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Functional mimicry of sea urchin biomineralization proteins with CaCO₃-binding peptides selected by phage display?

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The intricate process of biomineralization, e.g. in sea urchins, involves the precise interplay of highly regulated mineralization proteins and the spatiotemporal coordination achieved through compartmentalization. However, the investigation of biomineralization effector molecules, e.g. proteins, is challenging, due to their very low abundance. Therefore, we investigate the functional mimicry in the bioinspired precipitation of calcium carbonate (CaCO₃) with artificial peptides selected from a peptide library by phage display based on peptide-binding to calcite and aragonite, respectively. The structuredirecting effects of the identified peptides were compared to those of natural protein mixes isolated from skeletal (test) structures of two sea urchin species (Arbacia lixula and Paracentrotus lividus). The calcium carbonate samples deposited in the absence or presence of peptides were analyzed with a set of complementary techniques with regard to morphology, polymorph, and nanostructural motifs. Remarkably, some of the CaCO₃-binding peptides induced morphological features in calcite that appeared similar to those obtained in the presence of the natural protein mixes. Many of the peptides identified as most effective in exerting a structure-directing effect on calcium carbonate crystallization were rich in basic amino acid residues. Hence, our in vitro mineralization study further highlights the important, but often neglected, role of positively charged soluble organic matrices associated with biological and bioinspired CaCO₃ deposition.

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Introduction

Proteins are fundamental components of life that perform a wide range of functions. One of their key roles is to regulate

metabolic and cellular processes as enzymes, hormones, and antibodies through protein-protein interactions. In addition to these organic-organic interactions, proteins are also involved in the formation of biogenic minerals, referred to as biominerals, which characteristically represent organic-inorganic hybrid materials with complex hierarchical structures and remarkable properties.1 In biomineralization processes, the binding partners of the proteins are typically inorganic components including solvated ions, amorphous mineral species or the facets of crystals.2 Biominerals are found in a wide variety of organisms, ranging from unicellular microorganisms (e.g. coccolithophores3) to complex multicellular organisms, including humans. They play an important role in the formation of hard tissues such as vertebrate bones,⁵ teeth,⁶ and mollusk shells, thus providing structural support and protection to the organism. The formation of biominerals is a complex process that involves the molecular recognition of an inorganic target, such as a substrate surface, ion, or particle, by a protein. This molecular recognition is in many cases essential for the deposition, stabilization and assembly of inorganic mineral species and the transformation of precursor phases. As a tight

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interaction between water-soluble proteins and the (forming) mineral is often a prerequisite for a structure- and processdirecting effect, it is not unusual for the organic matrix to become occluded into the final crystalline product, 8,9 thus leading to the formation of organic-inorganic hybrid materials, whose mechanical properties are strongly affected by the incorporation of the soft organic component. 10,11 By studying biomineralization processes at the molecular level and translating fundamental concepts of biological crystallization control (e.g. mineral deposition directed by soluble and insoluble organic matrices, compartmentalization, isotropic amorphous species) into synthetic model systems, 12-15 scientists aim for a better understanding of how the underlying design principles can eventually be harnessed for biomedical applications, such as implants or bone substitutes.

Calcium carbonate (CaCO₃), in particular, is a ubiquitous inorganic component of many biominerals such as in sea urchin skeletons¹⁶ and mollusk shells.¹⁷ In a geological context, CaCO₃ occurs in three anhydrous crystalline polymorphscalcite, aragonite and vaterite - along with hydrated and amorphous modifications. The selectivity of the polymorph largely depends on environmental parameters (e.g. pressure, temperature), with calcite being the thermodynamically most stable crystalline phase under ambient conditions. Mineral deposition in the presence of biomineralization proteins, by contrast, can lead to the stabilization of polymorphs that would usually not form under the given environmental conditions or even induce the formation of stable or transient amorphous phases. 18,19 The specific protein-inorganic interactions along with the physicochemical boundary conditions for mineral deposition often lead to a hierarchical organization spanning several length scales from the macroscopic level down to the nanoscale. In this way, biomineralization can also result in nanogranular crystalline structures with long-range texture, such that the protein-interconnected crystallites are in nearperfect crystallographic register and the superstructures show a quasi-single crystalline character in its entity. Biologically, such a highly ordered arrangement of the mineral building blocks, termed mesocrystal,20 has been observed, for example, in sea urchin test plates and spines,21 which show single-crystalline behavior in polarized light microscopy and XRD analysis.²²

Sea urchins have long been used as model organism to study biomineralization processes. The biological deposition of mineral species is a spatiotemporally regulated process, i.e. sea urchins are capable of precipitating calcium carbonate phases with different structural features (e.g. spine, test, teeth) optimized for specific functions spatially separated from each other and at different stages of development (larvae, mature). 23,24 The resulting mineralized structures consists of $CaCO_3$ (>95 wt%), MgCO₃ (5 wt%) and biomolecules (0.1 wt%), mainly proteins but also glycoproteins and polysaccharides.²⁵ Although, more than 100 proteins are known to be associated with mineralized sea urchin tissues, their exact function in directing the crystallization process often remains obscure,²⁶ mostly because the low concentration of the organic matrix in mineralized tissues makes it difficult to extract proteins in

sufficient quantities for their characterization. However, some examples of proteins involved in biomineralization have indeed been isolated and the corresponding genes were identified by reverse genetics of CaCO₃-mineralizing organisms, including sea urchin species. 27,28 Moreover, it was shown that sea urchins from different orders, namely the black sea urchin (Arbacia lixula, Arbacioida) and the purple sea urchin (Paracentrotus lividus, Camarodonta), show similarities in their protein composition, ²⁹ thus pointing to a potential correlation between the amino acid sequence and the functionality in terms of interfering with calcium carbonate mineralization. Different protein families were shown to be associated with the biomineralization processes. 26,29-32 While the majority of proteins that were identified in biogenic calcium carbonate, specifically mollusk shell proteins^{33–35} and sea urchin proteins,²⁹ are negatively charged (i.e. rich in Asp and Glu residues), basic protein sequences (i.e. positively charged domains) have also been reported.30

The fundamental principles of biomineralization, such as confinement, crystallization pathways via amorphous precursors, and in particular, the control of mineralization by soluble and insoluble organic matrices, have been successfully transferred to synthetic systems. To investigate the effects of artificial soluble structure-directing additives, 36 in vitro model systems for the mineralization of CaCO3 have been designed. As the literature on biomineralization-associated proteins has long been dominated by reports on acidic protein domains, bioinspired synthetic approaches aiming to mimic structural features of biological crystallization often use negatively charged polyelectrolytes to control mineralization pathways and products.37 However, there are a growing number of publications discussing the effects of cationic (i.e. aminefunctionalized) polymer additives and their role is increasingly being recognized. 38-40 In bioinspired polymer-mediated mineralization, synthetic polyelectrolytes have been shown to exert multiple effects. Examples include the stabilization of transient amorphous precursors of hydrated colloids even in the presence of minute amounts of organic molecules, 41 promotion of the formation and stabilization of an amorphous precursor phase by suppressing crystalline growth, 42 altering the pH value of the reaction solution, 41 or affecting the composition and water content of the mineral phase, resulting in a manifold of different crystal morphologies.41

