
Theses and Dissertations

Spring 2010

Experimental and theoretical investigations of active center generation and mobility in cationic and free-radical photopolymerizations

Cynthia Caroline Hoppe
University of Iowa

Follow this and additional works at: <https://ir.uiowa.edu/etd>

 Part of the [Chemical Engineering Commons](#)

Copyright 2010 Cynthia Caroline Hoppe


This dissertation is available at Iowa Research Online: <https://ir.uiowa.edu/etd/516>

Recommended Citation

Hoppe, Cynthia Caroline. "Experimental and theoretical investigations of active center generation and mobility in cationic and free-radical photopolymerizations." PhD (Doctor of Philosophy) thesis, University of Iowa, 2010.

<https://doi.org/10.17077/etd.l9x6gylr>

Follow this and additional works at: <https://ir.uiowa.edu/etd>

 Part of the [Chemical Engineering Commons](#)

EXPERIMENTAL AND THEORETICAL INVESTIGATIONS
OF ACTIVE CENTER GENERATION AND MOBILITY
IN CATIONIC AND FREE-RADICAL PHOTOPOLYMERIZATIONS

by

Cynthia Caroline Hoppe

An Abstract

Of a thesis submitted in partial fulfillment
of the requirements for the Doctor of Philosophy
degree in Chemical and Biochemical Engineering
in the Graduate College of
The University of Iowa

May 2010

Thesis Supervisor: Professor Alec B. Scranton

ABSTRACT

Photopolymerization is considered an attractive alternative in many industries to traditional polymerization processes. The advantages of photopolymerization over other types of polymerization include elimination of heat sources, faster cure times, and reduction in the use of volatile organic solvents. Despite these environmental and cost-saving advantages, photopolymerizations have several limitations. Light attenuation can be a problem for systems containing pigments or fillers. The radiation source penetrates only to a shallow depth beneath the surface, limiting the thickness of strongly pigmented or filled coatings and films. Photopolymerization is also generally limited to systems with simple geometries that can be uniformly illuminated. Coatings on three-dimensional substrates, or other systems with complex geometries, are difficult to uniformly cure. These problems can be solved by “shadow cure,” which is defined as the reactive diffusion of photoinitiated active centers into regions of a polymer that are unilluminated. In this contribution, the generation and subsequent spatial and temporal evolution of the active center concentrations during illumination are described using the differential equations that govern the light intensity and photoinitiator concentration for polychromatic illumination. Reactive diffusion of the active centers during the post-illumination period is characterized, and shown to result in cure of unilluminated regions. A kinetic analysis is performed by coupling the active center concentration profiles with the propagation rate equation, yielding predicted cure times that are compared with experimental results. This analysis is used for the evaluation of cationic shadow cure in pigmented photopolymerization systems, and systems with complex geometries. The

extensive characterization of cationic systems is then applied to free-radical photopolymerization to examine the potential of shadow cure for active centers with much shorter lifetimes. An example of a free-radical photopolymerization system is presented in which the dimensional scales are small enough to utilize the short lifetimes of free-radical active centers for shadow cure. The results for both free-radical and cationic shadow cure indicate that the reactive diffusion of photoinitiated active centers may be used for effective cure in unilluminated regions of a photopolymer. This research will potentially allow photopolymerization to be used for applications in industries where it has never before been utilized.

Abstract Approved: _____
Thesis Supervisor

Title and Department

Date

EXPERIMENTAL AND THEORETICAL INVESTIGATIONS
OF ACTIVE CENTER GENERATION AND MOBILITY
IN CATIONIC AND FREE-RADICAL PHOTOPOLYMERIZATIONS

by
Cynthia Caroline Hoppe

A thesis submitted in partial fulfillment
of the requirements for the Doctor of Philosophy
degree in Chemical and Biochemical Engineering
in the Graduate College of
The University of Iowa

May 2010

Thesis Supervisor: Professor Alec B. Scranton

Graduate College
The University of Iowa
Iowa City, Iowa

CERTIFICATE OF APPROVAL

PH.D. THESIS

This is to certify that the Ph.D. thesis of

Cynthia Caroline Hoppe

has been approved by the Examining Committee
for the thesis requirement for the Doctor of Philosophy
degree in Chemical and Biochemical Engineering at the May 2010
graduation.

Thesis Committee: _____
Alec B. Scranton, Thesis Supervisor

Julie L. P. Jessop

Tonya Peeples

C. Allan Guymon

Ned Bowden

For my dearest family and friends, for providing mirrors into my soul.

ACKNOWLEDGMENTS

I would like to take this opportunity to thank the people without whom I would not have survived my graduate experience. First, I would like to thank Professor Alec Scranton, my academic advisor, for accepting me with open arms into his research group at my darkest hour. Photopolymerization was the best thing that ever happened to my graduate career, and I will be forever grateful for having had the opportunity to explore and contribute to such an exciting and cutting-edge field of research. Professor Scranton has become an invaluable mentor and friend, and could not have been more supportive and encouraging of me. I would not have learned a new research area in such a short time if it were not for his direction and creativity. I would also like to thank all of the past and present graduate students in our research group, as well as Chris Coretsopoulos, for helping me along the way.

Because of Professor Scranton's leadership, I have had the unique opportunity to work on several different research collaborations with industrial partners. I would like to thank all the collaborators I have worked with for their funding and support, as well as for the ideas and inspiration they have provided for this research.

I would like to thank the National Science Foundation and the University of Iowa Graduate College both for providing me with generous fellowship funding.

There are countless people to thank within the Chemical and Biochemical Engineering Department. I would like to say a special thanks to Associate Professors Julie Jessop and Tonya Peoples for being mentors and friends to me over the years. Thanks to Linda Wheatley for keeping me organized and on-track. All of my fellow graduate students who I have come

to know have touched my life in some way. I think I could name a student in every single one of our department research laboratories who has helped me learn an experimental or computational technique. I feel very fortunate to have found a home in such an open and collaborative department. I have had the opportunity to work with several talented undergraduate engineering students, including Amber Thiesen, Brandon Robson, Mike Baker, and Tim Hullermann. Thank you for your hard work and dedication.

I could not complete this list without saying a special thanks to my previous academic advisor, Professor John Wiencek, my first mentor at the University of Iowa who welcomed me the day I walked in the door. I appreciate his guidance and friendship during the first few years of my graduate experience. I will always be proud to have been a member of his group, and grateful for my experience working with protein expression, purification, and characterization.

Of course this would not be complete without thanking my family, starting with my parents who instilled in me a belief that I could achieve anything I set my heart and mind to. I would not be where I am today without their belief in me. I have cherished every day throughout my graduate experience because of my children, who provide me with the motivation to persevere. And finally, I must thank my soul mate and loving husband more than anyone, for believing in me, for having infinite patience with me, and for providing the biggest mirror of all.

TABLE OF CONTENTS

LIST OF TABLES.....	vii
LIST OF FIGURES.....	viii
CHAPTER 1 BACKGROUND AND SIGNIFICANCE	1
1.1 Introduction	1
1.2 Photopolymerization Background	1
1.3 Cationic Photopolymerization and Shadow Cure	4
1.4 A Case for Shadow Cure in Free-Radical Photopolymerizations.....	6
CHAPTER 2 OBJECTIVES.....	9
CHAPTER 3 CATIONIC PHOTOPOLYMERIZATION OF SYSTEMS PIGMENTED WITH CARBON BLACK NANOPARTICLES	11
3.1 Introduction	11
3.2 Experimental.....	13
3.2.1 Materials.....	13
3.2.2 Pigment Dispersion and Size Characterization.....	13
3.2.3 UV/Visible Spectroscopy	13
3.2.4 Determination of Diffusion Coefficient	14
3.2.5 Raman Spectroscopy for Characterization of the Propagation Rate Constant	15
3.2.6 Photopolymerization of Coatings.....	16
3.3 Results and Discussion	16
3.3.1 Photoinitiator, Photolysis Product, and Pigment Absorptivities	16
3.3.2 Determination of the Active Center Concentration Profiles Produced During Illumination.....	20
3.3.3 Post-illumination Diffusion of Active Centers in Pigmented Coatings	28
3.3.4 Kinetic Analysis for Prediction of Cure Times.....	31
3.4 Conclusions.....	36
CHAPTER 4 CATIONIC ACTIVE CENTER MOBILITY IN PHOTOPOLYMERIZATION SYSTEMS WITH COMPLEX GEOMETRIES.....	38
4.1 Introduction	38
4.2 Shadow Cure in Two Spatial Dimensions.....	40
4.2.1 Governing Equations	40
4.2.2 Results and Discussion	44
4.3 Shadow Cure Perpendicular to the Direction of Illumination.....	50
4.3.1 Materials and Experimental Methods	51

	4.3.2	Results and Discussion	52
4.4		Shadow Cure Through Multiple Coating Layers	60
	4.4.1	Experimental Methods.....	61
	4.4.2	Results and Discussion	63
4.5		Conclusions.....	65
CHAPTER 5		THE POTENTIAL OF FREE-RADICAL SHADOW CURE.....	67
	5.1	Introduction.....	67
	5.2	Selection of Photoinitiator and Light Source for a Photopolymerizable ACF	70
	5.2.1	Materials and Experimental Methods	70
	5.2.2	Results and Discussion	72
	5.3	Characterization of Shadow Cure	82
	5.3.1	Materials and Experimental Methods	82
	5.3.2	Results and Discussion	82
	5.4	Conclusions.....	94
CHAPTER 6		CONCLUSIONS AND RECOMMENDATIONS.....	95
	6.1	Cationic Photopolymerization of Systems Pigmented with Carbon Black Nanoparticles	95
	6.2	Cationic Active Center Mobility in Photopolymerization Systems with Complex Geometries	97
	6.3	The Potential of Free Radical Shadow Cure	98
REFERENCES.....			101

LIST OF TABLES

Table 3-1. Napierian molar absorptivities (L/mol-cm) for photoinitiator (ϵ_i) and photolysis products (ϵ_p) for incident wavelengths.....	17
Table 4-1. Results of shadow cure through multiple layers: bottom layer illuminated only.....	64
Table 4-2. Results of shadow cure through multiple layers: top layer illuminated only.....	64
Table 5-1. Overall conversions by FTIR for various visible light photoinitiators illuminated with green LED for one minute 8 cm from light source.	74

LIST OF FIGURES

Figure 3.1. DLS histogram with mean hydrodynamic radius = 29.2 nm.	19
Figure 3.2. Transmittance vs. concentration of CB-35 in propylene carbonate at 300nm. Effective absorptivity = 22.9 L/g-cm.	19
Figure 3.3. Profiles of the initial total light intensity summed over initiating wavelengths (295-308 nm) a) no pigment, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.....	23
Figure 3.4. Profiles of photoinitiator concentration after 2 minutes illumination. a) no pigment, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.....	24
Figure 3.5. Profiles of active center concentration for infinite thickness during 5 minutes of illumination. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.....	25
Figure 3.6. Profiles of active center concentration for 80 μm thick coatings during 5 minutes of illumination. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.	26
Figure 3.7. Profiles of active center concentration for 2 wt% CB-35 pigmented coatings ranging from 40 μm to infinitely thick. a) 2 minutes of illumination, b) 5 minutes of illumination. Monomer: EEC, Initiator: 1 wt% IPB.	27
Figure 3.8. Shadow cure distance versus square root of time for a 1 wt% carbon black pigmented system. Monomer: EEC, Initiator: 1 wt% IPB, Exposure Time: 5 min.	29
Figure 3.9. Active center concentration profiles diffusing post-illumination in an 80 μm thick pigmented coating. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 min.	30
Figure 3.10. Active center concentration profiles diffusing post-illumination in pigmented coating (40 μm thick). a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 3 min.	31
Figure 3.11. Active center profiles for various CB loadings at bottom interface of 80 μm thick coating. Monomer: EEC, Initiator: 1 wt% IPB, Pigment: 1-3 wt% CB-35, Exposure Time: 5 min.	33
Figure 3.12. Comparison of predicted with experimental cure times (t_{mpd}) at bottom interface of 40 μm and 80 μm thick coatings for varying illumination times. Monomer: EEC, Initiator: 1 wt% IPB, Pigment: 2 wt% CB-35.	35

Figure 3.13. Predicted cure times (t_{mpd}) at bottom interface of coatings pigmented with 1-3 wt% CB-35. a) 40 μm thickness, b) 80 μm thickness. Monomer: EEC, Initiator: 1 wt% IPB.	36
Figure 4.1. Active center concentration profile at the end of the illumination period. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 minutes.	45
Figure 4.2. Active center concentration profiles diffusing with increasing shadow cure time. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 min.	47
Figure 4.3. Geometrical configuration for shadow cure in two spatial dimensions. Rectangular cross-section of sample with depth z and width x	47
Figure 4.4. Active center concentration profiles in two spatial dimensions (x and z). a) end of illumination period, b) 2.5 minutes shadow cure, c) 5 minutes, d) 10 minutes, e) 25 minutes, and f) 50 minutes. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 minutes.	49
Figure 4.5. Schematic representation of shadow cure experiments used to measure cationic active center migration perpendicular to the direction of illumination.	50
Figure 4.6. Photo showing illuminated region and shadow cure region of cured epoxide coating on aluminum panel. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.	52
Figure 4.7. Experimental measurements of shadow cure distance into shaded region over 30 minutes post-illumination time. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.	53
Figure 4.8. Active center concentration profiles throughout the 80 μm coating depth throughout the 3 second illumination time.	54
Figure 4.9. Active center concentration profiles diffusing into the shadow region (x -direction) with increasing shadow cure time. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.	55
Figure 4.10. Cationic active center concentration as a function of time at various positions in the shadow cure region (x -direction). Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.	57
Figure 4.11. Experimental measurements of shadow cure distance into shaded region over 30 minutes post-illumination time. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.	58
Figure 4.12. Active center concentration profiles diffusing into the shadow region (x -direction) with increasing shadow cure	

time for a) 1 cm illuminated width, and b) 4 cm illuminated width. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.	59
Figure 4.13. Pictorial representation of the experimental setup for bottom layer only illumination. a) the bottom layer, containing monomer and photoinitiator, is illuminated. b) the second layer of monomer is added, which cures with no further illumination. Arrows indicate diffusion of active centers into unilluminated layer.....	62
Figure 4.14. Pictorial representation of the experimental setup for top layer only illumination. a) a layer of monomer only is applied to a substrate. b) a second layer of monomer containing photoinitiator is added and illuminated, resulting in cure of the entire film. Arrows indicate diffusion of active centers into unilluminated layer.	63
Figure 5.1. Flexible printed circuit assembly using heat-curable ACF.....	69
Figure 5.2. Flexible printed circuit assembly using proposed photopolymerizable ACF.	69
Figure 5.3. Absolute irradiance of 150 W xenon lamp and high intensity green LED compared with absorbance of polyimide film.	73
Figure 5.4. Eosin Y (spirit soluble) absorbance spectrum and molecular structure (0.001 wt% Eosin Y in methanol)	75
Figure 5.5. Direct reaction between Eosin Y spirit soluble and MDEA [56].....	75
Figure 5.6. Summary of FTIR conversion results for HEA films using Eosin Y/MDEA photoinitiator system with cumene hydroperoxide.	77
Figure 5.7. Real-time FTIR conversion during 2 minute illumination with 150 W xenon lamp.....	78
Figure 5.8. FTIR conversion for HEA films using 3-component Eosin Y photoinitiator system.	79
Figure 5.9. Optimization of 2-component Eosin Y system. Conversions calculated by FTIR. 1 minute illumination with 150 W Xe lamp.....	80
Figure 5.10. Real-time FTIR conversion during 1 minute illumination with 150 W xenon lamp. HEA monomer, Eosin Y photoinitiator system with and without cumene hydroperoxide.	81
Figure 5.11. Example of an electrode imbedded polyimide film with 100 x 100 μm electrode spacing.	83

Figure 5.12. Polarized light micrographs of HEA films containing optimized 2-component Eosin Y photoinitiator system. Illuminated for 1 minute with 150 W xenon lamp through a PET film with 50 x100 μm electrode spacing.	84
Figure 5.13. Polarized light micrograph showing cross-section of HEA film containing optimized 2-component Eosin Y photoinitiator system. Illuminated for 1 minute with 150 W xenon lamp through PET film containing 50x100 μm gold electrode spacing.....	85
Figure 5.14. Light intensity profile of a 3-component Eosin Y system initiated with a 150 W xenon lamp ($I_0 = 60 \text{ mW/cm}^2$ in the 425 - 575 nm spectral region).....	87
Figure 5.15. Photoinitiator concentration profile for a 3-component Eosin Y system initiated with a 150 W xenon lamp ($I_0 = 60 \text{ mW/cm}^2$ in the 425 - 575 nm spectral region).	88
Figure 5.16. Geometrical configuration for analysis of free-radical active center diffusion.	89
Figure 5.17. Post-illumination concentration profiles for active centers diffusing from illumination region (left side: 0-50 μm) to unilluminated region (right side: 50-100 μm) for initial concentration of 10^{-8} mol/L . a) diffusion only (Equation 5.5), b) diffusion and termination (Equation 5.7).....	92
Figure 5.18. Post-illumination concentration profiles for active centers diffusing from illumination region (left side: 0-50 μm) to unilluminated region (right side: 50-100 μm) for initial concentration of 10^{-6} mol/L . a) diffusion only (Equation 5.5), b) diffusion and termination (Equation 5.7).....	93

CHAPTER 1

BACKGROUND AND SIGNIFICANCE

1.1 Introduction

Radiation initiated polymerization of multifunctional monomers in the presence of a photoinitiator is one of the most efficient methods for producing highly crosslinked polymer materials. Photopolymerizable coatings and films have gained acceptance in many multi-million dollar industries. Some examples of applications for radiation curable coatings and films include protective coatings for plastics, metals, and woods such as flooring and cabinetry [1,2]; *in-situ* photopolymerizable bioadhesives such as dental sealants [3-5]; UV-curable ink-jet printing inks [6,7]; adhesives for flat panel displays and other microelectronics [8-10]; printed circuit boards and other optical lithography applications [11-13]; optical fibers and optical data storage such as CDs and DVDs [14,15]; and many others. In this chapter, the current state of technology for photopolymerizable coatings and films is described within the context of some applications of photopolymers, and the limitations to the use of photopolymerization for these applications are explored. The need to overcome these limitations forms the basis for the research that is presented in this contribution.

1.2 Photopolymerization Background

Photopolymerization can be defined as the effectively instantaneous conversion of a liquid resin into a solid, insoluble polymer by exposure to ultraviolet or visible radiation [16]. Radiation curing is considered an attractive alternative to traditional polymerization processes due to many advantages. Polymer coatings are typically produced using thermal polymerization, which uses heat to generate active centers that polymerize

the coating after a liquid monomer has been applied to a substrate. Industrial heat curing requires large amounts of energy to operate high temperature ovens for extended periods of time. Photopolymerization eliminates the need for heat curing, resulting in significant energy and cost savings. As an additional consequence of eliminating the thermal sources, heat sensitive substrates such as printed circuit boards and other electronic assemblies that can be produced by photopolymerization are not damaged by heat curing. Any heat buildup that may be caused by the radiation source in the photopolymer or substrate is minimal, and is quickly dissipated for thin film applications. Improved manufacturing efficiency also results from photopolymerization processes due to the comparatively short time scales required for radiation curing. Higher line speeds can be achieved and multiple curing steps can be performed in a single manufacturing line. Finally, the use of volatile organic compounds (VOCs) is reduced or eliminated because solvent-free resins are generally used in photopolymerizable formulations. Before environmental regulation began in the 1970s, many industrial coatings contained as much as 80% solvent, and large quantities of solvent were released into the atmosphere [14]. Again, manufacturing efficiency can be improved by eliminating the need for solvent vapor handling systems. These advantages make photopolymerization better for the environment than other types of polymerization processes.

Despite all of these advantages, photopolymerizations have some disadvantages that cause their use to be limited in some industries. Light attenuation can be a problem for coatings or films containing pigments or fillers. Pigments are often added to coatings to provide color or to hide the surface of the substrate, but can compete with the photoinitiator by absorbing the initiating light. For this reason, free-radical photopolymerizations of

pigmented coatings are typically limited to extremely thin applications, such as printing inks. The same light attenuation problem exists for other types of filled systems, such as nanocomposites [17].

Another barrier to the use of photopolymerization in many applications is the inability to cure systems with complex geometries. Photopolymerization is commonly used for curing coatings on two-dimensional substrates. However, when the need arises for coating substrates with complex geometries, traditional photopolymerization suffers several disadvantages. Photopolymerizations are typically dependent upon the generation of short-lived free-radicals that are sensitive to termination by oxygen. Oxygen reacts with free-radical active centers, prevents the polymerization from proceeding until all of the oxygen is consumed, and results in the formation of harmful peroxides and hydroperoxides. Oxygen inhibition can lead to incomplete polymerization, slow reaction rates, and tacky surfaces. Expensive methods are used to overcome this problem, such as blanketing a photopolymer system with an inert gas such as nitrogen.

Furthermore, free-radical photopolymerization reactions proceed only in the presence of UV irradiation, so the reactions terminate when the UV irradiation ceases. For these reasons, free-radical photopolymerizations fail to provide a practical and economical method for coating surfaces with irregular or intricate shapes. Some of the methods being developed to overcome these problems for curing coatings on three-dimensional objects include plasma curing [18], dual-cure systems [19-21], and robotic UV curing [22]. In this contribution, cationic photopolymerization is presented as a solution to some of these limitations to photopolymerization.

1.3 Cationic Photopolymerization and Shadow Cure

Cationic photopolymerization has been shown to overcome some of the limitations of the more common free-radical photopolymerization. Cationic photopolymerization is an alternative to free-radical photopolymerization offering several unique advantages. Photo-generated cationic active centers exhibit extremely long lifetimes. They are not sensitive to free-radical scavengers, such as oxygen, and do not terminate by a radical-radical termination mechanisms. For these reasons, the long-lived cationic active center lifetimes can result in photopolymerization reactions that proceed long after irradiation has ceased, until the monomer is consumed or the active centers become entrapped in the polymer matrix.

Despite these advantages over free-radical systems, the development of efficient cationic photopolymerization systems has been challenging. Cationic photoinitiators originally developed to generate cationic active centers, such as aryldiazonium salts, were very costly, had poor stability, and would produce inferior physical properties in the resulting polymer. Diaryliodonium salt photoinitiators and triarylsulfonium salt photoinitiators were developed in the 1980s in a joint patent between 3M and General Electric [23]. These cationic photoinitiators, commonly used today, are thermally stable over a wide range of temperatures, absorb light very efficiently with an optical yield of approximately 0.7, are inexpensive to manufacture, and result in improved polymer physical properties. The commercial availability of these newer cationic photoinitiators has spurred research into the kinetics, mechanisms, and physical properties of cationic photopolymerization over the past two decades. Cationically polymerizable monomers were subsequently developed to provide reaction rates and physical properties that could rival those of free-radically polymerized

monomers. Vinyl ethers typically have high polymerization rates, but can result in runaway reactions. Epoxide reactions are easily controlled, but the polymerization rate is slow in comparison to vinyl ethers and acrylates. Finally, high ring-strain epoxides, such as 3,4-epoxy-cyclohexylmethanyl 3,4-epoxy-cyclohexanecarboxylate (EEC) have become an industrial standard for their high reactivity and excellent physical properties. Cationic photopolymerization can also be used for other important classes of monomers, such as oxetanes and siloxanes. Many cationically polymerizable monomers exhibit exceptional physical properties, such as clarity, adhesion, abrasion resistance, and chemical resistance. Some cationic photopolymers exhibit less shrinkage than their free-radical counterparts, such as acrylates.

Several investigators have shown cationic active centers to lead to dark cure, or post-polymerization [24-26]. Due to their long lifetimes, cationic active centers can also be responsible for a process known as shadow cure, or cure in regions that have never been illuminated [27]. The cationic active center lifetimes are long enough for diffusion to take place, either during the illumination period or after the illumination has ceased. This mobility of the cationic active centers facilitates cationic cure in recessed or shadow areas of a substrate that have not been directly exposed to irradiation. This contribution provides a theoretical and experimental investigation of cationic shadow cure in pigmented systems and in systems requiring cure in multiple spatial dimensions. In the period of time during which the photopolymer is illuminated, the cationic active centers are preferentially generated at the surface to produce a sharp concentration gradient that provides a driving force for diffusion into regions of unexposed monomer. An analysis of the active center diffusion that takes place post-illumination is used to characterize shadow cure in these systems. Cationic

active center mobility is shown to have the potential to make photopolymerization possible in some applications where it has not been commonly utilized.

1.4 A Case for Shadow Cure in Free-Radical Photopolymerizations

Free-radical active centers have significantly shorter lifetimes than cationic active centers, and tend to terminate immediately upon cessation of the illuminating light source. However, some cutting edge applications for photopolymers may require the development of shadow cure in free-radical systems. Advances in microelectronics have led to an increase in demand for circuits with increasingly small pitch, or spacing between interconnects. Despite the extremely short lifetimes of the free-radical active centers, mobility of free-radicals within the increasingly small dimensional scales required by microelectronics is becoming a possibility.

One application in which free-radical shadow cure in photopolymerizations may become feasible is polymer-based conductive adhesives, which are substitute for lead-based solders [9]. Electrically conductive adhesives provide an environmentally friendly solution for interconnections in many current electronics applications. Isotropic conductive adhesives (ICAs) are heat-curable materials containing an isotropic concentration of conductive particles, typically silver-filled epoxides that allow current to flow in all directions through the cured polymer. These adhesives can be used to electrically interconnect non-solderable substrates such as ceramics or plastics, or to replace solder for thermally-sensitive components. A second type of electrically conductive adhesive is anisotropic conductive adhesive, which allow current to flow along a unidirectional axis [8]. This type of adhesive is a composite containing fine electrically

conductive particles uniformly dispersed in a polymer matrix designed to provide electrical interconnection only at planned sites where the particles come into contact with a conductive substrate. The most common form these adhesives is a heat-curable, thermoplastic, anisotropic conductive film (ACF) used in flexible printed circuits. Ninety percent of all ACFs are currently sold for use in flat panel displays such as LCDs [10]. They are also used in flexible printed circuits, in which electronic circuits are mounted on flexible plastic substrates. Flexible printed circuits are being used in many applications, such as electronic books and other forms of electronic paper, displays for cameras and cell phones, and computer keyboards.

