Optimizing the concentration of 2,4,6-trimethylbenzoyldiphenylphosphine oxide initiator in composite resins in relation to monomer conversion

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The aim of this study was to optimize the concentration of 2,4,6-trimethylbenzoyldiphenylphosphine oxide (Lucirin® TPO) in unfilled and filled composite resins in relation to the degree of conversion (DC). Increasing concentrations of Lucirin® TPO between 0.05–4.97 wt% were added to equimolar mixtures of Bis-GMA/TEGDMA. Filled resins contained 75 wt% fillers. Standardized samples were cured using a polywave LED light-curing unit (bluephase® G2, Ivoclar Vivadent). Increased initiator concentrations increased logarithmically the DC of unfilled and filled resins. The DC of unfilled resins was in the range of 73–91% at the top and 63–81% at the bottom surfaces. The DC in unfilled and filled resins reached a plateau at 1.08 wt% and 1.50 wt% Lucirin® TPO, respectively. Fillers significantly reduced conversion but had no effect on the logarithmic relationship between initiator concentration and the DC.

Keywords: Lucirin® TPO, Composite, Degree of conversion, Raman, Light curing unit

INTRODUCTION

Polymerization of dental adhesives and resin-based composites (RBCs) is still generally based on light activation of a camphorquinone (CQ)-tertiary amine system. CQ alone has been shown to have a clinically unacceptable low rate of polymerization and significantly lower final conversion rate compared to the CQ-amine system. The use of the CQ-amine system is limited by aesthetic requirements due to the yellow, unbleachable, chromophore group in CQ as well as the by-products of the addition of fillers have an effect on the DC.

Alternative initiators, such as α-diketone phenylpropanedione, acylphosphine oxide derivatives, camphorquinone derivative with an acylphosphine oxide group and octyloxy-phenyl-phenyl iodonium hexafluorantimonate have been used in experimental dental resins.

It has been stated that 2,4,6-trimethylbenzoyl diphenylphosphine oxide (Lucirin® TPO) is already present in some commercial RBCs. Lucirin® TPO has been studied as an alternative to the CQ-amine system in experimental resin-based composites (RBCs) as well as RBC formulations. Unfilled methacrylate mixtures containing Lucirin® TPO showed comparable or higher degrees of conversion (DC) and color stability than CQ-amine and phenylpropanedione containing resins. Lucirin® TPO resulted in higher DC than CQ-amine in composite resins cured with halogen or polywave light-curing units (LCUs). In dental adhesives, the use of Lucirin® TPO instead of CQ-amine results in similar DC and mechanical properties when cured with a polywave LED LCU.

A disadvantage of Lucirin® TPO is that its absorption spectrum extends from 380 nm to about 425 nm and does not match the 430–490 nm emission spectra of most monowave LED LCUs. Recently marketed polywave LED LCUs have two emission peaks, one around 467 nm to cover the absorption spectrum of CQ and another around 400 nm to match Lucirin® TPO.

No study could be found on the effect of varying concentrations of Lucirin® TPO on the DC of experimental composite resins. The aim of this study was to optimize the concentration of Lucirin® TPO in unfilled and filled composite resin mixtures for maximum DC in clinically relevant curing conditions. The null hypotheses were that neither different concentrations of the initiator nor the addition of fillers have an effect on the DC.

MATERIAL AND METHODS

The following materials were used in the study: bisphenol A bis(2-hydroxy-3-methacryloxypropyl)ether (Bis-GMA), triethylenglycol dimethacrylate (TEGDMA), (Esstech Inc., Essington, PA, USA), Lucirin® TPO and fillers used in Tetric EvoCeram, both supplied by Ivoclar Vivadent (Schaan, Liechtenstein). According to the manufacturer's data, the fillers are a mixture of barium oxide, ytterbium trifluoride and mixed oxides, 40–3000 nm in size with a mean particle size of 550 nm. All materials were used as received. Bis-GMA and TEGDMA, 1 mmol each, were measured in a dark eppendorf on an analytical balance (Sartorius AG, Goettingen, Germany, d=0.01 mg), and homogenized for 24 h on a rotator wheel (Stuart® SB3, Bibby Scientific Ltd, Staffordshire, UK). Lucirin® TPO was added in concentrations of 0.00125–0.12 mmol that corresponded to 0.05–4.97 wt%.

