RELATIONSHIP BETWEEN THE DEGREE OF CONVERSION, SOLUBILITY AND SALIVARY SORPTION OF A HYBRID AND A NANOFILLED RESIN COMPOSITE: INFLUENCE OF THE LIGHT-ACTIVATION MODE

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Received: November 27, 2007 - Modification: December 03, 2007 - Accepted: January 18, 2008

ABSTRACT

This study analyzed the relationship between the degree of conversion (DC), solubility, and salivary sorption of a hybrid (Filtek P 60) and a nanofilled resin composite (Filtek Supreme), and evaluated the influence of the light-activation mode on these properties. Two light-activation modes were used: Conventional (C; 850 mW/cm² for 20 s) and Soft-start (SS; 100-1,000 mW/cm² for 10 s + 1,000 mW/cm² for 10 s). The DC (%) was evaluated by FT-Raman spectroscopy. The solubility and salivary sorption were measured after immersion in artificial saliva for 7 days. Data were analyzed by ANOVA and Student-Newman-Keuls’ test and linear regression analysis (α = 0.05). The DC varied from 50.52% (nanofilled composite) to 57.15% (hybrid composite), and was influenced by the light-activation mode: C > SS. The solubility (0.45 µg/mm³) and salivary sorption (8.04 µg/mm³) of the nanofilled composite were greater than those of the hybrid composite (0.40 µg/mm³ / 6.87 µg/mm³), and were influenced by the light-activation mode: SS > C. Correlation was found between DC and solubility (r = -0.89, p<0.05), as well as between solubility and salivary sorption (r = 0.95). These findings suggest that nanofilled composites may present higher degradation in the oral environment than hybrid ones. Soft-start light-activation mode may increase the solubility of resin composites.

Key words: Composite resins. Degree of conversion. Solubility. Salivary sorption. Light-activation mode.

INTRODUCTION

During the last few years, resin composites have been classified according to their filler particle size, as hybrid, microhybrid and microfilled. More recently, however, with the introduction of nanotechnology in dentistry¹⁸, a new class of resin composites, the so-called nanocomposites, is available to clinicians. Filtek Supreme (3M ESPE) is a typical nanofilled composite that has a filler particle system that combines silica nanofillers with a primary particle size of 20 or 75 nm and zirconia-silica nanoclusters, 0.6-1.4 µm in diameter. Its filler load is 59.5 vol%, which is close to several hybrid and microhybrid composites.¹⁵ Several published studies have shown that this resin composite has good mechanical properties³,²¹. However, there is a lack of data about the sorption and solubility phenomena of this new class of restorative materials.

Resistance to degradation in the oral environment is essential to the longevity of resin composite restorations. From this point of view, it is crucial to understand the solubility and salivary sorption phenomena, in order to predict the behavior of resin composite restorations. Water sorption by resin composites is a diffusion-controlled process that may cause chemical degradation of the material, leading to several drawbacks, such as filler-polymeric matrix debonding²⁶ and residual monomer release. This process can decrease the mechanical properties of the material²² and reduce the longevity of resin composite restorations. The solubility of resin composites is reflected by the amount of leachable unreacted monomers. During polymerization, these unreacted monomers are either trapped in micropores between the polymer chains and adsorbed to the surrounding network, or trapped in nanopores. The monomers in the nanopores are more prone to leaching out from the material than those inside the micropores²⁴.

Several factors, such as the polymeric matrix
composition\(^{24}\), filler particle type and content\(^{27}\), and the DC reached after the polymerization reaction\(^{24}\) can influence the solubility and sorption behavior of resin composites. Some light-activation modes can be used to start the polymerization reaction of resin composites.\(^{20}\) Modes that use a high initial irradiance provide a higher DC. However, it has been shown that this activation mode may induce a high shrinkage stress in the material, thus increasing marginal gap formation at the cavity wall-resin composite interface.\(^{25}\) Soft-start light-activation modes have been introduced in attempt to overcome this drawback in composite polymerization. These polymerization modes, in which the composite is first submitted to a low irradiance followed by higher irradiance, allow shrinkage stress relief by polymer chain relaxation\(^{15,25}\). It has been demonstrated that these light-activation modes do not reduce the DC of dental composites or dental polymers\(^{2,15}\).