In this contribution, one possible formulation that could be used in a photopolymerizable ACF assembly is presented in which free-radical shadow cure is proposed as a method for attaining full polymerization in unilluminated regions, behind electrodes and around conductive particles. Considerations for the selection of a specific formulation for this application are explored, and light sources required for this application are evaluated. The motivation behind this research is to eliminate the heating step traditionally required for curing a conductive adhesive film. The advantages of replacing the heat-curing process with photopolymerization in this type of microelectronics application include protecting the heat-sensitive circuits from elevated temperatures, minimizing cure times, reducing energy usage, and eliminating the use of VOCs.

Aside from the specific scenario for free-radical shadow cure outlined above, applications have generally not yet been developed in which diffusion of free-radical active centers could be used to cure unilluminated regions of a photopolymer. In typical applications currently employing photopolymerization for microelectronics, such as printed circuit boards and other

types of microlithography, the goal is usually to attain high resolution. Deep UV wavelengths are utilized in these industries to enhance resolution, and diffusion of active centers presents a limitation rather than an asset. Therefore, the concept of free-radical shadow cure is extremely novel, has likely never before been utilized, and represents a great potential for the use of photopolymerization in the cutting-edge field of microelectronics.

CHAPTER 2

OBJECTIVES

The previous chapter illustrates that there is a compelling motivation for the development of shadow cure in photopolymerizations. The mobility of active centers can overcome many of the current limitations to the use of photopolymerization in some industries. This research addresses this need, and provides an increased fundamental understanding of the diffusional and kinetic mechanisms of shadow cure. The hypothesis of this research is that the mobility of active centers can lead to polymerization in regions of a photopolymer that have not been illuminated.

The broad objective of this research is to utilize a fundamental set of differential equations that govern the light intensity gradient and photoinitiator concentration gradient for polychromatic illumination to describe the spatial and temporal evolution of active center concentration profiles generated during illumination. These profiles of the active center concentrations are then used to characterize the reactive diffusion process which takes place during the post-illumination period. Consequently, a fundamental understanding of the active center migration into unilluminated regions is attained.

This broad objective is achieved by accomplishing the following three specific goals:

- 1) to demonstrate the generation and mobility of cationic active centers in systems pigmented with carbon black, specifically to understand the post-illumination diffusion of the active centers to depths greater than the light penetrates, and to support with

experimental results the idea that polymerization can occur in regions that have not been illuminated.

2) to illustrate the effects of shadow cure, or post-illumination diffusion, of cationic active centers in multiple spatial dimensions for the purpose of utilizing shadow cure for systems with complex geometries, again supporting with experimental results the hypothesis that shadow cure can result in polymerization of unilluminated regions.

3) to investigate the potential of free-radical shadow cure, specifically the ability of free-radical active centers to migrate post-illumination over the short dimensional scales required for some microelectronics applications, resulting in free-radical polymerization of unilluminated regions.

CHAPTER 3

CATIONIC PHOTOPOLYMERIZATION OF SYSTEMS PIGMENTED WITH CARBON BLACK NANOPARTICLES

3.1 Introduction

Photopolymerization is well established as an effective method for curing transparent films. However, the need for pigmented coatings has limited the use of photopolymerization in many applications. Pigments are often added to coatings to provide color or to hide the surface of the substrate. Photopolymerization reactions are hindered by pigments and fillers, which directly compete with the photoinitiator for incident photons [28]. This results in increased light attenuation in pigmented coatings, especially with increasing film thickness. For this reason, free-radical photopolymerizations of pigmented coatings are typically limited to very thin applications, such as inks ($\sim 10 \mu\text{m}$), with the thickest systems approaching $100 \mu\text{m}$ [29,30].

Unlike free-radicals, cationic active centers are not inhibited by oxygen and are essentially non-terminating, and therefore have been shown to remain active long after irradiation has ceased and may lead to further polymerization in the illuminated region (dark cure). Dark cure in cationic photopolymerizations of epoxide monomers has been characterized by a number of investigators [24-26]. Due to their long lifetimes and tendency to diffuse, cationic active centers can also be responsible for “shadow cure” in regions that have never been illuminated such as unexposed depths and regions shaded by opaque constituents or fillers. In a recent contribution, Ficek *et al.* [27] characterized the diffusion of the cationic active centers in photopolymerizations of cycloaliphatic epoxides, and demonstrated that the cure can continue for several hours to extend deep below the illuminated

surface in unpigmented systems. Sangermano and collaborators [17,21] recently reported hybrid cationic photopolymerization/sol-gel reaction processes to produce coatings containing nano-scale inorganic domains. In this clever approach, the inorganic domains are produced *in situ*, after the UV curing step, and the investigators demonstrated that coatings with desirable scratch resistance, toughness, clarity, and refractive index could be obtained.

This contribution provides a theoretical and experimental investigation of cationic photopolymerizations of epoxide coatings pigmented with carbon black nanoparticles. The fundamental differential equations describing the polychromatic photoinitiation process [31,32] are solved to obtain profiles of the concentration of active centers as a function of time and depth. Here the differential reaction/diffusion equations which describe the consumption of photoinitiator and production of active centers are coupled to the differential absorption equation which accounts for the polychromatic absorption by all system components, including the pigment. During the illumination period the active centers are preferentially generated at the surface to produce a concentration gradient that leads to diffusion into the regions of unexposed monomer. Using the active center concentration gradient at the end of the illumination period as the initial condition for the differential diffusion equation allows the active center concentration profile to be determined long after the illumination has ceased. Coupling these concentration profiles with the propagation rate equation allows the cure time, or time for macroscopic property development, to be predicted for a required conversion. In this manner, experimental cure times for carbon black pigmented coatings are compared with theoretical predictions of cure time.

3.2 Experimental

3.2.1 Materials

The cationically polymerizable monomer 3,4-epoxy-cyclohexylmethanyl 3,4-epoxy-cyclohexanecarboxylate (EEC, Sigma Aldrich) was used in these experiments. The photoinitiator used in these experiments was (tolycumyl)iodonium tetrakis (pentafluorophenyl) borate (IPB, Secant Chemicals). The carbon black pigment studied was CB-35 (NIPex 35, Degussa Engineered Carbons, LP). Methanol and propylene carbonate solvents were used for UV/Visible spectroscopy.

3.2.2 Pigment Dispersion and Size Characterization

The carbon black used in this study is a commercial furnace black designed for chemically prepared toner applications because of its ease of dispersion. Furnace blacks generally exhibit a hydrophobic, non-polar, basic character. Propylene carbonate was selected as a solvent for carbon black spectral measurements because it is a polar aprotic solvent in which the pigment exhibits good dispersion. The size and monodispersity of CB-35 carbon black in propylene carbonate was characterized using dynamic light scattering (DLS). Experiments were carried out with a DynaPro 99P instrument (Protein Solutions) equipped with a 1 cm pathlength cell. Samples were analyzed using the Dynamics software provided with the instrument.

3.2.3 UV/Visible Spectroscopy

The absorbance spectra for the monomer, photoinitiator, photolysis products, and pigments were determined in one nanometer increments using an 8453 UV-Visible spectrophotometer (Agilent Technologies). For the

monomer and photoinitiator, the spectra were obtained for dilute solutions (10^{-2} M and 10^{-3} M respectively) in methanol placed in an air-tight, quartz cell to prevent any changes in concentration due to evaporation of the solvent. To obtain the absorbance spectra after photolysis, the photoinitiator samples were illuminated with a 200 Watt Hg-Xe arc lamp (Oriel Light Sources) until there was no change in the absorbance spectrum. For carbon black spectra, the pigment was dispersed in propylene carbonate (~ 1 g/L) by mixing on a stir plate for at least 24 hours prior to analysis.

3.2.4 Determination of Diffusion Coefficient

A method previously documented for determining the diffusion coefficient for cationic active centers in cycloaliphatic diepoxide systems [27] was used to measure distance over which shadow cure occurred for a carbon black pigmented system, and to calculate the effective diffusion coefficient. Solutions containing 98 wt% EEC, 1 wt% IPB, and 1 wt% pigment were mixed together for 24 hours in dark conditions. The monomer mixture was placed in disposable 4.5 ml polystyrene cuvettes, which were chosen because they are transparent to the wavelengths of interest and readily dissolve in a number of solvents. Each cuvette, filled to a level of 3 cm with monomer solution, was illuminated from underneath with the light from the 200 W Hg-Xe arc lamp for 5 minutes. Since the density of the polymer is higher than that of monomer, illumination from below avoids polymerization-induced convection or mixing. After exposure, the system was maintained at 50 °C for the prescribed shadow cure time. At the prescribed shadow cure time, the sample was placed in THF to dissolve the cuvette and monomer from the uncured region of the sample. The insoluble polymer matrix was washed with acetone to remove any remaining THF and excess monomer. The polymer

sample was dried thoroughly and its weight was recorded. The polymerized thickness was determined by dividing the weight of the polymer sample by the product of the polymer density and the area of illuminated surface (the cross-sectional area of the cuvette, 1 cm²). At each shadow cure time, an unilluminated control sample was prepared to verify that thermally-induced polymerization did not occur.

3.2.5 Raman Spectroscopy for Characterization of the Propagation Rate Constant

The propagation rate constant for cationic polymerization of the cycloaliphatic diepoxide used in this study was determined independently for the neat monomer using Raman Spectroscopy. Raman spectra were collected using a holographic fiber-coupled stretch probehead (Mark II, Kaiser Optical Systems, Inc.) attached to a modular research Raman spectrograph (HoloLab 5000R, Kaiser Optical Systems, Inc). A sample containing EEC monomer with photoinitiator was placed inside a sealed 1 mm ID quartz capillary tube. A 200 mW 785 nm near-infrared laser through a 10x non-contact sampling objective with 0.8 cm working distance was directed into the sample to induce the Raman scattering effect. Photopolymerization was initiated by simultaneously illuminating the sample with a 100 W high pressure mercury lamp (Acticure Ultraviolet/Visible Spot Cure System, EXFO Photonic Solutions, Inc.) filtered for 250 – 450 nm wavelengths. The Raman peak at 790 cm⁻¹ was used to determine the epoxide conversion [33]. The effective propagation rate constant, k_p , was determined using a previously published procedure [34]. The value of k_p for EEC was determined to be 0.1 L/mol-sec, which is consistent with literature reported k_p values for ring-opening polymerizations [35].

3.2.6 Photopolymerization of Coatings

The time required for the CB-35 pigmented coatings to reach macroscopic property development was investigated over a range of pigment loadings and illumination times. For these experiments, solutions containing 96 - 99 wt% EEC, 1 wt% IPB, and 0-3 wt% pigment were mixed together for 24 hours in dark conditions. The solutions were then spread onto aluminum substrates using a draw bar to achieve the desired uniform coating thickness (40 or 80 μm). The coated panels were then illuminated for various times, using a 200 W Hg-Xe arc lamp. The output of the lamp was passed through a water filter to eliminate infrared light, resulting in an overall irradiance of 50 mW/cm^2 . The wavelength range of interest was determined to be 295-307 nm, corresponding to the overlap between the photoinitiator absorbance spectrum and the lamp emission spectrum. The irradiance in this range was determined to be 5 mW/cm^2 , measured using a calibrated miniature fiber optic spectrometer (USB4000, Ocean Optics, Inc.) The photopolymerization was carried out under atmospheric conditions and at room temperature. After exposure, the panels were stored at room temperature. The cure time required for macroscopic property development was determined by characterizing the surface tack and the adhesion to the substrate at regular intervals. Once full property development was achieved, the thickness of the coating was obtained by a micrometer (micro-TRI-gloss μ , BYK Gardner).

3.3 Results and Discussion

3.3.1 Photoinitiator, Photolysis Product, and Pigment Absorptivities

A number of investigators have shown that free-radical photoinitiators typically exhibit significant photo-bleaching [36-38] since the absorption of

the photolysis products is lower than that of the original photoinitiator. Photobleaching is important for photo-curing of thick systems since it allows light to penetrate deeper beneath the illuminated surface upon production of active centers [39-43]. Photobleaching during cationic photopolymerizations has received considerably less attention in the literature, and the degree of photobleaching of most cationic photoinitiators has not been characterized. Table 3-1 shows the molar absorptivity as a function of the incident wavelength (for the wavelengths of interest in this study) for the both the original cationic photoinitiator (IPB) and the photolysis products. This table illustrates that, although the IPB photoinitiator exhibits some photobleaching upon photolysis, the molar absorptivity of the photolysis products is significant (average 34% of the photoinitiator absorptivity).

Table 3-1. Napierian molar absorptivities (L/mol-cm) for photoinitiator (ϵ_i) and photolysis products (ϵ_p) for incident wavelengths

λ (nm)	ϵ_i	ϵ_p	ϵ_p/ϵ_i
295	2,736	1,046	0.38
296	2,537	936	0.37
297	2,352	839	0.36
298	2,175	753	0.35
299	2,005	677	0.34
300	1,844	610	0.33
301	1,688	549	0.33
302	1,544	496	0.32
303	1,409	450	0.32
304	1,282	410	0.32
305	1,164	373	0.32
306	1,054	342	0.32
307	954	315	0.33

The optical properties of the pigments play an important role for the potential photopolymerization of pigmented coatings. In general, pigments may reduce light penetration by absorption, scattering or reflection of the incoming light, therefore the radiative flux in a pigmented sample can be characterized by accounting for these effects. This has been accomplished for highly reflective pigments, such as titanium dioxide, by describing the specular and diffuse reflectance using a four-flux model [44]. In one study of Pigment Red 254, Jahn and Jung [45] found the reflectance contribution to be negligible for pigment particles smaller than 100 nm. Similarly, for a matrix containing carbon black pigments, Tesfamichael *et al.* [46] found that the contribution of the reflectance was insignificant for incident wavelengths below 500 nm. The system under investigation meets both of these criteria since the mean particle size is less than 30 nm and the initiating light falls in a narrow wavelength region where the UV photoinitiator absorption overlaps with the 200 W Hg-Xe arc lamp emission (295-308 nm).

To confirm the particle size distribution of the carbon black nanoparticles used in this study, dilute solutions of CB-35 in propylene carbonate solvent were evaluated using DLS. These results showed the samples to exhibit a monodisperse and unimodal distribution of particle sizes. The DLS histogram shown in Figure 3.1 gives a mean hydrodynamic radius of 29.2 nm.

These observations suggest that the reflectance and scattering effects can be neglected for the small, monodisperse carbon black particles used in this study. To confirm that the absorption is linearly dependent upon the CB-35 pigment loading, the direct transmittance in the wavelength range of interest (295-308 nm) was measured as a function of the pigment mass concentration. A plot of the transmittance at 300 nm as a function of CB-35

concentration in propylene carbonate is shown in Figure 3.2. The effective absorptivity was found to be 22.9 L/g-cm. This value for the pigment absorptivity remains constant over the incident wavelength range, and was used throughout the remainder of this chapter.

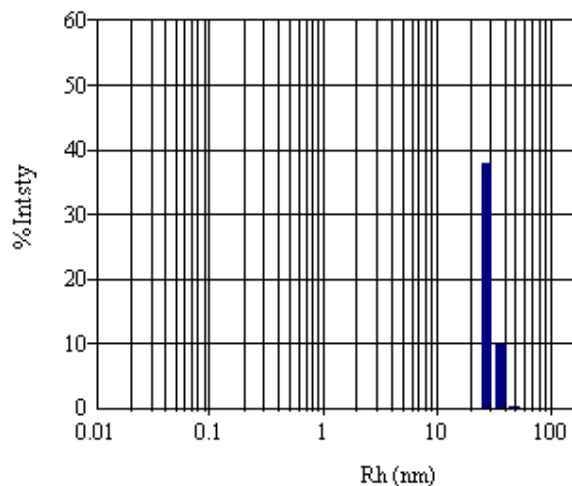


Figure 3.1. DLS histogram with mean hydrodynamic radius = 29.2 nm.

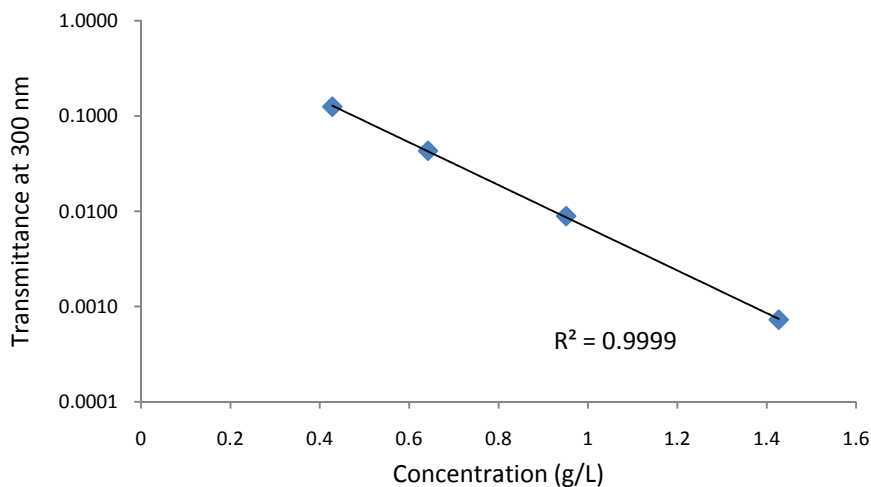


Figure 3.2. Transmittance vs. concentration of CB-35 in propylene carbonate at 300nm. Effective absorptivity = 22.9 L/g-cm.

3.3.2 Determination of the Active Center Concentration Profiles Produced During Illumination

An accurate description of the spatial photoinitiation profiles produced during the illumination step is necessary to predict the depth of cure in a pigmented system. The evolution of the light intensity gradient and the corresponding active center concentration profiles were found using the following set of differential equations for polychromatic illumination, including diffusion of the initiator and photolysis products:

$$\frac{\partial C_i(z, t)}{\partial t} = -\frac{C_i(z, t)}{N_A h} \sum_j \left(\frac{\varepsilon_{ij} \phi_j I_j(z, t)}{v_j} \right) + D_i \frac{\partial^2 C_i(z, t)}{\partial z^2} \quad (3.1)$$

$$\frac{\partial C_p(z, t)}{\partial t} = \frac{C_i(z, t)}{N_A h} \sum_j \left(\frac{\varepsilon_{ij} \phi_j I_j(z, t)}{v_j} \right) + D_p \frac{\partial^2 C_p(z, t)}{\partial z^2} \quad (3.2)$$

$$\frac{\partial I_j(z, t)}{\partial z} = -[\varepsilon_{ij} C_i(z, t) + \varepsilon_{pj} C_p(z, t) + a_{CBj} C_{CB} + A_m] I_j \quad (3.3)$$

Here, z is the direction perpendicular to the illuminated surface, with $z=0$ at the illuminated edge. The subscript j is an index with a different value for each wavelength of light under consideration; $C_i(z, t)$ is the initiator molar concentration at depth z and time t ; $C_p(z, t)$ is the photolysis product molar concentration at depth z and time t ; $I_j(z, t)$ is the incident light intensity of a specific wavelength at depth z and time t with units of mW/cm^2 ; ε_i is the initiator Napierian molar absorptivity at a specific wavelength with units of $\text{L}/\text{mole}\cdot\text{cm}$; ε_p is the photolysis product Napierian molar absorptivity at a specific wavelength with units of $\text{L}/\text{mole}\cdot\text{cm}$; a_{CB} is the carbon black pigment Napierian absorptivity at a specific wavelength with units of $\text{L}/\text{g}\cdot\text{cm}$; C_{CB} is the carbon black pigment mass concentration in units of g/L ; ϕ_i is the quantum yield of the initiator at a specific wavelength, defined as the fraction of absorbed photons that lead to fragmentation of the initiator; N_A is

Avogadro's number; h is Plank's constant; ν is the frequency of light in units of inverse seconds; D_i is the diffusion coefficient of the initiator in units of cm^2/sec ; D_p is the diffusion coefficient of the photolysis products; and A_m is the absorption coefficient of the monomer and the polymer repeat unit with units of $1/\text{cm}$. Note that the Napierian absorptivities are used because they are most natural for the differential version of the absorption equation (Equation 3). The quantum yield for IPB is 0.7 and the diffusion coefficients are $1 \times 10^{-7} \text{ cm}^2/\text{sec}$ [47].

For a polymerization system of thickness z_{max} which is illuminated at the planar surface in which $z = 0$, the following initial and boundary conditions apply [31,32]:

$$C_i(z,0) = C_o \quad (3.4)$$

$$C_p(z,0) = 0 \quad (3.5)$$

$$\frac{\partial C_{i,p}}{\partial z} = 0 \text{ at } z = 0 \text{ and } z = z_{max} \quad (3.6)$$

$$I(0, t) = I_o \quad (3.7)$$

Equation 3.4 states that the initial initiator concentration, C_o , is uniform throughout the depth of the sample. Similarly, Equation 3.5 indicates that the initial photolysis product concentration is zero throughout the sample. Equation 3.6, the no-flux boundary condition, indicates that there is no transport of initiator or photolysis product across the illuminated surface or the opposite boundary (typically an interface with a substrate). Finally, Equation 3.7 states that the light intensity on the illuminated surface is constant and equal to the initial intensity, I_o .

3.3.2.1 Active Center Concentration Profiles for Infinitely Thick Systems

To demonstrate the effect of the carbon black pigment on the photoinitiation process, it is useful to examine simulation results for an infinitely thick system where $z_{max} = \infty$. Simultaneous solution of Equations 3.1-3.7 yields profiles of light intensity and initiator concentration as functions of depth at various instants in time for an infinitely thick system. Figure 3.3 contains plots of the light intensity as a function of depth for two different EEC systems: unpigmented (Figure 3.3a) and pigmented with 3 wt% CB-35 (Figure 3.3b). The figure illustrates that the presence of the carbon black has a marked effect on the initial light intensity gradient in the sample. In the pigmented case, the light intensity drops to a value of essentially zero in less than 60 μm . In contrast, for the unpigmented case the initial light intensity is still 75% of the incident value at a depth of 60 μm , and retains more than 10% at a depth of 500 μm . The photobleaching described in Table 1 leads to the change in the gradient with increasing illumination time for the unpigmented system (Figure 3.3a), but is found to be negligible for the pigmented system (Figure 3.3b) due to the strong absorption by the pigment.

Since the rate of consumption of the photoinitiator at a given depth increases with increasing total light intensity (as described in Equation 3.1), the initial rate is highest at the illuminated surface, and is zero anywhere in which the total light intensity is zero. Therefore, a photoinitiator concentration gradient will be established immediately upon illumination, and will evolve with time in a manner described by the simultaneous solution of Equations 3.1-3.7.

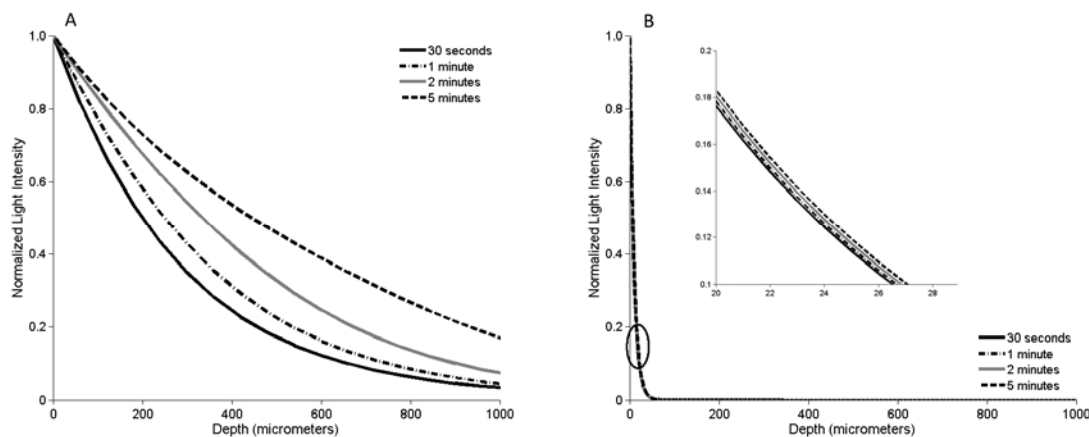


Figure 3.3. Profiles of the initial total light intensity summed over initiating wavelengths (295-308 nm) a) no pigment, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.

For example, Figure 3.4 shows profiles of the photoinitiator concentration as a function of depth with increasing illumination time for the cationic photopolymer system with and without 3 wt% CB-35 pigment. These results illustrate that in the case of the pigmented system (Figure 3.4b), the photoinitiator is depleted rapidly at the surface of the sample where the light intensity is highest, but the photoinitiation reaction does not extend much beneath the surface due to the light attenuation caused by the strongly absorbing pigment. As a result, the pigmented system exhibits sharp gradients in both the light intensity (Fig. 3.3b) and the initiator concentration (Figure 3.4b). Compared to the unpigmented system (Figure 3.4a), the steep concentration gradient results in a stronger driving force for diffusion of the initiator and photolysis products, therefore the diffusive contributions during illumination are much more important for the pigmented systems. Diffusion of the photoinitiator during the illumination

period is responsible for the concentration change that takes place at depths where the light intensity is zero (depths greater than 60 μm).