Filled resins were prepared as 25 wt% resin and 75 wt% fillers. Appropriate amounts of fillers were measured on the same analytical balance and added...
to each resin mixture in small amounts followed by vortexing and centrifuging (Heraeus Pico, DJB Labcare Ltd, Buckinghamshire, UK) until homogenous mixtures were achieved. Prepared filled resins were stirred by hand using a metal spatula immediately before sample preparation.

The absorption spectrum of Lucirin® TPO in toluene (0.00152 M ⁻¹) was obtained in the range of 380–530 nm using a UV-Vis spectrophotometer (Agilent 8453 UV-Visible spectrophotometer, Agilent Technologies UK Limited, Stockport, UK).

Each sample was prepared by filling a plastic mould (Felix, Novi Sad, Serbia), 5 mm in diameter and 2 mm thick, placed on a Mylar strip on a microscopic slide. The material was covered by another Mylar strip and light cured for 20 s using a polywave LED LCU (bluephase® G2, Ivoclar Vivadent, Schaan, Liechtenstein). The tip-to-surface distance of 1 mm was maintained by a custom-made spacer. The curing parameters, light irradiance, energy and emission peaks were monitored using MARC™ (BlueLight analytics Inc, Halifax, NS, Canada) which incorporates a spectroradiometer (USB 4000, Ocean Optics, Dunedin, FL). The samples were wet polished with superfine polishing discs (Buehler, Lake Bluff, IL, USA) for 10 s. Three samples per group were prepared for unfilled and six per group for filled resins according to the sample size calculation based on the number of groups and estimated standard deviation values.

Micro-Raman analysis was done within 10 min post-curing with the samples being retained in the moulds. Standard micro-Raman parameters of the spectrometer used in this study (LabRam 300, Horiba JobinYvon, Stanmore, Middlesex, UK) were: 20 mW HeNe laser with 632.817 nm wavelength, spatial resolution ~1.5 μm, spectral resolution ~2.5 cm⁻¹, ×100 magnification.

The DC was calculated according to the following formula:

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

where R is the ratio of peak heights at 1639 cm⁻¹ and 1609 cm⁻¹ in cured and uncured material which served as reference. Three point spectra each were taken from the top and bottom surface of each sample.

Statistical analysis was done in Minitab 15 (Minitab Inc., State College, PA, USA). The assumptions required for parametric testing were validated using Kolmogorov-Smirnov and Bartlett’s tests. The differences in DC values were evaluated using a general linear model ANOVA for the factors “concentration” and “surface”. Further testing was done using one-way ANOVA with the Bonferroni correction and Tukey’s post hoc comparisons for each surface, top or bottom, of unfilled and filled resins. The level of significance was α=0.05. Regression analysis was performed to establish the relationship between the initiator concentration and the resultant DC of resins.

RESULTS

Figure 1 shows the absorption spectrum of Lucirin® TPO and emission spectrum of bluephase® G2. The mean irradiance of the bluephase® G2 LCU was 1208.94 mW/cm² and the total energy was 2.33 J/cm² and 23.40 J/cm² in the 380–420 nm and 420–540 nm regions, respectively.

Figures 2 and 3 show plotted DC values against initiator concentrations. R² values of 0.9552 and 0.9423 for the top surfaces in unfilled and filled resins, respectively, indicated a better logarithmic fit compared to the bottom surfaces. The DC values at the bottom surfaces of unfilled resins showed a better logarithmic fit (R²=0.8525) than the filled resins which showed least predictability (R²=0.7716).

