INTRODUCTION

Secondary caries at the tooth-restoration margins is the most-frequent reason for replacement of restorations (Sakaguchi, 2005; Sarrett, 2005). Replacement dentistry accounts for 70% of all operative work and costs $5 billion/year in the U.S. (Jokstad et al., 2001; CDC, 2005). Several calcium-phosphate phases are regarded as biological precursors that form initially and then transform to apatites (LeGeros, 1991). Recent studies showed that methacrylate-based composites containing calcium-phosphate fillers released calcium (Ca) and phosphate (PO4) ions to supersaturated levels for apatite precipitation, and effectively remineralized tooth lesions in vitro (Skrtic et al., 1996a, 2000; Dickens et al., 2003).

However, the Ca-PO4 fillers did not reinforce the resin as do glass fillers (Söderholm et al., 1984; Goldberg et al., 1994; Bayne et al., 1998; Ferracane et al., 1998; Drummond and Bapna, 2003). Ca-PO4 composites had flexural strengths half that of unfilled resin (Skrtic et al., 1996b). Such low strengths were "inadequate to make these composites acceptable as bulk restoratives" (Skrtic et al., 2000).

Whiskers were used as fillers to reinforce dental composites (Xu, 1999). Silica nano-particles were fused onto the whiskers to facilitate silanization and enhance retention in the resin. These composites possessed strength and toughness nearly two-fold greater than those of several commercial composites (Xu et al., 2002a).

Calcium phosphate nano-particles were recently developed and incorporated into resins (Chow et al., 2004; Xu et al., 2006). A recent study investigated the effects of different resins/cure conditions with a single filler level (Xu et al., 2006). In the present study, two hypotheses were tested: (1) DCPA-whisker composites with filler levels from 0-75% would possess strengths matching/exceeding those of a commercial non-releasing, stress-bearing composite; and (2) the Ca-PO4 release would be proportional to the DCPA nano-particle filler level in the resin. The first hypothesis was tested because low filler levels could be envisioned for Ca-PO4-releasing sealant applications, medium filler levels for Ca-PO4 flowable composites, and high filler levels for stress-bearing and caries-inhibiting restorations. The purpose of the second hypothesis was to establish a relationship between release and volume fraction, to guide the tailoring/processing of composites.

MATERIALS & METHODS

ABSTRACT

Nano-particles of dicalcium phosphate anhydrous (DCPA) were synthesized for the first time. The objectives of this study were to incorporate DCPA nano-particles into resin for Ca-PO4 release to combat dental caries, and to investigate the filler level effects. Nano-DCPA and nano-silica-fused silicon nitride whiskers at a 1:1 ratio were used at filler mass fractions of 0-75%. The flexural strengths in MPa (mean ± SD; n = 6) of DCPA-whisker composites ranged from (106 ± 39) at 0% fillers to (114 ± 23) at 75% fillers, similar to (112 ± 22) of a non-releasing composite (TPH) (p > 0.1). The composite with 75% fillers in a NaCl solution (133 mmol/L, pH = 7.4, 37°C) yielded a Ca concentration of (0.65 ± 0.02) mmol/L and PO4 of (2.92 ± 0.07) mmol/L. Relationships were established between ion-release and DCPA volume fraction VDCPA: Ca = 4.46 VDCPA1.6, and PO4 = 66.9 VDCPA2.6. Nano-DCPA-whisker composites had high strength and released high levels of Ca-PO4 requisite for remineralization. These new nano-composites could provide the needed combination of stress-bearing and caries-inhibiting capabilities.

KEY WORDS: dental composite, nano-particles, whisker reinforcement, tooth caries inhibition, Ca and PO4 ion release.

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Silicon-nitride whiskers (β-Si₃N₄, UBE, New York, NY, USA) with diameters of 0.1-2 μm (mean = 0.4 μm) and lengths of 2-30 μm (mean = 5 μm) were mixed with silica (Aerosil-OX50, Degussa, Ridgefield, NJ, USA; diameter = 40 nm) at a whisks: silica mass ratio of 5:1. The mixture was heated in a furnace at 300°C for 30 min to fuse the silica onto the whiskers. The powder was silanized with 4% 3-methacryloxypropyltrimethoxysilane and 2% N-propylamine (mass fractions) (Xu, 1999).

