TOPICAL REVIEW

Hydroxylapatite nanoparticles: fabrication methods and medical applications

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Abstract

Hydroxyapatite (or hydroxyapatite, HAp) exhibits excellent biocompatibility with various kinds of cells and tissues, making it an ideal candidate for tissue engineering, orthopedic and dental applications. Nanosized materials offer improved performances compared with conventional materials due to their large surface-to-volume ratios. This review summarizes existing knowledge and recent progress in fabrication methods of nanosized (or nanostructured) HAp particles, as well as their recent applications in medical and dental fields. In section 1, we provide a brief overview of HAp and nanoparticles. In section 2, fabrication methods of HAp nanoparticles are described based on the particle formation mechanisms. Recent applications of HAp nanoparticles are summarized in section 3. The future perspectives in this active research area are given in section 4.

Keywords: hydroxyapatite, nanoparticle, composite, coating, medical device

1. Introduction

Apatite is a general term for the crystalline minerals that can be represented by the formula \( \text{M}_{10}(\text{ZO}_4)_6\text{X}_2 \). Each component (M, ZO_4 and X) in the formula can be replaced by a large number of different ions as listed in table 1. The most common apatite found in the nature is calcium phosphate apatite, where M and ZO_4 are Ca\(^{2+}\) and PO\(_4^{3-}\), respectively. When X is OH\(^-\) (i.e. Ca\(_{10}(\text{PO}_4)_6(\text{OH})_2\), stoichiometric Ca/P molar ratio 1.67), the mineral is called hydroxyapatite [1, 2] or hydroxyapatite (HAp), which is the main subject of this review. HAp is the inorganic component of the hard tissues of vertebrates. Biological apatite is a non-stoichiometric Ca\(^{2+}\)-deficient form of HAp containing trace ions. The trace ions include cations (such as Mg\(^{2+}\), Na\(^+\) and K\(^+\)) and anions (such as CO\(_3^{2-}\), Cl\(^-\) and F\(^-\)). The most common substituting ion is carbonate (CO\(_3^{2-}\)) that can replace OH\(^-\) and PO\(_4^{3-}\) ions via the so-called A-type and B-type replacements, respectively. Detailed chemical structures of HAp and other ion-substituted HAp can be found in recent reviews [3–9] and books [10–13].

Traditional HAp sintered ceramics (in dense, porous and powder forms) have been used in medical and dental fields [13–16], and the applications include alveolar ridge reconstruction and augmentation [17, 18], fillers for bone defects [19–22], middle ear implant [23], etc. HAp is bioactive (osteconductive), that is, HAp can encourage bone growth along its surface when placed in the vicinity of viable bone or differentiated bone-forming cells [11]. HAp composites have been developed to overcome the inferior mechanical properties (brittleness and low stiffness) of traditional HAp sintered ceramics (or to improve...
bioactivity of the composite matrix). The original concept of bioceramics/polymer composite was introduced by Bonfield et al. [24], and the idea was based on the concept that cortical bone itself comprises an organic matrix reinforced with a mineral component. The bone analogue HAp/polymer composite (commercialized under the trade name HAPEX) consists of HAp (~50 vol%) mixed in a polyethylene matrix, and provides the stiffness and bioactivity of HAp and the toughness of polyethylene.

Table 1. Major ions that can be part of apatite, M₁₀(ZO₄)₆X₂.

| Component | Ions |
|-----------|------|
| M         | Ca²⁺, Mg²⁺, Sr²⁺, Ba²⁺, Mn²⁺, Fe²⁺, Zn²⁺, Cd²⁺, Pb²⁺, H⁺, Na⁺, K⁺, Al³⁺, etc |
| ZO₄       | PO₄³⁻, AsO₄³⁻, VO₂⁺, SO₄²⁻, CO₃²⁻, SiO₃²⁻, etc |
| X         | OH⁻, F⁻, Cl⁻, Br⁻, O²⁻, CO₃²⁻, etc |

HAp nanoparticles can be obtained via a variety of methods including dry-synthesis (solid-state) and wet-synthesis methods (table 3). Other attempts at classification of HAp nanoparticle fabrication techniques can be found elsewhere [3–7, 53]. Dry synthesis usually yields stoichiometric and well-crystallized products, but they require relatively high temperatures (typically above 700 °C) and long treatment time. Besides, HAp is generally obtained in a bulk form by dry synthesis, and therefore grinding or milling is necessary to obtain nanoparticles. Wet-synthesis methods require relatively low temperatures and easily produce nanoparticles, but their crystallinity and Ca/P ratio are relatively low.