The solubility and sorption of resin composites and dimethacrylate-polymeric matrices have been extensively investigated\(^{5,11,12,17,24,26}\). Few of these studies, however, have analyzed the material’s behavior when immersed in artificial saliva\(^{11,26}\), which is a more compatible model with the oral environment conditions and would therefore supply a more realistic knowledge of these phenomena. Indeed, the effect of saliva on composites can be more deleterious than that of water\(^{26}\).

The purposes of this in vitro study were to analyze the correlation between the DC, solubility and salivary sorption of a hybrid and a nanofilled composite with similar polymeric matrices, and to evaluate the influence of the light-activation mode on these properties. The research hypotheses of the study were: (1) the light-activation mode does not influence the DC of resin composites, (2) nanofilled composite presents higher solubility and salivary sorption than the hybrid composite, and (3) there is a positive correlation between DC and solubility, as well as between DC and salivary sorption.

**MATERIAL AND METHODS**

Two commercially available resin composites with different types of filler particles were selected: a hybrid (Filtek P 60) and a nanofilled composite (Filtek Supreme – Su). Both materials have the same polymeric matrix (Bis-GMA, Bis-EMA, TEGDMA and UDMA). Their compositions are described in Table 1.

A quartz-tungsten-halogen unit (Optilux 501; Kerr, Danbury, CT, USA) was used to photoactivate the materials. Two light-activation modes were used: Conventional (C) – 850 mW/cm\(^2\) for 10 s (17 J/cm\(^2\)); and Soft-start (SS) – from 100 to 1,000 mW/cm\(^2\) for 10 s and 1,000 for 20 s (≈17 J/cm\(^2\)). The radiant exposure was calculated as the product of the irradiance of the curing unit, as measured with a curing radiometer (model 100; Demetron Inc. Danbury, CT, USA) and the time of irradiation. For the Soft-start mode, the radiant exposure was obtained by the sum of mean irradiance over the first 10 s multiplied by 10 s with 10 J/cm\(^2\), corresponding to the radiant exposure in the last 10 s of light exposure.

**Degree of Conversion (DC%)**

Raman spectra of the uncured and cured specimens of each resin composite were recorded by a Raman spectrometer (Nicolet FT-Raman 950, Thermo Nicolet Inc., Madison, WI, USA) operating with 120 scans at a resolution of 2 cm\(^{-1}\). Increments of each material were compressed between two polyester strips and two glass slides to produce a thin film (thickness of approximately 60 µm). Five films of each material were then photactivated according to the light-activation modes, with the light tip in contact with the glass slide. Raman spectra were recorded after a 24-h dry storage at 37ºC. The DC (%) was calculated from the ratio between the peak intensities of the aliphatic C=C bond (1638 cm\(^{-1}\)) to the aromatic C=C bond (1608 cm\(^{-1}\)), obtained from the cured and uncured specimens by the following equation:

$$DC(\%) = 100 \times \left[1 - \left(\frac{R_{cured}}{R_{uncured}}\right)\right]$$

where R = peak at 1638 cm\(^{-1}\) / peak at 1608 cm\(^{-1}\)

**Solubility and Salivary Sorption**

Disc-shaped specimens (1 mm thick and 6 mm in diameter) were fabricated using an aluminum mold. After filling the mold to excess, the material surface was covered with a polyester strip and a glass slide, and compressed with a device (500 g) for 20 s to avoid porosities. The discs were
photoactivated from the top. Ten discs were prepared for each studied material. The discs were placed in a desiccator containing freshly dried silica gel and transferred to an oven at 37°C. After 24 h, the discs were repeatedly weighed (AX 220 Shimadzu, Tokyo, Japan) until a constant mass (m₁) was obtained (i.e., disc mass variation was less than ±0.2 mg). The discs were immersed in artificial saliva (KCl, NaCl, MgCl₂, CaCl₂, Nipagin, CNC, Sorbitol, and deionized water)¹¹, neutral pH, at 37°C for 7 days, removed, washed in distilled water, dried at room temperature for 15 min and weighed (m₂). Thereafter, the discs were reconditioned to a constant mass (m₃), using the same protocol as for m₁. The thickness and the diameter of the discs were measured at four points with a digital caliper (MPI/E-101 Mitutoyo, Tokyo, Japan), and the volume (V) was calculated in mm³. The values of solubility (SL) and salivary sorption (SS) were obtained in µg/mm³ using the following equations:

\[ SL = \frac{m_1 - m_3}{V} \]  
\[ SS = \frac{m_2 - m_3}{V} \]