While a large variety of soluble additives, have been investigated as structure-directing agents in calcium carbonate precipitation, including ions like $\mathrm{Mg}^{2+},^{43}$ small molecules such as individual amino acids,44 and polyelectrolytes,45 block copolymers, 42 as well as acidic and basic (poly) amino acids, 43 a specific molecular interaction between the additive and the inorganic phase is often not directly addressed experimentally. To gain access to a sufficient amount of biomolecules with a specific binding capacity to inorganic materials, so called inorganic-binding peptides, the phage display method has been successfully applied.46,47 Phage display uses a pseudorandomized peptide library presented on the surface of M13 phages (bacterial viruses). Peptides are selected from this

peptide library based on their ability to bind to a target material, where the interaction is based on the specific amino acid sequence of the peptide. Such inorganic-binding peptides have been shown to mineralize engineering materials such as zinc oxide, thereby controlling material properties. ^{13,14} In a biological context, peptides that show a significant effect on the morphology or polymorphism of the synthetic organic/inorganic hybrid materials may be indicative of the sequence of natural mineralization proteins. In this way, a feedback loop between the biogenic minerals and the synthetic model systems is generated, which may help to identify previously unknown proteins or protein domains in biomineralization proteins.

In our study, we present a systematic investigation into the bioinspired precipitation of calcium carbonate using selected peptides obtained from a peptide library through phage display, based on their binding affinity to geological calcite and aragonite. To assess their structure-directing effects, we compared them with natural protein mixes extracted from the calcitic tests of two distinct sea urchin species, Arbacia lixula and Paracentrotus lividus, belonging to different orders. We applied the calcite- and aragonite-binding peptides, along with the biologically derived soluble proteins, as organic additives in the in vitro mineralization of CaCO3 and determined their influence on the (nano-)morphology and polymorphism of the resulting crystals with scanning electron microscopy (SEM), X-ray diffraction (XRD), and small-angle X-ray scattering (SAXS). The most effective peptides that led to calcite structures as seen for the natural biomineralization proteins, were basic peptides. This observation further emphasizes the importance of basic soluble organic molecules in biomineralization.

Materials and methods

Materials

Agar agar, bovine serum albumin (BSA), $CaCl_2 \cdot 2H_2O$, glycine, isopropyl β -d-thiogalactoside (IPTG), $(NH_4)_2CO_3$, NaCl, NaOCl, polyethylene glycol-8000, Tris–HCl, tryptone, yeast extract, 5-Bromo-4-chloro-3-indolyl- β -d-galactoside (XGal) were purchased from Roth (Germany) or Sigma–Aldrich (Germany) and were used as received. All solutions were prepared using ultrapure deionized water (ddH₂O, 18.2 M Ω , Arium 611UF, Sartorius).

Phage display

Aragonite- and calcite-binding peptides were isolated from a random 12-mer peptide library (New England Biolabs, Inc.) by the phage display technique. The peptide library is expressed as N-terminal fusion protein with the minor coat protein p3 of M13 phages. The peptides are separated by a short spacer sequence (GGGS) from the p3 coat protein. General phage methods were conducted according to the manufacturer's recommendations.

For phage display experiments, geological aragonite (Alpha Aesar, Product no. 42523, LOT N04A029) and calcite (Alpha

Aesar, Product no. 44520, LOT N04A030) substrates were applied as received from the manufacturer. Aragonite and calcite substrates of approximately 9 mm² were applied for the biopanning procedure. The target substrates were washed twice in Tris buffered saline (TBS), pH 7.5 supplemented with 0.1% Tween 20. The random phage library (1.5×10^{11}) phages was incubated with the aragonite and calcite substrates, respectively, under light agitation in TBS, 0.1% Tween 20 for one hour at RT. Unbound phages were removed from the substrate by washing 10 times in TBS, 0.5% Tween 20. The remaining phages were eluted applying 1 mL Glycine Elution Buffer. The eluate was neutralized with 150 µL of 1 M Tris-HCl, pH 9.1. The eluted phages were amplified in E. coli ER2738 and purified by polyethylene glycol-8000/sodium chloride (PEG/NaCl) precipitation. In the following bio panning rounds at least 1.3 \times 10¹⁰ phages of the restricted library were applied. To enrich the specificity of aragonite and calcite-binding sequences, in total 5 panning rounds were carried out. From each experiment, 50 randomly selected phage clones were analyzed by DNA sequencing.

Extraction of CaCO₃-associated proteins from sea urchin skeletons

The extraction of CaCO₃-associated protein was performed according to Kanold et al.25 Skeletons from the sea urchin strains Arbacia lixula and Paracentrotus lividus were mechanically cleaned. Adherent organic debris was removed by incubation under agitation in NaOCl (0.26% active chlorine) for 72 h. Afterwards, skeletons were extensively rinsed in ddH₂O and dried at 37 °C and were pulverized in a mortar. For decalcification, 30 g of skeleton powder were suspended in 70 mL ddH₂O and 400 mL of cold 10% acetic acid were drop-wise added with an electronic burette, flow rate of 100 µL every 5s, at 4 °C to the final pH 4 to 5. Non-solubilized material, acid insoluble matrix (AIM), was removed by filtering (pore size 200 µm) and centrifugation 30 min, 4500 rpm at 4 °C. The AIM pellet was washed in ddH₂O and stored at −80 °C. The soluble fraction, acid soluble matrix (ASM), was filtered (pore size 0.45 µm) and concentrated with a centrifugal concentrator with 10 kDa molecular weight cut off size (MWCO) (Vivaspin 20, Sartorius, Göttingen, Germany) for 45 min at 4500 rpm and 4 °C. The concentrated ASM sample was dialyzed 2 kDa MWCO at 4 °C for 40 h and lyophilized at 1 mbar and -70 °C within three days. The extraction of acid-soluble matrix proteins (ASM) from sea urchin skeletons was conducted through four repetitions, yielding consistent results in terms of protein extraction efficiency. The protein extraction process was carried out using 15 to 30 grams of initial material. Specifically, from A. lixula, 1.4, 2.0, and 6.1 milligrams of ASM were successfully extracted, representing a range of 0.01 to 0.02 weight percent (wt%). In the case of P. lividus, 3.5 milligrams of ASM (0.01%) were obtained from 33 grams of starting material. These results align well with previous studies focused on the extraction of sea urchin proteins, both in terms of the total matrix protein content and the yield of ASM. 16,25,29

ASM and AIM protein fractions were analyzed by 1D SDS/PAGE (12% acrylamide). 10 mg AIM sample was mixed with

100 μL 2x Laemmli buffer and 100 μL ddH₂O. The AIM-PAGE samples represented only a fraction of the AIM sample since some proteins were not solubilized in Laemmli buffer. ASM samples were resuspended in 2x Laemmli buffer:ddH₂O (1:1) to a final concentration of 2 mg mL⁻¹. Gels were silver stained afterwards.