2. Fabrication methods for HAp nanoparticles

HAp nanoparticles can be obtained via a variety of methods including dry-synthesis (solid-state) and wet-synthesis methods (table 3). Recent applications of HAp nanoparticles are one of the topics of this paper. A major application of HAp is filling materials to impart bioactivity to various composite biomaterials, which can be found in recent reviews [47–52].

2.1. Dry synthesis

Stoichiometric and well-crystallized HAp can be prepared by solid-state reactions of calcium orthophosphates with calcium oxide or related salts such as Ca(OH)₂ (e.g. CaHPO₄ + CaO [54], Ca₃P₂O₇ + CaO [55, 56], β-Ca₃(PO₄)₂ + CaO [57, 58]) at a stoichiometric Ca/P mixing ratio for HAp formation at high temperatures (typically above 700 °C). Calcination of chlorapatite in a steam also leads to the formation of a stoichiometric HAp [59]. Non-stoichiometric HAp can be derived from natural materials such as bovine bones [60–62], fish bones [63], coral [64–66] and eggshell [67, 68] after thermal treatments. By controlling the parameters of the solid-state reactions, the morphological control of the products in the nanoscale can be achieved. For example, nanostructured biphasic calcium phosphate (HAp and β-Ca₃(PO₄)₂) particles in the form of plates with nanosized hollows have been prepared by a solid-state reaction [69].

These HAp materials are generally obtained in the bulk form, and therefore grinding or milling is necessary to obtain HAp as nanoparticles. Trakhtenberg et al. [70] recently reported the effects of grinding in planetary mills on the phase composition, morphology and water content of HAp powder.

Mechanochemical processing (or mechanical alloying) through direct ball milling for the above mixtures (that is, calcium orthophosphates and calcium oxide) is another way to fabricate HAp nanoparticles in a dry state [71–76]. Otsuka et al. [71] reported the effect of environmental conditions on the crystalline transformation of metastable calcium phosphates during grinding, and found that a mixture of CaHPO₄ · 2H₂O and Ca(OH)₂ transformed into HAp after grinding in air. The crystallinity of the products was low and hence post-synthesis annealing at temperatures between 500 and 1000 °C was required. Yeong and Wang [72] used CaHPO₄ and CaO as raw materials and fabricated well-crystallized HAp nanoparticles (particle size 25 nm,
2.2.1 Wet chemical precipitation. Among the wet-synthesis methods, the precipitation process (wet chemical precipitation) has been widely used [6, 46, 81] because of its simplicity. Mixing of two aqueous solutions of calcium carbonate (at pH > 7) results in the formation of highly supersaturated solutions for HAp, which induces a fast precipitation process of nanoparticles [81] according to the following reactions:

\[ 10\text{Ca(OH)}_2 + 6\text{H}_2\text{PO}_4 \rightarrow \text{Ca}_{10}\text{(PO}_4\text{)}_6\text{(OH)}_2 + 18\text{H}_2\text{O}, \]

\[ 10\text{Ca(NO}_3\text{)}_2 + 6(\text{NH}_4\text{)}_2\text{HPO}_4 + 8\text{NH}_4\text{OH} \rightarrow \text{Ca}_{10}\text{(PO}_4\text{)}_6\text{(OH)}_2 + 20\text{NH}_4\text{NO}_3 + 6\text{H}_2\text{O}. \]

Some researchers conducted this precipitation process in a simulated body fluid (SBF [82]) to obtain biomimetic apatite nanoparticles [83, 84].

The aqueous precipitation process yields calcium-deficient HAp (CDHA) via intermediate phases (precursors): amorphous calcium phosphate (ACP) is formed over a broad
range of precipitation conditions [85–95] and octacalcium phosphate (OCP) was formed in some cases [46, 85, 96, 97]. Alternatively CDHA may precipitate without going through an ACP precursor [98]. Recently, Mossaad et al. [99] reported the formation of 5-nm-sized HAp particles from a Ca(C$_2$H$_3$O$_2$)$_2$–K$_2$PO$_4$–H$_2$O system at room temperature: the obtained nanoparticles were very unstable and, interestingly, a crystalline to amorphous phase transformation was observed when the nanoparticles were aged for 5 months in 30% relative humidity.

