CHARACTERIZATION OF DYNAMIC AND STATIC MECHANICAL BEHAVIOR OF POLYETHERIMIDE

by

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ABSTRACT

Polymers are increasingly being used in engineering designs due to their favorable mechanical properties such as high specific strength, corrosive resistance, manufacturing flexibility. The understanding of the mechanical behavior of these polymers under both static and dynamic loading is critical for their optimal implementation in engineering applications. One such polymer utilized in a wide variety of applications from medical instrumentation to munitions is Polyetherimide, referred to as Ultem. This thesis characterizes both the static and dynamic mechanical behavior of Ultem 1000 through experimental methods and numerical simulations. Standard compression experiments were conducted on and MTS test frame to characterize the elastic-plastic behavior of Ultem 1000 under quasi-static conditions. The dynamic response of the material was investigated at very high strain rates using a custom built miniaturized Kolsky bar apparatus. The smaller Kolsky bar configuration was chosen over the conventional Kolsky device to increase the maximum capable strain rates and to reduce common experimental problems such as wave dispersion, friction, and stress equilibrium. Since a universal test standard for this apparatus is not available, the details of the design, construction, and experimental procedures of this device are provided. The results of the high strain rate testing revealed a bilinear relationship between the material yield stress and strain rate. This relationship was modeled using the Ree-Eyring two stage activation process equation.

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LIST OF NOMENCLATURE

ε	True strain
Ė	Strain rate
$\varepsilon_I, \varepsilon_R, \varepsilon_T$	Incident, reflected, and transmitted strain pulses
θ	Absolute temperature
$ ho_b, ho_s$	Density of bar and specimen
σ	True stress
σ_y	Yield stress
τ	Ring up period
A_1, A_2	Ree-Eyring coefficients
A_b, A_s	Area of bar and specimen
C_1, C_2	Ree-Eyring coefficients
C_b, C_s	Elastic wave speed of bar material and specimen
D_b, D_s	Diameter of bar and specimen
е	Engineering stress
E_b, E_s	Elastic modulus of bar and specimen
L_b, L_s, L_{st}	Length of impact/transmission bar, specimen, and striker bar
Q_1, Q_2	Activation energies
R	Universal gas constant
Т	Period of incident pulse
v_{st}	Velocity of striker bar

1. INTRODUCTION AND MOTIVATION

It is well known that the mechanical properties of many engineering materials are dependent on the rate of deformation to which the material is subjected. This dependence is more pronounced for certain materials such as polymers and at very high strain rates $(10^2 - 10^4 \text{ s}^{-1})$. Due to the increase of use of polymers in many engineering designs, it is desirable to accurately characterize the behavior of these polymers at very high strain rates. One widely employed method of experimentally determining the high strain rate response of materials is by using a Split Hopkinson Pressure Bar (SHPB) or Kolsky bar device. Such pressure bar devices have been used by various researchers since the pioneering work of Bertram Hopkinson in 1914 [Hopkinson, 1914]. The present work investigates the high strain rate mechanical behavior of Polyetherimide, also referred to as Ultem, through use of a miniaturized Split Hopkinson Pressure Bar (mSHPB). Ultem is a thermoplastic which is used in many static and dynamic engineering applications throughout various industries due to its high strength to weight ratio and other favorable characteristics, but has yet to be analyzed under dynamic conditions. In the defense industry, Ultem is employed in guided projectile designs, illustrated in Figure 1, where circular Ultem plates are designed to fail in a predictable manner under dynamic axisymmetric pressure.



Figure 1. Guided projectile application of Ultem 1000

Although Kolsky bars have been successfully used for decades, the devices are not commercially available and there is no formal standardized test method to guide researchers on the proper experimental setup, procedure, or data analysis. Furthermore, only a small fraction of the available literature on Kolsky bar testing is related to the miniaturized version of the Kolsky bars, which possess many advantages over the full-scale Kolsky bar system. As such, this thesis is also intended to serve as a detailed guide on the design, construction, calibration, and experimental procedure for a custom built miniaturized Kolsky bar device. All of the needed parts and materials are listed along with detailed engineering drawings of the various custom made components. The goals of this project were to design and fabricate an accurate Kolsky bar system within a relatively modest budget, characterize the strain rate dependence of Ultem 1000 under compression, and to contribute to the existing body of literature a guide for research groups to be able to design and construct their own Kolsky bar device more efficiently.

A review of the literature regarding the development of the Kolsky bar device, its improvements throughout the years, and its application to the characterization of polymers is given in Chapter 2. Chapter 3 covers the design, construction, and calibration of the miniaturized Kolsky bar system. The experimental setup and specific procedures for the quasistatic and high strain rate experiments are provided in Chapter 4. The results of the experiments are presented in Chapter 5 and discussed in Chapter 6. Finally, Chapter 7 contains the conclusions as well as planned future work regarding the miniaturized Kolsky bar system. Appendix A contains a detailed bill of materials and engineering drawings for the construction of the Kolsky bars. The codes developed for measuring the striker velocity (LabVIEW) and processing the experimental data (MATLAB) are provided in Appendix B and Appendix C, respectively. Photographs of the specimens used in the experiments as well as experimental data are provided in Appendix D.