CaCO₃-binding peptides

CaCO₃-binding peptides for mineralization experiments were purchased from EMC microcollections GmbH (Germany) and used as received from the supplier. The purity of the peptides was assessed using high-performance liquid chromatography (HPLC) and was found to be greater than 80%.

Secondary structure prediction of CaCO₃-binding peptides

Starting from the amino acid sequence in fasta format three secondary structure prediction servers are used. 48-51 De novo secondary structure predictions are performed with the PEP-Fold server. 52-54 PEP-Fold predicts first structural alphabet (SA) letters from the sequence using a hidden Markov model approach.55 These fragments are assembled by a greedy procedure driven by a modified optimized potential for efficient structure prediction (OPEP) coarse grained force field⁵⁶ energy score followed by a clustering procedure.

In vitro mineralization of CaCO₃

CaCO3 was precipitated by slow diffusion of gaseous NH3 and CO₂ formed by the decomposition of solid ammonium carbonate into a 10 mM CaCl₂ solution according to a protocol described by Albeck et al. 57 The aqueous CaCl₂ stock solution was freshly prepared in ddH2O and sterile-filtered (pore size 0.2 µm) prior to use. The mineralization was performed in 16 well chamber slides in a total volume of 500 µL per well (Lab-Tek, Thermo Fisher Scientific, USA). To ensure gas exchange, a hot cannula was used to poke one hole per well into the lid of the chamber slide. Organic additives, ASM and CaCO₃-binding peptides, were diluted to final concentrations of 500, 100, 30, 20, 10, 5, and 1 μg mL⁻¹. Solutions with 2 mg mL⁻¹ of the structure-directing organic matrix were additionally prepared for the CaCO3-binding peptides. The chamber slides were placed under vacuum in a sealed desiccator, containing a 5 cm Petri dish with solid ammonium carbonate, and the prepared crystallization solutions were incubated in the forming vapor phase at 4 °C for three days. Afterwards, the chamber slides were removed from the desiccator, the supernatant was decanted and the CaCO₃ crystals were washed twice with ddH2O and dried at 37 °C. All mineralization samples were prepared in triplicate.

Mineralization products were analyzed by scanning electron microscopy (SEM), X-ray diffraction (XRD), and small-angle X-ray scattering (SAXS).

CaCO₃ sample characterization

Scanning electron microscopy (SEM). Mineralization products deposited on the glass support of chamber slides or immobilized on Si-wafers, respectively, were sputtered with

0.5 nm platinum/palladium. Scanning electron microscopy (SEM) (Zeiss, Oberkochen, Germany) images were recorded with a Zeiss GEMINI scanning electron microscope applying in general 3 kV acceleration voltage and secondary electron (SE) detection.

X-ray diffraction (XRD). The phase analysis of the in vitro synthesized CaCO₃ particles was performed on powder X-ray diffraction patterns obtained with a Rigaku SmartLab 3 kW X-ray diffraction system equipped with a copper anode (Rigaku Europe SE, Neu-Isenburg, Germany). The samples were measured on a low-background silicon plate, to avoid any reflexes of the sample holder or in a capillary. The diffractograms of samples were compared to diffraction patterns of reference crystals (without peptide) and geological calcite and aragonite, respectively.

Small angle X-ray scattering (SAXS). CaCO₃ powders of selected specimen types (control calcite and CaCO3 deposited in the presence of peptide A-4-09) were sandwiched between two strips of adhesive tape (Scotch Magic) and mounted on a sample holder. Scattering patterns were collected using a Nanostar instrument (Bruker AXS, Karlsruhe, Germany) equipped with a single photon counting area detector (Hi-Star) accumulating the scattered intensity obtained from each sample over a period of 1 hour. Radiation with a wavelength of $\lambda = 1.54 \text{ Å}$ $(Cu-K_{\alpha})$ was produced by a copper sealed tube fine focus source and monochromatized with a parabolically bent graded multilayer Göbel mirror. To cover a sufficiently broad angular range, measurements were generally carried out in two configurations, for which the detector was placed at a distance of either 105 cm or 26 cm to the sample. Calibration was performed using a silver behenate standard.

All two-dimensional data sets were reduced to onedimensional traces of the modulus of the scattering vector q $(q = 4\pi \sin(2\theta/\lambda))$, where 2θ represents the scattering angle) vs. the intensity I(q) and corrected for sample transmission and background intensity (by subtraction of an empty tape pocket). Additionally, the Laue background (I_{Laue}) was subtracted from the data based on a Porod fit $(I(Q) = \frac{P}{Q^4} + I_{\text{Laue}}, P \text{ being the})$ Porod constant) prior to T-parameter analysis as previously reported $(T = \frac{4 \cdot \tilde{I}}{\pi \cdot P} = \frac{4\phi \cdot (1 - \phi)}{S}$, where $\tilde{I} = \int_0^\infty I(q) \cdot q^2 dq$ represents the integral intensity, and S and ϕ being the specific surface area and the volume fraction of the scattering objects).⁵⁸⁻⁶⁰ Due to the strong scattering contribution of the large external surfaces of the powder grains in the low q limit, data points with $q < 0.3 \text{ nm}^{-1}$ were approximated by a rectangle curve when calculating the integral intensity \tilde{I} . For dilute platelet-shaped heterogeneities within a sample, the Tparameter, which generally represents an average measure for the volume fraction per surface area, can be interpreted in terms of the thickness D of the discs with T = 2D.⁶¹

Results

Natural sea urchin proteins are often present in low abundance within the hard tissue, and their isolation can be challenging,

potentially leading to loss or degradation. To circumvent these limitations, we turned to synthetic $CaCO_3$ binding peptides that exhibit mineralization properties similar to those of natural sea urchin proteins. These peptides were carefully selected from a random peptide library based on their strong binding affinity to mature crystal material, effectively mimicking the biomineralization function observed in sea urchins. By characterizing and studying these synthetic peptides, we aim to gain insights into the nature and composition of the elusive natural biomineralization proteins.

Phage display and selection of peptides

Peptides, binding to geological aragonite or calcite were selected from a peptide library by phage display. Phage clones, expressing individual peptides, were analyzed after the 4th and 5th biopanning round. In total, 33 aragonite-binding peptides and 39 calcite-binding peptides were identified from each 50 analyzed phage clones (Tables S1 and S2, ESI†).

