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Effects of absorption and inhibition during grating formation in photopolymer materials

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Photopolymer materials are practical materials for use as holographic recording media, as they are inexpensive and self-processing (dry processed). Understanding the photochemical mechanisms present during recording in these materials is crucial to enable further development. One such mechanism is the existence of an inhibition period at the start of grating growth during which the formation of polymer chains is suppressed. Some previous studies have indicated possible explanations for this effect and approximate models have been proposed to explain the observed behavior. We examine in detail the kinetic behavior involved within the photopolymer material during recording to obtain a clearer picture of the photochemical processes present. Experiments are reported and carried out with the specific aim of understanding these processes. The results support our description of the inhibition process in an acrylamide-based photopolymer and can be used to predict behavior under certain conditions. © 2006 Optical Society of America

1. INTRODUCTION

Photopolymer materials have the ability to optically record high-diffraction-efficiency, low-loss, volume holographic gratings in self-processing materials and are of ever-increasing commercial importance. Improving their characteristics, to attain the full potential of these materials, requires the development of accurate models, validated using reproducible experimental data sets, which can ultimately provide clear insight into the photochemical processes involved during recording. Exploration of the kinetics involved in the recording of holographic gratings is an integral part of this work. In this paper the photochemical kinetics involved during recording in our acrylamide-based photopolymer material is examined. Specifically we aim to better understand what takes place inside the material during exposure, i.e., to explain the effects of the variation of the absorbance of the photosensitive dye with time and the suppression of radical production due to the presence of inhibitors. By improving the physical accuracy of our model, and understanding the relationship between the rate of polymerization and the concentrations of monomer, polymer, dye, initiator, and inhibitor, we aim in this paper to improve the validity of the one-dimensional polymerization-driven diffusion (PPD) model. Development of this model will facilitate the eventual full nonlocal polymerization-driven diffusion (NPDD) modeling of such materials.

We proceed as follows. First we examine the photochemical processes involved during grating formation and incorporate the suppression of radical production into the rate equations, which form the basis of our models. We then incorporate the effects of changes in the absorbance of the photosensitive dye during exposure. Having developed appropriate rate equations, we derive and solve the resulting coupled equations numerically. A set of experiments is carried out that, following comparison with the model, support the assumptions made in deriving our equations and illustrate the effects associated with the inhibition and bleaching processes.

2. PHOTOCHEMICAL PROCESSES

Assuming bimolecular termination, we begin by discussing radical chain polymerization. In particular we examine polymerization due to photoinitiation and examine the effect on excited dye molecules due to the presence of inhibitors such as oxygen. Furthermore we include the effect of a time-dependent transmittance, which describes the change in the material absorption during grating growth.

Free radical polymerization is a chain reaction involving three steps: initiation, propagation, and termination. Initiation involves the production of free radicals,

\[ I \rightarrow R^* \]

where \( k_d \) is the radical generation rate constant. The rate of this reaction is given by

\[ R_d = d[R^*]/dt = k_d[I] \]

where \([R^*]\) is the free radical concentration and \([I]\) is the initiator concentration. The free radicals then bind to a monomer \( M \) to form the chain initiation species \( M_1^* \),

\[ R^* + M \rightarrow M_1^* \]

where \( k_i \) is the initiation rate constant. The radical \( M_1^* \) then propagates by bonding with monomer molecules to form long polymer chains with an active tip known as a macroradical.
where $k_p$ is the propagation rate constant and $M_n^*$ is a macroradical of $n$ monomeric units where a monomeric unit is the largest constitutional unit contributed by a single monomer molecule. The initiator consists of a photosensitive dye and a reducing agent. The dye can become excited in the presence of a photon and when excited can accept an electron from the reducing agent, i.e., a tertiary amine (triethanolamine), and can then produce a free radical $R^*$:

$$
dye + h\nu \rightarrow dye^*,$$
$$
dye^* + M \rightarrow R^*.$$  

As the radicals are produced, some of the dye molecules, which are excited, return to their ground state or move to a triplet state:

$$
dye^* \xrightarrow{k_f} dye + h\nu,$$  

$$
dye^* \xrightarrow{k_{ic}} dye^T,$$  

$$
dye^T + C_I \xrightarrow{k_R} dye^* + C_I^* + H^*.$$  

An excited dye molecule, which has been promoted to that state due to a photon interaction, can either decay by fluorescence at a rate of $k_f$ back to its ground state or can be excited to a triplet state dye $^T$ by intersystem crossing at a rate of $k_{ic}$. The excited triplet state can then undergo an electron transfer reaction (rate constant $k_R$) with the coinitiator $C_I$, which creates the primary radical $C_I^*$, a dye radical $dye^*$, and a free proton $H^*$. The dye radical eventually reacts with other constituents and forms a bleached state. This process results in dye molecules being consumed and can lead to cessation of polymerization if the dye is consumed too quickly.