Statistical analysis is summarized in Tables 1 and 2. In both types of resins very low initiator concentrations showed significantly lower DC values at the top and bottom surfaces compared to higher
initiator concentrations. Lucirin® TPO in excess of 0.43 wt% resulted in the DC values that started to level off with very small increase with each subsequent initiator concentration. Unfilled resins containing more than 1.08 wt% Lucirin® TPO showed similar DC values (p>0.05), except for the top surface of unfilled resin containing the highest concentration of initiator which resulted in the highest DC values. The addition of fillers reduced the DC by about 10–20%. There were no statistically significant differences in the DC values between filled resins containing more than 1.50 wt% Lucirin® TPO (p>0.05).

Figure 4 shows the average DC ratios between bottom and top surfaces and between filled and unfilled resins. In most cases, higher bottom-to-top ratios were observed for filled compared to unfilled resins. Similarly, higher filled/unfilled ratios were found for bottom compared to top surfaces.

DISCUSSION

Generally, increased concentrations of Lucirin® TPO resulted in higher DC of both unfilled and filled resins. On the other hand, the addition of fillers reduced the DC of the resins irrespective of the amount of Lucirin® TPO. Therefore, both null hypotheses were rejected.

Light irradiance of the polywave bluephase® G2 LCU obtained using MARC™, corresponded to manufacturer’s technical data. Further analysis of light curing parameters revealed that energy of about 2 J/cm² was delivered in the range of 380–425 nm. This means that less than 10% of the total energy of this LCU initiated Lucirin® TPO. On the other hand, previous studies have shown that substantially higher energy is required for sufficient polymerization of RBCs containing a CQ-amine system. The differences in the initiating potential between Lucirin® TPO and CQ-amine and the
### Table 1  Summary of statistical analysis for unfilled resins: upper and lower 95% confidence intervals (CI) and statistically significant differences between groups

<table>
<thead>
<tr>
<th>Lucirin® TPO concentration</th>
<th>Top surface</th>
<th>Bottom surface</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>95% CI</td>
<td>Statistical significance</td>
</tr>
<tr>
<td>mmol</td>
<td>wt%</td>
<td>95% CI</td>
</tr>
<tr>
<td>0.00125</td>
<td>0.05</td>
<td>72.5–73.7</td>
</tr>
<tr>
<td>0.0025</td>
<td>0.11</td>
<td>76.7–80.1</td>
</tr>
<tr>
<td>0.005</td>
<td>0.22</td>
<td>80.5–82.5</td>
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<tr>
<td>0.01</td>
<td>0.43</td>
<td>81.2–84.5</td>
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<tr>
<td>0.015</td>
<td>0.65</td>
<td>86.1–87.9</td>
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<td>0.86</td>
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<tr>
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<td>1.29</td>
<td>88.6–89.8</td>
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<tr>
<td>0.035</td>
<td>1.50</td>
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<tr>
<td>0.04</td>
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<td>87.3–90.2</td>
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<td>0.06</td>
<td>2.55</td>
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<tr>
<td>0.12</td>
<td>4.97</td>
<td>90.7–92.7</td>
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Same numbers indicate no significant difference (p>0.05)

### Table 2  Summary of statistical analysis for filled resins: upper and lower 95% confidence intervals (CI) and statistically significant differences between groups

<table>
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<tr>
<th>Lucirin® TPO concentration</th>
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<th>Bottom surface</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>95% CI</td>
<td>Statistical significance</td>
</tr>
<tr>
<td>mmol</td>
<td>wt%</td>
<td>95% CI</td>
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<tr>
<td>0.12</td>
<td>4.97</td>
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Same numbers indicate no significant difference (p>0.05)
lack of data for polywave LED LCUs justify the need to systematically assess the effect of varying concentrations of Lucirin® TPO on the DC of unfilled and filled resins cured with such LCUs.

In the present study, a wide range of initiator concentrations was used, from 0.05 wt% to 4.97 wt% Lucirin® TPO, in an equimolar Bis-GMA/TEGDMA mixture. Concentrations between 0.037 wt% and 1.5 wt% Lucirin® TPO were used in previous studies which evaluated various properties of resins containing different initiators2,12,18,19. Since manufacturers state that initiator concentration in their materials is up to 1 wt%, small increments of about 0.2 wt% Lucirin® TPO were tested in this range.