The morphology (shape and size) of HAp nanoparticles depends on precipitation conditions such as concentration of reactants, ionic strength, pH and temperature [100]. The mixing condition for two (calcium and orthophosphate) aqueous solutions is also important. A microreactor is a device that has microchannels on the order of micrometers and that enables chemical reactions to be performed in a space several orders of magnitude smaller than conventional batch reactors [101]. The potential advantages of using a microreactor, rather than a conventional batch reactor, include high-speed mixing, better control of reaction conditions and improved yield. Yang et al. [102] prepared rod-like HAp nanoparticles (particle size 58 nm) using a tube-in-tube microchannel reactor. They reported that the particle size decreased sharply with increasing the continuous-phase flow rate, and a narrow size distribution was achieved at a high flow rate. Kandori et al. [103] recently prepared smaller HAp nanoparticles (2 nm in width, 15 nm in length) by using a multichannel microreactor.

The morphology of HAp nanoparticles also depends on additives, such as surfactant molecules [104, 105] (cf. micelle-templated method described in the next paragraph, and emulsion methods described in section 2.2.5), alcohol [106], amino acids [107–109], citrates [110, 111], poly(acrylic acid) [112, 113], poly(ethylene imine) [114, 115], poly(ethylene oxide) (poly(ethylene glycol)) [116, 117], poly(vinyl alcohol) [118, 119] and biopolymers [119], mainly due to the inhibition of crystal growth by preferentially adsorbing the additives onto a HAp surface (a or c plane).

Micelles form when the concentration of surfactant molecules is greater than the critical micelle concentration (CMC). The shape (sphere, cylinder or lamella) and size of a micelle can be controlled by the surfactant molecular geometry and solution conditions (surfactant concentration, temperature, pH and ionic strength). The addition of surfactant micelles as a template is beneficial for the fabrication of nanostructured HAp particles. Yao et al. [120] utilized surfactant micelles for preparing mesoporous HAp nanoparticles. They prepared HAp nanoparticles with channels approximately 3 nm in diameter by precipitation of HAp on rod-like micelles of cationic surfactants (cetyltrimethylammonium bromide, CTAB) in an aqueous medium. Wu and Bose [121] have also reported the preparation of HAp nanoparticles by the precipitation process in the presence of anionic dodecyl phosphate micelles. Ye et al. [122] reported micelle-templated synthesis of HAp hollow nanoparticles or nanotubes (outer diameter ~35 nm, inner diameter 13 nm, length 50–250 nm) with nonionic surfactants (EO$_{20}$PO$_{70}$EO$_{20}$; EO = ethylene oxide, PO = propylene oxide). Surfactant molecules can also stabilize oil droplets (or air bubbles) in aqueous solutions, and these droplets can be used for the fabrication of nanostructured HAp hollow or capsule particles having larger cavities as compared with the use of micelles alone. For example, He et al. [123] reported the formation of HAp nanocapsules (size 50 nm–1 μm) by precipitation of HAp on hexane droplets stabilized with bis(2-ethylhexyl)sulfosuccinate.

2.2.2. Hydrothermal process. Hydrothermal processes include several techniques for crystallizing materials in aqueous media at high temperatures (typically 100–250 °C, and hence under high vapor pressure when the temperature is above 100 °C). Microwave irradiation [124–127] can be used instead of conventional heating in pressure-resistant vessels. The hydrothermal process usually produces HAp particles with larger size (up to millimeter range), higher degree of crystallinity and a Ca/P ratio close to the stoichiometric value compared with those obtained by the precipitation processes at lower temperature (see section 2.2.1). However, the size distribution of the produced particles is usually broad. Hydrothermal processes can be further classified into two categories: (i) hydrothermal conversion and (ii) hydrothermal crystal growth techniques.