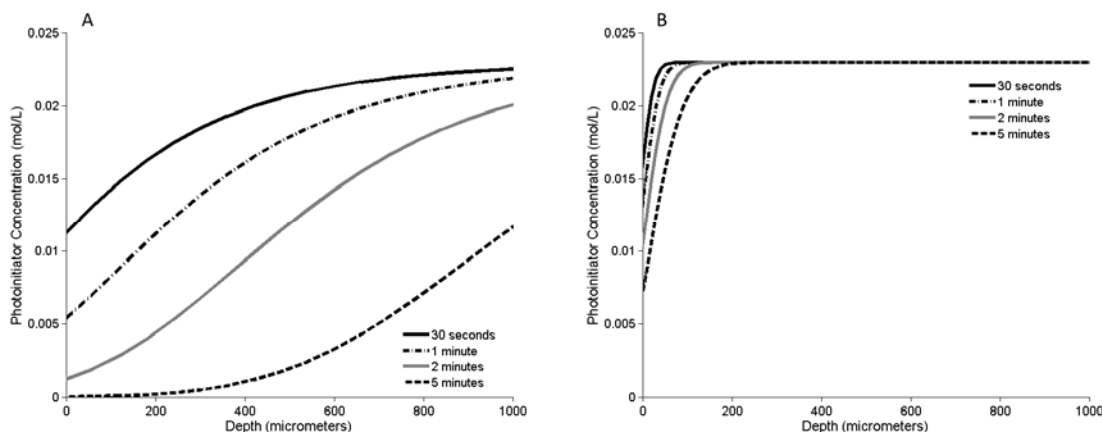


Figure 3.4. Profiles of photoinitiator concentration after 2 minutes illumination. a) no pigment, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.

The local rate of active center generation is equal to the product of the local initiator concentration and the local light intensity summed over the initiating wavelengths [31,32]. Since the cationic active centers are essentially non-terminating, and each photoinitiator molecule leads to the formation of a single active center molecule, the cationic active center concentration, C_{AC} , at a given depth, z , and time, t , can be determined from the integrated form of the rate equation:

$$C_{AC}(z,t) = \int_0^t C_i(z,t) \sum_j [I(z,t)]_j \phi_j \varepsilon_{ij} dt \quad (3.8)$$

Figure 3.5 shows the evolution of the active center concentration profiles during the 5 minute illumination time for two different pigment loadings. The 3 wt% pigmented system in Figure 3.5b shows the active center concentration profiles within the first 300 μm of an infinitely thick

system, resulting from the light intensity shown in Figure 3.3b and the photoinitiator concentration shown in Figure 3.4b. These results are compared with the active center concentration profiles for a 2 wt% pigmented system shown in Fig. 3.5a. The concentration of active centers generated at the illuminated surface of the sample is slightly higher for the 2 wt% system. The concentration drops off quickly to a value of zero within the first $\sim 200 \mu\text{m}$ of sample depth for the 3 wt% pigmented system.

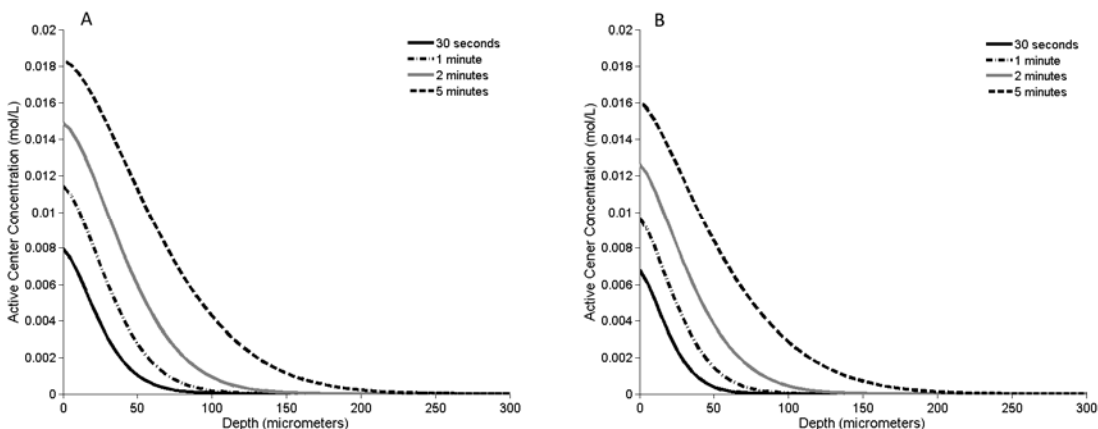


Figure 3.5. Profiles of active center concentration for infinite thickness during 5 minutes of illumination. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.

3.3.2.2 Active Center Concentration Profiles for Pigmented Coatings

For coatings of a finite thickness, the active center concentration profiles predicted during the illumination period differ from those for an infinitely thick system due to the no-flux boundary condition at the interface between the coating and the substrate. Figure 3.6 shows the active center concentration profiles throughout the depth of an $80 \mu\text{m}$ thick coating during 5 minutes of illumination for 2 wt% CB-35 (Figure 3.6a) and 3 wt% CB-35 (Figure 3.6b).

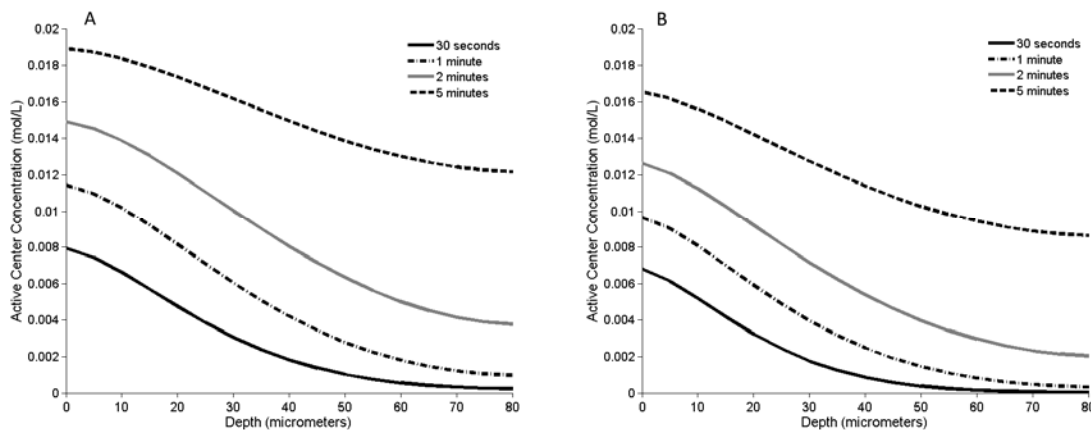


Figure 3.6. Profiles of active center concentration for 80 μm thick coatings during 5 minutes of illumination. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB.

In contrast to the infinitely thick case (Figure 3.5), the concentration gradients for the 80 μm coatings are not as steep at a given illumination time. This arises from the fact that the diffusion of the active centers is confined to the finite thickness of the coating, therefore the concentration at the interface increases more rapidly with time. For example, the active center concentration at the bottom of the 80 μm thick 2 wt% pigmented coating after 5 minutes of illumination is 0.012 mol/L (from Figure 3.6a), whereas the value at the position of 80 μm depth in the infinitely thick system (Figure 3.5a) is 0.005 mol/L. The comparison between Figures 3.6a and 3.6b reveals the effect of the pigment loading on the active center concentration profiles. An increased pigment loading decreases the active center concentration at a given location and time, due to the effect of the pigment on the light intensity gradient.

The effect of the interface no-flux boundary condition on resulting active center profile becomes more pronounced as the coating thickness is reduced, as illustrated in Figure 3.7.

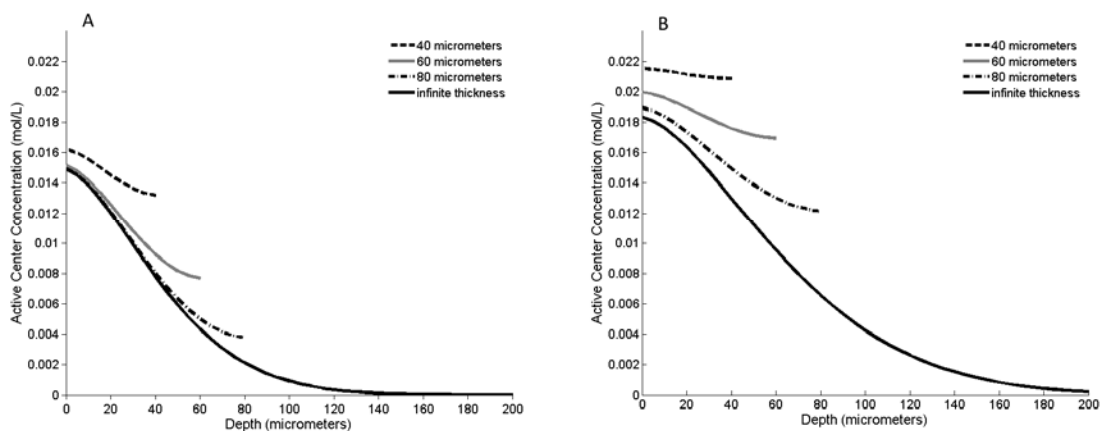


Figure 3.7. Profiles of active center concentration for 2 wt% CB-35 pigmented coatings ranging from 40 μm to infinitely thick. a) 2 minutes of illumination, b) 5 minutes of illumination. Monomer: EEC, Initiator: 1 wt% IPB.

Figures 3.7a and 3.7b show the active center concentration profiles at specific illumination times (2 minutes and 5 minutes respectively) for coating thicknesses ranging from 40 μm to infinitely thick, for 2 wt% CB-35. The figure illustrates that the active center concentration profiles for the infinitely thick systems decrease relatively sharply and reach a value of zero concentration at a depth of 140 μm for a two minute illumination time and 220 μm for an illumination time of five minutes. The profiles for the infinitely thick case provide an asymptotic limit for the coatings of finite thickness. Specifically, the active center concentration profiles approach those of the infinitely thick case as the coating thickness is increased or the illumination time is decreased. The figure also illustrates that the active center concentration profile becomes more uniform throughout the thickness

of the coating as the thickness is decreased or the illumination time is increased. These trends arise from the fact that the diffusion of the active centers is confined to the finite thickness of the coating. In the case of the 40 μm thick coating, the concentration is nearly uniform throughout the thickness of the coating after 5 minutes of illumination.

3.3.3 Post-illumination Diffusion of Active Centers in Pigmented Coatings

During the illumination period, active center profiles decrease sharply with depth, resulting in a concentration gradient and therefore a driving force for diffusion. Diffusion of the active centers during the post-illumination period is described by Fick's Second Law

$$\frac{\partial C_{AC}(z, t)}{\partial t} = D_{AC} \frac{\partial^2 C_{AC}(z, t)}{\partial z^2} \quad (3.9)$$

where C_{AC} corresponds to the concentration of active centers, and D_{AC} is the diffusion coefficient of active centers in cm^2/sec . The initial condition for the active center concentration as a function of depth is the profile obtained by applying Equation 8 at the end of the illumination period for each desired depth increment. In addition, the no-flux boundary condition indicates that there is no transport of initiator or photolysis product across the illuminated surface ($z = 0$) or the substrate boundary ($z = z_{max}$).

$$\frac{\partial C_{AC}(z, t)}{\partial z} = 0 \text{ at } z = 0 \text{ and } z = z_{max} \quad (3.10)$$

The diffusion coefficient for active centers generated by the IPB photoinitiator in the cycloaliphatic diepoxide containing 1 wt% carbon black was determined using the method described in the experimental section. Experimental cure depths for a 1 wt% carbon black pigmented system are shown in Figure 3.8. The effective diffusion coefficient was determined by

fitting this data to the diffusion equation, and was found to be 1×10^{-7} cm^2/sec , which is a reasonable value for reactive diffusion in which the active centers migrate by propagating with unreacted monomers [27]. Reactive diffusion has been identified as the primary mode for active center mobility in free-radical polymerizations of multifunctional acrylates [48] and cationic polymerizations of divinyl ethers [34,49].

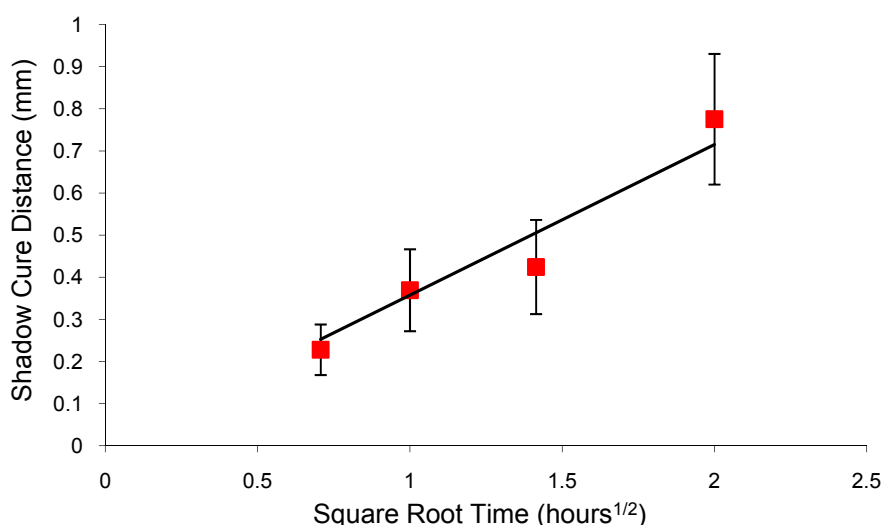


Figure 3.8. Shadow cure distance versus square root of time for a 1 wt% carbon black pigmented system. Monomer: EEC, Initiator: 1 wt% IPB, Exposure Time: 5 min.

Numerical solution of Equations 3.9 and 3.10 yields profiles of the active center concentration diffusing with increasing post-illumination time into a pigmented coating of finite thickness. Figures 3.9a and 3.9b show the active center profiles for an 80 μm coating pigmented with 2 wt% and 3 wt% CB-35, respectively, with increasing post-illumination time. As shown previously in Figure 3.6, the active center concentration profile exhibits a gradient at the end of the illumination period. Figure 3.9 indicates that the active center

concentration becomes uniform throughout the thickness of the 80 μm coating within 10 minutes post-illumination due to diffusion of the active centers. The comparison between Figures 3.9a and 3.9b illustrates that the final uniform active center concentration increases as the pigment loading is decreased since a higher fraction of the photoinitiator undergoes photolysis as the competitive absorption by the pigment is decreased.

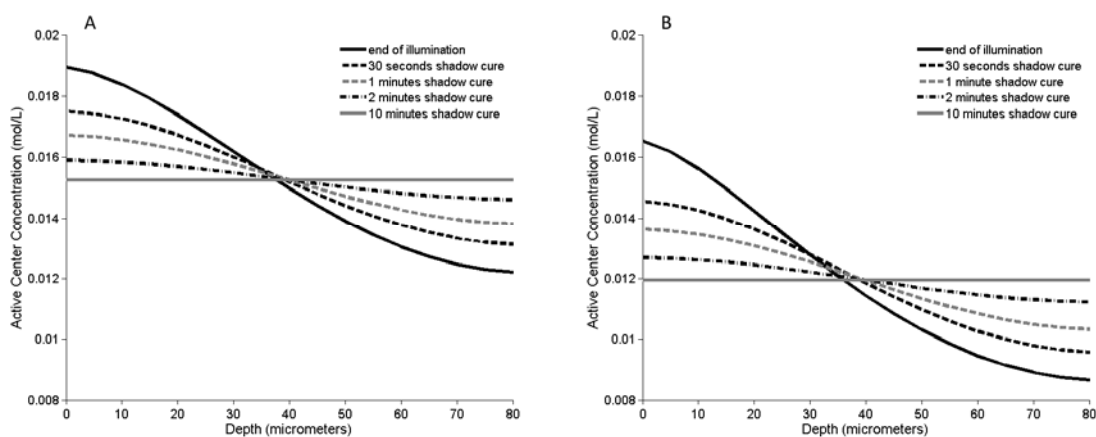


Figure 3.9. Active center concentration profiles diffusing post-illumination in an 80 μm thick pigmented coating. a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 min.

Shorter illumination times were used to demonstrate the post-illumination diffusion of active centers in a 40 μm pigmented coating. According to the results shown previously in Figure 3.7, the active center concentration profile was nearly uniform throughout the thickness of the coating after 5 minutes of illumination. But after only 3 minutes of illumination, the active centers produced near the surface of the coating were able to diffuse post-illumination, as shown in Figures 3.10a and 3.10b, for 2 wt% and 3 wt% pigment loadings, respectively.

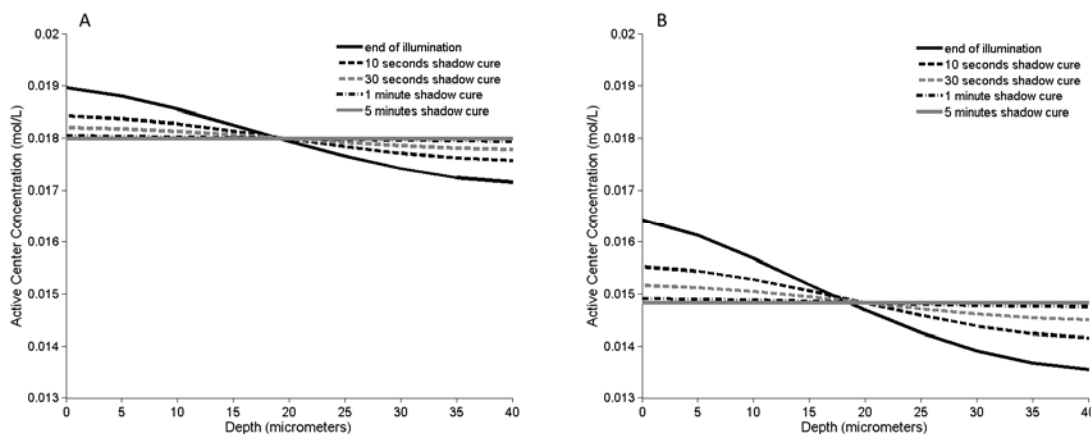


Figure 3.10. Active center concentration profiles diffusing post-illumination in pigmented coating (40 μm thick). a) 2 wt% CB-35, b) 3 wt% CB-35. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 3 min.

For the thinner 40 μm coatings, the active centers can only diffuse over half the distance. The concentration profile therefore becomes uniform within half the time (5 minutes post-illumination) compared with the 80 μm coatings in Figure 3.6 (10 minutes post-illumination).

3.3.4 Kinetic Analysis for Prediction of Cure Times

In pigmented coating applications, the cure time required to reach macroscopic property development (t_{mpd}) is especially important since it determines when a coated substrate may undergo additional process steps that involve contact with the surface. At this time, the exposed surface of the coating must be tack-free, and the cure on the bottom of the coating, where it interfaces with the substrate, must be sufficient to ensure effective adhesion. For pigmented systems polymerized cationically, the time required to achieve

cure at the bottom determines the t_{mpd} , since there is no oxygen inhibition at the exposed surface and the light intensity is the lowest at the bottom of the sample. For this reason, a conservative criterion of a 35% epoxide conversion at the bottom interface was established to predict the t_{mpd} . The t_{mpd} was estimated by obtaining the active center concentration profiles at the bottom interface using Equations 3.8 (for the illumination period) and 3.9 (for the shadow cure period) and entering these profiles into the integrated form of the polymerization rate equation for cationic polymerization, shown below.

$$\int_0^{t_{mpd}} C_{AC} dt = -k_p \ln\left(\frac{m_f}{m_i}\right) \quad (3.11)$$

where t_{mpd} is the cure time required to reach macroscopic property development, k_p is the propagation rate constant, and m_f/m_i is the ratio of final monomer concentration to initial monomer concentration.

Figure 3.11 contains plots of the active center concentration at the bottom interface as a function of time for 80 μm EEC coatings containing three different pigment loadings. In this figure, the system is illuminated for the first five minutes (active center concentration given by Equation 3.8), while the final ten minutes correspond to shadow cure (the active center concentration is determined by solving Equations 3.9 and 3.10). The figure illustrates that the active center concentration reaches a plateau when the active center concentration becomes uniform throughout the thickness of the coating. Increasing the pigment concentration leads to a reduced active center concentration at the bottom interface at any given time, including the plateau value when the active center concentration is uniform throughout the depth. As explained previously, this trend arises from the competitive absorption by the carbon black pigment which reduces the total number of active centers created during illumination.

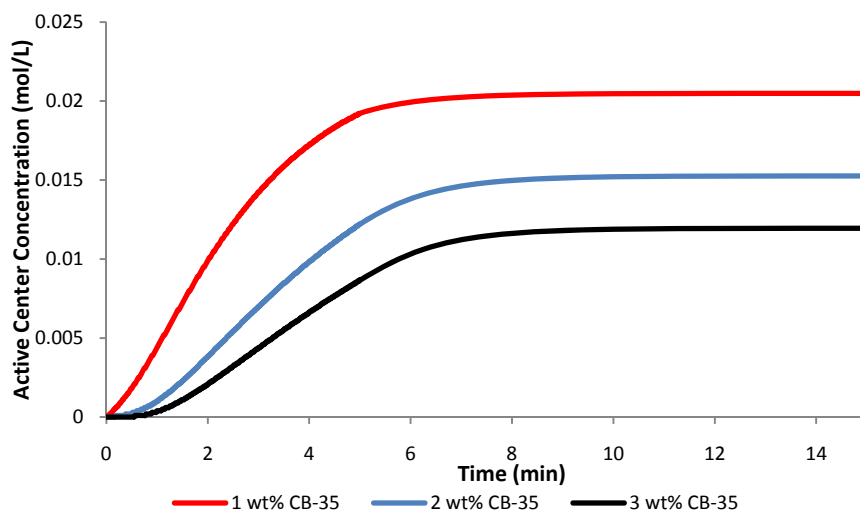


Figure 3.11. Active center profiles for various CB loadings at bottom interface of 80 μm thick coating. Monomer: EEC, Initiator: 1 wt% IPB, Pigment: 1-3 wt% CB-35, Exposure Time: 5 min.

The data shown in Figure 3.11 were integrated numerically to estimate the post-illumination cure time (t_{mpd}) by applying Equation 3.11, using the independently measured propagation rate constant described in the experimental section. However, Equation 3.11 is only valid if the carbon black neither catalyzes nor inhibits the reaction. Depending upon the method of preparation and surface treatment, the surface electronegativity of the pigment can vary from strongly acidic to strongly basic. Based upon a standard test for characterizing the acidity or basicity of pigments, the CB-35 pigment used in this study was reported to be basic (a pH of 9 from the ISO 787-9 test method was reported by the manufacturer [50]). For cationic photopolymerizations of pigmented coatings, a basic carbon black is preferable to ensure that the system has a desirable shelf-life. Formulations

used in these experiments were stable over several months. However, due to this basicity, the CB-35 carbon black acts as an inhibitor to the cationic photoinitiator in addition to absorbing the initiating light. Therefore, the active center concentration used in Equation 11 was taken to be the generated concentration shown in Figure 3.11 minus an inhibited active center concentration which is proportional to the carbon black loading. The value of the inhibited concentration was found to be 0.003 mol/L for each 1 wt% loading of carbon black.

Figure 3.12 shows a comparison between the experimentally determined t_{mpd} values and the theoretical values calculated using the procedure described above. In this figure, each data point corresponds to at least three independent experiments with the standard deviation indicated by the error bars. Recall that the t_{mpd} corresponds to the time after the illumination has ceased during which the long-lived active centers continue to react and diffuse into the thickness of the coating. Figure 3.12 indicates good agreement between the experimental and theoretical values of the post-illumination time required for macroscopic property development, indicating that the criterion of 35% cure at the bottom of the sample is reasonable. In addition, the data illustrate some interesting effects of illumination time and coating thickness. The illumination time is an important process variable since it determines the number of active centers produced, and therefore available for diffusion and cure. For this reason, the t_{mpd} decreases with increasing illumination time, especially at short illumination times. As the coating thickness is increased, the required cure time increases significantly at a given illumination time.

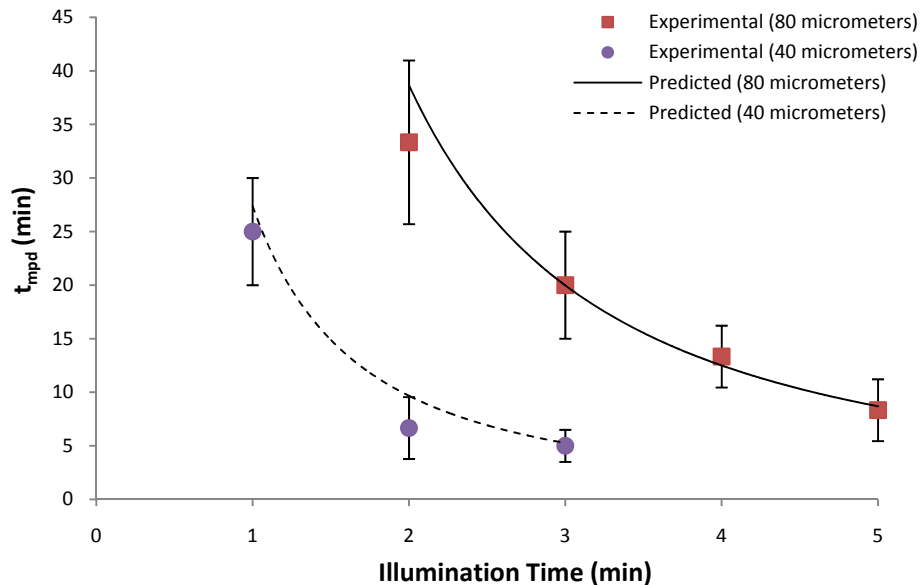


Figure 3.12. Comparison of predicted with experimental cure times (t_{mpd}) at bottom interface of 40 μm and 80 μm thick coatings for varying illumination times. Monomer: EEC, Initiator: 1 wt% IPB, Pigment: 2 wt% CB-35.

