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Program Manager:

Dr. Charles Lee

AFOSR/NL 801 N. Randolph St., Ste 732 Arlington, VA 22203-1977 Phone: 703-696-7779 Fax: 703-696-8449 Email: **charles.lee@afosr.af.mil**

Prof. Madhu S. Madhukar

316 Perkins Hall The University of Tennessee Knoxville TN 37996-2030 Phone: 865-974-7676 Fax: 865-974-7663 Email: **mmadhuka@utk.edu**

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14. ABSTRACT

This work includes experimental and theoretical study of the effect of cure cycles on the development of cure-induced stresses in thermoset polymer composites. The experimental part includes modifying a newly developed test method to monitor the development of cure-induced stresses in thermoset polymer composites. Test results gave a clear understanding of the mechanisms of stresses development. The test method was further modified with a closed loop feedback control system to modify the cure cycles to reduce cure-induced stresses. Test results show that the modified cure cycles reduce residual stresses in composite laminates. The theoretical part included the development of a mechanical optimization model to study the effect of changing the cure cycle on the development of residual stresses in composite laminates. The model was used to generate cure cycles that reduce residual stresses in thermoset polymer composites. Findings of the theoretical modeling were found to agree well with those of the experimental study.

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Cure Cycle Optimization in Polymer Composites

Madhu S. Madhukar Mechanical and Aerospace Eng. The University of Tennessee, Knoxville, TN 37996-2030 (865) 974-7676 **mmadhuka**(a)utk.edu

ABSTRACT

This work concerns the study of the effect of cure cycles on the development of cure induced stresses in thermoset polymer composites. The study includes experimental testing as well as theoretical modeling. The experimental part included modifying a newly developed test method to monitor the development of cure induced stresses in thermoset polymer composites. Test results were shown to be highly reproducible and gave a clear understanding of the mechanisms of stresses development as well as cancellation. Good qualitative agreement between test results and the independently measured polymer volume change data was seen. The test method was further modified with a closed loop feedback control system to modify the cure cycles to reduce cure induced stresses. Several composite material systems were studied. Test results show that the modified cure cycles reduce residual stresses in composite laminates compared to the cure cycles recommended by materials manufacturers while maintaining the same glass transition temperature. It was shown that the modified cure cycles may also enhance the matrix-dominated mechanical properties, maintain better dimensional stability, and reduce cure time. The theoretical part of the study included the development of a mechanical optimization model and a computer program, OPTICURE, to study the effect of changing the cure cycle on the development of residual stresses in composite laminates. The model and the program were used to generate cure cycles that reduce residual stresses in thermoset polymer composites and hence enhance the dimensional stability and mechanical properties. Findings of the theoretical modeling were found to agree well with those of the experimental study.

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1-1 INTRODUCTION

The inhomogeneous structure of thermoset polymer composites causes cure shrinkage and thermal volume changes during processing to produce residual stresses. These stresses may be high enough in epoxy polymer composites to cause microdamages such as ply cracking and delamination which may affect their mechanical properties and durability (1-3). In addition, these stresses result in geometrical distortion of composite parts which increases as the part size increases. This goes against the current trend in aerospace and other industries to develop large integrated structures to reduce the expensive cost of assembling small parts. Part assembly and fit up are known as the major cost item of producing tight-tolerance parts as those used in stealth technology.

Residual stresses in polymers and polymer composites have been the subject of many previous studies. These stresses develop during cure process as well as during cooldown after cure is complete. Contradictory findings and assumptions have been found regarding the contribution of cure shrinkage to the residual stresses. Also, little work was done to modify cure cycles to reduce residual stresses

1-2 EXPERIMENTAL STUDIES

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Several experimental methods have been used to measure residual stresses on polymers and polymer composites. This includes double beam method [4-6], photoelasticity [7-10], confinement tubes with different geometry [11-18], fiber optics [19], piezoresistivity [20], peel-ply [21], unsymmetric laminates [21-23], single fiber stress test [24], and others [25]. Double beam method is simple and can be used to study the development of stresses in thin polymer films but not in composites or polymers under constraints. Using this method, Dannenberg studied development of residual stresses in Epon 828 and 1002 epoxies cured with different agents [4]. It was shown that residual stresses increase with curing temperature and depend on the curing agent. Also, it was shown that stresses start to develop significantly when temperature drops below glass transition temperatures. Using similar test setup, Wang and Yu studied residual stresses in diglycidyl ether of bisphenol A [5]. They used different cure cycles and showed that lowering cure temperatures may reduce stresses build up. However, longer cure time is needed at lower cure temperatures. Using a film coat on quartz beam, Feger and his coworkers studied the development of residual stresses in PMDA-ODA polyamic acide polyimide films [6]. They showed that the removal of solvent and changes in moisture content cause stresses to develop in the polyimide films. Results indicate that the polyimide films maintained about 25% of total stresses in the vicinity of glass transition temperatures and disappeared only after heating to higher temperatures.

Pawlak and Galeski [7] used three-dimensional photoelasticity to study residual stresses in polymer matrix with spherical inclusions. It was shown that residual stresses increase significantly as the space between inclusion decreases. Similar conclusions were shown by Theocaris and Paipetis [8]. Asamoah and Wood used photoelasticity to study residual stresses in fiber reinforced epoxies [9]. They showed that stresses in resin increases significantly as fiber spacing decreases. A tri-con of matrix material is created between fibers which is subjected to high level of constraining by the fibers. The authors used elastic finite element

analysis and confirmed their conclusions qualitatively. Cunningham and his coworkers studied residual stresses in resin materials between fibers in fiber reinforced composites inside a rigid outer shell [10]. The constrained resin was shown to undergo significant residual stresses.

Shimbo and his coworkers measured the effect of modifying epoxide resin with a spiro ortho-ester on residual stresses [11]. They measured stresses by measuring strains in a ring surrounded by the curing polymer. It was found that the modified resin has a lower glass transition temperature and hence less build up of residual stresses. In another paper, they modified the polymer to reduce the polymer stiffness while maintaining the glass transition temperature [12]. Reducing stiffness causes residual stresses to decrease. In both cases, reduction of stresses is accompanied by scarifying desirable properties. In a later study, Shimbo and his coworkers studied the mechanism of stresses development in epoxide resins [13]. It was shown that contribution of cure shrinkage to final residual stresses increases significantly when the cure is carried below glass transition temperature.

Lee and Schile tried to simulate development of stresses in fiber reinforced epoxy composites. They arranged glass rods inside a curing polymer and measured strains in the rods during cure and cooldown [14]. Also, they conducted an elastic finite element analysis. They studied different epoxy/hardener systems used in composites. The results show that the glass rods constraint the polymer volume changes. Residual stresses due to thermal shrinkage during cooldown assuming elastic analysis were smaller than measured values. The difference was attributed to cure shrinkage. Different systems of epoxy-hardener produced different relative values of stresses due to cure shrinkage to cooldown thermal shrinkage. Plepys and his coworkers studied the effect of degree of confinement of the polymer during cure on the development of residual stresses used in Epon 828 epoxy cured with different curing systems [15,16]. Test data show that when one-dimensional constraints exists, the contribution of cure shrinkage to the final residual stresses is much smaller than that of thermal shrinkage. When three-dimensional constraints were considered, stresses due to cure shrinkage were significantly high and of comparable values to that of thermal stresses during cooldown. Similar findings were shown by Korothov and his coworkers [17,18]. According to the principles of mechanics, the behavior of polymer approaches pure elastic behavior under three-dimensional confinement. In this case, stress relaxation of stresses due to cure shrinkage may be ignored and cure shrinkage, which is usually of large value, will contribute significantly to residual stresses. Interestingly, Plepys showed that for certain cure schedules, stresses due to cure volume changes (thermal expansion during heating and cure shrinkage) after the cure is complete may be in the opposite direction of stresses due to cure shrinkage and hence will reduce final residual stresses.

Lawrence and his coworkers used embedded fiber optics to measure strains in composites due to residual stresses [19]. It was shown that significant strains are developed during processing of cross ply laminates made of HyE 6049U carbon/epoxy composites. Results indicate that strains start to build up during cure due to chemical shrinkage. These strains may constitute about 25% of the total mechanical strains and hence residual stresses. Crasto and Kim tried to utilize carbon fiber piezoresistivity to measure residual

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stresses in composites [20]. It was shown that, as already established, that increasing cure temperature results in increasing fiber stresses as measured by fiber piezoresistivity.

In another study, Crasto and Kim measured stress free temperature in AS4/3501-6 graphite/epoxy as an indication on residual stresses [21]. A definition for the stress free temperature is the temperature at which a freestanding unsymmetric laminate will be flat. The higher the residual stresses, the higher the stress free temperature will be. The unsymmetric laminates were cured at different temperatures and produced different glass transition and stress free temperatures. Test data showed that stress free temperature is higher than both curing temperature and glass transition temperature. The difference between stress free temperature and cure temperature, about 15°C, was attributed to cure shrinkage stresses. Stress free temperature was higher than glass transition temperature with about 8°C. It was expected that above glass transition temperatures. However, when the temperature was raised above stress free temperature the direction of the laminate curvature was reversed indicating that the composite maintained high stiffness. This can be explained by the fact the behavior of the polymer in composites in the existence of fiber confinement may be different than that of neat polymers.

Some case studies have been conducted to investigate the effect of cure cycle modification on the cure induced residual stresses [22,23]. In these studies, the cure cycles were modified arbitrarily to see the effect on residual stresses. In one study [22], the two dwell time temperature standard cycle for IM6/3100 graphite/epoxy composite was modified by introducing a third dwell temperature. Different locations and duration of the third dwell temperature were investigated (see Figure 1.1). One of the modified cycles reduced the stresses by up to 25% while maintaining the mechanical properties as that of the specimens cured with the standard cure cycle (cure cycle recommended by the prepreg manufacturer). There were no criteria for choosing the location and duration of the additional dwell temperature. In another study [23], the two dwell cure cycle of 976 epoxy was modified by extending the first dwell temperature. As can be seen from Figure 1.2 such extended cycles can reduce spring back of unsymmetric laminates by more than 25%. While these studies showed that modifying cure cycles can reduce residual stresses in composites, they are not effective because of the trial and error methods involved. These experimental methods to modify the cure cycles to reduce the residual stresses demonstrated the validity of such approach.

A new test method was developed by Madhukar and his coworkers which is based on curing the polymer around a pretensioned fiber [24]. After the polymer develops significant stiffness, volumetric changes in the polymer will change fiber tension which is a measure of the residual stresses. They demonstrated that changing the cure cycle of Epon 828 cured with mPDA will change the contribution of cure volume changes to the residual stresses in the fibers.

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1-3 THEORETICAL STUDIES

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In earlier studies, stresses developed due to thermal shrinkage during cooldown were evaluated using linear elastic analysis [26]. Stresses due to cure shrinkage were neglected assuming that it will be relaxed during cure. In another study, the effect of swelling due to moisture absorption was shown to reduce residual stresses [27]. The author indicated that residual stresses are higher than 50% of the transverse strength. Based on the elastic analysis, he concluded that stress-free temperature is lower than the curing temperature. Excluding the effect of moisture effect which depend on environmental conditions considered, this conclusion can not be generalized in the view of experimental data discussed above. In a later study by the author, effects of residual stresses on ply cracking and warping in unsymmetric laminates were discussed [1]. It was shown experimentally that stress free temperature is higher than curing temperature and the cooldown rate apparently has no effect on residual stresses. Also, data shows that changing gelation temperature may reduce residual stresses.