2.2.2.1. Hydrothermal conversion. The hydrothermal conversion technique involves the hydrolysis of other calcium phosphates (CaHPO$_4$, CaHPO$_4$·2H$_2$O, Ca$_8$(HPO$_4$)$_2$(PO$_4$)$_4$·5H$_2$O, Ca$_3$(PO$_4$)$_2$, fluorapatite and chlorapatite [128]) into HAp in aqueous media, usually with the aid of calcium, phosphate and/or alkaline sources to control the Ca/P ratio of HAp particles [129–135]. Hydrothermal conversions of poorly soluble calcium salts such as CaCO$_3$ (from coral [136–138], seashells [139, 140], aragonite or calcite crystals [141, 142]) and CaSO$_4$·2H$_2$O [124] into HAp have also been studied, and the conversion mechanism—ion exchange, dissolution and recrystallization—has been discussed. Although hydrothermal conversion techniques generally lead to the formation of large crystalline HAp particles due to the small number of nucleation sites, Shih et al. [132] prepared stoichiometric HAp nanoparticles (50 nm in width and 100 nm in length) by hydrolysis of a mixture of CaHPO$_4$·2H$_2$O and CaCO$_3$, performed with 2.5 M aqueous solution of NaOH at a relatively low temperature (75 °C). Rahaman and co-workers reported the fabrication of hollow HAp microspheres with nanosized porous shells (~13 nm) via a conversion of Li$_2$O–CaO–B$_2$O$_3$ glass microspheres in a K$_2$HPO$_4$ solution [143, 144].

As is the case in the wet chemical processes, the morphology of HAp particles can be varied by adding chemicals, such as carboxylic acids [145] and methanol [146], during the hydrothermal conversion.

2.2.2.2. Hydrothermal crystal growth. The hydrothermal crystal growth technique involves post-hydrothermal
treatment for as-prepared low-crystallinity HAp nanoparticles (prepared by wet chemical precipitation in many cases) in aqueous media. The dimensions of the precipitated HAp nanoparticles increase by the Ostwald ripening (maturation) under boiling or ambient aging in the mother liquid [147–150]. Pang and Bao [148] investigated the effects of temperature and ripening time on the crystallinity and morphology of the HAp nanoparticles, and found that the crystallinity and crystallite size increased with the increase in the treatment temperature and time. Wei and co-workers also reported the same tendency, but they observed that the length of the HAp decreased when the temperature was raised above 170 °C [26]. Pathi et al [151] recently reported a two-step hydrothermal method to obtain HAp nanoparticles with varying crystallinity and average length (32–103 nm). Although general hydrothermal treatment is a batch process, Darr and co-workers developed a continuous hydrothermal flow process [152–154] and prepared 8-g crystalline HAp nanoparticles (39 nm in width, 166 nm in length) within 1 h (yield > 95%).

Many investigations have been carried out to understand the effect of organic additives. Additives such as surfactants [155–158], amino acid [159, 160], dicarboxylic acid [161] and citrates [162] as well as pH [162] affect the morphologies of the particles treated. Wu et al [163] recently revealed that the chirality of the additives plays an important role in the asymmetric crystal growth of apatite by using L- or D-form of glutamic acid or aspartic acid.

Some researchers succeeded in preparing nanostructured HAp hollow particles without templates. For examples, Nathanael et al [164] reported a template-free formation of HAp nanorings with an inner diameter of 70 nm by a combined high gravity and hydrothermal crystal growth approach. Jiang et al [165] reported a template-free route to fabricate nanostructured hollow HAp microspheres assembled with nanorods (50 nm in thickness, 0.5–1 μm in length) by hydrothermal approach using poly(aspartic acid) as both a chelating and a capping agent. Ma and Zhu [166] reported the fabrication of hierarchically nanostructured HAp hollow spheres (assembled from nanorods) using solvothermal method at 200 °C in water/N,N-dimethylformamide (DMF) mixed solvents. They proposed a self-assembly/base-erosion mechanism for the formation of the nanostructured hollow spheres [167].