Statistical Analysis

Statistical analysis was performed using Statgraphics 5.1 Software (Manugistics, Rockville, MD, USA). After verifying normal distribution of errors and the homogeneity of variance using Shapiro-Wilk’s test and Levene’s test, respectively, each test variable (DC, solubility and salivary sorption) was analyzed separately by two-way analysis of variance and Student-Newman-Keuls’ test for multiple comparisons. Data were also submitted to regression analysis, with salivary sorption and solubility as the dependent variables and the DC as the independent variable. All statistical analyses were performed at a significance level of α = 0.05

RESULTS

The results of the DC, solubility and salivary sorption are presented in Table 2. Regarding DC (%), two-way ANOVA detected a significant influence for resin composite independent factor (p = 0.0071). The DC of Filtek P60 (54.22 ± 3.64) was higher than that of Filtek Supreme (51.70 ± 1.95) (p<0.05). Light-activation mode factor was proved significant (p=0.0001), with C mode presenting the highest DC (55.01 ± 3.15). The double interaction (resin composite vs. light-activation mode) was not statistically significant (p>0.05).

Regarding solubility, two-way ANOVA detected a significant influence for the two main factors: resin composite and light-activation mode (p<0.05). The solubility of P60 (0.40 ± 0.03 µg/mm³) was lower than that of Su (0.45 ± 0.05 µg/mm³). Furthermore, the solubility with C mode (0.39 ± 0.03 µg/mm³) was lower than that presented by SS mode (0.46 ± 0.04 µg/mm³), (p<0.05). The double interaction was also proved significant (p = 0.0138).

For salivary sorption, two-way ANOVA revealed statistically significant difference for both main factors, as well as for the double interaction (p<0.05). The salivary sorption of P60 (6.87 ± 0.31 µg/mm³) was lower than that presented by Su (8.04 ± 0.77 µg/mm³). Indeed, the sorption of specimens photoactivated with C mode (7.02 ± 0.43 µg/mm³) was lower than that of specimens activated with SS mode (7.89 ± 0.91 µg/mm³)(p<0.05).

Correlation was found between solubility and DC, r = -0.89 (Figure 1), as well as between solubility and salivary sorption, r = 0.95 (Figure 2). No correlation was found between DC and salivary sorption.

![Figure 1](image_url)

**FIGURE 1**- Regression line (linear model) of solubility plotted against the degree of conversion (r = -0.89, p < 0.05)

| Groups  | DC%          | Solubility (µg/mm³) | Salivary sorption (µg/mm³) |
|---------|--------------|---------------------|----------------------------|
| P60C    | ⁰57.15 ± 2.64 | ⁰0.38 ± 0.02        | ⁰6.70 ± 0.32               |
| P60SS   | ⁰51.28 ± 1.11 | ⁰0.43 ± 0.02        | ⁰7.04 ± 0.19               |
| SuC     | ⁰52.88 ± 1.96 | ⁰0.41 ± 0.03        | ⁰7.35 ± 0.24               |
| SuSS    | ⁰50.52 ± 1.13 | ⁰0.49 ± 0.02        | ⁰8.74 ± 0.35               |

Within each column, values with same superscript letters are not statistically different (α = 0.05).
DISCUSSION

As in the present study the DC obtained with C light-activation mode was higher than that obtained with SS mode, the first research hypothesis of the present study was rejected. On the other hand, as the solubility and salivary sorption of the nanofilled composite was higher than that of the hybrid one, the second research hypothesis was supported. Finally, as only the correlation between the DC and solubility was proved, the third research hypothesis was partially accepted.

The properties of resin composites as well as their DCs are influenced by both phases of their composition, i.e. polymeric matrix and filler particle system. Both resin composites analyzed in the present study are representative of Filtek series (3M ESPE, St Paul, MN, USA). According to this manufacturer, all composites from this series have the same composition regarding polymeric matrix. Based on this, all findings about the properties assessed in the present study were discussed taking into account only the influence of the filler particle systems.