2. BACKGROUND

2.1 Kolsky Bar Background

2.1.1 Kolsky Bar Theory

The original Kolsky bar device, constructed in 1949, was designed to subject specimens to high rate compression deformation. Typically, a compression Kolsky bar apparatus is comprised of three bars, each of the same material and diameter ranging between 10 and 25 mm. The three bars are referred to as the striker bar, the impact bar, and the transmission bar and are shown schematically in Figure 2. Prior to the start of an experiment, the specimen is sandwiched between the impact and transmission bars. Then the striker bar is given a velocity (v_{st}), usually by the expansion of compressed gas, and will strike the impact bar sending a compressive pulse through the impact bar. This compressive pulse, known as the incident pulse (ε_I), will travel through the impact bar at the elastic wave speed as defined by

$$C_b = \sqrt{\frac{E_b}{\rho_b}} \tag{1}$$

where E_b is the elastic modulus and ρ_b is the density of the bar material. The magnitude of this incident pulse is determined by

$$\varepsilon_I = \frac{1}{2} \frac{v_{st}}{C_b} \tag{2}$$

and the width of the pulse, T, is

$$T = \frac{2L_{st}}{C_b} \tag{3}$$

where L_{st} is the length of the striker bar. Once the pulse reaches the interface of the impact bar and specimen, a portion of the pulse will pass through the specimen and into the transmission bar and is referred to as the transmitted pulse (ε_T). The remainder of the original pulse will be reflected back through the impact bar as a tensile pulse and is called the reflected pulse (ε_R). An example of these pulses for a striker velocity of 23.8 m/s and striker length of 76.2 mm is shown in Figure 3. The incident and reflected pulses are measured by a strain gauge on the impact bar and the transmitted pulse is measured by a strain gauge mounted on the transmission bar.



Figure 2. Schematic of Kolsky bar apparatus



Figure 3. Example of incident, reflected, and transmitted pulses

A detailed derivation of the equations used to compute the instantaneous stress and strain within the specimen as a function of the incident, reflected, and transmitted pulses can be found elsewhere [Gray III, 2000; Chen, 2011]. The key relations used for the data reduction in Kolsky bar experiments for specimen strain rate ($\dot{\varepsilon}$) and stress (σ) as functions of time are

$$\dot{\varepsilon}(t) = 2\frac{C_b}{L_s}\varepsilon_R(t) \tag{4}$$

$$\sigma(t) = \frac{A_B}{A_S} E_B \varepsilon_T(t)$$
(5)

where L_S is the original specimen length, and A_B and A_S are the cross sectional areas of the bars and specimen respectively. For Eqs. (4) and (5) to be valid, the specimen must be in stress equilibrium where the stresses at both ends of the specimen are equivalent. When the specimen is in stress equilibrium, the following relation is true.

$$\varepsilon_I(t) + \varepsilon_R(t) = \varepsilon_T(t) \tag{6}$$

However, there is a period of time for which the pulses have not yet traversed the entire specimen resulting in different values of stress at the specimen ends. During this time, known as the "ring-up period," the specimen is not in stress equilibrium and Eq.(4) and Eq. (5) are not valid. Therefore, Eq. (6) is employed to check for which times the specimen is in stress equilibrium. An example is presented in Section 5.2.1 detailing the procedures used for analyzing the strain history data and checking for stress equilibrium validity.

2.1.2 Kolsky Bar Evolution

Since the original implementation of the pressure bar technique to characterize the dynamic behavior of materials by Bertram Hopkinson in 1914, the devices and techniques used have been significantly modified and improved. One such improvement to the apparatus was made by Kolsky in 1949 where the single impact bar was replaced by two bars with the specimen placed in between them [Kolsky, 1949]. This device was known as the "split-Hopkinson pressure bar" or simply the Kolsky bar and is the foundation for many of the designs still in use today. The Kolsky bar apparatus has been further modified to facilitate loading modes other than compression, such as tension [Harding, 1960; Nicholas, 1981; Staab, 1991], torsion [Gilat, 2000], shear, triaxial [Nemat-Nasser, 2000] and unique combinations of these loading modes [Lewis, 1973]. The load history subjected to the specimen can be further controlled by a technique in which the momentum traveling through the bars is trapped after either one or multiple wave reflections [Nemat-Nasser, 1991]. Such a technique facilitates precise control of the load pulse shape, duration, and the number of loading cycles which is

critical for dynamic recovery experiments. Furthermore, material properties other than the stress-strain response have been investigated under high strain rate conditions such as dynamic indentation and fracture toughness properties [Klepaczko, 1980].

The Kolsky bar apparatus has also been modified to test a wide range of materials including concrete [Zhao, 1998], ceramics [Subhash, 2000], and soft materials such as foam, and polymers [Gray III and Blumenthal, 2000]. The testing of soft or low impedance materials requires special considerations, especially when using polymer materials for the bars, and a detailed discussion on the topic is provided in [Gray III and Blumenthal, 2000]. Researchers have also developed methods to heat or cool the specimen just prior to testing so the dual effects of temperature and strain rate can be investigate [Frantz, 1984; Gray, 1997].

In addition to modifications made to the physical device, improvements have been made to the data acquisition and analysis processes for increased accuracy in experimental results. Since in an actual Kolsky bar experiment a longitudinal pulse is traveling through a 3D rod, a phenomenon known as wave dispersion was found to adversely affect the accuracy of the collected data [Davies, 1948]. Pochhammer [Pochhammer, 1876] and Chree [Chree, 1889], were the first to solve for the propagation of waves through cylindrical bars, but Davies [Davies, 1948] was the first to apply their solutions to the Kolsky bar technique. Since then various researchers have developed mathematical techniques to correct the measured pulses for wave dispersion [Follansbee, 1983; Gong, 1990; Gorham, 1983; Lifshitz, 1994; Tyas, 2005; Yew, 1978].

