Rowan University

Rowan Digital Works

Theses and Dissertations

5-8-2002

Characterization of structural changes in thermally enhanced Kevlar-29® Fiber

James W. Downing Jr. Rowan University

Follow this and additional works at: https://rdw.rowan.edu/etd



Let us know how access to this document benefits you share your thoughts on our feedback form.

Recommended Citation

Downing, James W. Jr., "Characterization of structural changes in thermally enhanced Kevlar-29® Fiber" (2002). *Theses and Dissertations*. 1429.

https://rdw.rowan.edu/etd/1429

This Thesis is brought to you for free and open access by Rowan Digital Works. It has been accepted for inclusion in Theses and Dissertations by an authorized administrator of Rowan Digital Works. For more information, please contact LibraryTheses@rowan.edu.

CHARACTERIZATION OF STRUCTURAL CHANGES IN THERMALLY ENHANCED KEVLAR-29® FIBER

by James W. Downing Jr.

A Thesis

Submitted in partial fulfillment of the requirements of the Master of Science Degree of The Graduate School at Rowan University 5/8/02

Approved by		
Date Approved_	May 8	2002

ABSTRACT

James W. Downing Jr. CHARACTERIZATION OF STRUCTURAL CHANGES IN THERMALLY ENHANCED KEVLAR-29® FIBER 2002/03

Dr. James Newell
Master of Science in Chemical Engineering

The purpose of this exploratory investigation was to elucidate the structural mechanism accounting for the enhanced compressive properties of heat-treated Kevlar-29® fibers. A novel theory was set forth that hydrogen bond disruption and concurrent misorientation of crystallites may account for the observed augmentation of compressive properties. To examine the said theory, virgin Kevlar-29® fibers were characterized by Thermogravimetric analysis (TGA) and Differential Scanning Calorimetry (DSC) in an effort to determine if crosslinking and/or hydrogen bond disruption was responsible for the improved behavior in compression. Additionally, Kevlar-29® fibers that had been exposed to treatment temperatures of 400, 440, and 470 °C were profiled by Fourier-Transform Infrared Spectrophotometry (FTIR) to determine if crosslinking and/or hydrogen bond obfuscation had been promoted. The results indicate that both mechanistic changes are occurring within the Kevlar-29®, albeit in different regions of the rigid-rod polymer. In particular, heat-treatment of poly-p-phenylene terephthalamide results in crosslinking of its skin region and hydrogen bond disruption within the core realm.

MINI-ABSTRACT

James W. Downing Jr. CHARACTERIZATION OF STRUCTURAL CHANGES IN THERMALLY ENHANCED KEVLAR-29® FIBER 2002/03

Dr. James Newell
Master of Science in Chemical Engineering

The purpose of this exploratory investigation was to elucidate the structural mechanism accounting for the enhanced compressive properties of heat-treated Kevlar-29® fibers. To this end, PPTA fibers were examined using Thermogravimetric Analyzer, Differential Scanning Calorimeter, and Fourier Transform Infrared Spectrophotometer. Summarily, the heat-treatment of poly-*p*-phenylene terephthalamide results in crosslinking of its skin region and hydrogen bond disruption within the core realm.

Acknowledgements

Grateful acknowledgement is given to the following persons/organizations for their significant contributions to this thesis or support of the author thereof:

To my lovely wife, Lori M. Faer-Downing, for her unyielding love and support throughout this academic journey

To my employer, 3M, for allowing me to utilize its equipment to study a competitive polymer

To my thesis advisor, Dr. James Newell, Ph.D, for sharing my passion for high-performance polymers

To the other members of my thesis committee, Dr. Stephanie Farrell, Ph.D and Dr. Kevin Dahm, Ph.D, for sharpening and refining this opus

Table of Contents

1.0 Introduction
2.0 Literature Review
3.0 Mechanistic Discussion
4.0 Experimental
5.0 Results and Discussion
6.0 Conclusions
7.0 Recommendations

List of Illustrations

- Figure 1. The Primary Structure of Kevlar®
- Figure 2. The Secondary Structure of Kevlar®
- Figure 3. The Pleated Sheet Microstructure of Kevlar®
- Figure 4. Thermogravimetric Analyzer
- Figure 5. Differential Scanning Calorimeter
- Figure 6. Fourier Transform Infrared Spectrophotometer
- Figure 7. Allylic Site for Free Radical Formation
- Figure 8. Thermogravimetric Isotherms of Kevlar-29® Fiber
- Figure 9. DSC Thermogram of Kevlar-29® Fiber
- Figure 10. FTIR Spectra of Heat-treated Kevlar-29® Fiber

List of Tables

- Table 1. Tensile and Compressive Strengths of Several Rigid Rod Fibers
- Table 2. Tensile and Recoil Compressive Strengths of E-beam Treated Kevlar-29® Fiber
- Table 3. Effects of Temperature, Ramp, and Dwell on Ratio of σ_c/σ_t of Kevlar-29® Fiber

1.0 Introduction

Poly-p-paraphenylene terephthalamide (PPTA) is produced in the U.S. by DuPont under the trade name Kevlar®. Kevlar® is an organic fiber with a distinct chemical composition of wholly aromatic polyamides (aramids). Since its development in 1965 by DuPont research scientist Stephanie Kwolek, the commercial and scientific interest in poly-p-phenylene terephthalamide has increased exponentially. This can be attributed to Kevlar's® unique combination of high tensile strength and modulus, toughness, and thermal stability. In air, PPTA demonstrates seven times the tensile strength of steel on an equal weight basis. In sea water, this advantage in tension increases by a factor of twenty. Thus, applications utilizing this rigid-rod polymer abound in nearly every realm of industry. Some examples of the diverse products that incorporate Kevlar® fiber, staple, or floc include brake pads, bullet-proof vests, kayaks, mooring lines, and high performance tires. Moreover, the potential use of Kevlar® in structural composites is only now being realized, and investigations into this newest group of engineering materials are being pursued with great zeal.

The exceptional strength of Kevlar® fiber in tension is a direct result of its primary, secondary, and tertiary chemical structure. The primary, or molecular structure of the Kevlar® extended chain can be classified as monoclinic with the following lattice parameters: $\mathbf{a} = 7.87$ angstroms, $\mathbf{b} = 5.18$ angstroms and $\mathbf{c} = 12.9$ angstroms with a 90°

unit cell angle.² Its "rigid-rod" characteristic is a function of the para substitution of the benzene ring,¹ as well as the covalent bond strength in the **c** direction,⁴ which allows an axial stress to be distributed evenly throughout the highly linear macromolecular chain. Kevlar's® primary structure is illustrated in Figure 1.