In a later study, White and Hahn studied the development of residual stresses through the whole cure cycle [28]. It was shown that transversal stiffness and strength, which are matrix dependent, develop only after polymer degree of cure reached 0.83 in IM6/CYCOM 3100 graphite/BMI composites. It was shown that postcure causes increase in the residual stresses which was attributed to shrinkage due to additional cure and loss of moisture or volatiles.

Other models were developed to estimate stresses development during cure cycle [29-35]. Modeling of the mechanical properties as a function of degree of cure and temperature have been the most difficult because of the viscoelasticity involved. Assumptions involved in modeling the mechanical properties development in the composite directly affect the validity of the predicted stresses. A model to evaluate residual stresses in polymer composites predicted that chemical shrinkage contributes less than 5% to final residual stresses in IM6/3100 graphite/bismaleimide composites during cure [29,30]. The model included viscoelastic effects to account for stress relaxation. It was assumed that longitudinal stiffness increases about 50% when the cure is complete while transverse modulus increases significantly only after certain degree of cure. However, test data as shown in Figure 1.3, indicates that the model underestimates the contribution of chemical shrinkage [22]. This can be recognized considering that for thermosetting polymers, the cure-induced stresses by chemical shrinkage in some cases may exceed 30% of final stresses [15-19, 25]. This is supported by the fact that changing the cure cycle for some composite systems (with the same cure temperature and cooldown rate) affects the final residual stresses significantly [22,23]. This is mainly attributed to the change in the contribution of chemical shrinkage to the final stresses.

A model was developed to evaluate cure stresses in thick composites [31-35]. Stress relaxation was not considered and resin modulus was evaluated using a mixing rule depending on polymer degree of cure. As shown above, a threshold value of degree of cure must exist before some of the mechanical properties are developed significantly [28]. Mechanical properties of the composite were estimated from properties of matrix and fibers using micromechanics neglecting any interaction. As discussed above, behavior of resin

may be significantly different than the behavior of resin with high fiber volume fractions as in composites. Full cure shrinkage was used to estimate stresses. Experimental results indicate that only part of cure shrinkage may contribute to residual stresses [4-18,28]. As expected high values of cure stresses were estimated. No experimental verification was presented to support the model.

Weitsman developed a rigorous viscoelastic optimization model to maximize stress relaxation during cooldown for a given cooling down time [36,37]. The model predicts 10% reduction in residual stresses in epoxy-resin composites if optimum cooldown path is applied although this has not yet been supported by experimental data. Harper and Weitsman conducted an experimental and theoretical investigation on the effect of cure shrinkage, moisture, and cooldown path on residual stresses in polymer composites during cooldown [38-40]. The presence of moisture was shown to enhance viscoelastic behavior. Predictions of stresses due to moisture change were in reasonable agreement with experimental data. Based on their optimal cooldown path for AS4/3502 graphite/epoxy, a 10-15% reduction in residual stresses may be obtained, however, it was not seen experimentally. Using double beam method, Harper showed that cure shrinkage may contribute up to 25% of residual stresses in 3502 epoxy [41].

Several models were developed to evaluate residual stresses in thermoset resin [42-50]. Levitsky and Shaffer developed an elastic model to evaluate stresses in a solid sphere cast from a thermoset polymer during curing [42-43]. The formulation is rigorous, however results are not directly applicable to composites. Lange and his coworkers studied stresses build-up in thermoset films cured below their ultimate glass transition temperatures [44,45]. They used a Maxwell model to simplify viscoelastic analysis. Double beam method was used to verify analysis predictions. It was shown that contribution of cure volume changes to final residual stresses depends on amount of resin shrinkage beyond gelation and density of crosslinking. They indicated that stresses from cure shrinkage vary from 1 to 30 % of the final residual stresses depending on the polymer under consideration and on cure cycle. Mallick and Krajcinovic developed a micromechanical model to study stress and strain fields in resin slab [46]. They demonstrated the use of aggregation-percolation models to obtain the constitutive equations of the curing resin. Stresses were shown to develop significantly only after gelation. A similar approach was considered earlier by Adolf and Martin [47-49]. Adolf suggested the use of time-cure superposition to account for nonisothermal cure viscoelasticity. They indicated that changing cure cycle for a given polymer may affect significantly the contribution of cure shrinkage to final residual stresses. Monk and his coworkers developed a viscoelastic model for residual stresses in spun polyimide films [50]. They used fundamentals of chemical reactions and macromolecules to predict macro-properties such as strains. Shrinkage due to evaporation was included in the analysis. Given the formulation approach, many parameters were difficult to measure and/or verify. These parameters were estimated by fitting model predictions with test results obtained using double beam method.

Finite element analysis (FEA) was used to study development of residual stresses in composites [51-56]. Jain and Mai used linear FEA to study development of residual stresses in T300/934 carbon/epoxy composite cylinders including the effect of cure shrinkage (which was taken arbitrarily as 1.5%) [51,52].

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Measured spring-in values for channel sections were found to change with tooling materials. The predicted results based on the assumed cure shrinkage value appear to agree with measured values. Sala and Di Landro used linear FEA and classical laminate theory to study residual stresses in T800/5250-2 carbon/BMI composites [53]. In their formulation, they considered stress build-up form stress free temperature. In that study, stress free temperature was measured experimentally and was found to exceed cure temperature by about 5365°C. For a given measured stress free temperature, the elastic solution was shown to overestimate curvature of unsymmetric laminates. In a later study [54], the authors reported that based on the comparison between analytical and experimental results, the transverse modulus of the composite was found to change with time and temperature as an indication of the presence of viscoelasticity.

Wang and Daniel developed a linear viscoelastic model to study residual stresses and warpage in wovenglass/epoxy laminates [55]. They considered thermorheologically simple model and time-temperature superposition in their formulation. Chemical shrinkage in addition to thermal expansions were considered in calculating stresses. Because of the woven nature of the fabric both transverse modulus and longitudinal modulus were found to exhibit time-temperature dependency. The viscoelastic analysis was found to yield satisfactory results when qualitatively compared to experimental data.

Mohan and Grentzer used linear FEA to simulate processing and predict residual stresses development in glass/vinyl ester composites [56]. They applied the method to study stresses development during cure of a composite nozzle. The authors indicated that the contribution of cure shrinkage is significant although it needs more investigation to be quantified.

1-4 PROBLEM STATEMENT

From the above discussion, it can be seen that a substantial amount of research work was done to investigate the development of residual stresses in thermoset polymers and polymer composites. However, contradictory assumptions and conclusions still exist regarding the effect of major parameters on the development of stresses. For example, the extent of cure shrinkage contribution to residual stresses and effect of stress relaxation are not clear yet. In addition, very little work was done to reduce these stresses without scarifying desirable properties or considerable elongating the cure cycle. This necessitates a fundamental understanding for the process of cure stresses development. This fundamental understanding can then be used to study the possibility of reducing residual stresses.

The approach used here is mainly experimental using single fiber cure induced stress test (section 1.2) [24]. This new method was developed to monitor stresses developed in fibers during cure of a single fiber composite. The method was shown to be both simple and effective to study mechanisms of stresses development. In the current study, this method was used to investigate stresses development in single fiber thermoset polymer composites. Different material systems were considered to obtain general understanding. Then the method was modified as necessary to optimize cure cycles to reduce residual stresses.

2. EXPERIMENTAL METHODS

2-1 INTRODUCTION

A test procedure to study the development of cure induced stresses in thermoset polymer composites was developed at the Composite Materials Laboratory at the University of Tennessee as discussed in Chapter 1 [24]. The method was applied to a simple epoxy system (Epon 828/mPDA system). The test proved to be a simple and effective method to study the development of cure induced stresses. Preliminary results showed how polymer volume changes during cure affect fiber stresses. Also, it was shown that changing the cure cycle changes the stresses developed during cure. However, the early version of the test setup was basic and not suitable to run tests on more complicated polymer systems requiring higher cure temperature and more uniform heating. In addition, test components and procedure were not yet well developed to eliminate the effects of parameters other than polymer volume changes. This includes friction between fiber and test components, degassing technique that takes long time that may advance cure before resin is poured around fiber, shape of the mold which result in constraining the specimen during shrinkage, and nonuniform temperature along the specimen. For example, results of the early version of the test setup for AS4/3501-6 graphite/epoxy composite were not always consistent.

In the current research work, this test method was modified to resolve the above problems. The modified test setup and the testing procedure produced consistent results for several materials systems. The modified setup and procedure were used to study the effect of the cure cycles on the cure induced stresses. A closed loop feedback control system was also developed to cure cycles that reduce stresses. In this chapter, the modified test method and typical results are shown and discussed. Feedback control system and its applications are also described.

2-2 CURE INDUCED STRESS TEST (CIST)

Figure 2.1 shows the conceptual mechanism assumed for the development of fiber stresses in CIST due to polymer volume changes. The polymer is cured around a portion of a pretensioned fiber. Once the polymer develops sufficient stiffness, thermal expansion in the polymer (as a result of heating) will result in decreasing fiber tension. On the other hand, shrinkage in the polymer (as a result of crosslinking or cooling down) will result in increasing fiber tension. The change in the fiber tension will decrease with time because of stress relaxation whenever the polymer exhibits viscoelastic behavior. Correspondingly, it can be seen that polymer shrinkage will result in compressive stresses in the fiber portion embedded inside the polymer whereas the expansion will result in tensile stresses.

All volume change data presented here for comparison were obtained at Air Force Research Laboratory. The volumetric dilatometer used was GNOMIX, Inc. PVT Apparatus (Figure 2.2). More details about the test setup and procedure can be found in reference [57]. Figure 2.3 shows the volume change data of 3501-6 epoxy during a two dwell cure cycle recommended by resin manufacturer (standard cure cycle). There is expansion during heating from ambient temperature to the first dwell temperature and during heating from

the first dwell temperature to the second dwell temperature. On the other hand, there is shrinkage due to crosslinking during the temperature holds and also upon cool down after cure is complete. Hence, there are two types of volume changes occurring during the cure of thermoset composites; first type is the expansion during heating and the other is the shrinkage due to the chemical reaction (crosslinking shrinkage) and during cooling down (thermal shrinkage). Volume changes prior to polymer gelation are not expected to contribute to residual stresses since the melted polymer acts like a fluid with low viscosity and stress relaxation will impede stresses build up. Gelation is defined by the point at which infinite network of molecules is developed inside the polymer. Beyond gelation, volume changes produce proportional stresses in the fibers and stresses may start to build up.

Through the rest of this report, the term *effective* volume will be used to refer to the volume changes that produce stresses in the fibers. *Effective* expansion occurs during any heating beyond the polymer gelation, and *effective* shrinkage results from polymer crosslinking beyond gelation and during cool down. *Effective* expansion will produce tensile stresses in the fiber portion embedded inside the polymer, whereas *effective* shrinkage will produce compressive stresses. This can easily be seen from the test mechanism shown in Figure 2.1. In typical cure cycles, gelation temperature is close to maximum cure temperature and hence *effective* expansion (which corresponds to the increase in the temperature from gelation to the maximum cure temperature) is typically small, if any.