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2.2 Material Background

2.2.1 Polyetherimide

The material under investigation, Ultem 1000, is an amorphous thermoplastic without fillers or other fiber reinforcements. Ultem 1000 is composed of repeating units of $C_{37}H_{24}O_6N_2$ which is shown in Figure 4. Amorphous refers to the random orientation of polymer chains shown schematically in Figure 5. The molecular organization and interaction of these chains determines the mechanical response of polymers much like the crystalline structures in metallic materials. Since Ultem 1000 is used in a range of applications ranging from aerospace to medical industries, the quasi-static mechanical properties as well as fatigue, creep, and wear properties have been previously characterized [Bijwe, 1990; Facca, 2006; Smmazcelik, 2008; Stokes, 1988; Tou, 2007]. Selected mechanical and thermal properties of Ultem 1000 are provided in Table 1.



Figure 4. Chemical composition of Ultem 1000 [Pecht, 1994]



Figure 5. Schematic of amorphous polymer chains

Mechanical Properties	Value (English)	Value (SI)
Tensile Modulus, E_t	475 ksi	3.28 GPa
Compressive Modulus, E_{o}	480 ksi	3.31 GPa
Flexural Modulus, E_{f}	500 ksi	3.45 GPa
Poisson's Ratio, v	0.36	-
Elongation (Yield), ϵ_{y}	7.0 %	-
Tensile Strength, σ_{ut}	16.5 ksi	113.8 MPa
Compressive Strength, σ_{uc}	22 ksi	151.7 MPa
Shear Strength, σ_{su}	15 ksi	103.4 MPa
Flexural Strength, σ_{uf}	20 ksi	137.9 MPa
Elongation (Fracture), $\epsilon_{\!f}$	60 %	-
Izod Impact Strength, Notched	1.0 ft-lbs/in	0.034 J/m
Rockwell Hardness	109 ("M" Scale)	-
Physical Properties	Value (English)	Value (SI)
Specific Gravity	1.28	-
Thermal Properties	Value (English)	Value (SI)
CLTE - Flow, af	0.000031 in/in/°F	-
CLTE - Tansverse, a_t	0.000030 in/in/°F	-

Table 1. Mechanical and thermal properties of Ultem 1000 [Mutter, 2010]

2.2.2 Rate Dependent Polymer Characterization

The rate-dependent behavior of polymers has received relatively little attention in comparison to engineering metals over the years; however, since plastic materials are replacing metals in many engineering designs, the dynamic stress-strain response of these materials is of considerable interest. A brief review of past investigations of polymer rate dependency is presented here, but the discussion is limited to amorphous polymers as their temperature and rate dependent mechanical response differ from crystalline or semicrystaline polymers and since the candidate material (Ultem 1000) is an amorphous polymer.

Before the strain rate effects on material behavior can be understood, it is enlightening to review the typical quasi-static response of an amorphous polymer and the underlying molecular interactions. Also, it is helpful to clarify the definitions of some of the mechanical properties as determined from a stress-strain curve since these definitions differ from those commonly used in metals. To illustrate, a quasi-static compressive stress strain curve of Ultem 1000 is shown in Figure 6.



Figure 6. Quasi-static compressive stress-strain curve of Ultem 1000

The stress-strain curve in Figure 6 is marked with four distinct regimes classified by their macro-mechanical response, but they will be explained in terms of their molecular-level interactions. Unlike most metals, the slope of the elastic portion of the stress-strain curve for amorphous polymers is not constant, hence polymers are said to exhibit nonlinear elasticity. So the elastic region is simply divided into a small portion where the slope is constant (linear elastic) and another portion where the slope is constantly changing (nonlinear elastic). On a molecular level, the linear elastic phase is caused by the van der Waal forces between polymer chains resisting the deformation. Once the strain reaches a certain level (in the case of Figure 6, approximately 0.01) the polymer chains begin to slide with respect to one another and the response appears nonlinear. The most important characteristic of the stress-strain curve for the

purposes of the present work is the yield stress denoted in Figure 6. The yield stress defined for amorphous polymers is the local maximum in stress just after the elastic portions where the material deforms or flows without an increase in stress. This definition differs from that used in the analysis of metallic materials, but is widely accepted for polymers. The third phase (strain softening) has been the source of some debate as to whether it is caused by a local temperature rise [Marshall, 1954] or a permanent rearrangement of polymer chains with respect to one another [Brown, 1968]. However, for quasi-static testing conditions, the time scale needed for the temperature to equilibrate is smaller than the loading rate and the strain softening can be attributed to a permanent molecular rearrangement [Vincent, 1960]. Lastly, the strain hardening phase is a result of the once randomly oriented polymer chains aligning themselves in such a way requiring increased levels of stress for continued deformation.

Now that the quasi-static response is well defined, a brief review of the rate-dependent investigations of amorphous polymers is presented. The first researcher credited for conducting such investigations is Chou et al. [Chou, 1973]. He employed a Kolsky bar device and custom medium-strain rate apparatus to subject polymethylmethacrylate (PMMA), cellulose acetate butyrate (CAB), polypropylene, and nylon 6-6 specimens to strain rates ranging from 10^{-4} to 10^3 s⁻¹. Some of the results from this work are shown in Figure 7 for PMMA. As expected, the yield stress exhibited by the material increases with increasing strain rate (ε). Another example of this relationship between yield stress and strain rate is in Figure 8 where the rate dependent stress strain curves for polycarbonate (PC) are shown for rates from 10^{-5} to 10^4 s⁻¹.

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Figure 7. Rate dependent stress-strain curves for PMMA [Chou, 1973]



Figure 8. Rate dependent stress strain curves for PC [Siviour, 2005]

Furthermore, Walley and Field and others [Field, 1994; Walley, 1989; Walley, 1991] investigated a wide range of polymers, both at room temperature and elevated temperature, from rates of 10^{-2} to 10^4 s⁻¹. They, too, observed the relationship between the yield stress and the strain rate, but went further to classify materials into three groups based on the type of relationship observed between yield stress and log($\dot{\epsilon}$). The three groups are (1) materials that exhibit a linear relationship between yield stress and log($\dot{\epsilon}$), (2) materials that exhibit a bilinear relationship between yield stress and log($\dot{\epsilon}$), and (3) materials that exhibit a decrease in yield at approximately 10^3 s⁻¹ [Siviour, 2005]. An example of the bilinear dependence on log($\dot{\epsilon}$) is shown in Figure 9 for PC tested at room temperature.



