Effect of Hydroxyapatite Nano-Particle on Properties of Modified Tricalcium Silicate Bone Cements

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Abstract

Solid-state reactions were used to synthesize bone-cements based on zinc modified tricalcium silicate (C₃S: Ca₃SiO₅). C₃S is the main self-setting and the fastest hydration constituent in conventional cements. Zinc oxide was added to replace part of CaO in order to stabilize the C₃S phase. 5 mole % ZnO addition (Zn-C₃S) reveals the best compressive strength that reaches 166 MPa, compared with those without ZnO, 100MPa. In vitro direct-contact-tests using primary cultured osteoblast are persuasive of negligible cytotoxicity. Hydroxyapatite nano-particles prepared through hydrothermal methods were added as strengthener and resulted in the compressive strength increment of the Zn-C₃S up to 175 MPa at an HA addition amount 7.5 wt%. The setting mechanism of C₃S and Zn-C₃S seems to be the same except varied gel properties that extend the fluidity of the slurry. These cements show remarkable self-setting properties and comparable strength with natural bones hence are potential in bony and dental restorations.

Keywords: Bone-cement, Tricalcium Silicate, Self-setting, Zinc oxide, Hydroxyapatite, Nano-particles

Introduction

Acrylic cements have been widely used in orthopedics and dentistry as fixation agents and fillers [1,2]. However, they have some disadvantages such as thermal necrosis of cells, shrinkage during polymerization, monomer releasing, and mechanical mismatch with the implant site, among others [3].

It has been known that glasses and derivatives based on the CaO-SiO₂ system show excellent biocompatibility and form apatite layer on their surface in vivo as well as in vitro [4-5]. Tricalcium silicate (Ca₃SiO₅, C₃S in short) is the main self-setting and the fastest hydration constituent in cements. C₃S swells during setting and gets very high strength after being set [6]. From the phase diagram, it is manifested that C₃S is a single phase at very high temperature, but will decompose into dicalcium silicate (C₂S, Ca₂SiO₄) and calcium oxide (CaO) at temperatures below 1250 oC. Therefore, it’s difficult to obtain pure C₃S phase by solid-state reaction. It is reported that some transition metals are able to stabilize the C₃S phase [7]. Hydroxyapatite (HA) being the main inorganic content of human bone gets excellent biocompatibility and osteoconductivity, and accelerates the restoration and growth of injured bone [8,9]. In this study, our aim was to synthesize quality improved C₃S with high phase purity and mechanical properties, aiming to apply the aforesaid characteristics in bony and dental restorations. We chose zinc as a phase-stabilizing additive as well as making use of the special bio-effects of zinc. Zinc was excepted to substitute calcium in the crystal structure forming (Ca₁₋ₓZnx)₃SiO₅. The phase purity of developed materials was analyzed by XRD. Besides, hydroxyapatite nano-particle was added as a strengthener aiming also to benefit the biocompatibility. The setting time of developed cements were evaluated by the Gillmore needle method [10]. Mechanical properties were measured by an Instron machine. The pH value and in vitro biocompatibility were also tested.

Materials and Methods

1. Material preparation and analyses

The precursory powders (CaCO₃+ZnO):SiO₂= 3:1 were mixed by ball milling the weighed constituents. Molar percentage ZnO to replace CaO was from 5 to 50%. After heating and soaking the mixtures at 1400 °C, they were quenched in liquid nitrogen. The products were ground to smaller than 37 µm (400mesh sieved). Powders were mixed with de-ionized water (in weight%) by firstly rapid shaking.

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(3000 rpm) and self-setting. The slurry was filled into a stainless mold (10mm in diameter), pressed by a pressure 8.3 kgf/cm², incubated at 37°C under saturated steam for curing.

2. Preparation of hydroxyapatite nano-particles through a hydrothermal route

Hydroxyapatite nano-particle was prepared by an autoclave hydrothermal process. Reagent grade Ca(H₂PO₄)₂·H₂O and Ca(OH)₂ powders were mixed by a conventional ball-milling technique using zirconia balls immersed in ethanol for 1h. The molar ratio of Ca(H₂PO₄)₂·H₂O to Ca(OH)₂ was 3:7, which is based on the hydroxyapatite (HA) stoichiometric composition. After mixing, the slurry was dried by vacuum extractor. The HA powder was synthesized by heating mixtures of powders in 100~200 g batch with 2000 ml distilled water under 121 °C for 1~2 h in a conventional autoclave used for sterilization. The reaction product was filtered and washed several times with DI water for the complete removal of PO₄³⁻ and Ca²⁺ ions, then frozen dried.

3. Cells and cell culture

The primary cultural osteoblast was used for in vitro biocompatibility evaluation. Calvaria from Wistar rat (born in three days) were excised aseptically. Soft tissues on the surface were removed carefully. After rinsing the calvaria with PBS, the calvaria were minced and incubated in collagenase (1mg/ml PBS, Sigma Co.) at 37 °C for 10 min. The isolated osteoblasts were centrifuged for 5min at 1500 rpm and then cultured in DMEM. The medium was refreshed every two days. Each cement disc was gamma ray irradiation sterilized (10 kGy) before biocompatibility tests.

4. Characterization methods

XRD patterns were obtained from samples using a wide-angle x-ray powder diffractometer (model D/max-IIB, Rigaku Co., Tokyo, Japan). X-ray radiation CuKα₁ (λ = 1.5405Å) was set at 30 kV and 20 mA. Detailed peak features of the diffraction angles, 2θ, ranging from 24° to 38° were obtained by a 0-20 scan rate of 0.4° per minute. The crystalline phases were identified using JCPDS files (HA: 24-0033 and Si: 03-0529). Setting properties were evaluated by the Gillmore needle method. When the needle can no longer penetrate the surface of cement it is meant that setting time for proper strength is reached. The compressive strength of the cylinder was measured using an Instron 4505 Universal testing instrument.