2.2.3. Homogeneous precipitation. The homogeneous precipitation starts with a homogeneous, acidic calcium phosphate solution, and nucleation/growth of HAp is induced by thermal decomposition (hydrolysis) of urea ((NH₄)₂CO + H₂O → CO₂ + 2NH₃) [168–171] or acetamide (CH₃CONH₂ + H₂O → CH₃COOH + NH₃) [172, 173]. The resultant NH₃ raises the solution pH (and degree of supersaturation for HAp), leading to the formation (precipitation) of HAp particles. Slow hydrolysis of the molecules at a high temperature leads to the formation of large and well-crystallized HAp particles. The hydrolysis of urea can be accelerated by the addition of enzyme (urease), especially at a low temperature of 37 °C [174]. Zhuang et al [175] recently synthesized plate-shaped HAp particles with a preferred orientation to the c plane by a homogeneous precipitation method via an enzyme reaction of urea. Yu et al fabricated willow-leaf-like nanorods (diameter 25 nm, length 120 nm) in a water/ethanol mixed solvent using urea as additive at 80 °C [176].

Hydrothermal treatment of the homogeneous calcium phosphate solution, in which calcium ions are dissolved by a chelating agent ethylenediaminetetraacetic acid (EDTA), also leads to the formation of HAp particles due to thermal dissociation of calcium-EDTA chelates in phosphate solutions [177, 178]. From the Ca–EDTA chelate solutions, HAp particles are obtained in the micrometer-size range after hydrothermal treatments, because EDTA ions do not inhibit HAp crystal growth [179]. López-Macipe et al [180] used citrate ions, which inhibit HAp crystal growth, as a calcium-chelating agent, and rod-like HAp nanoparticles (30–60 nm) were obtained after microwave heating. These calcium-chelated homogeneous precipitation methods utilize the promotion of Ca²⁺ dissociation by the calcium chelates (and solubility decrease of HAp) at an elevated temperature above 100 °C. Recently, He et al [181] and Cheng et al [182] fabricated nanostructured HAp porous microspheres by employing homogeneous precipitation methods with calcium-chelated phosphate solution in the presence of CO₂ bubbles as a hollow template.

2.2.4. Sol–gel method. The sol–gel method has been used for fabricating fine ceramics in wet conditions for a long time [183]. A typical precursor is metal alkoxide, which undergoes hydrolysis and polycondensation reactions to form a solid phase. In this process, the sol (i.e. solution) dissolving precursors evolves gradually towards the formation of a gel-like network of the solid phase. The solid phase can also be deposited on a substrate to form a film. By controlling the reaction parameters, the solid phase can also be obtained as nanoparticles dispersed in media [184].

The sol–gel method is also an effective way to fabricate nanostructured HAp sintered ceramics [185–187] and coating layers on several substrates [188–190] with a number of precursor combinations (calcium alkoxides and phosphorus alkoxides). Some researchers reported the effect of the starting precursor structure on the product (such as size, structure and crystallinity) [191, 192].

In the sol–gel method for HAp preparation, the as-obtained solid phase is generally amorphous Ca–P intermediates (and/or the mixture of unreacted precursors) and hence a thermal treatment (typically at 400–500 °C, which is lower than the sintering temperature of HAp powders, ∼800–1000 °C) is necessary to obtain well-crystallized HAp. The products are obtained in a sintered polycrystalline form and hence grinding or milling is usually necessary to obtain HAp nanoparticles (see also section 2.1). Recently, Costa et al [193] prepared microspheres consisting of nanosized HAp wires (25–800 nm in thickness) by a combination of sol–gel and hydrothermal (conversion) processes.
2.2.5. Emulsion method (surfactant-based biphasic process). This process involves the crystal nucleation/growth in a restricted space [194–202], which is generally prepared with inverse (water-in-oil type) emulsion droplets, inverse microemulsion droplets or inverse micelles (cf precipitation of HAp 'on' the normal micelles formed in aqueous media, described in section 2.2.1). The morphology control of HAp nanoparticles can be achieved by restricting the crystal growth, and this surfactant-based process can also inhibit excessive agglomeration of the particles. That is, this process has a lot of promise to synthesize nanocrystalline material without aggregation, but it has a drawback of contamination by the surfactants. For example, Lim et al [194] have prepared HAp powders by reacting CaCl₂ and (NH₄)₂HPO₄ in an inverse microemulsion formed with non-ionic C₁₅H₃₁O(CH₂O)₉·H (x = 5 or 9) in cyclohexane. They found that the microemulsion route led to a significant refinement in the particle size and the degree of particle agglomeration as compared with the particles obtained by a direct reaction in the absence of the emulsifiers and oil (that is, by a conventional wet chemical precipitation). The effect of reaction temperature (25–70°C) on the morphology of HAp nanoparticles formed in non-ionic C₁₅H₃₁O(CH₂O)₉·H-stabilized inverse emulsion droplets in dodecane media was investigated in terms of the cloud point (the temperature at which dissolved molecules are no longer completely soluble) of the non-ionic emulsifier [203, 204]. Sun et al [205] demonstrated the morphology control of HAp nanoparticles using inverse microemulsion systems (aqueous microemulsion in n-butanol/cyclohexane media stabilized with nonionic C₁₅H₃₁O(CH₂O)₉·H and cationic CTAB) under hydrothermal conditions at 160°C. Cationic CTAB micelles are also used to modulate the formation of HAp nanoparticles [206, 207]. Cao et al [208] reported the preparation of ultrahigh-aspect-ratio HAp nanofibers in inverse CTAB micelles under hydrothermal conditions. They showed that after the HAp nucleation in the inverse micelles, the surfactant headgroups preferentially adsorb on the surface planes parallel to the c-axis of HAp, resulting in the formation of anisometric nanofibers. Saha et al [209] compared the surfactant types (anionic dioctyl sulfosuccinate sodium salt (AOT) and dodecyl phosphate (DP); and nonionic C₁₅H₃₁O(CH₂O)₉·H (x = 5 or 12)) in terms of the morphology of the particles obtained in inverse microemulsions [209].