Figure 9. Strain rate versus peak stress for PC [Siviour, 2005]

Unlike in the quasi-static case, the dynamic response of amorphous polymers requires the careful examination of both the temperature rise due to rapid plastic deformation and the

molecular rearrangement of polymer chains. The rate dependence of yield stress in amorphous polymers has been successfully modeled by the Eyring activation theory [Eyring, 1936] by many investigators [Bauwens-Crowet, 1969] for strain rates up to 10^{-2} s⁻¹. This model predicts a linear relationship between yield stress and $log(\dot{\epsilon})$ which fit the experimental data satisfactorily for Bauwens and others at the time. However, for the materials that exhibit a bilinear relationship between yield stress and $log(\dot{\epsilon})$, a modified model known as the Ree-Eyring equation was developed [Ree, 1955]. The Ree-Erying model, presented in Eq. 7, takes into account the simultaneous effect of two activation processes (denoted by the subscripts 1 and 2) on the yield stress.

$$\frac{\sigma_{y}}{\theta} = A_{1} \left(\ln(2C_{1}\dot{\varepsilon}) + \frac{Q_{1}}{R\theta} \right) + A_{2} \sinh^{-1} \left(C_{2} \dot{\varepsilon} e^{\frac{Q_{2}}{R\theta}} \right)$$
(7)

Here, A_i is a material parameter with units [Pa/°K], C_i is a material parameter with units [s], Q_i are the activation energies associated with each process [kcal/mol], R is the universal gas constant, and θ is the absolute temperature of the material. This dual-process characteristic of the Ree-Erying model predicts the bilinear behavior by assigning a greater weight to the contribution of the second process after a threshold strain rate has been reached. The threshold strain rate and the relative weight of the second process contribution are dictated by the values of C_2 and A_2 , respectively, and are determined from fitting experimental data.

3. DESKTOP KOLSKY BAR DESIGN, CONSTRUCTION, AND CALIBRATION

Although Kolsky bars have been used successfully for decades, there exists no standardized (ASTM, ISO, etc.) method for designing, constructing, and carrying out Kolsky bar experiments. This is because the design of the device is highly dependent on the specific application and because there is still much on-going research on how to solve many of the complications associated with this type of testing. Perhaps the most extensive set of guidelines are found in the ASM Handbook [Gray III, 2000] and by Chen and Song [Chen, 2011]. Even in these relatively comprehensive works, numerous references are made to other works which focus on specific technical issues such as wave dispersion, lubrication methods, and momentum trapping techniques. A discussion is presented in this chapter on the methodologies followed to design and construct the miniaturized Kolsky bar used for the experiments in the present work. Detailed explanations are given regarding design choices and fabrication techniques with the goal of providing a template for others to build and modify Kolsky bar devices. Additionally, an approach to modify the current Kolsky bar design to test other classes of materials, such as metals, is provided at the end of the chapter.

3.1 Design

Configuring the desktop Kolsky bar device is a process which is dependent on the specimen strength, length scale, strain rate, loading mode, and environment. The Kolsky bar device constructed for the present work is much smaller than the typical setup and is referred to as a miniaturized or desktop Kolsky bar. The overall length of the desktop apparatus is

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approximately 1 meter compared to approximately 6 meters for a full-scale setup and the bar diameters are 3.175 mm compared to the conventional diameter of 10-25 mm [Gray III, 2000]. There are numerous advantages to miniaturizing the Kolsky bars such as increasing the upper limit on strain rate, and reducing the negative effects of wave dispersion, friction, and inertia in the bars and specimen. A thorough investigation of these benefits was conducted by Jia and Ramesh [Jia, 2004].

The primary design consideration is the yield strength and stiffness of the candidate material as this will dictate the choice of bar material that must be used. The derivation of Eqs. (4) and (5) requires that the impact and transmission bars deform elastically. Hence, the yield strength of the bar material must be greater than the stress generated by the initial impact of the striker and impact bars which is given by

$$\sigma_I = \frac{1}{2} \rho_b C_b v_{st} \tag{8}$$

In addition to the bar strength, the bar stiffness must be considered with respect to the stiffness of the materials to be tested. If the bars are too stiff in comparison to the specimen material, the sensitivity of the output signals from the gauges will be reduced and complications in data reduction may arise due to the low signal magnitude. To increase the resolution of the output signals, investigators have used low impedance bar materials such as titanium alloys [Field, 2004], aluminum alloys [Chen, 1999], and polymeric materials [Sawas, 1998; Wang, 1994; Zhao, 1997]. The viscoelastic behavior of polymeric bar materials complicates the data analysis because the simple linear-elastic wave propagation equations are not directly applicable. Therefore, for the Kolsky bar apparatus under consideration, Aluminum 7075-T6 was chosen as the bar material with a yield strength of 190 MPa and a modulus of elasticity of 69.0 GPa. An

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aluminum alloy provides a sufficient signal to noise ratio for accurate data acquisition as well as satisfying the linear elastic assumption. The elastic modulus of the aluminum results in a ratio of bar stiffness to specimen stiffness of less than 20 for many engineering polymers which is what this device was designed to test.