The secondary structure of Kevlar®, depicted in Figure 2, can be characterized as a pleated sheet configuration that is oriented perpendicular or transverse to the fiber axis. Pleats form within the core region when PPTA enters the coagulation bath, in response to the relaxation of the local stress field at the onset of coagulation. ¹² This pleated conformation of the polymer chains is primarily governed by a consortium of intra- and intermolecular interactions between the conjugated groups within the PPTA molecular structure. These interactions include (1) the resonance effect attempting to stabilize coplanarity of the amide groups and the phenylene groups, (2) the counteracting steric hindrance found between the oxygen and an ortho-hydrogen of the p-phenylene moieties, as well as between the amide hydrogen and an ortho-hydrogen of the terephthalic segment, and (3) the centro-symmetric pairs of hydrogen bonds between the amide and carbonyl groups. 16 In a PPTA crystal, intermolecular hydrogen bonding between the C=O and the N-H acts along the **b** direction.⁵ The pleating of the fibrils is superimposed on the fibrillar structure of PPTA, with a variation from linearity of approximately 5° and a periodicity of 500 nm as determined by optical microscopy. ¹⁷ A schematic of the pleated nature of the PPTA microstructure is illustrated in Figure 3. Other analytical methods, including X-ray diffraction and electron microscopy, have corroborated the existence of the pleated sheet secondary structure of Kevlar® fibers. 16, 18 It is this secondary structure that establishes the nearly perfect uniaxial linearity that defines Kevlar®. However,

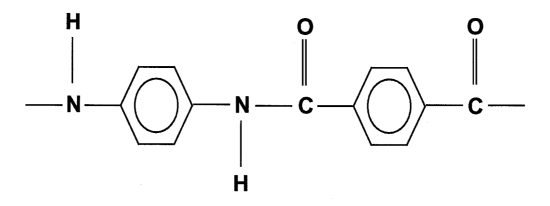


Figure 1. The Primary Structure of Kevlar®

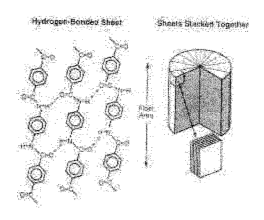


Figure 2. The Secondary Structure of Kevlar®

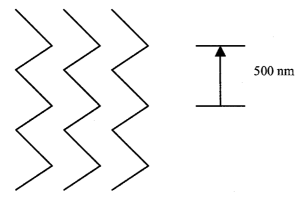


Figure 3. The Pleated Sheet Microstructure of Kevlar®

Kevlar's® microstructure can not be described as perfectly linear due to the existence of microvoids and other crystalline defects located mainly around the periphery of the fibers. 19

Additionally, Kevlar® possesses a clearly discernible tertiary structure. During the course of the spinning and post-treatment stages of Kevlar® processing, a skin-core morphology develops. The major distinction between skin and core regions is the higher degree of order and intermolecular bonding in the core region as compared to the skin. Consequently, the core region possesses an elastic modulus of 60.8 GPa. In contrast, the skin region of Kevlar's® tertiary structure is virtually non-crystalline, and thus possesses a significantly lower elastic modulus of approximately 13.4 GPa. However, the skin may be critical in the prevention of crack propagation within the crystalline core region.

The tremendous behavior of Kevlar® in tension can be attributed to structural characteristics including its stiff, extended chain conformation and nearly perfect uniaxial orientation. Despite the superlative tensile strength and modulus of PPTA in its fiber form, it possesses a comparatively low compressive strength due to its highly linear and regular microstructure. Kevlar's® compressive strength is merely 1/10 of its ultimate tensile strength. Additionally, Kevlar® exhibits a compressive to tensile strength ratio of between .13 and .25, while that of carbon fiber often exceeds 1.8 This imbalance of the high tensile and low compressive properties has proved to be a limiting factor in its widespread incorporation into many structural composites. Compressive weakness relative to behavior in tension is not unique to Kevlar®, but is quite pervasive within high performance fibers as illustrated in Table 1.

Table 1

Tensile and Compressive Strengths of Several Rigid Rod Fibers

High Performance Fiber	Tensile Strength (GPa)	*Compressive Strength (GPa)
Polybenzoxazole (PBO)	5.7	0.28
Polybenzothiazole (PBZT)	4.1	0.2
Kevlar 29	2.8	0.35
Kevlar 49	2.9	0.37
Carbon Fiber (AS-4)	3.6	1.44
*Data from recoil testing		

Aircraft fabricators find that advanced carbon composites more than not win out over Kevlar®, based largely on its inferior compressive strength and the greater stiffness that carbon fibers afford the structure. Clearly, enhancement of the compressive properties of Kevlar® would markedly improve the utility and commercialization of this rigid-rod polymer.

2.0 Literature Review

Efforts to improve the compressive strength of Kevlar® fibers have been conducted by several groups. Approaches to enhance the axial compressive strength of Kevlar® may be classified into two main techniques. The first entails the incorporation of an additive or modifier into the Kevlar® microstructure during the wet spinning or polymerization process with the proposed goal of facilitating the formation of intermolecular covalent crosslinks. The second involves achieving the same goal by subjecting the PPTA fibers to a post-spinning treatment, such as elevated temperature or electron bombardment.

Sweeny et al.¹⁰ attempted to improve the compressive modulus of Kevlar-29® by incorporating an activated aryl halogen into the molecular sequence of PPTA. The activated aryl halogen functioned as a copolymer and thus was introduced during the polymerization of the aromatic polyamides. The resultant prefiber was comprised of both PPTA and halogenated aryl moieties. The extruded, halogen-bearing polymer was then subjected to various temperatures for various durations in order to develop aryl free-radicals. The aryl radicals would then facilitate the formation of crosslinks via a ring-coupling phenomenon. The degree of crosslinking could be quantified as a function of percentage of polymeric halide retained. Sweeny et al. ¹⁰ recognized that a high degree of crosslinking could not be accommodated without affecting the interchain hydrogen

bonding. Moreover, it was concluded that the crystal structure of Kevlar® would not accommodate a high level of crosslinking.

Jiang et al.⁷ also developed an activated form of PPTA by incorporating XTA, a benzocyclobutene-modified derivative of terepthalic acid, into the polymer backbone during prefiber production. Various concentrations of the XTA were spun into the Kevlar® fibers during the dry-jet wet spinning stage of the production process in order to control the degree of crosslinking. The XTA was triggered into reactive status via heat treatment at temperatures within the range of 325 and 425 °C. Summarily, the XTA-rich PPTA fibers appeared to exhibit crosslinking as verified by swelling assays.

Accordingly, recoil tests showed a slight improvement in the compressive strength of these modified Kevlar® fibers, accompanied by a decrement in tenacity.

An investigation by Newell et al.³ provides an example of enhancement of asspun, unmodified Kevlar® fiber. Unlike more common methods including thermal treatment or the addition of a crosslinking agent during spinning, this paper describes the use of electron beam bombardment to induce what is purported to be crosslinking in Kevlar-29®. The recoil compressive strength of the fiber increased significantly within increasing radiation exposure, reaching 487 MPa at a dose of 1100 KGy. Other key findings are depicted in Table 2. The augmentation of axial compressive strength adversely affected the tensile strength of the poly-*p*-phenylene terephthalamide by approximately twenty percent.

