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Utilization of Compounds of Phosphorus

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Abstract

The last chapter of this book provides brief description of utilization of apatites and phosphorus-bearing compounds in industry and material science research. Since the chemistry of phosphorus is quite complicated and a quickly developing field of science, the topics described below are only limited insights to chemically bonded ceramics and refractories, dental phosphate cements, oil-well cements, phosphate glasses and glass ceramics. Chapter continues with description of functional phosphate materials applied as solid oxide fuel cells electrolytes, sensors, phosphors, catalysts and coatings. The chapter ends with introduction of basic ideas for biological apatite in bone tissue engineering, collagen apatite composites, apatite layers and biocoatings.

Keywords: Apatite, Chemically Bonded Ceramics, Phosphate Cement, SFOC Electrolytes, Sensors, Biological Apatite, Collagen, Biocoatings

In the previous chapter, the utilization of phosphate ore was described. Elemental phosphorus (Section 9.1) and phosphoric acid (Section 9.2) are used in the manufacturing of other phosphorus compounds (as illustrated in Fig. 1) and materials, which are briefly reported in this chapter. Since the chemistry of phosphorus is quite complicated and a quickly developing field of science, the topics described below are only limited insights to phosphorus-bearing compounds utilized in the industry and material science research.
10.1. Chemically bonded phosphate ceramics

Ceramics are formed by the compaction of powders and their subsequent fusion at high to very high temperatures, ranging somewhere from ∼700° to 2000°C. Once fused, the resulting ceramics are hard and dense and exhibit very good corrosion resistance [2]. Among the conventional ceramic bonds, the chemical (organic and inorganic) and hydraulic bonds were used during the preparation of ceramic materials.

Phosphate bond can be utilized for the preparation of hard and quick-setting ceramic materials, chemically bonded phosphate ceramics (PCBC) and ceramic composites [3],[4] via the reaction of metal cation with phosphate anions (Fig. 2). The reaction is attained by mixing a cation donor, generally an oxide (CaO [5],[6],[7],[8], MgO [2],[6],[9], ZnO [2],[8], Al₂O₃ [2], Fe₂O₃ [2],...
etc.), raw materials [10], or secondary raw materials (fly ash [11] and blast furnace slag [12]) with either phosphoric acid or acid phosphate such as ammonium phosphate solution or magnesium dihydrogen phosphate (Mg(H$_2$PO$_4$)$_2$·2H$_2$O) and aluminum hydrogen phosphate (AlH$_3$(PO$_4$)$_2$·H$_2$O). Typical acid-base reaction between a metal oxide and phosphoric acid can be written as [2],[13]:

$$\text{MO}_x + n \text{H}_3\text{PO}_4 + m \text{H}_2\text{O} \rightarrow \text{MH}_{3n-2x} (\text{PO}_4)_n + (m + x) \text{H}_2\text{O}$$  \hspace{1cm} (1)

where $x$ denotes half of valence of M, $n \geq 2x/3$, and $m$ is an arbitrary integer that decides the amount of water to be added in the reaction.

In the case of magnesium phosphate chemically bonded ceramics (MPCBC), the acido-basic reaction 1 of orthophosphoric acid with MgO yields the following products [2]:

$$\text{MgO} + 2 \text{H}_3\text{PO}_4 \rightarrow \text{Mg} (\text{H}_2\text{PO}_4)_2 \cdot \text{H}_2\text{O} \quad \Rightarrow \quad x = 1, \; n = 2, \; \text{and} \; m = 0$$  \hspace{1cm} (2)

$$\text{MgO} + \text{H}_3\text{PO}_4 + 2 \text{H}_2\text{O} \rightarrow \text{Mg} (\text{H}_2\text{PO}_4)_2 \cdot 3\text{H}_2\text{O} \quad \Rightarrow \quad x = 1, \; n = 1, \; \text{and} \; m = 2$$  \hspace{1cm} (3)

$$3 \text{MgO} + 2 \text{H}_3\text{PO}_4 \rightarrow \text{Mg}_3 (\text{PO}_4)_2 + 3 \text{H}_2\text{O} \quad \Rightarrow \quad x = 1, \; n = \frac{2}{3}, \; \text{and} \; m = 0$$  \hspace{1cm} (4)

If acidic phosphate is used, the reaction proceeds as follows [2]:

$$\text{MgO} + \text{Mg} (\text{H}_2\text{PO}_4)_2 \rightarrow 2 \text{MgHPO}_4 + \text{H}_2\text{O}$$  \hspace{1cm} (5)

and similar reaction can be also written for the reaction of Al(H$_3$PO$_4$)$_3$ with AlO$_{3/2}$:

$$2 \text{AlO}_{3/2} + \text{Al} (\text{H}_3\text{PO}_4) \rightarrow 3 \text{AlPO}_4 + 3 \text{H}_2\text{O}$$  \hspace{1cm} (6)

Reactions 5 and 6 indicate that partially acidic phosphate salt is only an intermediate phase. The dissociation of orthophosphoric acid and the pH stability range of formed ionic species are given by Eqs. 11–13 in Chapter 9. General dissociation reaction for acidic dihydrogen phosphate and its dissociation constant in the form of $pK_{\text{dis}}$ ($pK_{\text{dis}} = -\log K_{\text{dis}}$) can be written as follows:
\[ \text{M(H}_2\text{PO}_4\text{)}_n \leftrightarrow \text{M}^{n+} + n \text{(H}_2\text{PO}_4\text{)}^{n-} \quad (7) \]

and

\[ pK_{\text{dis}} = -\log \left( \left[ \text{M}^{n+} \right] \left[ \text{H}_2\text{PO}_4\text{)}^{n-} \right] \right) \quad (8) \]

The values of \( pK_{\text{dis}} \) for KH\(_2\)PO\(_4\), (NH\(_4\))H\(_2\)PO\(_4\), Mg(H\(_2\)PO\(_4\))\(_2\):2H\(_2\)O and Ca(H\(_2\)PO\(_4\))\(_2\):H\(_2\)O are 0.15, -0.69, 2.97 and 1.15, respectively [2].

The preparation scheme for the synthesis of aluminum phosphate binder is shown in Fig. 3. The evolution of chemical composition of phosphate binder with Al:P ratio of 1.4:3 with temperature is introduced by Table 1.

![Fig. 3. Schematic procedure for the synthesis of aluminum phosphate binder [14].](image)

Initially, the motivation for the development of these ceramic materials was the preparation of dental cements. Phosphate chemically bonded ceramics (PCBC) find recently the application in diverse fields (Fig. 2), which include structural ceramics, refractory materials [15], toxic, radioactive [16],[17],[18],[19] or asbestos-containing [20],[21],[22] waste management [23], oil drilling and bioceramics, pigments [1], etc. [2].

10.1.1. Ceramics and refractories

The broad term “ceramics” refers to any of a large family of materials that are usually inorganic and require high temperatures in their processing or manufacture. They are generally classified into glass, whitewares, including artware and structural ceramics, and refractories. In general,
The term “glass” refers to an amorphous solid with non-directional properties, characterized by its transparency, hardness, rigidity at ordinary temperatures and capacity for plastic working at elevated temperatures. Major commercial uses of glass include plate or “float” glass, as used for windows and windshields; glass tubing, or formed shapes, used for electric lighting envelopes; and glass containers such as tumblers and bottles. Whiteware is characterized by a crystalline matrix held together by a glassy phase and usually covered by a glazed coating. Major classifications of whiteware are ceramic tiles, glazed and unglazed, for floors, walls and external use; sanitaryware in the form of toilets and lavatories; and tableware, from earthenware to fine china [24].

Kaolin is major raw material used for the fabrication of conventional ceramics. It is obtained from the alteration of granitoid rocks [25]. It consists mostly of kaolinite and a small amount of impurities such as quartz, micas and other phyllosilicates. Firing of kaolinite induces numerous complex structural and microstructural transformations leading to the formation of various phases.

| Temperature [°C] | Dominant phase | Secondary phase |
|------------------|----------------|----------------|
| 60               | AlH₂(PO₄)₂·3H₂O | —             |
| 105              | AlH₂(PO₄)₂·3H₂O (monoclinic) | Al(H₃PO₄)₃ (hexagonal) |
| 200              | Al(H₃PO₄)₃ (hexagonal) | —             |
| 220              | AlPO₄ (trigonal) | AlH₂P₃O₁₀·2H₂O |
| 250              |                 |                |
| 300              |                 |                |
| 400              | Al(PO₃)₅       | AlH₂P₃O₁₀·2.5H₂O |
| 500              | Al₂P₆O₁₈       | Al(PO₃)₅ (cubic) |
| 600              |                 |                |
| 700              |                 |                |
| 800              | Al(PO₃)₅ (cubic) | AlPO₄ (rhombic system) |
| 900              |                 |                |
| 1000             |                 |                |

Table 1. The alteration of composition of phosphate binder with increasing temperature [14].

The alteration of composition of phosphate binder with increasing temperature [14].

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2 SEM picture of thin long fibrous structure of chrysotile (H₃Mg₃Si₂O₃ [20],[21],[22]) asbestos under the magnification of 98× (a), 5000× (b) and 10,000× (c).
of mullite and silica (cristobalite) phase. Mullite phase is characterized by some advantageous properties such as good corrosion resistance, low dilatation coefficient, good creep and thermal shock resistances, thermal stability and high strength. These advantages make this phase favorable for different applications. Orthophosphoric acid reacts with aluminum from kaolin to provide new compounds, which are Al(H$_2$PO$_4$)$_3$ at room temperature and AlPO$_4$ when heated to temperatures above 800°C [10].

Approximately 70% of all refractories used in industry are in the form of bricks, which are cast in the shapes such as straights, soaps, splits, arches, wedges, keys, skews, jambs or other special and frequently patented shapes. Most industrial refractories are composed of metal oxides or of carbon, graphite or silicon carbide. Most refractory bricks are shaped by combining the size-graded refractory aggregate with a small amount of moisture and casting in a dry press. The important properties of any refractory, including its high-temperature strength, depend on its mineral makeup, the particle-size distribution of minerals and the way these materials react at high temperatures and in furnace environments. When a refractory is chosen for a particular service, the service conditions must be considered in the design. In proper selection of a refractory, these factors, together with an economic balance, must be considered so that the refractory ultimately produces the lowest cost per unit weight of product per unit weight of refractory consumed [26].