All of these effects tend to suppress the creation of radicals and therefore slow the rate of polymerization (of monomer). Other contributors to this radical suppression are inhibitors, which are present within the chemicals in our material, and the presence of oxygen, which also acts as an inhibitor. We assume that the effect of inhibition during the fabrication process is primarily due to the oxygen present in our material. Therefore we suggest that the oxygen reacts with the excited dye and deactivates it into a passive state:

$$
dye^* + \rho \xrightarrow{k_p} Y + \rho^*.$$  

where $Y$ is the photoreduction product and is assumed to only participate once (i.e., its relaxation time is long compared with the exposure time), $\rho$ is the oxygen, and $k_p$ is the deexcitation rate of the dye. This effect is present throughout exposure but is most noticeable at the start. The equation governing the removal of excited dye is

$$
M_n^* + M \xrightarrow{k_p} M_{n+1}^*,$$  

where $R_\rho$ is the rate of removal of free radicals due to the removal of excited dye molecules by oxygen. This process reduces the concentration of excited dye molecules available for creating free radicals. Thus it ultimately reduces the monomer polymerization rate.

The frequency of encounters between the free radicals is another factor determining the rate of polymerization. It can be accounted for using the cage effect. It is assumed that only some fraction $f$ of the free radicals produced will react with the monomer (in the starting reaction).

From Eqs. (2) and (8) we see that the rate of production of free radicals $R_r$ that are responsible for the production of monomer radicals, with the inclusion of the presence of oxygen, can then be given by

$$
R_r = f k_d [I] - R_\rho,$$

with the initiator concentration $[I]$. From relation (3) the rate of production of monomer radicals $R_i$ can be written as

$$
R_i = d [M_n^*] / dt = k_i [R^*] [M],$$

where $[M]$ is the monomer concentration and $[R^*]$ is the primary radical concentration. In general the rate of production of monomer radicals $R_i$ is much greater than the rate of production of free radicals $R_r$; therefore initiator radicals are consumed as fast as they are generated. The rate-determining step is thus the decomposition of the initiator. Initiator radicals are consequently formed with a rate

$$
d[R^*] / dt = R_r - R_i = (f k_d [I] - R_\rho) - k_i [R^*] [M] = 0.$$  

Thus the rate of monomer radical generation $R_i$ is equal to the chain initiation rate and

$$
R_i = k_i [R^*] [M] = f k_d [I] - R_\rho.$$  

Chain growth would continue in this way until the supply of monomer is exhausted were it not for the strong tendency of radicals to react in pairs to form paired-electron covalent bonds with the loss of radical activity. At sufficiently low initiator concentrations, chain termination will occur mainly by combination,

$$
M_n^* + M_m^* \xrightarrow{k_{t_c}} M_{n+m},$$

or by disproportionation,

$$
M_n^* + M_m^* \xrightarrow{k_{t_d}} M_n + M_m.$$  

On the basis of these observations, the rate of termination $R_t$ can then be given by

$$
R_t = k_t [M^*]^2,$$

where $k_t = k_{t_c} + k_{t_d}$ is the termination constant, and $[M^*]$ is the total concentration of all chain radicals of size $M_n$ and larger. For low-monomer conversions we assume that the rate of radical formation equals the rate of radical disappearance (Bodenstein steady-state principal); this is
therefore steady state in this process implies that the rate of initiation $R_i$ and the rate of termination $R_t$ are equal, yielding\
\[ \frac{f k_p [I] - R_p}{k_t} = k_p [M^*]^2. \] (16)

Solving for $[M^*]$ we obtain\
\[ [M^*]_{stat} = \left( \frac{f k_p [I] - R_p}{k_t} \right)^{1/2}, \] (17)

where $[M^*]_{stat}$ is the total concentration of all chain radicals of size $M_1^*$ and larger at steady state. Assuming that much more monomer is consumed due to propagation polymerization than in the initiation reaction, the propagation rate $R_p$ [the rate of reaction of relation (4)] is approximately equal to the polymerization rate $R_p$, therefore\
\[ R_p = -d[M]/dt = k_p[M^*][M]. \] (18)

Substituting into Eq. (18) from Eq. (17) for $[M^*]$ yields\
\[ R_p = k_p \left( \frac{f k_p [I] - R_p}{k_t} \right)^{1/2} [M]. \] (19)

Examining the photochemical formation of free radicals, there are a number of initiation mechanisms, many involving a photochemical electron transfer reaction. If we reexamine the way in which the monomer radicals are formed, we see that they are dependent on the quantity of free radicals formed per photon absorbed and the intensity of the light used for recording. As the inhibitor present indirectly consumes some of the absorbed photons, there is a reduction in the number of free radicals available for the initiation of monomer radicals and thus for the formation of polymer chains. The rate of initiation $R_i$ can therefore be given by\
\[ R_i = f(\Phi' - R'_p) I_a = \Phi I_a - R_p, \] (20)

where $\Phi'$ is the total number of radicals produced per photon absorbed and $R'_p$ is the decrease due to the presence of oxygen. $I_a$ is the intensity of light absorbed in moles of light quanta per liter per second and $\Phi$ is the total number of propagating chains that would be initiated per light photon absorbed if no inhibition took place. As only a fraction $f$ of the free radicals, which are produced, reacts with the monomer in the starting reaction, the cage effect has been included. $R'_p$ indicates that the process of production of radicals, which cannot produce chains due to the cage effect, may still undergo inhibition.