Sweeney et al. 11 achieved similar results through thermal treatment of an unmodified form of Kevlar-29® fiber. This group, whose results are summarized in

Table 2

Tensile and Recoil Compressive Strengths of E-beam Treated Kevlar-29® Fiber

Radiation Level (kGy)	Mean Tensile Strength (MPa)	Mean Recoil Compressive Strength (MPa)		
0	2160 +/- 60	365 +/- 6		
100	2043 +/- 120	368 +/- 8		
200	1996 +/- 86	381 +/- 9		
500	1891 +/- 71	404 +/- 8		
1000	1786 +/- 93	472 +/- 7		
1100	1723 +/- 87	487 +/-7		

Table 3, developed a central composite design to study the orthogonal and interactive effects of three factors on compressive and tensile strength. Three levels of maximum treatment temperature, soak rate, and soak time (dwell) were examined within the scope of this study. It was concluded that maximum treatment temperature had a marked effect on the compressive strength of Kevlar® fiber, with the higher treatment temperature resulting in a significant enhancement of compressive strength. Moreover, as seen in the aforementioned studies, this improvement in compressive behavior was accompanied by a compromise in strength in tension.

Table 3 Effects of Temperature, Ramp, and Dwell on Ratio of $\sigma_c/\,\sigma_t$ of Kevlar-29® Fiber

Average Ratio	Ramp	Average Ratio	Dwell	Average Ratio
	°C/min		min.	
0.46	2.5	0.67	5	0.49
0.63	5	0.62	7	0.71
0.90	7.5	0.60	10	0.67
	0.46 0.63	Ratio Ramp °C/min 0.46 0.63 5	Ratio Ramp Ratio °C/min 0.46 2.5 0.67 0.63 5 0.62	Ratio Ramp Ratio Dwell °C/min min. 0.46 2.5 0.67 5 0.63 5 0.62 7

3.0 Mechanistic Discussion

Clearly the enhancement of compressive properties can be realized in both modified and unmodified Kevlar-29® fibers by employing a post-spinning stimulus such as thermal or ionization treatment. However, the structural mechanism accounting for the observed increase in compressive properties and accompanying decrement in tenacity has largely been a source of debate. In particular, unresolved is whether this compressive enhancement was due to a crosslinking phenomenon or hydrogen bond obfuscation and concurrent crystallite misorientation.

It was originally assumed that the effects of crosslinking within the PPTA could provide the only explanation accounting for the enhanced compressive strength. Such crosslinking would provide a covalent bond lattice in the a direction. There is strong empirical reason to support this theory; however many of these studies relied upon an activated, or modified, form of poly-*p*-phenylene terephthalamide that would readily form a chemically labile moiety in order to promote free radicals upon heat treatment. The Kevlar-29® studied in the study by Sweeney et al. was devoid of such activated species, and whether or not such crosslinking could occur within chemically unmodified PPTA specimens has not yet been validated. But if not due to the predominance of transaxial intermolecular covalent crosslinks within the core region, another structural

phenomenon must account for the markedly improved compressive properties of the Kevlar® fibers that had been treated at 400, 440, and 470° C.

Sweeny ¹⁰ claimed that a high degree of crosslinking could not be accommodated within the current crystal structure and without disruption of intermolecular hydrogen bonding. It is likely that the secondary, or radially oriented pleated sheet, structure of Kevlar® poses a steric limitation to the development of crosslinks within the core region of the PPTA. Moreover, in a study which characterized the pyrolytic behavior of aramid fibers, Mosquera et al. ¹³ concluded that below 500 °C, only minor changes associated with loss of hydrogen bonds is prevalent, and that no noticeable framework changes take place until 545 °C - the temperature at which notable chain scission begins.

As mentioned earlier in this work, Jiang et al. ⁷ concluded that lateral covalent crosslinks had been established via thermal treatment based upon swelling trials. This group concluded that the enhancement of compressive properties was a direct result of crosslinking, exclusively. This swelling study involved placing a small section of a Kevlar® fiber on a glass slide and introducing the polymer to a few drops of sulfuric acid. Solubility of the PPTA fiber was then evaluated. Fibers treated above 330 °C were characterized as insoluble and this finding was attributed to bulk crosslinking within the fiber. However, this conclusion does not take into consideration the tertiary structure of the Kevlar® fiber. It is quite possible that the skin region, devoid of a highly crystalline packing order, was crosslinked while the core region was not. A crosslinked skin region could possibly render the fiber insoluble by creating an impermeable sheath around a highly linear and uncrosslinked core region.

The correlation between crystallite perfection and tensile behavior of Kevlar® is well documented. Barton 14 has studied x-ray peak shape parameters for PPTA fibers variously treated to produce a range of tensile moduli. He demonstrates an excellent correlation between the axial paracrystalline distortion parameter and the tensile modulus. From this work, he concludes that overall crystal perfection is the controlling feature for the tensile modulus. Additionally, Hindeleh and cohorts 15 report a correlation between tensile modulus and transverse crystallinity, which is highly dependent upon hydrogen bonding. These studies illustrate the strong relationship between the tensile behavior of the PPTA fibers and its transverse and axial crystalline perfection.

Accordingly, the relationship between crystallite perfection and behavior in compression should not be overlooked.

Lee et al. ¹² validated that when Kevlar® fibers are subjected to a temperature of 400 °C under an applied low tensile force (.1 g/d), misalignment of the crystallites ensues. Secondly, the author asserts that misalignment of the crystallites increases with temperature and time of heat treatment, accompanied by a decrement in tensile modulus. Logically, misalignment would be even more pronounced when such a tensile force is absent. Lee et al. ¹² also discovered that upon heat treatment under tension, the pleating conformation begins to vanish, the de-pleated region working inward from the skin, or boundary region. Thus, the decrease in tensile strength and improvement in compressive behavior observed by Sweeney et al. ¹¹ could potentially be attributed to the aforementioned spatial distortion of the crystallites resulting from the destruction of interchain hydrogen bonding and concurrent destruction of pleated sheets within the core.

While the theory that covalent crosslinks between crystallites would limit structural maneuverability and therefore reduce creep abounds, the role of hydrogen bonding has been largely ignored in articles addressing induced thermal crosslinking of Kevlar® microfibrils. Microfibril/fibril buckling within high performance fibers is directly related to compressive strength (σ_c) according to the following equation:

$$\sigma_c = C \pi E (R/L)^2$$

where C is an empirical constant; E, the axial rigidity of the test specimen; R, the radius of the fiber; and L, the length of the fiber.⁸

It is reasonable to believe that any compromise of Kevlar's® nearly perfect uniaxial orientation, in particular the misalignment of crystallites, would correlate with an increase in axial rigidity (E) and, ultimately, an increase in compressive strength. Since transaxial and axial alignment of crystallites is contingent upon interchain hydrogen bonding, the destruction of these bonds would likely yield an improvement in compressive strength. Without the existence of transaxial bonds, the polymer chains are unlikely to orient uniformly into pleated sheets, and would consequently develop a chiasmatic or crossing orientation. The resulting chiasmas, or plexuses formed at the treatment temperatures studied may very well account for the observed increase in structural rigidity and compressive strength. Moreover, misorientation of the crystal structure comprising the pleated sheet configuration within the core structure would result in an increase in the radius of the fiber, thereby decreasing compressive

deformation by increasing the diametric area upon which to distribute a compressive load.