The desired strain rate and total strain accumulation subjected to the specimen is another design variable which must be considered and is dependent on the bar and specimen dimensions. Based on the conservation of momentum, the theoretical upper limit on strain rate is determined by the velocity of the striker, v_{st} , and initial length of the specimen, L_s , i.e.,

$$\dot{\varepsilon}_{max} = \frac{v_{st}}{L_s} \tag{9}$$

This strain rate will not be realized in an actual experiment, but it serves a starting point during the design phase. To achieve very high strain rates, a balance must be struck between increasing the striking velocity and reducing the length of the specimen. The upper limit of striker velocity will most likely be dictated by the strength of the bar material and so it can be readily seen that reducing the length of the specimen will facilitate increased strain rates without having to manipulate the bar material. The theoretical maximum strain rate for the miniaturized Kolsky bar device used in the present work is $2.5 \times 10^4 \text{ s}^{-1}$ based off a specimen length of 1 mm and maximum striker velocity of 25 m/s.

The design of the specimen affects the maximum achievable strain rate as well as the accuracy of the Kolsky bar experimental results. Due to ease of machining, right cylindrical specimens are most commonly used. The end surfaces of the specimen must be flat and perpendicular to the axis of the specimen to maintain the validity of the one dimensional wave propagation equations. The ratio of the specimen length to specimen diameter (L_s/D_s) should be

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between 0.5 and 1.0. This range of ratios was determined as the optimal range to keep the adverse effects of radial inertia and specimen-bar interface friction to a minimum [Gray III, 2000]. Additionally, the diameter of the specimen is typically chosen to be no greater than 80% of the bar diameter to allow sufficient expansion in the radial direction during a test [Gray III, 2000]. The specimen dimensions chosen for the Kolsky experiments discussed here are a length of 1.0 mm and a diameter of 1.83 mm resulting in a length to diameter ratio of 0.55. These dimensions are similar to those employed by Jia and Ramesh [Jia, 2004].

3.2 Construction

The construction of the desktop Kolsky bar can be broken up into five main parts; the air delivery system, the alignment system, the momentum trapping system, the bars, and the data acquisition system. Figure 10 shows the desktop Kolsky bar with each of the sub-systems highlighted. A precision optical table is used as the base of the Kolsky bar to facilitate accurate component alignment and stability. The air delivery system, shown in Figure 11, consists of a compressed nitrogen tank, an air storage chamber, an electric solenoid valve, a safety release valve, and a gas gun barrel. The compressed nitrogen can be regulated between 0 and 3447 kPa depending on the desired striker bar velocity. A pressure of approximately 700 kPa is sufficient to attain striker velocities of 50 m/s, but all of the air deliver components were chosen to operate safely with pressures well above the maximum regulated pressure of 3447 kPa. The purpose of the small air chamber is to store compressed air at the desired pressure so that the air can be quickly released into the barrel once the solenoid valve is activated. The internal volume of the air storage chamber was designed to be approximately five times greater than the internal volume

of the barrel to ensure a near constant pressure expansion in the tube. A pressure indicator and safety release valve are attached to the storage chamber so the internal pressure can be accurately measured and so the pressure can be released from the chamber without activating the solenoid and releasing air into the barrel. The solenoid valve chosen is a SB051 (STC) because it has a fast reaction time and a maximum pressure rating of 6895 kPa. The gas gun barrel was machined from Stainless Steel 316 for its high strength and corrosion resistance. The bore of the barrel was machined to a diameter of 6.35 mm using a gun drilling process to ensure a straight hole with a smooth wall finish. Four air relief holes were drilled at one end of the barrel to ensure a constant striker velocity beyond these holes. A detailed drawing of the launch tube is provided in Appendix A in Figure 43.



Figure 10. Desktop Kolsky bar apparatus

The fifth hole is for the laser of the photogate [Pasco ME-9498A] to accurately measure the striker velocity immediately before impact. The photogate works by simply outputting a constant voltage when the laser receiver is not blocked and then outputting a null voltage when the receiver is blocked. The square voltage pulse created by the striker bar momentarily blocking the laser is acquired by a NI 9215 DAQ. A LabVIEW program computes the striker bar velocity based on the striker length and width of the timing pulse. The program is provided for reference in Appendix B.



Figure 11. Air delivery system

Precise alignment of the impact and transmission bars is essential for accurate Kolsky bar data. As such, the alignment system was designed from components commonly used in optical experiments since these components are precision machined and easily mount to the optical table. The bar alignment stanchions were constructed by affixing low-friction polymer bushings in the center of adjustable lens mounts (LH1 Thor Labs). Six posts were placed at intervals of 76.2 mm to provide adequate support and alignment for the impact and transmission bars. Precise alignment of the bushings was achieved by first attaching the lens mounts to the optical table and loosely placing the bushings in the center of each mount. The arms of the adjustable lens mounts were adjusted individually until the bars could easily translate from one stanchion to the next without binding and with minimal friction. The alignment posts are shown in detail in Figure 12. The alignment of the interface between the impact and transmission bars can be seen in Figure 13. The mounting blocks for the launch tube were machined from Aluminum 6061-T6 to maintain alignment between the striker bar and impact bar. A detailed drawing of the striker bar alignment blocks is provided in Appendix A in Figure 44.



Figure 12. Alignment stanchion



Figure 13. Interface between impact and transmission bars

The momentum trapping system is designed to quickly slow down the bars without damaging or misaligning the transmission bar. A tough rubber is attached to an energydissipative foam to act as a dashpot to remove the momentum from the bars. A hole slightly larger than the diameter of the transmission bar is drilled in the surrounding foam approximately 12 mm deep which forms a housing for the end of the transmission bar. The foam maintains the proper alignment of the transmission bar upon impact of the dashpot assembly. The momentum trapping system is shown in Figure 14.