Let us assume cosinusoidal spatially modulated illumination, i.e., $I(x,t) = I_0[1 + V \cos(Kx)]$ where $V$ is the fringe visibility, $K = 2\pi/\lambda$, the grating vector magnitude, and $\lambda$ is the grating period. The concentration of photosensitizers is related to the absorbed intensity by Beer’s law:\
\[ I_a(x,t) = I(x)[1 - \exp(-\epsilon Z(t)d)] = I(x)[1 - T(t)], \] (21)

where $\epsilon$ is the molar absorptivity, $Z(t)$ is the time-dependent concentration of the photosensitizers (initiators), and $d$ is the photopolymer layer thickness. Since the concentration of photosensitizers is a function of time, the transmittance of the layer $T(t)$ (Refs. 7 and 8) also depends on time and is discussed below. The concentration of free radicals given in Eq. (17) can now be written as\
\[ [M^*] = \left( \frac{f(\Phi' - R'_p) I_a [1 - T(t)]}{k_t} \right)^{1/2}. \] (22)

Therefore the polymerization rate from Eq. (19) is given by\
\[ R_p = k_p[M] \left( f(\Phi' - R'_p) I_a [1 - T(t)] \right)^{1/2} = \kappa(t)[M][I(x)]^{1/2}, \] (23)

where $\kappa(t) = \kappa_0(t) A(t)^{1/2}$, $A(t) = 1 - T(t)$, $\kappa_0(t) = k_p \left( \Phi(t)/k_t \right)^{1/2}$, and $\Phi(t) = f(\Phi' - R'_p)$. We can summarize the above as follows: $A(t)$ tells us the fraction of the incident light absorbed, $\Phi'$ tells us the fraction of this absorbed light that results in initiation, and finally $\Phi(t)$ tells us the fraction of photons that lead to polymerization.

The inclusion of the transmittance $T(t)$ allows a time-varying model for the polymerization rate $R_p$ to be obtained. At the start of exposure the transmittance of the exposing light will be small due to the high absorbance of the photosensitive dye. This high absorption results in large values of $I_a(x,t)$ and of the polymerization rate factor $\kappa(t)$, and therefore a high rate of polymerization $R_p$. Similarly, as the transmittance of the layer increases during the exposure, both the light absorbed by the photopolymer material layer and the polymerization rate constant decrease, resulting in a lower rate of polymerization $R_p$.

To estimate $T(t)$, several transmittance curves were experimentally obtained for differing exposure intensities in our standard material layer.7 There is an initial nonzero transmittance at the beginning of exposure as the material is never completely opaque. The transmittance function $T(t)$ was then determined based on fits to the experimental data. In Section 3 we introduce a loss fraction $B$ to allow for nonabsorptive losses, i.e., Fresnel boundary reflections. $T(t)$ is estimated by fitting with the function\
\[ T(t) = E + G[1 - \exp(-a_0[\epsilon + a_1(t)^2])], \] (24)

where $E$, $G$, $a_0$, and $a_1$ are constant parameters related to the exposure intensity and the initial transmittance of the layer. Equation (24) is used in our numerical simulations, and some typical parameter results (and fitting errors) for different exposing intensities are given in Table 1.

In this section we have derived, for the first time to our knowledge, the rate equations governing the photochemical processes involved in grating formation that include a term to explicitly account for radical suppression due to the effect of inhibition, which we have assumed to be caused mainly by oxygen. Furthermore, following the work of Blaya et al., time dependence for the absorbance of the photosensitive dye and photoinitiators is included in our rate equations.
3. POLYMERIZATION-DRIVEN DIFFUSION

In this section the basic theory behind grating growth will be presented. Following Galstyan et al., 14 we incorporate the presence of inhibition and time-varying absorption effects in the material, as discussed in Section 2, into a local PDD model.2

A dry photopolymer layer typically consists of a monomer, binder, cross-linker, an electron donor, and a photoinitiator. As the material is exposed to the recording beams, the monomer is polymerized, and the amount of polymerized monomer increases with the exposure. In our material more monomer is polymerized in the bright fringes of the interference pattern than in the dark fringes. This results in a higher concentration of monomer in the dark regions than in the bright, and therefore a spatial monomer concentration gradient. The excess monomer will tend to diffuse into the bright regions.2–6,19