Clearly the theory that hydrogen bond disruption may account for measurable crystal misalignment and, consequently, enhanced compressive strength and lower tensile strength, warrants legitimate consideration. The objective of this work was to determine which structural change, crosslinking and/or hydrogen bond disruption, was responsible for the enhancement in compressive strength of the Kevlar® fibers heat-treated by Sweeney et al.¹¹

4.0 Experimental

Treated and untreated Kevlar-29® fibers were subjected to a variety of analyses in order to elucidate the structural mechanism by which the compressive strength of the Kevlar-29® studied by Sweeney et al. 11 had been enhanced. The treated Kevlar-29® fibers were those prepared by Sweeney et al. 11 in a Thermodyne oven under nitrogen purge at temperatures of 400, 440, and 470 °C. These PPTA fibers were characterized by the Thermogravimetric Analyzer (TGA), Differential Scanning Calorimeter (DSC), and Fourier-transform Infrared Spectrophotometer (FTIR) illustrated in Figures 4, 5, and 6.

Thermogravimetric experiments were conducted using a Mettler-Toledo TG-50 TGA under an inert nitrogen purge in an effort to quantify mass loss associated with crosslinking at elevated temperatures. Differential Scanning Calorimetry assays were performed using a Mettler-Toledo DSC25 with a TC15 TA controller in order to observe thermal transitions (endo or exotherms) within the polymer. A Fourier Transform infrared spectrophotometer was used to investigate specific peak intensities of the heat treated Kevlar-29® fibers. The FTIR was a Perkin-Elmer Spectrum One spectrophotometer with HATR assembly. All requisite calibrations were performed prior to characterization of the heat-treated Kevlar-29®.

The purported thermally induced crosslinking phenomenon would most likely occur through a free radical reaction. Based upon bond dissociation energies, the most

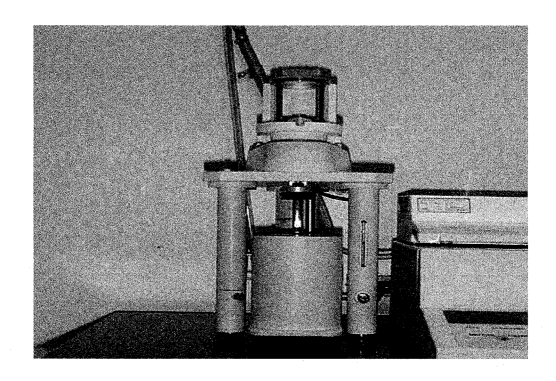


Figure 4. Thermogravimetric Analyzer

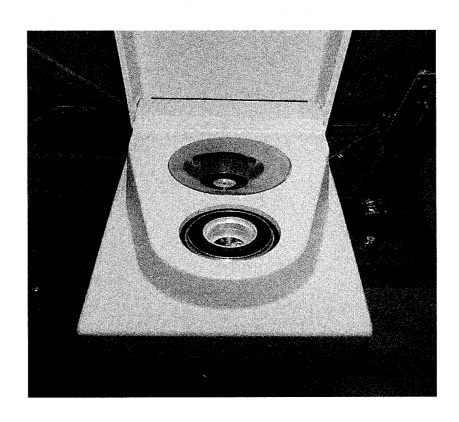


Figure 5. Differential Scanning Calorimeter

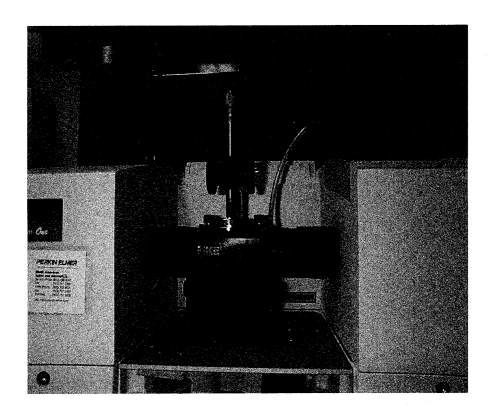


Figure 6. Fourier Transform Infrared Spectrophotometer

likely location of the free radical formation is at the ortho and meta positions of the aryl group. This site is indicated in Figure 7.

In order to establish a single covalent crosslink between the Kevlar® macromolecules, dehydrogenation must occur. This would involve the loss of two hydrogen atoms from an aryl ring component. In order to achieve the level of lateral enhancement witnessed in the unmodified, heat-treated Kevlar® study, dehydrogenation would need to occur on a widespread scale within the fiber. Quantification of this statement is difficult. However, at least one crosslink per nine repeat units could account for a crosslink density capable of achieving improved compressive strengths. Hydrogen losses at these crosslink densities should result in a detectable mass loss that may be captured via TGA measurements. Detection is likely given the sensitivity of the microgram TGA balance, which is capable of detecting a mass loss as miniscule as one-millionth of a gram.

Untreated Kevlar-29® fibers were subjected to TGA isotherms at select temperatures in order to quantify mass loss and gain insight into the initiation temperature necessary to promote significant levels of crosslinking within the PPTA fibers. The ramping rate used in these TGA experiments was 100 °C/minute. During the ramp, an empty crucible was situated upon the microbalance. When the target temperature was reached, the balance was zeroed and a 2 – 5 mg. sample of Kevlar-29® was then placed within the crucible. The time at temperature for isothermal experiments was invariably 23 minutes. Recall that the earlier clinic experiment which served as a background for this paper had utilized three levels for the time factor: 5, 7, and 10 minutes soak times. Therefore, the selected isothermal treatment times would be more

$$+C \longrightarrow C \longrightarrow H \longrightarrow H$$

Figure 7. Allylic Site for Free Radical Formation

than adequate to initiate the structural changes induced within the earlier experiment by Sweeney et al.¹¹ All isotherms were performed in an inert nitrogenous environment to preclude side reactions such as oxidation.

Hopes of improving the compressive properties of Kevlar® must recognize, of course, that a high degree of crosslinking could not be accommodated without affecting the efficiency of the interchain hydrogen bonding. ¹⁰ In order to discover what structural change (crosslinking and/or hydrogen bond disruption) was occurring within the PPTA fibers at elevated temperatures, a DSC scan was performed on 5 mg. of untreated Kevlar-29® fibers. The temperature sweep began at 25 °C and ended at 600 °C, and entailed a slow heating rate of 5 °C/minute in order to capture the structural change. Both the reference and sample DSC pans were heated under an inert nitrogen purge of 1 L/minute.