![Fig. 4. Cutting plane throughout the block of chemically phosphate bonded refractory material containing large grains of calcined bauxite (a) and electron microscopy (SEM) picture of ceramic body (b).](image)

The materials must be chosen with generous safety margins in the temperature capability and the refractory construction system, whether brick, monolithic or fiber, should be suitable not only for the operating conditions but also for the type of equipment concerned and the construction conditions. Refractory concretes and castables (monolithic refractories) are especially suitable for these small burner quarls. Fibers and refractory ceramic fibers (glass, 3Granular crystalline rock consisting essentially from quartz, orthoclase-feldspar and mica. Usually is light gray, white or light in color [25].
mineral, ceramic fibers and whiskers) become an important construction as well as insula-
tion material, although the brickwork has the longest history of the development [27].

Aluminosilicate refractories are manufactured using refractory clays, sillimanite minerals,
bauxite (Fig. 4) and mixtures of alumina and silica sand. They will refer, somewhat arbitrarily,
to common crystalline compounds with melting temperatures of at least 1500°C. The major
categories of traditional refractories are fire clays, high alumina and silica. The choice of
material for traditional refractory applications, as well as for advanced material applications,
was and is based on balancing the cost and the performance lifetime [28].

Industries involved in steel melting and casting of special alloys are always interested in using
the maintenance-free, functional refractories to achieve energy-efficient metal processing.
They seek for the development of new and advanced thermal ceramics for protective ther-
mal insulation linings and molten metal handling crucibles. Conventionally, porous to dense
Al₂O₃, SiO₂, MgO, ZrO₂, SiC [29] and fireclay ceramic bodies were used as thermal insula-
tion refractory liners [30]. Phosphate bond can also be utilized in manufacturing of refracto-
ry and wear-resistant and of protective coatings on metal [14] or ceramic surfaces [31].

Intensive development of ceramic materials has increased the availability of well-character-
ized engineering ceramics capable of the utilization over the range of temperatures and
atmospheres [32].

Refractory castables can be classified according to different aspects including the content of
calcium, binder source, overall chemical composition, bulk density, application method and
others. Binders commonly used in monolithic products may be of various classes [33]:

1. Hydraulic (cement and hydratable alumina) [34]
2. Chemical (phosphoric acid, silicates, geopolymers, etc.) [33],[35]
3. Sol-gel (colloidal silica or alumina) [33]

Regarding the chemical ones, the bond strength can be provided by the addition of phos-
phates (dry or solution) or by in situ generation of phosphates (formed via the reaction with
added H₃PO₄) in the refractory structure. The reaction of Al₂O₃ or Al(OH)₃ with orthophos-
phoric acid can be described by reactions [33]:

\[6 \text{H}_3\text{PO}_4 + \text{Al}_2\text{O}_3 \rightarrow 2 \text{Al}[(\text{H}_2\text{PO}_4)_3 + 3 \text{H}_2\text{O}] \quad (9)\]

\[3 \text{H}_3\text{PO}_4 + \text{Al(OH)}_3 \rightarrow \text{Al}[(\text{H}_2\text{PO}_4)_3 + 3 \text{H}_2\text{O}] \quad (10)\]

A cold-setting refractory material was developed by HîPEDINGER et al [36] via the magnesia-
phosphate reaction. A cement paste based on alumina, silica fume, magnesia and orthophos-
phoric acid or monoaluminum phosphate was designed to form cordierite-mullite during

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1 It was proved that H₃PO₄ is effective binder for SiC [29].
heating. This cement paste set at room temperature and MgHPO₄·3H₂O phase (newberyite) was observed, but amorphous phases were predominant. Two exothermic effects were detected during the setting process corresponding to the acido-basic reaction of magnesia with phosphates and to the formation of bonding hydrates. At 1100°C, c-AlPO₄ was formed by the reaction of alumina with orthophosphoric acid or monoauminum phosphate. At 1350°C, the dominant crystalline phases were cordierite and mullite. A refractory concrete with obtained cement paste and a cordierite-mullite aggregate (scrap refractory material) was prepared.

The acid phosphate impregnation, with ozone pretreatment, improves the oxidation resistance of carbon materials (polycrystalline graphite and pitch-based carbon fiber), as shown by the weight measurement in air up to 1500°C. The impregnation involves using phosphoric acid and dissolved aluminum hydroxide in the molar ratio of 12:1 and results in rough, white and hard aluminum metaphosphate coating of the weight of about 20% of that of the carbon before the treatment. Without ozone pretreatment, the impregnation is not effective. Without aluminum hydroxide, the impregnation even degrades the oxidation resistance of carbon [37].

10.1.2. Dental phosphate cement

A variety of cements are used in modern clinical dentistry, such as glass ionomers, zinc phosphate and zinc polycarboxylate [38],[39]. Dental zinc phosphate cement is primarily used for the cementation of indirect restorations, such as crown and bridges. It has the longest record of any cement, approximate 100 years, and has remained popular throughout this time. Zinc phosphate cements are also considered the strongest among the dental cements. However, it is also applied for temporary fillings, cavity bases and buildings of teeth beneath crowns. Zinc phosphate cement is primarily in contact with the pulp-dentin system and in certain cases (e.g. temporary fillings) with the gingiva. A variety of cementing materials are currently used as the bases and luting agents, but zinc phosphate cement has been used for many decades. Phosphoric acid-based cements originated from OSTERMANN’s formula from 1832, which was composed of calcium oxide and anhydrous phosphoric acid. In 1902, FLECK established a formula that is similar to that being in use today [40],[41],[42],[43],[44].

The powder is mainly a mixture of zinc oxide and up to 13% magnesium oxide. The liquid is an aqueous solution of phosphoric acid containing 38 – 59% H₃PO₄, 30 – 55% water and 0 – 10% zinc. Aluminum is essential to the cement-forming reaction, and zinc moderates the reaction between powder and liquid, allowing adequate working time and sufficient quantity of powder to be added for optimum properties of cement. When the powder is mixed with liquid, phosphoric acid attacks the surface of particles, dissolving zinc oxide, which releases zinc ions into the solution. Aluminum in the liquid reacts with phosphoric acid to form zinc aluminophosphate gel on remaining portion of particles. Thus, the cement reveals a cored structure consisting primarily of non-reacted zinc oxide particle core embedded in a cohesive amorphous matrix of zinc aluminophosphate (glasslike phosphate). Aluminum phos-

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5 The word ‘luting’ implies the use of a molded or moldable article to seal a space or to cement two components together [44].
phate, in addition to its role as a retarder, contributes to the increase in mechanical hardness of cement [40],[45].

Setting reactions and resultant structure of zinc phosphate cements are largely based on the formation of hopeite \(\text{Zn}_3(\text{PO}_4)_2\cdot4\text{H}_2\text{O}\) and/or zinc phosphate hydrate \(\text{Zn}_2\text{P}_2\text{O}_7\cdot3\text{H}_2\text{O}\) when using orthophosphoric acid (OPA) cement-forming liquids. OPA solutions buffered with aluminum and zinc ion produced better mechanical properties than non-buffered OPA solutions because of the formation of hopeite and amorphous phase. The development of crystalline forms of phosphate hydrates of zinc and magnesium was retarded and/or prevented by the incorporation of aluminum and zinc ion in the cement-forming liquid [43].

This mechanism is similar to the cement-forming reaction described by \textit{Wilson} [43],[46] in a dental silicate cement (Fig. 5). When \(\text{H}^+\) ions attack the glass powder, \(\text{Al}^{3+}, \text{Ca}^{2+}, \text{Na}^+\) and \(\text{F}^-\) are liberated from the glass, leaving behind an ion-depleted layer of silicate gel at the surface of glass particles. Liberated ions migrate and react with \(\text{H}_2\text{PO}_4^-\), and salts precipitate. The principal reaction is the formation of an insoluble aluminum phosphate, the gel matrix. Associated side reactions are the precipitation of calcium fluoride and the formation of soluble sodium dihydrogen phosphate.

Fig. 5. Setting reaction of dental silicate cement [43].

**10.2. Oil-well cement**

The main application of the cement in an oil well is to stabilize the steel casing in the borehole and to protect it from corrosion. The cement is pumped through the borehole and is pushed upwards through the annulus between the casing and the formation. The cement is exposed to the temperature and pressure gradients of the borehole [47].
Various types of oil-well cements are distinguished. The most important is the phase composition of cement, primarily the C₃A phase content, which causes quick paste thickening. The main feature of oil-well cement is that it must remain sufficiently fluid for a long period, required for its pumping to deep well. Simultaneously, the temperature in the borehole increases with depth [48],[49]. The types and properties of oil-well cements specified by the American Petroleum Institute (API) are introduced by Clinkers containing limestone loess, diatomite, pyritic ash and sand modified with gypsum and apatite were used for the manufacture of heat resistant oil-well cements. Apatite is also a good stabilizer for high belite cements [50].

| API class | Special properties | Intended use |
|-----------|-------------------|--------------|
| A         | Same as ASTM Type I | Well depths up to 1800 m and temperatures of 27 – 77°C. |
| B         | Similar to ASTM Type II, low C₃A, high sulfate resistance | |
| C         | Similar to ASTM Type III, low C₃A, high sulfate resistance | Well depths up to 1800 m and temperatures of 80 – 170°C. |
| D         | Low C₃A with set retarder | Depths up to 1800 – 3600 m and temperatures of 77 – 138°C. |
| E         |           | Depths up to 1800 – 4200 m and temperatures of 77 – 138°C. |
| F         |           | Depths up to 3000 – 4800 m and temperatures of 127 – 160°C. |
| G and H   | Coarse-ground ASTM Types II and IV | Temperatures of 27 – 93°C. |
| J         | Essentially β-C₂S and pulverized silica sand | Depths below 6000 m and temperatures > 177°C |

Table 2 Types and properties of oil-well cements [49].

