Biomineralized hybrid nanoparticles for imaging and therapy of cancers

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Abstract: In this review, we describe the research trends of hybrid nanocarriers developed based on a biomimetic mineralization process, and their recent applications in imaging and therapy of cancers. Organic-inorganic hybrid nanostructures formed by diverse biomimetic mineralization approaches are briefly reviewed, and particularly, the biomedical applications of these hybrid nanocarriers for the diagnosis and therapy of cancers are discussed. Biomineralization is an important process in which living organisms produce biominerals, such as calcium phosphate (CaP), calcium carbonate (CaCO₃), and silica (SiO₂), to strengthen their tissues, as found in the formation of bone and teeth. Introducing the artificial biomimetic mineralization process to nanobiotechnology has inspired researchers to develop smart stimuli-responsive nanoparticles for multiple purposes, such as improved therapeutic activity and activatable imaging of cancers.

Keywords: Biominerals; mineralization; organic-inorganic hybrid nanoparticles; cancer; therapy; imaging; theranostics

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Introduction

Recent advances in the field of nanomaterials exhibiting excellent therapeutic activity have attracted significant attention for the extended applications of novel drug delivery systems (1-3). For a successful treatment, it is recognized that nanodelivery systems should satisfy a number of prerequisites (4). In particular, the research needs to improve the biosafety, bioavailability, and biostability of nanocarriers are one of the main issues for the successful delivery of drugs, genes, proteins, and imaging agents to targeted disease sites. To meet the above requirements, various smart nanocarriers such as liposomes, polymeric micelles, and amphiphilic polymers have been constructed with stimuli-responsive functions, including labile cross-links and core-blockers, in pH, light, or redox environments (5-7). Furthermore, to date, bioinspired mineralized hybrid materials have expanded the development of smart nanocarriers in biotechnology and nanomedicine for advanced biomedical applications (8,9).

Biomineralization is a naturally important biological process by which living organisms develop biominerals [calcium carbonate (CaCO₃), silica (SiO₂), calcium phosphate (CaP), etc.] to strengthen their structures, as found in the formation of bone, teeth, shell, and diatoms (10). A distinct property of biominerals, the aqueous solubility of CaCO₃ or CaP minerals largely depends on pH (11-14) (Figure 1). For instance, the minerals maintain their crystalline structure at physiological pH, whereas they start dissociating into nontoxic ionic species around or below weak acidic pH levels (11). It is also known that CaCO₃ mineral reacts with protons and decomposes in an acid environment, generating carbon dioxide (CO₂) gas (12). Thus, these unique characteristics of biominerals can offer researchers...
with interesting and promising strategy for the development of stimuli-responsive hybrid biomaterials. In this article, we address the understanding of biomineral formation and its mechanism in nature. Then, the biomedical applications of biomimetic mineralized hybrid nanocarriers are presented.

**Biominerals and mineral characteristics for nanomedicine**

It is known that living organisms produce a wide range of specially designed inorganic-organic hybrid structures through the mineralization process in nature. The biomineralization process is applied for a variety of purposes, such as photonics, mechanical support, and protection of the soft parts of the body (15). The formation process of the bone or shell is vital for building the skeleton of the body, which uses both inorganic (CaCO₃, and CaP) and organic (proteins, lipids, and carbohydrates) materials (16-18). CaCO₃ and CaP are used in combination with organic polymers, such as chitin and collagen, to give structural support to bones and shells (19,20). The hard parts of these minerals are living structures that undergo progressive demineralization and remodeling in response to environmental influences and biological environments. A representative example to explain the biomineralization is the structure of the abalone shell. The structures of the inner and outer shells of abalone are formed by CaCO₃, proteins, and natural organic materials that exist between the CaCO₃ gaps (Figure 2). The organic template consists of chitin layers between the alanine-rich proteins and the silk-like glycine-rich protein, and the outer surface of the template is coated by hydrophilic poly(aspartic acid) macromolecules. Crystallization of CaCO₃ mineral is formed by the binding of CO₃²⁻ ions to the surface of the anionic poly(aspartic acid) and subsequent binding of Ca²⁺ ions. In addition, as another example, natural bone is formed from apatite crystals (CaP) having a specific orientation by biomineralization at the surface of the collagen fibril, which consists of self-assembled collagen triple helices (21).