The time for macroscopic property development is of significant practical importance since it corresponds to the earliest time in which the polymerized ink or coating can be subjected to further processing steps without risk of damaging the surface or losing adhesion to the substrate. Using the analytical approach described above, the effects of the illumination time and the pigment loading on the post-illumination t_{mpd} were investigated more thoroughly. Figure 3.13 contains plots of the t_{mpd} as a function of the illumination time for three different pigment loadings (1, 2, and 3 wt.%) and two different thicknesses (40 and 80 μm in Figure 3.13a and b respectively). Recall that active centers are produced (and polymerization occurs) during the illumination time, and that the long-lived active centers continue to

propagate after the illumination has ceased. The post-illumination t_{mpd} will have a value of zero if the system cures during the illumination time (for example, the 1 wt% pigment, 40 μm , five minute illumination time case). For this reason, all of the plots will approach a value of zero as the illumination time is increased. If the illumination time is too short to produce enough active centers, the post-illumination t_{mpd} will go to infinity. The threshold illumination time required to cure the coating increases with increasing pigment loading due to the inhibitory effect of the basic carbon black pigment. Comparison between Fig. 13a and b shows that the thicker coatings require longer illumination times for a given post-illumination t_{mpd} .

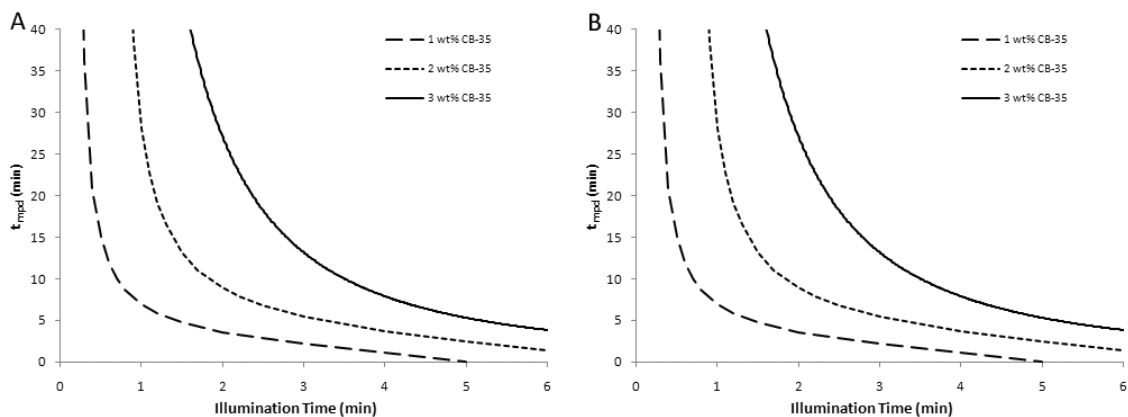


Figure 3.13. Predicted cure times (t_{mpd}) at bottom interface of coatings pigmented with 1-3 wt% CB-35. a) 40 μm thickness, b) 80 μm thickness. Monomer: EEC, Initiator: 1 wt% IPB.

3.4 Conclusions

In this contribution, the ability of long-lived cationic active centers to effectively cure coatings pigmented with carbon black has been investigated. The slightly basic, monodisperse carbon black pigment with a mean hydrodynamic radius of 29.2 nm used in these studies was found to act as a

mild inhibitor of the cationic photopolymerization. The light intensity gradient and photoinitiator concentration gradient for polychromatic illumination were determined for the pigmented system. The strong absorption by the carbon black resulted in sharp gradients in the pigmented systems. Consequently, the photoinitiator diffusion during the illumination period was found to have a marked effect on the resulting active center concentration profiles. Analysis of the active center reactive diffusion during the post-illumination period revealed that migration of the active centers leads to cure beyond the illuminated depth. The propagation rate equation coupled with the active center concentration profiles yielded theoretical cure times for the pigmented coatings. The coating thickness and pigment loading were found to be important variables in the time required for macroscopic property development. The long lifetimes and mobility of cationic active centers result in effective photopolymerization of carbon black pigmented coatings, and this comprehensive approach could be applied to other types of pigmented systems.

CHAPTER 4
CATIONIC ACTIVE CENTER MOBILITY IN
PHOTOPOLYMERIZATION SYSTEMS
WITH COMPLEX GEOMETRIES

4.1 Introduction

Photopolymerization is becoming a state-of-the-art technology for curing coatings on two-dimensional substrates. However, when the need arises for coating complex, three-dimensional objects, traditional photopolymerization suffers several disadvantages. Photopolymerizations are typically dependent upon the generation of extremely short-lived free-radicals that are sensitive to termination by oxygen. And since the free-radical photopolymerization reaction proceeds only in the presence of UV irradiation, the reaction terminates when the UV irradiation ceases. In order to successfully use free-radical photopolymerization for coating a three-dimensional substrate, such as an automobile body, first, the environment would need to be made inert to prevent oxygen inhibition. Secondly, complete irradiation of the entire substrate surface would be required, including recessed or shaded regions of the three-dimensional object. For these reasons, free-radical photopolymerizations fail to provide a practical and economical method for coating surfaces with irregular or intricate shapes.

Some of the methods being developed to overcome these problems for curing coatings on three-dimensional objects include plasma curing [18], which involves placing the coated substrate effectively within the light source; dual-cure systems [19-21], which require simultaneous thermal and photochemical curing mechanisms; and robotic UV curing [22], which uses

mechanical means of directing irradiation into recessed or shaded regions of the substrate. In this chapter, the use of cationic photopolymerization for multi-dimensional cure will be investigated. Cationic photopolymerization is an alternative to free-radical photopolymerization offering several unique advantages, especially for use in curing coatings on substrates with complex geometries. The cationic active centers generated in this type of photopolymerization exhibit extremely long lifetimes. They are not sensitive to free-radical scavengers, such as oxygen, and do not terminate by a radical-radical termination mechanisms. The long-lived cationic active center lifetimes can result in photopolymerization reactions that proceed long after irradiation has ceased, until the monomer is consumed or the active centers are entrapped in the polymer matrix.

Several investigators have shown cationic active centers to lead to dark cure, or post-polymerization, especially in the types of epoxide monomers used for cationic photopolymerization [24-26]. Due to their long lifetimes and tendency to diffuse, cationic active centers can also be responsible for a process known as shadow cure, or cure in regions that have never been illuminated [27]. The mobility of the cationic active centers facilitates cationic cure in recessed or shadow areas of a substrate that have not been directly exposed to irradiation.

This chapter provides a theoretical and experimental investigation of shadow cure in multiple spatial dimensions in cationically photopolymerized systems. The fundamental differential equations describing the polychromatic photoinitiation process are solved to obtain profiles of the active center concentration as a function of time and depth in the illuminated regions of the coating. During the illumination period, the active centers are preferentially generated at the surface to produce a sharp concentration

gradient that provides a driving force for diffusion into regions of unexposed monomer. Using the active center concentration profile at the end of the illumination period as the initial condition for the differential diffusion equation allows the spatial evolution of the active centers to be determined after the illumination has ceased. First, this analysis is used to describe the cationic active center concentration profiles in two spatial dimensions: the direction of the illuminating light, as well as a direction perpendicular to the light source. Second, photopolymerizations of cationically polymerizable monomer coatings are performed, with a portion of a two-dimensional substrate shaded from the initiating light. The coatings are shown to shadow cure in the shaded region, and experimental measurements are compared with theoretical predictions of cure distance by coupling the post-illumination diffusion analysis with the propagation rate equation. Finally, a novel method is presented in which cationic shadow cure is utilized to cure coatings with multiple layers using a single illumination step. The results of this research show that photopolymerization of systems with complex geometries can be made possible by the mobility of cationic active centers.

4.2 Shadow Cure in Two Spatial Dimensions

4.2.1 Governing Equations

For a photopolymerization system of rectangular cross-section subject to uniform polychromatic illumination normal to the top surface, the set of differential equations which govern the evolution of the light intensity gradient and initiator concentration gradient for polychromatic illumination [31,32] are shown below:

$$\frac{\partial C_i(z, t)}{\partial t} = -\frac{C_i(z, t)}{N_A h} \sum_j \left(\frac{\varepsilon_{ij} \varphi_j I_j(z, t)}{\nu_j} \right) + D_i \frac{\partial^2 C_i(z, t)}{\partial z^2} \quad (4.1)$$

$$\frac{\partial C_p(z, t)}{\partial t} = \frac{C_i(z, t)}{N_A h} \sum_j \left(\frac{\varepsilon_{ij} \varphi_j I_j(z, t)}{\nu_j} \right) + D_p \frac{\partial^2 C_p(z, t)}{\partial z^2} \quad (4.2)$$

$$\frac{\partial I_j(z, t)}{\partial z} = -[\varepsilon_{ij} C_i(z, t) + \varepsilon_{pj} C_p(z, t) + A_m] I_j \quad (4.3)$$

Here, the subscript j is an index with a different value for each wavelength of light under consideration; $C_i(z, t)$ is the initiator molar concentration at depth z and time t ; $C_p(z, t)$ is the photolysis product molar concentration at depth z and time t ; $I_j(z, t)$ is the incident light intensity of a specific wavelength at depth z and time t with units of energy/(area*time); ε_i is the initiator Napierian molar absorptivity of a specific wavelength with units of volume/(length*mole); ε_p is the photolysis product Napierian molar absorptivity of a specific wavelength with units of volume/(length*mole); φ_i is the quantum yield of the initiator at a specific wavelength, defined as the fraction of absorbed photons that lead to fragmentation of the initiator; N_A is Avogadro's number; h is Plank's constant; ν is the frequency of light in units of inverse seconds; D_i is the diffusion coefficient of the initiator in units of length²/time; D_p is the diffusion coefficient of the photolysis products; and A_m is the absorption coefficient of the monomer and the polymer repeat unit with units of inverse length. Note that this is the Napierian molar absorptivity because it is most natural for the differential version of the absorption equation (Equation 4.3).

For a photopolymerization system of thickness z_{max} which is illuminated at the planar surface where $z = 0$, the following initial and boundary conditions apply:

$$C_i(z,0) = C_o \quad (4.4)$$

$$C_p(z,0) = 0 \quad (4.5)$$

$$\frac{\partial C_{i,p}}{\partial z} = 0 \text{ at } z = 0 \text{ and } z = z_{\max} \quad (4.6)$$

$$I(0, t) = I_o \quad (4.7)$$

Equation 4.4 states that the initial initiator concentration, C_o , is uniform throughout the depth of the sample. Similarly, equation 4.5 indicates that the initial photolysis product concentration is zero. Equation 4.6, the no-flux boundary condition, indicates that there is no transport of initiator or photolysis product across the illuminated surface or the opposite boundary (typically an interface with a substrate). Finally, Equation 4.7 states that the light intensity on the illuminated surface is constant and equal to the initial intensity, I_o .

The solution to this set of equations provides detailed information regarding the time-evolution of the light intensity gradient and the initiator concentration gradient. For an accurate description of initiation with polychromatic illumination, the light intensity gradient at each incident wavelength must be individually described. As shown in equation 4.3, the intensity of an individual wavelength is attenuated by absorption of the initiator, monomer and polymer repeat units, and the photolysis product. Since the local initiator concentration depends upon all of the incident wavelengths, and the local light intensity of each wavelength depends upon the initiator concentration, the time-evolution of all of the light intensities are coupled to one another, and therefore the complete set of differential equations must be solved simultaneously. Unlike the previous chapter where the photoinitiator diffusion was found to be important for highly pigmented systems, in transparent systems the diffusion effects have been shown to be

negligible for relatively low initiator concentrations of photobleaching initiators. Therefore the diffusion terms in Equations 4.1 and 4.2 were neglected for the purposes of this chapter.

The local rate of active center generation is equal to the product of the local initiator concentration and the local light intensity summed over the initiating wavelengths. Since the cationic active centers are essentially non-terminating, and each photoinitiator molecule leads to the formation of a single active center molecule, the cationic active center concentration, C_{AC} , at a given depth, z , and time, t , can be determined from the integrated form of the rate equation:

$$C_{AC}(z,t) = \int_0^t C_i(z,t) \sum_j [I(z,t)]_j \phi_j \varepsilon_{ij} dt \quad (4.8)$$

In this manner, active center concentration profiles as a function of time and depth are determined during the illumination period. These resulting active center concentration profiles can then be used as an initial condition for determining the mobility of active centers during the post-illumination period. Post-illumination diffusion of the active centers in one spatial dimension (the direction of the illuminating light) during the post-illumination period is described by Fick's Second Law:

$$\frac{\partial C_{AC}(z,t)}{\partial t} = D_{AC} \frac{\partial^2 C_{AC}(z,t)}{\partial z^2} \quad (4.9)$$

where C_{AC} corresponds to the concentration of active centers, and D_{AC} is the diffusion coefficient of active centers in cm^2/sec . The initial condition for the active center concentration as a function of depth is the profile obtained by applying Equation 4.8 at the end of the illumination period for each desired depth increment. In addition, the no-flux boundary condition indicates that

there is no transport of initiator or photolysis product across the illuminated surface ($z = 0$) or the substrate boundary ($z = z_{max}$).

$$\frac{\partial C_{AC}(z,t)}{\partial z} = 0 \text{ at } z = 0 \text{ and } z = z_{max} \quad (4.10)$$

Post-illumination diffusion of cationic active centers into a second spatial dimension (both in the direction of the illuminating light and in a perpendicular direction) can be described using the following form of Fick's law in two spatial dimensions:

$$\frac{\partial C_{AC}(x,z,t)}{\partial t} = D_{AC} \left[\frac{\partial^2 C_{AC}(x,z,t)}{\partial x^2} + \frac{\partial^2 C_{AC}(x,z,t)}{\partial z^2} \right] \quad (4.11)$$

In this equation, x represents the direction perpendicular to the illuminating light, or the width of the sample. $C_{AC}(x,z,t)$ is the active center molar concentration at depth z , width x , and time t ; and D_{AC} is the diffusion coefficient of the active center in units of cm^2/sec . Again, the initial condition for this equation is the active center concentration profile obtained at the end of the illumination period. Again, no flux boundary conditions were used for the illuminated surface ($z = 0$) and the edge of the sample width ($x = 0$):

$$\frac{\partial C_{AC}(x,z = 0,t)}{\partial z} = 0 \quad \frac{\partial C_{AC}(x = 0,z,t)}{\partial x} = 0 \quad (4.12)$$

4.2.2 Results and Discussion

4.2.2.1 *Active Center Concentration Profile Produced During Illumination*

Equations 4.1 – 4.8 were solved numerically using the method of finite differences to obtain a profile of the active center concentration as a function of depth during a five minute illumination period for a system containing the monomer EEC and 1 wt% of the photoinitiator IPB. Molar absorptivity

values for the monomer, photoinitiator, and photolysis products were determined using methods described in Chapter 3. Detailed spectral information for the light source used in this simulation study (200 W Hg/Xe arc lamp) was also obtained using methods described Chapter 3. Figure 4.1 shows the active center concentration profile obtained for the EEC/IPB system as a function of sample depth after 5 minutes of illumination with the Hg/Xe lamp. This profile falls off rapidly with depth, and therefore exhibits a sharp gradient and a considerable driving force for diffusion. Due to light attenuation by the photoinitiator, the active centers are only produced up to a depth of approximately 0.3 cm.

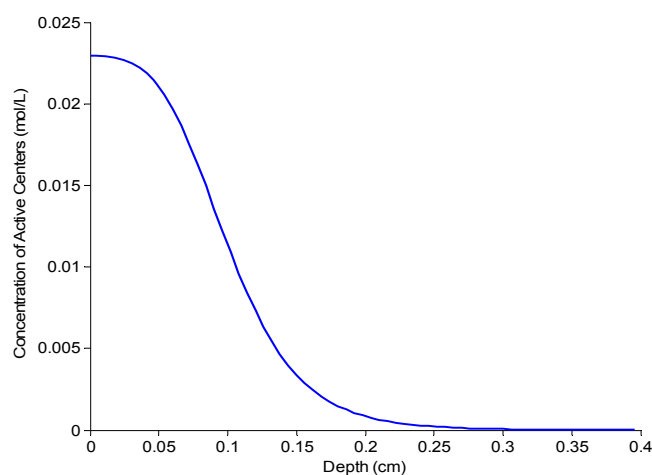


Figure 4.1. Active center concentration profile at the end of the illumination period. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 minutes.

4.2.2.2 *Post-illumination diffusion of cationic active centers*

It is convenient to consider the active center generation step and the active center diffusion step separately since each step is driven by different fundamental processes and occurs on different timescales. In the one-

dimensional case in which the concentration depends only upon depth, the diffusion and shadow cure occur in the same direction as the initial illumination, and the “shadow region” begins at the depth at which the light does not reach due to optical density of the system. Therefore, the one dimensional shadow cure case is relevant for situations in which the light does not penetrate through the entire depth of the sample, for example thick systems and pigmented systems. Note that for the EEC/IPB system described in Figure 4.1, the lower boundary condition ($z = z_{max}$) does not come into play for the sample thickness of 0.4 cm or greater, as shown in the figure.

The effective shadow cure diffusion coefficient for active centers generated by the IPB photoinitiator in the cycloaliphatic diepoxide EEC was measured for thick systems in one spatial dimension using a previously described experimental protocol [27], and was determined to be 3×10^{-6} cm²/sec, which is a reasonable value for reactive diffusion in which the active centers migrate by propagating with unreacted monomers. Reactive diffusion has been identified as the primary mode for active center mobility [34, 48, 49]. Numerical solution of the second-order partial differential diffusion equations (Equations 4.9 and 4.10) yields profiles of the active center concentration diffusing with increasing post-illumination time, or shadow cure time, in the z -direction. The resulting active center concentration profiles shown in Figure 4.2 demonstrate that there is a steep gradient at the end of the illumination period (the initial condition shown in Figure 4.1) which becomes more uniform throughout the thickness of the sample with increasing shadow cure time.

This theoretical description of shadow cure can be extended to the multidimensional case in which diffusion can occur in directions perpendicular to the initial illumination. Figure 4.3 shows a schematic of the

representative geometrical configuration that was chosen to investigate shadow cure in two spatial dimensions.

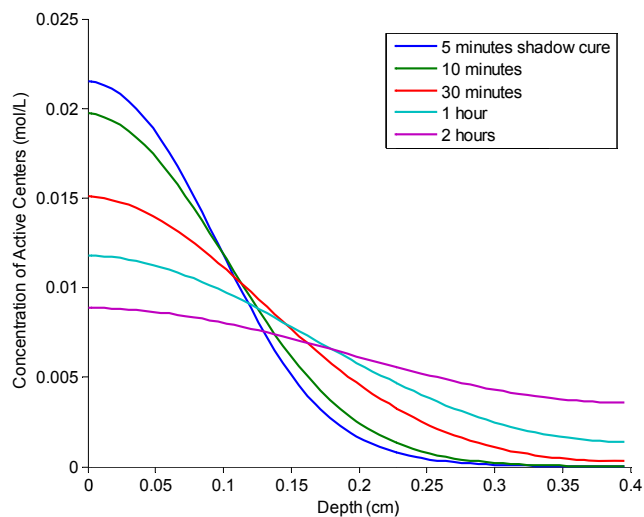


Figure 4.2. Active center concentration profiles diffusing with increasing shadow cure time. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 min.

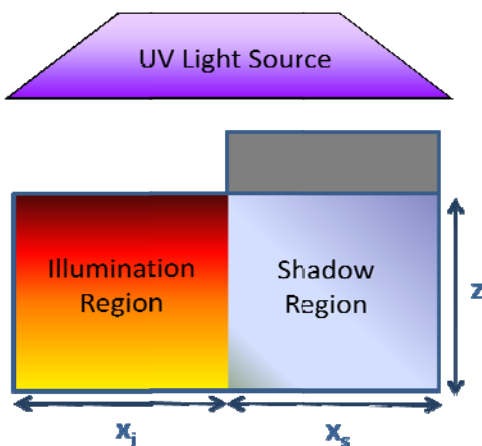


Figure 4.3. Geometrical configuration for shadow cure in two spatial dimensions. Rectangular cross-section of sample with depth z and width x .

The configuration shown in Figure 4.3 is a simple geometry for a system with depth z and width x . In this configuration, the illumination area of width x_i is exposed to the light source, and the shadow area of width x_s is masked.

Profiles of the cationic active centers diffusing into the shadow region were determined by applying Equations 4.11 and 4.12, again to the initial condition shown in Figure 4.1. Figures 4.4a through 4.4f illustrate the cationic active center concentration profiles obtained for the configuration illustrated in Figure 4.3. The simulation results for this representative case of two-dimensional shadow cure illustrate that as the shadow cure time increases from zero to 50 minutes, the active center profile broadens and extends deeper into the sample in the z -direction, as it did in the one-dimensional shadow cure case described in Figure 4.1. In addition, Figure 4.4 illustrates that active centers diffuse into the unilluminated region behind the mask in the x -direction with increasing post-illumination time. Figure 4.4a shows the vertical concentration gradient at the end of the illumination period between the illuminated region and the shaded region (where $x = 0.4$). Since this concentration gradient is steeper at the interface between the illuminated and masked regions, the driving force for diffusion is higher in the x -direction than it is in the z -direction. This method for describing shadow cure in multiple spatial dimensions can be applied to other configurations in which a region of the photopolymer is shaded from the illuminating light.

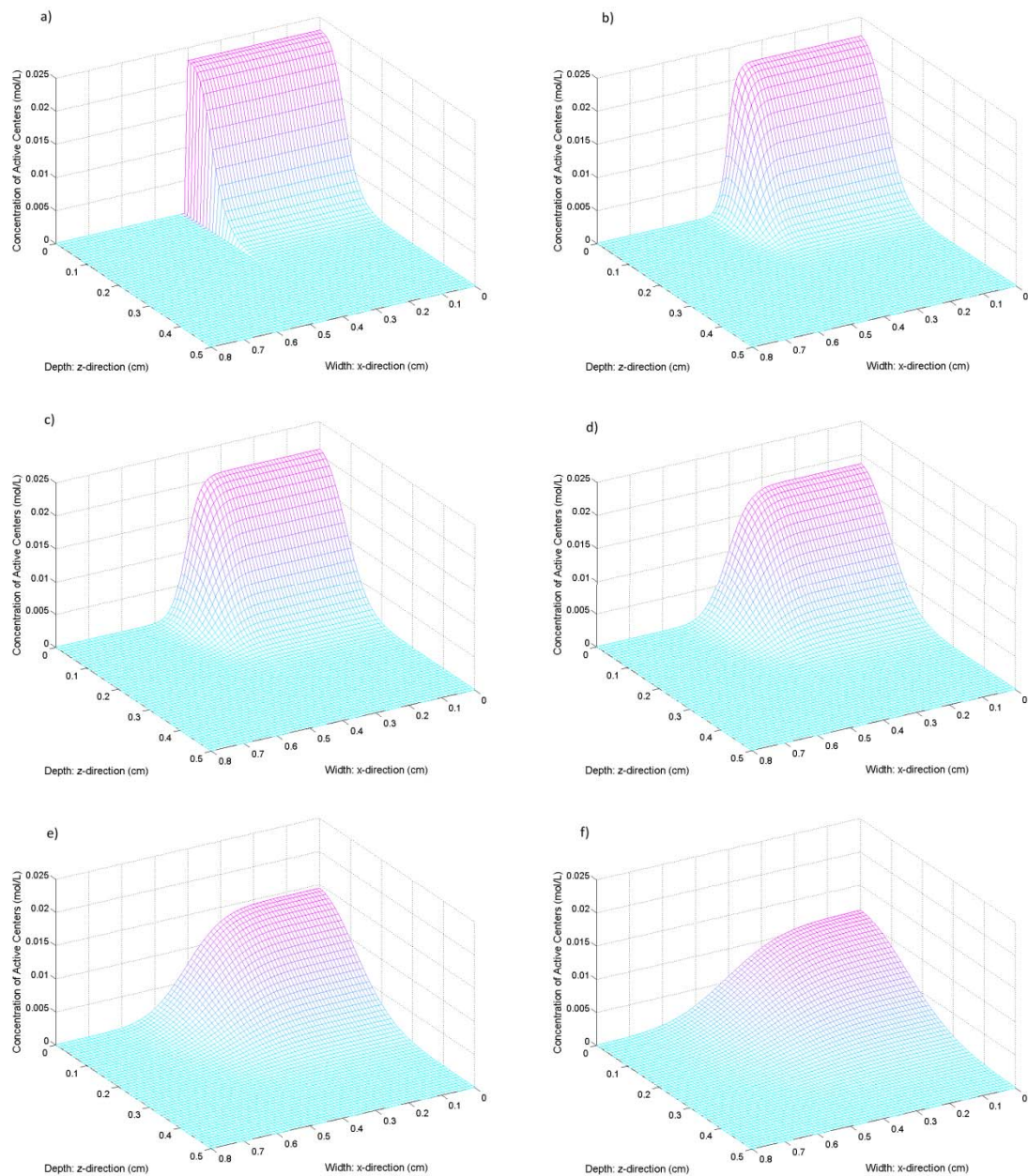


Figure 4.4. Active center concentration profiles in two spatial dimensions (x and z). a) end of illumination period, b) 2.5 minutes shadow cure, c) 5 minutes, d) 10 minutes, e) 25 minutes, and f) 50 minutes. Monomer: EEC, Initiator: 1 wt% IPB, Exposure time: 5 minutes.

4.3 Shadow Cure Perpendicular to the Direction of Illumination

Experiments were conducted to demonstrate the ability of cationic active centers to migrate into unilluminated regions, resulting in shadow cure. In order to measure the distance over which the cationic active centers migrate, or diffuse, post-illumination in a second spatial dimension (perpendicular to the direction of illumination), thin epoxide coatings were photopolymerized on aluminum panels with a portion of the coating shaded from the illuminating light source, as shown in the schematic in Figure 4.5. The governing equations for active center generation and diffusion into a second spatial dimension (Equations 4.11-4.12) were then solved to obtain the concentration profiles representative of this experimental setup, and theoretical results were compared with experimental findings.