Cosurfactants are often used to increase the stability and the solubilizing capacity of microemulsions. An example of such a cosurfactant is a long-chain alcohol, which is also known to reduce the rigidity of water-in-oil interfaces [149]. García et al [210] studied the effect of cosurfactant (n-butanol) in a CTAB/toluene/water microemulsion system under hydrothermal conditions. They found that the cosurfactant to surfactant molar ratio also played an important role in controlling the morphology of HAp nanoparticles.

2.2.6. Calcined HAp nanoparticles. When low-crystallinity HAp nanoparticles prepared by precipitation processes (see section 2.2.1) or precursor nanoparticles prepared by sol–gel methods (see section 2.2.4) are calcined to improve their crystallinity (and hence their thermal and chemical stabilities), the nanoparticles typically sinter into large polycrystals (figure 1(a)) [211–215]. Consequently, calcined HAp crystals dispersed in a liquid on the nanoscale have been difficult to obtain. Hydrothermal crystal growth (see section 2.2.2) of HAp particles in aqueous medium is effective for preparing well-crystallized single crystals, but generally leads to an increase in crystal size and is restricted to laboratory-scale products as it is a high-pressure process.

A novel calcination method was developed to prepare well-crystallized HAp nanoparticles [216–218]. In this calcination method, an anti-sintering agent covering the nanoparticles is used to prevent the contact (and hence to prevent the calcination-induced sintering) between the nanoparticles; the agent is removed after calcination. Calcium salts (such as Ca(OH)₂ and Ca(NO₃)₂) were selected as the anti-sintering agent, and poly(acrylic acid) (PAA) was used to cover the HAp nanoparticles with the anti-sintering agent. The morphology of the calcined nanoparticles corresponds to that of as-prepared low-crystallinity nanoparticles (prepared by a wet chemical process) before calcination. Spherical or rod-like single HAp nanocrystals were fabricated by the novel calcination methods (figures 1(b) and (c)). Calcination with an anti-sintering agent has potential application to a wide range of calcined nanoceramic powders, such as alumina, titania and magnesia.

3. Applications of HAp Nanoparticles

HAp nanoparticles received considerable attention due to their large surface area, improved sinterability

![Figure 1. SEM images of HAp calcined (a) without additives and (b, c) with anti-sintering agents for (a, b) spherical and (c) rod-shaped nanoparticles.](image-url)
3.1. Carriers for drug, protein and gene delivery