2-2-1 Test Setup

Figure 2.4 shows a schematic diagram of CIST. Figure 2.5 shows a photograph of the test setup. The fiber is fixed to a rigid support at one end, and the other end is passed through a cavity in a silicone mold and then glued to a load cell (250 gm load capacity and 0.01 gm sensitivity). A ceramic shield is used to form a cure chamber around the mold. The shield has two narrow slots, one at each end to pass the fiber. The sizes of these slots were kept to a minimum to reduce the heat loss at the ends and to avoid significant variation in the temperature inside the cure chamber. The dimensions of the cure chamber are approximately 160 mm long, 70 mm wide, and 76 mm high. A ceramic strip heater (from Research Inc., Model 4184) is placed at the top to apply the time-temperature cycle. A thermocouple is located beside the middle of the silicon mold to monitor temperature. The thermocouple is connected to a data acquisition system. A computer program was found to produce a heating profile within $\pm 0.25^{\circ}$ C of a given input time-temperature cycle. The dimensions of the silicon mold are 76 mm long, 3.5 mm wide, and 2 mm high. The thickness of the base of the silicon mold was kept as small as possible (about 0.3 mm) to minimize the effect of silicon volume changes and constraints applied by mold on the polymer on test results.

After positioning the fiber, it is glued at both ends. The glue is allowed to cure according to the recommendations of its manufacturer. Then, the fiber is pretensioned to about 5 gm. The load cell reading should be stable for at least 5 minutes before pouring the polymer around the fiber to ensure proper fiber fixation and the cure of the glue. About 1 gm of the resin is melted and degassed in a high temperature

syringe using a vacuum oven. The degassing temperature is chosen to be the minimum temperature that is high enough to melt the resin and enable degassing while not causing significant cure in the polymer. After degassing, the resin is injected inside the silicon mold around the fiber and the time-temperature cycle is started.

CIST results will be shown as the fiber tension data obtained from the load cell reading. A comparison between fiber tension data and stresses in fiber portion embedded in the resin shows that any increase in the fiber tension more than its initial value (as a result of the resin shrinkage) corresponds to the application of compressive stresses on the fiber portion embedded inside the polymer. On the other hand, any drop in the fiber tension below its initial value (as a result of the resin expansion) corresponds to the application of tensile stresses on the fiber portion embedded inside the polymer. Under no external loads, the matrix in the vicinity of the fiber experiences tensile or compressive stresses when the fiber experience compressive or tensile stresses, respectively.

2-2-2 Typical Results

The material used in this section is AS4/3501-6 graphite/epoxy composite system. The standard cure cycle of 3501-6 consists of: heating the sample from room temperature to 116°C in 30 minutes; holding the temperature at 116°C for 60 minutes; raising the temperature from 116°C to 177°C in 25 minutes; holding the temperature at 177°C for 240 minutes; followed by cool down to room temperature.

Figure 2.6 shows the fiber tension data during such a cycle. The polymer volume change data for this cure cycle is shown in Figure 2.3. A good correlation between fiber tension and volume change can be seen. During the first part of the cure cycle fiber tension remains constant because the viscosity of the resin is small. As the resin viscosity increases, the expansion during the heating ramp causes the fiber tension to decrease and the crosslinking shrinkage causes the fiber tension. Finally, thermal shrinkage during shrinkage during the second temperature dwell increases the fiber tension. Finally, thermal shrinkage during cool down causes increase in the fiber tension. The drop in the fiber tension during the heating ramp corresponds to the development of tensile stresses in the fiber portion embedded in the polymer. The fiber tension at the end of the cure, just before cool down starts, is higher than the initial fiber tension indicating the development of compressive stresses in the fiber portion embedded in the polymer. These stresses are undesirable because they will be added to compressive stresses.

In the CIST, the composite system is essentially a single fiber in an infinite polymer. To simulate a more representative fiber volume fraction as in composite laminates, an experiment was performed with fiber coated with a thin layer of resin. The coated fiber was subjected to the standard cure cycle and changes in the fiber tension were measured (Figure 2.7). The shape of the change of fiber tension is the same as obtained with the fiber running inside an infinite polymer (Figure 2.6) but with smaller values. This

indicated that the mechanisms that are identified in the single model composites are present in composite laminates as well.

2-2-3 Parameters of CIST

To study the reproducibility of the test results and determine the effect of different parameters including initial fiber tension and fiber embedded length, several CIST experiments were conducted on AS4/EPON 828 graphite/epoxy single fiber composites. This composite system was chosen because it is easy to work with and the duration of the cure cycle is relatively short. The EPON 828 is a diglycidyl ether of bisphenol-A and was cured with 14.5 parts per hundred by weight of meta-phenylene diamine (mPDA). The matrix was mixed, degassed under vacuum for 10 minutes, and cured with different cycles. The typical cure cycle used for this resin consists of: heating from ambient temperature to 75°C in 5 minutes then the temperature is held at 75°C for 2 hrs; then it is raised to 125°C and held at 125°C for two hours followed by cool down to ambient temperature.

Figure 2.8 shows typical CIST results for Epon 828/mPDA epoxy. The fiber tension is unchanged for the first 1.5 hours at 75°C. Some increase in the fiber tension occurs during the last 0.5 hour at 75°C. The fiber tension drops rapidly when the temperature is increased from the first dwell at 75°C to the second dwell at 125°C. Then, again there is an increase in the fiber tension during the first half-hour at 125°C temperature hold. During cooling to ambient temperature, the fiber tension again increases and stabilizes at the completion of cooling.

In Figure 2.8, the rapid heating from room temperature to the first dwell temperature just melts the resin without allowing significant resin cure to occur. Thus, it is expected that the resin just flow around the fiber without causing fiber stresses. During the temperature hold at 75°C, the viscosity of the resin increases with the advancement of resin cure. At that stage, volume changes start to be *effective* and the cure shrinkage starts to increase fiber tension. During the temperature ramp between first and second dwell temperatures both thermal expansion and crosslinking shrinkage exists. However, it appears that the rapid heating rate causes thermal expansion to dominate producing net expansion. This expansion causes fiber tension to drop. The chemical shrinkage during second temperature hold causes fiber tension to increase again. Finally, thermal shrinkage during cool down increases the tension.

The changes in the fiber tension shown in Figure 2.8 are primarily due to polymer volume changes because the effect of fiber expansion/shrinkage is comparatively small. This can be seen from CIST results shown in Figure 2.9. In the set of tests, the silicon mold cavity was not filled with resin and only the fiber was running through. The cure cycle was applied and the changes in fiber tension were recorded. The test was repeated with different types of fibers. Results show that fiber tension does not change significantly during the cure cycle because of fiber volume changes.

Figure 2.10 shows results of CIST under different values of initial fiber tension. Results indicate that the initial fiber tension has no effect on the changes in the fiber tension during the cure cycle. Figure 2.11

shows the effect of embedded fiber length on the variation in the fiber tension. The tension in the free fiber length is proportional to the deformation at the end embedded in the polymer. This deformation increases as the embedded fiber length increases. Thus the fiber tension in the free length increases as the embedded length increases. This can be seen from CIST results in Figure 2.11. In all cases, test results were highly reproducible.

To further understand the effect of polymer volume change and stress relaxation on the fiber stress, the cure cycle was modified as shown in Figure 2.12. A heating ramp from 65°C to 149°C was divided into 3 equal steps of 16.7°C and one last step of 33.3°C. Similar to the case of typical two dwell cycle, there is no change in the fiber tension during first heating ramp from ambient temperature to the first dwell temperature. The fiber tension remains unchanged until the end of the second temperature ramp suggesting that the polymer has low viscosity and/or insignificant crosslinking shrinkage. The fiber tension starts to increase rapidly during the second temperature hold indicating that the resin now has significant stiffness. The fiber tension drops during the third temperature ramp due to thermal expansion. Beyond this stage, the fiber tension increases during the temperature holds and drops during the following temperature ramps.

A comparison among the change in the fiber tension during different temperature steps in Figure 2.12 shows that after the resin has sufficient stiffness, the rate of the fiber tension increase decays with advancement in the cure cycle. This is expected since the rate of the chemical reaction and hence the crosslinking shrinkage decreases after it reaches certain extent. Also, the drop in the fiber tension due to thermal expansion during heating increases with the increase in the polymer cure. This is expected since as the degree of cure increases, the resin attains higher stiffness. The drop in the fiber tension during the last temperature ramp (with 33.3°C temperature increment) is almost twice the drop in the ramp just before it (with 16.7°C increment). This indicates that polymer stiffness reaches somewhat constant value beyond certain extent of cure. Results in Figure 2.12 suggest that the thermal expansion when developed towards the end of the cure cycle will produce significant fiber stresses.

Average stresses in the embedded part of the fiber during a cure cycle were calculated from simple mechanics of materials. Assuming a uniform fiber stress distribution along the embedded fiber length and perfect bond between the fiber and the matrix, the fiber stresses in the free part of the fiber, $\sigma'_f(t)$ (Figure 2.13), at time t, can be related to the fiber tension as

$$\sigma_f'(t) = \frac{T(t) - T(0)}{A_f} \tag{1}$$

where T(t) is the fiber tension at any time t, T(0) is the initial fiber tension, and A_f is the cross sectional area of the fiber. The corresponding strain $(\varepsilon'_f(t))$ can be written as

$$\varepsilon_f'(t) = \frac{T(t) - T(0)}{A_f E_f} \tag{2}$$

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where E_f is the Young's modulus of the fiber. The total deformation in the free fiber length $(\Delta'(t))$ is given by:

$$\Delta'(t) = \frac{T(t) - T(0)}{A_f E_f} 2L_l$$
(3)

where L_1 is as shown in Figure 2.13. Since the fiber length is fixed during the test, the deformation in the embedded part equals that of the free part with a negative sign. Thus, the strain in the embedded length $(\varepsilon_i(t))$ is given as

$$\varepsilon_f(t) = -\frac{T(t) - T(0)}{A_f E_f} \frac{2L_l}{L_2}$$
(4)

where L_2 is the embedded fiber length as shown in Figure 2.13. Thus, stresses in the embedded fiber length can be written as

$$\sigma_{f}(t) = -2\frac{T(t) - T(0)}{A_{f}} \left(\frac{L_{1}}{L_{2}}\right)$$
(5)

The effect of the change in the fiber length due to temperature change was neglected in Equation 5 since the thermal expansion coefficient of fibers is generally very small relative to that of the resin. The fiber stresses calculated from Equation 5 for a single fiber graphite/epoxy composite are plotted in Figure 2.14. The fiber stresses shown in this figure are due to matrix volume changes only, i.e., the fiber stresses resulting from the initial fiber tension T(0) are not included. An important observation in Figure 2.14 is that when the temperature is raised from the first dwell to the second dwell, the embedded fiber experiences a tensile stress peak (~1.2 GPa). To verify the results on cure induced fiber stress obtained from Equation 5, independent tests were conducted with different values of T(0). The average tensile strength of AS4 fibers ranged from 2.5 to 3.0 GPa. Thus, if a fiber is prestressed to about 1.4 GPa, the cure induced tensile stress (1.2 GPa) superimposed on the fiber prestress value will cause the fiber stress to approach its failure stress. Tests showed that when CIST conducted with the fiber prestressed to \sim 1.4 GPa, a fiber crack was detected as shown in Figure 2.15. When the fiber was prestressed to less than 1.4 GPa, no fiber crack was detected. These observations suggest that, although the analysis used to calculate the fiber stress during the curing is based on some gross assumptions (such as uniform fiber stress along the entire embedded length), it still agree quite well with the experimental observations.