Five aragonite-binding peptide sequences were repeatedly isolated. The calculated isoelectric points (pI) of the isolated aragonite-binding peptides ranged from 4.47 to 12.3, with two accumulations at pH 8 (22.7% of the peptides) and pH 10 (29.5% of the peptides). The calcite-binding peptides were more diverse compared to the aragonite-binding peptides. The pI values of the calcite-binding peptides spanned a range from 3.81 to 11.12, with an accumulation at pH 7 (28.2% of the peptides). The amino acid residue frequencies of both, the aragonite- and the calcite-binding peptides, were referenced to the frequencies of the naïve peptide library (Table 1). The quotient of an amino acid frequency in the isolated peptide pool and in the initial peptide library is indicative of its enrichment (criterion: quotient > 1.25) or depletion (criterion: quotient < 0.75). Enriched amino acid residues may have a beneficial effect on the binding of peptides, while depleted amino acid residues might have an adverse effect on the interaction. Aragonite-binding peptides showed an enrichment of lysine (K) and to a lesser extend of histidine (H). The incorporation of lysine into CaCO₃ had been shown previously in in vitro mineralization experiments, where only lysine was incorporated to detectable amounts, while histidine and arginine were below the detection limit.⁶² Furthermore, negatively charged amino acids were depleted in both, aragonite- and

calcite-binding peptides. This is remarkable, since acidic proteins are known to control CaCO₃ mineralization in biogenic systems,⁶³ including marine organisms, like sponges⁶⁴ and mollusks.³⁴ Moreover, the majority of the proteins associated with the tests of *A. lixula* and *P. lividus* are also mainly acidic.^{16,25} We also identified some similarities between the selected peptides and natural biomineralization proteins. Glycine (G) and tyrosine (Y) were enriched in aragonite- and calcite-binding peptides. These amino acids are also present in increased amounts in the proteins^{16,25,29} of *Pinctada fucata*, the Akoya pearl oyster, a marine mollusk species.^{65,66}

Calcium carbonate precipitation in the presence of biogenic protein mixtures and peptides

In a next step, the structure-directing effect of biomineralization-associated proteins isolated form sea urchin tests and CaCO₃-binding peptides on the in vitro mineralization of CaCO₃ were investigated and compared. In the mineralization experiments, only the designed peptides with demonstrated mineral interaction capabilities were included. The preselection process via phage display enabled to avoid the extensive screening of random peptides, ensuring that only effective peptides were utilized. The acid-soluble biomineralization proteins of sea urchin tests were characterized by 1D polyacrylamide gel electrophoresis (SDS-PAGE) and 2D PAGE. 16,25,29 Briefly, the proteins accumulated in a range between 25 kDa and 40 kDa with distinct focal points at 40 kDa, 36 kDa, and 27 kDa. The isoelectric points of the majority of ASM from A. lixula were within the acidic to neutral range (pH 3-6.5), with only a very small fraction of low molecular weight ASM having a pI > pH 9.29 In case of P. lividus more proteins in the basic pH regime were identified.¹⁶ In the experimental set up, CaCO₃ was mineralized in vitro from aqueous CaCl2 solutions where the precipitation was induced by the influx of atmospheric CO2 and NH₃ through the slow decomposition of ammonium carbonate in an enclosed chamber.57

(i) Biogenic protein mixtures. The morphology of calcium carbonate particles mineralized in the presence of protein mixtures form *A. lixula* and *P. lividus* tests was investigated (Fig. 1 and 2). The protein mixtures were applied in a wide concentration range (1 to 500 μg mL⁻¹) and the particle morphologies were analyzed by X-ray diffraction (XRD) and

Table 1 Frequency of amino acid residues in aragonite- and calcite-binding peptides. The inorganic-binding peptides were analyzed after the 4th and 5th biopanning round and are separately indicated in the table. The frequencies are given as the quotient of the amino acid frequency in the selected peptides and the initial peptide library (NEB). Enriched amino acid residues (criteria: quotient > 1.25) are highlighted dark grey, depleted amino acid residues (criteria: quotient < 0.75) are highlighted light grey. Amino acids are represented in one-letter code

Clb. a			basic		sm	nall	hydrophobic			bic		acidic		amide		nucleophilic			aromatic		
Substrate		Н	K	R	G	Α	V	L	ı	Р	М	D	Ε	Ν	Q	S	Т	C a)	Υ	F	W
Aragor	nite, 4 th	1.23	2.88	1.18	1.71	0.74	0.57	0.87	1.39	1.16	1.18	0.50	0.26	0.79	0.60	0.97	0.73	-	1.31	0.84	1.14
Aragor	nite, 5 th	1.61	3.40	0.89	1.60	0.40	0.31	0.58	1.05	1.41	1.60	0.64	0.47	1.55	0.35	1.01	0.54	-	0.83	0.36	1.89
Calcit	te, 4 th	0.83	0.74	1.48	2.40	1.10	0.89	0.82	0.92	0.71	0.67	1.61	0.33	1.21	1.36	1.15	0.88	-	1.06	0.53	1.10
Calcit	te, 5 th	1.06	0.40	0.95	1.71	1.57	1.28	0.42	0.82	1.05	1.07	0.60	0.53	1.09	1.20	1.06	0.75	-	1.54	0.51	2.02

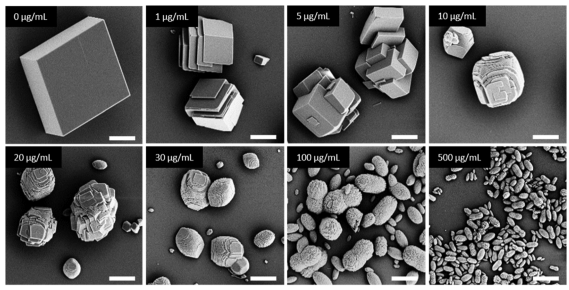


Fig. 1 Effect of skeletal proteins from Arbacia lixula on the in vitro mineralization of CaCO3. Extracted CaCO3-associated proteins from A. lixula skeletons were applied in the bioinspired mineralization of CaCO₃ via a slow diffusion method. The mineralization was performed in the presence of protein concentrations ranging from 1 μ g mL⁻¹ to 500 μ g mL⁻¹. In protein-mediated precipitation, the deposited CaCO₃ particles showed a complex morphology, whereas regular, rhombohedral particles were synthesized in the absence of proteins. Scale bars 10 µm.