HAp surfaces can bind various kinds of molecules (both acidic and alkaline proteins), because HAp belongs to a hexagonal crystal system and possesses different properties on its a and c planes [44]. HAp nanoparticles, therefore, have been investigated as a carrier for delivery of drugs such as growth factors [219, 220], antibiotics [5, 221–225] and anticancer drugs [5, 226, 227]. The adsorption/desorption characteristics [44, 220, 228–230] and conformation changes [230–235] of various kinds of proteins (or peptides) on HAp surfaces have been studied. Mukhopadhyay et al [236] recently reported that HAp nanoparticle supplementation increases thermal stability of pectate lyase from Bacillus megaterium, that is, this enzyme can retain high activity at elevated temperatures (up to 90 °C) in the presence of HAp nanoparticles. Recently, computational (in silico) studies have been gradually progressed to clarify the interaction between proteins (or peptides) and HAp surfaces [237–239]. Detailed information on the protein/peptide adsorptions can be found in the original papers or a recent review by Goobes et al [235].

Calcium phosphate (CaP)-deoxyribonucleic acid (DNA) co-precipitation method has been used for in vitro gene transfection because of the biocompatibility, biodegradability and ease of handling of CaP [240, 241]. To achieve effective gene transfection, the conditions for CaP-DNA co-precipitation method have been studied [242–244]. HAp nanoparticles can also be utilized as gene carriers because of their capability to absorb DNA molecules [245, 246].

Because of their small size, inhaled nanoparticles can have a high deposition rate in alveolar regions of the lung. Fan et al [247] investigated a time-dependent effect of spherical HAp nanoparticles (∼90 nm, prepared by a wet chemical process) on a pulmonary surfactant protein. They found that at a low concentration of 50 μg ml⁻¹ HAp nanoparticles inhibited the biophysical function of the surfactant protein due to adsorption of surfactant proteins onto the nanoparticles, whereas the nanoparticle concentrations up to 100 μg ml⁻¹ did not elicit any significant toxicological effects on human bronchial epithelial BEAS-2B cells in vitro. Fan et al claimed that the conventional cytotoxicological test alone may not be sufficient in evaluating the toxicological effect of inhaled nanoparticles, but also claimed that the interaction between HAp nanoparticles and pulmonary surfactant proteins shed light on the feasibility of HAp nanoparticle-based pulmonary drug delivery.

3.2. Reparative materials for damaged enamel

HAp nanoparticles have been investigated as fillers for dental composites [248–250] and glass-ionomer cements to improve their mechanical properties [251]. The application of HAp to repair the damaged enamel has recently attracted considerable attention in dental field because of the chemical and structural similarities of HAp to tooth minerals (20–40 nm particles of HAp [252, 253]). Li et al [254] reported a higher remineralization effect when using 20-nm-sized HAp compared with several hundred nanometer-sized HAp or 20-nm-sized ACP. Huang et al [255] demonstrated that HAp nanoparticles have a similar re-mineralization effect on an initial caries lesion to fluoride. Kim et al [256] examined the effect of carbonate HAp nanoparticles to prevent re-staining and the change of enamel surface after dental bleaching.

3.3. Particulate emulsifier (Pickering emulsion stabilizer)

General emulsions are stabilized by adsorption of surfactant molecules onto oil/water interfaces. Pickering emulsions are solid-particle-stabilized emulsions, in which solid particles are adsorbed onto oil/water interfaces [257, 258]. Inorganic particles (such as silica [259–263], clay [264, 265] and carbon black [266, 267]), organic particles (such as latex [268–270] and microgels [271, 272]) and Janus particles [273–276] have been used as particulate emulsifiers. These solid-stabilized emulsion droplets can be used as polymerization vessels; therefore, Pickering-type suspension polymerization [277, 278], inverse suspension polymerization [279, 280], and mini-emulsion polymerization [281–283] methods can be used to prepare nanocomposite particles.

HAp nanoparticles have been tested as a particulate emulsifier [284]. Stable oil-in-water type emulsions were readily obtained using oils containing an ester group (e.g. methyl myristate or methyl trimethyl acetate, see figure 2). Although no stable emulsion was obtained using oils without an ester group (e.g. dichloromethane), dichloromethane droplets could be stabilized by dissolving polymers having ester groups (or carboxyl terminal group) via the interaction between polymer and HAp nanoparticles at the oil/water interfaces [285, 286]. It was also demonstrated that emulsification–demulsification cycles could be repeatedly achieved by pH adjustment, because the HAp-adsorbed emulsion droplets became unstable at lower pH (< 4) and the emulsion stability was recovered by raising pH. The HAp-nanoparticle-stabilized emulsions are pH-responsive emulsions, and can be used for preparing porous HAp ceramics and HAp-coated microspheres (see section 3.4.2).