10.3. Phosphate glasses and glass-ceramics

In the past decades, optical waveguides have raised great interest, as they are the most fundamental and integral part of integrated optic circuits. Glass-based integrated optical devices have several obvious advantages over other technologies such as low intrinsic absorption in near-infrared region of the spectrum, minimized coupling losses to optical fibers and no intrinsic material birefringence compared to crystalline semiconductors. Phosphate glasses are regarded as excellent glass host for the waveguide laser fabrication mainly because of high solubility of rare-earth ions compared to other oxide glasses, which allows for high doping concentrations without significant lifetime reduction, resulting in high gain in short waveguides or cavities and a desirable feature in single-frequency lasers. High-performance

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\[^{a}\] The rate depends on the geothermal degree, which in Europe is about 33 m·°C⁻¹ [48].
waveguide amplifiers or lasers are fabricated in various earth-ion-doped phosphate glasses and commercial phosphate glasses such as Kigre Q89 and Schott-IOG 1 [51].

Glass-ceramics are polycrystalline materials with an inorganic–inorganic microstructure, which are prepared from the base glass by controlled crystallization. This can be achieved by subjecting glasses to regulated heat treatment, which results in the nucleation and growth of one or more crystal phases within the glass [52]. Once a stable crystal nucleus has formed and begun to grow, there are a number of possible crystal growth mechanisms and these determine the final crystal morphology [53],[54]:

a. **Faceted crystal growth**: the favored growth sites on an atomic scale are the steps in rows of atoms. However, these sites can be easily eliminated by the very growth that they promote. Despite this, several types of lattice or crystal defects were shown to provide the growth sites that are impossible to be eliminated by the growth.

b. **Dendritic crystal growth**: formed dendritic crystals are characterized by their tree-like appearance. This type of solidification usually takes place in metals and sometimes occurs naturally in silicate minerals, especially in olivine during rapid cooling of lavas.

c. **Spherulitic crystal growth**: is most commonly associated with organic polymeric materials. The geological definition of a spherulite is: “A crystalline spherical body built of exceedingly thin fibers radiating outwards from a center and terminating on the surface of the sphere...”.

Bioactive glass-ceramics are an alternative to synthetic HAP (Section 10.9) for the use in vivo both in restorative dental applications and in bone implantation [54],[55]. Artificial materials implanted into the bone defects are generally encapsulated by fibrous tissue isolating them from the surrounding bone. This is the normal response of the body towards inert artificial materials. However, some ceramics, such as bioglass, A-W glass-ceramics and sintered hydroxyapatite form a bone-like apatite on their surfaces in the living body and bond to living bone through this apatite layer. This bone-bonding ability is called the bioactivity [56].

These bioactive ceramics are already used clinically as important bone-repairing materials. Their bone-bonding ability is achieved by the formation of a biologically active apatite layer after the reaction of the ceramics with surrounding body fluid. Controlled surface reaction of the ceramics is an important factor governing its bioactivity as well as its biodegradability [56]. The preparation of glass-ceramics with high CaO/P₂O₅ ratio, containing large amounts of calcium phosphate crystals, is believed to be one of the best approaches to obtain the bioceramic implants suitable for bone replacement/regeneration [57].

### 10.3.1. Alkaline aluminum phosphate glasses

Alkaline aluminum phosphate glasses (NMAP) with excellent chemical durability for thermal ion-exchanged optical waveguide were investigated by Wang et al [51]. The transition temperature \( T_g = 470 \degree C \) is higher than the ion-exchange temperature \( 390 \degree C \), which is favorable for sustaining the stability of the glass structure for planar waveguide fabrication. The glass-forming region in the \( \text{Na}_2\text{O}-\text{Al}_2\text{O}_3-\text{P}_2\text{O}_5 \) system is shown in Fig. 6 [51],[58].
10.3.2. Iron phosphate glasses

Iron phosphate glass is a versatile matrix for the immobilization of various radioactive elements found in high-level nuclear waste (HLW). Among various compositions of iron phosphate glass, the one with 40 mol.% Fe₂O₃-60 mol.% P₂O₅ was found to be chemically durable. It also has the ability to accommodate large amounts of certain nuclear wastes, especially those that are not well suited for borosilicate glasses. Better chemical durability of iron phosphate glass is attributed to the presence of more hydration-resistant Fe-O-P bonds compared to P-O-P bonds available in other phosphate glasses [59],[60],[61].

10.3.3. Lithium vanado-phosphate glasses

Lithium vanado-phosphate (LiVP) glasses have been largely studied due to their potential application as cathode materials as a result of mixed electronic-ionic conductivity character. Furthermore, lithium and vanadium structural rearrangements in the glass matrix could modify the transport properties of the systems. Interestingly, the modifier ions depolymerize the glass network, creating useful channels that enhance the ionic conductivity, but they can also break some V⁴⁺/O/V⁵⁺ linkages that are essential to the electronic conductivity because they are supposed to be the preferential path for small polaron hopping. The population of

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7 Fast ion conducting (FIC) phosphate glasses have become very important due to a wide range of applications in solid-state devices [63].
V$^{4+}$/O/V$^{5+}$ paths depends even on the V/P ratio and it can modify the main structure role of vanadium ions in glasses [62],[63],[64].

10.3.4. Apatite-wollastonite glass-ceramics

Wollastonite (CaSiO$_3$) is white glassy silicate mineral that occurs as masses or tabular crystals of metamorphosed limestone. A silica chain GC that contained crystalline apatite and wollastonite (AW) was introduced in MgO-CaO-SiO$_2$-P$_2$O$_5$ glassy matrix and it showed excellent bioactivity, biocompatibility, machinability and adequate mechanical properties such as Young’s modulus (117 GPa), compressive strength (1080 MPa) and bending strength (215 MPa) [52],[65],[66],[67].

Wollastonite-2M and pseudowollastonite (low- and high-temperature forms of wollastonite, respectively) are the most common calcium silicate biomaterials proposed for bone tissue regeneration [68]. The major drawback of the CaSiO$_3$ bioceramics is their relatively fast dissolution rate that could reduce their mechanical strength. In addition, the pH of surrounding medium significantly increases, which could affect the osseointegration of the substitute material within the natural bone. The problem can potentially be solved by the development of multiphase materials containing highly dissolvable phases such as wollastonite, on one hand, and stable phases such as HAP, on the other hand [69].

From the bioactive ceramics, the A/W glass-ceramics show high bioactivity and high mechanical strength [56]. The A/W glass-ceramics composed of apatite and wollastonite crystalline phases in a glassy matrix was developed by KOKUBO [70]. This bioceramics is highly bioactive and also mechanically strong in comparison with other glasses and glass-ceramics because of wollastonite and apatite crystals’ presence.

The A/W glass-ceramics is used in some medical applications, either in powder form as a bone filler or as a bulk material. These materials are currently manufactured by the powder processing methods, providing uniform crystallization of apatite and wollastonite phases in the glassy matrix, as the crystallization of parent glass in a bulk form leads to the appearance of large cracks [69],[71]. The glass-ceramics exposed to the SBF releases predominantly Ca and Si ions, due to the dissolution of amorphous phases and wollastonite-2M, leading to the formation of an apatite-like layer on the surface of the material [69].

The addition of ZnO increased the chemical durability of A-W glass-ceramics, resulting in a decrease in the rate of apatite formation in simulated body fluid. On the other hand, the release of zinc from the glass-ceramics increased with increasing ZnO content. The addition of ZnO may provide bioactive CaO-SiO$_2$-P$_2$O$_5$-CaF$_2$ glass-ceramics with the capacity for appropriate biodegradation as well as the enhancement of bone formation [56]. The effect of MgO was investigated by Ma et al [72]. As the MgO content increased, the glass crystallization temperature increased and the crystallization of the glass-ceramics was changed from the bulk crystallization to the surface crystallization. The addition of MgO slowed down the rate of dissolution and retarded the formation of apatite layer.
10.3.5. Apatite-mullite glass-ceramics

Apatite-mullite glass-ceramics crystallize from the glass of generic composition SiO$_2$-Al$_2$O$_3$-P$_2$O$_5$-CaO-CaF$_2$ to form an osseoconductive apatite phase existing as spherulites within a mullite matrix. The formation of apatite spherulites is accompanied by a depletion zone, from which calcium, phosphate and fluorine is taken and added to the growing crystal. The depletion zone also inhibits the formation of further apatite crystals in the immediate vicinity due to the glass’ compositional similarity with a mullite composition. As mullite begins to crystallize, there is interdependence between the growth of apatite and mullite. Spherulitic grain boundaries in partially devitrified apatite-mullite ceramics act as crack promoters, offering preferential paths to propagation due to the grain boundary interfacial surface energy [54],[73].

10.3.6. FAP-anorthite-diopside glass-ceramics

There is a considerable interest in oxyfluoride glasses and glass-ceramics for laser amplifiers and up-conversion processors. Fluoride-containing crystals have low phonon energies and apatite crystals, in particular, are good host phases to adsorb rare-earth ions. Fluoride-containing glass-ceramics also often crystallize on a nanoscale, which is an added advantage, since optically transparent materials are required for the applications such as fiber amplifiers. The evidence of the nanoscale crystallization in an FAP-anorthite-diopside-based glass-ceramics was found by Hill et al [74].

10.3.7. Apatite-wollastonite ceramics

Wollastonite-hydroxyapatite ceramics was successfully prepared by a novel method, corresponding to the thermal treatment of a silicone embedding micro- and nanosized fillers in air. CaCO$_3$ nanosized particles, providing CaO upon the decomposition, acted as “active” filler, whereas different commercially available or synthesized hydroxyapatite particles were used as “passive” filler. The homogeneous distribution of CaO, at a quasi-molecular level, favored the reaction with silica derived from the polymer, at only 900°C, preventing extensive decomposition of hydroxyapatite. Open-celled porous ceramics suitable for scaffolds for bone-tissue engineering applications were easily prepared from the filler-containing silicone resin mixed with sacrificial PMMA microbeads as the templates [75].