Fourier Transform Infrared Spectrophotometry analysis was also performed in an effort to elucidate the structural mechanism accounting for the compressive improvements of Kevlar-29® fibers treated at 400, 440, and 470 °C. According to the chemical formula and structure of PPTA, the following IR band assignments can be made: 1018 cm⁻¹ for an in-plane, C-H vibration, characteristic of para-substituted aromatic compounds, particularly polyaramids. Given the location of free-radical formation of Kevlar-29® fiber, significant crosslinking should be manifested as a decrease in intensity for this band. The band assignment for out-of-plane C-H vibrations of two adjacent hydrogens in an aromatic ring (para substitution of the aromatic) would be 827 cm⁻¹. The loss of hydrogen-bonded amide functionality (3432 cm⁻¹) was also assessed to estimate the degree of interchain hydrogen bond destruction. FTIR scans were performed in reflectance mode, applying a wavenumber range of 650 – 3600 cm⁻¹.

64 total scans were performed for each of the three assays, corresponding to the three different maximum treatment temperatures employed by Sweeney et al. 11

5.0 Results and Discussion

As detailed earlier, if free radicals were being created on a widespread scale within the core region of the Kevlar® fibers at temperatures of 400, 440, and 470 °C, the TGA with its sensitive microbalance should be able to capture and quantify such a mass loss. This mass loss would correlate to the dehydrogenation of aryl hydrogens during the formation of free radicals that serve as a precursor to interchain crosslinks. Greater mass loss would correspond to higher imposed crosslink densities.

Interestingly, the isotherm at 400 °C indicated that no detectable decrement in mass had occurred during the 23 minute interval within the untreated, as-spun Kevlar-29® fibers. This result is illustrated in Figure 8. Accordingly, an isotherm temperature was then selected beyond 470 °C, the highest temperature level enacted by Sweeney et al. ¹¹ An isotherm at 480 °C demonstrated that no mass loss of untreated Kevlar-29® fiber had occurred within the 23 minute interval. Thus neither the isotherm at 400 or at 480 °C, each with an aggressive holding time of 23 minutes, resulted in substantive mass losses according to the TGA analyses performed at these conditions. It is worth noting, however, that the samples that had undergone these isotherms experienced a noticeable color change, from yellow (virgin Kevlar-29® fiber) to dark brown/black (post-isotherm Kevlar-29® fibers).

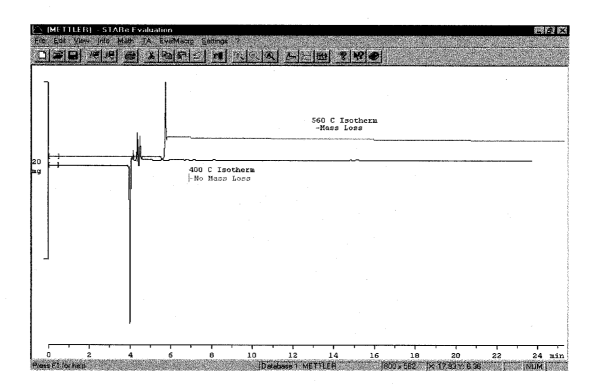


Figure 8. Thermogravimetric Isotherms of Kevlar-29® Fiber

Recall that Mosquera¹³ had purported that Kevlar® is thermally stable up to about 545 °C; below this temperature only minor change associated with the loss of hydrogen bonds occur. In order to test the validity of this claim, an isotherm at 540 °C was first performed. The results demonstrated that no significant mass loss had occurred within the PPTA fibers during the 23 minute hold time. Secondly, a temperature beyond the critical 545 °C mark was selected, utilizing the same treatment interval. Remarkably, an isotherm at 560 °C did indicate a mass loss of 0.1 mg., or 4.5%. The thermogram depicting this higher-temperature isotherm is also included in Figure 8.

These thermogravimetric findings support the claim by Mosquera¹³ that no significant mass loss occurs within Kevlar® fibers below 545 °C. If crosslinking is initiated at treatment temperatures of 400, 440, and 470 °C, this structural change is not accompanied by appreciable or detectable mass loss, according to the predescribed results obtained on the Mettler-Toledo TG-50 TGA. This raises doubt as to whether the structural mechanism accounting for enhanced compressive strength within heat-treated Kevlar® fibers is primarily, or exclusively, crosslinking.

If crosslinking was the predominant mechanistic change accounting for the enhancement of compressive properties within thermally-treated Kevlar-29®, the formation of crosslinks should result in a detectable exotherm within the DSC profile. Crosslink formation is exothermic due to the thermodynamically favorable conformation that such interchain covalent bonds would promote. Conversely, the destruction of interchain hydrogen bonding and thus the pleated sheet structure would be captured as a DSC endotherm. As a result, the DSC serves as an excellent tool for determining which

structural change, crosslinking or hydrogen bond destruction, is accounting for improved compressive properties and compromised behavior in tension.

The DSC scan, shown in Figure 9, depicts a gradual exotherm that begins at approximately 380 °C and continues until a temperature of 540 °C is achieved. It is likely that this exotherm corresponds to a crosslinking event within the Kevlar-29® fibers. According to the DSC thermogram generated within this study, the maximum crosslinking rate commences at approximately 470 °C.

Above 540 °C, a large endotherm was observed in the DSC scan, most likely corresponding to degradation of the PPTA and the formation of pyrolytic products such as hydrogen cyanide, benzene, toluene and benzonitrile. The endotherm present at 540 °C, presumably representative of the onset of pyrolysis of the PPTA fibers, mirrors the earlier TGA result that also indicated degradation of the aramid fiber accompanies this treatment temperature.

Crosslinking in polymers is an exothermic phenomenon due to the thermodynamically favorable conformation achieved by the formation of an interchain lattice structure. When order within the polymer structure is decreased as a result of thermal treatment, as is the case during polymer melting, an endothermic event ensues. Therefore, the destruction of interchain hydrogen bonds can be characterized as an endothermic event within the DSC spectra. No such endothermic event was detected in the DSC spectra of untreated Kevlar-29® fiber. Perhaps hydrogen bond destruction was below the level of detection of the calorimeter, or simply not prevalent to such a degree as to elicit a distinct endotherm. Conversely, a clear exotherm was identified and supports the theory that crosslinking is occurring within the polymer chain. It is possible

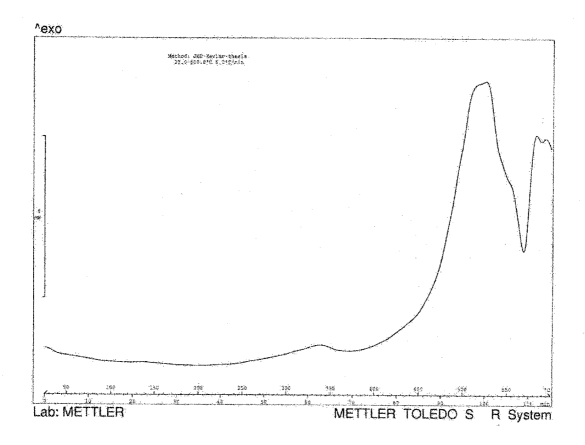


Figure 9. DSC Thermogram of Kevlar-29® Fiber

that an endo- and exothermic event is occurring simultaneously, with the dominant thermodynamic change being detected by the DSC.