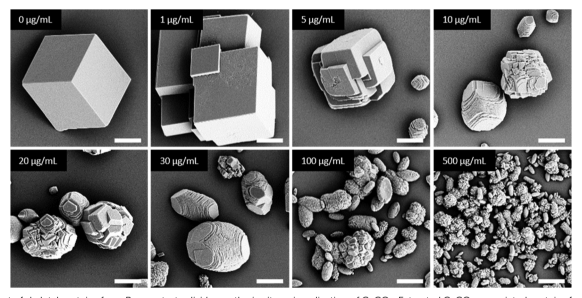


Fig. 2 Effect of skeletal proteins from Paracentrotus lividus on the in vitro mineralization of CaCO₃. Extracted CaCO₃-associated proteins from P. lividus skeletons were applied in the bioinspired mineralization of CaCO₃ via a slow diffusion method. The mineralization was performed in the presence of protein concentrations ranging from 1 μ g mL⁻¹ to 500 μ g mL⁻¹. In protein-mediated precipitation, the deposited CaCO₃ particles showed a complex morphology, whereas regular, rhombohedral particles were synthesized in the absence of proteins. Scale bars 10 µm.

scanning electron microscopy (SEM). Reference samples, without protein additives, showed rhombohedral calcite crystals with particle sizes between 50-100 μm. In the presence of sea urchin proteins, the mineralized CaCO₃ polymorph remained calcite (Fig. S1, ESI†). Moreover, the particle morphology drastically changes with the addition of the protein mixtures. Both protein mixtures had similar effects on the CaCO₃ morphology. The size of CaCO₃ particles was inversely related to the protein concentration, i.e. as the protein concentration increased the

particle sizes decreased, resulting in a particle size of around 5 μm in samples with the highest protein concentration (500 $\mu g \; mL^{-1}$). The reduction in particle size was accompanied by an increase in the number of particles. At low protein concentrations up to 5 µg mL⁻¹ the rhombohedral shape of the inorganic particles was still evident (Fig. 3A). Protein concentrations from 10 µg mL⁻¹ and higher induced the elongation in c-axis direction of the particles (Fig. 3B). The particles expressed a step structure on the surface, which was

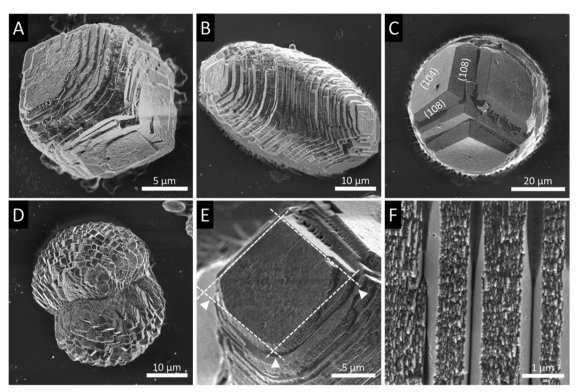


Fig. 3 Details of the surface topography of CaCO₃ particles obtained by in vitro mineralization in the presence of sea urchin protein mixtures. (A) The formation of pronounced steps blurs the sharp-edged rhombohedral particle morphology (probably initial stage). (B) Elongated, cigar-shaped particles are formed. (C) Top view of an elongated particle oriented along the principle axis. (D) Dumbbell-shaped particles with a pronounced constriction in the centre. (E) The (104) rhombic surfaces (white frame) usually have an intact obtuse angle, which is opposed by a corner that is not completely expressed or has been degraded. Likewise, the acute angles are not fully developed (arrowheads). (F) The particle surface appears to consist of co-aligned platelets with a nanogranular substructure.

again more pronounced at high protein concentrations. The morphological changes were most pronounced at the highest protein concentrations (100 and 500 µg mL⁻¹), where elliptical particles were formed, exhibiting a cigar-like shape (Fig. 3B). At the particles' tips three planes converged (Fig. 3C). In addition, dumbbell-shaped CaCO3 crystals were obtained, which exhibited a prominent separation line in the center (Fig. 3D). These dumbbell-shaped particles were composed of the same rhombohedral CaCO₃ crystallites. The stair-like morphology of all CaCO₃ crystals became more pronounced with increasing protein concentration. The morphology of the steps showed an intact edge at one of the acute angles of the rhombohedral (104) plane, while the opposite obtuse angle and the included acute angles showed a distinct degradation or an incomplete/suppressed growth characterized by rounded corners (Fig. 3E). The adjacent (108) side planes of the steps below were apparently composed of layered CaCO₃ platelets (Fig. 3F). In general, the (104) planes had a smoother surface appearance, but also showed to a lesser extent pores and irregular growth lines. The irregular mineral growth might be attributed to nonuniform crystal growth rates,⁶⁷ probably induced by the attachment of proteins or the etching of calcite from dissolution processes in aqueous solutions.68

(ii) Aragonite- and calcite-binding peptides. Next, the effect of aragonite- and calcite-binding peptides on the mineralization of CaCO₃ particles was analyzed (Fig. 4, 5, Table 2 and Fig. S3, S4, ESI†). Five aragonite-binding peptides and four calcite-binding peptides were selected. These peptides cover a broad range of isoelectric points, from 4 to 12, thus providing peptides with different net charges for mineralization.

The conformation of the most effective peptides (A-4-17, A-4-09, A-5-13, and C-4-19; Table 2) was assessed using different secondary structure prediction servers (see Fig. S2, ESI†). The modern machine learning-based techniques NetSurfP 3.0⁵⁰ and PSSpred,⁵¹ applied here, as well as *de novo* predictions, offer significantly high accuracy and showed that all investigated peptides predominantly adopt an unordered conformation.

In addition, to the afore mentioned sea urchin protein concentrations, the peptides were applied at a concentration of 2 mg mL⁻¹. All peptides had at least one charged amino acid side chain (Table 2). The aragonite-binding peptides applied in the mineralization experiments exhibit a higher proportion of charged amino acids compared to the calcite-binding peptides (Table 2), thus facilitating electrostatic interactions. In contrast, calcite-binding peptides contained a higher proportion of amino acids with hydroxyl groups (i.e. serine, threonine, tyrosine) that may promote the formation of hydrogen bonds.

In general, in the in vitro precipitation system, the calcitebinding peptides showed a minor impact on particle morphology compared to the aragonite-binding peptides. The aragonite-

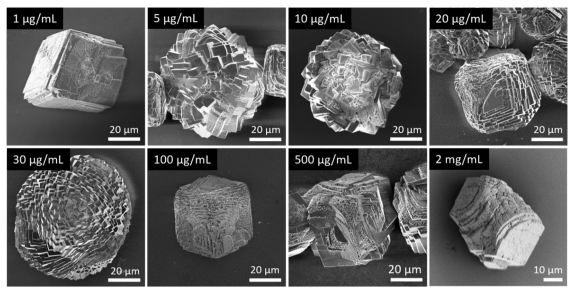


Fig. 4 Effect of peptide A-04-09 on the in vitro mineralization of CaCO₃. The aragonite-binding peptide was applied in the bioinspired mineralization of CaCO₃ via a slow diffusion of NH₃ and CO₂ induced by the decomposition of ammonium carbonate into a CaCl₂ solution. The mineralization was performed in the presence of protein concentrations ranging from 1 μ g mL⁻¹ to 2 mg mL⁻¹.