10.3.8. Oxyapatite glass-ceramics

The crystallization of oxyapatite of the composition Na$_x$Y$_y$(SiO$_4$)$_6$O$_2$ (P6$_3$/m, a = 9.334 Å and c = 6.759 Å, $c/a = 1:0.7241$, $V = 509.97$ Å, Fig. 7) from three different glasses from the SiO$_2$-B$_2$O$_3$-Al$_2$O$_3$-Y$_2$O$_3$-CaO-Na$_2$O-K$_2$O-F glass-ceramics system with different F and B$_2$O$_3$ content after the
heat treatment was observed by Van’t Hoff et al [76]. The formation of oxyapatite proceeds according to the mechanism of controlled surface nucleation and crystallization. Therefore, applying suitable chemical composition and heat treatment, it is possible to produce glass-ceramics containing oxyapatite crystals. This type of crystal exhibits similar crystal structure as that of fluorapatite. Therefore, there is an isotype relationship between those two phases: the differences in the crystal structures are found in the structural units, with fluorapatite containing [PO$_4$]$^{3-}$ and [F]$^-$, while oxyapatite containing [SiO$_4$]$^{4-}$ and [O]$^{2-}$. Because of its specific optical properties (high opacity), this new glass-ceramic material may be used as layering material for dental restoration [76].

10.4. Solid oxide fuel cells

Solid oxide fuel cell (SOFC) is an electrochemical energy-conversion device, which offers tremendous promise for delivering high electrical efficiency and significant environmental benefits in the terms of fuel flexibility (hydrocarbons and municipal waste) as well as clean and efficient (>70% with fuel regeneration) electric power generation. SOFC produces useful
electricity by the reaction of fuel with an oxidant via the diffusion of oxide ions (or protons) through an ion-conducting solid-electrolyte layer [77].

SOFC is composed of a dense electrolyte layer that is sandwiched between two porous electrodes (i.e. cathode and anode) as shown in Fig. 8. SOFC can use either oxide ion (a) and/or proton conduction through the electrolyte (b). Electrons generated through the oxidation of fuel on anode are accepted for the oxygen reduction on cathode, which completes the external circuit. The electricity is, thus, produced by the flow of electrons in the external circuit (from the anode to the cathode). Since the current is obtained via the diffusion of oxide ions (or protons) through a solid electrolyte, it becomes imperative to use high operating temperatures (~800 – 1000°C) for achieving high ionic conductivity (of ~0.1 S·cm$^{-1}$) [77],[78].

The first fuel cell was invented in 1838 by an English scientist, William Grove. He named it “wet cell battery” or “Grove cell”, which operated by reversing the electrolysis phenomena of water [77],[79]. The fuel cell, invented in 1839 by Grove, is an electrochemical device that converts the chemical energy of fuels directly into electricity and heat by electrochemically combining H$_2$, CO/H$_2$ or reformed hydrocarbons in fuel and an oxidant gas transported via an ion-conducting electrolyte. Direct combustion of fuels is eliminated here, which renders the fuel cells much higher conversion efficiencies compared to other conventional thermomechanical methods. Moreover, with fuel cells, the power generation is virtually noise-free and can produce 0.9 times lower emissions of NO$_X$ and SO$_X$ per unit of power output compared to that of conventional technologies. Additionally, it is possible to use the fuel cells for combined heat and power (CHP or cogeneration) generation [77],[80].

The development of high-performance SOFC involves the material selection and operation-related issues (of anode, cathode, electrolyte, sealant and interconnects). These challenges open up the myriad research opportunities for researchers in the field of SOFC. A list of various materials used in SOFC is presented in Fig. 9.
Apatite-type silicates are considered as promising electrolytes for solid oxide fuel cells. Lanthanum silicate with the composition of \( \text{La}_{10.5}\text{Si}_{5.5}\text{Al}_{0.5}\text{O}_{26.75} \) was evaluated as an electrolyte with the electrode materials commonly used in SOFC, i.e. manganite, ferrite and cobaltite as cathode materials and NiO-CGO composite, chromium-manganite and \( \text{Sr}_2\text{MgMoO}_6 \) as anode materials by MARRERO-LÓPEZ et al [81]. This electrolyte has conductivity values higher than those of YSZ and comparable to most important solid electrolytes proposed for the intermediate-temperature range, such as doped ceria and lanthanum gallate-based electrolytes. The chemical compatibility did not reveal appreciable bulk reactivity between silicate and many electrodes up to 1300°C [81]. Among several reported rare earth apatites, lanthanum silicates exhibit the ionic conductivity higher than their germinate counterparts [77].

On examining and modeling the probable conduction mechanism in apatite silicates, the atomistic simulation results suggest that the conduction in \( \text{La}_{9.33}(\text{SiO}_4)_6\text{O}_2 \) and \( \text{La}_8\text{Sr}_2(\text{SiO}_4)_6\text{O}_2 \) takes place via interstitial and vacancy mechanism, respectively [82],[83]. The predicted pathway appears to be a complex nonlinear “sinusoidal-like” process for the interstitial oxygen migration along the c-axis (Fig. 10), while a direct linear pathway is predicted for oxygen migration via the vacancy mechanism [77].

ZENG et al [84] developed a model capable of prediction of oxygen ion conduction from relative Coulomb electronic interactions in oxyapatites and rationalized observed experimental trends reported in the literature. Two types of fundamental chemical property, i.e. the electronegativities and ionic radii of the constituents, control oxygen ionic conduction in oxyapatites. Those two properties were used to represent the relative charge densities and the distances between charged units, respectively, and then to formulate the relative Coulomb energy. It was found that this relative Coulomb energy is linearly correlated to the oxygen ionic conductivity.
conductivity (in logarithmic form) in the oxyapatite systems. Doping a cation with large ionic radius and low electronegativity tends to increase the ionic conductivity of oxyapatite.

The c-axis-oriented apatite-type lanthanum silicate (La$_{10}$Si$_6$O$_{27}$) ceramics was prepared by NAKAYAMA et al [85] via the sintering process under high magnetic field. The degree of orientation in the La$_{10}$Si$_6$O$_{27}$ ceramics sintered at 1700°C was 48.1%. The conductivity of the c-axis-oriented ceramics is about 0.5 orders of magnitude higher than that of non-oriented ceramics. Higher conductivity is caused by the orientation of oxide ions in the grains composing the ceramics, which are located along the c-axis and are responsible for the ionic conduction.

10.5. Sensors

Compacted sinters of Ln$_{9.33+x/3}$Si$_{6-x}$Al$_x$O$_{26}$ (0 ≤ x ≤ 2.0, Ln = La, Nd and Sm) are composed of an apatite-like phase with a hexagonal structure. Compacted sinters were used as potentiometric oxygen sensors (Fig. 11). The concentration dependence of EMF was well expressed by the Nernst equation:

$$E_{\text{obs}} = \frac{RT}{nF} \ln \frac{p_{O_1}}{p_{O_2}} \quad (11)$$

Furthermore, the electron number $n$ is comparable to the theoretical value of 4. The sensing characteristics of the sinters are comparable to those of the sensors with 3 and 8 mol.% YSZ [86].
Electrical properties and humidity sensor characteristics of lead hydroxyapatite material were reported by Tudorache et al [87]. The electrical characteristics of lead hydroxyapatite material treated at different temperatures made us focus on the analysis of the influence of water vapors upon the electrical characteristics. Thus, the electrical response to humidity adsorptive processes of lead hydroxyapatite material suggested that we analyze the material characteristics in terms of its use as a humidity sensor.

The humidity-sensitivity of yttrium-substituted calcium oxyhydroxyapatites was studied by Owada et al [88]. The logarithm of electrical resistance of the present sensors decreased linearly with increasing relative humidity (RH) from 30 to 65%. The resistance of $[Ca_{9.0}Y_{1.0}] (PO_4)_{6}(O_1.5\square_{0.5})$ with the highest OH vacancy content was about one order of magnitude lower than that of calcium hydroxyapatite. It was found that the larger ratio of surface hydroxyl groups per unit surface area in the sample, the lower the resistance and the higher the amount of OH vacancies.

Using the hydroxyapatite ceramics as CO$_2$ sensor, which is based on electrical conductivity changes, was investigated by Nagai et al [89]. Starting powders prepared by usual wet process were cast in film with an organic vehicle and fired on alumina substrates after the electrodes had been arranged. It was necessary to soak the samples in a CaCl$_2$ solution in order to make them reactive with CO$_2$. Both D.C. and A.C. measurements were carried out in various atmospheres including air, CO$_2$ and air containing different amounts of CO$_2$.

10.6. Phosphors

Fluorescent lamps typically have transparent glass envelope enclosing sealed discharge space containing an inert gas and mercury vapor. When subjected to a current provided by electro-
des, mercury ionizes to produce the radiation having the primary wavelengths of 185 nm and 254 nm. This ultraviolet radiation, in turn, excites phosphors on the inside surface of the envelope to produce visible light that is emitted through the glass. Generally, the fluorescent lamp for illumination uses a phosphor that absorbs the 254 nm Hg-resonance wave; the phosphor is activated so as to convert the ultraviolet light into the visible light. In order to improve the color-rendering properties and the emission output of fluorescent lamps, efficient illumination of a white color has been recently provided using a three-band-type fluorescent lamp, which employs a mixture of red, green and blue-emitting phosphors. In such three-band-type phosphor lamp, the emitting colors of the respective phosphors are considerably different from one another. Therefore, if the emitting intensity of any of the three corresponding phosphors is decreased, the color deviation occurs, degrading the color-rendering properties of the lamp [90]. The literature dedicated to the preparation of apatite-type light-emitting phosphors is really abundant.

A series of orange-red-emitting \( \text{Ba}_2\text{Y}_3(\text{SiO}_4)_3\text{F}:\text{Sm}^{3+} (0.003 \leq x \leq 0.08) \) fluorosilicate apatite phosphors were synthesized via the conventional solid-state reaction by Yu et al [91]. The emission spectra of the \( \text{Ba}_2\text{Y}_3(\text{SiO}_4)_3\text{F}:\text{Sm}^{3+} \) phosphors contained some sharp emission peaks of \( \text{Sm}^{3+} \) ions centered at 564, 601, 648 and 710 nm. The strongest one is located at 601 nm. The optimum dopant concentration of \( \text{Sm}^{3+} \) ions in \( \text{Ba}_2\text{Y}_3(\text{SiO}_4)_3\text{F}:\text{Sm}^{3+} \) is around 3 mol.\% and the critical transfer distance of \( \text{Sm}^{3+} \) was calculated to be 26 Å. The quenching temperature is above 500 K.