A monomer consisting of 48.975% Bis-GMA (bisphenol glycidyl methacrylate), 48.975% TEGDMA (triethylene glycol dimethacrylate), 0.050% 2,6-di-tert-butyl-4-methylphenol, and 2.000% benzoyl peroxide formed part I, the initiator, of a two-part chemically activated resin. Part II, the accelerator resin, consisted of 49.5% Bis-GMA, 49.5% TEGDMA, and 1.0% N,N-dihydroxyethyl-p-toluidine.

The fillers consisted of nano-silica fused whiskers and nano-DCPA at a DCPA:whisker mass ratio of 1:1 (Xu et al., 2006). The (DCPA+whiskers)/(DCPA+whiskers+resin) mass fractions were: 0%, 30%, 40%, 50%, 60%, 65%, 70%, and 75%. Filler levels ≥ 80% resulted in a dry paste. Equal masses of parts I and II were mixed and filled into 2x2x25 mm³ molds. The specimens were incubated at 37°C and 50% humidity for 24 hrs prior to being tested.

A hybrid composite (TRIP, Caulk/Dentsply, Milford, DE, USA) was used as a non-releasing control. It consisted of 0.8-μm silicate fillers at 78% level in a urethane-modified Bis-GMA-TEGDMA resin. The specimens were photo-cured (Triad-2000, Dentsply, York, PA, USA).

**Flexural Strength and Nano-indentation**

Flexural strength was measured by a three-point test with a 10-mm span at a crosshead speed of 1 mm/min on a Universal Testing Machine (5500R, MTS, Cary, NC, USA) at approximately 25°C and 50% relative humidity. Nano-indentation (Nano-Instruments, Knoxville, TN, USA) was used to measure elastic modulus and hardness at a peak-load of 1 N (Xu et al., 2002a).

**Ca and PO₄ Release**

A NaCl solution (133 mmol/L) buffered with 50 mmol/L HEPES (pH = 7.4; 37°C) was used. Following a previous study (Xu et al., 2006), we immersed 3 specimens of approximately 2x2x12 mm³ in 50 mL of solution, yielding specimen volume/solution = 2.9 mm³/mL. This was similar to the specimen volume/solution of approximately 3.0 mm³/mL in a previous study (Skrtic et al., 1996b). The concentrations of Ca and PO₄ released from the specimens were measured vs. immersion time (in days): 1, 2, 4, 7, 14, 21, 28, 35, 42, 49, and 56. At each time period, aliquots of 0.5 mL were removed and replaced by fresh solution. The aliquots were analyzed with a spectrophotometer (DMS-80 UV-visible, Varian, Palo Alto, CA, USA) according to established standards and calibration methods (Skrtic et al., 1996b; Dickens et al., 2003).

**Potential Diagram**

The remineralization potential of a Ca- and PO₄-releasing composite can be described via a potential diagram (Dickens et al., 2003). For a solution saturated with respect to hydroxyapatite \(\text{Ca}_{10}(\text{PO}_4)_{6}(\text{OH})_2\), the solubility constant relationship, \(K_{SP} = (\text{Ca}^{2+})(\text{OH}^-)^2 = (\text{H}^+)3(\text{PO}_4)^3-\), applies (Chow and Brown, 1984). Rearranging the equation leads to:

\[
K_{SP} = \left[\left(\text{Ca}^{2+}\right)\left(\text{OH}^-\right)^2\right]^{10} \times \left[\left(\text{H}^+\right)^{3}\left(\text{PO}_4\right)^{3-}\right]^6 / K_w^9
\]

where \(K_w = (\text{H}^+)(\text{OH}^-)\) is the dissociation constant of water. Taking log of both sides and rearranging the equation lead to:

\[
\log\left[(\text{H}^+)3(\text{PO}_4)^3-\right] = -1.67x\log[\text{Ca}^{2+}(\text{OH}^-)^2] + K
\]

where \(K = (1/6)(\log K_{SP} + 9\log K_w)\) is a constant. Eq. (2) shows that, for a solution saturated with respect to hydroxyapatite, the logarithms of the activities of \(\text{H}_3\text{PO}_4\) and \(\text{Ca(OH)}_2\) are linearly related. The activities \((\text{H}^+)3(\text{PO}_4)^3-\) and \((\text{Ca}^{2+})3(\text{OH})^2\) for a solution can be calculated from the pH, the measured Ca-P0₄ concentrations, and the ionic strength. Hence, a solution can be represented as a point in the potential diagram. Solutions located to the left of the hydroxyapatite line (e.g., Fig. 8, Dickens et al., 2003) are undersaturated, and those to the right are supersaturated, with respect to hydroxyapatite.