The physical and mechanical properties of resin composites are strongly influenced by the degree of conversion, defined as the percentage of reacted aliphatic C=C bonds from the dimethacrylate monomers present in their polymeric matrices. Thus, the extent of this response plays a crucial role in the clinical performance of resin composite restorations. In the present study, the values of DC ranged from 50.52% (nanofilled composite) to 57.15% (hybrid composite). These values are in agreement with those of other studies that have used Raman spectroscopy to evaluate the degree of conversion of resin composites. The DC obtained with C light-activation mode was higher than that obtained with SS mode. Given that the radiant exposures generated by both light-activation modes were close to the same value, this result was unexpected. Several published studies have shown that light-activation modes that used similar radiant exposures produced statistically similar DCs. However, the data in Table 2 show that this result was strongly influenced by the behavior of hybrid resin composite. The only reasonable explanation for this result could be based on light-scattering phenomenon. A previous published study showed that at a wavelength of 470 nm a resin composite with mean filler particle size in the nanoscale, i.e. Silux Plus (3M, St. Paul, MN, USA), showed a significantly lower light transmittance than a hybrid composite. According to the authors, the light that passed through the resin composite was scattered by the small filler particles and light transmittance was reduced. It is possible that in the presented study the non-agglomerated silica nanoparticles with mean size of 20 nm may have caused a light-scattering effect in nanofilled composite. Thus, the light intensity might have been attenuated and the DC decreased. On the other hand, in the hybrid composite, light transmittance was probably higher, and so was its DC.

Previous studies have shown that soft-start light-activation modes improved the cavity wall-resin composite interface sealing. As in the present study the nanofilled composite presented similar DCs with either of the light-activation modes, it is reasonable to consider SS light-activation mode more advantageous for this type of resin composite.

The mean solubility values presented by the tested resin composites varied from 0.38 to 0.49 µg/mm³. This range is in agreement with the findings of a previous study. As expected, the nanofilled composite presented a higher solubility than the hybrid one. This result may be attributed to the differences in the filler particle systems of the two resin composites. Although both materials have similar volumetric filler content, i.e. 59.5 vol% for nanofilled and 61 vol% for hybrid, the filler particles of nanofilled composite will theoretically present a greater total surface area, due to nonagglomerated 20 nm silica filler. Two factors derived from this fact can increase the solubility. Firstly, a great amount of ions will be released from the surfaces of filler particles. Secondly, coupling agents associated with the filler particles, e.g. g-methacryloxypropyltrimethoxysilane, are prone to hydrolysis via ester linkages within the molecules or siloxane links that are formed with the filler particles. Water in contact with silica filler surfaces breaks siloxane bonds to form silanol groups and facilitate particles debonding. Based on this, it is reasonable to speculate that more filler particles were eluted from the nanofilled composite, thus increasing its solubility. Water in contact with silica filler surfaces breaks siloxane bonds to form silanol groups and facilitate particles debonding. Based on this, it is reasonable to speculate that more filler particles were eluted from the nanofilled composite, thus increasing its solubility. To emphasize this discussion, Hofmann, et al. (2002) analyzing the leachable components from several resin composites showed that Silux Plus (a microfill composite with silica filler particles similar to those present in Supreme) had a higher solubility than Z250 (a resin composite similar to P60). Obviously, among other aspects, this behavior could also be associated with differences in filler particles between these resin composites. As shown in Figure 1, correlation was found between DC and SL. This outcome is supported by the findings of previous studies. With regard to salivary sorption, differences were also found between resin composites, with the nanofilled (8.04 ± 0.77 µg/cm³) presenting a worse behavior than hybrid composite (6.87 ± 0.31 µg/cm³). The sorption phenomenon in polymer-based restorative materials, such as resin composites, is mainly dependent on the hydrophilicity of
At sufficiently high water sorption, macromolecular polymer chains undergo a relaxation process as they swell to absorb the water. Initially, the presence of water softens the polymer by swelling the network and reducing the frictional forces between the polymer chains. After the relaxation process, unreacted monomers trapped in the polymer network are released to the surroundings at a rate that is controlled by the polymer’s swelling and relaxation capacity. Taking into account that linear polymer chains have a greater relaxation capacity, this explanation could justify the greater SL obtained with the SS light-activation mode. Figure 2 shows the correlation between SL and SS. It is clear that the greater the amount of saliva absorbed, the greater the amount of components that could leach out from resin composites.

**CONCLUSIONS**

In conclusion, the results of the present study showed that the SS light-activation mode produced a lower DC, indicating that, instead of the final radiant exposure, the initial low irradiance may affect the DC of resin composites. The nanofilled composite presented higher solubility and salivary sorption than the hybrid composite. This suggests that the total surface area of the filler particle systems plays a key role in the properties of these resin composites. Therefore, light-activation modes that lead to a higher DC may diminish the solubility of resin composites. From a clinical standpoint, these results suggest that nanofilled composites may present higher degradation in the oral environment than hybrid ones. Moreover, soft-start light-activation mode may increase the solubility of resin composites.

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