To verify that the fiber fracture occurs within a short time interval when the temperature is raised from 75°C to 125°C, two additional experiments were conducted. In both of these specimens, the fibers were pretensioned to about 70% of their expected failure load. Such pretension loads are expected to produce fiber failure during the standard cure cycle. In one of the specimens the test was terminated after the first temperature hold, i.e. before starting the temperature ramp. The specimen was removed and immediately examined under an optical microscope. No fiber crack was detected in this specimen. The other specimen was removed at the end of the heating ramp and was immediately examined under the microscope. This specimen did show a fiber crack as that shown in Figure 2.15. This suggests that the fiber failure occurred during the heating ramp from 75°C to 125°C.

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For a given upper cure temperature limit, there may be many cure cycles which complete the cure. The resulting residual stresses will not be the same since different cycles mean different amounts of effective volume changes and different extents of stress relaxation. The following example shows how changing the cure cycle can change cure induced stresses. Figure 2.16 shows the change in the fiber stresses during a modified cure cycle. The heating rate during the heating ramp from the first to the second dwell temperatures was reduced arbitrarily to allow for more balance between chemical shrinkage and the thermal expansion. Also, the slow heating rate allows for more stress relaxation. Moreover, the gradual heating applies more thermal expansion towards the end of the cure cycle which causes thermal expansion to be effective. The modified cycle results in more uniform profile for the fiber stress. The maximum value of tensile stress was lower than that in the typical two dwell cure cycle. To further verify the results of Figure 2.16, another CIST was conducted with the fiber prestressed to 1.8 GPa. No fiber cracks were detected when cure cycle shown in Figure 2.16 was used. The final value of tensile stresses in the fiber at the end of the cure, just before cool down, was higher than that in the standard cure cycle. As discussed earlier, tensile stresses that develop during cure counteract part of the stresses developed due to the thermal shrinkage and thus reduce the final residual stresses. In this case, the new cycle is expected to produce less residual stresses after cool down is completed. This can be seen by comparing final value of fiber stresses at the end of each cycle ($\Delta \sigma$ in Figures 2.14 and 2.16).

2-3 MODIFICATION OF CURE CYCLES USING TRIAL AND ERROR METHOD

To study the effect of changing the cure cycle on fiber tension profile, several tests were performed on AS4/3501-6 graphite/epoxy. The objective was to find a modified cure cycle that minimizes the stress development in the fiber during cure. Such a cure cycle produces ultimately a flat fiber tension curve. The parameters are, initial heating temperature and the subsequent heating rate/s. Maximum temperature was set as that of the standard cure cycle. Trial and error approach was used to explore the existence of such a cycle. Figure 2.17 shows corresponding fiber tension profile during one of the intermediate cycles. It is clear that changing the cure cycle affect the variation in the fiber tension. The trial and error was continued until the cycle shown in Figure 2.18 was obtained. The obtained cure cycle almost completely eliminates the variation in the fiber tension during cure. The cure cycle was terminated when the temperature reached the highest cure temperature hold and the fiber tension remained constant for 30 minutes during temperature hold. The new cure cycle produces an almost flat fiber tension curve and it is shorter than the standard cure cycle.

Figure 2.19 shows volume change data during such a cycle. There is a good agreement between polymer volume change and fiber tension. There is no change in the fiber tension during the initial part of the cure cycle where the resin viscosity is low. As the polymer viscosity increases, the thermal expansion combined with stress relaxation act to cancel stresses due to crosslinking shrinkage and thus fiber tension remains constant. To check the completeness of resin cure, the glass transition temperature of the specimens cured using the standard and the modified cure cycles were measured. Results indicate that the glass transition

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temperature is almost the same, Table 2.1. Also, the T_g remained unchanged for specimens cured for 30 and 60 minutes longer at the highest temperature in the modified cycle. This indicates that the modified cure cycle satisfies the cure requirement while keeping cure induced stresses to a minimum.

Cure Cycle	Standard	Modified	Modified Cycle +	Modified Cycle +
	Cycle	Cycle	30 min.	60 min.
Glass Transition Temperature	170°C	171°C	172°C	171°C

Table 2.1 Glass transition temperature for 3501-6 epoxy specimens cured using different cure cycles.

In the above modified cycle, it was intended to cancel stresses developed during cure cycle. Another approach is to develop tensile stresses inside the composite during cure and maintain these stresses until the cool down. These tensile stresses will counteract part of the compressive stresses resulting from the thermal shrinkage. To develop tensile stresses during cure, the effective thermal expansion should be increased. This can be done by developing more thermal expansion after the polymer attains high viscosity. This implies that cure should be carried at relatively low temperature for long time before heating to a higher temperature to complete the cure. Thermal expansion that occurs during this part of the cure cycle will produce larger tensile stresses than if the heating was applied early in the cure cycle. Figure 2.20 shows typical results of 3501-6 epoxy when this approach is considered. This cure cycle was chosen arbitrarily with the first dwell extended for three hours (versus 1 hr in the standard cure cycle). The drop in the fiber tension during the heating ramp between first and second dwell temperatures in this cycle is larger than that in the standard cycle (Figures 2.6 and 2.20). As a result of this large drop, the fiber tension at the end of the cure cycle, prior to cooling down, was still smaller than the initial fiber tension (ΔF in Figure 2.20). This decrease in the fiber tension corresponds to the development of net tensile stresses in the fiber running inside the polymer. These tensile stresses will offset part of the compressive stresses produced during cool down and hence reduce the final residual stresses. It should be noted that this type of cycle is longer than the standard cycles. The parameters in this approach include, first dwell temperature, number of dwells, duration of these dwells, and heating rates between dwells. The proper choice of these parameters can result in cycles with lower final residual stresses but the cure cycle will be longer.

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2-4 MODIFICATION OF CURE CYCLES USING CLOSED LOOP FEEDBACK CONTROL

SYSTEM (CLFS)

Effects of residual stresses generated during the cure are reflected in the mechanical properties and dimensional stability of the composite parts. Different cure cycles result in different patterns of volume changes in the polymers during cure. Thus, changing cure cycles can affect residual stress development in composites. In section 2-3, CIST was used to determine modified cure cycles which reduced cure induced stresses in single fiber composites. The approach was to use trial and error to obtain such a cycle. Although the trial and error approach answered the question regarding the existence of such cycles, it was not practical to use with different composites systems since many trials may be involved. This calls for the development of a feedback control system. The idea is to reschedule the volume changes in the polymer and utilize the stress relaxation to minimize the variation and the values of the cure induced stresses.

As mentioned before, volume expansion and shrinkage occur in the polymer during different periods of the cure process. Before the polymer gelation, stress relaxation is very effective and fiber stresses are not likely to build up. As the polymer approaches gelation, the increase in the viscosity causes relaxation of stresses to take longer times and stresses may start to build up due to cure shrinkage. These stresses can be reduced by causing thermal expansion in the polymer. In the current study, the CLFS was developed to determine modified cure cycles that reduce cure shrinkage stresses for various composite systems. In this feedback system, the unrelaxed part of stresses developed due to polymerization shrinkage is simultaneously canceled by proportional thermal expansion.

Since *effective* expansion produces tensile stresses, it will counteract a fraction of the compressive stresses due to shrinkage, and thus it reduces final residual stresses. Therefore, it is desirable to increase the *effective* thermal expansion. *Effective* thermal expansion is not a constant for a given resin. It depends mainly on gelation temperature. The objective of this research work is to develop a simple procedure to reschedule cure cycle to produce maximum cancellation of stresses due to *effective* shrinkage with *effective* expansion and stress relaxation.

2-4-1 Optimization Procedure

A computer program was developed to automatically adjust the heating rate during cure to minimize the variation in the fiber tension. The method is based on searching for a heating path that results in a maximum cancellation of the unrelaxed stresses. Thus, when the CLFS detects an increase in fiber tension (as a result of crosslinking shrinkage in the resin), it will increase the temperature such that the resulting thermal expansion is enough to cancel the stresses produced by the chemical shrinkage. On the other hand, a decrease in fiber tension implies excessive thermal expansion in the resin due to rapid heating. In such cases, the program will start to decrease heating rate proportionally. The check for the change in the fiber tension is done at certain time intervals that are long enough to allow for some stress relaxation. The test is

considered complete when the fiber tension does not change for a certain period during a temperature hold at the maximum cure temperature. This period (typically from 20 to 40 minutes) varies according to the resin type. A block diagram of the CLFS is shown in Figure 2.21.

2-4-2 Optimization Parameters

The time interval and the heat increment applied by CLFS determine the heating rate at a particular moment in the cure cycle. Time interval is chosen by the user. The longer the time interval, the more stress relaxation is allowed (for same temperature increment). However, longer time interval means longer cure cycle. For most polymers at high temperature, stress relaxation is inevitable but its extent may vary depending on time and temperature. Stress relaxation can be utilized together with *effective* thermal expansion to minimize stresses due to chemical shrinkage. In practical cure cycles, most of thermal expansion is consumed during initial heating and hence there is a small amount of *effective* thermal expansion. In such cases, stress relaxation should be allowed as much as possible (without significantly prolonging the cure cycle) together with increasing thermal expansion to balance shrinkage stresses. The time interval should not be shorter than that required to allow the effect of the heat increment on matrix thermal expansion to occur completely. The heat increment is proportional to the change in the fiber tension from a reference value, which is taken as the initial fiber tension.

The proportionality constant that determines the extent of heat increment corresponding to a certain change in the fiber tension can be determined approximately by running the CIST with a cure cycle in which temperature is increased in steps. In this study, the initial value of this constant was taken as the ratio of the temperature increase during the ramp between the first and second dwells and the corresponding drop in the fiber tension of the standard cure cycle ($\Delta T/\Delta F$ in Figure 2.6). The proportionality constant was found to change with the degree of cure. The larger the degree of cure, the larger the value of the proportionality "constant". This may be attributed to the decrease in the contribution of stress relaxation at higher degree of cure to cancel the change in stresses as the degree of cure increases. In the CLFS, only approximate initial value is required, then the program uses continuously measured fiber tension and temperature change data to adjust this ratio to accommodate for the possible effects of degree of cure and temperature.

The test starts with raising the temperature from room temperature to an initial temperature after which the feedback system starts to control the heating rate. The program scans the load cell reading 1000 times every minute and records their average, which is then compared to a reference value to determine the change in the fiber tension. A threshold value of the fiber tension change, below which no adjustment in the heating rate is applied, was taken as 0.05 gm to account for the possible small environmental fluctuations.

2-4-3 Typical Results

Figure 2.6 shows the fiber tension profile during the standard cure cycle of AS4/3501-6 graphite/epoxy composites. A noteworthy observation is that the final value of the fiber tension prior to cool down is higher than the initial fiber tension value. As discussed in earlier sections, this net increase in the fiber

tension corresponds to the development of compressive stresses in the fibers. These stresses will be added to the stresses produced during cool down resulting in more residual stresses. In the following discussion, results of the modified cure cycles obtained by CLFS are presented.