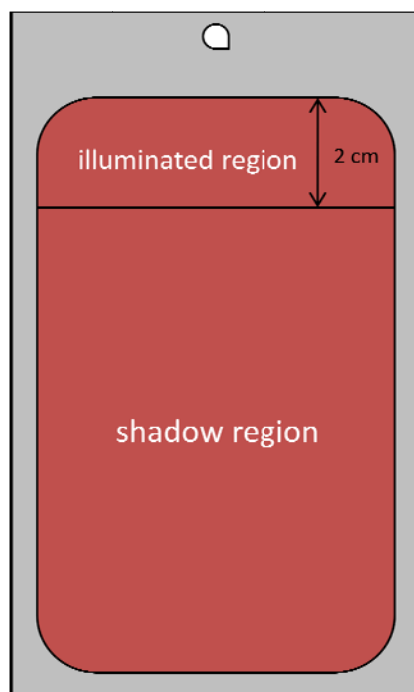


Figure 4.5. Schematic representation of shadow cure experiments used to measure cationic active center migration perpendicular to the direction of illumination.

4.3.1 Materials and Experimental Methods

The cationically polymerizable monomer 3,4-epoxy-cyclohexylmethanyl 3,4-epoxy-cyclohexanecarboxylate (EEC, Sigma Aldrich) was used in these experiments. The photoinitiator used was (tolycumyl) iodonium tetrakis (pentafluorophenyl) borate (IPB, Secant Chemicals). Methanol solvent was used for UV/Visible spectroscopy. The absorbance spectra for the monomer, photoinitiator, and photolysis products were determined in one nanometer increments using an 8453 UV-Visible spectrophotometer (Agilent Technologies). For the monomer and photoinitiator, the spectra were obtained for dilute solutions (10^{-2} M and 10^{-3} M respectively) in methanol placed in an air-tight, quartz cell to prevent any changes in concentration due to evaporation of the solvent. To obtain the absorbance spectra after photolysis, the photoinitiator samples were illuminated with a 200 Watt Hg-Xe arc lamp (Oriel Light Sources) until there was no change in the absorbance spectrum. Molar absorptivities before and after photolysis for IPB are reported in Chapter 3.

Solutions containing 97 wt% EEC and 3 wt% IPB were mixed together for 24 hours in dark conditions. The solutions were then spread evenly onto 3 in. by 6 in. aluminum substrates (Q-Lab) using a draw bar to achieve the desired uniform coating thickness. The coated panels were then covered with a coverplate to shade part of the panel, and irradiated using a LC6B bench top conveyer equipped with a F300S lamp system (Fusion UV Systems, Inc.). This lamp was equipped with a 13 mm "H" type bulb, and the irradiance was measured at the surface of the conveyer using UV integrating radiometer (EIT, Inc.) which gave an irradiance measurement in the UVB range of 1.9 W/cm². The irradiation time was controlled by the speed of the conveyer. The photopolymerization was carried out under atmospheric conditions and

at room temperature. The distance to which the coating cured within the shaded region was determined by characterizing the surface tack and the adhesion to the substrate at regular intervals using a dental probe. The cure distance was then measured with calipers. Once full property development was achieved, the thickness of the coating was obtained using a micrometer (micro-TRI-gloss μ , BYK Gardner).

4.3.2 Results and Discussion

4.3.2.1 *Experimental Results*

Aluminum panels coated with 80 μm of the EEC/IPB mixture were irradiated using the experimental configuration described above. The conveyor belt speed was set at 4 ft/min, giving an approximate illumination time of 3 sec. The photo in Figure 4.6 shows an example of the resulting shadow cure in the unilluminated region of a panel that progressively cured

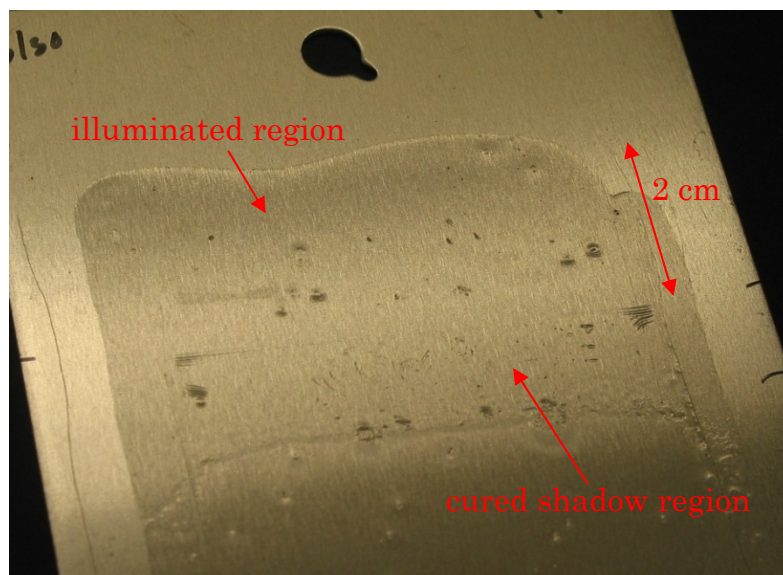


Figure 4.6. Photo showing illuminated region and shadow cure region of cured epoxide coating on aluminum panel.
Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.

over 30 minutes post-illumination. The edge of the illuminated region can be seen as a faint line corresponding to the black mark on the side of the panel which marked the position of the cover plate. As shown in the photo, the photopolymer cured almost 2 cm into the shaded region of the coating after 30 minutes. The shadow cure distance was measured as a function of post-illumination time using the method described. The numerical values recorded for shadow cure distance are shown in Figure 4.7. The shadow cure progressed in a linear fashion for approximately 30 minutes (18 mm) after which time the shadow cure front stopped. The unreacted monomer beyond the shadow cure front remained unpolymerized for days/weeks post-illumination.

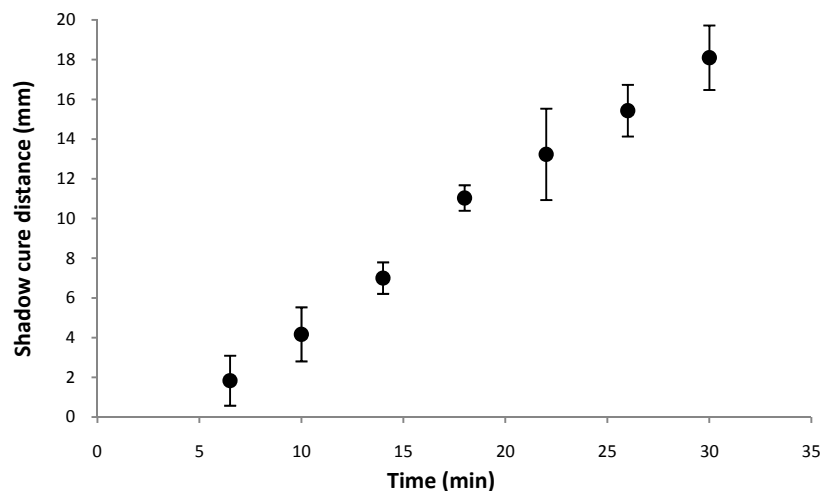


Figure 4.7. Experimental measurements of shadow cure distance into shaded region over 30 minutes post-illumination time.
Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.

Control experiments were performed to verify that no polymerization took place on a panel coated with the EEC/IPB mixture which was completely shaded from the light source, yet processed through the fusion lamp

conveyor. No noticeable heat was generated in the panels, and a temperature indicator on the UV radiometer used to measure the light source intensity showed an increase in temperature through the conveyor/lamp system of only 3-5 °C, indicating that very little thermal polymerization, if any, could have taken place.

4.3.2.2 Post-illumination Diffusion and Kinetics

Equations 4.1-4.8 were solved numerically to determine the active center concentrations produced in the illuminated region of the coating. Figure 4.8 shows the active center concentration as a function of the depth during the 3 second illumination period. Due to the high intensity of the light source, the photoinitiator is completely consumed, resulting in the active center concentration reaching a maximum value (equal to the initial photoinitiator concentration) within the 3 seconds of illumination. Therefore the active center concentration is considered to be uniform throughout the 80 μm thick coating after the 3 seconds of illumination with the Fusion lamp.

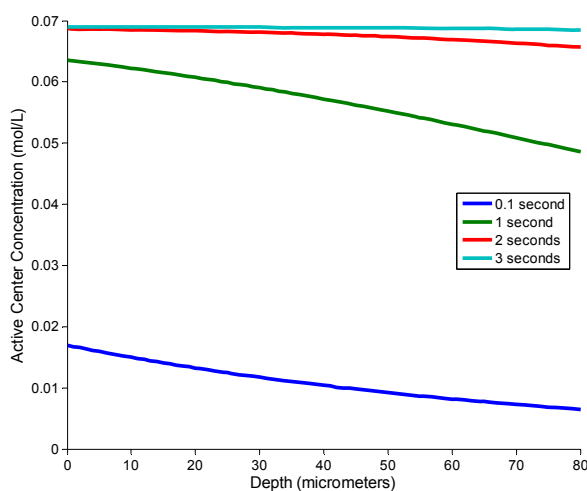


Figure 4.8. Active center concentration profiles throughout the 80 μm coating depth throughout the 3 second illumination time.

Since the active center concentration profile in the illuminated region is uniform throughout the thickness of the coating, diffusion of the active centers in the direction of the illuminating light (z -direction) does not occur. Therefore, this coating configuration can now be modeled as a one-spatial dimension problem. Using Equations 4.11 and 4.12 (with the z -terms in the equations removed), the active centers diffusing into the shaded regions in the direction perpendicular to the illuminating light (the x -direction) were described. The effective shadow cure diffusion coefficient used for this analysis was $1 \times 10^{-4} \text{ cm}^2/\text{sec}$, which was determined by fitting the experimental data shown in Figure 4.7 to the diffusion equation. Figure 4.9 shows the resulting active center concentration profiles diffusing during 30 minutes post-illumination for a 2 cm illuminated region (left side of Figure 4.9) and the first 2 cm of the shaded region (right side of Figure 4.9).

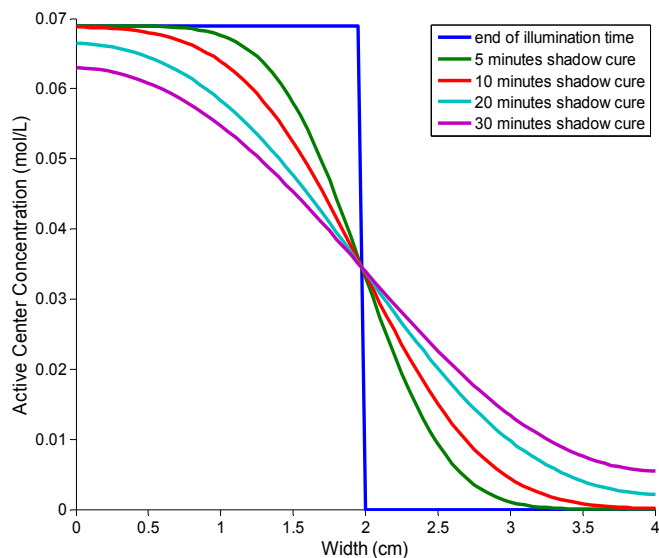


Figure 4.9. Active center concentration profiles diffusing into the shadow region (x -direction) with increasing shadow cure time. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.

The active center concentration profile begins as a vertical gradient at the end of the illumination time, and progressively becomes more uniform as the active centers from the illuminated region diffuse into the shaded region with increasing post-illumination time.

In coating applications, the cure time required to reach macroscopic property development (t_{mpd}) is especially important since it determines when a coated substrate may undergo additional process steps that involve contact with the surface. At this time, the exposed surface of the coating must be tack-free, and the cure on the bottom of the coating, where it interfaces with the substrate, must be sufficient to ensure effective adhesion. The t_{mpd} was estimated by obtaining the active center concentration profiles at given positions within the shadow cure region, and entering these profiles into the integrated form of the polymerization rate equation for cationic polymerization, shown below.

$$\int_0^{t_{mpd}} C_{AC} dt = -k_p \ln\left(\frac{m_f}{m_i}\right) \quad (4.13)$$

The t_{mpd} is the cure time required to reach macroscopic property development, k_p is the propagation rate constant, and m_f/m_i is the ratio of final monomer concentration to initial monomer concentration. A conservative criterion of a 35% epoxide conversion was established to predict the t_{mpd} , and the k_p for EEC was determined previously to be 0.1 L/mol-sec (see Chapter 3 for details).

Figure 4.10 contains plots of the active center concentration as a function of post-illumination time at various positions in the shadow cure region (x -direction). The figure illustrates that the active center concentrations increase rapidly at positions close to the illuminated edge (2-4

mm into the shadow cure region) and increase very slowly at positions much farther from the illuminated region (16-20 mm into the shadow cure region).

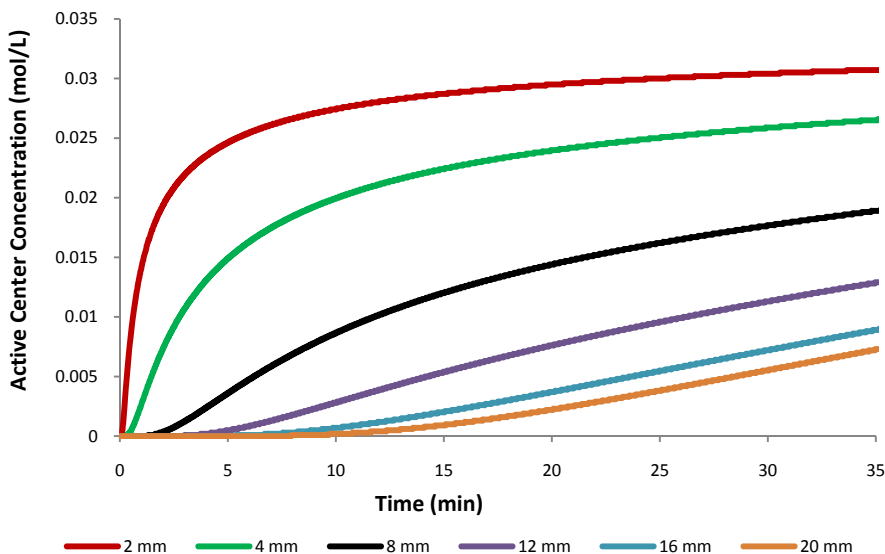


Figure 4.10. Cationic active center concentration as a function of time at various positions in the shadow cure region (x-direction). Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.

The data shown in Figure 4.10 were integrated numerically to estimate the post-illumination cure time by applying Equation 4.13. Figure 4.11 shows a comparison between the experimentally determined cure times (from Figure 4.7) and the theoretical values predicted using this method. This figure indicates very good agreement between the experimental findings and theoretical predictions. The shape of the plot shown in Figure 4.11 is very interesting because both the experimental data and model predictions demonstrate a linear dependence of shadow cure distance on time. The shadow cure front progresses linearly with time until diffusing active centers are depleted or until they become trapped in the polymer matrix.

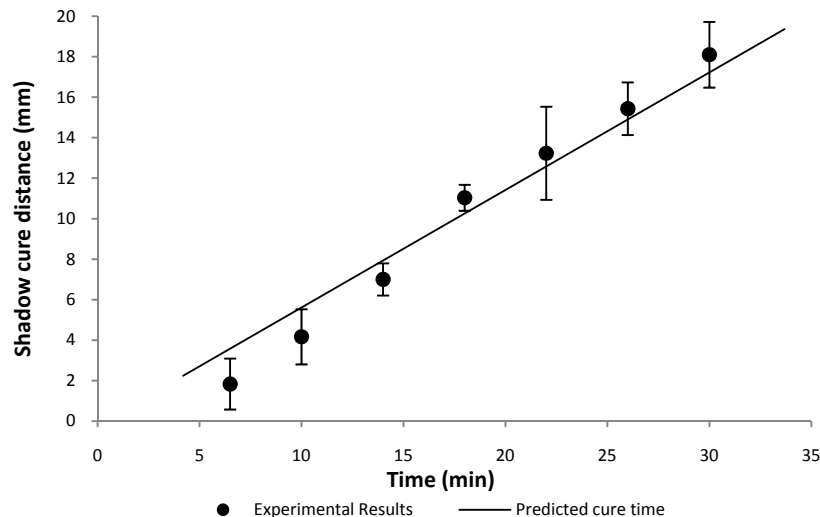


Figure 4.11. Experimental measurements of shadow cure distance into shaded region over 30 minutes post-illumination time. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 seconds.

The data in Figure 4.9 indicate that the active centers are diffusing from the entire 2 cm width of the illuminated region of the 80 μm thick coatings. The diffusion analysis was again employed to determine whether the shadow cure distance is dependent upon the width of the illuminated region, which can be considered a “reservoir” of active centers. Figures 4.12a and b illustrate the effect of the illuminated width. Figure 4.12a shows active center concentration profiles as a function of the x -direction for a 1 cm wide illumination width. The active centers diffuse approximately 3 cm into the shaded region (4 cm on the x -axis), dropping off to a value of zero at this position after approximately 30 minutes post-illumination. Figure 4.12b shows the active center concentrations migrating over the same shadow cure distance, 3 cm into the shaded region after 30 minutes post-illumination, for a much wider illumination width of 4 cm. These results suggest that unless

the width of the active center reservoir is so narrow that the active centers are completely depleted (less than a few millimeters) the shadow cure distance remains independent of the illuminated width.

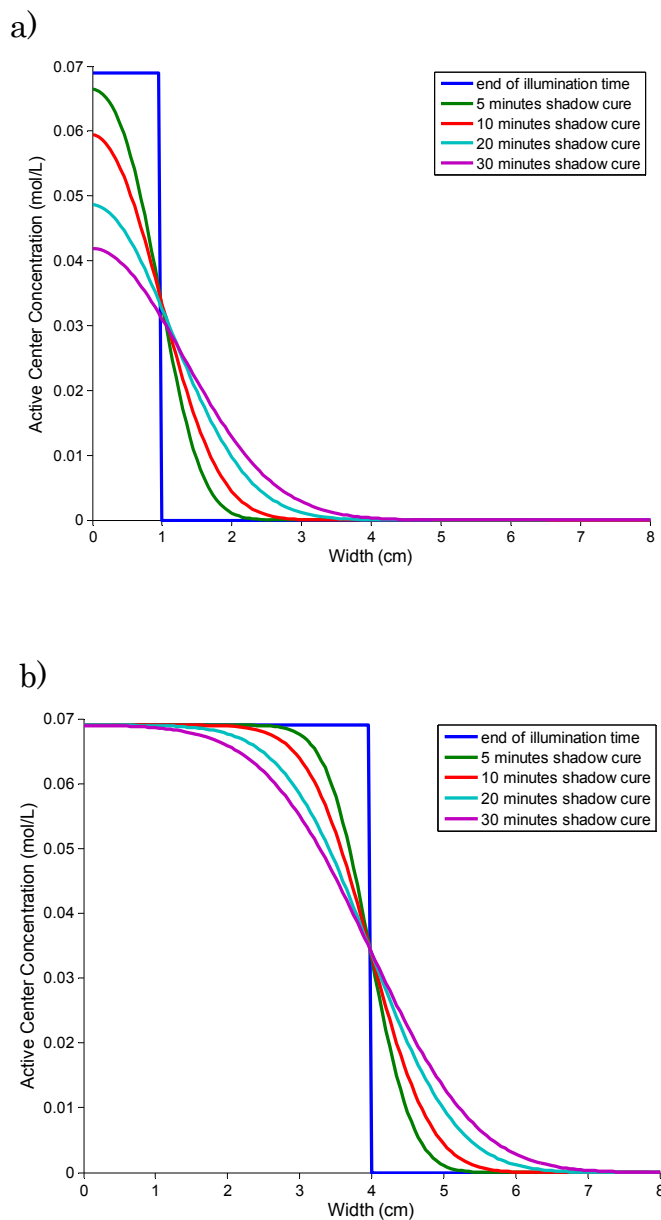


Figure 4.12. Active center concentration profiles diffusing into the shadow region (x -direction) with increasing shadow cure time for a) 1 cm illuminated width, and b) 4 cm illuminated width. Monomer: EEC, Initiator: 3 wt% IPB, Exposure time: 3 sec.

4.4 Shadow Cure Through Multiple Coating Layers

Many commercial coatings contain multiple layers with the composition of each layer selected to promote the properties required for that layer. For example, some layers may primarily promote adhesion, impart color, or change optical properties, while other layers may primarily impart scratch resistance or other desirable mechanical properties. Curing multiple layers using thermal polymerization generally requires a complete cure cycle, which can include application of a monomer followed by heating for an extended period of time for each individual layer. This process is cumbersome, time intensive, and increases the cost of the coating. Examples of industries which use multi-layered coatings include automotive bodies, furniture and cabinetry, and resilient flooring.

The remainder of this chapter presents a method for photopolymerizing multi-layered coatings. In this method, long-lived cationic centers that are photo-generated in only one layer of a multi-layered coating lead to cure in multiple layers, even if the layers are added after illumination. Rather than illuminate each layer individually after application of the monomer, this method requires only one illumination step to cure a coating containing multiple layers. For example, when a second layer of monomer is applied to a previously photopolymerized sub-layer, cationic active centers created in the sub-layer can migrate into the monomer coating applied on top. Therefore a fully cured coating comprised of two or more layers can be completely cured using only one illumination step.

An additional advantage of this method is that cationic active centers can be used to cure pigmented coatings. In the method described, one or more of the layers in a multi-layered coating may contain pigments. Long-lived cationic active centers have been shown to fully polymerize single-

layered pigmented coatings (see Chapter 3). However, pigmented coatings often require multiple layers to be polymerized. For example, automotive body coatings can require application of a pigmented layer followed by a protective clear coat. This method could be used to photo-cure a pigmented layer containing cationic photoinitiator, followed by a clear coat application that can be fully cured without further illumination. Because this method reduces the number of illumination steps required and has the potential for use in pigmented coatings, it will be of potential commercial value in many industries where multiple layer protective coatings are utilized.

4.4.1 Experimental Methods

The experimental setups for two examples of the method introduced above are illustrated in Figures 4.13 and 4.14. In the first set of experiments, Figure 4.13, the bottom layer only of a two-layer coating was illuminated. A mixture of typical epoxide monomers commonly used for cationic photopolymerization was mixed with 1 wt% of either one of two cationic photoinitiators: (tolycumyl) iodonium tetrakis (pentafluorophenyl) borate (IPB, Secant Chemicals) or diaryliodonium hexafluoroantimonate (IHA, Sartomer). The monomer mixture contained 70 wt% 3,4-epoxycyclohexyl-methanyl 3,4-epoxycyclohexane-carboxylate (EEC, Sigma Aldrich) and 29 wt% 2-butoxymethyl-oxirane (BMO, Hexion Specialty Co.). BMO was added to reduce viscosity so that monomer solutions could be sprayed onto the aluminum substrate using an airbrush. These experiments were conducted both with and without an additional 1 wt% of the pigment Titanium Dioxide (TiO₂, Dupont) added to the monomer/ photoinitiator mixture in the bottom layer. This layer was then irradiated for 10 minutes using a 200 Watt Hg-Xe arc lamp (Oriol Light Sources) with a measured irradiance of 50 mW/cm². A

layer of the same monomer mixture containing no photoinitiator was then sprayed on top of the cured layer. The coated panels were stored at atmospheric conditions and room temperature, and monitored to determine the time required to obtain a tack-free surface. Once polymerized, the thickness of the coating was measured using a micrometer (micro-TRI-gloss μ , BYK Gardner).

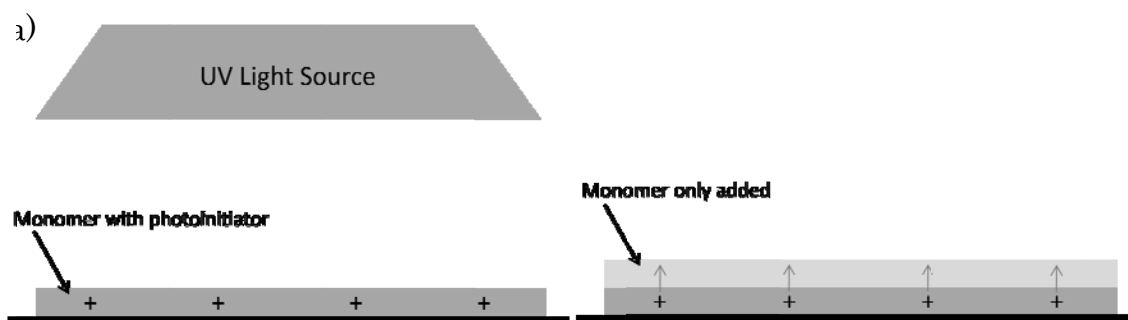


Figure 4.13. Pictorial representation of the experimental setup for bottom layer only illumination. a) the bottom layer, containing monomer and photoinitiator, is illuminated. b) the second layer of monomer is added, which cures with no further illumination. Arrows indicate diffusion of active centers into unilluminated layer.

In a second set of experiments, Figure 4.14, the top layer only of a two-layer coating was illuminated. A mixture containing 70% by weight CDE and 30% by weight BMO was sprayed onto an aluminum substrate using an airbrush. A second layer containing the same monomer mixture with 1% by weight IPB photoinitiator was sprayed over the top of the unpolymerized bottom layer. This experiment was conducted both with and without an additional 1 wt% of the pigment TiO_2 added to the monomer/photoinitiator mixture in the top layer. The two-layer coating was then irradiated for 10 minutes using the Hg-Xe arc lamp. The photopolymerization was carried out

under atmospheric conditions and at room temperature. The coated panels were monitored to determine the time required to obtain a tack-free surface. Once polymerized, the thickness of the coating was measured using a micrometer.