Figure 14. Momentum trapping system

The striker, impact, and transmission bars are made from Aluminum 7075-T6. They were machined with tight tolerances on the diameter (\pm 0.01 mm) and straightness (0.1 mm per 300 mm). The diameter of all three bars is 3.175 mm. The length of both the impact and transmission bars is 254 mm. The striker bar length was chosen to be 76.2 mm to facilitate relatively high strain rate experiments ($1.5 \times 10^4 \text{ s}^{-1}$). The ends of all the bars were polished to ensure excellent flatness and perpendicularity to the axis of the bars. The striker bar is press fitted with custom made Turcite bushings, as shown in Figure 15, to provide an air tight seal in the bore of the gas gun with minimal friction. A detailed drawing of the bushing is provided in Appendix A in Figure 48.


Figure 15. Striker bar assembly

Perhaps one of the most challenging aspects of the desktop Kolsky bar design is the data acquisition system (DAQ). The extremely fast nature of the experiment requires the implementation of high-speed signal amplification and recording devices which are available commercially but are costly. Strain gauges [EA-06-031DE-350/LE] manufactured by MicroMeasurements with a gauge length of 0.79 mm and a grid width of 0.81 mm and a resistance of 350Ω were used to measure the strain pulses. One gauge was mounted longitudinally on the impact bar 165 mm from the specimen interface and another gauge was mounted longitudinally on the transmission bar 102 mm from the specimen interface. The procedures for preparing the bar surface, bonding the gauges to the bars, and soldering the lead wires were closely followed in accordance with those provided by the manufacturer. Bonded solder terminals were used as a junction between the small jumper leads of the strain gauges and the larger wires leading to the DAQ. The terminals protect the gauge wires from the tugging of the lead wires under high acceleration forces. The bonded terminal and strain gauge configuration is shown in Figure 16 (a). It was observed, however, that after numerous impacts the bonded terminal would de-bond from the bars thus leading to premature gauge lead wire

failure. The bonded terminal joint and lead wires were reinforced with a thin strip of electrical tape as shown in Figure 16 (b). The tape effectively protected the lead wires and added strength to the bonded terminal joint preventing gauge failure.



(a)



Figure 16. Bonded terminal and strain gauge configuration (a) and reinforced gauge (b)

Each of the gauges is connected to a quarter Wheatstone bridge completion module (BCM-1 Omega) employing the three lead wire configuration to increase gauge sensitivity and provide automatic temperature compensation. To amplify the signal from the Wheatstone

bridge, a custom circuit, shown in Figure 17, was designed and fabricated to provide the necessary gain at a sufficient bandwidth. The circuit is comprised of an instrumentation amplifier (INA111) set to a gain of 10 connected in series with two operational amplifiers (LF411) each set to a gain of 10 to give a final amplification of 1000. The limiting frequency response of the amplification circuit is approximately 500 kHz dictated by the operational amplifiers. A 25 MHz digital storage oscilloscope (2530B BK Precision) was used to acquire and record the amplified pulses for data processing. A flow chart illustrating the path of the signals from the strain gauges and photogate to the computer is shown in Figure 18. The typical range of either the change in resistance or change in voltage is denoted for each component.



Figure 17. Strain gauge amplification circuit



Figure 18. System signal flow chart

3.3 Calibration

Before experiments can be performed on a newly built testing apparatus, a comprehensive calibration procedure must be performed to verify the accuracy of the results. Calibration procedures should be repeated routinely as the material properties of the bar materials and strain gauge bond can change over time. Additionally, the device should be calibrated after any changes to the mechanical or electrical components are made to maintain confidence in the accuracy of the results. The following is a discussion of the methodology procedures employed on the miniaturized Kolsky bar device.

3.3.1 Strain Gauge Gain Calibration

Although the amplification circuit was designed to provide a total gain of 1000, small deviations in resistor values in addition to sources of error inherent to the operational and instrumentation amplifiers cause the actual gain to slightly differ. The relationship between the output voltage from the Wheatstone bridge and the final amplified voltage recorded by the oscilloscope was determined by offsetting the bridge by a known amount and measuring the amplified voltage. This was repeated for a range of bridge output values from approximately - 2.0 mV to 2.0 mV. The relationship is shown graphically for each of the strain gauge amplification circuits in Figure 19. Linear regressions were fit to both lines which provide the equations for converting the voltage recorded by the oscilloscope to the actual voltages being output from the Wheatstone bridges. Knowledge of the actual bridge output voltage is needed for the data analysis discussed later. The equation can be stated as follows

$$V_B = \frac{V_A - I}{G} \tag{10}$$

Here, V_B is the voltage from the bridge, V_A is the amplified voltage, I is the intercept, and G is the gain given by the slope of the line. The gain and intercept values for each of the strain gauges are given in Table 2.



Figure 19. Strain gauge gain calibration plot

Table 2. Strain gauge gain calibration values

Strain Gauge	Gain	Intercept (mV)
Impact Bar	1208.5	78.5
Transmission Bar	1203.8	-174.2

3.3.2 Strain Pulse Calibration

The calibration method discussed here compares the magnitude of the strain pulse measured by the gauges to the theoretical magnitude for the case where the impact and transmission bars are initially in direct contact. This is termed a "bars together" calibration. For such a condition, the magnitude (ε_I) and period (*T*) of the incident pulse can be analytically determined by Eqs. (2) and (3), respectively. If the bars are properly aligned and the bar surfaces are sufficiently flat, the entire pulse should transfer into the transmission bar and no reflection should be recorded by the gauge on the impact bar. Pulses generated by such a calibration test are illustrated in Figure 20. It should be noted that the magnitude of the measured pulse is less than the theoretical pulse denoted on the graph. Also, the entire incident wave is transmitted to the transmission bar with the exception of a small reflection pulse. This reflection is due to either a small misalignment in the bars or the bar faces are not exactly flat and parallel to each other. There are two important calibration values that can be determined from Figure 20; the elastic bar wave speed calibration and the strain pulse magnitude calibration.