The governing one-dimensional diffusion equation is

$$\frac{\partial u(x,t)}{\partial t} = \frac{\partial}{\partial x} \left[ D(x,t) \frac{\partial u(x,t)}{\partial x} \right] - F(x,t) u(x,t),$$  \hspace{1cm} (25)

where $u(x,t)$ is the monomer concentration, $D(x,t)$ is the diffusion constant, and $F(x,t)$ is the polymerization rate. Equation (25) is Fick’s law with the addition of a driving function representing the physical effects of the photopolymerization. The light intensity in the material is assumed periodic and is described by

$$I(x,t) = I_0 [1 + V \cos(Kx)],$$  \hspace{1cm} (26)

with $I_0$ the average irradiance. In Section 2 we derived an equation governing the resulting polymerization rate. It is proportional to the exposure irradiance raised to $\gamma = 1/2$ (Ref. 21):

$$F(x,t) = F_0(t) [1 + V \cos(Kx)]^2 A(t),$$  \hspace{1cm} (27)

where $A(t)$, the time-varying absorbance of the material, has been included in the expression for the polymerization rate to account for the change in absorbed intensity during exposure. To account for the nonabsorptive losses present, a loss fraction $B$ has been included by defining the polymerization rate parameter as $F_0(t) = \kappa_0(t) I_0^2 (1 - B)$. The value of the loss fraction $B$ is empirically obtained by repeated measurement of the losses in the layers and plates when all the monomer is polymerized and the dye is bleached, assuming that the nonabsorptive losses can be obtained. The expression for the polymerization rate parameter also includes the time-varying function $\kappa_0(t)$, which we introduce to model the inhibition period present at the start of exposure.16–18,22,23

For the $I(x,t)$ in Eq. (26), the monomer concentration can be written as a cosine series

$$u(x,t) = \sum_{i=0}^{\infty} u_i(t) \cos(iKx).$$  \hspace{1cm} (28)

This is substituted into Eq. (25) with the initial condition that $u(x,0) = U_0$, where $U_0$ is the initial uniform monomer concentration in the material. In our analysis we assume that harmonics of an order greater than 2, $i > 2$, can be neglected, i.e., their contributions are assumed negligible in comparison with that of the first three terms.2 A set of first-order coupled differential equations, in terms of the monomer concentration harmonic amplitudes, is obtained.2,14

We now examine the $F_0(t)$ term appearing in Eq. (27) using the result from Section 2. From repeated observations of our experimental results, we know that grating growth is negligible for some period of time at the start of exposure. This appears to be primarily due to the action of inhibitors, such as oxygen,14 which suppress the creation of free radicals. The process is as follows:

(i) the photosensitive dye is excited by the exposing photons;
(ii) the excited dye then reacts with the monomer generating free radicals;
(iii) this leads to the formation of polymer chains,14 and then
(iv) the inhibitor acts to deactivate the dye from its excited state, stopping radical creation.

Although this process continually takes place during grating formation, it is most obvious (and most easily observed) at the beginning of exposure due to the high inhibitor concentration and the low concentration of excited dye.22 As exposure proceeds, the suppression of radicals becomes less visible. We model this sharp temporal-state transition using a step function14 $\Theta(x)$, where

| Parameter | Case 1 | Case 2 | Case 3 |
|-----------|--------|--------|--------|
| $I_0$ (mW/cm$^2$) | 3.5 | 4.5 | 5.0 |
| $E$ (mW/cm$^2$) | 0.857 | 1.124 | 1.636 |
| $G$ (mW/cm$^2$) | 1.517 | 1.831 | 2.143 |
| $\sigma_0$ ($\times 10^{-2}$) | 6.238 | 7.321 | 8.397 |
| $\sigma_1$ ($\times 10^{-4}$) | 0.891 | 1.211 | 3.623 |
| MSE ($\times 10^{-3}$) | 4.48 | 5.09 | 5.15 |
| $t_i$ (s) | 0.41 | 0.21 | 0.13 |
| $\kappa$ (cm$^2$/mW s$^{-1}$) | 0.109 | 0.027 | 0.039 |
| $D$ (cm$^2$/s) | 3.0 | 2.5 | 3.5 |
| $C$ (cm$^3$/mol) ($\times 10^{-12}$) | 3.1 | 5.5 | 6.1 |
| $\text{MSE}$ ($\times 10^{-10}$) | 0.32 | 1.72 | 1.26 |

*See also Fig. 3.
*Mean square error.
Here \( \rho(t) \) is the concentration of oxygen present in the material, and \( i_d \) specifies by how much the monomer concentration must be greater than that of the oxygen for polymerization to occur. The fraction \( U_0/i_d \) acts to define a threshold (time after onset of exposure), at which polymerization begins to take place. The polymerization rate in Eq. (27) becomes

\[
F(x,t) = κI_0[(1-B)\Theta[U_0/i_d-\rho(t)](1+V \cos(Kx))]A'(t),
\]

where \( κI_0[(1-B)\Theta[U_0/i_d-\rho(t)]=F_0(t)=κ_0(t)I_0'(1-B)\). 

