Irradiation Modes’ Impact on Radical Entrapment in Photoactive Resins

INTRODUCTION

In modern dentistry, photopolymerizable resin-based composites have become the choice material for the direct restoration of damaged teeth. However, despite significant material improvements since their introduction, several drawbacks—such as inadequate curing depths and deleterious effects of polymerization shrinkage stress—remain. In an attempt to overcome these deficiencies, various material aspects have been investigated, such as inorganic filler content, resin formulation, curing light technology, and photoinitiation protocols. In the latter field, despite the large number of studies analyzing the effects of various initiation modes on the properties of photoactive materials, discrepancies within the literature remain. Globally, two main goals are pursued. First, there exists a demand for reduction in curing exposure to minimize chairside procedure times of multi-increment resin composite restorations. Accordingly, several researchers support the exposure reciprocity law (Sakaguchi and Ferracane, 2001; Halvorson et al, 2002; Price et al, 2004; Emami and Söderholm, 2003), claiming that radiant exposure (RE, J/cm²), the product of irradiance (I, mW/cm²) and exposure time (t, s), is the main determining factor of the degree of conversion and mechanical properties of the photoactive material. This supposes reciprocity between I and t, and encourages a tendency among dentists and manufacturers to use or suggest a high-power illumination to reduce curing time. However, other studies (Musanje and Darvell, 2003; Asmussen and Peutzfeldt, 2005; Peutzfeldt and Asmussen, 2005; Dewaele et al, 2009) have reported that although RE plays an important role, I and t independently influence polymer chain length, extent of crosslinking, and mechanical properties, and as such, exposure reciprocity does not hold under all conditions. Second, the use of “soft-start” curing protocols was proposed to delay the onset of polymer gelation. Notably, pulse-delay modes (consisting of a short flash of light, followed by a delay before the final polymerization) seem to contribute to a reduction of polymerization shrinkage stress (Kanca and Suh, 1999; Soh et al, 2004; Chye et al, 2005; Cunha et al, 2008). Nevertheless, some authors have also highlighted lower conversion values and/or inferior mechanical properties of composites cured using those modes (Asmussen and Peutzfeldt, 2001, 2003; Lu et al, 2005; Benetti et al, 2007; Dewaele et al, 2009). Although the use of alternative illumination methods may provide certain advantages, their effects on polymerization mechanisms and final polymer characteristics are poorly understood.
One interesting approach to a better grasp of the chemical process of photopolymerization is the observation of trapped free-radical concentration. It has been previously demonstrated that vitrified dental resins contain 2 types of trapped free radicals (R\textsuperscript{•}\textsubscript{Trapped}), which are detectable by electron paramagnetic resonance (EPR), namely, a “propagating” radical and an allylic radical type (Fig. 1) (Truffier-Boutry et al., 2003). The study of R\textsuperscript{•}\textsubscript{Trapped} is particularly relevant regarding dimethacrylate-based resins, since their polymerization leads to highly crosslinked networks, where monomolecular termination, as a consequence of radical entrapment, becomes the dominant pathway (Andrzejewska, 2001). A strong relationship between autoacceleration and radical trapping was also suggested (Zhu et al., 1990). This highlights the importance of considering the quantity of trapped radicals in the study of the impact of photoirradiation protocols, which, to our knowledge, has never been investigated with clinically relevant irradiation parameters. For that reason, the aim of this paper was to test the null hypothesis that no difference in concentrations of R\textsuperscript{•}\textsubscript{Trapped} existed between samples cured at similar RE but different I and t, with or without a delay.