Red-emitting phosphors \( \text{Ba}_2\text{Gd}_8(\text{SiO}_4)_6\text{O}_2:\text{Eu}^{3+} \) (BGS:Eu\(^{3+}\)) with silicate apatite structure were prepared by Liu et al [92] via the conventional high-temperature solid-state reaction method. There are two different sites (4\(f \) and 6\(h \) [93]) for Eu\(^{3+}\) occupying the host. It was found that the phosphors BGS:Eu\(^{3+}\) exhibit red emission with high quenching concentration at \( \sim70.75 \) at.\%, and the critical transfer distance of Eu\(^{3+}\) in BGS:Eu\(^{3+}\) was calculated to be \( \sim12.3 \) Å. More importantly, it has better CIE chromaticity coordinate for white light-emitting diode (w-LED) application in comparison with commercial phosphor (Y,Gd)BO\(_3\):Eu\(^{3+}\) (YGB:Eu\(^{3+}\)) under near-ultraviolet (n-UV) 393 nm excitation [92],[94]. White Tb\(^{3+}/\text{Sm}^{3+}\) ions co-doped \( \text{Ca}_2\text{La}_8(\text{GeO}_4)_6\text{O}_2 \) (CLGO) phosphors prepared by Jeon et al [95] show observable emission spectra under 374 nm excitation.

A novel blue-emitting phosphor \( \text{Sr}_8\text{La}_2(\text{PO}_4)_6\text{O}_2:\text{Eu}^{2+} \) was synthesized by Liu et al [96] via conventional high-temperature solid-state method and its photoluminescence (PL) properties were investigated for the applications in white light-emitting diodes. The phosphor exhibited strong broad absorption band in the near-ultraviolet (n-UV) range and generated bright-blue emission centered at 442 nm upon 365 nm excitation light. The critical Eu\(^{2+}\) quenching concentration (QC) mechanism was verified to be the dipole-dipole interaction.

A green-emitting phosphor of Eu\(^{2+}\)-doped \( \text{Ca}_9(\text{PO}_4)_2\text{SiO}_4 \) was prepared via a solid-state reaction by Roh et al [97]. The phosphor was excited at the wavelengths of 220 – 450 nm, which was suitable for the emission band of near-ultraviolet (n-UV) light-emitting diode (LED) (350 – 430 nm). In \( \text{Ca}_9(\text{PO}_4)_2\text{SiO}_4:\text{Eu}^{2+} \) phosphor, there were three distinguishable Eu\(^{2+}\) sites, which resulted in a strong green emission peaking at 530 nm and broad bands up to 700 nm.
10.7. Catalysts

Catalysts are usually defined as the substances that increase the rate at which a chemical reaction approaches the equilibrium without becoming permanently involved in this reaction. Basically, the catalysis can be divided to [98]:

i. Homogeneous catalysis,

ii. Heterogeneous catalysis.

Heterogeneously catalyzed process is more complex because the catalyst is not uniformly distributed throughout the reaction medium. Considering a two-phase system, either vapor/solid or liquid/solid, with the catalyst in the solid phase, the several steps need to be realized to complete the catalytic cycle [98]:

1. Transport of the reactant to the catalyst;
2. Adsorption, i.e. the interaction of the reactant with the catalyst;
3. Reaction of adsorbed species to the products;
4. Desorption of the products from the catalyst;
5. Transport of the product away from the catalyst surface.

Due to their versatility in anionic and cationic composition and their ability to adsorb organic and organometallic molecules as well as metallic salts, the surface properties of apatites can be tuned and they can behave as powerful catalysts in a wide range of organic reactions. In many cases, the apatite-based catalysts can be used without a solvent and show good recycling capacity. The catalytic properties mainly arise from the acid-base character of the apatite’s surface. In some cases, adsorbed moieties are responsible for the catalytic properties, apatites playing the role of a solid support. Finally, the catalytic activities can result from the combination of properties of the apatite’s surface and of the adsorbed or anchored moieties [99].

The oxidative Glaser-Hay coupling reaction of terminal alkynes is a very important reaction in organic chemistry to achieve the synthesis of diyne compounds. In general, the reaction is performed under homogeneous conditions using Cu(I) or Cu(II) salts in the presence of a reagent such as tetramethylethylene diamine (TMEDA), which can bind to copper ions, an organic base and dioxygen. Although this reaction is known for a long time, the mechanism is still under the discussion. It is possible to catalyze the Glaser-Hay reaction under heterogeneous conditions using Cu-modified hydroxyapatite (Cu-HAp). With several para-substituted phenyl-acetylenes and alkynols, we can show that Cu-HAp acts as a catalyst for single-bond coupling reactions leading to diyne derivatives in high yields without using auxiliary chelating molecules and organic bases. These heterogeneous conditions allow easy recovery of the catalyst and simplify the purification work-up.

The oxidative Glaser-Hay coupling reaction of terminal alkynes (acetylenes) is a very important reaction in organic chemistry to achieve the synthesis of diyne compounds. In general, the reaction is performed under homogeneous conditions using Cu(I) or Cu(II) salts in the
presence of base (ethanolic ammonia solution, tetramethylethylenediamine, pyridine, …), which can bind to copper ions, an organic base and dioxygen [100],[101]:

\[
\text{CuCl}_\text{base} \quad \text{R} - \text{C} = \text{CH} \rightarrow \text{R} - \text{C} = \text{C} = \text{–R} \quad (12)
\]

Although this reaction is known for a long time, the mechanism is still under the discussion. It is possible to catalyze the Glaser-Hay reaction under heterogeneous conditions using Cu-modified hydroxyapatite (Cu-HAp). With several para-substituted phenyl-acetylenes and alkynols, where Cu-HAp acts as a catalyst for single-bond coupling reactions leading to diyne derivatives in high yields without using auxiliary chelating molecules and organic bases. These heterogeneous conditions allow easy recovery of the catalyst and simplify the purification work-up [100].

The apatite catalyst was utilized for the catalysis of the synthesis of n-butanol, 1,3-butadiene and high octane fuel from bioethanol. The process requires relatively low temperature. The synthesis shows significantly lower cost compared to n-butanol derived from petroleum-based processes. The technology offers a closed-loop system with no waste or emissions [102].

### 10.8. Phosphate conversion coatings

Conversion coatings provide the resistance to corrosive environments. Phosphate conversion coatings (PCC) bring about the transformation of metal substrates into new surfaces having non-metallic and non-conducting properties. The transformations occur in phosphating solution containing divalent metal phosphates and, in some cases, in solutions containing monovalent metal phosphates. Generally, the solutions are prepared from liquid concentrates containing one or more divalent metals (zinc, magnesium, calcium, etc., phosphates), free phosphoric acid and an accelerator. Three types of phosphate conversion coatings are currently being used [103],[104]:

1. **Zinc phosphate coatings** are often used as a pretreatment for painted parts. They are also used to impart the corrosion resistance and to aid in cold-forming operations.
2. **Iron phosphate coatings** are primarily used to form a passive substrate under paints.
3. **Manganese phosphate coatings** are primarily on machined parts such as gears and internal combustion engine components as an anti-scuff film for the break-in wear.

The addition of metal ions, such as cupric ions, to a conversion bath greatly reduces the formation time and the size and nonuniformity of coating crystals. Copper, which is cathodic to dissolving metals, deposits on the base metal to form many local cells and thus to increase the potential difference between local anode and cathode site. Nickel ions behave differently than cupric ions, and their benefit results from the catalytic action associated with the release of molecular hydrogen. Furthermore, the addition of Ni$^{2+}$ and Mn$^{2+}$ into the treating solution
refines the grain size and reduces the porosity of phosphate conversion coatings on electro-galvanized steels. While Ni exists in both the zero-valance state and the two-valance state, Mn is mainly present in the two-valance state in the phosphate conversion coating [103],[105].

In fact, all of the chemical reactions of phosphating process are based on mutual interactions of metal immersed in phosphate bath and redox of the accelerators. Generally, phosphating proceeds in the acidic solution containing Zn$^{2+}$, Mn$^{2+}$, Ca$^{2+}$, Na$^{+}$, Fe$^{2+}$ and Mg$^{2+}$. Phosphating with different metal substrates and types is not of the same reaction mechanism. For example, when a pure iron is immersed in the phosphating solution, iron dissolves on the micro-anodes through the following reaction [106]:

$$\text{Fe} + 2 \text{H}^+ \rightarrow \text{Fe}^{2+} + \text{H}_2$$  \hspace{1cm} (13)

The hydrogen evolution occurs at the micro-cathodic sites resulting in an increase of pH value at the metal-solution interface. This change in pH alters the dissociation equilibrium, which leads to the formation of PO$_4^{3-}$:

$$\text{H}_3\text{PO}_4 \rightarrow \text{H}_2\text{PO}_4^- + \text{H}^+ \rightarrow \text{HPO}_4^{2-} + 2 \text{H}^+ \rightarrow \text{PO}_4^{3-} + 3 \text{H}^+$$  \hspace{1cm} (14)

When PO$_4^{3-}$ and Me$^{2+}$ (metal ion, e.g. Zn$^{2+}$, Mn$^{2+}$, Ca$^{2+}$ and Fe$^{2+}$) in the solution reach the saturation, the deposition of insoluble phosphate will be achieved. Then, it can crystallize in PCC coating, as shown in Fig. 12.

---

**Fig. 12.** Schematic representation of the deposition process of PCC coating on the surface of pure iron [106].
In the zinc phosphating solution, the addition of Zn$^{2+}$ supports the formation of crystals of zinc phosphate. Zinc ions combine with phosphate ions to form an insoluble film. The formation of hopeite is described by the reaction [106]:

\[
3 \text{Zn}^{2+} + 2 \text{H}_2\text{PO}_4^- + 2 \text{H}^+ + 4 \text{H}_2\text{O} + 6 \text{e}^- \rightarrow \text{Zn}_3(\text{PO}_4)_2 \cdot 4\text{H}_2\text{O} + 3 \text{H}_2
\]

(15)

In some cases, Zn and ZnO were found in the phosphating process in reactions:

\[
\text{Zn} + 2 \text{e}^- \rightarrow \text{Zn}
\]

(16)

\[
\text{Zn} + 2 \text{H}_2\text{O} \rightarrow \text{Zn(OH)}_2 + \text{H}_2
\]

(17)

\[
\text{Zn(OH)}_2 \rightarrow \text{ZnO} + \text{H}_2\text{O}
\]

(18)

In the phosphate solution of coexisting Zn$^{2+}$ and Ca$^{2+}$, zinc calcium phosphate can be formed by the reactions:

\[
\text{Ca}^{2+} + \text{HPO}_4^{2-} \rightarrow \text{CaHPO}_4
\]

(19)

\[
\text{Ca}^{2+} + 2 \text{Zn}^{2+} + 2 \text{H}_2\text{PO}_4^- + 2 \text{H}_2\text{O} \rightarrow \text{CaZn}_2(\text{PO}_4)_2 \cdot 2\text{H}_2\text{O} + 4 \text{H}^+
\]

(20)

It is notable that the phosphating mechanism varies in different phosphating systems and materials [106].