The extent of this supersaturation can be quantified via the saturation ratio. The saturation index is (Margolis et al., 1999):

\[
I = (\log[IAP] - \log[K_{SP}])/q
\]

where \(q\) is the number of ions in the ion activity product expression (for hydroxyapatite, \(q = 10 + 6 + 2 = 18\)). \(IAP = (\text{Ca}^{2+})^{10}(\text{PO}_4)^{3-}\) is the ion activity product, which can be calculated from the measured concentrations with specific software (Chemist, Micromath Research, St. Louis, MO, USA). The saturation ratio

\[
SR = 10^{(\text{saturation index})} = 10^I
\]

where SR < 1 means that the solution is undersaturated, and SR > 1 means that the solution is supersaturated, with respect to hydroxyapatite (Margolis et al., 1999).

Furthermore, the Gibbs free energy can be used to quantify the thermodynamic driving force for remineralization (Dickens et al., 2003):\[\Delta G^0 = -2.303(R/T)\log(IAP/K_{SP})\]

where \(R\) is the ideal gas constant, and \(T\) is absolute temperature.

We performed one- and two-way ANOVA to detect the significant effects of the variables. We used Tukey’s multiple-comparison test to compare the measured data at a p value of 0.05.

**RESULTS**

**Mechanical Properties**

Increasing the filler level did not significantly change the strength (\(p > 0.1\)) (Fig. 1A). Flexural strength in MPa (mean ± SD; \(n = 6\)) at 60% and 75% fillers were (117 ± 17) and (114 ± 23), respectively, not significantly different from (106 ± 39) of the unfilled resin and (112 ± 22) of the hybrid control (\(p > 0.1\)).

Elastic modulus (Fig. 1B) and hardness (Fig. 1C) increased with filler level. Modulus in GPa (mean ± SD; \(n = 10\)) at 75% fillers was (14.9 ± 0.7), higher than (11.7 ± 0.4) for the hybrid control and (3.9 ± 0.1) for the unfilled resin (\(p < 0.05\)).

**Ca and PO₄ Release**

The release increased with time and then started to plateau (Figs. 2A, 2B). At 56 days, the Ca concentration in mmol/L (mean ± SD; \(n = 3\)) was (0.65 ± 0.02) with 75% fillers, significantly higher than (0.59 ± 0.02) with 70% fillers, and (0.39 ± 0.03) with 65% fillers (\(p < 0.05\)). The corresponding PO₄ concentrations were (2.29 ± 0.07), (1.92 ± 0.14), and (1.26
± 0.09), significantly different from each other (p < 0.05).

Potential Diagram
In the chemical potential diagram (Fig. 3A), the solution of the composite with 30% fillers was undersaturated, that with 40-50% fillers was supersaturated after 2 days, and those of all other composites were supersaturated from 1 day, with respect to hydroxyapatite. The saturation ratio increased with increasing filler levels (Fig. 3B). The larger the negative value of $\Delta \tilde{G}$ (Fig. 3C), the lower the energy state for hydroxyapatite precipitation, and the higher the remineralization potential (Dickens et al., 2003).

DISCUSSION
Synergistic Effects of Nano-particles/Whiskers
Previous studies have measured fluoride release from dental
materials (Geurtsen et al., 1999; Anusavice et al., 2005). Other studies developed Ca-PO₄ composites with flexural strengths of about 55 MPa (Skrtic et al., 1996b). This led to the observation that “all the amorphous calcium phosphate fillers yielded polymerized materials weaker than unfilled polymers” (Skrtic et al., 1996b). Another composite, with micron-sized DCPA, had a flexural strength of 40-50 MPa (Dickens et al., 2004). In the present study, the nano-DCPA-whisker composites had flexural strengths of about 110 MPa. The DCPA-whisker composite at 75% fillers had an elastic modulus of 14.9 GPa. It was lower than the 18 GPa of dentin, but higher than the 11.7 GPa of the commercial stress-bearing, non-releasing composite control.