Figure 2.22 shows the cure cycle obtained from CLFS for the AS4/3501-6 graphite/epoxy. Recall that the feedback system constantly adjusts the heating rate to keep the changes in the fiber tension to a minimum. The fiber tension curve is almost flat. Figure 2.23 shows the volume change data obtained from the volumetric dilatometer for the 3501-6 epoxy for a cycle in which the obtained modified cure cycle was approximated with linear segments. The resin volume change curve is almost flat after about 1.5 hours. It is believed that the polymer volume change before 1.5 hours has minor effect on the fiber tension because polymer viscosity is low and, hence, stress relaxation is very effective to cancel developed stresses. After 1.5 hours, the thermal expansion appears to balance the crosslinking shrinkage to keep the polymer volume almost constant. The initial temperature beyond which the computer starts to change the heating rate was 138°C which was taken from the trial and error results. Starting at a lower temperature will increase the duration of the cure cycle. On the other hand, starting from higher temperature will reduce the total amount of thermal expansion that may be needed to the balance the crosslinking shrinkage. The effect of initial temperature will be discussed in Chapter 3. It should be noted that the total duration of the modified cure cycle.

Table 2.2 shows the Glass Transition Temperature (T_g) obtained using TMA for 3501-4 epoxy specimens cured using the standard and the modified cure cycles. The values of the T_g for the standard and the modified cycles are almost the same. The modified cure cycle was extended by 30 minutes to study the effect of such increase. The extension of the modified cure cycle does not increase the T_g and hence the cure was considered complete.

Table 2.2 Glass transition temperature for 3501-6 epoxy specimens cured using different cure cycles.

Cure Cycle	Standard Cycle	Modified Cycle	Modified Cycle + 30 min.
Glass Transition Temperature	170°C	171°C	172°C

2-5 SUMMARY

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A new test method to monitor the development of cure induced stresses in thermoset polymer composites was modified to study several materials systems. Test results were shown to be highly reproducible. Good qualitative agreement between test results and independently measured polymer volume change data was seen.

The test method was modified with a closed loop feedback control system to modify the cure cycles to produce less cure induced stresses. The modified cure cycles were found to produce less cure induced stresses compared to standard cure cycles while maintaining same glass transition temperature and in some cases shorter cure time.

3. EFFECT OF CURE CYCLE ON CURE INDUCED STRESSES

3-1 INTRODUCTION

In chapter 2, CIST method was presented. Sample results were demonstrated and it was shown that the test method can be used to monitor the cure induced stresses in single fiber thermoset polymer composites. The test method was modified with a closed loop feedback control system (CLFS) to alter the time-temperature schedule to reduce the cure induced stresses.

In this chapter, results of applying CIST and CLFS to different thermoset systems are shown. The CIST was used to study the development of fiber stresses during the cure of several thermoset polymers using cure cycles recommend by resin manufactures. The resins included were 3501-6 and 934 epoxies, 977-3 toughened epoxy, LTM-45 ELD and X343-59 low cure temperature epoxies, and 5250-4 bismaleimide (BMI). Fibers mainly were AS4 graphite and, for comparison, S2 glass in some cases. The CLFS was used to modify such standard cure cycles to reduce cure induced stresses. The results were analyzed and cure cycles were classified according to their effect on the cure induced stresses. The cure cycles obtained from the CIST were applied to cure composite laminates in vacuum-hot-press to demonstrate the applicability of such cure cycles to processing of composites.

3-2 STANDARD CURE CYCLES

In this section, fiber tension data are shown for different materials during their standard cure cycles. Volume change data whenever available are also shown for verification.

3-2-1 3501-6 and 934 Epoxies

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Volume change data during standard cure cycle of 3501-6 epoxy was shown in Figure 2.3. Typical fiber tension data for AS4/3501-6 graphite/epoxy single fiber composite was shown in Figure 2.6. For comparison purpose, the test was repeated with S2 glass fibers instead of the AS4 graphite fibers. The resulting fiber tension profile is shown in Figure 3.1. The trends of the results are similar for the two different fibers. The values of the fiber tension changes are different which may be attributed to the difference in the fiber dimensions, mechanical properties, and interface properties.

Standard cure cycle of 934 epoxy resin consists of: heating from room temperature to 116°C in 30 minutes; holding at 116°C for 60 minutes; raising temperature from 116°C to 177°C in 25 minutes; holding at 177°C for 240 minutes followed by cooling down to room temperature. Figure 3.2 shows the volume change data of this resin during such a cure cycle. There is crosslinking shrinkage during the first dwell temperature indicating the onset of cure. The crosslinking shrinkage increases significantly during the second dwell. Figure 3.3 shows the variation in an AS4 fiber tension during such a cure cycle. The trend of the results is similar to that 3501-6 epoxy. However, the values of the fiber tension change are different for these two

resins as should be expected because of the differences in the volume change characteristics of the resins and fiber/matrix interface properties. A comparison between Figures 3.2 and 3.3 shows that volumetric changes occurring during the first dwell have no effect on fiber stresses. In this case, cure induced stresses are governed by the chemical shrinkage during the second dwell only. There is a good agreement between volume change data (Figure 3.2) and fiber tension data (Figure 3.3), i.e. after the polymer has achieved relatively high viscosity, the thermal expansion causes a drop in the fiber tension where as shrinkage causes an increase. Figure 3.3 shows that the value of the fiber tension prior to the onset of the cool down (point B) is larger than that of the initial tension (point A) indicating the development of net compressive stresses in the fiber due to cure shrinkage.

3-2-2 974-3 Toughened Epoxy

Standard cure cycle of 977-3 toughened epoxy consists of: heating from room temperature to 180°C in 30 minutes; holding at 180°C for 360 minutes followed by cooling down to room temperature. Volume changes of this resin during the standard cure cycle are shown in Figure 3.4. There is volume expansion during the initial heating from the room temperature to the maximum cure temperature. The crosslinking shrinkage dominates the volume changes during the temperature hold at 180°C. Figure 3.5 shows fiber tension profile during the same cycle. Clearly, thermal expansion during the initial heating does not change fiber tension due to low polymer viscosity during this stage. As the resin starts to cure, its stiffness develops during the temperature hold at 180°C and chemical shrinkage causes fiber tension to increase. The rate of increase in the fiber tension during cool down due to thermal shrinkage.

3-2-3 5250-4 BMI

For the 5250-4 BMI resin, the standard cure cycle is divided into two parts: initial cure and post cure. Only the first part will be presented here (section 3-4 shows results for an alternative cure cycle for 52550-4 BMI including post cure). This cycle consists of: heating the sample from room temperature to 121°C in 35 minutes; holding at 121°C for 45 minutes; raising temperature from 121°C to 177°C in 25 minutes; holding at 177°C for 360 minutes followed by cooling down to room temperature. Figure 3.6 shows the volume change of 5250-4 BMI during the first part of the standard cure cycle. Figure 3.7 shows the fiber tension profile for AS4 in 5250-4 BMI matrix during the same cure cycle. The trends of the results are similar to those of the epoxy resins shown above. A comparison between Figures 3.6 and 3.7 shows that there is a good correlation between polymer volume change and fiber tension change during cure, i.e. once the polymer has developed significant stiffness, an increase or decrease in the polymer volume produces a decrease or increase, respectively in the fiber tension.

3-2-4 X343-59 and LTM-45 ELD Low Cure Temperature Epoxies

The standard low temperature cure cycle for X343-59 resin consists of: heating from 30°C to 66°C at 1.5°C/min; holding at 66°C for 840 minutes; cooling to 30°C at 2.5°C/min; heating to 177°C at 1.5°C/min; holding for 120 minutes; and cooling to 30°C at 2.5°C/min. The standard low temperature cure cycle for LTM-45 ELD consists of: heating from 30°C to 60°C at 0.5°C/min; holding at 60°C for 960 minutes; cooling to 30°C at 1.5°C/min; heating to 177°C at 0.25°C/min; holding for 120 minutes; and cooling to 30°C to 60°C at 0.5°C/min; holding for 120 minutes; and cooling to 30°C to 60°C at 0.5°C/min; holding for 120 minutes; and cooling to 30°C to 60°C at 0.5°C/min; holding for 120 minutes; and cooling to 30°C at 1.5°C/min. In a different set of experiments, these two resins were cured using an autoclave cure cycle that used for 3501-6 epoxy. That cycle consists of: heating from room temperature to 116°C in 30 minutes; holding at 116°C for 60 minutes; heating from 116°C to 177°C in 25 minutes; holding at 177°C for 240 minutes; and cooling to room temperature in 50 minutes. This cycle was proposed by the industry as an alternative to cut cure time of the standard low temperature cure cycles whenever autoclave cure is justified.

Figure 3.8 shows the volume change data of X343-59 during a low temperature cure cycle. Cure shrinkage occurs during the 66°C hold and continues during the 177°C hold. Figure 3.9 shows the variation in the fiber tension during the same cycle. There is a slight gradual increase in fiber tension during the 66°C hold due to cure shrinkage. During cool down from 66°C to 30°C at the end of the 66°C hold, another increase in the fiber tension occurs due to thermal contraction. As the resin is heated to post cure temperature, 177°C, the fiber tension decreases as a result of the thermal expansion. The tension drops below the initial tension value at the end of the heating ramp to 177°C. Some increase in the fiber tension can be seen during the 177°C hold time due to remaining cure shrinkage and stress relaxation as discussed below. Finally, fiber tension increases during cool down from 177°C to room temperature.

Figure 3.9 shows that fiber tension drops below its initial value during heating to post cure temperature. It should be noted that stress relaxation causes decay in any change in fiber tension (increase or decrease) from its initial value. In case of fiber tension drop below its initial value, stress relaxation causes fiber tension to increase again towards its initial value. Therefore, the increase in the fiber tension during the 177°C hold in Figure 3.9 is due to a combination of chemical shrinkage and stress relaxation (since fiber tension is below its initial value).

During the above cycle (Figure 3.9), curing at a relatively low temperature for a long time increases the viscosity of the resin. This causes the thermal expansion occurring during heating to post cure temperature, at 177°C, to be *effective* expansion. The term *effective* volume changes was defined in Section 2-2. This *effective* expansion causes a drop in the fiber tension below its initial value. This drop is larger than the increase in the fiber tension during the 177°C hold (compare thermal expansion during the 177°C ramp to that of the cure shrinkage during the 177°C hold in Figure 3.8). The final value of the fiber tension before cool down is still smaller than the initial fiber tension. This indicates that net tensile stresses are developed in the fiber prior to cool down. These tensile stresses will cancel part of the compressive stresses developed

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during the cool down and thus reduce residual stresses. In this particular case, stress free temperature will be below the post cure temperature. Stress free temperature was defined in Chapter 1.