The formation of conversion coating on zinc-coated samples under cathodic conditions was studied by Perrin et al. [107] in a chromating bath containing phosphate (phosphate-chromate solution). Thick chromium phosphate (Cr-P) coating has two distinct layers: an outer porous layer and an inner and thinner pore-free adherent one. Both layers contain chromium phosphate as the main constituent and, to a lower extent, zinc phosphate species, the concentrations of which decrease from the metal-coating interface outwards. Formed zinc phosphates have a general formula of $x\text{CrPO}_4 \cdot y\text{Zn}_3(\text{PO}_4)_2 \cdot z\text{H}_2\text{O}$ where $x > (y, z)$.

### 10.9. Synthetic apatite analogues in tissue engineering

Biomaterials in general are based on the materials such as metals, polymers, and ceramics. Typical metallic biomaterials are based on stainless steel, cobalt-based alloys, titanium or
titanium alloys and amalgam alloys. Polymeric biomaterial composites from monomers are based on amides, ethylene, propylene, styrene, methacrylates and/or methyl methacrylates. Biomaterials based on ceramics are found within all classical ceramic families (Table 3) including traditional ceramics, special ceramics, glasses, glass-ceramics, coatings and chemically bonded ceramics (CBCs) [13].

| Ceramic material classification | Example                                      |
|-------------------------------|----------------------------------------------|
| Traditional ceramics          | Dental porcelain, leucite-based ceramics     |
| Special ceramics              | Al, Zr and Ti oxides                         |
| Glass                         | Bioglass (Na, O-CaO-P2O5-SiO2)               |
| Glass-ceramics                | Apatite-wollastonite, Li-silicate-based      |
| CBCs                          | Phosphates, aluminates, silicates and sulfates |

Table 3 Examples of biomaterials based on ceramics [13].

Whereas many chemists and materials scientists consider the biomaterial to be synthetically produced material, most biologists, geologists and mineralogists consider the materials such as bone and tooth, which are biologically produced, to be the biomaterials. Also a common reference to Ca:P (1.67, Table 7 in Chapter 1) ratio is usually used in the biomaterials literature, which disregards the fact that different calcium phosphate phases have different crystalline structures.

There are many phosphate minerals and salts (Table 4) that do not have the crystalline structure of apatite [108], [109]. Apatite-based materials have attracted a considerable interest for orthopedic and dental applications because of their biocompatibility and tight bonding to bone, resulting in the growth of healthy tissue directly onto their surface. Several combinations of apatite and other phases were proposed in order to improve poor mechanical properties of apatite [110].

| Typical acronym | Chemical name                  | Chemical formula | Mineral name       | Structure     | Ca/P ratio |
|-----------------|--------------------------------|------------------|--------------------|---------------|------------|
| HAP, HA         | Tribasic calcium phosphate     | Ca₅(PO₄)₃OH     | Hydroxylapatite    | Apatitic      | 1.67       |
| ACP             | Amorphous calcium phosphate    | ?                | —                  | —             | ?          |
| PCHA, PCA       | Poorly crystalline hydroxylapatite | Ca₅(PO₄)₃OH     | Hydroxylapatite    | Apatitic      | 1.67       |
| CAP             | Carbonated apatite             | Refer to Sections 2.6 and 4.6 |                |               |            |
| TCP             | Tricalcium phosphate           | Ca₃(PO₄)₂       | Whitlockite        | Non-apatitic  | 1.5        |
| β-TCMP          | Magnesium-substituted TCP      | Ca₅(PO₄)₃OH     | Whitlockite        | Non-apatitic  | ≤1.5       |
Typical acronym | Chemical name | Chemical formula | Mineral name | Structure | Ca/P ratio
---|---|---|---|---|---
? | “Tricalcium phosphate” Ca$_3$(Mg,Fe$^{2+}$)(PO$_4$)$_2$(HPO$_4$) | Geologically occurring whitlockite | Non-apatitic | 1.28
CPPD | Calcium pyrophosphate dihydrate | Ca$_3$P$_2$O$_7$·2H$_2$O | — | Non-apatitic | 1.0
γ-CCP | γ-Calcium pyrophosphate | Ca$_3$P$_2$O$_7$ | — | Non-apatitic | 1.0
OCP | Octacalcium phosphate | Ca$_8$H$_2$(PO$_4$)$_6$·5H$_2$O | — | Non-apatitic | 1.33
MON | Dibasic calcium phosphate | Ca(HPO$_4$)$_2$·5H$_2$O | Monetite | Non-apatitic | 1.0
DCPD | Dicalcium phosphate dihydrate | Ca(HPO$_4$)$_2$·2H$_2$O | Brushite | Non-apatitic | 1.0

| Chemical name | Chemical formula | Mineral name | Structure | Ca/P ratio |
|---|---|---|---|---|

| | | | | |

Table 4 Different apatitic and non-apatitic calcium phosphates [108].

Small organic molecules incorporated in apatite crystals act as porogens that control the porous structure of apatite single crystal. The presence of amino acid under the apatite synthesis conditions leads to firm bindings and encapsulation of amino acid within apatite single crystals. The amino acid elimination by heating or electron beam irradiation enhances the pore formation in apatite single crystals. Moreover, the incorporation of acidic amino acid into apatite induces the peapod-like nanotubes in apatite single crystals. That suggests the potential of using small organics for nanostructural control of apatite single crystals, which would be valuable for enhancing the drug loadings or for modulating the material digestion in vivo [111].

10.9.1. Biological apatite in bone tissue engineering

Tissue engineering (TE) techniques were developed to recover or enhance lost tissue function and structure. In biological hard tissues, for example, lost portions can be effectively reconstructed by the control of environmental factors, physical stimulation, addition of growth factors and by the use of degradable materials. These factors strongly facilitate the regeneration of macroscopic shape of defected hard tissues. Nevertheless, the differences in microstructure, and also in mechanical and physical properties, between regenerated and original hard tissues must be examined prior to clinical application [112],[113],[114].

For in vitro engineering of living tissues, cultured cells are grown on bioactive degradable substrates (scaffolds) that provide the physical and chemical cues to guide their differentiation and assembly into three-dimensional structures. One of the most critical issues in TE is the realization of scaffolds with specific physical, mechanical and biological properties. Scaffolds act as substrate for cellular growth, proliferation and the support for new tissue

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Scaffolds might be defined as artificial structure capable of supporting the three-dimensional tissue formation, which allows the cell attachment and migration, the delivery and retaining of cells and biochemical factors and enables the diffusion of vital cell nutrients and expressed products. In the case of bone, scaffolds should replicate its architecture and three-dimensional structure with predetermined density, hierarchical pore distribution and interconnected pathways [109],[115].
Biomaterials and fabrication technologies play a key role in tissue engineering. Materials used for tissue engineering applications must be designed to stimulate specific cell response on the molecular level. They should elicit specific interactions with the cell and thereby direct cell attachment, proliferation, differentiation and extracellular matrix production and organization [109],[115]:

- **Biocompatibility** is defined as the ability to perform its function in the host tissue without eliciting any immune response.

- **Biodegradability** denotes tunable rate of degeneration to match the growth of new bone tissue as scaffold gets replaced by new bone.

- **Mechanical properties** include the properties such as sufficient mechanical strength to provide temporary support to the defect region and withstand in vivo loading forces.

- **Microarchitecture** is an interconnected scaffold structure that uniformly distributes stresses throughout the scaffold.

- **Osteoinductivity** includes osteoinductive properties that enable to recruit and differentiate the osteoprogenitors to the defect region.

- **Porosity** is large surface area, where the volume and the pore size allow the tissue in-growth, neovascularization, mass transport and the osteogenesis.

- **Surface properties** include appropriate chemical and topographical properties for influencing cellular adhesion, proliferation and differentiation.

Inorganic-organic composites aiming at mimicking the composite nature of real bone combine the toughness of the polymer phase with the compressive strength of an inorganic one to generate bioactive materials with improved mechanical properties and degradation profiles. Hydroxylapatite (HAP) is widely used as a biocompatible ceramic material in many areas of medicine, but mainly for the contact with bone tissue, due to its resemblance to mineral bone [115].

Under normal conditions, human body fluid is supersaturated with respect to apatite, so that, when apatite nuclei form, crystals can grow spontaneously. Chemical species that are capable of supporting the nucleation of apatite are calcium and silicate ions, while, in contrast, phosphate ion does not affect this process. Hence, ceramics that release the former mentioned ions are capable of developing the surface apatite layer when exposed to human body fluids. This explains why glasses that do not contain P$_2$O$_5$ are able to develop this surface layer and show greater bioactivity than glasses containing P$_2$O$_5$ even if the latter compositions approximate to that of hydroxyapatite [116].

**10.9.2. Preparation of nanocrystalline apatites**

Nanocrystalline calcium phosphate apatites play an important role in biomineralization and in the field of biomaterials. Biological nanocrystalline apatites are the main inorganic compo-

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*The enamel of vertebrate teeth, vertebrate bone and tooth-like microfossils of conodonts.*
ments of hard tissues in mammals (bone\textsuperscript{10} and tooth\textsuperscript{11} (Fig. 13)) with the exception of enamel (which is closer to stoichiometric hydroxylapatite) and are involved in several pathological calcifications such as dental calculi, salivary stones and blood vessel calcification. In comparison with hydroxylapatite, which is a stoichiometric apatitic phase and is the most stable and the least soluble calcium phosphate at ambient conditions, nanocrystalline apatites are nonstoichiometric and calcium (and OH-) deficient and may incorporate substituted ions in their nanosized crystals. Their calcium and hydroxide deficiencies are responsible for higher solubility than HA. Besides, they have the ability to mature when submitted to humid environments [108],[112],[117],[118].