Previous Ca-PO₄ composites released PO₄ to concentrations of 0.1-0.7 mmol/L, and Ca to 0.3-1.0 mmol/L, measured with a similar method (Skrtic et al., 1996b; Dickens et al., 2003). These composites remineralized tooth lesions in vitro (Skrtic et al., 1996a; Dickens et al., 2003). The nano-DCPA-whisker composites released PO₄ with concentrations up to 2.2 mmol/L, and Ca up to 0.65 mmol/L (at 75% total fillers), even when half of the fillers were non-releasing whiskers. This was likely because the DCPA nano-particles had a high surface area, A = 18.6 m²/g (Xu et al., 2006).

In a previous study (Dickens et al., 2003), the DCPA particle size, d, was 1.1 μm and the TTCP (tetracalcium phosphate) particle size was 16 μm. The density, ρ, is 2.89 g/cm³ for DCPA and 3.07 g/cm³ for TTCP. Hence, A = 6/(pd) = 1.9 m²/g for DCPA, and A = 0.12 m²/g for TTCP. These traditional particles had surface areas 1-2 orders of magnitude less than the new DCPA nano-particles. As a result, these traditional composites needed to be fully filled with Ca-PO₄ fillers to have significant release. Replacing part of these Ca-PO₄ fillers with reinforcing fillers would substantially reduce the release. Even if only 10% of the ACP fillers had been replaced by reinforcing fillers, the Ca-PO₄ release would have been decreased from about 0.75 to only 0.1 mmol/L (Skrtic et al., 1996b). Therefore, there was little room left in traditional Ca-PO₄ composites for reinforcement fillers without diminishing the ion release capability.

In contrast, with nano-DCPA, high release could be achieved with less filler, thus making room available in the resin for reinforcement fillers. This synergistic releasing-filler/reinforcing-filler approach helped achieve a flexural strength of 110 MPa for the Ca-PO₄-releasing composites, matching that of a commercial stress-bearing, non-releasing composite.

**Effect of Nano-DCPA Volume Fraction**

The masses used in making the composites and the density were used to calculate the volume fraction of DCPA in the composite. The density of DCPA, d_{DCPA}, is 2.89 g/cm³. The density of the unfilled resin was measured to be d_{resin} = (1.19 ± 0.04) g/cm³. The density of silicon nitride whiskers is 3.34 g/cm³ (manufacturer’s data), and d_{fumed-silica} = 2 g/cm³. Hence, d_{silica-fused-whisker} = 2.97 g/cm³ at a whisker:silica ratio of 5:1. At total filler mass fractions of 30%, 40%, 50%, 60%, 65%, 70%, and 75%, the volume fraction V_{DCPA} was calculated to be 0.075, 0.108, 0.147, 0.192, 0.218, 0.247, and 0.279, respectively.

There appear to be two main factors influencing Ca-PO₄ release: (1) the amount of the source of release, V_{DCPA}; with the amount of release expected to increase with increasing V_{DCPA}; and (2) the resin polymerization conversion. Increasing the filler level usually decreases the polymerization conversion (Xu, 1999), because a higher concentration of air in the heavily filled composite may adversely affect the conversion. In addition, the fillers may partially absorb the heat of polymerization, thereby moderating the exotherm of polymerization. Therefore, with higher V_{DCPA} in the composite, there is not only more DCPA for release, but also the diffusion

**Figure 3. Remineralization potential.** (A) Diagram demonstrating the potential of DCPA-whisker composites to form hydroxyapatite. The straight line in the middle represents the hydroxyapatite solubility isotherm line. The line at the upper right corner represents the solubility isotherm for DCPD (dicalcium phosphate dihydrate, CaHPO₄·2H₂O). The dots represent log([Ca²⁺][OH⁻]²) vs. log([H⁺][PO₄³⁻]) from the measured Ca and PO₄ concentrations. (B) Degree of saturation SR < 1 means that the solution is undersaturated, and SR > 1 means supersaturated, with respect to hydroxyapatite. (C) Thermodynamic driving force for remineralization. The larger the negative value of the Gibbs free energy ΔG⁰, the lower the energy state for hydroxyapatite precipitation, and the higher the remineralization potential. Values near the right axis are filler level mass fractions.
of water and ions through the resin may be somewhat enhanced, due to the decreased polymerization conversion. If only factor (1) had been operative, the relationship between $V_{DCPA}$ and Ca-Po$_4$ release might have been simply linear. However, these two factors may both be operative. Hence, the release may increase with increasing $V_{DCPA}$ at a rate faster than linear. Based on these reasons, the following empirical relationships are proposed:

$$Ca = k_1 V_{DCPA}^{\alpha}$$  \hspace{1cm} (6)

$$Po_4 = k_2 V_{DCPA}^{\beta}$$  \hspace{1cm} (7)

$$SR = k_3 V_{DCPA}^{\gamma}$$  \hspace{1cm} (8)

where Ca and Po$_4$ (mmol/L) are concentrations, and $k$ and $\alpha-\gamma$ are coefficients. Fitting the above equations to the measured data (at 56 days) yielded the equations in Fig. 4.

Regarding potential applications, nano-DCPA-whisker composites with 30-50% fillers may be suitable for use as Ca-Po$_4$-releasing tooth cavity liners, adhesives, and pit-and-fissure sealants. Flowable DCPA-whisker composites with 50-60% fillers may be used as crown cements and orthodontic bracket cements, and to repair defective margins. Composites with 70-75% fillers may be useful in stress-bearing and caries-inhibiting restorations. The Ca-Po$_4$ release from the DCPA-whisker composites matched/exceeded those of previous composites known to remineralize tooth lesions (Skrtic et al., 1996a; Dickens et al., 2003). Further research is needed to investigate their potential applications.

The dimensional change of the composite with time is another issue that needs to be investigated. Previous studies on a Ca- and F-releasing composite (Ariston pHc) for buffering the local pH in plaque retention areas showed considerable enamel cracks after 2 yrs, which were related to the water-expansion of the restorations (Braun et al., 2001; van Dijken, 2002; Krämer et al., 2005). Ariston was completely covered with cracks after 24 mos (Frankenberger et al., 2005). It exhibited the highest dimensional expansion among the materials immersed for 2 mos (Martin et al., 2003), and a high wear-rate (Manhart et al., 2000). These failure phenomena may be related to its hydrophilic monomer (Table I, Manhart et al., 2000) and the formation of calcium carbonates with an expanding effect (van Dijken, 2002). One advantage of the new nano-composite was that it released high levels of Ca-Po$_4$ without the use of a hydrophilic monomer. The Bis-GMA/TEGDMA resin for the nano-composite was similar to the resin in previous composites, showing no significant degradation in thermal-cycling and water-aging for 2 yrs (Xu et al., 2002b; Xu, 2003). However, studies are needed to evaluate the long-term performance of the new nano-composites \textit{in vitro} and \textit{in vivo}.

In summary, we developed novel nano-composites using DCPA nano-particles with Ca-Po$_4$ release to combat dental caries. The effects of nano-DCPA filler level were systematically investigated and correlated with Ca-Po$_4$ release for the first time. The Ca-Po$_4$ release from DCPA-whisker composites matched/exceeded those of previous composites known to remineralize tooth lesions, while the strengths of the DCPA-whisker composites were two-fold those of previous Ca-Po$_4$ composites. Relationships between Ca-Po$_4$ release and nano-DCPA volume fraction, $V_{DCPA}$, were established: Ca = $4.46 V_{DCPA}^{1.6}$, and Po$_4$ = $66.9 V_{DCPA}^{2.6}$. This suggests that the filler volume fraction is a key factor, and the release increases.
with \( V_{DCPA} \) at a rate greater than linear. The new nano-composites, with substantial Ca and \( PO_4 \) release, possessed mechanical properties matching those of a commercial stress-bearing, non-releasing composite. Hence, the nano-DCPA-whisker composites may have both stress-bearing and caries-inhibiting capabilities, a combination not yet available in current dental materials.

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Certain commercial materials and equipment are identified to specify the experimental procedure. This does not imply recommendation or endorsement by NIST or ADAF, or that the material or equipment identified is necessarily the best available for the purpose. The standard uncertainty of the flexural strength measurement was estimated to be 1%. The standard uncertainty for the Ca and \( PO_4 \) release measurements was estimated to be 3%. Unless otherwise specified in the paper, one standard deviation was used as the estimated standard uncertainty of the measurements.

**REFERENCES**