In describing the behavior of the dye, we make three assumptions about the kinetics:

(a) The diffusion rate of oxygen within the material and from the atmosphere into the layer is much higher than the rates of diffusion of all other materials present. Therefore the oxygen is able to instantaneously maintain a constant distribution.

(b) After the reaction with the dye it is assumed that the active inhibiting oxygen becomes an inert component, referred to in relation (7) as the photoreduction product \( Y \), which has no further influence on the polymerization of the monomer.14

(c) The rate of change of the concentration of the active oxygen during exposure is assumed to be directly proportional to the total concentration of active oxygen.

From assumption (c), the differential equation governing the concentration of inhibiting oxygen can be written as

\[
\frac{d\rho(t)}{dt} = -\beta' \rho(t),
\]

where \( \beta'=k' I_0(1-B) \) is a constant that is proportional to the average recording intensity, where \( k' \) is a material constant at a fixed temperature. The solution to Eq. (31) is

\[
\rho(t) = ρ_0 \exp[-k' I_0(1-B)t],
\]

where \( ρ_0 \) is the initial oxygen concentration. We substitute this into Eq. (30), which is in turn substituted into Eq. (25) along with Eq. (28) giving a set of first-order coupled equations.2–6,14 Truncated equations, governing the monomer harmonic amplitudes, are obtained, i.e.,

\[
\frac{du_0(ξ)}{dξ} = -H(ξ)A(ξ)u_0(ξ) - \frac{1}{2}H(ξ)A(ξ)V_0u_1(ξ),
\]

\[
\frac{du_1(ξ)}{dξ} = -H(ξ)A(ξ)V_0u_0(ξ) - [H(ξ)A(ξ) + R
\]

\[
\times \exp[-aH(ξ)ξ]\cosh[aH(ξ)V_0ξ]u_1(ξ),
\]

where

\[
ξ = κI_0^{1/2} t, \quad R = \frac{DK^2}{f_0},
\]

\[
H(ξ) = \Theta \left[ \frac{U_0}{i_d} - ρ_0 \exp \left( - \frac{k'(1-B)}{κ} ξ \right) \right],
\]

\[
A(ξ) = 1 - T(ξ).
\]

We have assumed that \( D \) is the initial unchanged diffusion constant and \( κ=0.2–6 \).

The rate at which the concentration of polymer varies is proportional to the instantaneous rate of removal of monomer; thus \( N(x,t) \), the polymerized monomer concentration at time \( t \), is

\[
N(x,t) = \int_{t_i}^{t} F(x,t')u(x,t')dt'.
\]

We have observed that \( N(x,t=0) = 0 \), due to inhibition, gives the lower limit on the integral \( t_i \). While some photo-polymerization may occur when \( 0 < t < t_i \), it is assumed negligible; thus the step function provides a useful first-order description of the inhibition process. We substitute Eqs. (27) and (28) into Eq. (35) and write \( N(x,t) \) as

\[
N(x,t) = \sum_{i=0}^{2} N_i(t) \cos(iKx).
\]

For volume (thick) holographic gratings replayed on Bragg,24–26 it is the \( N_1 \) harmonic that diffracts the incident light. Therefore

\[
N_1(ξ) = \int_{0}^{ξ} H(ξ')A(ξ') \left[ Vu_0(ξ') + u_1(ξ') + \frac{1}{2} Vu_2(ξ') \right] dξ'.
\]

Coupled-wave theory predicts a relationship between the diffraction efficiency and the refractive index modulation of a sinusoidal thick transmission grating. Assuming that \( N_1 \) determines the amplitude of the grating refractive index modulation, \( Δn = CN_1 \) for some constant \( C \). The diffraction efficiency \( η(ξ) \) of the grating is

\[
η(ξ) = \sin^2 \left[ \frac{πd}{λ \cos θ} CN_1(ξ) \right],
\]

where \( d \) is the thickness of the material layer, \( λ \) is the wavelength of the probe–replay laser, and \( θ \) is the Bragg angle associated with that wavelength.28

We note that a much more complex relationship exists between the \( N_i \) and the \( Δn_i \) values,29–30 and that while Eq. (38) holds for thick volume gratings, in general a rigorous electromagnetic model is needed to calculate the diffraction efficiency.6

### 4. POLYMERIZATION-DRIVEN DIFFUSION SIMULATIONS

We wish to apply our new model to characterize our material behavior by fitting experimental growth curves. Before carrying out this procedure we wish to examine the general behavior predicted by our model.