Fig. 13. The cross-section of lower jaw bone (a) and tooth (b) of European roe deer (\textit{Capreolus capreolus}) and human tooth (c, d): enamel (1), dentin (2), pulp chamber (3), cementum (4) and root canal (5).

\textsuperscript{10}Bones are rigid organs that form a part of the endoskeleton of vertebrates. Their function is to move, support and protect various organs of the body, produce red and white blood cells and store minerals. Bones appear in a variety of shapes and have a complex internal and external structure described by various hierarchical models [117]. Bone is a complex and hierarchical tissue consisting of nano-hydroxylapatite (70 wt.\%) and collagen (30\%) as major portions [113].

\textsuperscript{11}Teeth consist of a bulk of dentin covered with (inorganic) enamel on the crown and cementum on the root surface. Thick collagen bundles, called periodontal ligaments (PDL), attach to cementum at one end and to the alveolar bone at the other end. The alveolar bone is supported by the jaw [117].
Scanning electron microscopy of deer bone (Fig. 14) shows a spongiform texture \((a, b)\) formed by crystals of carbonated hydroxylapatite with the size below 100 nm \((e, f)\).

The inorganic portion of bone contains two major mineral phases

i. Non-crystalline calcium phosphate;

ii. Crystalline bone apatite.

In amorphous calcium phosphate, the Ca:P ratio is about 1.33. The non-crystalline or amorphous bone mineral is metastable with respect to bone apatite. Bone apatite crystals are calcium deficient due to the defects in the crystalline lattice and isomorphous substitutions, that is, the replacement of some ions by others in the crystal without disrupting the general symmetry. Young bone tissue was found to be richer in amorphous mineral than crystalline apatite [119].

In the body, only the bone collagen has the property of inducing the mineralization through in vitro; collagens of other tissues do not possess this property. One of the differences between nucleating and non-nucleating collagens might be the presence of a sort of inhibitor bound to
the latter. This inhibiting substance was shown to be pyrophosphate. Calcium is accreted in bone tissue in the process of new bone formation or remodeling and resorbed from the bone tissue in the process of bone destruction. The loss of endogenous calcium in urine and feces is compensated for by an equivalent intake of this element (Fig. 15) [119].

![Diagram of calcium metabolism]

**Fig. 15.** General scheme of metabolism of calcium [119].

Enamel (Fig. 17) is normally the best preserved from hard tissues. It is almost completely a mineral, so the decomposition of organic matter has little effect on it. Archeological enamel nearly always yields good microscope sections, often indistinguishable from fresh enamel [120]. The structure of carbonate dental enamel refined by Wilson et al [121] is shown in Fig. 16.

![Crystallographic structure of human dental enamel apatite]

**Fig. 16.** Crystallographic structure of human dental enamel apatite (perspective view along the c-axis): P63/m, \(a = 9.4081 \text{ Å}, b = 6.8887 \text{ Å}, c:a = 0.7322\) and \(V = 528.05 \text{ Å}^3\) [121].
Among carbonate apatites, which are formed in mouth cavity, only dental enamel mine belongs to the B-type of carbonate-apatite (Section 4.6) and the others (of dentin, salivary and dental stones) belong to the AB-type (B > A). According to the variations in unit-cell parameters, isomorphic replacements in crystal structures of apatites of pathogenic origin (renal, salivary and dental stones) are more intensive in comparison with physiogenic dental enamel apatites. Among pathogenic apatites, the most considerable compositional variations are observed in renal stone apatites. That indicates strong variability of conditions of their formation. The changes in unit-cell parameters of bone apatites are not completely interpretable, because these apatites consist of nanosized crystals that are smaller than those of other biological apatites [122].
The age variations of the crystal lattice parameters of human enamel apatites are related to complicated processes of de- and remineralization, which result in the increase or reduction of vacancies in Ca positions and in the respective changes of \( \text{CO}_3^{2-}, \text{H}_2\text{O} \) and \( \text{HPO}_4^{2-} \) contents in the unit cell:

\[
\begin{align*}
(n + m / 2) \text{Ca}^{2+}n \text{PO}_4^{3-} + m \text{OH}^- & \rightarrow \nonumber \\
(n + m / 2) \left[ \text{Ca}^{2+} + n \text{A}^{2+} + m \left[ \text{OH}^- , \text{H}_2\text{O} \right] \right] & \end{align*}
\]

where \( \text{A}^{2+} = \text{CO}_3^{2-} \) or \( \text{HPO}_4^{2-} \), and \( \left[ \text{Ca}^{2+} , \text{OH}^- \right] \) are the vacancies. Until the age of 50 years, the values of \( a \) and \( c \)-parameter of enamel apatites change considerably without any dependence of particular age that may be explained by essential fluctuations of the content of Ca in human organism. After 50 years of age, significant direct correlation between the age and the \( a \)-parameter appears [122].

The surface of apatite nanocrystals is possibly doped with foreign elements or functionalized with organic molecules [117],[123],[124]. The course of facile synthesis of B-type carbonated nanoapatite with tailored microstructure is described by Gualtieri et al [125].

### 10.10. Collagen apatite composites

Tissue engineering techniques have been developed to recover or enhance lost tissue function and structure. Collagen-apatite (or collagen/apatite, Col-AP) composite resembling the composition of natural bone has been studied extensively and considered as a promising bone tissue engineering material, which can be used to replace or regenerate damaged tissue, resulting from an accident, trauma or cancer. Such synthetic or hybrid biomaterials must have high porosity with interconnected pores to allow the vascularization as well as the nutrients and gases diffusion. Moreover, they should be biodegradable to act as temporary cellular support [114],[126],[127],[128].

The defining feature of collagen is a structural motif in which three parallel polypeptide strands in a left-handed, polyproline II-type (PPII) helical conformation coil around each other with a one-residue stagger to form a right-handed triple helix. Tight packing of PPII helices within the triple helix mandates that every third residue be Gly, resulting in a repeating XaaYaaGly sequence, where Xaa and Yaa can be any amino acid. This repeating occurs in all types of collagen, although it is disrupted at certain locations within the triple-helix domain of nonfibrillar collagens. The amino acids in the Xaa and Yaa positions of collagen are often (2S)-

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12 One of the first polymers used for bone tissue engineering was based on a hydrolytically copolymer of polylactic-co-glycolic acid (PLGA) but its use for large bone defect regeneration was controversial as inflammatory events were observe. The utilization of chitosan and alginate was also investigated [128].

13 Names such as collagen-hydroxylapatite or collagen-hydroxyapatite composite were also often applied in published literature.
proline (PRO, 28%) and (2S,4R)-4-hydroxyproline (HYP, 38%), respectively. The PROHYPGLY is the most common triplet (10.5%) in collagen (Fig. 18) [129].

![Fig. 18. The triple helix of collagen formed from (PROHYPGLY)_n-(PROHYPALAA)- (PROHYPGLY)_n (a), the view down the axis of a (PROHYPGLY)_n triple helix (b) and the segment of triple helix (c) with hydrogen bonds (—) [129].](image)

The categories of collagen include the classical fibrillar and network-forming collagens, the FACITs (fibril-associated collagens with interrupted triple helices), MACITs (membrane-associated collagens with interrupted triple helices) and MULTIPLEXINs (multiple triple-helix domains and interruptions). The collagen of type I is the most abundant type used in tissue engineering. Natural polymer collagen that represents the matrix material of bone, teeth and connective tissue can be extracted from animal or human sources (skin, bones, tendons, ligaments and cornea). The treatment includes the separation and isolation (in soluble or insoluble form), decalcification, purification (purification is required to eliminate the antigenic component of protein), sterilization and chemical modification process to achieve polyanionic or purified protein. Type I polyanionic collagen was found to improve the cell adhesion [129],[130].

Collagen, the most abundant protein in extracellular matrix, is chemotactic to fibroblasts. It shows high affinity to cells and good resorbability in vivo. Nevertheless, its poor mechanical properties have restricted its usage in load-bearing applications. Carbonated apatite and collagen interact to form a composite material, the mechanical, physicochemical and biological properties of which differ considerably from those of either constituent considered separately. Collagen and non-collagenous proteins (NCPs) are thought to control the crystal deposition, size, crystallization and multiplication/maturation. On the other hand, the crystal deposition in intrafibrillary spaces is likely to modify the three-dimensional conformation of collagen [114],[131],[132],[133]. The solubility of apatite-collagen composites is significantly reduced by UV radiation [134],[135].

14 There are 27 types of collagen described in literature composed of at least 46 distinct polypeptide (CRP, collagen-related peptide) chains [129], but the types I – V are the most common. More than 90% of collagen in human body is the fibrillar type I.
Calcium phosphates are available commercially, as hydroxylapatites are extracted from bones or they can be produced wet by direct precipitation from pH-adjusted solutions of calcium and phosphate salts [130]. The crystallographic c-axes of the plate-shaped apatite crystals are well aligned with long axes of collagen fibrils (Fig. 19), and this preferred orientation between the mineral and the organic framework is assumed to be the general feature of the calcium phosphate biomineralization process. Several attempts were made to mimic this lowest level of hierarchical organization of bone by using proteins as a site for the heterogeneous nucleation and subsequent growth of stoichiometric hydroxylapatite crystals. In special approaches, the biomimetic apatite coatings on surfaces were prepared by soaking the materials in simulated body fluid (SBF) solutions, which contained ions in the concentrations similar to those in inorganic part of human blood plasma. It is generally accepted that the in vitro apatite growth during the exposure to SBF is an indicator for the in vivo bioactivity of materials surface [136],[137].