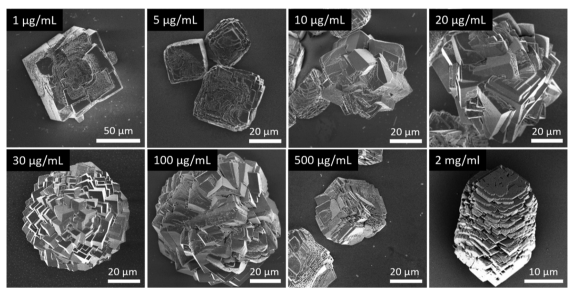


Fig. 5 Effect of peptide A-04-17 on the in vitro mineralization of CaCO₃. The aragonite-binding peptide was applied in the bioinspired mineralization of CaCO₃ via a slow diffusion of NH₃ and CO₂ induced by the decomposition of ammonium carbonate into a CaCl₂ solution. The mineralization was performed in the presence of protein concentrations ranging from 1 μg mL⁻¹ to 2 mg mL⁻¹.

binding peptides A-04-09 (Fig. 4) and A-04-17 (Fig. 5) demonstrated the largest effect on particle morphology at almost all concentrations tested. Furthermore, four peptides showed a strong effect on the morphology of the CaCO₃ particles at a peptide concentration of 2 mg mL⁻¹ (Fig. 6 and Table 2). Among these most potent structure-directing additives, were three aragonite-binding peptides (A-4-09, A-5-13, and A-4-17) and one calcite-binding peptide (C-4-19). In the presence of these peptides, calcium carbonate particles with shapes derived from a rhombohedron with one elongated axis were formed, similar to the crystal morphologies obtained in in vitro mineralization experiments with sea urchin proteins. At low peptide concentrations (5 µg mL⁻¹ to 500 µg mL^{-1}) the two aragonite-binding peptides A-4-09 and A-4-17 still had a visible effect on the particle morphology (Fig. 4 and 5). Furthermore, the c-axis elongation was reduced at low peptide concentration compared to particles synthesized at high peptide concentrations, thus almost globular particles with a staircasemorphology were formed. The greatest influence on the particle morphology was achieved with peptides with a large number of charged amino acids and hydroxyl-functionalized amino acids

Table 2 Biochemical parameters and mineralization effect of aragonite- and calcite-binding peptides applied in biomineralization experiments. The inorganic-binding peptides are sorted according to a mineralization factor. The mineralization factor was computed using the count of charged amino acids and hydroxyl groups in the amino acid residues. Each charged amino acid contributed 1 point, while each hydroxyl group contributed 0.5 points to the peptide's score. The cumulative points represented the mineralization factor. Peptide hydrophilicity was determined using the Vector NTI Software (Thermo Fisher). The effect of peptides on mineralization was assessed visually by SEM. Colour code of the amino acid (aa) residues, red: basic aa, blue: acidic aa, yellow: aa with hydroxyl group

Peptide	AA sequence / functionality	Charges (-/+)	OH- groups	Mineralization factor	Hydro- philicity	pl	Effect
A-4-17	HKHNQRPR <mark>S</mark> QKQ	0/6	1	6.5	2.000	12.02	+++
A-4-09	<mark>DH</mark> QV <mark>YSE</mark> N <mark>K</mark> P <mark>T</mark> I	2/2	3	5.5	2.067	5.32	+++
A-5-13	DSNVKNHNNKPE	2/3	1	5.5	2.000	6.75	+++
A-4-10	K <mark>GTS</mark> PQV <mark>TST</mark> HN	0/2	5	4.5	0.667	9.06	+
A-4-11	LP <mark>K</mark> W <mark>K</mark> AP <mark>H</mark> PFF <mark>S</mark>	0/3	1	3.5	0.867	10.00	+
C-4-19	<mark>S</mark> PL <mark>TRH</mark> VA <mark>D</mark> FQ <mark>T</mark>	1/2	3	4.5	0.700	6.46	+++
C-5-02	G <mark>S</mark> LV <mark>YRS</mark> G <mark>SH</mark> AP	0/2	4	4.0	1.100	8.75	-
C-5-03	<mark>S</mark> V <mark>S</mark> VGM <mark>K</mark> PSPRP	0/2	3	3.5	1.100	11.00	-
C-5-13	Q <mark>YT</mark> F <mark>E</mark> NPQ <mark>T</mark> NVA	1/0	3	2.5	1.067	4.00	-

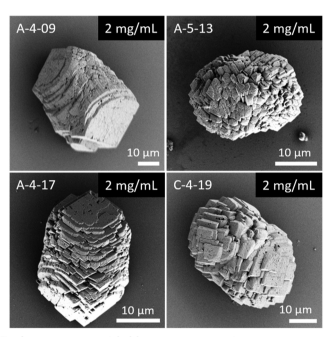


Fig. 6 Mineralization of $CaCO_3$ in the presence of peptides selected by phage display. Four of the investigated peptides (A-4-09, A-5-13, A-4-17, and C-4-19) showed a significant impact on the particle morphology similar to the effects of sea urchin proteins (compare Fig. 3). Selected micrographs are recorded from reaction products obtained with 2 mg mL $^{-1}$ peptide.

(Fig. 6 and Table 2). The mineralization factor was calculated using the count of charged amino acids and hydroxyl groups in the amino acid residues. Each charged amino acid contributed 1 point, while each hydroxyl group contributed 0.5 points to the peptide's score. The cumulative points represented the mineralization factor, a parameter representing an estimate for the capacity of the peptides to interact with ionic species or mineral facets, thus reflecting the effectiveness of the peptides on *in vitro* mineralization. The higher the value, the stronger the impact on particle

morphology. Only peptide C-4-19 deviated from this finding and showed a great influence on the morphology even at a lower mineralization factor. A correlation of pI and changes in morphology was not found, as reported for other mineralization proteins. ⁶⁹ As seen at the mineralization with sea urchin proteins, the surfaces of the particles sometimes showed voids and gaps that might be attributed to similar organic–inorganic interaction mechanisms. The peptides, however, usually required a higher concentration than the natural proteins to be effective. This difference might be in part due to the fact that the peptides were selected solely on the basis of their binding capacity to an inorganic substrate, while other potentially important interactions, *e.g.*, with a liquid mineral precursor or cooperative effects between different proteins, cannot be considered due to the experimental design.

The polymorphs of the crystalline CaCO₃ precipitates were determined by XRD measurements (Fig. 7). Samples with low (10 $\mu g \text{ mL}^{-1}$) and high (500 $\mu g \text{ mL}^{-1}$) peptide concentrations in the mineralization solutions were examined. The XRD diffraction patterns of all the *in vitro* CaCO₃ mineralization products showed calcite as the only polymorph regardless of the nature and concentration of the peptide additives. Although the peptides A-4-09 and A-4-17 were selected for their binding to aragonite, the polymorph of the synthesized particles was calcite. The observation that polymorph selectivity was independent of the concentration of the peptide, i.e. 10 $\mu g \text{ mL}^{-1} \nu s$. 500 μg mL⁻¹, clearly demonstrates that the peptides had a big influence on the morphology but not on the crystallographic modification of the synthesized CaCO₃. Since calcite was also precipitated in samples without peptide additives, we conclude that the polymorphism in this experimental setup is mainly driven by thermodynamic boundary conditions and not affected by the interaction with the peptides and proteins.