The TGA and DSC results indicate different, although perhaps not mutually exclusive, information as to the structural mechanism accounting for improved compressive properties in heat-treated Kevlar-29® fibers. The FTIR, however, is arguably the most sensitive technique for discerning if crosslinking and/or the entropic effect of hydrogen bond obfuscation are taking place within the polymeric fibers.

The peak intensity of Kevlar-29® fibers at approximately 824 cm⁻¹, corresponding to the out-of-plane C-H vibration, decreased with increasing treatment temperatures. Similarly, the peak intensity at 1018 cm⁻¹, reflecting the in-plane aryl C-H vibration, decreased within increasing treatment temperatures. This indicates that the loss of hydrogen atoms from the benzene ring increases with increasing treatment temperature. This finding may be explained by the formation of free radicals in the initiation stage of crosslink formation.

A clear decrement in peak intensity and area was observed with increasing treatment temperatures at an approximate wavenumber of 3432 cm⁻¹, corresponding to the loss of hydrogen-bonded amide functionality. This result indicates that interchain hydrogen bond destruction is occurring within the Kevlar-29® fibers that had been treated at 400, 440, and 470 °C. The FTIR scan illustrates that both hydrogen bond destruction and crosslinking is taking place simultaneously at the treatment temperatures utilized by Sweeney et al.¹¹ A split view of the FTIR spectra is depicted in Figure 10.

Summarily, the results of the TGA thermogram indicate that no significant mass loss corresponding to dehydrogenation of aryl hydrogens is realizable at temperatures

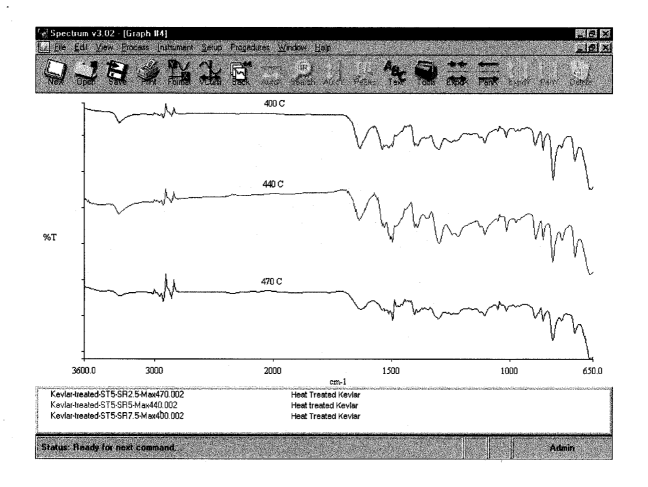


Figure 10. FTIR spectra of Heat-treated Kevlar-29® Fibers

below 560 °C, supporting the theory that hydrogen bond destruction is accounting for the observed increased compressive and decreased tensile properties within heat-treated PPTA fibers. Conversely, the results of the DSC thermal profile indicate that crosslinking is the mechanism accounting for the observed changes in mechanical properties. The FTIR scans indicate that both structural changes arise when Kevlar-29® is heat treated at thermal treatments of 400, 440, and 470 °C.

Initial impressions may lead one to conclude that these results are irreconcilable. However, an understanding and appreciation of the tertiary structure of poly-*p*-phenylene terephthalamide may provide a plausible interpretation for the findings detailed within this work.

It is likely that both structural changes are promoted within different regions of the polymer when exposed to elevated temperatures. The microstructure comprising the skin region may be crosslinking, while the inner core region experiences an entropic effect of hydrogen bond destruction and misorientation of the crystallites within the pleats.

It is feasible that crosslinking is not detected through TGA analysis because the bulk of the fiber is experiencing hydrogen-bond destruction. Thus, dehydrogenation of the aryl hydrogen is below the level of detection of the TGA. The FTIR analysis, when performed in reflectance mode, would detect the crosslinking within the skin region as the beam must penetrate the skin. Additionally, the infrared beam would partially penetrate the core region as well before experiencing full reflection, thus capturing the structural change within the core region as well. During thermal profiling via DSC analysis, heat would penetrate the fiber from the boundary inward, and therefore

crosslinking within the skin region is the dominant thermodynamic change captured by this instrument. These analyses have demonstrated that thermal enhancement of Kevlar-29® is achieved through both structural changes, crosslinking and hydrogen bond destruction, working in concert to promote the augmentation of compressive strength.

6.0 Conclusions

The thermal enhancement of compressive properties in Kevlar-29® fibers is achieved through two mechanisms: (1) free-radical formation within the external skin region of PPTA, resulting in the formation of interchain crosslinks and (2) hydrogen-bond disruption resulting in the destruction of the highly ordered, pleated sheet configuration within the core region of PPTA. Interchain covalent crosslinks improve the compressive strength of the aramid fiber by limiting the structural maneuverability of the fiber skin, thus augmenting its rigidity. Hydrogen bond disruption within the core region results in misorientation of crystallites within the core region, thus enhancing the compressive properties of Kevlar® by compromising its nearly perfect linearity.

Several high-temperature Thermogravimetric isotherms were undertaken in order to characterize the degree of dehydrogenation within the PPTA fibers, corresponding to the formation of covalent, interchain crosslinking between the aryl moieties. No significant mass loss was detected at a temperature of 540 °C or below. Substantive mass loss was discernible at a temperature of 560 °C, which can be attributed to pyrolysis of the Kevlar-29® fiber. The TGA results indicate that no detectable mass loss related to the bulk development of crosslinking throughout the fiber occurs at elevated temperatures as high as 540 °C.

The Differential Scanning Calorimetry profile did not indicate the presence of a distinct endotherm at elevated temperatures below 540 °C. Therefore, a distinct event relating to hydrogen bond disruption within Kevlar-29® fiber was not captured by the DSC. In contrast, DSC indicates that a crosslinking event is occurring within heat-treated Kevlar-29® fiber. This is evidenced by a gradual exothermic event that develops at a treatment temperature of approximately 380 °C and continues until a temperature of 540 °C is achieved. At a temperature of 540 °C, a clear endotherm is observable, most likely corresponding to degradation of the polymer into volatiles such as hydrogen cyanide, benzene, toluene, or benzonitrile. The DSC endotherm supports the earlier pyrolytic characterization of aramid fiber by the TGA thermogram.