Biomimetic materials are able to mimic the morphological and physicochemical features of biological apatite compounds, i.e. they are synthetic analogues of inorganic part of hard tissues [112],[117]. The biomimetic deposition is considered as an ideal method to produce calcium phosphate ceramics such as apatite coatings on titanium and its alloys for medical applications. It has also been proved that the chemical pretreatment in alkali solution can improve the bonding between the titanium substrate and calcium phosphate coatings fabricated by subsequent biomimetic deposition in simulated body fluid (SBF) [138],[139],[140],[141]. The biomimetic growth of apatite was also described on hydrogen-implanted silicon [142], polyvinyl alcohol (PVA) [143], TiO$_2$ nanotubes [144], alumina [145], zirconia ceramics (Y-TZP) [146], forsterite [147], akermanite [148], magnetite [149], glasses [150] and

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13 Titanium and its alloys are widely used as orthopedic and dental implant materials because of their high mechanical strength, low modulus and good corrosion resistance [139].
bioglasses [151],[152], cements [153], geopolymers [154], carbon nanotubes [155] and microspheres [156].

The ability to form apatite [150],[153] from supersaturated solution has been widely used to imply the bioactivity of an implant in vivo. However, the method itself may provide at best incomplete information, primarily because it is determined only by the solution supersaturation, irrespective of biological processes. The bone regeneration is triggered mainly by the vitality of osteoblasts and regulated by the expression of growth factors such as estrogen, parathyroid hormone and bone morphogenetic proteins, while ions or other species released from an implant may affect the expression of such growth factors and so the bone resorption or formation. The misinterpretation of the outcome of such tests must result in the misunderstanding of the true effects and behavior of materials intended for use in embedded biological applications. Moreover, SBF may not be able to mimic properly the physiological conditions because it is based on analytical concentrations and not on the activity of key components [157].

The methods of the preparation of collagen-hydroxyapatite composites include the production of composite gels, films, collagen-coated ceramics, ceramic-coated collagen matrices and composite scaffolds for bone substitutes and hard tissue repair via the following techniques [130],[158],[159]:

- **In vitro collagen mineralization**: the method is based on direct mineralization of a collagen substrate (film) through which calcium and phosphate ions diffuse into the fibrils or as phosphate-containing collagen solution (in situ precipitation [159]). **Fig. 20** introduces the experimental set-up for direct mineralization of a collagen sheet. The growth of HAP crystals with c-axis oriented along the collagen fibrils requires the pH in the range from 8 to 9 and the temperature of 40°C. These conditions promote the accumulation of calcium ions on the carboxyl groups of collagen molecules, leading to the nucleation of hydroxyapatite.

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**Fig. 20.** Simplified scheme of apparatus for the preparation of collagen-hydroxyapatite composite by in vitro collagen mineralization method: modified scheme according to Wahl and Czernuszka [130] for in situ collagen/HAP precipitation (a); modified scheme according to Wang and Liu [159]: pH meter (1), pH electrode (2), stirrer (3), peristaltic pump (4), hot-plate magnetic stirrer (5a) or stirrer (5b), thermocouple (6), hot plate (7) and peristaltic pump driven by pH controller (8) (b).

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These techniques usually continue with freeze-drying.
• **Thermally triggered assembly of HA/collagen gels**: the solution of calcium and phosphate ions is encapsulated within the liposomes and next inserted into the acidic suspension of collagen. After the injection into a skeletal defect, the increasing temperature due to body heat initiates the gelation process, which leads to fibrous network. The mineralization occurs after reaching the liposome’s transition temperature of 37°C.

• **Vacuum infiltration of collagen into a ceramic matrix**: ceramic scaffold is prepared by heating aqueous hydroxylapatite slurry containing poly(butyl methacrylate) (PBMA) spheres to high temperatures. The pyrolysis of PBMA particles leads to porous HAP green body. Pores in this matrix are then filled with collagen suspension under vacuum. The final composite was then freeze-dried to produce the microsponges within.

• **Enzymatic mineralization of collagen sheets**: is a method based on the cycle (Fig. 21) where collagen-containing alkaline (basic) phosphatase (ALP) is treated by aqueous solution of calcium ions and phosphate ester. The enzyme provides a reservoir for PO$_4^{3-}$ ions for calcium phosphate to crystallize and the mineralization occurs on coated area. The sample is then coated again with collagen suspension, air-dried and cross-linked with UV irradiation. Repeating this cycle results in multilayered composite sheets of calcium/phosphate and collagen.

![Fig. 21. The cycle of enzymatic mineralization of collagen sheets [130].](image)

• **Water-in-oil emulsion system**: purified collagen suspension mixed with HAP powders at the temperature of 4°C is next dispersed in olive oil and stirred at 37°C. The collagen aggregates and reconstitutes into the aqueous droplets. The addition of phosphate-buffered saline¹⁷ (PBS) leads to the gelation of Col-HAP microspheres (gel beads) of bone filler. The main disadvantages of this method are the problem with complete removing of the oil content from the composite and too low viscosity of the mixture.

• **Freeze-drying and critical point drying (CPD) scaffolds**: the ice crystals with collagen fibers at the interstices can be formed by freezing the suspension of collagen and HAP in water under controlled conditions. In the case when the freeze-drying is applied (temperature and pressure corresponding to CPD), ice crystals sublime to water vapor.

Under the condition of critical point,¹⁸ the density of liquid and gas phase converge and become identical (supercritical fluid) as well as the surface tension is negligible.

¹⁷ Aqueous solution of sodium phosphate and chloride, where the ion concentrations and osmolarity (osmotic concentration) correspond to human body. In some cases, potassium phosphate and chloride were used.

¹⁸ Critical point is defined by the value of three parameters including critical temperature, critical pressure and critical volume. Critical temperature is the highest temperature at which pure matter can exist as liquid.
• Col-HAP was cast into the mould and frozen, ice crystals replaced with ethanol, ethanol-liquid CO\textsubscript{2} exchanged and critical point dried to finally arrive with an exact porous replica of the original bone. Solid freeform fabrication techniques have recently been developed with artificial polymers and ceramic materials. These have the ability to change the pore interconnectivity, pore size and pore shape but have the disadvantage of not having the affinity of collagen to cell attachment. Another major advantage of Col-HAP scaffolds produced through the SFF method is the ability to control variables such as the control of external and internal structure, porosity and cross-linking [130].

The collagen-hydroxyapatite/pectin (Col-HA/pectin) composite was prepared in situ by the introduction of pectin, a kind of plant polysaccharide, into the collagen-hydroxyapatite composite. The structure of composite consisted of hydroxylapatite of low crystallinity with particles uniformly dispersed in organic materials. There is strong bonding interaction between HAP, collagen and pectin. The mechanical properties, water absorption, enzyme degradation and cytotoxicity indicate a potential use in bone replacement for the new composite [160].

Most mammalian biofluids are supersaturated with respect to the bone and tooth mineral hydroxylapatite. Nevertheless, the biofluids, which are in contact with soft tissues, especially like milk that must be stored for any length of time, should be highly stable with respect to calcium phosphate precipitation. In contrast, saliva and the extracellular matrix of hard tissues, especially near to the sites of mineralization or remineralization, are required not only to maintain the mineral phase with which they are in contact but also to deposit calcium phosphate in a highly controlled manner [161].

Collagen composites were also used as a scaffold for the repair of soft tissues. These materials provide analogous environment to extracellular matrix (ECM) and induced rate of synthesis or growth of new tissues. Several natural polymers as collagen, chitosan,\textsuperscript{19} gelatin\textsuperscript{20} and keratin\textsuperscript{21} possess the ability to induce the proliferation of cells and hence find their use as biomaterials for a wide range of biomedical applications. Among all biopolymers, collagen is a widely accepted material for the tissue engineering applications in the view of its low antigenicity, excellent biocompatibility and biodegradability [162].

The preparation of collagen-based biocomposite constructed from micro-crimped long collagen fiber bundles extracted from a soft coral embedded in alginate hydrogel matrix was described by Sharabi et al [163]. This biocomposite demonstrated the hyperelastic behavior similar to human native tissues.

\footnotesize{\textsuperscript{19} Chitosan is (1–4)-linked 2-amino-2-deoxy-b-glucan, a byproduct of N-deacetylation of chitin. It is a major constituent of crab and shrimp shells and of cuticles of insects.

\textsuperscript{20} Gelatin is a denatured form of collagen. Gelatin has low antigenicity and promotes the cell adhesion, differentiation and proliferation. Gelatin also possesses high cytocompatibility, which makes it a potential candidate as a biomaterial for various tissue engineering applications [162].

\textsuperscript{21} Keratin is a family of fibrous proteins, which is found abundantly in nature. It is the main constituent of hair, wool, nails, horns and hooves of mammals, birds and reptiles [162].}
10.11. Layers and biocoatings

The nanostructured coatings are expected to enhance the mechanical properties and improve the strength of bonding between coating and implant. These materials can also promote the deposition of calcium-containing minerals on their surface (bioactivity). Some typical examples of the research carried out in recent years, in the field of fabrication of nanostructured glass-ceramic coatings, utilize various techniques such as conventional enameling, sputtering, sol-gel processing, ion beam deposition, plasma spraying, electrophoretic deposition and pulsed laser deposition [164].

As was mentioned earlier, among the bioactive ceramics, the apatite/wollastonite (A/W) glass-ceramics, containing apatite and wollastonite crystals in the glassy matrix, has been largely studied because of good bioactivity and was used in some fields of medicine, especially in orthopedics and dentistry. However, medical applications of bioceramics are limited to non-load-bearing applications because of their poor mechanical properties. The solution of this problem can be the preparation of layers on the titanium alloy base material. Thermally sprayed layers by APS (atmospheric plasma spraying) on Ti-6Al-4V substrates combine good bioactivity of the bioceramics and good mechanical strength. The microstructure and the resulting properties were evaluated depending on the processing parameters and postprocessing thermal treatments. Thermal treatments decreased the bioactivity of the coatings, and after specific treatments, some bioactive materials were transformed into inert materials [67].

The biological performance of a porous apatite-mullite glass-ceramics, manufactured via the selective laser sintering (SLS) method, was investigated by Goodridge et al [165]. Laser-sintered A-M has shown similar cytotoxicity, in vitro bioactivity and in vivo results to cast A-M, indicating that the laser-sintering processing route does not alter the behavior of the material. The porous structure produced by SLS was seen by in vivo testing to be excellent for bone in-growth. However, the inability of the material to form apatite in vitro raised the concerns over the material’s ability to prove bioactive in vivo, and further assessment is required to confirm whether this material is not bioactive or whether it is just not particularly suited for the characterization through SBF testing [165].

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