Figure 3.10 shows the volume change data of X343-59 during the standard autoclave cure cycle of 3501-6 epoxy. A large amount of shrinkage occurs during the first temperature hold and additional shrinkage occurs during the second hold. Figure 3.11 shows the corresponding fiber tension variation. The shrinkage during the first temperature hold causes a significant increase in the fiber tension. The thermal expansion during the second temperature ramp decreases this tension. A slight increase in fiber tension can be seen during the second hold. Finally, cool down shrinkage increases fiber tension. Figure 3.11 shows that the final value of the fiber tension prior to cool down is more than its initial value. In this typical autoclave cycle, heating to the relatively high first temperature hold in short time just melts the resin and causes thermal expansion but does not develop tensile stresses in the fiber. This causes the chemical shrinkage to dominate the volume change and its effect will finally be added to cool down shrinkage causing more residual stresses.

Figure 3.9 shows that when the low temperature cure cycle is applied, the fiber is in a tensile stress state just before cool down. These tensile stresses will offset some of the compressive stresses that occur during cool down resulting in less final residual stresses. On the other hand, when typical autoclave cure cycle is applied, the fiber is in a state of compressive stress prior cool down. These compressive stresses combined with stresses developed during cool down produce higher residual stresses. A comparison between the two cycles suggests that the autoclave type cure cycle will result in more residual stresses than the low temperature cure cycle.

Figure 3.12 shows the volume change data of LTM-45 ELD epoxy during its standard low temperature cure cycle. Figure 3.13 shows the corresponding fiber tension profile. Compared to X343-59 resin (Figure 3.9), more stresses are developed during the 60°C hold. There is not much change in the volume during the 177°C hold. This suggests that the cure was almost completed during the slow heating ramp from 30°C to the post cure temperature. Fiber tension data agrees with the volume changes data showing only a slight increase during the post cure temperature hold (which may be due to stress relaxation). Similar to X343-59, the *effective* thermal expansion developed during heating to post cure temperature lowered the fiber tension below the initial value until the end of the cycle, i.e. fiber is under tensile stresses prior to cool down.

Figure 3.14 shows the volume change data of the LTM-45 ELD epoxy under the autoclave type cycle. The behavior of LTM-45 ELD under this cycle is similar to X343-59 under the same cycle, where most of the cure occurs during the first temperature hold. Figure 3.15 shows the corresponding fiber tension profile. Compressive stresses are developed in the composite prior to cool down. Hence, similar to X343-59, low temperature cure cycle is expected to produce less residual stresses than the autoclave type.

3-3 MODIFIED CURE CYCLES

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As discussed in Chapter 2, a closed loop feedback control system (CLFS) was developed to obtain modified cure cycles that reduce cure induced stresses. The method is based on searching for a heating path

that results in maximum cancellations of the unrelaxed stresses. Typical results of applying the method were demonstrated in Section 2-4-3. Results of applying the CLFS to different composite systems are given below.

3-3-1 3501-6 and 934 Epoxies

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Fiber tension data for an AS4 fiber in 3501-6 epoxy during standard cure cycle was shown in Figure 2.6. Results of CLFS for AS4 fibers were shown in Figure 2.22 where a modified cure cycle was shown to minimize change in the fiber tension during cure. The standard cure cycle was repeated again with S2 glass fibers instead of AS4 graphite fibers and results are shown in Figure 3.1. Figure 3.16 shows the modified cure cycle obtained for the glass fiber in 3501-6 epoxy resin as obtained from CLFS. The obtained cure cycle is similar to the modified cycle of the graphite fiber/3501-6 epoxy system. The slight differences are attributed to the differences in the interface properties. Since the method is based on reducing the strains through rescheduling of volume changes, the fiber properties do not seem to have much influence on the modified cycle.

Figure 3.17 shows a modified cure cycle obtained from the feedback control system for the AS4/934 graphite/epoxy system with the initial temperature taken as 138°C. By comparing the modified cycle with the standard cure cycle (Figure 3.3) we see that the modified cycle reduced fiber stresses during cure. However, the modified cycle is longer than the standard cycle because the cure cycle was extended by CLFS to allow for more stress relaxation of stresses due to chemical shrinkage. Also, results in Figure 3.17 indicate that a significant part of the thermal expansion was used early in the cure cycle (as can be seen from the rapid heating during first 60 minutes beyond initial heating). Applying thermal expansion early in the cure cycle is not recommended from stress cancellation point of view since stresses due to chemical shrinkage are more likely to relax in short time during this period without much temperature increase needed. Consuming thermal expansion during the early part of the cycle may cause chemical shrinkage to dominate the volume changes towards the end of the cycle where stress relaxation may not be very effective to cancel stresses in short time. This suggested introducing a temperature hold before the feedback starts. Another cycle was generated using CLFS with lower initial temperature to increase the available thermal expansion (rather than extending the cure cycle to have more stress relaxation) and with a temperature hold.

The approach of lowering the initial temperature and introducing the temperature hold leads to a shorter cure cycle with minimum cure-induced stresses as shown in Figure 3.18. The initial temperature was taken as 132°C. The new modified cycle results in an almost flat fiber tension curve. Also, the duration of this cycle is within that of the standard cure cycle. Stresses developed during the temperature hold beyond the initial heating are small because of rapid stress relaxation.

3-3-2 977-3 Toughened Epoxy

Figure 3.19 shows one of the modified cycles obtained from the CLFS. Stresses developed during cure are significantly less than that developed in the standard cycle. However, it was not possible to cancel all the stresses developed during cure. For 977-3 epoxy resin, the rate of the chemical reaction is only significant at high temperatures. The initial temperature was chosen as 149°C, which is higher than that used with other resin. In this case, the amount of thermal expansion available at the later part of the cure cycle to counteract the chemical shrinkage will be relatively small. This can be seen from Figure 3.19 where the maximum temperature was reached before the cure was completed (fiber tension still increasing). This causes chemical shrinkage to finally dominate and produce an increase in the fiber tension during the temperature hold at 180°C. At this late stage in the cure cycle, stress relaxation time is long and stresses were not relaxed (as can be seen, the fiber tension prior to cool down is still higher than the initial tension). An alternative approach utilized was to increase the time interval used in the CLFS to allow for more stress relaxation. In addition, the temperature was kept constant for 3 hours after the initial temperature to allow for more stress relaxation and increase the effective thermal expansion (by lowering gelation temperature). Stresses developed at early part of the cure cycle are more likely to relax during cure than those stresses developed at the later part. Figure 3.20 shows the results of the second modified cycle. It was possible to cancel most of the stresses due to chemical shrinkage. In addition, the duration of the new modified cure cycle is still within that of the standard cure cycle.

3-3-3 5250-4 BMI

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Figure 3.21 shows a cure cycle and the corresponding fiber tension profile obtained from the CLFS. The initial temperature was taken as 138°C. The fiber tension profile during the modified cure cycle is much uniform compared to that developed during the standard cycle. Figure 3.22 shows the volume change data of the 5250-4 BMI during the modified cycle. The cure cycle was approximated with linear segments. A good correspondence between fiber tension and volume change data can be seen after the viscosity of the polymer has increased.

To study the effect of the initial temperature, up to which the specimen is heated before enabling the CLFS, tests with different initial temperatures were conducted. Figure 3.23 shows a modified cycle in which the initial temperature was 143°C. The new cycle resulted in a more uniform fiber tension profile during cure. However, compared to the modified cycle shown in Figure 3.21, the new cycle is longer. Starting at high temperature reduces the total thermal expansion available to cancel shrinkage stresses. In this case, slower heat rate has to be applied by the feedback to allow for more stress relaxation. The slower heating rate resulted in longer cure cycle to complete the cure. A comparison between the two modified cycles shows the importance of choosing the initial temperature since it affects the combination of thermal expansion and stress relaxation required to cancel the chemical shrinkage.

The T_g data for 5250-4 BMI resin cured using the different cure cycles are shown in Table 3.1. For few experiments, the modified cure cycle was extended to study the effect of cure time increase. The value of

 T_g increases as the cure cycle is extended indicating that the cure is not completed. As was mentioned earlier, the 5250-4 BMI resin requires post cure at high temperature to complete the cure. Because the resin cure and hence shrinkage continues until the end of the cycle, Figures 3.21 and 3.22 show a steady continuous decrease in the resin volume and corresponding increase in the fiber tension towards the end of the cure cycle.

Table 3.1 Glass transition temperature for 5250-4 BMI specimens cured using different cure cycles.

Cure Cycle	Standard Cycle	Modified Cvcle	Modified Cycle + 30 min.
Glass Transition Temperature	165°C	153°C	163°C

3-3-4 X343-59 and LTM-45 ELD Low Cure Temperature Epoxies

The CLFS was used to modify the cure cycles of single fiber composites with X343-59 and LTM-45 ELD resins and AS4 fibers to reduce cure induced stresses. It should be noted that a major assumption in determining the required thermal expansion to cancel unrelaxed stresses is that the applied temperature increase will not change the polymer chemical reaction rate greatly. For low cure temperature epoxies, the chemical reaction is very sensitive to the increase in temperature. Hence, the temperature increment applied by the feedback control may greatly increase the rate of the reaction causing more chemical shrinkage. To cancel the additional shrinkage, the feedback control will apply more temperature. The new temperature increment will further increase the reaction rate and generate more shrinkage and so on. In such cases, equilibrium at each point may not be possible. In this situation, cancellation of stresses may be achieved by allowing more stress relaxation rather than heating. However, increasing time intervals to allow for more stress relaxation slows down the chemical reaction significantly and as a result the CLFS will stop raising the temperature (since there is no shrinkage to cancel). In such cases, the cure cycle will be stopped before the cure is complete.

For X343-59 resin, Figure 3.9, most of stresses were developed during the post cure, so the feedback control was enabled only during this part of the cure cycle. The objective of the feedback control here is to apply the heat incrementally in such a way that the fiber tension remains constant at a certain value until the resin is cured. This kind of incremental heating prevents *effective* cure shrinkage from developing compressive stresses in the fibers during cure. Also, applying the temperature incrementally prevent large decay of tensile stresses due to thermal expansion by the stress relaxation. Reducing stress relaxation of tensile stresses should finally reduce residual stresses.

The long hold at low temperature was applied as is. Then the temperature was increased linearly to 130°C. At this point, feedback control was enabled. This temperature was high enough to start continue the

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chemical reactions and also to reach the 177°C limit in reasonable time. Figure 3.24 shows the obtained cure cycle and the corresponding fiber tension. As can be seen, controlling the temperature through the feedback control does not cancel shrinkage stresses completely and the curve is not flat. As discussed above, the increase in the temperature appears to also increase the chemical shrinkage through enhancing the reaction rate. This will result in net increase in the fiber tension instead of canceling it. As the reaction advances, its rate appears to decrease allowing the thermal expansion to start canceling the chemical shrinkage. However, the 177°C limit is reached before the tension returns to the initial value. The completion of the cure was checked by measuring Tg. The obtained cycle gives 171°C versus 174°C for the standard low temperature cure cycle. The difference appears to be statistically insignificant since the highest temperature 177°C is applied in both cases for similar time.

Figure 3.25 shows the case where feedback control was enabled at arbitrarily chosen lower temperature $(110^{\circ}C)$. Better control over the reaction was possible resulting in an almost flat curve. So starting at low temperature gives the advantage of slower reaction rate but the reaction may stop before the cure is completed because of vitrification. It should be noted that starting at higher temperature is expected to increase the reaction rate and complete it in shorter time but offers less control as was seen when the feedback control was enabled at $130^{\circ}C$.