Recently, CaP- and CaCO₃-mineralized hybrid systems have attracted increasing interest in the field of nanotechnology and biomedical science for the development of novel nanocarriers system because introduction of bioinspired mineralization enables nanocarriers to maintain structural robustness under physiological pH (pH 7.4) and selectively dissolve under the acidic pH of the intracellular endo/lysosome or tumoral region. Therefore, from the knowledge of the natural hybrid structures and biomineral characteristics, we describe a novel concept of mineralized hybrid nanocarriers system for efficient drug delivery to and imaging of the targeted disease in next each section.

**Mineralized liposomal nanocarrier systems**

As a pioneering work, a biomimetic mineralization process for hybrid nanosized liposomes covered with CaP layers was firstly reported by Schmidt et al. (22,23), 2-dioleoyl-sn-glycero-3-phosphate (DOPA)-based liposomes were used as a template for CaP mineral deposition. CaP mineralization on the surface of the liposomes was initiated by electrostatic localization of calcium ions at anionic phosphate moieties of liposomal surfaces, and the subsequent addition of phosphoric acid induced the formation of the CaP mineral layer. As a result, liposomal nanoparticles uniformly coated with an outer layer of CaP were formed, and the thickness of the CaP mineral layer gradually increased with increasing

**Figure 1** pH-dependent solubility of biominerals. Solubility of calcium phosphate according to changes in pH. Reproduced with permission from (13).

**Figure 2** Hybrid structure of natural biominerals. Organic-inorganic hybrid structures of the abalone shell. Reproduced with permission from (20).
time of the CaP mineralization process.

In addition, Xu et al. reported the kinetics of pH-controlled drug release from hydroxyapatite (HA) \([\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2]\)-mineralized liposome (HACL) (24) (Figure 3). Although these authors used indomethacin (IMC) as a model drug for this mineralized liposomal delivery system, this work is a valuable guide for the development of diverse pH-responsive mineralized nanocarriers for therapy and imaging of cancers. The IMC-loaded liposomes were formed by 1,2-dimyristoyl-sn-glycero-3-phosphate (DMPA) and 1,2-dimyristoyl-sn-glycero-3-phosphocholine (DMPC), and \(\text{CaCl}_2\) and \(\text{KH}_2\text{PO}_4\) were induced for the HA mineralization layer. After mineralization, the resulting particles showed that the thickness of the inorganic mineral layer was dependent on the reaction time. The particle size increased from 163 to 585 nm, and the zeta potential was changed from −38 to −4 mV after HA mineralization. For the result of the IMC drug release behavior of the liposome, the nonmineralized liposome demonstrated fast drug release (70%) within 5 h. In contrast, the mineralized liposome showed sustained drug release of over 20 h. Because the dissolution of the inorganic HA minerals was also accelerated under acidic pH conditions, a faster drug release rate for HACL was observed at pH 4.0 than at pH 7.4. These results indicated that the mineralized layer of the liposome controlled the kinetics of drug release by adjusting the pH levels.