3.4. Coating agent

HAp composites are generally fabricated by mixing HAp particles with a matrix. However, the bioactivity of HAp is sometimes hindered because most of HAp is buried inside the composite matrix by the simple mixing methods. HAp coatings are employed to improve the bioactivity of the matrix (table 4).

HAp coating in the dry state (especially by plasma spraying) is a well-accepted and widely used technique, but it has some intrinsic drawbacks related to the extremely high processing temperature (above 10 000 °C): coating on polymer substrates and onto intricate shapes [363, 364], as well as incorporation of biological molecules (such as growth
The surface modification of the substrate is essential to determine the bonding strength of HAp [354]. In the nanocrystal coating method, the substrates were modified by graft polymerization of functional monomers having alkoxysilyl, isocyanate or carboxyl groups. The HAp surface possesses hydroxyl groups and calcium and phosphate ionic sites that can react with several functional groups. For example, alkoxysilyl groups (silane coupling agents) [210, 366] and isocyanate groups [367] can covalently react with surface hydroxyl groups of HAp, and carboxyl groups can interact ionically with calcium ions on the HAp surfaces. Polymer and metal substrates have been used [346–355]. After adsorption of HAp nanocrystals and reaction at the interface between HAp nanocrystals and substrate surfaces, the HAp-nanocrystal-coated substrates retained the mechanical properties of the substrates and showed improved cell adhesion properties owing to the presence of HAp crystals on the surface [349–353, 355]. The HAp nanocrystal coating is applied to percutaneous devices [349,356], artificial blood vessels [357, 358] and stents [352].

### 3.4.2. Nanoparticle coating via Pickering emulsion route.

Solid-particle-stabilized emulsions (see section 3.3) have been utilized for the preparation of HAp-coated biodegradable polymer microspheres by evaporation of solvent (oil) from HAp-nanoparticle-stabilized oil droplets dissolving polymer (figure 4). By using a water-in-oil-in-water multiple emulsion, multiholow microspheres can also be prepared in the same manner [359]. The HAp-nanoparticle-coated microspheres showed improved cell adhesion and spreading compared with bare biodegradable microspheres [285]. Recently, Mima et al [368] showed the effectiveness of HAp-coated biodegradable polymer microspheres as an injectable cell scaffold for cell-based therapeutic angiogenesis.

### 4. Summary and future perspectives

This paper reviewed the fabrication methods of HAp nanoparticles, highlighting recent improvements for morphology control. Numerous methods for fabricating HAp nanoparticles have evolved in the past few decades to control biological effects of nanostructured HAp. However, the development of facile fabrication methods for monodispersed HAp nanoparticles (with very narrow size distribution) is still
a challenging task. The monodispersity of nanoparticles is important to guarantee the homogeneity of the nanoparticles in a batch, and hence to define the biological effects of HAp nanoparticles. Monodispersed HAp nanoparticles are now fabricated in a limited size range \[102\] because of the fabrication strategy to eliminate the particle growth stage. The fabrication of monodispersed HAp nanoparticles in a broad size range will be realized by precise adjustment (separation) of the nucleation and growth stages \[369\] or by entirely different approaches \[370\]. Dispersion of HAp nanoparticles in liquid media (or in solid matrices) is also important for the applications of HAp nanoparticles and for the development of well-designed HAp-filled composites \[53\]. Although the dispersion is now improved by adding surface modifiers, another approach will be proposed because modifying pristine HAp surfaces generally hinders their bioactivity.

Recent applications of HAp nanoparticles are also summarized in this review. The greatest motivation behind the use of HAp is to accurately mirror the chemistry of natural minerals. The recent progress in computational analyses for the conformation changes, which alter structure and function at a distant active site, of the proteins interacting with HAp surfaces \[235, 237–239\] will provide a guide for designing nanostructured HAp surfaces to transcend the chemistry of natural minerals. Although nanostructured biomaterials have many potential advantages, it is important to remember that the effects of nanoparticle exposure on human health are not well understood. Understanding the biological effects of nanosized HAp is essential for the applications.

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