Other optimization runs were performed to find autoclave type cure cycles with minimum residual stresses. Figures 3.26 and 3.27 show the obtained fiber tension and resulting volume change, respectively. Again, at one stage in the cure cycle there is a region in which the feedback control does not cancel stresses because the chemical reaction rate also increases greatly as the temperature increases. However, the final value of the fiber tension is almost identical to initial value. This indicates that there are no cure induced stresses developed during such a cycle in the composite. Such an optimized cycle will produce less residual stresses than typical autoclave cure cycle shown in Figure 3.4 in which final value of the fiber tension is more than the initial tension value. The increase in the fiber tension above the initial value indicates compressive cure stresses in the composite. These compressive stresses will be added to the compressive stresses developed during cool down producing more residual stresses. Glass transition temperature for such a cycle is 176°C which means that the reaction is complete

Using the optimized autoclave type cycle (Figure 3.26), the resin can be cured in a shorter time than for the optimized low temperature cure cycle, Figure 3.24. However, a comparison between the two cycles shows that the optimized low temperature cure cycle has the advantage of developing tensile stresses in the composite during cure versus almost zero stresses in the optimized autoclave type cycle. These tensile stresses will decrease the final residual stresses in the composite.

The feedback control was also used to obtain an optimized low temperature cure cycle for LTM-45 ELD resin with AS4 fibers. Similar to the case with X343-59, during the feedback control, the temperature increase also increases the reaction rate significantly preventing canceling chemical shrinkage stresses completely. After some trials, the cycle shown in Figure 3.28 appears to decrease stresses during the first

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stage of the cure cycle. The post cure was used as is since there is no significant variation in fiber tension. Figure 3.29 shows the corresponding volume change data.

The optimized cure cycle shown in Figure 3.28 produces almost zero stresses during the first part of the low temperature cure cycle. This may be helpful to maintain dimensional stability of the parts during this stage (before post cure at higher temperature). Also, when this optimized cycle is compared with typical low temperature cure cycle (Figure 3.13), the typical cycle results in compressive stresses at the end of the first stage of the cycle (as indicated by the increase in the fiber tension prior to cool down from 60°C to 30°C). These compressive stresses will reduce the tensile stresses that will be developed during the heating ramp to post cure temperature. As discussed earlier, tensile stresses help to reduce residual stresses. The optimized cycle has the advantage of almost zero stresses developed during the first stage. Thus, it keeps more tensile stresses in the composite (which would be used to cancel stress due to shrinkage during the first stage) and hence less final residual stresses.

3-3-5 Applications

Results of the CIST and CLFS indicate that the modified cycles can effectively reduce stresses produced during cure of single fiber model composites. However, composite laminates contain high fiber volume fraction. Also, during manufacturing there may exist gradients in the temperature and degree of cure across laminate thickness especially in thick parts. These factors and others should be considered to modify cure cycles for composite laminates to reduce residual as will be addressed in chapter 4. In this part of the study, the modified cure cycle as obtained from the CLFS was applied to composite laminates without any changes.

The curvature of unsymmetric laminate was used as a measure of the total residual stresses. The material used was 3501-6/AS4 graphite/epoxy prepreg from Hercules. The composite laminates were cured in a vacuum hot press. Six unsymmetric 1X10 inches $[0^{\circ}_{4}/90^{\circ}_{4}]$ strips were cured using the standard cure cycle (Figure 2.2) and another six strips were cured using the modified cycle as obtained from the CLFS (Figure 2.22). The same values of the pressure, vacuum, and cool down rate were used in both the cure cycles. The specimens were made narrow to limit the effect of the curvature in the 90° direction. No machining on the specimens was done to avoid any stress relaxation. The dimensionless curvature of the specimens was calculated as follows,

Dimensionless Cuvature =
$$\frac{8tD}{4D^2 + L^2}$$
 (6)

where D, L, and t are the midpoint deflection, projected length of the specimen, and thickness of the specimen. A photograph of two representative specimens cured using the two cycles is shown in Figure 3.30 (a). The modified cycle reduces the specimen curvature. A plot of the dimensionless curvature (Equation 6) shows that the modified cycle reduces the curvature of the unsymmetric laminates by about 14% (Figure 3.30 (b)). The modified cycle was also shown to complete the cure as measured by the glass

transition temperature. Additional modification of the cure cycle to account for the effect of possible temperature and cure gradients and fiber volume fraction in the laminates may further reduce the final residual stresses.

3-4 EFFECT OF POST CURE AND COOL DOWN RATES

Stresses may develop in fibers during cure, post cure, and cool down. In the above sections, results of CIST and CLFS were shown mainly for stresses develop during cure. In this section, results of CIST for fiber tension changes during post cure and cool down are presented. The materials used in this section are AS4 fibers in 5250-4 BMI. Standard cure cycle for 5250-4 BMI was presented in section 3-2-3. In this section, an alternative cure cycle will be used. This cycle consists of: heating the sample from room temperature 177°C in 30 minutes; holding at 177°C for 360 minutes; cooling down to room temperature in 100 minutes. The post cure consists of: heating from room temperature to 205°C in 30 minutes; holding at 205°C for 360 minutes.

Figure 3.31 shows the variation in the fiber tension during such a time-temperature cycle. During the first part of the cure cycle, the fiber tension increases due to chemical shrinkage and during cool down due to thermal shrinkage. During post cure temperature hold, there is a slight increase in the fiber tension. This increase is the resultant of two opposite actions. The first is the decrease in fiber tension because of heating above previous cure temperature to post cure temperature (from177°C to 205°C). The second is the increase in fiber tension due to additional cure at high temperature. It seems that the increase in the fiber tension due to additional cure shrinkage and thermal expansion due to additional heating are almost of equal value and as a result, the fiber tension is almost unaffected. The final value of the fiber tension after cool down from posture (point B) is similar to that obtained after cool down from curing at 177°C (point A). Similar findings were reported by White and Hahn for composite laminates subjected to post cure [22].

To study the effect of cool down rate on final fiber tension, after cool down from post cure, the specimen was heated to the same temperature (205°C) and cooled to room temperature at the same previous rate. Same value of final fiber tension was obtained (point C). In the next step, the specimen was heated to the same temperature (205°C) followed by cool down at slower rate. The final fiber tension at room temperature (point D) is approximately 7% lower than the previous recorded values. To verify that this reduction is due to the more stress relaxation allowed by slower heating rate, the specimen was heated again followed by cool down at same rates as from post cure twice. The reduction in the fiber tension disappeared when cool down rate was increased again (points E and F). The test was repeated and similar results were recorded. Results indicate that slower cool down rate may reduce fiber stresses due to cool down shrinkage. It should be noted that these results are for 5250-4 BMI and can not be generalized. Also, because of the dominant polymer volume fraction in CIST (a single fiber in infinite polymer), polymer viscoelasticity may be more effective than it will be in case of composites with high fiber volume fraction. Constraints applied by fibers on polymer reduce the extent of viscoelasticity as seen in resins.

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3-5 CLASSIFICATION OF CURE CYCLES

The results presented in the above section suggest that the major factors affecting such contribution are the amount of *effective* shrinkage and amount of *effective* expansion generated and extent of stress relaxation allowed. *Effective* shrinkage and expansion. For a given thermoset polymer and a given maximum cure temperature, the location of the gelation point is the governing factor that determines the amount of *effective* thermal expansion. The lower the gelation temperature, the larger the *effective* thermal expansion. *Effective* cure shrinkage produces compressive stresses in the fibers. The unrelaxed part of these stresses will be added to the cool down stresses increasing the residual stresses in the composite. The location of the gel point can be lowered to allow for more *effective* thermal expansion. The cure time beyond gelation can be extended by slowing down the heating rate. This may allow for more stress relaxation to overcome the stresses generated by the *effective* shrinkage. Depending on the balance between the three factors, the cure cycles can be classified into three types:

Type I: In this case the stresses generated by *effective* shrinkage are larger than the combined effect of the *effective* thermal expansion and stress relaxation. Figure 3.32 (a) shows a schematic of such a cycle as monitored by CIST. The tension prior to the onset of cool down is larger than the initial tension indicating net compressive fiber stresses due to *effective* shrinkage.

Type II: In this case the stresses generated by *effective* shrinkage are equivalent to the combined effect of the *effective* thermal expansion and stress relaxation. Figure 3.32 (b) shows a schematic of such a cycle as obtained by CLFS. The tension prior to the onset of cool down is the same as the initial tension indicating zero net fiber stresses due to cure volume changes. It should be noted that the balance can be generated through the whole cure cycle or just prior to onset of cool down, i.e. allow for increase or decrease in the fiber tension and then cancel these changes through the cycle.

Type III: In this case the stresses generated by *effective* shrinkage are smaller than the combined effect of the *effective* thermal expansion and stress relaxation. Figure 3.32 (c) shows a schematic of such a cure cycle as monitored by CIST. The tension prior to the onset of cool down is smaller than the initial tension indicating net tensile fiber stresses due to *effective* expansion.

Most of the standard cure cycles belong to Type I. This is attributed to the rapid heating to the maximum cure temperature or using short dwell at low temperature. In most of these cycles the gelation temperature is very close to the maximum cure temperature which allows for a small amount of *effective* thermal expansion. Type II cycle can be generated using CLFS as explained above. Type III cycles can be obtained by lowering the gelation temperature to increase *effective* expansion and allowing more stress relaxation. It should be noted that increasing the *effective* thermal expansion by increasing the cure temperature will not help in terms of the final residual stresses since this will increase the cool down shrinkage. So, Type III cycle is basically generated by lowering the gel point and allowing for more stress relaxation. Results for

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two typical resin systems, 3501-6 and 977-3 epoxies, are presented here. However, the method can be applied to other resin systems.

3-5-1 3501-6 Epoxy

Figure 3.33 (a) shows a Type I cure cycle of 3501-6 epoxy with corresponding fiber tension changes in an -AS4 graphite fiber. This cycle is the standard cure cycle recommended by resin manufacturer. The fiber tension prior to cool down is higher than the initial tension indicating net compressive fiber stresses. Figure 3.33 (b) shows a Type II modified cycle which was obtained using CLFS. The change in the fiber tension through the cure cycle is close to zero. The duration of the modified cycle is shorter than the standard cycle while producing the same glass transition temperature as was shown before in Section 2-4-3. These results were shown before and presented here for convenience.

Figure 3.33 (c) shows a Type III cure cycle. The cure was carried out at a low temperature for longer time to reach gelation at lower temperature. The effect of thermal expansion is larger than that in the standard cure cycle as can be seen by comparing the drop in the fiber tension corresponding to the increase in the temperature from the first to the second dwell temperature. The fiber tension prior to cool down is larger than the initial fiber tension indicating net tensile fiber stresses. These stresses will reduce the residual stresses developed during cool down by thermal shrinkage. The duration of this cycle is longer than the standard cure cycle.

A comparison between the three types of cycles shows that Type II cycle is the shortest and produces less residual stresses than the standard cure cycle. Type III cycle results in minimum residual stresses however it is the longest. The three types of cycles produce the same glass transition temperatures (170, 171, and 170°C respectively).

3-5-2 974-3 Toughened Epoxy

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The standard cure cycle of 974-3 (Figure 3.34 (a)) is a Type I cycle. The CLFS was applied to obtain Type II and III cycles. The objective in Type II cycle is to have a balance between *effective* shrinkage and the summation of *effective* expansion and stress relaxation. In this case there will be no cure induced stresses and residual stresses will that produced only during the cool down. For Type III cycle, the objective is to increase the summation of the *effective* thermal expansion and stress relaxation more than the *effective* shrinkage.