Figure 20. Pulses for bars together calibration

It has been previously determined [Lifshitz, 1994] that a deviation of only 1% in the value of the bar wave speed (C_b) can significantly affect the accuracy of the specimen stress-

strain plot determined from Kolsky bar experiments. Therefore, instead of using handbook values for the elastic modulus and density of the bars to calculate the wave speed, the wave speed was determined by measuring the time it takes for the pulses to travel from the gauge on the impact bar to the gauge on the transmission bar. Specifically, the time between the two pulses at a strain value of 0.001, as denoted in Figure 20 as Δt , was measured for 20 experiments with the bars initially in direct contact. The travel time was determined to be 55.2 µs. The distance between the two gauges was measured at 273 mm. Thus, the calibrated value of elastic wave speed for the aluminum bars used in the miniaturized Kolsky bar apparatus is 4943 m/s. This value is approximately 2% from the theoretical wave speed value of 5051 m/s calculated from the handbook values of aluminum 7075-T6.

The calibration of the strain pulse magnitude is also critical for accurate specimen stressstrain results. The difference between the recorded and theoretical pulses arises from a number of errors characteristic of Kolsky bar experiments. Examples include the deviation from the assumed one-dimensional wave propagation behavior and the effects of the strain gauge bond. To compensate for these errors, the magnitude of the theoretical pulse based on the striker length and calibrated wave speed is compared to the recorded pulse. This magnitude is denoted in Figure 20 as a horizontal line. For each of the 20 "bars together" experiments, the ratio between the theoretical pulse amplitude and the measured amplitudes for each strain gauge were calculated. Even over the wide range of striker velocities at which the experiments were conducted (9-25 m/s), the calibration values were nearly constant. The average and standard deviations for the strain pulse amplitude calibration constants are provided in Table 3. The pulses from Figure 20 are shown again in Figure 21 after the calibration correction was applied.

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Figure 21. Calibrated pulses for bars together

Table 3. Strain pulse amplitude calibration constants

Strain Gauge	Calibration Constant	Standard Deviation
Impact Bar	1.15	0.07
Transmission Bar	1.19	0.07

4. EXPERIMENTAL SETUP

4.1 Specimen Preparation

The specimens employed for both the quasi-static and dynamic experiments were fabricated from unfilled Ultem 1000 which was received in an extruded 6.35 mm rod form. The cylindrical specimens were turned down to their final diameter ($D_s = 1.83$ mm) using a conventional high-speed steel lathe cutting tool. The circumferences and ends of the specimens were wet sanded with 600 grit silicon carbide sand paper to ensure a smooth and uniform finish. A micrograph of a finished specimen is shown Figure 22 with the final dimensions and estimated tolerances. A custom made specimen polishing jig, shown in Figure 23 was used to sand the specimen length down to $L_s = 1.0$ mm while ensuring the ends of the specimen were flat and perpendicular to the axis of the specimen. The jig allows up to19 specimens to be fabricated from one polishing operation which facilitates rapid and uniform specimen production. Prior to testing, the specimens were conditioned for at least 48 hours according to ASTM D695-08 Standard Test Method for Compressive Properties of Rigid Plastics. A detailed discussion of the effects of the polishing procedure is left for future investigation.



Figure 22. Micrograph of Ultem 1000 test specimen



Figure 23. Specimen polishing jig

4.2 Quasi-Static and Medium Strain Rate Compression Experiments

Compression experiments were carried out under quasi-static loading conditions $(\dot{\epsilon} = 2.1 \times 10^{-2} \text{ s}^{-1})$ and at a medium strain rate $(\dot{\epsilon} = 2.1 \times 10^{0} \text{ s}^{-1})$ for comparison to the high strain

rate mechanical behavior of Ultem 1000. The compression experiments were conducted on an electromechanical MTS Insight 5kN universal testing frame and an MTS 634.11E-25 axial extensometer was used to record the deflection of the specimens. As shown in Figure 24, the extensometer is contacting the surface of the compression platens instead of the specimen because the length of the specimen was below the minimum measurable length of the extensometer. Since the stiffness of the platens is much greater than that of the Ultem specimens, it was assumed that any deflection measured by the extensometer was due to the specimen only. A molybdenum disulfide based lubricant was used to reduce the friction between the platen and specimen surfaces.



Figure 24. Test configuration used in quasi-static testing

4.3 Kolsky Bar Experiments

To characterize the high strain rate behavior of Ultem 1000, Kolsky bar experiments were conducted at a range of strain rates from 8.7×10^3 to 1.5×10^4 s⁻¹. A minimum of three experiments were performed at each strain rate. The velocity of the striker bar can be used to estimate the strain rate of an experiment, but the actual strain rate, which is averaged over the stable duration of the test, can only be determined after processing the data and employing Eq. (4).

Before an experiment is performed on the miniaturized Kolsky bar apparatus, power to the Wheatstone bridges and amplification circuits is supplied for at least 20 minutes to allow the circuitry to warm up and stabilize. The excitation voltage powering the two Wheatstone bridges is 3.3 V which is the optimal voltage based on the strain gauge size, type, and bar. The length and diameter of each specimen is measured just prior to the experiment using a micrometer with a resolution of 0.0254 mm. The same molybdenum disulfide based lubricant used in the quasistatic experiments is applied to the ends of both the impact and transmission bars as shown in Figure 25. The specimen is carefully placed in between the two bars as they are brought together and the excess lubrication is wiped away. Once the specimen is centered such that it is coaxial with the bars, a clear box is placed around the specimen designed to contain the specimen during the experiment so that it can be recovered. A properly aligned specimen about to be tested is shown in Figure 26.