Fourier Transform Infrared Spectrophotometric assays of heat-treated Kevlar-29® fiber provide the greatest insight into what structural changes account for the improved behavior of the polymer in compression. The FTIR indicates that both structural mechanisms, crosslinking and hydrogen bond destruction, account for the improved compressive properties of the polymer. FTIR spectra of Kevlar-29® subjected to temperatures of 400, 440, and 470 °C demonstrate a decrease in hydrogen bond and aryl hydrogen functionality with increasing temperature.

A cursory evaluation of these results may lead the reader to deem them incompatible. However, with an appreciation of the tertiary structure of PPTA and test methodology utilized within the scope of this text, these results are resolvable. It is likely that both the entropic effect of hydrogen bond disruption and interchain crosslinking are promoted within heat-treated poly-*p*-phenylene terephthalamide, albeit in distinct regions

of the polymer matrix. The anisotropic microstructure comprising the skin is crosslinked, while the core region characterized by a pleated sheet configuration experiences the entropic effect of hydrogen bond disruption and a concurrent misorientation of the crystallites. Crosslinking of the skin region and misorientation of core crystallites serve as the structural mechanisms that promote the enhancement of compressive strength of heat-treated Kevlar-29® fiber.

7.0 Recommendations

This work highlights the need to differentiate between the distinct morphologies of the core and skin regions of heat treated Kevlar-29® fiber. The author utilized various thermal and non-thermal analytical techniques to elucidate the structural changes occurring within the PPTA fibers. Therefore, a logical extension of this work would employ other characterization techniques, such as Scanning Electron Microscopy, Optical Microscopy, or X-ray Diffraction, to characterize the core and skin regions individually. Most of the non-thermal characterization of Kevlar® has focused on the bulk polymer as an isotropic material rather than dissecting the polymer in terms of a skin and core region with unique morphologies. The current work demonstrates the need to treat and examine the skin and core regions of heat-treated Kevlar® fibers as separate morphological entities, each providing a unique contribution to the improved compressive properties of the aramid fiber.

Works Cited

- 1. DuPont Kevlar® Technical Guide (2000) [Brochure]. DuPont Advanced Fiber Systems.
- Tanner D, Fitzgerald J A and Phillips B R. "The Kevlar® Story-an
 Advanced Materials Case Study." <u>Advanced Materials</u> 5 (1989): 151-156.
- Newell J A, Puzianowski A A, and Schmidt L R. "The Influence of Electron Beam Radiation on the Tensile and Recoil Compressive Strengths of Kevlar-29". Accepted for Publication in <u>High Performance</u> <u>Polymer</u>, 2002.
- 4. Zhang Q, Liang Y and Warner S B. "Partial Carbonization of Aramid Fibers." Journal of Polymer Science 32 (1994): 2207-2220.
- 5. Rao Y, Waddon A J and Farris R J. "The Evolution of Structure and Properties in Poly(*p*-phenylene terephthalamide) Fibers." Polymer 13,42 (2000): 5937-5953.

- 6. Graham J F, McCague C, Warren O L and Norton P R. "Spatially
 Resolved Nanomechanical Properties of Kevlar® Fibers." Polymer 41,12
 (2000): 4761-4768.
- 7. Jiang T, Rigney J, Jones M, Markoski L, Spilman G, Mielewski D, and Martin D C. "Processing and Characterization of Thermally Cross-Linkable Poly(p-phenyleneterephthalamide-*co-p-*1,2-dihydrocyclobuta-phenyleneterephthalamide) Copolymer Fibers." <u>Macromolecules</u> 28 (1995): 3301-3312.
- 8. Kumar S and Helminiak T E. "Compressive Strength of High Performance Fibers." SAMPE Journal 26,2 (1990): 51-61.
- 9. Demmler A W. "Kevlar® and Carbon Composites Compared."

 <u>Automotive Engineering</u> 93,2 (1985): 44-52.
- Sweeny W. "Improvements in Compressive Properties of High Modulus
 Fibers by Crosslinking." <u>Journal of Polymer Science: Part A: Polymer</u>
 <u>Chemistry</u> 30 (1992): 1111-1122.
- 11. Sweeney D, Newell J, Picerno S, and Kurzeja T. "Influence of Thermal Treatment Conditions on the Recoil Compressive Strength of Kevlar-29® Fibers." Accepted for Publication in <u>High Performance Polymer</u>, 2002.

- 12. Lee K G, Barton Jr. R, and Schultz J M "Structure and Property

 Development in Poly(p-phenyleneterephthalamide) During Heat

 Treatment under Tension." Journal of Polymer Science: Part B: Polymer

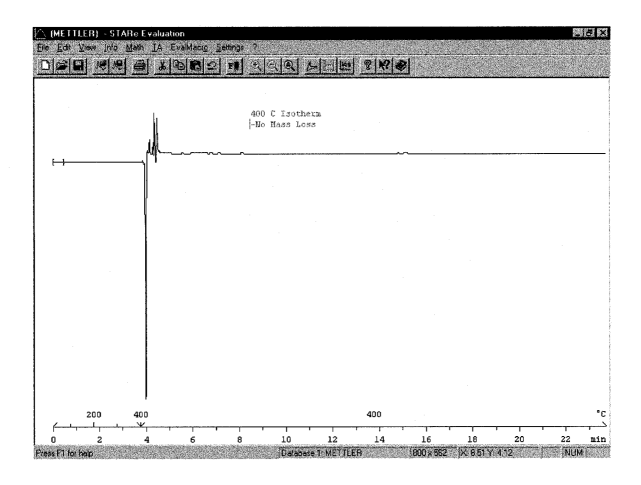
 Physics 33 (1995) 1-14.
- M Mosquera, Jamond M, Martinez-Alonso A, Tascon J. "Thermal
 Transformations of Kevlar® Aramid Fibers during Pyrolysis." <u>Chem.</u>
 <u>Mater.</u> 6,11 (1994) 1918-1924.
- 14. R Barton "Paracrystallinity-Modulus Relationships in Kevlar® Aramid Fibers." J. Macromol. Science Physics 24, 1-4 (1985) 119-130.
- 15. Hindeleh A M, Halim N A, and Ziq K A "Solid-State Morphology and Mechanical Properties of Kevlar-29® Fiber." J. Macromol. Science Physics 23, 3 (1984) 289-309.
- Northolt, M G "X-Ray Diffraction Study of Poly(p-PhenyleneTerephthalamide) Fibers." <u>European Polymer Journal</u> 10 (1974) 799-804.
- 17. Panar M, Avakian P, Blume R C, Gardner H, Gierke T D, and Yang H H

 "Morphology of Poly(p-Phenylene Terephthalamide) Fibers." <u>Journal of</u>

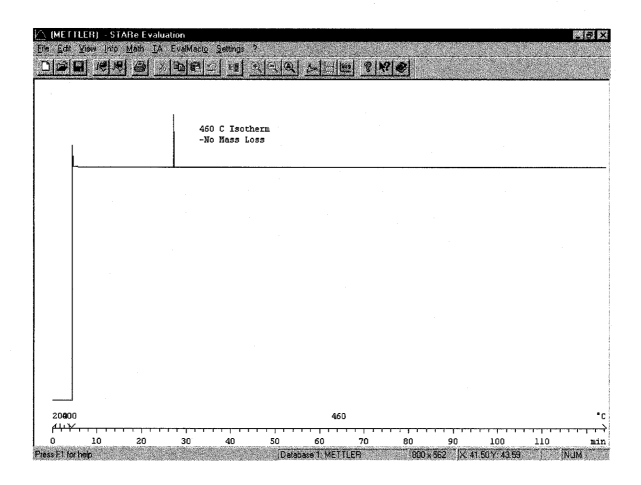
 <u>Polymer Science: Polymer Physics Edition</u> 21 (1982) 1955-1969.