First we examine how the initial inhibition period is affected by changes in the oxygen concentration and the exposing intensity. Combining Eqs. (29) and (32) we obtain
We can find $t_i$ using the experimentally reasonable parameter. Examining Eq. (39) it is clear that $t_i$ decreases as the concentration of oxygen present in the material decreases. By increasing the average exposure intensity $I_0$, the concentration of oxygen can be reduced at a quicker rate and this will result in a reduction of $t_i$. Equation (39) suggests that by using sufficiently strong exposure intensities, the presence of the initial dead band could be completely removed. Furthermore Eq. (39) can be used to determine the initial concentration of oxygen $\rho_0$, which is present in the material.

Second we examine the behavior of the harmonics of monomer concentration, and plots of the first three harmonics were generated. The results are plotted in Fig. 1 for an average exposure intensity of $I_0 = 3.5$ mW/cm$^2$. The following parameter values were used: $R = 1$, $V = 1$, $\alpha = 0$, $\kappa = 0.3$ cm$^2$ mWs$^{-1}$, and $k' = 0.1$ cm$^2$ mWs$^{-1}$. The initial monomer concentration was taken as $U_0 = 3.2$ g/l and $i_d = 100$. A value for the loss fraction was empirically found to be $B = 0.2 \pm 0.05$. The absorbance function $A(t)$ is defined using Eq. (24) and the transmittance parameter values used can be seen in case 1, Table 1. The initial oxygen concentration present in the layer was chosen to be $\rho_0 = 0.03$ g/l. This estimated value appears reasonable relative to the densities and volumes of the other materials present. We note the presence of the initial dead band in Fig. 1 with $t_i \sim 1$ s.

Figure 2 shows the predictions for the first harmonic of the polymer concentration $N_1$ for the cases when $R = 1$ and 10. These values were chosen to illustrate the experimentally observed range of $R$ values obtained. As can be seen, the larger the $R$ value, the larger the $N_1$ saturation value. Once again we note the presence of the inhibition period at the start of exposure due to inhibition.

5. EXPERIMENTAL RESULTS, NUMERICAL FITTING, AND PARAMETER ESTIMATION

There are three parts to this section. First, in Subsection 5.A, we attempt to examine the effects of exposure intensity changes on the inhibition period to verify the predictions made using Eq. (39). Second, in Subsection 5.B, we discuss preexposure and note that a minimum dead-band duration is found to exist independent of the exposing intensity. Finally, in Subsection 5.C, we examine the effects of cover plating (sealing) on the inhibition effects. In all cases standard materials and holographic exposing setups were used. We added 16 cm$^3$ of Methylene Blue of concentration $1.25 \times 10^{-3}$ M to the standard polyvinyl alcohol–acrylamide mix to produce dry layers of thickness $d \sim 100 \pm 10$ $\mu$m.

Fits to the experimental data presented in this section were performed using the model described in Section 3 applying the numerical fitting algorithm described in Refs. 1 and 6.

### A. Intensity Effects

Figure 3 contains three growth curves for three different average exposure intensities $I_{01} = 3.5$ mW/cm$^2$, $I_{02} = 4.5$ mW/cm$^2$, and $I_{03} = 5$ mW/cm$^2$. The gratings were all recorded using a He–Ne laser with $\lambda = 633$ nm, $\Lambda = 1$, $\mu$m, and probing at $\lambda = 532$ nm (Refs. 1 and 7) with the same setup as that used in Refs. 7 and 28. As expected, the rate of polymerization increases with increasing exposure intensity. It is also noticeable that there is a decrease in the inhibition period. The corresponding inhibition periods for the three intensities are $t_{i1} = 0.41$ s, $t_{i2} = 0.21$ s, and $t_{i3} = 0.13$ s, respectively. The results are in agreement with the predictions of Eq. (39). The dye molecules within the material are excited at a faster rate due to the increase in the intensity entering the material. This causes the oxygen present in the material to be removed at a quicker rate, as described by the rate equations in...
Section 2. This reduces the radical suppression, caused by the oxygen, and hence there is a decrease in the duration of the dead band caused by inhibition.

Since in Eq. (39) the $\beta' = k' I_{0}(1 - B)$ term is a function of the average exposure intensity $I_{0}$, we can estimate values for $i_d$ for different exposure intensities. From the experimental data presented in Fig. 3 we can extract values for the inhibition times; and by substituting reasonable values for $U_{0}$, $k'$, $\rho_{0}$, $I_{0}$, and $B$, we can estimate $i_d$. These values can then be used during the data fitting procedure.

Applying a numerical least-squares algorithm we obtain best fits to the experimental data for the mean squared errors (MSEs) shown in the lower part of Table 1. We fit by first substituting known parameter values, i.e., $U_{0} = 3.2 \text{ g/l}$, $i_{d} = 85$, $\rho_{0} = 0.03 \text{ g/l}$, $k' = 0.1 \text{ cm}^{2} \text{ mW}^{-1}$, and $B = 0.2$, into our equations and then interactively testing the fit quality over reasonable ranges of the unknown parameters. In this way estimates for $D$ (the diffusion coefficient), $\kappa$ (the polymerization rate constant), and $C$ are extracted. In Fig. 3 the solid curves are the theoretical fits to the data points. Typically the loss fraction is measured to have values in the range $B = 0.2 \pm 0.05$. We note that using values over this range does not appreciably vary the quality of the fits achieved (the calculated MSE values).