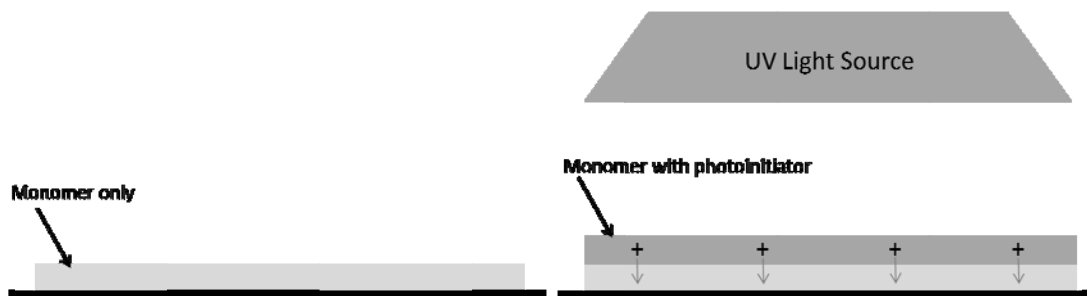


Figure 4.14. Pictorial representation of the experimental setup for top layer only illumination. a) a layer of monomer only is applied to a substrate. b) a second layer of monomer containing photoinitiator is added and illuminated, resulting in cure of the entire film. Arrows indicate diffusion of active centers into unilluminated layer.

4.4.2 Results and Discussion

In the first set of experiments illustrated in Figure 4.13, the bottom layer was illuminated for 10 minutes with the Hg-Xe lamp. The active centers then diffused into the top layer, which was applied post-illumination. This resulted in overall cure and full property development for both layers. Table 4.1 summarize the experimental results for various combinations of monomer, photoinitiator, and pigment, and gives the overall thickness of the final cured coatings. These results illustrates that the overall shadow cure of the unilluminated layer is complete within 1-2 hours regardless of the constituents of the layers.

Table 4-1. Results of shadow cure through multiple layers:
bottom layer illuminated only

Bottom Layer	Top layer	Exposure time (min)	Time to tack-free (hrs)	Average thickness (μm)
EEC/BMO/1 wt% IHA	EEC/BMO only	10	1.3 \pm 0.2	51.4
EEC/BMO/1 wt% IPB	EEC/BMO only	10	1-2	51.8
EEC/BMO/1wt% IPB	EEC/BMO/ 1wt% IPB	10	1-2	59.0
EEC/BMO/1wt% IPB/ 1wt% TiO ₂	EEC/BMO only	10	1-2	61.1
EEC/BMO/1wt% IPB	EEC/BMO/ 1wt% TiO ₂	10	1-2	49.0

In the second set of experiments, illustrated in Figure 4.14, two successive layers were sprayed onto the panels, one on top of the other using an airbrush. The two layered coating was then illuminated for 10 minutes with the Hg-Xe lamp. In these experiments, the bottom layer contained only monomer, with no photoinitiator. Both layers cured within the illumination time regardless of the constituents of the layers. Table 4.2 lists the combinations of monomer, photoinitiator, and pigment that were used in the two layers, as well as the overall thickness of the final coatings.

Table 4-2. Results of shadow cure through multiple layers:
top layer illuminated only

Bottom Layer	Top layer	Exposure time (min)	Time to tack-free (hrs)	Average thickness (μm)
EEC/BMO	EEC/BMO/ 1wt% IPB	10	0	60.4
EEC/BMO	EEC/BMO/ 1wt% IPB/ 1 wt% TiO ₂	10	0	82.9

In both sets of experiments, the cationic active centers were able to diffuse into an unilluminated layer containing no photoinitiator. The driving force for this phenomenon is considered to be the concentration gradient between the two layers. The films cured completely within 1-2 hours regardless of the constituents of the layers, and the cure time was not affected by the presence of pigment in the illuminated layer. The results of these experiments have shown this method to be a novel use of cationic photopolymerization for curing multi-layer coatings.

4.5 Conclusions

The long lifetimes and mobility of cationic active centers have been shown in this chapter to provide a solution to some of the limitations to photopolymerization of systems with complex shapes. First, the theoretical description for post-illumination diffusion of cationic active centers was applied in two spatial dimensions. This information provides a fundamental understanding of shadow cure in multiple dimensions. Secondly, this approach was used to predict shadow cure distance in a direction perpendicular to the illumination for cationically polymerized epoxide coatings by coupling the diffusion analysis with the kinetic rate equation. These results showed very good agreement between theoretical predictions and experimental findings which showed that the coatings were able to shadow cure over substantial distances. Finally, a novel method was introduced in which multiple layers of epoxide coatings can be cured using only a single illumination step. This method is made possible by the mobility of the cationic active centers, which diffuse into the unilluminated layers. All of these aspects of shadow cure show that cationic photopolymerization may

have tremendous potential in many applications where curing systems with complex geometries are required.

CHAPTER 5

THE POTENTIAL OF FREE-RADICAL SHADOW CURE

5.1 Introduction

The long-lived cationic active centers discussed previously in this contribution have shown great potential for mobility, resulting in cure of unilluminated regions of a photopolymer. Free-radical active centers have significantly shorter lifetimes, and tend to terminate immediately upon cessation of the illuminating light source. However, some cutting edge applications for photopolymers may necessitate the use of shadow cure in free-radical systems, specifically in some electronics applications. Advances in microelectronics have led to an increase in demand for fine pitch technology, which is defined as electronic surface mount components that contain interconnections whose pitch, or spacing, is on the order of 100 to 600 μm . Despite the extremely short lifetimes of the free-radical active centers, mobility of free-radicals within these increasingly small dimensional scales is becoming a possibility.

An example of an application in which free-radical shadow cure in photopolymerizations may become feasible is polymer-based conductive adhesives. These types of adhesive materials are becoming more widely used in the electronics industry as a substitute for lead-based solders [9]. Electrically conductive adhesives provide an environmentally friendly solution for interconnections in many electronics applications requiring fine pitch. Isotropic conductive adhesives (ICAs) are heat-curable materials containing an isotropic concentration of conductive particles, typically silver-filled epoxides, that allow current to flow in all directions through the cured polymer. These adhesives can be used to electrically interconnect non-

solderable substrates such as ceramics or plastics, or to replace solder for thermally-sensitive components. ICAs can be used to replace tin/lead solder for surface mounting on printed circuit boards, where the increasing density of discrete components soldered onto the surface is reaching a limit.

A second type of electrically conductive adhesive is anisotropic conductive adhesive, which allow current to flow along a unidirectional axis [8]. This type of adhesive is a composite containing fine electrically conductive particles uniformly dispersed in a polymer matrix. The design of these anisotropic conductive adhesive materials results in no direct contact between the conductive particles, but allows for electrical interconnection only at planned sites where the particles come into direct contact with a conductive substrate. The most common form these adhesives is a heat-curable, thermoplastic, anisotropic conductive film (ACF) used for flexible printed circuits. Ninety percent of all ACFs are currently sold for use in flat panel displays such as LCDs [10]. They are also used in flexible printed circuits, in which electronic circuits are mounted on flexible plastic substrates. Flexible printed circuits are being used in many applications, such as electronic books and other forms of electronic paper, displays for cameras and cell phones, and computer keyboards.

Figure 5.1 shows a schematic of the interconnection process for a typical flexible printed circuit assembly containing an ACF. A polymer resin containing conductive particles is laminated between two printed circuits. A specified temperature and pressure are applied until contact is made between the electrodes on the printed circuits and the conductive particles between them. As a result, current is allowed to travel in the z -direction only, and not in the x/y plane. Mechanical integrity of the assembly is maintained by the cured adhesive film.

Figure 5.2 shows a schematic for a proposed photopolymerization process that could be used to cure an ACF. In this case, the heating step can be eliminated by photocuring through a transparent substrate using an appropriate light source. Advantages of replacing the heat-curing process with photopolymerization include: protecting the heat-sensitive circuits from elevated temperatures; minimizing cure times; eliminating the heat source, resulting in energy savings; and eliminating the use of solvents, thereby minimizing emissions of volatile organic compounds and reducing the need for solvent vapor handling systems.

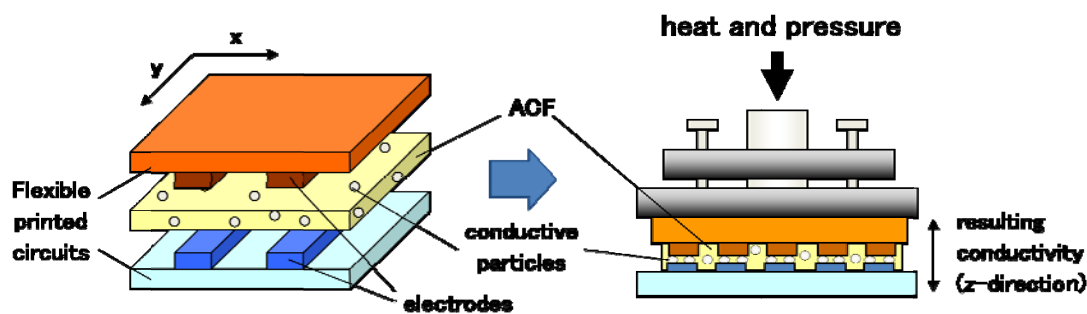


Figure 5.1. Flexible printed circuit assembly using heat-curable ACF.

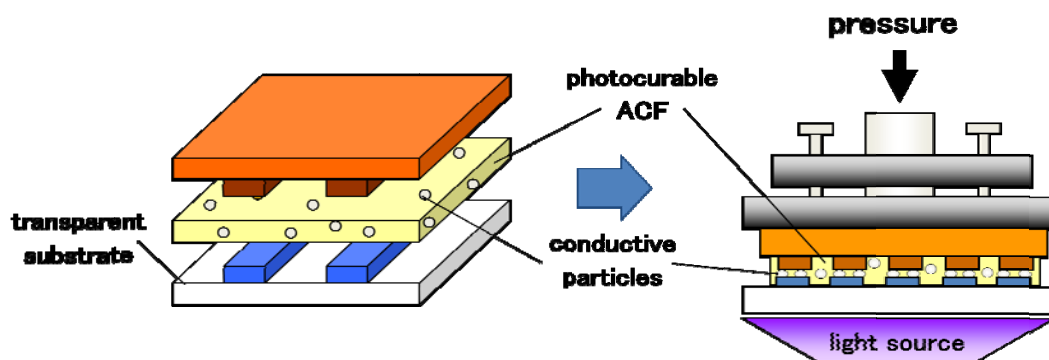


Figure 5.2. Flexible printed circuit assembly using proposed photopolymerizable ACF.

The remainder of this chapter discusses an investigation of one possible formulation that could be used in a photopolymerizable ACF assembly, such as the one proposed in Figure 5.2. One primary consideration in selecting a photopolymer system for this application is to attain full polymerization in unilluminated regions, behind electrodes and around conductive particles. These regions are shaded from the initiating light source, and it is important that these shadow regions cure to an acceptable conversion. A second consideration is that the substrate through which the ACF is illuminated may be only partially transparent, and may limit the initiating wavelengths. This chapter first discusses the selection process for both the photoinitiator and illuminating light source chosen for this application, and then characterizes shadow cure behind the electrodes for the chosen system.

5.2 Selection of Photoinitiator and Light Source for a Photopolymerizable ACF

The majority of flexible circuits are printed on polyimide films, which are chosen for their superior thermal stability [51,52]. Using polyimide film as a substrate through which an ACF can be cured by photopolymerization presents a particular challenge since this material absorbs wavelengths below 500 nm. This problem prohibits the use of UV light for this application, and requires selection of a visible light photoinitiator system for this application. Appropriate light sources and photoinitiation systems were selected for effective photopolymerization through polyimide film.

5.2.1 Materials and Experimental Methods

The monomer used in these experiments was 2-hydroxyethyl acrylate (HEA, Sigma Aldrich). The photosensitizers evaluated in this study included

Eosin Y (spirit-soluble), 3-hydroxy-2,4,5,7-tetraiodo-6-fluorone (H-Nu 535, Spectra Group, Inc.), Erythrocin B, Rose Bengal, and Camphorquinone. Except for the H-Nu 535, all photoinitiators were acquired from Sigma Aldrich. These photosensitizers were used in combination with the electron donor N-methyl-diethanolamine (MDEA, Sigma Aldrich). The absorption spectra for the monomer and photoinitiators were determined in one nanometer increments using an 8453 UV-Visible spectrophotometer (Agilent Technologies). The emission spectra of the illuminating light sources were collected using measured using a calibrated miniature fiber optic spectrometer (USB4000, Ocean Optics, Inc.)

Monomer conversion was measured using Fourier transform infrared spectroscopy (FTIR), which has become a standard method for analyzing photopolymerization systems [53,54]. For each of these studies, a droplet of monomer was placed between two rectangular IR grade sodium chloride salt crystal slides, with 15 μm Teflon beads placed between both ends of the slides to serve as spacers. Infrared spectra were collected using a modified Bruker 88 FTIR spectrometer designed to accommodate a horizontal sample. The infrared absorption spectra were obtained before, during, and after illumination. The =C–H infrared absorbance peak at 812 cm^{-1} , which is associated with an out-of-plane vibration, has traditionally been used to monitor acrylate conversion [53]. Conversion was calculated using Equation 5.1:

$$\% \text{ Conversion} = \left[\frac{A_0 - A_t}{A_0} \right] \quad (5.1)$$

where A_0 is the absorbance at 812 cm^{-1} before irradiation, and A_t is the absorbance at time t . The absorbance is found by measuring the baseline corrected peak height for the =C–H out-of-plane band at 812 cm^{-1} .

5.2.2 Results and Discussion

5.2.2.1 *Selection of Initiating Light Source for Photo-curing Through Polyimide Films*

In order to effectively cure acrylate resins through polyimide film, the emitted wavelengths of the illuminating light source must be matched with the absorptivity spectrum of visible light photoinitiator (which was in turn selected based upon the wavelengths transmitted through the film). Mercury arc lamps are by far the most common light source used in industrial photo-cure processes. However, an alternative light source was necessary for use in this investigation since the mercury arc lamps emit very little light above 500 nm. Significant advances have been made in developing light emitting diodes (LEDs) to provide high intensity in the visible region of the spectrum. LEDs offer many advantages over traditional broad-spectrum lamps, the most important of which is high efficiency resulting in relatively low energy consumption. In addition, LEDs can instantly be switched on and off, are ideal for heat sensitive materials (no stray IR emission), have long lifetimes (on the order of tens of thousands of hours of lamp life), do not contain hazardous vapors, and are light-weight/compact for customizable design and scaling. Since these advantages make LEDs ideal for photo-curing applications, a high-intensity green LED (PhlatLight, Luminus Devices, Inc.) was selected for illuminating the visible-light photopolymerization systems evaluated in this study. A 150 W xenon light source (MAX-150, Asahi Spectra) was also selected which provides higher intensity than the LED lamp. Both light sources provide illumination in the 400 – 800 nm range required for this application, and do not interfere with the absorptivity of the polyimide film. The emission spectra for both lamps, along with the polyimide film absorptivity, are shown in Figure 5.3. The emission spectra

were collected using the miniature fiber optic spectrometer positioned 8 cm from the light source. The total irradiance in this wavelength region (the integrated peak area from 400 – 800 nm) was 130 mW/cm² for the xenon lamp and 10 mW/cm² for the green LED. The 150 W xenon lamp is therefore a much more efficient and highly intense light source across the entire visible region than the LED.

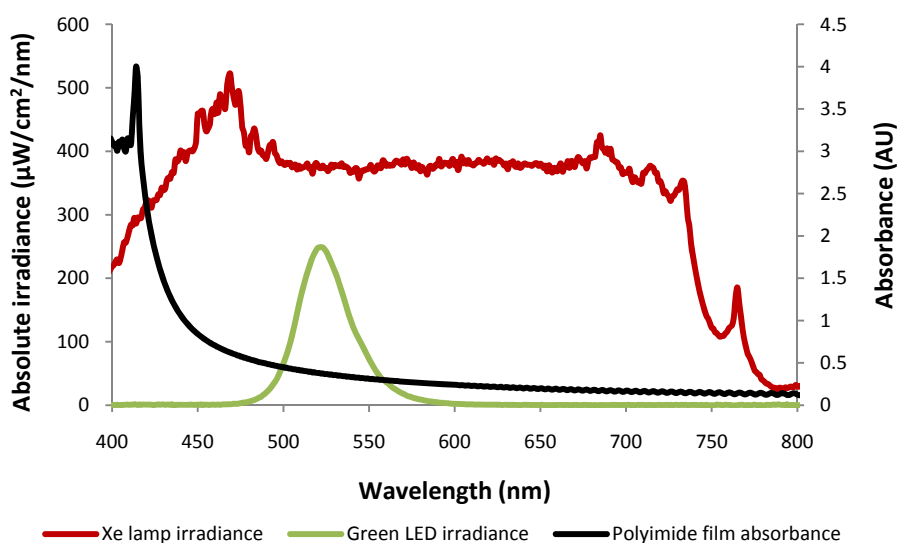


Figure 5.3. Absolute irradiance of 150 W xenon lamp and high intensity green LED compared with absorbance of polyimide film.

5.2.2.2 Visible Light Photoinitiator Systems Selected for Photo-curing Through Polyimide Films

Illumination through a polyimide film necessitates the use of visible light photoinitiators. Unlike typical ultraviolet photoinitiators, the energy of a photon in the visible region of the spectrum is generally less than the bond dissociation energy for most organic compounds, including photoinitiators. Therefore, visible-light-induced initiator systems are generally bimolecular initiator systems in which the active centers are produced via electron

transfer followed by proton transfer from the electron donor to the excited photosensitizer [55]. Five photosensitizers compatible with the wavelength range of the illuminating light required were screened for use in this study, in combination with the electron donor MDEA. FTIR conversion was measured for HEA monomer containing concentrations of 0.07 wt% of photosensitizer and 2.5 wt% MDEA. The 15 μm films were laminated between NaCl slides as described in section 5.2.1, and illuminated for 1 minute with the green LED held 4 inches from the sample.

Table 5-1. Overall conversions by FTIR for various visible light photoinitiators illuminated with green LED for one minute 8 cm from light source.

Photoinitiator	Monomer	Photoinitiator wt%	MDEA wt%	Conversion
Eosin Y	HEA	0.07	2.5	77.2 %
H-Nu 535	HEA	0.07	2.5	66.4 %
Erythrocin B	HEA	0.07	2.5	51.7 %
Rose Bengal	HEA	0.07	2.5	< 30 %
Camphorquinone	HEA	0.07	2.5	< 30 %

A comparison between the five photosensitizers shown in Table 5-1 reveals that Eosin Y was found to provide the highest HEA conversion as measured by FTIR. The absorption spectrum and molecular structure of the neutral form (spirit soluble) of Eosin Y is shown in Figure 5.4.

The electron-transfer/proton-transfer reaction between Eosin Y and MDEA shown in Figure 5.5 has been previously investigated [56]. The triplet state of Eosin Y, $[\text{EYss}]^*$, and the amine, A-H, form an exciplex similar to that described for other dyes and amines.

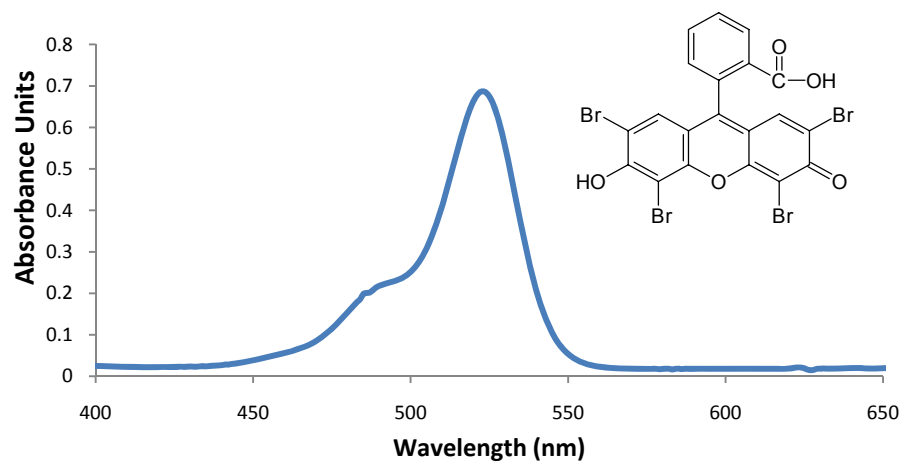


Figure 5.4. Eosin Y (spirit soluble) absorbance spectrum and molecular structure (0.001 wt% Eosin Y in methanol)

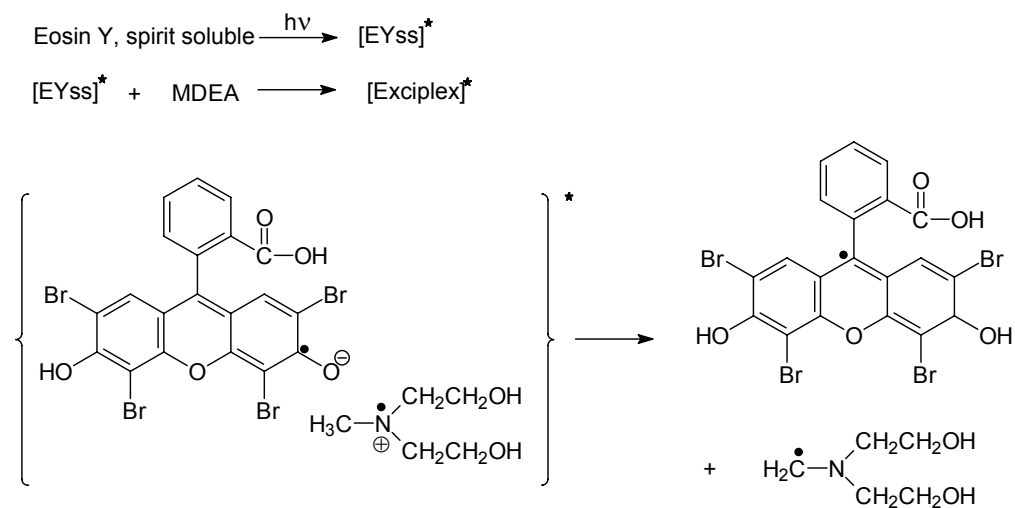
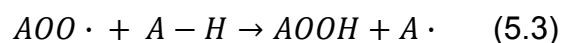
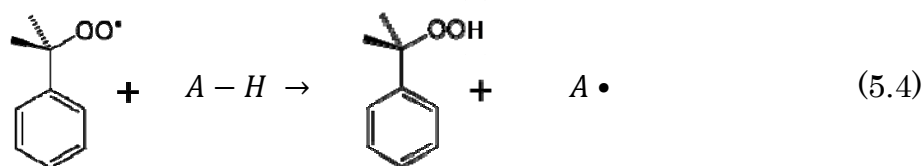


Figure 5.5. Direct reaction between Eosin Y spirit soluble and MDEA [56].

As shown in Figure 5.5, the exciplex decomposes in the last step of the mechanism by an electron-proton transfer step to form an amine radical, $A\cdot$, and a dye based radical. The amine radical, $A\cdot$, becomes the active initiator species. The amine radicals themselves suppress oxygen inhibition by the consumption of oxygen in a radical chain process:



Many investigators have illustrated that the effectiveness of a two-component electron transfer initiator system, such as the one shown in Figure 5.5, can be enhanced by the addition of a third component, such as an iodonium salt. However, ionic species may be undesirable for the photocurable ACF application proposed in this contribution due to the possible corrosive effects of the salt on the conductive elements of the flexible printed circuit. In order to enhance the conversion of the two-component Eosin Y/MDEA system, a nonionic hydroperoxide was investigated for use as a chemical scavenger of dissolved oxygen. Cumene hydroperoxide has been used to enhance the conversion of typical 3-component visible-light photoinitiation systems, and has been shown to increase conversion over the dye/amine system alone [57]. An example of a reaction mechanism by which cumene hydroperoxide may consume dissolved oxygen is shown in Equation 4, which shows the hydroperoxide reacting with the amine to form additional amine radicals, which in turn consume more dissolved oxygen according to Equation 5.2.



The addition of 0.15 wt% cumene hydroperoxide (Sigma Aldrich) to the two-component Eosin Y/MDEA visible light photoinitiator system resulted in a 10% increase in conversion measured by FTIR. The overall conversion for the 3-component system was further increased by replacing the LED light source with the 150 W xenon lamp. Figure 6 summarizes the average conversion for 3 or more experiments for 2 minute and 1 minute xenon lamp illumination times (95% and 92% for bars (a) and (b) respectively) compared with experiments illuminated using the green LED, both with and without the addition for the cumene hydroperoxide additive (87% and 77% for bars (c) and (d) respectively).

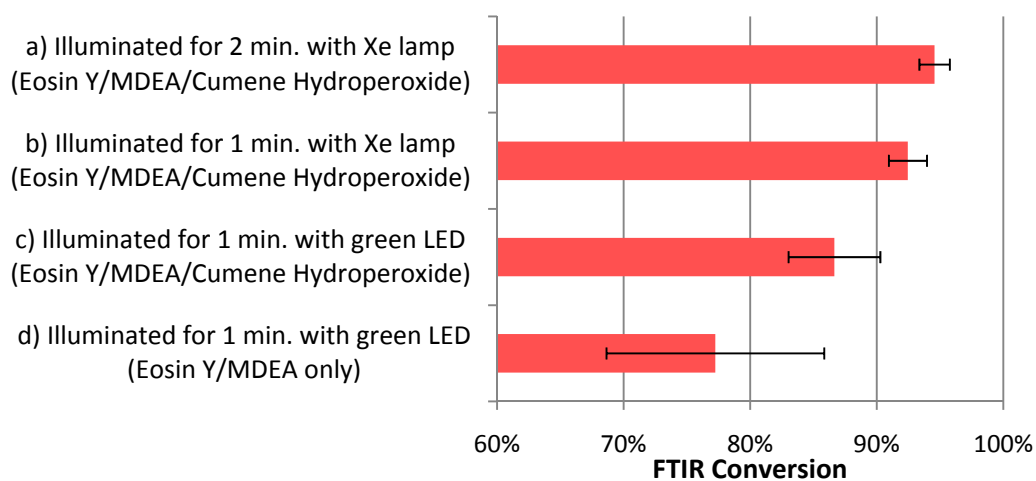


Figure 5.6. Summary of FTIR conversion results for HEA films using Eosin Y/MDEA photoinitiator system with cumene hydroperoxide.