MATERIALS & METHODS

Bisphenol A glycidyl dimethacrylate (Bis-GMA) (Heraeus Kulzer, Wehrheim, Germany) and triethylene glycol dimethacrylate (TEGDMA) (Aldrich, Belgium) were used without further purification from the supplier and mixed in a 50:50 wt% ratio. A photosensitizer system comprised of 0.5 mass% camphorquinone (CQ; Aldrich, Belgium) and 0.5 mass% of N,N-dimethyl-p-aminobenzoic acid ethyl ester (DABE, Aldrich) was incorporated into the resin. The resin was placed in a rectangular brass mould (7 x 1 x 1 mm), covered with Mylar film and cured (at 23 ± 1°C) with a halogen light (Optilux 501, Demetron/Kerr Co., Orange, CA, USA) with the curing tip (8 mm exit diameter) centrally placed on the mould, in contact with the film. Two different illumination modes were used: continuous and pulsedelay (one-second cure, one- to two- or three-minute delay, final cure), for which t and I were adapted so that only 2 different REs of 6 and 12 J/cm\textsuperscript{2} were obtained (Table). A rheostat was used to achieve 3 specific irradiances (150, 300, and 600 mW/cm\textsuperscript{2}) that were verified by differential scanning calorimetry (DSC822, Mettler Toledo, Zaventem, Belgium) equipped for in situ irradiation. An aluminum crucible filled with black graphite powder was irradiated to ensure complete absorption of emitted photons, while the other crucible was left empty and was not irradiated. Following irradiation, specimens (n = 3) were carefully stored in a lightproof container between subsequent measurements and recovered for analysis by EPR (MiniScope MS200 with rectangular resonator TE102, Magnettech, Berlin, Germany) 5 min, 1 to 6 hrs, 1 day, and 1, 2, and 3 mos post-irradiation. A similar EPR technique has been described in detail in other works (Ottaviani et al., 1992; Sustercic et al., 1997; Truffier-Boutry et al., 2003, 2006). An alignment device was used to ensure an identical position of the sample in the EPR cavity, enabling accurate comparisons of the peak intensities to be made. EPR parameters consisted of: center field, 3367.42 G; frequency, 9.30-9.55 GHz; sweep width, 198.40 G; modulation amplitude, 1 G; microwave power, 0.5 mW. Since peak width did not significantly vary with time, measurement of the height of some EPR peaks could be used to determine the relative concentrations of R\textsuperscript{•}\textsubscript{Trapped}, height of the fifth (from 334.9 to 335.5 mT) and fourth peaks (from 336.1 to 336.6 mT of the magnetic field) of the 9-line spectrum for allylic (R\textsuperscript{•}Allyl) and propagating radicals (R\textsuperscript{•}Propag), respectively (Fig. 1). These peaks are least influenced by the signal of the other radical types. Concentrations of R\textsuperscript{•}\textsubscript{Trapped} (in arbitrary units, a.u.) at each storage time were analyzed by one-way ANOVA and Tukey’s test (p = 0.05).

RESULTS

The evolution of R\textsuperscript{•}\textsubscript{Trapped} concentrations with time for the continuous modes for R\textsuperscript{•}Allyl and R\textsuperscript{•}Propag is presented in Figs. 2a and 2b, respectively. The curves of both radical species display similar profiles (logarithmic time scale), except during approximately the first 4 hrs, where concentrations of R\textsuperscript{•}Allyl decreased at higher rates compared with R\textsuperscript{•}Propag, which is further highlighted in Fig. 2c. The curves are parallel during the first 24 hrs (Figs. 2a, 2b). Therefore, the results and statistical analyses for both radical types will be displayed only for five-minute and 24-hour storage periods (Table). In the same way, since pulse-mode curves showed profiles similar to continuous-mode curves, but overlapped with them, they were not displayed here for more clarity. Nevertheless, values and statistical analyses that were not reported in the manuscript are available in the

![Nine-line EPR spectrum of experimental resin after polymerization and chemical structure of both radical species.](image)
Appendix. In the Table is also reported the mean degree of conversion of each mode (published data from Dewaele et al., 2009; data were obtained with the same resin and same irradiation protocols, and are reproduced here for reader convenience).

Generally, increased concentrations of R•Trapped were measured for continuous mode curing regimes following higher (12 J/cm²) compared with lower RE (6 J/cm²) (Figs. 2a, 2b, Table). However, the amounts of R•Trapped generated by 20 sec x 600 mW/cm² (12 J/cm²) and 40 sec x 150 mW/cm² (6 J/cm²) were not significantly different (p > 0.05). Moreover, while the increase in concentrations of both radicals is significant (p < 0.05) at twice the exposure time (40 sec x 150 mW/cm² to 80 sec x 150 mW/cm² and 20 sec x 300 mW/cm² to 40 sec x 300 mW/cm²), it is not significant (p > 0.05) at twice the irradiance (40 sec x 150 mW/cm² to 40 sec x 300 mW/cm² and 20 sec x 300 mW/cm² to 20 sec x 600 mW/cm²). When continuous modes with similar RE were compared, higher concentrations of R•Trapped were measured for longer exposure time with lower irradiance. There was no influence of the delay time on the concentration of R•Trapped, for either the application of a one-, two- or three-minute delay or when pulse delay was compared with continuous mode for the same RE (Table).

**DISCUSSION**

The two distinct phases of radical concentration, highlighting the decrease of R•Trapped following polymerization, are in accordance with previous work (Truffier-Boutry et al., 2006; Leprince et al., 2009). The decrease in R•Trapped from 0 to 24 hrs was attributed to post-cure volumetric shrinkage (Truffier-Boutry et al., 2006), whereas from 24 hrs to 3 mos, the effect of oxidation by atmospheric oxygen may play a more significant role in the
reduction of radical concentration (Leprince et al., 2009). Since 2 radical species are considered, the relationship between both species requires clarification. \(R_{\text{Propag}}\) concentrations decrease at higher rates than \(R_{\text{Allyl}}\) during the first 4 hrs. As mentioned above, during approximately the first 24 hrs after light cure, volumetric shrinkage occurs. As a result, \(R_{\text{Trapped}}\) move closer to each other, and the probability of termination increases due to shrinkage (Truffier-Boutry et al., 2006). But while \(R_{\text{Allyl}}\) are unable to react with species other than free radicals (higher stability due to a resonance phenomenon), \(R_{\text{Propag}}\) exhibit sufficient reactivity to interact also with remaining double-bonds. In this way, additional propagation reactions can repeat locally, slightly increasing DC (Truffier-Boutry et al., 2006) and local radical mobility, leading to some bimolecular radical termination through a reaction-diffusion-controlled termination mechanism (Anseth et al., 1995). The different behaviors of the 2 different radical species observed here are an additional proof in favor of the identification of \(R_{\text{Trapped}}\) proposed by Truffier-Boutry et al. (2003). Previous work (Ottaviani et al., 1992) also identified an EPR spectrum similar to that presented in Fig. 1, although in that study the signal disappeared several mos after irradiation, and other signals were detected in some commercial composites. Since unfilled experimental resins were tested in the present study, it could be hypothesized that fillers or other components of commercial resins may be responsible for the creation of additional minor signals.