Mineralized polymer nanocarriers for drug delivery

In recent years, functional polymeric micelles assembled by amphiphilic block copolymers have led to the fabrication of smart nanomaterials that can enhance the delivery efficacy of therapeutic drugs (25,26). In particular, polymeric micelles have diverse advantages for cancer treatments owing to their ability to efficiently escape from rapid renal clearance and nonspecific uptake by the reticuloendothelial system (RES). These unique properties enable the passive targeting and enhanced permeability and retention (EPR)
effect for effective tumor treatments (27). However, most polymeric micelles suffer from low structural stability in the bloodstream (28). In addition, drug release is induced upon intravenous administration, resulting in drug leakage at unwanted sites. By increasing the stability of micelles and minimizing the loss of therapeutic drugs at unwanted sites, Lee et al. demonstrated that spatially selective mineralization of polymeric micelles produced a highly biocompatible and robust hybrid structure that can controllably release anticancer drugs or therapeutic agents (28,29) (Figure 4). In this work, to prepare polymeric micelles as a nanotemplate for the mineralization process, the triblock copolymer of poly(ethylene glycol)-b-poly(L-aspartic acid)-b-poly(L-phenylalanine) (PEG-b-PAsp-b-PPhe) was utilized (29). Polyamino acids and PEG were selected as polymer components owing to their low toxicity and biocompatibility (28). CaP mineral deposition was processed by selective electrostatic localization of calcium ions at the anionic shell domain of Asp moieties. The subsequent addition of Na₂HPO₄ as a phosphate anion source initiated the formation of double ionic layers on the surface on the PAsp middle shells, thereby increasing the

Figure 4 Mineralized self-assembled polymeric nanocarriers for cancer therapy. (A) Illustration of spatial mineralization of drug-loaded polymeric micelles and drug release facilitated under specific intracellular conditions; (B) in vivo NIRF images of tumors from MDA-MB-231 tumor-bearing mice after intravenous injection of Cy5.5-DOX-NPM and Cy5.5-DOX-CaP-PM; (C) ex vivo images of normal organs and tumors after 3 days postinjection of DOX-NPM and DOX-CaP-PM. Reproduced with permission from (28,29).
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 Ionic concentration above the critical saturation point for CaP mineral growth. Doxorubicin (DOX), an anticancer drug, was simply loaded into the hydrophobic PPhe domain of polymeric micelles using the dialysis method before CaP mineralization. The mean hydrodynamic diameter was 31.2 nm, and X-ray spectroscopy (EDS) indicated that the mineralized micelles contained calcium and phosphate elements, which belong to CaP minerals. The DOX release rate from nonmineralized polymeric micelles (DOX-NPM) did not notably exhibit differences in release patterns between extracellular (pH 7.4) and lysosomal environments (pH 4.5). On the other hand, mineralized polymeric micelles (DOX-CaP-PM) effectively inhibited drug release in the extracellular environment at 12 h by 70%.

The *in vivo* tumor accumulation of the DOX-CaP-PM labeled with a near-infrared fluorescent (NIRF) moiety, Cy5.5, was observed in MDA-MB231 tumor-bearing mice. The result of optical imaging indicated that the DOX-CaP-PM showed enhanced tumor accumulation owing to prolonged systemic circulation in the bloodstream and an enhanced permeation and retention (EPR) effect compared with the DOX-NPM (29). Moreover, the DOX-CaP-PM exhibited an enhanced therapeutic effect in tumor-bearing mice compared with both free DOX and DOX-NPM.

For efficient photodynamic therapy (PDT), a similar mineralization approach was applied to the photosensitizer (PS)-loaded polymeric micelles (Figure 5) (30). Chlorin e6 (Ce6) was loaded into the polymeric micelles of PEG-PAsp-PPhe, and then, CaP minerals were deposited on the middle shell of the polymeric micelles. The Ce6-loaded CaP-mineralized micelles (Ce6-NP-CaP) displayed an enhanced stability of their structure and pH-dependent release patterns of Ce6. The MCF-7 cells treated with Ce6-NP-CaP with near-infrared (NIR) laser irradiation exhibited a dramatic decrease in cell viability. Moreover, after the Ce6-NP-CaP administered to the MCF-7 tumor-bearing mice by intravenous injection, *in vivo* biodistribution was monitored. NIRF images indicated that the Ce6-NP-CaP showed tumor-targeting ability compared to both free Ce6 and nonmineralized Ce6-loaded polymeric micelles (Ce6-NP).