Results and Discussion

In our past researches [11], through the synthetic procedures, C₃S phase with substantial amount of free CaO can be obtained routinely for unmodified 3CaO·SiO₂ compositions. And the amount of free CaO phase obviously decreases with increasing ZnO addition up to 7.5 mol%. The mechanical strength of the samples increases with curing time. After curing for 24 hours, the synthesized XRD-pure C₃S shows a setting time of 56.3 min, a compressive strength around 100MPa. Among those ZnO modified products, the (Caₙ₋₂Zn₇)₀SiO₅ samples have the least CaO impurity and show a compressive strength around 128MPa, yet with a worse handling property. The (Ca₉Zn₂)₀SiO₅ (Zn-C₃S) samples show the best mechanical property with a setting time 54 min and the compressive strength reaches 166MPa. More ZnO addition leads to a lengthened setting time and a decreased compressive strength. Besides, the un-modified C₃S slurry loses its fluidity in one minute, while that of the Zn-C₃S slurry sample maintains fluidity until 3 minutes, which may be more suitable in clinical usage. Samples of pure C₃S and Zn-C₃S were selected for biocompatibility evaluation. After direct contact tests, the samples didn’t show obvious cytotoxicity (Figure 1).

Because the calcium silicate bone cement was set with water, different weight ratios between water to cement powder lead to different working time and setting time. Figure 2 was the setting time of C₃S and Zn-C₃S cements with different amounts of water addition. It’s obviously that setting was positively dependent on water content. And when the water/powder weight ratio was 0.2, the C₃S and Zn-C₃S cements had the shortest setting time, however, the resulted solid got a loose structure. So the water/powder ratio was optimally set as 0.4 (200mg water versus 500mg powder) and applied for further studies. Figure 2 also tells that zinc addition extends the working time and setting time, and also provides better fluidity that was good for clinical usage. It is manifest that addition of zinc oxide varies the property of gel formed upon mixing with water as compared with that of un-modified C₃S. The modified gel would extend the time to the formation

![Figure 1.](image) (a) Pure C₃S sample shows no cytotoxicity; (b) Zn-C₃S sample shows no cytotoxicity.
Zn-modified 3CaO-SiO$_2$ with Hydroxyapatite Nano-Particle

Figure 2. Setting time of C$_3$S and Zn-C$_3$S cements with different water addition

Figure 3. (a) Setting time of Zn-C$_3$S with different amounts of HA addition. It’s obvious that setting time is prolonged upon HA addition less than 10 wt%, while decreased above 10 wt%. It’s suggested that when the HA is of small amount, the added HA will hinder the hydration of Zn-C$_3$S that otherwise self-sets as usual. While in case of high HA concentration, the HA nano-particles tend to incorporate into the hydration mechanism leading to a loose hydrated structure and an uneven local setting. The setting time of C$_3$S+HA cement shows a turning point at 5 wt% HA addition (Figure 3(b)). The difference between setting time of HA added Zn-C$_3$S and C$_3$S was due to fluidity as stated earlier.

Figure 4 presents the relationship between compressive strength and curing time of the cements. After curing for twenty days, Zn-C$_3$S has higher compressive than that of C$_3$S. The mechanisms are elucidated due to the following steps and reasons:

1. Zn-C$_3$S has better fluidity that is beneficial for early packing of the as hydrated components.
2. Zn$^{2+}$ inserts into the lattice of final set mass.
3. From the analyses of fine structure of the final products, Zn-C$_3$S shows more continuous and compact structure. There existed more boundaries and grains in final product of C$_3$S cement.

Besides, these two cements show similar time dependent curves of compressive strength during curing, it implies that the two cements have the same strengthening mechanism.

Figure 5(a) shows the effects on compressive strength of C$_3$S cements added with different HA: 5wt%, 7.5wt%, 10wt%, respectively. The results reveal that addition of HA decreases the final compressive strength. It’s due to the reason that added HA nano-particles might not be well separated and might react with the gel locally. Nevertheless the 7.5 wt% added one has relatively high strength than others. Figure 5(b) is same measurement upon the Zn-C$_3$S cement. For HA added Zn-C$_3$S, the compressive strength is raised, the best value as high as 175MPa is possible at an HA amount of 7.5 wt%. The reason
is due to the good fluidity that helps dispersion of HA nano-particles, so that HA could be competent for the strengthening. And the 7.5 wt% HA added one reaches the highest strength in around 5 days that are faster than the un-modified one.

Conclusions

In this study, a series of ZnO modified 3CaO-SiO$_2$ bone cements (C$_3$S) was successfully developed. The results showed that addition of up to 7.5 mol% ZnO to replace CaO in the C$_3$S composition would stabilize the C$_3$S phase, increase the phase purity, lengthen the setting time and also increase the final compressive strength. The (Ca$_{0.95}$Zn$_{0.05}$)$_3$SiO$_5$ samples show the best mechanical property with a compressive strength 166MPa. In vitro cell culture tests showed that the studied samples show negligible cytotoxicity. Zn-C$_3$S is concluded to be more suitable for clinical injecting. Besides, for Zn-C$_3$S added with 7.5 wt% HA, its compressive strength is further improved and setting time is lengthened. Concerning clinical usages, Zn-C$_3$S is more suitable and potential. The developed (C,Z)$_3$S bone cements in this study show promising applications in bony and dental restorations.

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