Samples were polished to remove the oxygen inhibition layer and obtain DC values that closely match the bulk polymer. Wet polishing was applied since it has been shown to have no effect on DC values20. Micro-Raman analysis was done within 10 min post-polymerization since studies on polymerization kinetics of various resin models and RBCs have shown a plateau in conversion within minutes of the start of polymerization21-23).

The present results have shown a logarithmic relationship between the amount of Lucirin® TPO and the resultant DC for both unfilled and filled resins. The greatest increase in the DC values occurred between the two groups containing the smallest amount of initiators. The increase in the DC values approached plateau for initiator concentrations exceeding 1.08 wt% in unfilled and 1.5 wt% in filled resins. Small rise in the DC beyond these concentrations had no statistically significant effect on monomer to polymer conversion and is not expected to have a significant effect on material properties. On the other hand, initiator concentrations of 0.22–0.43 wt% seem to be the lower cut-off points for acceptable DC values. Further studies are necessary to relate monomer conversion to various material properties as a function of Lucirin® TPO concentration. The present study may serve as an indicator of relevant initiator concentrations with regard to changes in monomer to polymer conversion.

Addition of fillers significantly reduced the DC irrespective of initiator concentrations. This is in agreement with a study by Leprince et al.12 who reported the same finding for resins initiated by CQ-amine or Lucirin® TPO. Previous studies have also reported lower DC values with increased filler loading24 and particle size25. Present findings have shown that fillers did not affect the logarithmic relationship between the initiator concentration and the DC. This suggests that the free radical polymerization occurs in much the same way across the entire range of tested initiator concentrations irrespective of the presence of fillers. Possible filler effect on light penetration and restricted mobility within the growing polymer may be the reasons why higher initiator concentrations in the filled resins were required to reach the conversion plateau compared to unfilled resins.

A previous study has reported that Lucirin® TPO results in lower depth of cure in unfilled and filled resins compared to the equimolar concentration of CQ-amine. This was attributed to a higher molar absorptivity of Lucirin® TPO which may lead to reduced penetration of light with depth25. In the present study, higher bottom-to-top ratios in filled resins indicate less difference in DC values across sample thickness compared to unfilled resins. Once fillers are added monomer conversion becomes restricted even in the surface layers closest to the light source. Similarly, higher DC ratios for bottom than top surfaces in filled and unfilled resin containing the same initiator concentrations indicate less difference in DC values in unfilled and filled resins with depth. The decrease in monomer conversion with depth is greater in unfilled resins probably because of higher consumption of photons of light by the initiator in the absence of fillers which reduces Lucirin® TPO initiation with depth.

The consensus DC values for most experimental and commercial RBCs are in the range of 40–75%26-29.
The present results are in agreement with these previously reported DC values for RBCs initiated by CQ-amine systems. Lucirin® TPO concentration in excess of 0.86 wt% resulted in 74% or higher DC in the filled resins. Moreover, as little as 0.22 wt% Lucirin® TPO resulted in 68% DC which was still higher than many of the previously tested RBCs. It should be noted that such high DC values were obtained with only about 2 J/cm² of delivered light energy in the absorption region of Lucirin® TPO. A high reactivity of Lucirin® TPO when initiated with a polywave LED LCU may be due to its higher molar extinction coefficient, polymerization quantum yield and, thus, polymerization efficiency than CQ-amine in the case of matching emission spectra of the LCUs and absorption spectra of initiators[19,30].

CONCLUSION

Lucirin® TPO has shown a high reactivity upon irradiation with the polywave bluephase® G2 LED LCU whose additional emission peak around 400 nm matches the absorption spectrum of this initiator. Increased concentrations of Lucirin® TPO increased the DC logarithmically in an equimolar Bis-GMA/TEGDMA mixture. Monomer to polymer conversion reached plateau values at 1.08 wt% and 1.5 wt% of Lucirin® TPO in unfilled and filled resins, respectively. High conversion with depth and in the presence of fillers in clinically relevant material thickness was maintained when Lucirin® TPO was in excess of 0.22 wt%. Fillers significantly reduced conversion but had no effect on the relationship between initiator concentration and the DC.

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