Standard cure cycle of 977-3 is a Type I cycle (shown in Figure 3.24 (a) for convenience). A Type II was developed using CLFS as shown in section 3-3-2 (shown in Figure 3.24 (b) for convenience). The CLFS was applied to produce Type III cycle with more *effective* thermal expansion and stress relaxation. The initial heating temperature was taken as 118°C. However, at this low temperature a longer temperature hold is required to advance the cure before increasing the temperature (such that thermal expansion will be *effective*). Figure 3.34 (c) shows the Type III cure cycle. The final fiber tension prior to cool down is lower than the initial fiber tension. This indicates the development of tensile stresses in the fiber prior to cool

down. These tensile stresses will cancel part of the cool down stresses and hence this type of cycle will produce less residual stresses than the Type I and II cure cycles. However, Type III cycle is considerably longer than the other two types of cycles.

3-5-3 Applications

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In this part of the study, the three types of cycles for 977-3 toughened epoxy are applied to cure composite laminates and residual stresses are compared. The composite system used here is IM7/977-3 graphite/toughened epoxy prepreg. The cure cycles are used to cure unsymmetric $[0^{\circ}/90^{\circ}]$ composite laminates. The curvature of the cured laminates was used as an indicator for residual stresses. The laminates were cut and laid up manually and cured in the same location of the single fiber composite specimen to reduce the variation between the cure parameters between CIST and CLFS and the laminates. The dimensions of the specimens are 3/8x3 inches and the lay up is $[0^{\circ}_{1}/90^{\circ}_{2}]$. The cure cycles are shown in Figure 3.34 (a), (b), and (c). The same cool down rate and pressure schedule were used in the three cure cycles. The specimens were made narrow to limit the effect of the curvature in the 90° direction. No machining on the specimens tend to have delamination which may affect the comparison. Curvature of the cured specimens was calculated using Equation 6.

A photograph of representative specimens cured using the three different cycles is shown in Figure 3.35 (a). The modified cycles reduced the specimen curvature. A plot of the dimensionless curvature is shown in Figure 3.35 (b) for the different specimens. Compared to the standard cure cycle, the first modified cycle (Type II) reduced the curvature by about 16% while the second modified cycle (Type III) reduced it by about 19%. The cure was completed in all cases since the glass transition temperatures in all three cases were statistically same (about 205-210°C).

3-6 EFFECT OF MODIFIED CURE CYCLES ON MECHANICAL PROPERTIES

The objective here is to investigate the effect of changing the cure cycle on the mechanical properties of the composite laminates which is a major concern for structural applications. The properties affected most by the change in the cure cycle are those which are matrix dependent. The inplan shear strength and shear modulus were chosen as representative mechanical properties.

The composite system used in this part of the study is IM7/977-3 graphite/toughened epoxy prepreg. Symmetrical specimens (10.5X10.5 inches; $(45^{\circ}_{2}, -45^{\circ}_{4}, 45^{\circ}_{2})$) were cut and laid and cured in a vacuum-hotpress. The cured laminates were then cut to 1X10 inches strips and tested in tension in a universal testing machine (MTS machine). Strain gauges were used to measure strains in 0° and 90° directions. Load and strain data were collected at close intervals to allow for the determination of the stiffness. Six specimens of each cycle were tested and shear strength and shear modulus were calculated.

Figure 3.36 (a) shows the shear strength data for the specimens cured with the three types of cycles. The Type II cycle increased the shear strength by about 15%. The increase in the shear strength may be due to the reduction in the residual stresses and micodamages caused by residual stresses. Type III cycle reduced

the failure shear stress by about 25%. This may be due to the long heating beyond cure of the specimens, which is similar to post curing that was shown to reduce shear strength in epoxy composites [58]. Figure 3.36 shows that the three cure cycles results in approximately the same shear modulus.

Among the three types of cycles, Type II cycle appears to produce the best results. While the duration is close to the standard cure cycle, it reduces the residual stresses and enhances the mechanical properties.

3-7 SUMMARY AND CONCLUSION

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In this chapter, results of a new approach to monitor and reduce cure induced stresses in thermoset polymer composites were presented. The approach utilizes a new test setup and a feedback control system. The method was applied to study the cure induced stresses in several thermoset polymer composites cured using the cure cycles recommended by prepreg manufacturers. Modified cure cycles that reduce cure induced stresses were demonstrated. It was shown that the modified cure cycles not only reduce residual stresses in composite laminates but might enhance the matrix-dominated mechanical properties as well, while maintaining the glass transition temperature as those of the standard cure cycles.

A general classification of the cure cycles according to their cure induced stresses was presented and demonstrated with two examples. The classification can be used to compare qualitatively between different cure cycles to reduce cure induced stresses.

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Figure 1.1 Cure cycle modification to reduce residual stresses.

Adapted from White, S. R. and H. T. Hahn. 1993. "Cure Cycle Optimization for the Prediction of Processing-Induced Residual Stresses in Composite Materials," J. of Composite Materials, 27(14), 1352-1378.



(b) Spring-back

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Figure 1.2 Another method for cure cycle modification to reduce residual stresses

Adapted from Sarrazin, H., B. Kim, S-H. Ahn, and G. S. Springer. 1995. "Effect of Processing Temperature and Layup on Springback," J. of Composite Materials, 29(10), 1278-1294.



Figure 1.3 Comparison between test results and model predictions

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Adapted from White, S. R. and H. T. Hahn. 1993. "Cure Cycle Optimization for the Prediction of Processing-Induced Residual Stresses in Composite Materials," J. of Composite Materials, 27(14), 1352-1378.



(a) Polymer shrinkage increases the fiber tension.



(b) Polymer expansion decreases the fiber tension.

Figure 2.1 Mechanism of cure induced stress test.

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Figure 2.2 Details of the PVT apparatus.

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Figure 2.3 Variation in polymer volume during standard cure cycle.

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Figure 2.4 Schematic diagram of single fiber test.



Figure 2.5 Cure induced stress test



Figure 2.6 Variation in the fiber tension during the standard cure cycle of 3501-6 epoxy.



Figure 2.7 Fiber tension during the standard cure cycle for a single fiber coated with a thin layer of 3501-6 epoxy.

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Figure 2.8 Fiber tension change during a typical two dwell cure cycle.



Figure 2.9 Effect of temperature change on the fiber tension (no polymer).



Figure 2.10 Effect of embedded fiber length on the variation in the fiber tension during cure.



Figure 2.11 Effect of initial fiber tension on the variation in the fiber tension during cure.

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Figure 2.12 Effect of changing cure cycle on fiber tension in AS4/Epon 828 single fiber composites.



Figure 2.13 Schematic of the model used to calculate fiber stress during cure.



Figure 2.14 Fiber stresses during a typical two dwell cure cycle.

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Figure 2.15 Cure induced stresses produced fiber fracture in a pretensioned fiber.



Figure 2.16 Variation in fiber stresses during the modified cure cycle.

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Figure 2.17 Fiber tension during an intermediate cure cycle obtained using trial and error modification method.



Figure 2.18 Fiber tension during a modified cure cycle of 3501-6 epoxy obtained using trial and error modification method.

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Figure 2.19 Volume changes during the modified cure cycle of 3501-6 shown in Figure 2.18.

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Figure 2.20 Variation in fiber tension during another modified cure cycle of 3501-6 epoxy.



Figure 2.21 Flow chart of the Closed Loop Feedback System (CLFS) used to obtain cure cycles with reduced cure induced stresses.

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Figure 2.22 Fiber tension data during a cure cycle modified using CLFS.



Figure 2.23 Volume change data during a cure cycle modified using CLFS.



Figure 3.1 Variation in the fiber tension during the standard cure cycle of 3501-6 epoxy.



Figure 3.2 Volume change of 934 epoxy during the standard cure cycle.



Figure 3.3 Variation in the fiber tension during the standard cure cycle of 934 epoxy.

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Figure 3.4 Volume change of 977-3 epoxy during the standard cure cycle.



Figure 3.5 Fiber tension data during the standard cure cycle of 977-3 epoxy resin.



Figure 3.6 Volume change data of 5250-4 BMI resin during the standard cure cycle.



Figure 3.7 Variation in the fiber tension during standard cure cycle of 5250-4 BMI.



Figure 3.8 Volume change of X343-59 during the standard low temperature cure cycle.



Figure 3.9 Variation in the fiber tension during the standard low temperature cure cycle of X343-59 epoxy.

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Figure 3.10 Volume change of X343-59 during an autoclave type cure cycle.



Figure 3.11 Variation in the fiber tension during an autoclave type cure cycle of X343-59.

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Figure 3.12 Volume change of LTM-45 ELD epoxy during the standard low temperature cure cycle.



Figure 3.13 Variation in the fiber tension during the standard low temperature cure cycle of LTM-45 ELD epoxy.

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Figure 3.14 Volume change of LTM-45 ELD during an autoclave type cure cycle.



Figure 3.15 Variation in the fiber tension during an autoclave type cure cycle of LTM-45 ELD epoxy.

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Figure 3.16 A modified cure cycle for S2/3501-6 glass/epoxy system.

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Figure 3.17 A modified cure cycle for AS4/934 graphite/epoxy.

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Figure 3.18 Another modified cure cycle for AS4/934 graphite/epoxy.



Figure 3.19 A modified cure cycle for AS4/977-3 graphite/epoxy.

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Figure 3.20 Another modified cure cycle for AS4/977-3 graphite/epoxy.

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Figure 3.21 A modified cure cycle for AS4/5250-4 graphite/BMI.



Figure 3.22 Volume change data of 5250-4 BMI during the modified cure cycle shown in Figure 3.21.

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Figure 3.23 Another modified cure cycle for 5250-4 BMI with higher initial temperature.

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Figure 3.24 Variation in the fiber tension during a modified low temperature cure cycle for AS4/X343-59 graphite/epoxy.



Figure 3.25 Variation in the fiber tension during another modified low temperature cure cycle for AS4/X343-59 graphite/epoxy.



Figure 3.26 Variation in the fiber tension during a modified autoclave type cure cycle for AS4/X343-59 graphite/epoxy.



Figure 3.27 Volume change of X343-59 during the modified cure cycle shown in Figure 3.26.



Figure 3.28 Variation in the fiber tension during a modified low temperature cure cycle for LTM-45 ELD.



Figure 3.29 Volume change of LTM-45 ELD epoxy during the modified cure cycle shown in Figure 3.28.

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(a) Photograph.



Figure 3.30 Curvature of unsymmetric AS4/3501-6 graphite/epoxy laminates cured with different cure cycles.





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Figure 3.32 Schematic showing the effect of the cure cycle selection on the contribution of cure volume changes to fiber stresses.

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Figure 3.33 Different types of cure cycles for AS4/3501-6 graphite/epoxy.

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Figure 3.34 Different types of cure cycles for AS4/974-3 graphite/epoxy.



(a) Photograph.



Figure 3.35 Curvature of unsymmetric IM7/977-3 graphite/epoxy laminates cured using different cure cycles.



(b) Shear Modulus (ksi)