(iii) Nanostructure of calcite crystals deposited in the presence of an aragonite-binding peptide. (Bio-)macromolecules may become occluded into both, biominerals^{70,71} and synthetic crystals⁵⁹ resulting in an oriented intracrystalline nanostructure, either by selective interaction with the facets

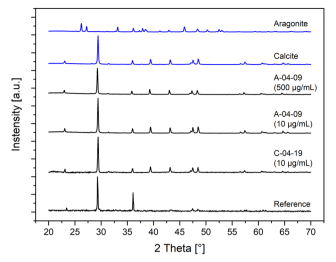


Fig. 7 Polymorphism of in vitro mineralized and geological CaCO₃. The polymorph of CaCO₃ mineralized in the presence of aragonite (A)- and calcite (C)-binding peptides was determined by capillary powder X-ray diffraction (XRD). For comparison, the diffractogram of a reference sample of calcium carbonate, which was mineralized without peptide additives, is shown. The comparison with diffractograms of geological (blue lines) aragonite (Alpha Aesar, Product no. 42523, LOT N04A029) and calcite (Alpha Aesar, Product no. 44520, LOT N04A030) indicated that calcite was deposited, both in the absence and presence of peptides independent of the original binding target of the peptides.

of crystalline building blocks or by incorporation into an amorphous precursor followed by facet-selective exclusion from the particle upon crystallization. Therefore, we analyzed the nanostructural features of the CaCO3 sample, which showed the most drastic morphological changes upon addition of an inorganic-binding peptide, by small-angle X-ray scattering (SAXS). Specifically, volume-averaging SAXS measurements were performed on in vitro mineralized CaCO3 powders deposited in the presence of peptide A-4-09 (concentration of 500 μg mL⁻¹) and compared to a reference sample, which had been prepared under the same reaction conditions but without a peptide additive (Fig. 8). Importantly, the XRD analysis revealed that both the sample and the reference are composed of phase-pure calcite (Fig. 7), thus excluding potential effects of polymorphism on the nanostructural motifs and facilitating the comparability between the peptide-mediated and the additive-free crystals. In addition, the overall particle morphology was homogenous throughout the entire sample, as exemplarily shown for the mineralization with peptide A-04-09 (SI Fig. 5).

In SAXS analysis, the control calcite sample exhibited a feature-less scattering profile following a Porod-like ($I(q) \propto q^{-4}$) behavior 72 over the entire investigated q range, which corresponds to larger crystals with smooth micron-sized external facets and provides no indication of an internal nanostructure (Fig. 8, grey symbols). In the presence of the peptide, by contrast, the scattering curve of the mineralized structures showed a more complex profile suggesting a pronounced substructure at the nanoscale presumably originating from internal porosity/organic occlusions within the crystals (Fig. 8,

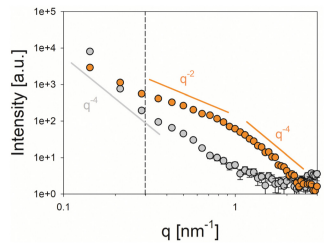


Fig. 8 Orientation-averaged plots of the small-angle X-ray scattering (SAXS) intensity versus the modulus of the scattering vector q (log-log representation) recorded for powdered samples of control calcite (gray symbols, no additives) and calcite crystals prepared in the presence of peptide A-04-09 ([peptide] = $500 \mu g mL^{-1}$, orange symbols). While the control sample shows an intensity decay of roughly $I(q) \propto q^{-4}$ according to Porod's law over the entire q-range covered by the experiment, the scattering profile of the peptide-mediated crystals provides indications for structural complexity at the nanometer scale. The dashed line indicates the cutoff $q = 0.3 \text{ nm}^{-1}$ between the low q regime, where the scattering profile of the peptide-mediated crystals is dominated by their external facets and the intermediate q-range, where a curve shape roughly following an exponential decay of $I(q) \propto q^{-2}$ is observed. The latter indicates platelike nanostructural motifs (presumably in the form of organic occlusions).

orange symbols). Even though the signal in the low q-regime (q < 0.3 nm, vertical dashed line) appeared to be dominated by the external facets of the micron-scale powder grains, a bent curve shape which can be described by a $I(q) \propto q^{-2}$ power law characteristic of oblate scattering objects is seen at larger values of q. T-Parameter analysis revealed a mean characteristic size of T = 2.28 nm of the nanostructural motifs, which can be associated with a thickness of D = 1.14 nm in the case of dilute (non-interacting) platelet-shaped scattering objects (Fig. S6, Porod and Kratky plots for the determination of the Porod constant P and the integral intensity \tilde{I} , respectively, ESI†).

The nanosized granularity of the peptide-mediated crystals may reflect on a formation mechanism that proceeds via the assembly of amorphous calcium carbonate (ACC) particles, which only subsequently crystallize. The peptide additives can interact with the amorphous CaCO3 and become occluded during crystallization. Such a process has been demonstrated for the occlusion of anionic polymers into calcite along so-called transition bars during the crystallization of CaCO₃ from ACC.⁷³ Therefore, we assume that the peptides are incorporated in the CaCO₃ crystals. This conclusion is further supported by the finding that such nanoscale structures are not detectable in the reference sample.

Discussion

Biomineralization proteins control the formation of inorganic materials at very low concentrations. In the natural system, the