The optimal 95% conversion shown in Figure 5.6a, for the 3-component system illuminated for two minutes with the xenon lamp, may be representative of a desirable conversion for an application such as ACF. Kinetic profiles of this optimized 3-component visible-light induced polymerization were investigated using real-time FTIR. FTIR spectra were

collected at 1 second increments throughout the two minute irradiation period, and Eqn. 1 was used to calculate the change in conversion as a function of time. The results are shown in Figure 5.7. The contribution from dark cure to the overall conversion after the light was shuttered off is about 1%.

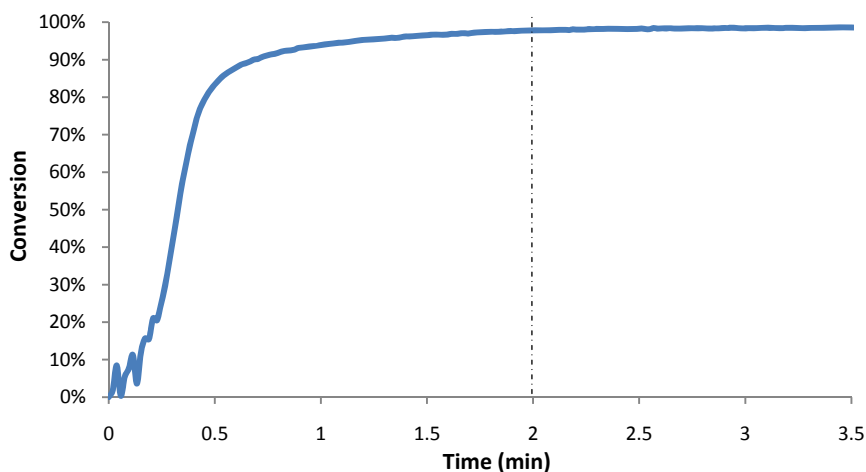


Figure 5.7. Real-time FTIR conversion during 2 minute illumination with 150 W xenon lamp.

Because the proposed ACF application may require illumination through a polyimide film, the effect of the polyimide absorbance on the overall conversion was investigated. FTIR was again used to measure the HEA conversion. In these experiments, a polyimide film of desired thickness was placed over the NaCl slides during the illumination. The IR spectra were measured before and after illumination, and overall conversion was calculated using Equation 5.1. Since the polyimide film absorbs strongly in the IR region of the spectrum, real-time FTIR measurements were not possible for these experiments. These results are summarized in Figure 5.8 below. When illuminated through polyimide films, the conversion for the

optimized 3-component photoinitiator HEA system containing cumene hydroperoxide was reduced to 90% and 80% for the 25 μm and 75 μm polyimide film thicknesses respectively.

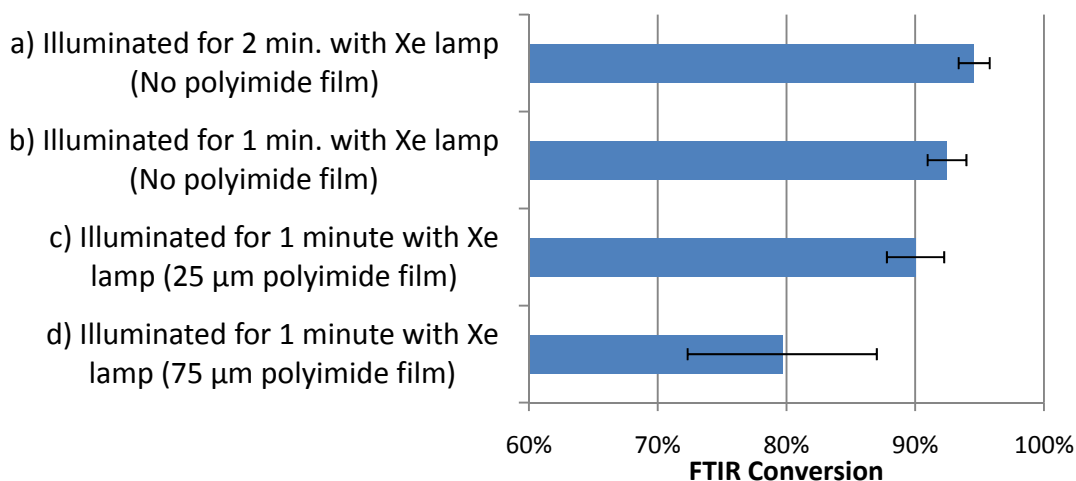


Figure 5.8. FTIR conversion for HEA films using 3-component Eosin Y photoinitiator system.

Although the addition cumene hydroperoxide was shown to enhance the overall conversion in HEA films over the two-component visible light photoinitiator system alone, the cumene hydroperoxide may possibly be an undesirable additive from a manufacturing perspective. Therefore, experiments were conducted to optimize the 2-component system containing only Eosin Y and MDEA, in an attempt to obtain the same level of conversion as the 3-component system (>90% conversion). To meet this objective, several molar concentration ratios of MDEA/Eosin were screened, ranging from 180 to 250 mol MDEA per mol Eosin Y. These studies revealed an optimal molar ratio of approximately 250 mol/mol, and this ratio was maintained for the next set of studies in which the overall concentration was varied to identify the global optimum. Figure 5.9 shows the results of the concentration study,

and indicates the optimum composition of the 2-component system: 0.074 wt% Eosin Y and 3.69 wt% MDEA.

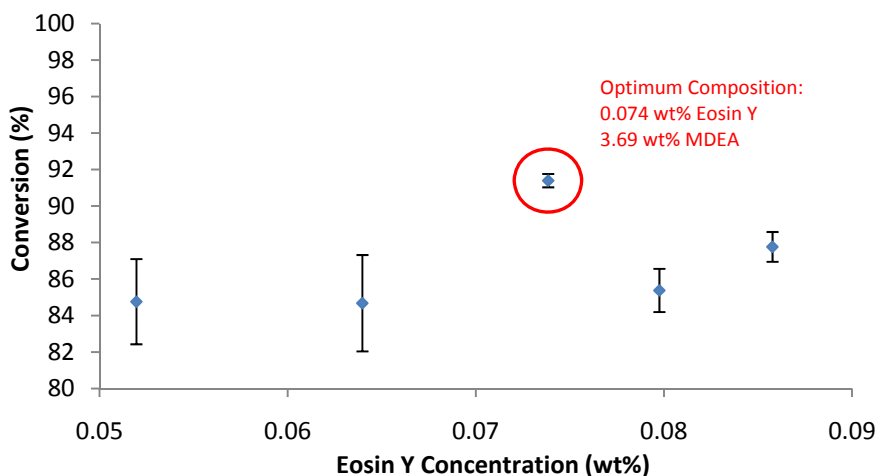


Figure 5.9. Optimization of 2-component Eosin Y system. Conversions calculated by FTIR. 1 minute illumination with 150 W Xe lamp.

Kinetic profiles of the optimized 2-component visible-light induced polymerizations were investigated using real-time FTIR. Real-time kinetic data were collected during illumination, and the corresponding profiles of the conversion as a function of time are shown in Figure 5.10. The data in the figure illustrate that the real time conversion profile for the optimized 2-component system at least as high as that of the original 3-component initiator system at all times. The overall conversion for both samples at one minute, when the light was shuttered off, was approximately 92%. The contribution from dark cure to the overall conversion after the light was shuttered off is about 0.5%.

These FTIR data reveal that the optimized 2-component initiator system results in an overall conversion of 92% (for one minute illumination) with no additional additives. Therefore, the same approximate level of conversion

that was achieved with the 3-component system including cumene hydroperoxide has been accomplished with the optimized 2-component system containing only Eosin Y and MDEA alone.

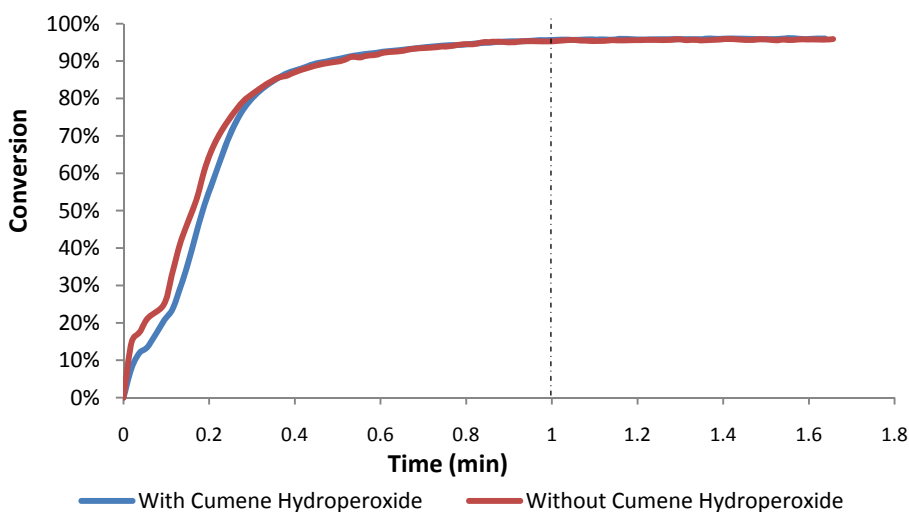


Figure 5.10. Real-time FTIR conversion during 1 minute illumination with 150 W xenon lamp. HEA monomer, Eosin Y photoinitiator system with and without cumene hydroperoxide.

Based upon the mechanistic considerations shown in Equations 5.2-5.4, it is possible to speculate on the reason that the 2-component system containing excess amine performs as well as the 3-component initiator system containing cumene hydroperoxide. Equation 5.4 indicates that the hydroperoxide reduces the oxygen inhibition by interacting with the amine to produce amine radicals. These amine radicals in turn consume oxygen as shown in Equations 5.2 and 5.3. However, Figure 5.5 illustrates that the Eosin dye also interacts with the amine to produce the same amine radicals. This is the reason that the increased concentrations of the Eosin Y and the amine can have the same effect on the reaction kinetics as the addition of the hydroperoxide.

5.3 Characterization of Shadow Cure

5.3.1 Materials and Experimental Methods

The optimized visible light photoinitiator systems described above were used to photopolymerize 100 μm thick HEA films representative of the proposed ACF system. Transparent electrode-imbedded polyimide films were used as a substrate through which an acrylate resin was photo-cured, similar to the ACF system depicted in Figure 5.2. Acrylate films were produced by placing a droplet of monomer sample into a 1 cm^2 PET spacer with a thickness of 100 μm . The spacer and monomer were laminated between an electrode imbedded polyimide film on the top surface, and a coated separation film on the bottom surface. The laminated film was then placed on a black surface, and illuminated from the top down for 1 minute using the 150 W xenon lamp 8 cm from the sample. To visually observe the degree of cure, the films were removed from the substrates, placed onto glass slides, and observed with a polarized optical microscope (Nikon Corporation).

5.3.2 Results and Discussion

5.3.2.1 Visualization of the Spatial Cure Profile for Illumination Through an Electrode-Containing PET Film

Since the electrodes imbedded in the substrate are opaque, they serve to shade, or mask, alternating regions of the resulting photopolymer from the initiating light. Figure 5.11 shows an example of the electrode imbedded polyimide film that was used as a substrate in these experiments, through which the HEA resin was photopolymerized using visible light. The electrodes imbedded in the film shown in Figure 5.11 are 100 μm wide and spaced 100 μm apart. This example is referred to 100 x 100 μm spacing.

Polyimide films with 50 μm wide electrodes were used for the results presented below (50 x 100 μm spacing).

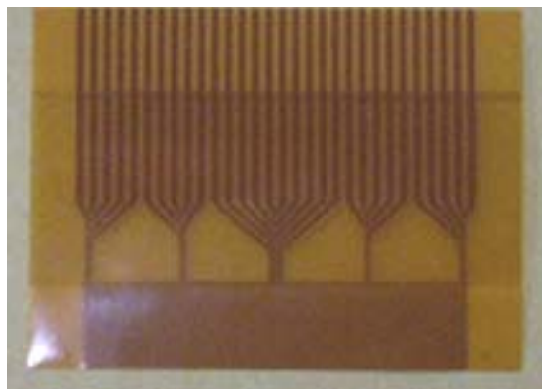


Figure 5.11. Example of an electrode imbedded polyimide film with 100 x 100 μm electrode spacing.

HEA monomer containing the optimized 2-component Eosin Y visible light photoinitiator system was laminated between the electrode imbedded polyimide film and the separation film substrates, and the acrylate was cured using the method described above. The resulting HEA films were easily removed intact from both the separation film and the polyimide film containing the electrodes, and appeared to be fully polymerized by macroscopic observation. Using the polarized light microscope, the appearances of the illuminated and unilluminated regions were compared. Figure 5.12 shows an optical micrograph of an HEA film containing the 2-component photoinitiator system photopolymerized by illumination with the 150 W xenon lamp for one minute through an electrode imbedded polyimide film. In the illuminated regions of the sample, the pale yellow color is indicative of a transparent region because the settings on the polarized light microscope resulted in a pale yellow background when there was nothing in

the field of view. Within these regions, the Eosin Y was photoreduced, thereby losing its color (photobleached). In contrast, the areas shaded by the gold electrodes appear as a dark pink or light brown color in the polarized light micrograph. The films appeared a pale pink when observed by the naked eye. This color likely arises from the presence of unreacted Eosin Y photoinitiator. Both the yellow and pink regions appear to be fully cured, indicating the possibility of shadow cure.

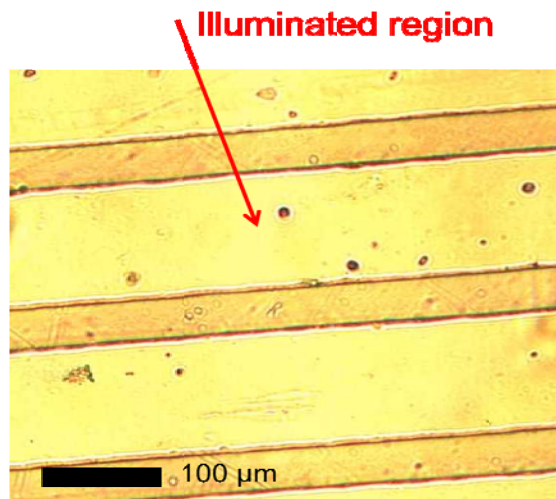


Figure 5.12. Polarized light micrographs of HEA films containing optimized 2-component Eosin Y photoinitiator system. Illuminated for 1 minute with 150 W xenon lamp through a PET film with 50 x100 μm electrode spacing.

Due to shadow cure, polymerization may occur even in the unilluminated regions. In order to further understand the effect of shadow cure, the films were visualized using cross-sectional micrographs. Small cross-sections of the acrylate films described above were cut using a razor blade and placed on microscope slides. Figure 5.13 shows the cross-sectional polarized light micrograph of a film produced by illuminating through a polyimide film with 50 x 100 μm gold electrode spacing.



Figure 5.13. Polarized light micrograph showing cross-section of HEA film containing optimized 2-component Eosin Y photoinitiator system. Illuminated for 1 mintue with 150 W xenon lamp through PET film containing 50x100 μm gold electrode spacing.

The shape of the top surface of the film arises from the imprint created by the gold electrodes, therefore the regions under the indentations were not illuminated by the xenon lamp. The textured appearance provided by the polarized light microscope indicates that this film may be fully cured polymer through the $\sim 100 \mu\text{m}$ depth in both the illuminated and unilluminated regions.

5.3.2.2 Predicting free-radical active center diffusion into unilluminated regions

The generation of free-radical active centers produced during illumination is well understood, and may be described by a previously reported multi-wavelength simulation [31]. For an accurate description of the spatial photoinitiation profiles produced during the illumination step, it

is important to account for changes in the photoinitiator concentration as the initiator is consumed in the illuminated region. Therefore, the active center generation as a function of position in a sample immediately after the illumination has ceased depends on a number of factors including the light intensity, initial photoinitiator concentration and absorbance, and exposure time. This model is based upon the following set of fundamental differential equations which govern the evolution of the light intensity gradient and initiator concentration gradient for multi-wavelength illumination.

The governing equations for light intensity, photoinitiator concentration, and photolysis product concentration were given in Chapter 4 for polychromatic light (Equations 4.1-4.7). The solution to this set of equations provides detailed information regarding the time-evolution of the light intensity gradient and the initiator concentration gradient. Simulation results for the 2-component Eosin Y photoinitiator system illuminated with the 150 W xenon lamp are shown in Figure 5.14 and Figure 5.15. Figure 5.14 shows the light intensity gradient as a function sample depth over a 1 minute illumination period. Initially, the light intensity drops off rapidly at the surface of the sample (where depth = 0). As the photoinitiator absorbs the initiating light and photobleaches, the light then penetrates farther into the depth of the system. At one minute of illumination, the light has fully penetrated up to a few millimeters in depth. The wavelength range used for this analysis was 425 - 575 nm, which is the range over which the photoinitiator Eosin Y absorbs.

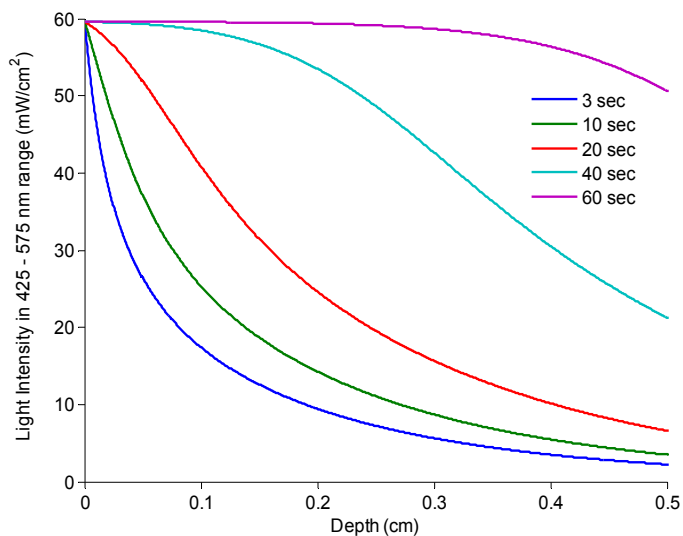


Figure 5.14. Light intensity profile of a 3-component Eosin Y system initiated with a 150 W xenon lamp ($I_0 = 60 \text{ mW/cm}^2$ in the 425 - 575 nm spectral region).

Figure 5.15 shows the depleting concentration of photoinitiator with illumination time. As the initiating light is absorbed at the surface (depth = 0), free-radical active centers are created, and the photoinitiator concentration diminishes. These results illustrate that for a one minute illumination time, the photoinitiator is completely consumed for the first 2 millimeters (2000 μm), indicating films with a thickness less than a few hundred micron will have the same conversion throughout their depth. Therefore, the FTIR studies of 15 μm films are equally valid for films over 100 μm thick.

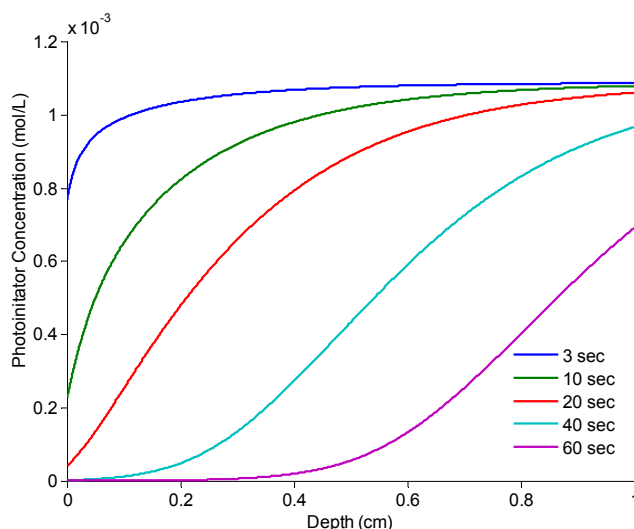


Figure 5.15. Photoinitiator concentration profile for a 3-component Eosin Y system initiated with a 150 W xenon lamp ($I_0 = 60 \text{ mW/cm}^2$ in the 425 - 575 nm spectral region).

Previously in this contribution, cationic active centers were shown to migrate after the illumination ceased, resulting in shadow cure in unilluminated regions. A similar analysis was used to predict the diffusion of free-radical active centers into the unilluminated regions of the masked polymer films. Figure 5.16 shows the geometrical configuration used to develop the free-radical diffusion analysis. The shadow regions indicated in the schematic are designed to represent the regions behind the electrodes for the conductive adhesive film application. Within the illumination region, the one-dimensional active center generation model was used to calculate the light intensity and photoinitiator concentration in the z direction, as shown in Figure 5.14 and Figure 5.15 above.

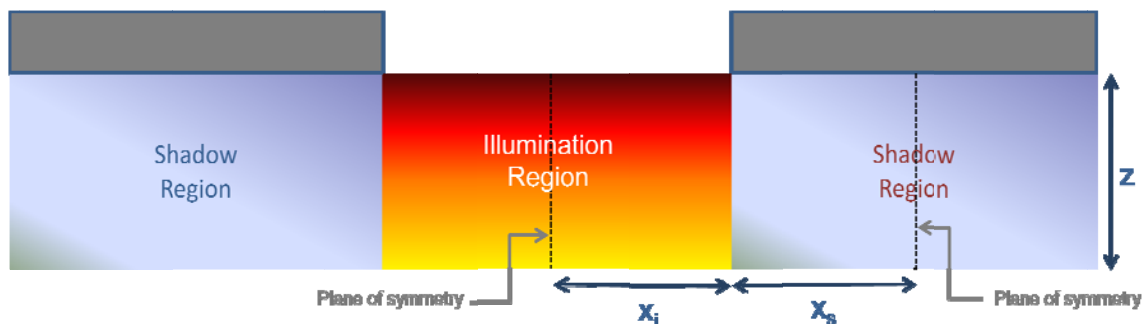


Figure 5.16. Geometrical configuration for analysis of free-radical active center diffusion.

Since the results in Figure 5.14 and Figure 5.15 have shown that the light intensity and photoinitiator concentration are uniform throughout the 100 μm thick films, the post-illumination diffusion analysis can be simplified to a one-dimensional case where free-radical active centers diffuse into the shadow regions in one direction only: the x -direction. In addition, due to the periodic nature of the sample geometry resulting from the evenly spaced electrodes, there is a plane of symmetry at the center of each illuminated region and each shaded region. Therefore, the system may be analyzed by considering a single repeat cell that extends from the middle of the illuminated region to the middle of the shaded region. Note that if the cell width of the illuminated region is x_i , and the cell shadow width x_s , a 100 x 100 μm electrode spacing would give $x_i = 50 \mu\text{m}$, and $x_s = 50 \mu\text{m}$.

Post-illumination diffusion of free-radical active centers from the illuminated region into the shadow region is governed by Fick's second law:

$$\frac{\partial C_{AC}(x,t)}{\partial t} = D_{AC} \frac{\partial^2 C_{AC}(x,t)}{\partial x^2} \quad (5.5)$$

In this equation, $C_{AC}(x,t)$ is the active center molar concentration at time t , and width x , and D_{AC} is the diffusion coefficient for the active centers in units of length²/time. The no-flux boundary condition for this system indicates that there is no transport of initiator or photolysis product across the planes of symmetry.

$$\frac{\partial C_{AC}(x,t)}{\partial x} = 0 \quad (5.6)$$

The active center concentration in free-radical polymerizations rapidly achieves a pseudo-steady-state value in the range of 10^{-7} to 10^{-9} mol/L [58]. The concentration of active centers remains very low because the rate of the termination reaction, which consumes two free-radical active centers, is proportional to the square of the free-radical concentration. Therefore, any increase in the radical concentration leads to a greater increase in the termination rate (for example doubling the radical concentration would quadruple the termination rate), thereby driving the radical concentration back down. The precise value of the active center concentration depends upon the initiator concentration and the light intensity as well as the termination rate constant.

At a given location in the reaction system, the local active center concentration changes with time due to two simultaneous processes: diffusion and chemical termination. The governing equation for the local active center concentration accounting for both of these processes is shown in Equation 5.7 below:

$$\frac{\partial C_{AC}(x,t)}{\partial t} = D_{AC} \left[\frac{\partial^2 C_{AC}(x,t)}{\partial x^2} \right] - 2k_t C_{AC}(x,t)^2 \quad (5.7)$$

Equations 5.6-5.7 were solved numerically to provide profiles of the active center concentration as a function of the shadow cure time in the x -

direction. To solve this equation for the system geometry shown in Figure 5.16, initial conditions and boundary conditions must be established. For the initial condition, the active center concentration is assumed to be uniform in the illuminated region, and zero in the shaded region. This initial condition allows for determination of the extent of diffusion as a function of time. Two boundary conditions are required, and the symmetry of the system geometry leads to a flat concentration profile at each plane of symmetry (the no flux boundary condition, Equation 5.6). The active center diffusion coefficient was estimated to be 1×10^{-5} cm²/sec based on literature reported values for acrylates [59]. The termination constant that was used was 1×10^5 L/mol-sec [58].

Results for representative values of the active center concentration profiles are shown in Figure 5.17 and Figure 5.18. To illustrate the effect of termination reactions on the potential for diffusion to lead to shadow cure, simulations were performed with and without termination for two different initial active center concentrations. Figure 5.17a shows the simulation results for 10^{-8} active center concentration generated in the illumination region, assuming no termination during the post-illumination diffusion process (Equation 5.5). These active center diffusion profiles can be compared to the case shown in Figure 5.17b where the termination rate is included (Equation 5.7). Since the active center concentration is very low, the termination term in Equation 5.7 is negligible compared to the diffusion term, and the profiles are indistinguishable from each other.

