1$_{LC}$ (black) and S2$_{LC}$ (red). (Down) Schematic representation of the ionic layers around arginine and aspartate. Color scheme for ball and stick representation: nitrogen (blue), carbon (gray), oxygen (red)

**Fig. 9**  Time correlation functions of the hydrogen bonds between the polar head groups of arginine with CO$_3^{2-}$'s in S1$_{LC}$ (red) and S2$_{LC}$ (blue) and with water in S1$_{LC}$ (black) and S2$_{LC}$ (gray). Each plot was fitted with multiexponential function (dashed lines) to calculate h-bond relaxation time.

<table>
<thead>
<tr>
<th>System</th>
<th>Function</th>
<th>$\tau_{hb}$ (ns)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1$_{LC}$</td>
<td>$C_{hb}(t)$</td>
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<td>S2$_{LC}$</td>
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<td>154.53</td>
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<td>$C_{ow}(t)$</td>
<td>0.12</td>
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around a Gdm⁺ functional group and that between an isolated arginine residue and ACC, which can be attributed to the inhomogeneous environment of the carbonate ion in the present study. The interactions between an anchoring agent (here, arginine) and the ions (Ca²⁺ and CO₃²⁻) play a crucial role in the initial stages of the ion aggregation, but further growth of ACC is generally governed mainly by the interactions between ions and water molecules. Thus, the barrier height reported here is an approximate estimate on the energetics for the growth of existing ion aggregates irrespective of the anchoring agent.

Further analysis of the metadynamics trajectory shows that the addition of a single CO₃²⁻ in the arginine environment increases the ion-cluster size by at least three more CaCO₃ pairs via near-neighbor electrostatic interactions (Fig. 10b).

**Conclusions**

In this work, we report protein-mediated formation of amorphous calcium carbonate (ACC) from the free ions using molecular dynamics (MD) simulations of a hen eggshell protein, lysozyme, in ionic solutions of calcium and carbonate. Our investigation provides comprehensive understanding on the role of selected side chain residues of lysozyme that facilitate the ion association to develop a dense amorphous phase of ions driven primarily by the strong electrostatic interactions between the functional group and its respective counter ions (Ca²⁺ or CO₃²⁻). The radial distribution functions (RDFs) among the ions and water in the ionic clusters around lysozyme are in good agreement with that in the hydrated ACC resembling a similar local environment in both the systems. Three key residues, arginine, glutamate, and aspartate, serve as potential nucleation sites for the crystallization of CaCO₃. However, the structure and energetics results indicate that the influence of arginine dominates over the rest, which could be a general scenario in all positively charged proteins, e.g. lysozyme, responsible for biomineralization.

Two model systems of lysozyme were generated with different ion concentrations, and both show essentially similar results, apart from a denser ACC phase in the solution with higher concentration of ions. The three dimensional density profile of ions around arginine shows that, on the average, each arginine is coordinated by at least two CO₃²⁻ ions forming four h-bonds with the Gdm–NH₂-groups making a twin-nitrogen-twin-oxygen (TNTO) geometry. This strong association with the nearest neighbors initializes the foundation for the successive alternating layers of positive and negative charges replacing the solvent water molecules. Moreover, the charge density plots also confirm this layering of charges suggesting that arginine can play a pivotal role in the nucleation of CaCO₃ crystals. To elucidate the growth of ionic layers around a given arginine residue, we performed metadynamics calculations to estimate the required free energy for ion association in the higher coordination shell. The computed free energy profile shows a barrier height of ~8 kcal mol⁻¹ between the bound and unbound states of a CO₃²⁻ ion. We also observed that the addition of a single CO₃²⁻ may fetch few more Ca²⁺ and CO₃²⁻ pairs via near-neighbor electrostatic interactions and thus increases the size of the ion cluster around arginine. In contrast, the local environment around glutamate and aspartate retain the strongly bound solvent water through hydrophilic interactions with Ca²⁺ that inhibit the formation of next layer of CO₃²⁻ ions.

**Conflicts of interest**

There are no conflicts to declare.

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**References**

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J. Lee, et al., CHARMM-GUI input generator for NAMD, GROMACS, AMBER, OpenMM, and CHARMM/OpenMM