- 18. Dobb M G, Johnson D J, and Saville B P "Supramolecular Structure of High-Modulus Polyaromatic Fiber (Kevlar-49®)" <u>Journal of Polymer Science</u> 15 (1977) 2201-2211.
- 19. Dobb M G, Johnson D J, and Saville B P "Microvoids in Aramid-Type Fibrous Polymers." Polymer 20 (1979) 1284-1288.

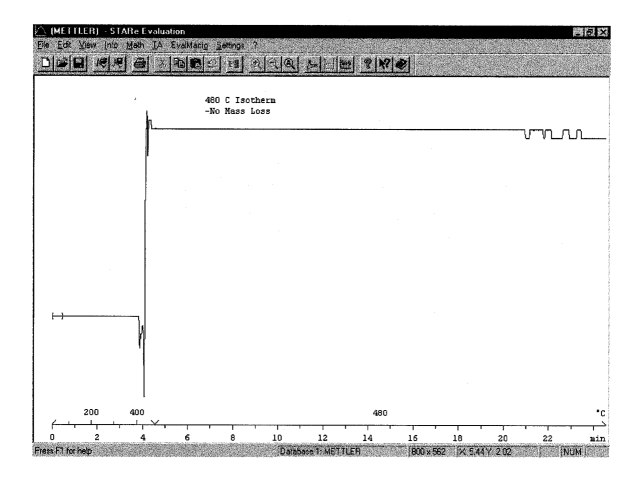
APPENDIX I



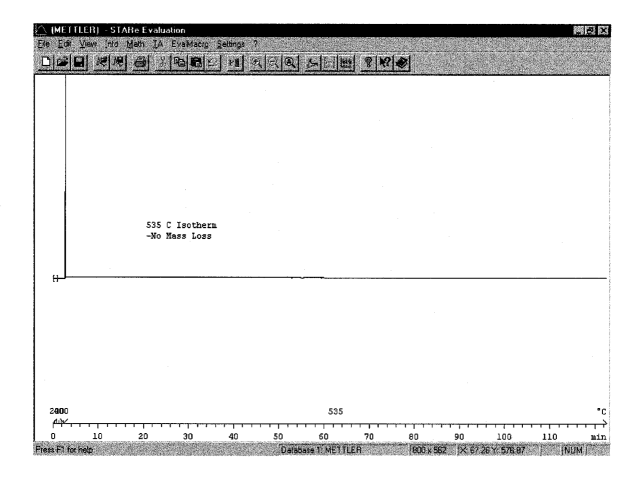
Thermogravimetric 400 °C Isotherm of Kevlar-29®



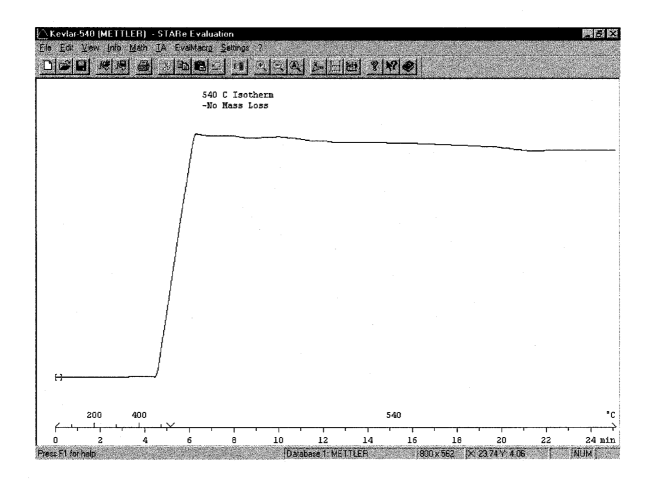
Thermogravimetric 460 °C Isotherm of Kevlar-29®



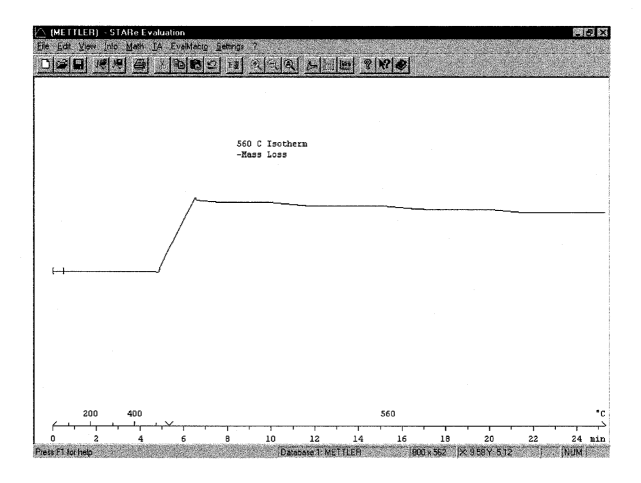
Thermogravimetric 480 °C Isotherm of Kevlar-29®



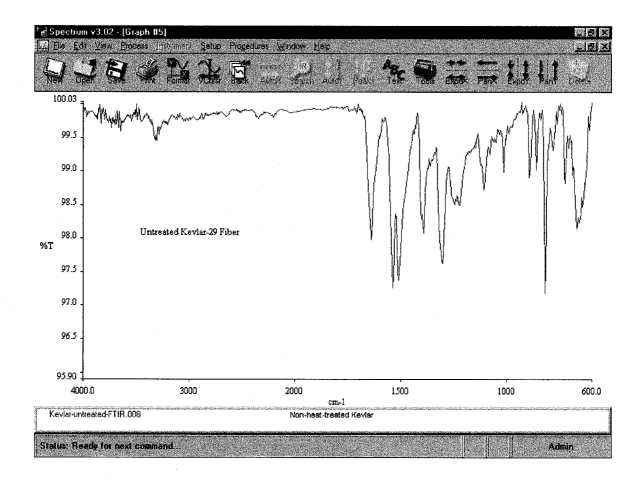
Thermogravimetric 535 °C Isotherm of Kevlar-29®



Thermogravimetric 540 °C Isotherm of Kevlar-29®



Thermogravimetric 535 °C Isotherm of Kevlar-29®



FTIR Spectrum of Untreated Kevlar-29®