For an increased initial active center concentration of 10^{-6} mol/L, the simulation results for diffusion only, shown in Figure 5.18a, are similar to those at the lower concentrations in Figure 5.17. However, due to the higher initial active center concentration, the termination rate is higher, resulting in

a reduction in active center concentration as it diffuses into the unilluminated region, as shown in Figure 5.18b.

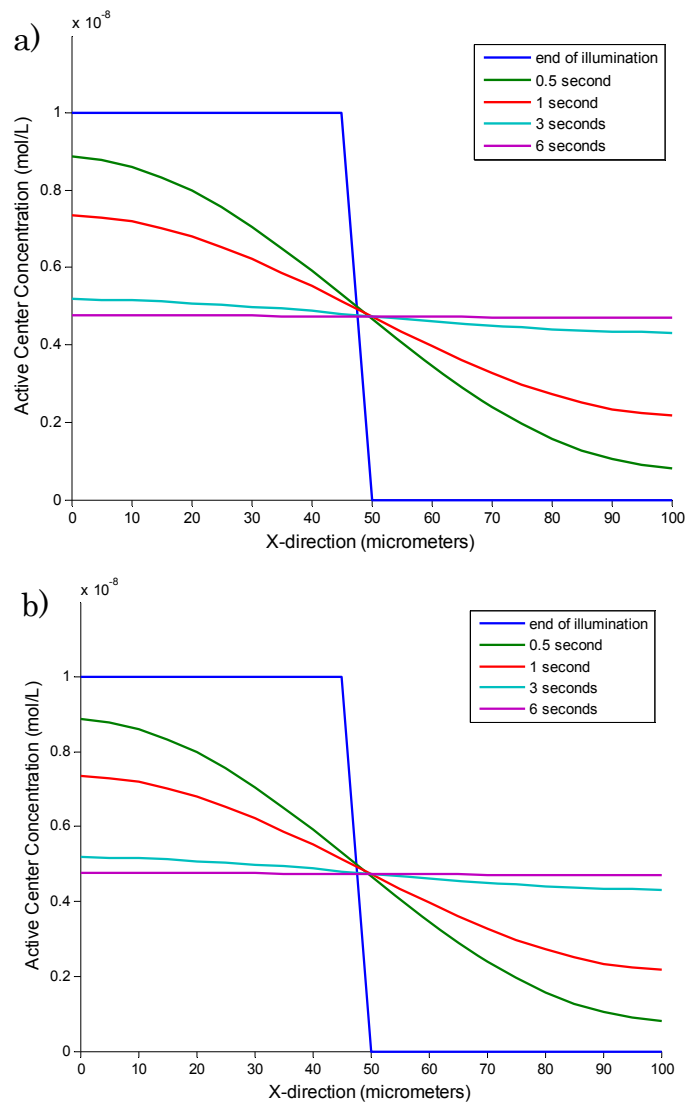


Figure 5.17. Post-illumination concentration profiles for active centers diffusing from illumination region (left side: 0-50 μm) to unilluminated region (right side: 50-100 μm) for initial concentration of 10^{-8} mol/L. a) diffusion only (Equation 5.5), b) diffusion and termination (Equation 5.7)

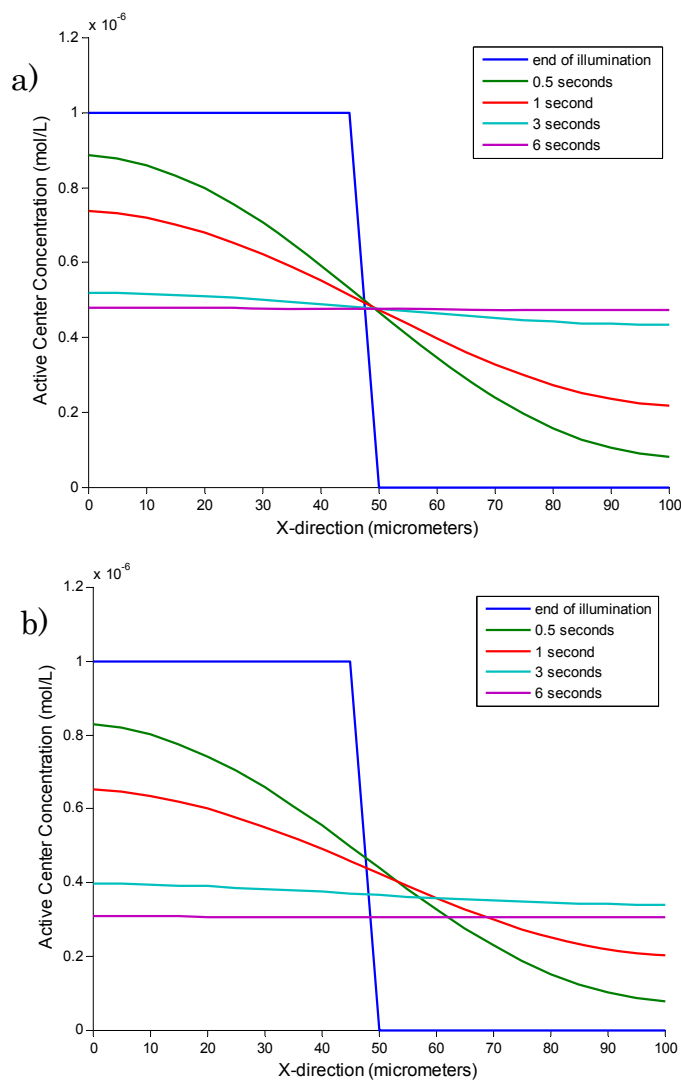


Figure 5.18. Post-illumination concentration profiles for active centers diffusing from illumination region (left side: 0-50 μm) to unilluminated region (right side: 50-100 μm) for initial concentration of 10^{-6} mol/L. a) diffusion only (Equation 5.5), b) diffusion and termination (Equation 5.7)

5.4 Conclusions

According to the theoretical results reported in section 5.3.2.2, the diffusion of free-radical active centers into the unilluminated regions takes place very rapidly. In all cases presented in Figure 5.17 and Figure 5.18, the diffusion is complete approximately 6 seconds after the illumination has ceased. Since these are conservative estimates of diffusion rates, assuming no diffusion during the illumination period, these results support the hypothesis that the shaded regions of the films are likely polymerized due to shadow cure. Macroscopic observation of the cured films using the polarized light microscope also support this, showing cured polymer in regions where unilluminated photoinitiator still exists.

In this detailed discussion of a specific application where shadow cure is important on an extremely small dimensional scale, the potential for free-radical active centers to provide shadow cure into unilluminated regions is demonstrated. In the future, as the size of microelectronic components becomes smaller, shadow cure in free-radical photopolymerizations may become feasible for a wide variety of applications.

CHAPTER 6

CONCLUSIONS AND RECOMMENDATIONS

In this contribution, the mobility of photo-generated active centers has been shown to result in shadow cure in a variety of applications. First, the ability of long-lived cationic active centers to effectively cure coatings pigmented with carbon black was investigated. Secondly, the mobility of cationic active centers was evaluated for use in curing photopolymers with complex geometries. Finally, the potential of shadow cure in free-radical systems was investigated for a specific microelectronics application in which radical active centers migrate over relatively small dimensional scales. In all of these applications, a theoretical description of active center generation and mobility was used to characterize shadow cure, and these findings were compared with experimental results. In this chapter, an overview is provided of the major conclusions from each section of the research on these different applications, and recommendations for future research into these areas are summarized.

6.1 Cationic Photopolymerization of Systems Pigmented with Carbon Black Nanoparticles

The active centers responsible for cationic photopolymerizations are essentially non-terminating, and continue to propagate after the illumination has ceased. The mobility of the long-lived cationic active centers was investigated for the cure of coatings pigmented with carbon black nanoparticles. The spatial and temporal evolution of the cationic active center concentration profile during illumination was analyzed using the set of differential equations that govern the light intensity gradient and photoinitiator concentration gradient for polychromatic illumination. Due to

the sharp light intensity and concentration gradients established in the pigmented systems, the photoinitiator diffusion during the illumination period was found to have a marked effect on the resulting active center concentration profiles. Analysis of the active center reactive diffusion during the post-illumination period revealed that migration of the active centers leads to cure beyond the illuminated depth. A kinetic analysis performed by coupling the active center concentration profiles with the propagation rate equation yielded estimates of cure time for coatings of varying thickness and pigment loading. These theoretical cure times showed good agreement with experimental results obtained for photopolymerizations of cycloaliphatic diepoxide coatings pigmented with a monodisperse carbon black with mean hydrodynamic radius of 29.2 nm. This slightly basic furnace black pigment was found to act as a mild inhibitor of cationic photopolymerization. The results presented indicate that the long lifetimes and reactive diffusion of cationic active centers may be used for effective curing of carbon black pigmented coatings.

This comprehensive approach could be applied to other types of pigmented or filled systems. The characterization techniques presented here for evaluating the carbon black nanoparticle dispersion and optical properties may not be applicable for other types of pigments or fillers. Some pigments are large aggregates that scatter and reflect light in addition to absorbing the UV irradiation necessary for photopolymerization. In order to completely characterize other types of pigments or fillers in a photopolymer, the particle size and dispersion would need to be rigorously evaluated. Other systems that could be evaluated in this manner for their ability to shadow cure include nanocomposites, conductive adhesives, and other types of filled photopolymer systems.

6.2 Cationic Active Center Mobility in Photopolymerization Systems with Complex Geometries

This section provided a theoretical and experimental investigation of shadow cure in multiple spatial dimensions for cationically photopolymerized systems. Cationic photopolymerization is an alternative to free-radical photopolymerization offering several unique advantages, especially for use in curing coatings on substrates with complex geometries. First, the fundamental set of governing differential equations was used to describe the generation and post-illumination diffusion of cationic active centers in two spatial dimensions. The results of this analysis provided a fundamental understanding of shadow cure in multiple dimensions so that these descriptions could be applied to systems with complex geometries. Secondly, this approach was used to predict shadow cure distance in a direction perpendicular to the illumination for cationically polymerized epoxide coatings by coupling the diffusion analysis with the kinetic rate equation. These results showed very good agreement between theoretical predictions and the experimental findings which showed that the coatings shadow cured over substantial distances. Finally, a novel method was presented in which long-lived cationic centers that are photo-generated in only one layer of a multi-layered coating lead to cure in multiple layers, even if the layers are added after illumination. Using this method, a fully cured coating comprised of two or more layers can be completely cured using only one illumination step. All of these examples of shadow cure in cationic photopolymerization indicate that there may be tremendous potential in many applications to use cationic photopolymerization to cure systems with complex geometries.

A more rigorous kinetic analysis would be required in order to completely characterize the extent of cure in unilluminated regions. For example, the effective diffusion coefficients reported in this contribution are

likely influenced by variables that have not yet been investigated. For example, the conversion in the illuminated region during the post-illumination diffusion period may have an effect on the diffusion rate as well as the overall extent of shadow cure. The detailed active center concentration profiling demonstrated here should be coupled with a more detailed kinetic analysis that includes evaluation of actual conversions in order to completely characterize the shadow cure process. Raman spectroscopy, FTIR, or thin film calorimetry could be utilized for this purpose.

The mobility of the cationic active centers responsible for curing the multiple-layer systems should be further investigated. For the purposes of the multiple coating layer application, the cycloaliphatic diepoxide used throughout this contribution (EEC) was co-polymerized with 2-butoxymethyl-oxirane (BMO) so that the viscosity could be reduced and the coating could be applied with an airbrush. A detailed kinetic analysis of the copolymer would need to be performed in order to fully characterize the extent of shadow cure in this system. Again, conversions in the multi-layered photopolymer films would need to be measured so that the reactive diffusion process could be completely described.

6.3 The Potential of Free-Radical Shadow Cure

Free-radical active centers have significantly shorter lifetimes than cationic active centers, and tend to terminate immediately upon cessation of the photo-curing light source. However, some cutting edge applications for photopolymers may necessitate the use of shadow cure in free-radical systems, specifically in some microelectronics applications. Advances in microelectronics have led to demand for devices with increasingly small components. Despite the extremely short lifetimes of the free-radical active

centers, mobility of free-radicals within these minute dimensional scales may become a possibility. This section presented one formulation that could be developed to utilize free-radical shadow cure for the application anisotropic conductive adhesive film in flexible printed circuits. The selection process for both the photoinitiator and illuminating light source for this application was described. Free-radical mobility into unilluminated regions behind the imbedded electrodes in an ACF application was characterized using an analysis of post-illumination diffusion and termination reactions. The results for free-radical shadow cure showed that the diffusion of active centers into the unilluminated regions takes place very rapidly, supporting the hypothesis that the shaded regions of the films are likely polymerized due to shadow cure. Macroscopic observation of cured films indicated that cured polymer existed in shaded regions containing unilluminated photoinitiator.

In order to completely characterize the extent of cure in unilluminated regions of a free-radical photopolymerization system, a more rigorous technique for spatial profiling would be required. Raman spectroscopy in combination with microscopy would provide a non-invasive method for extensively characterizing the extent of cure in the shaded regions of the film.

A novel concept for inducing shadow cure in free-radical systems was conceived during this research, but not thoroughly investigated. The idea is to introduce a fluorescent additive. The absorption of light by the fluorescent additive can trigger the emission of another photon with a longer wavelength. In this manner, the fluorescent additive could diffuse into unilluminated regions, producing a multidirectional fluorescence, potentially resulting in indirect illumination within shadow regions. Since visible light photosensitizers were used for this study of free-radical shadow cure, there is a potential that a photosensitizer could be found that would double as both a

photosensitizer and a fluorescent additive. Fluorescence spectroscopy would be required to begin to evaluate additives for such a system, and fluorescence monitoring would need to be coupled with spatial profiling to evaluate the effect on shadow cure.

More than any other area covered by this research, free radical shadow cure has the greatest potential to grow into uncharted territory. In typical applications currently employing photopolymerization for microelectronics, such as printed circuit boards, the goal is usually to attain high resolution. Therefore, shadow cure is something that tends to be avoided in these industries. Deeper UV wavelengths are used to enhance resolution, and diffusion of active centers presents a limitation rather than an asset. Applications have generally not yet been developed in which diffusion of free-radical active centers could be used to cure unilluminated regions of a photopolymer.

REFERENCES

1. Bongiovanni R, Montefusco F, Priola A, Macchioni N, Lazzeri S, Sozzi L, Ameduri B. High performance UV-cured coatings for wood protection. *Progress in Organic Coatings* 2002;45(4):359-363.
2. Weiss KD. Paint and coatings: A mature industry in transition. *Progress in Polymer Science* 1997;22(2):203-245.
3. Benson RS. Use of radiation in biomaterials science. *Nuclear Instruments and Methods in Physics Research Section B: Beam Interactions with Materials and Atoms* 2002;191(1-4):752-757.
4. Zou Y, Armstrong SR, Jessop JLP. Apparent conversion of adhesive resin in the hybrid layer, Part 1: Identification of an internal reference for Raman spectroscopy and the effects of water storage. *Journal of Biomedical Materials Research Part A* 2008;86A(4):883-891.
5. Zou Y, Jessop JLP, Armstrong SR. Apparent conversion of adhesive resin in the hybrid layer, Part II: In situ studies of the resin-dentin bond. *Journal of Biomedical Materials Research Part A* 2009;89A(2):355-362.
6. Hancock A, Lin L. Challenges of UV curable ink-jet printing inks - a formulator's perspective. *Pigment & Resin Technology* 2004;33(5):280-286.
7. Edison SE, Madhusoodhanan D, Nagvekar D, Wilson M, Ellison M. UV curable inkjet inks: Is there anything they can't do? *Proceedings of RadTech UV/EB Technology Expo & Conference, Chicago, IL, 2008.*
8. Lin YC, Zhong J. A review of the influencing factors on anisotropic conductive adhesives joining technology in electrical applications. *Journal of Materials Science* 2008;43(9):3072-3093.
9. Lu D, Wong CP. Novel conductive adhesives for surface mount applications. *Journal of Applied Polymer Science* 1999;74(2):399-406.
10. Mir I, Kumar D. Recent advances in isotropic conductive adhesives for electronics packaging applications. *International Journal of Adhesion and Adhesives* 2008;28(7):362-371.
11. Fang N, Sun C, Zhang X. Diffusion-limited photopolymerization in scanning micro-stereolithography. *Applied Physics A: Materials Science & Processing* 2004;79:1839-1842.
12. Schmitt H, Frey L, Ryssel H, Rommel M, Lehrer C. UV nanoimprint materials: Surface energies, residual layers, and imprint quality. *Journal of Vacuum Science & Technology B: Microelectronics and Nanometer Structures* 2007;25(3):785-790.

13. Truskett VN, Watts MPC. Trends in imprint lithography for biological applications. *Trends in Biotechnology* 2006;24(7):312-317.
14. Koleske JV. *Radiation Curing of Coatings*. West Conshohocken, PA: ASTM International, 2002.
15. Trout TJ, Schmiege JJ, Gambogi WJ, Weber AM. Optical Photopolymers: Design and Applications. *Advanced Materials* 1998;10(15):1219-1224.
16. Decker C. Light-induced crosslinking polymerization. *Polymer International* 2002;51(11):1141-1150.
17. Sangermano M, Messori M, Galleco MM, Rizza G, Voit B. Scratch resistant tough nanocomposite epoxy coatings based on hyperbranched polyesters. *Polymer* 2009;50(24):5647-5652.
18. Jung T, Simmendinger P, Tobisch W. Out of the shadows. Plasma process facilitates UV curing in three dimensions. *European Coatings Journal* 2005;4:138-143.
19. Studer K, Decker C, Beck E, Schwalm R. Thermal and photochemical curing of isocyanate and acrylate functionalized oligomers. *Eur Polym J* 2005;41(1):157-167.
20. Maag K, Lenhard W, Loffles H. New UV curing systems for automotive applications. *Progress in Organic Coatings* 2000;40(1-4):93-97.
21. Sangermano M, Voit B, Sordo F, Eichhorn KJ, Rizza G. High refractive index transparent coatings obtained via UV/thermal dual-cure process. *Polymer* 2008;49(8):2018-2022.
22. Mills P. Robotic UV curing: a cost-effective way to cure large 3D plastic parts. 2007. p. 174-183.
23. Crivello JV. The discovery and development of onium salt cationic photoinitiators. *Journal of Polymer Science Part A: Polymer Chemistry* 1999;37(23):4241-4254.
24. Decker C, Moussa K. Kinetic study of the cationic photopolymerization of epoxy monomers. *J Polym Sci Pol Chem* 1990;28(12):3429-3443.
25. Sipani V, Scranton AB. Kinetic studies of cationic photopolymerizations of phenyl glycidyl ether: termination/trapping rate constants for iodonium photoinitiators. *J Photochem Photobiol A-Chem* 2003;159(2):189-195.
26. Sipani V, Scranton AB. Dark-cure studies of cationic photopolymerizations of epoxides: Characterization of the active center lifetime and kinetic rate constants. *J Polym Sci Pol Chem* 2003;41(13):2064-2072.
27. Ficek BA, Thiesen AM, Scranton AB. Cationic photopolymerizations of thick polymer systems: Active center lifetime and mobility. *European Polymer Journal* 2008;44(1):98-105.

28. Wicks ZW, Jones FN, Pappas SP. Organic Coatings: Science and Technology. New York: John Wiley & Sons, Inc., 1994.
29. Catilaz-Simonin L, Fouassier JP. Investigation of a system capable of photoinitiating radical polymerizations in thick pigmented media. *Journal of Applied Polymer Science* 2001;79(10):1911-1923.
30. Danu S, Darsono, Marsongko. UV-curing of titanium dioxide pigmented epoxy acrylate coating on ceramic tiles. *Journal of the Ceramic Society of Japan* 2008;116(1356):896-903.
31. Kenning NS, Ficek BA, Hoppe CC, Scranton AB. Spatial and temporal evolution of the photoinitiation rate for thick polymer systems illuminated by polychromatic light: selection of efficient photoinitiators for LED or mercury lamps. *Polym Int* 2008;57(10):1134-1140.
32. Kenning NS, Kriks D, El-Maazawi M, Scranton AB. Spatial and temporal evolution of the photoinitiation rate for thick polymer systems illuminated with polychromatic light. *Polymer International* 2006;55(9):994-1006.
33. Cai Y, Jessop JLP. Decreased oxygen inhibition in photopolymerized acrylate/epoxide hybrid polymer coatings as demonstrated by Raman Spectroscopy. *Polymer* 2006;47:6560-6566.
34. Nelson EW, Scranton AB. Kinetics of cationic photopolymerizations of divinyl ethers characterized using *in situ* Raman spectroscopy. *Journal of Polymer Science Part A: Polymer Chemistry* 1996;34(3):403-411.
35. Odian G. Principles of Polymerization. New York: John Wiley & Sons, 1991.
36. Terrones G, Pearlstein AJ. Effects of optical attenuation and consumption of a photobleaching initiator on local initiation rates in photopolymerizations. *Macromolecules* 2001;34(10):3195-3204.
37. Ivanov V, Decker C. Kinetic study of photoinitiated frontal polymerization. *Polym Int* 2001;50(1):113-118.
38. Asmussen S, Arenas G, Cook WD, Vallo C. Photoinitiation rate profiles during polymerization of a dimethacrylate-based resin photoinitiated with camphorquinone/amine. Influence of initiator photobleaching rate. *European Polymer Journal* 2009;45(2):515-522.
39. Baikerikar KK, Scranton AB. Photopolymerizable liquid encapsulants for microelectronic devices. *Polymer* 2001;42(2):431-441.
40. Baikerikar KK, Scranton AB. Photopolymerizable liquid encapsulants for microelectronic devices: Thermal and mechanical properties of systems with reduced in-mold cure times. *Journal of Applied Polymer Science* 2001;81:3449-3461.
41. Narayanan V, Scranton AB. Photopolymerization of composites. *Trends in Polymer Science* 1997;5:415-419.

42. Coons LS, Rangarajan B, Godshall D, Scranton AB. Photopolymerizations of vinyl ester: glass fiber composites. Photopolymerization, ACS Symposium Series 673 1997:203-218.
43. Miller GA, Gou L, Narayanan V, Scranton AB. Modeling of photobleaching for the photoinitiation of thick polymerization systems. J Polym Sci, Polym Chem Ed 2002;40(6):793-808.
44. Azan V, Lecamp L, Lebaudy P, Bunel C. Simulation of the photopolymerization gradient inside a pigmented coating - Influence of TiO₂ concentration on the gradient. Prog Org Coat 2007;58(1):70-75.
45. Jahn R, Jung T. Relationship between pigment properties and UV-curing efficiency. Prog Org Coat 2001;43(1-3):50-55.
46. Tesfamichael T, Hoel A, Wackelgard E, Niklasson GA, Gunde MK, Orel ZC. Optical characterization and modeling of black pigments used in thickness-sensitive solar-selective absorbing paints. Sol Energy 2000;69(1-6):35-43.
47. Cussler EL. Diffusion: Mass transfer in fluid systems. New York: Cambridge University Press, 1984.
48. Anseth KS, Wang CM, Bowman CN. Kinetic evidence of reaction diffusion during the polymerization of multi(meth)acrylate monomers. Macromolecules 1994;27(3):650-655.
49. Nelson EW, Jacobs JL, Scranton AB, Anseth KS, Bowman CN. Photo-differential scanning calorimetry studies of cationic polymerizations of divinyl ethers. Polymer 1995;36(24).
50. Tauber G, Hasenzahl S, Johnson RE. Evonik Industries. NIPex Pigment Blacks for Toner: Technical Information No. 1025., 2008.
51. Schuessler F, Feldmann K, Bigl T. Selection and qualification of polymers for rigid and flexible interconnect applications. Circuit World 2007;33(2):36-42.
52. Kamiya S, Furuta H, Omiya M. Adhesion energy of Cu/polyimide interface in flexible printed circuits. Surface and Coatings Technology 2007;202(4-7):1084-1088.
53. Lee TY, Roper TM, Jonsson ES, Kudryakov I, Viswanathan K, Nason C, Guymon CA, Hoyle CE. The kinetics of vinyl acrylate photopolymerization. Polymer 2003;44(10):2859-2865.
54. Dietz JE, Elliott BJ, Peppas NA. Real-Time Attenuated Total Reflectance-Fourier Transform Infrared Spectroscopy To Monitor Multiacrylate Polymerization Reactions. Macromolecules 1995;28(15):5163-5166.
55. Kim D, Scranton AB. The role of diphenyl iodonium salt (DPI) in three-component photoinitiator systems containing methylene blue (MB) and

- an electron donor. *Journal of Polymer Science Part A: Polymer Chemistry* 2004:42(23):5863-5871.
56. Padon KS, Scranton AB. A mechanistic investigation of the three-component radical photoinitiator system Eosin Y spirit soluble, N-methyldiethanolamine, and diphenyliodonium chloride. *Journal of Polymer Science Part A: Polymer Chemistry* 2001:39(5):715-723.
 57. Nie J, Andrzejewska E, Rabek JF, Lindén LÅ, Jean PF, Paczkowski J, Scigalski F, Wrzyszczyński A. Effect of peroxides and hydroperoxides on the camphorquinone-initiated photopolymerization. *Macromolecular Chemistry and Physics* 1999:200(7):1692-1701.
 58. Odian G. *Principles of Photopolymerization*, 2nd Edition. New York: John Wiley & Sons, 1991.
 59. Griffiths MC, Strauch J, Monteiro MJ, Gilbert RG. Measurement of Diffusion Coefficients of Oligomeric Penetrants in Rubbery Polymer Matrixes. *Macromolecules* 1998:31(22):7835-7844.