The resulting physical parameters are summarized in the lower part of Table 1. It can be seen that, as the exposure intensity is increased, the inhibition time decreases as expected. We note that the values for the diffusion constant $D$ are comparable with previous results, and the diffusion constant values of water and propanol reported recently. The $C$ values estimated are also comparable with previous results. As can be seen in the lower part of Table 1, the resulting MSE values are small, which is also evident in the quality of the fits in Fig. 3. However, the estimated values for the polymerization rate constant $\kappa$ vary significantly from case to case. This variation occurs due to the breakdown of the model. For example, all nonlocal material effects (temporal and spatial) have been neglected. A step function has been used as a first-order approximation to model the threshold effect created by the inhibition process. Furthermore the mechanism of chain termination is assumed to be homolecular ($\beta = 1$). Recently, we have indicated that better fits can be achieved when a primary termination mechanism is used ($\beta = 2$).

B. Minimum Dead Band and Preexposure

We have observed experimentally that increasing the exposure intensity above some fixed value results in no further discernible reduction in the dead band. Where we increased the average holographic exposure intensity above 6 mW/cm$^2$ in our experiments, no further reduction of the initial inhibition period took place. We will discuss some possible explanations for the existence of this minimum dead band in Subsection 5.C. But first we note that it would seem reasonable to expect that by preexposing the material, dye molecules will become excited and reduce the concentration of active oxygen causing radical suppression. At this point, with all inhibition processes (i.e., active oxygen) eliminated, it should be possible for uninhibited holographic recording to take place.

In Fig. 4 the grating refractive index modulation, extracted from three different diffraction-efficiency grating-growth curves, are presented. In each case the polymer layer was preexposed, prior to holographic exposure, by a uniform plane wave with a preexposure intensity of $I_{PE} = 1.5 \text{ mW/cm}^2$. The preexposed region on the plate was twice that of the area undergoing holographic grating exposure. This was done to ensure that there was no oxygen in the surrounding layer to diffuse into the area in which grating formation is taking place. The holographic exposure started almost immediately after the preexposure was complete, a typical delay of 0.5 s occurred, and in all cases a holographic exposure intensity of 3.5 mW/cm$^2$ was used. Curves 3, 2, and 1 are the growth curves of the gratings recorded with preexpose durations of 3, 2, and 1 s, respectively.

It can be seen in Fig. 4 that preexposure reduces the rate of polymerization. This may arise because preexposure bleaching means that less dye is available at the start of the holographic exposure. Importantly we note from Fig. 4 that the inhibition period does not seem to change significantly with the preexposure. This would appear to suggest that the active oxygen, which is removed before the start of the holographic exposure by the preexposure, is being rapidly replaced from the air above the plate. If oxygen is constantly and rapidly diffusing into the layer, then the suppression of radicals is continuously taking place and reducing the rate of photopolymerization at all times during the exposure.

Fits to the experimental data presented in Fig. 4 were obtained (solid curves). The values for the transmittance and material parameters used are those given in Subsection 5.A and Table 1 for the holographic exposure intensity, $I_{0} = 3.5 \text{ mW/cm}^2$. This intensity was chosen to allow the inhibition period to be easily observed. The resulting extracted physical parameters obtained for the data in Fig. 4 are summarized in Table 2. Once again disagreements between the experimental and theoretical results can be explained in terms of the deficiencies of our models.

C. Cover Plating

From the experiments described above we have identified the effects on the inhibition period and on the rate of polymerization of the average holographic exposure inten-
sity and of preexposure. However, to explain the continuing existence of a minimum dead band, which is observed in Fig. 4, we have proposed that a significant amount of oxygen is diffusing into the material from the air. In an effort to eliminate the effects of such diffusion, we cover plated the material using an index-matched fluid, silicon oil, and a glass plate. Figure 5 contains two growth curves. Curve (a) is for a sealed plate that has been pre-exposed for 1 s with a preexposure intensity of $I_{PE} = 1.5 \text{ mW/cm}^2$ and then, within 0.5 s, holographically exposed with an average exposure energy of $I_0 = 3 \text{ mW/cm}^2$. Curve (b) is for an unsealed plate, which was not preexposed. In both cases identical holographic exposure intensities, the inhibition period can be reduced to a minimum induction period value by a combination of preexposure and cover plating. This reduction is accompanied by an increase in the rate of polymerization, which arises due to the reduced concentration of unbleached dye, appears to be significantly less than the rate reduction arising due to the presence of oxygen.