More recently, Nam *et al.* developed a novel polymer nanotemplate for CaP-mineralized nanocarriers composed of a poly(3,4-dihydroxy-l-phenylalanine) (PDOPA) core and a poly(ethylene glycol) (PEG) shell (Figure 6) (31). The catecholic amino acid DOPA originates from the adhesive foot protein (Mefp-5, *Mytilus edulis* foot protein-5) of marine mussels (32). It has been reported that the catecholic OH of DOPA can bind to various metal ions (Ca, Mn, Fe, Zn, and Ti) to form a strong coordinate complexation (33,34). The strategy for successful CaP mineralization onto PEG-PDOPA polymer nanocarriers was based on the strong binding of Ca$^{2+}$ ions with OH of DOPA, which serves as a stable mineralization site of nucleation and growth for the CaP mineral core. DOX was loaded into the core of polymer nanocarriers (NPs) during self-assembly of PEG-PDOPA. In addition, CaP minerals were effectively induced

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**Figure 5** CaP-mineralized polymeric nanocarriers for photodynamic therapy. (A) Fabrication of CaP-Ce6-loaded polymeric micelles; (B) *ex vivo* images of organs and tumor tissues at 1 day postinjection of Ce6, Ce6-NP, and Ce6-NP-CaP. Reproduced with permission from (30).
on the PDOPA core of the PEG-PDOPA nanocarriers. The release of DOX from DOX-loaded CaP-mineralized nanocarriers (DOX-CaP-NPs) was triggered at endosomal pH (5.0), whereas the CaP mineral core stably maintained the structure of DOX-CaP-NPs, which resulted in the inhibition of drug leakage at extracellular pH. These DOX-CaP-NPs enhanced cellular uptake and cell cytotoxicity in SCC-7 head and neck cancer cells.

**Mineralized hybrid nanocarriers for protein delivery**

Despite their excellent therapeutic effect, target specificity, and biosafety, protein drugs have limitations in clinical applications because they are degraded by proteolytic enzymes and have difficulty in permeating into targeted cells. To overcome these limitations of protein drugs, Koo et al. introduced cytochrome c-loaded CaCO$_3$-mineralized nanocarriers (Cyt c MNPs) for effective intracellular protein drug delivery (35). They synthesized a block copolymer PEG-PAsp for a nanotemplate CaCO$_3$ mineralization approach. Cytochrome c (Cyt c), an apoptosis-inducing protein (36), was loaded in situ into the CaCO$_3$ mineral core during the nucleation and growth of CaCO$_3$ minerals via PEG-PAsp-template mineralization in the presence of Ca$^{2+}$ and CO$_3^{2-}$. The resulting mineralized nanoparticles had a spherical shape with a size of approximately 277.1 nm. The mineral core of the Cyt c MNPs sufficiently inhibited the release of Cyt c at physiological pH (7.4), but the dissolution of CaCO$_3$
triggered drug release at endosomal pH (5.0). With the capability of efficient cellular uptake and endosomal escape, this system successfully induced apoptosis and killed the MCF-7 cells.

**Mineralized hybrid nanocarriers for nitric oxide delivery**

Very recently, nitric oxide (NO)-generating CaCO$_3$ mineralized nanocarriers were developed to treat targeted cancer cells (Figure 7) (37). It has been known that NO is an endogenously produced free radical that plays multiple roles, such as neurotransmission, vasodilation, and regulation of proliferation and apoptosis of cells (38,39). In addition, NO has been reported to have an inhibitory effect on tumor growth by inducing apoptosis (40,41). However, despite the therapeutic potential of NO, the clinical applications of NO are limited by its instability and short