total organic content in tests of the sea urchin A. lixula is approximately 0.12% w/w, 25 and the sea urchin proteins have been reported to intercalate into calcite crystals.⁷⁰ A morphological effect at low protein doses was also shown in the in vitro mineralization of CaCO₃ particles presented here. Interestingly, not only the proteins from sea urchin tests affected the morphology but, in addition, CaCO3-binding peptides selected by phage display. The effect of proteins on the mineralization is connected to their physicochemical properties, in particular to their net charge and hydrophilicity.⁷⁴ Early mineralization processes in sea urchin larvae were reported to occur at neutral to light alkaline conditions, 75 similar to the pH value of phage display selection. Therefore, no fundamental differences in the amino acid composition of the interacting proteins and peptides are expected as a function of pH. Differences between natural mineralization proteins and phage-display selected inorganic-binding peptides may account on other reasons. The lattice constants of geological calcite and the sea urchin test calcite are slightly different, 76 which is due to integrated organic molecules and, more importantly, to the incorporation of Mg²⁺ ions into the biogenic CaCO₃. These lattice differences may also contribute to variations in binding sequences. The strong influence of sea urchin matrix proteins on in vitro mineralization could be due to the glycosylation of these proteins. Up to 1.8% of the matrix proteins are glycosylated, ²⁵ and glycosylation may enhance the effect on mineralization, e.g. by increasing the hydrophilicity of proteins. The majority of sea urchin biomineralization proteins identified so far are acidic. However, basic mineralization proteins are also found in sea urchins associated with tests and spicules. 16,25,29,77,78 A prominent example is the spicule matrix protein SM50 from the purple sea urchin Strongylocentrotuts purpuratus⁷⁹ that stabilizes amorphous calcium carbonate.⁷¹ Interestingly, in vitro, the basic glycine rich region (GRR) domain (pI 10.48) of SM50 protein interferes with calcite formation by stabilization of prenucleation clusters,⁷¹ thus controlling the mineralization process. The exact mechanisms of crystallization are still under debate with two different theories are discussed, i.e. the classical theory of ionby-ion crystallization and the non-classical crystallization via an amorphous precursor.80 Accordingly, the organic-inorganic interaction in biomineralization was reported to occur either with an amorphous precursor or with crystalline particles. The nonclassical crystallization mechanism via an ACC precursor is supported by some findings. In natural sea urchin biomineralization, the formation of skeletons progresses through a transient phase of ACC. 81,82 In vitro, it was shown that protein domains interact with ACC and stabilize these structures.71 Inspired from such basic biomineralization proteins, the positively charged polymer poly(allylamine hydrochloride) (PAH) was shown to induce a liquid-like mineral precursor phase in the early stages of polymer-mediated in vitro crystallization of calcite, resulting in distinct particle morphologies.³⁸ However, it has also been shown that negatively charged polymers, which are considered standard as structure-directing additives in mineralization processes, can sequester and interact with ions in order to initiate mineralization. Polystyrene sulfonate (PSS) was shown to complex Ca²⁺ to form the primary species for crystallization in solution.⁸³ In addition, other negatively charged polymers like polyacrylic acid and polyaspartic acid were shown to interact with precursor species and step edges of CaCO₃ crystals.^{74,84,85}

In the designed mineralization peptides, one of the prerequisites for effectively influencing the mineralization process was found to be the presence of charges within the peptide. In addition, polar hydroxyl groups also play a minor role in this context. The peptides are identified as predominantly adopting unordered conformations. The conformational instability was reported as a prevalent feature of bio-molecules involved in the recognition and binding to solid materials. ^{86–89} Moreover, the inherent binding affinity of peptides is determined by the amino acid sequence and the secondary structure. It is assumed that conformationally labile peptides enhance their adaptability to interfacial features on surfaces, leading to the creation of accessible side chain regimes. ⁸⁶

The experimental approach for bioinspired calcium carbonate mineralization is significantly simplified compared to the biological situation. Many conditions cannot be exactly replicated in the experiment, e.g. due to the incomplete identification of all involved components, and the inability to reproduce the temporal and spatial distribution of precursor molecules. The proteins extracted from sea urchin tests strongly suggest a function of this organic matrix in the mineralization process. This is particularly evident in the impact of these proteins on the morphology of the in vitro mineralization product as compared to pure calcite. The structure-directing effects of CaCO₃-binding peptides can be inferred in the context of the formation of structural motifs reminiscent of those observed in the presence of sea urchin proteins. However, since the peptide selection is based on the binding affinity of peptides to calcite or aragonite, the peptide selectivity has to be carefully interpreted. Consequently, a distinction should be made between the selectivity regarding peptide binding to target substrates and the effects of the additives on structural parameters e.g. the morphology in the bioinspired mineralization of CaCO3. In the experimental set up, calcite was formed as the only polymorph, which does not necessarily hint towards a poor peptide selectivity but may rather be explained by the reaction conditions in the artificial model system, e.g. the thermodynamic boundary conditions.

Biomolecules and also high concentrations of ions (Mg²⁺) can decelerate biomineralization in the beginning by inhibiting the transformation of ACC into the crystalline phase.^{71,90} This temporal delay may allow control over the mineralization process. Furthermore, it was suggested that the conformation of peptides may play a role in mineralization⁷¹ and that structurally disordered proteins may match unstructured ACC precursors.^{91,92} In the progress of biomineralization, the ACC is subsequently transformed into calcite, ⁴⁵ aragonite and vaterite, ⁸⁰ respectively. Contrasting the interaction of proteins and peptides with ACC, it has been reported that *in vitro* rhombohedral calcite crystals are formed first and grow to a size of about 100 to 500 nm and only then the interaction with bioorganic polymers changes their morphology.⁴⁵ In the further

course of in vitro mineralization, positively and negatively charged additives bind to different sites of the nascent rhombohedral calcite crystals with distinct effects on particle morphology. Rhombohedral calcite exposing (104) faces have acute and obtuse step edges that are possible sites of interaction. Mg²⁺ ions interact with growing calcite crystals preferentially at acute step edges, 93 leading to the elongation along the c-axis. 45 Negatively charged polymers like PSS interact with the obtuse step of calcite and with the (001) calcite surface which leads to the formation of truncated edges and the formation of polyhedral particles. 45,83 Accordingly, negative and positive protein domains could simultaneously interact with calcite crystals, each in their own mode of action.

The morphological changes of calcite particles are correlated with the concentration of peptides, as shown for the most effective peptides A-04-09 and A-04-17 (Fig. 4 and 5). Such a correlation has also been demonstrated for the polymer PSS. 45 Proteins as well as other soluble additives, e.g. block copolymers and polyelectrolytes, showed control over the polymorph as well as the morphology and the texture of the crystal.⁵⁹ The concentration of mineralization peptides also influences the kinetics of the mineralization processes, where low peptide concentrations promote and high concentrations repress growth rate. 74,84 The adsorption of proteins could accelerate the growth rate in biomineralization. 74,94,95 It was speculated, that the interaction reduces the diffusion barrier by disturbing the layer of water molecules adsorbed on the surface in order to lower the energy barrier.⁷⁴ Furthermore, a positive effect on the nucleation was also shown for synthetic macromolecules.83

Conclusion

Inorganic-binding peptides, which interact with calcite and aragonite, respectively, were selected by phage display. While most of the biomineralization proteins associated with sea urchin calcite up to now were found to be acidic in nature, many CaCO3-binding peptides identified in our study are enriched in basic amino acid residues. We showed that the peptides, obtained by phage display, were able to control the morphology of nascent CaCO₃ crystallites in in vitro mineralization experiments. Moreover, the influences over crystallization exerted by CaCO3-binding peptides and natural biomineralization proteins from sea urchins lead to similar particle morphologies. In this *in vitro* mineralization approach, CaCO₃-binding peptides with a significant number of positively charged amino acids effectively change particle morphologies towards more elongated shapes. However, the sea urchin proteins have a greater influence on mineralization, as they provoke pronounced morphological modulations even at low protein concentrations. Although some of the peptides were isolated on aragonite binding, the polymorph of the crystalline phase is not affected by the inorganic-binding peptides, as reported for some natural systems. Additional prerequisites present in the natural systems, e.g. compartmentalization, mineralization vesicles, and cooperative interaction of several proteins, might be necessary to control the polymorphism.

Conflicts of interest

There are no conflicts to declare.

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