Regarding the comparison of free-radical concentrations, the period from 5 min to 1 day post-cure should be considered, since radical concentration within this timeframe remains proportional between curing regimes. For continuous curing modes, the increase in \(R_{\text{Trapped}}\) concentration following twice the \(RE\), by doubling either \(I\) or \(t\), can be explained by the creation of a higher concentration of free radicals (generated by an increase in number of photons), which are more likely to become trapped. However, for similar \(RE\), higher concentrations of \(R_{\text{Trapped}}\) were measured for longer \(t\) and lower \(I\). This observation is in apparent contradiction to the reciprocity observed between \(I\) and \(t\) for CQ quantum yield of conversion (number of converted photoinitiator molecules per number of absorbed photons) (Chen et al., 2007). However, even if the same total amount of radicals is created, it does not necessarily result in the same number entrapped. This can be explained by a consideration of both types of termination mechanisms co-existing during polymerization (Lovell et al., 1999): While bimolecular termination probably prevails during the very initial stages of the reaction, monomolecular termination (first-order) dominates when molecular diffusion becomes limited (auto-acceleration). There is probably a loss of a given amount of radicals during the early stages by bimolecular termination. This loss (or early termination) is higher for high-irradiance protocols, since termination is proportional to \(\left[R_{\text{Trapped}}\right]^2\). As a result, at constant \(RE\) but higher \(I\) and shorter \(t\), the amount of growth centers created during auto-acceleration is lower, leading to a lower final concentration of \(R_{\text{Trapped}}\). In other words, the amount of \(R_{\text{Trapped}}\) measured in the present work may provide an indication of the quantity of growth centers created during auto-acceleration. In light of these results, the use of EPR to measure \(R_{\text{Trapped}}\) gives valuable information on monomolecular termination, which is critical in highly crosslinked systems. Other recent work has suggested that the applicability of exposure reciprocity law depends on factors other than irradiation parameters alone, such as the Bis-GMA/TEGDMA ratio (Feng and Suh, 2007). Besides, the efficiency of the photoinitiator or the amounts and types of inorganic fillers can also play a part in the resin reactivity, which can also result in different concentrations of \(R_{\text{Trapped}}\). For example, previous work has demonstrated that the concentration of \(R_{\text{Trapped}}\) is about 3 times higher in the organic fraction of a filled
resin than in an unfilled resin of similar composition (Leprince et al., 2009). As a consequence, the influence of filler content, monomer ratio, and photoinitiator might explain differences in previous results on commercial composites obtained by a similar EPR technique (Ottaviani et al., 1992). Accordingly, the measurement of R•_{Trapped} will be very useful in further understanding the impact of resin, photoinitiator chemistry, and filler morphology on complex photopolymerization processes, and related experiments are ongoing by the authors.

For pulse-delay modes, no influence of any delay time (1, 2, or 3 min with no irradiation) was observed on the concentration of R•_{Trapped} or compared with continuous mode curing with similar RE. However, DC values associated with pulse-delay protocols were lower than those associated with the continuous mode of similar RE (Dewaele et al., 2009), which can be explained by differences in polymer chain growth. The initial pulse probably results in microgels with relatively linear polymer chains (few pendant double-bonds available) (Poshusta et al., 2002; Dewaele et al., 2009) and where most free radicals probably terminated. During final irradiation, these microgel regions, in which unreacted double-bonds are less available, are certainly less reactive. Therefore, they may disturb the polymerization reaction compared with a continuous irradiation, leading to reduced DC. However, as discussed above, most of the free radicals are trapped when propagation becomes diffusion-controlled. Therefore, it might be assumed that the irradiation parameters at the time of auto-acceleration govern the final amount of trapped radicals, leading to comparable amounts of R•_{Trapped} with or without a delay.

In conclusion, for continuous modes, the null hypothesis must be rejected, since the concentrations of R•_{Trapped} between samples depend not only on RE but also on I and t. Conversely, regarding pulse-delay mode, the null hypothesis cannot be rejected, since the presence of a delay does not have a significant influence on the concentrations of R•_{Trapped} when compared with the equivalent continuous mode of the same irradiance and total irradiation time.

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REFERENCES