![Figure 7 Mineralized nanoparticles for intracellular nitric oxide delivery. (A) Illustration of the fabrication process for GSNO-loaded CaCO$_3$-mineralized nanoparticles (GSNO-MNPs) and intracellular NO release; (B) TEM images of MCF-7 cells treated with the GSNO-MNPs for 30 min (red arrows: individual GSNO-MNP, blue arrow: aggregates of GSNO-MNPs); (C) CLSM image of MCF-7 cells treated with GSNO-MNPs and DAF-FM DA. Reproduced with permission from (37).](image-url)
To increase the sensitivity and specificity of magnetic resonance imaging (MRI) probes for tumor imaging, Mi et al. designed CaP-mineralized nanoparticles into which diethyleneetriaminepentaacetic acid gadolinium (III) (Gd-DTPA) was incorporated (Figure 8) (45). Gd-DTPA-incorporated CaP nanoparticles (Gd-DTPA/CaP) were fabricated via a simple two-step method, which combined the PEG-PAsp block copolymer-templated CaP mineralization step with Gd-DTPA, followed by a hydrothermal process. In particular, the hydrothermal treatment endowed Gd-DTPA/CaP with enhanced structural stability in aqueous environments. In a molecular relaxivity ($r_1$) study, the $r_1$ value of Gd-DTPA from Gd-DTPA/CaP was 6 times higher than that of free Gd-DTPA. $T_1$-weighted MR images indicated that at the same concentration, Gd-DTPA/CaP showed much higher contrast enhancement than free Gd-DTPA. Furthermore, Gd-DTPA/CaP gradually increased the MR contrast of the tumor position after intravenous injection of Gd-DTPA/CaP to C-26 tumor xenograft mice, while the MR image of the tumor position from the free Gd-DTPA/CaP-treated group with 4 times higher dose than that of Gd-DTPA/CaP showed no signal enhancement. At 4 h postinjection, the tumor-to-muscle signal ratio of Gd-DTPA/CaP increased more than 40%.

Based on the results of these mineralized nanoparticles, the same group developed an improved mineralized hybrid MRI agent system. Mn$^{2+}$-doped CaP-mineralized nanoparticles (PEGMnCaP) were developed using poly(ethylene glycol)-b-poly(glutamic acid) copolymers [PEG-b-P(Glu)] with mineralization of Mn$^{2+}$, Ca$^{2+}$ and HPO$_4^{2-}$ and subsequent hydrothermal treatment (Figure 9) (46). The resulting particles showed that the release of Mn$^{2+}$ was pH dependent and accelerated in an acidic environment, resulting from the dissolution of the CaP mineral core. In $T_1$-weighted MRI, higher contrast enhancement of tumor regions was obtained for the PEGMnCaP-treated group compared with groups treated with both Gd-DTPA and PEGylated Mn$_3$O$_7$ nanoparticles (PEGMn$_3$O$_7$). The partially higher MR contrast regions than other regions in the tumors were correlated with both the hypoxic area and the high lactate level in tumor tissues. This is because the production of lactate and deficiency in environmental oxygen in the tumor tissue induced partial hypoxia, which resulted in an acidic microenvironment in tumors (47). Furthermore, PEGMn$_3$O$_7$ detected metastatic tumors in the liver induced by spreading C26 colon cancer cells compared to gadolinium ethoxybenzyl dimeglumine (Gd-EOB-DTPA), used for clinical hepatobiliary magnetic resonance probes.
Figure 8 Mineralized hybrid nanoparticles for enhanced MR imaging. (A) Preparation of Gd-DTPA/CaP; (B) in vivo MR images of C-26 xenograft tumor-bearing nude mice after intravenous injection of Gd-DTPA/CaP; (C) in vivo MR images of C-26 xenograft tumor-bearing nude mice after intravenous injection of free Gd-DTPA; (D) tumor-to-muscle signal ratio from MR images. Reproduced with permission from (45).
Mineralized nanoparticles for ultrasound imaging (US) and US imaging-guided therapy of cancers