For both sets of data, fitting is performed using the same material parameter values used in Fig. 3. However since $I_0 = 3 \text{ mW/cm}^2$, the transmittance parameters used are different: $E = 0.7656 \text{ mW/cm}^2$, $G = 1.45 \text{ mW/cm}^2$, $a_0 = 4.45 \times 10^{-3} \text{ s}^{-1}$, and $a_1 = 3.61 \times 10^{-4} \text{ s}^{-2}$ extracted from a fit with an MSE $= 4.01 \times 10^{-3}$. The loss fraction value used to produce the fits shown in Fig. 5 are (a) $B = 0.3$ and (b) $B = 0.2$. The reason for the difference in the loss fraction for the two cases is the change in the amount of light lost due to the extra reflective surfaces in the cover plate. Once again errors in the $B$ values ($\pm 0.05$) do not effect the quality of the resulting growth curve fits.

The resulting extracted physical parameters are summarized in Table 3. Clear differences between the numerical estimates of the various polymerization rates and inhibition periods are found. The agreement between the experimental data and theoretical fits can be seen in Fig. 5 and are quantitatively supported by the low MSE values presented in Table 3.

We believe that Fig. 5 illustrates most clearly the significance of effects studied in this paper. Comparing the two growth curves, it can be seen that for particular holographic exposure intensities, the inhibition period can be reduced to a minimum induction period value by a combination of preexposure and cover plating. This reduction is accompanied by an increase in the rate of polymerization.

While the values obtained for the diffusion rate $D$ and the constant $C$ are consistent with values presented in the literature, once again it can be seen that the polymerization rate constant values vary significantly. Once again we attribute this to the assumptions made in deriving our model and we are currently working to generalize our model.

### Table 2. Physical Parameter Values Obtained from Fits to the Data in Fig. 4

| Time (s) | $\rho_0$ (g/l) | $i_d$ | $t_i$ (s) | $\kappa$ (cm$^2$ mW$^{-1}$) | $D$ (cm$^2$/s) | $C$ (cm$^3$/mol) | MSE ($\times 10^{-6}$) |
|----------|----------------|-------|-----------|----------------------------|---------------|-----------------|---------------------|
| 1        | 0.0400         | 87.9  | 0.30      | 0.0100                     | 4.54          | 2.6             | 25                  |
| 2        | 0.0376         | 92.0  | 0.22      | 0.0070                     | 4.52          | 0.94            | 3.8                 |
| 3        | 0.0355         | 95.2  | 0.15      | 0.0067                     | 4.52          | 0.69            | 2.2                 |

Fig. 5. Fit to the experimentally obtained growth curve of refractive index amplitude versus exposure time for (a) cover-plated material layer preexposed for 1 s and (b) uncover-plated material layer without preexposure: theoretical fit (solid curves) and experimental data (dotted curves).
Experimentally we have explored the behavior of the inhibition period. We have shown that this initial dead-band period can be reduced, although not eliminated, using strong holographic exposing intensities. We have experimentally observed both a decrease in the inhibition period and an increase in the rate of polymerization using a combination of preexposures and cover plating to seal the dry layer. We have explained the minimum dead band as a photopolymerization induction period.

The model presented here has many limitations and these have been discussed. However, despite these drawbacks, we have succeeded in applying the model to provide plausible explanations for the observed results and to numerically extract reasonable estimates for several material parameters.

Much work remains to be done. One fundamental assumption made in this paper is that it is the reaction of oxygen with the excited dye molecules that leads to inhibition. It is also possible that inhibition may occur primarily at the radicalized monomer stage. While much work is still clearly needed in either of these cases, our simplified mathematical model, which stresses the suppression of chain initiation, will be of value.

The model for the time-varying absorption of the material also needs to be improved to include variations with depth. The theoretical models for inhibition and absorption must be integrated into the nonlocal polymerization-driven diffusion model for both nonlocal time and spatial responses. A more exact expression for the refractive index modulation such as the Lorentz–Lorenz equation should be included. Other effects such as the chain termination mechanisms and material shrinkage must be included to enable a more accurate physical picture to emerge. In particular we note that our use of a step function to model the inhibition period must be reexamined. However, given the variety of material in which inhibition effects and threshold effects have been observed, even the incomplete results presented here are significant for a wide range of applications, including holographic data storage, diffractive optical element fabrication, and photoembossing.

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|                  | $t_i$ (s) | $\kappa$ (cm$^2$/mW s) | $i_d$ | $\rho_0$ (g/l) | $D$ (cm$^2$/s) ($\times 10^{-11}$) | $C$ (cm$^2$/mol) ($\times 10^{-5}$) | MSE ($\times 10^{-5}$) |
|------------------|----------|------------------------|-------|---------------|-----------------------------------|-----------------------------------|-----------------------|
| Cover plated and preexposed (1 s) | 0.25     | 0.200                  | 92    | 0.0355        | 2.0                               | 6.06                             | 0.95                  |
| Not cover plated, no preexposure | 1.10     | 0.055                  | 120   | 0.0400        | 2.5                               | 2.04                             | 1.50                  |

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