Theranostics that integrate imaging and therapeutic functions in one system have emerged in the field of nanomedicine for developing novel nanoparticles (48,49). In recent years, CaCO$_3$-mineralized theranostics nanoparticles with spherical shape and sizes of approximately 240 nm have been developed for ultrasound (US) imaging and cancer therapy (Figure 10) (50). The block copolymer PEG-PAsp was applied as a CaCO$_3$ mineralization template. In this study, at physiological pH (pH 7.4), the DOX-loaded CaCO$_3$-mineralized nanoparticles (DOX-CaCO$_3$-MNPs) maintained their robust mineral structures, which inhibited the release of DOX owing to the solubility of CaCO$_3$ mineral. However, at tumoral acidic pH (pH 6.8), the DOX-CaCO$_3$-MNPs showed rapid DOX release patterns, which well correlated with the dissolution of Ca$^{2+}$ ions from the DOX-CaCO$_3$-MNPs. Moreover, US images from the DOX-CaCO$_3$-MNPs were significantly enhanced at tumoral acidic pH (pH 6.8, 7.0, and 7.2) because of sufficient CO$_2$ bubble generation from the DOX-CaCO$_3$-MNPs (50). In contrast, the DOX-CaCO$_3$-MNPs did not produce sufficient CO$_2$ gas bubbles, which resulted in no significant contrast in US signals at pH 7.4. For the in vivo US imaging experiment, after the intratumoral injection of DOX-CaCO$_3$-MNPs to the tumor tissue of SCC-7 tumor-bearing nude mice, the US contrast enhancement of the tumor site was detected up to 120 min. To confirm the in vivo antitumor effect of the DOX-CaCO$_3$-MNPs, DOX-CaCO$_3$-MNPs were injected intratumorally. The
Figure 10 Mineralized theranostic nanoparticles for ultrasound imaging and cancer therapy. (A) In vitro US images from DOX-CaCO$_3$-MNPs at various pH; (B) in vivo US images of the SCC-7 tumor via intratumoral injection of DOX-CaCO$_3$-MNPs. Reproduced with permission from (50).
DOX-CaCO₃-MNPs suppressed the tumor weight by 85% compared to the saline-treated group at 16 days after injection.

For successful tumor treatment by photodynamic therapy (PDT), it is of great importance to identify and detect the exact size and location of tumors before laser irradiation to the target site. Furthermore, during the therapy of cancers, real-time monitoring of the therapeutic response is needed (51). To achieve this purpose, US serves as an imaging guidance tool. Park et al. prepared a theranostics system of US imaging and PDT using mineralized hybrid nanocarriers for US image-guided photodynamic therapy (Figure 11) (52). Ce6-loaded bubble-generating mineralized nanoparticles (Ce6-BMNs) were fabricated by a polymeric nanotemplate-CaCO₃ mineralization process and provided potential for ultrasound imaging and US image-guided photodynamic therapy in response to a tumoral acidic pH. The Ce6-BMNs showed enhanced structural stability under serum condition. The Ce6-BMNs generated CO₂ gas bubbles at tumoral acidic pH, which resulted in excellent US

**Figure 11** Mineralized bubble-generating theranostic mineralized nanoparticles for US image-guided photodynamic therapy. (A) Fabrication of Ce6-loaded CaCO₃-mineralized nanoparticles (Ce6-BMNs); (B) generation of CO₂ bubbles with Ce6 release and subsequent PDT upon laser irradiation after tumor accumulation of Ce6-BMNs. Reproduced with permission from (52).
signals. After NIR laser irradiation of Ce6–BMNs-treated MCF-7 cells, cell viability dramatically decreased. As a result, the Ce6–BMNs exhibited much higher phototoxicity compared with that of the free Ce6-treated group.

**Conclusions**

We have introduced the recent research trends of mineralized hybrid nanocarriers system and the understanding of the process of biomimetic mineralization. This mineralization approach plays a vital role in the development of nanocarriers that can prevent the release of drugs and bioactive agents in the blood circulation and can release drugs at the targeted sites. Currently, the mineralized hybrid system is still at a challenging stage of development for biomedical applications, and it continues to have great potential over a wide range of applications and can overcome the disadvantages and limitations of the existence used nanocarriers system.

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**Footnote**

_Conflicts of Interest:_ The authors have no conflicts of interest to declare.

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