Figure 25. Lubrication applied to the impact bar

To acquire the strain pulses, a trigger is setup on the oscilloscope to collect data at 50 MHz after a rising edge is detected. The gas gun chamber is pressurized according to the desired strain rate and the apparatus is then ready to conduct the experiment. After the experiment is completed, the specimen length and diameter are measured and the specimen is labeled and bagged. Once the circuitry is warmed up, a complete experiment takes approximately 10 minutes to complete including the pre and post specimen measurements.



Figure 26. Specimen in Kolsky bar experiment

5. EXPERIMENTAL RESULTS

5.1 Quasi-Static and Medium Strain Rate Compression Results

The stress-strain curves discussed in the ensuing sections are in terms of the true rather than the engineering values of stress and strain. The conversion from engineering stress (σ_{eng}) and engineering strain (*e*) to true stress (σ) and true strain (ε) is given by

$$\varepsilon = \ln(1+e) \tag{11}$$

$$\sigma = \sigma_{eng}(1+e) \tag{12}$$

The above equations are only valid for the case of constant volume deformation which was verified for both the quasi-static and Kolsky experiments by measuring the specimen dimensions before and after each test. Stress-strain curves representative of the average response for both the quasi-static and medium strain rate compression experiments are shown in Figure 27. The curves exhibit a four stage response (linear elastic, nonlinear elastic, strain softening, strain hardening) typical of amorphous polymers. The average yield stress was found to be 166 MPa and 193 MPa at strain rates of 2.1×10^{-2} and 2.1×10^{0} , respectively.



Figure 27. Quasi-static stress strain curves

5.2 Kolsky Bar Results

5.2.1 Data Analysis

Ideally, the Kolsky bar apparatus will provide the stress-strain curve for the specimen throughout the entire loading duration, which is governed by Eq. (3). Also, under ideal conditions, the strain rate would be constant during the entire loading phase. From these stress-strain curves, a relationship between strain rate and the mechanical properties such as elastic modulus, yield strength, ultimate strength, and the hardening/softening behavior can be determined. However, due to some of the limitations inherent to the Kolsky bar apparatus, the strain rate is not exactly constant throughout the test and there is a finite time during which the specimen is not in stress equilibrium. During this time, known as the "ring-up period," Eq. (4)

and Eq. (5) are not valid. The ring-up period duration, τ , is the time it takes for π reverberations to traverse through the specimen [Davies and Hunter] and is expressed

$$\tau = \pi \frac{L_s}{C_s} \tag{13}$$

where L_s is the initial length of the specimen and C_s is the elastic wave speed of the specimen. Therefore, the stress-strain response of the material before τ seconds cannot be considered valid. Often times this means that the elastic portion of the stress strain curve cannot be accurately determined. Methods have been developed to decrease the ring up period and to increase the duration at which the strain rate is constant through a technique referred to as "pulse shaping." These techniques were not employed in the current Kolsky bar apparatus, but are saved for future work and are discussed in Chapter 6

To illustrate how the pulses recorded during a Kolsky bar experiment are analyzed and how they are checked for validity, an example is provided here detailing the steps taken to fully process the experimental data. The data analysis procedure includes data smoothing, pulse synchronization, and algebraic manipulation of the pulses. A program was developed in MATLAB (see Appendix C) to automatically perform the smoothing and algebraic operations. Determining the starting and ending points of the incident, transmitted, and reflected pulses was done manually, which if done incorrectly introduces error in the stress-strain curve [Chen, 2011]. The pulses that are acquired from the oscilloscope are in terms of an amplified voltage and are slightly offset from zero due to the Wheatstone bridges being initially unbalanced. Before the pulses can be converted from voltage to strain, the amplified voltage is converted into the actual bridge output voltage by employing the strain gauge gain calibration equation discussed in Section 3.3.1. The voltage now is on the order of a few millivolts instead of on the order of volts

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which was recorded by the oscilloscope. The baselines of these pulses are then shifted to zero by taking the average of the output voltage for 20 μ s before the impact is recorded and subtracting this offset value from the magnitude of the pulses. The voltages are then converted into strain using the one-quarter Wheatstone bridge equation below

$$\varepsilon = \frac{4\Delta V}{G_f V_E} \tag{14}$$

where ΔV is the change in output bridge voltage, G_f is the gauge factor for the gauges used and V_E is the bridge excitation voltage. The pulses, now in terms of strain, are multiplied by the strain amplitude calibration constants discussed in Section 3.3.2. Finally, the strain pulses are smoothed using a local weighted regression algorithm. The data analysis code provided in Appendix C performs all of these operations automatically for each experimental data set. The example experiment was conducted on an Ultern specimen with dimensions $L_s = 1.039$ mm and $D_s = 1.778$ mm and with a striker velocity of 23.8 m/s. Therefore, according to Eq. (3), the pulse width should be approximately 31 µs. The processed pulses for this example case are illustrated in Figure 28.



Figure 28. Post-processed strain pulses from example Kolsky bar experiment

The next procedure is to synchronize the three pulses by first determining their start and end points and then eliminate the time variable between them as shown in Figure 29. To check the validity of the stress equilibrium assumption, the incident and reflected pulses are added together and compared to the transmitted pulses, as shown in Figure 30. Only for times at which the addition of the incident and reflected pulses is sufficiently close to the transmitted pulse is the specimen in stress equilibrium. The percent difference between the transmitted pulse and the addition of the incident and reflected pulses are shown graphically in Figure 31. Although the ring-up duration given by Eq. (13) for Ultem is approximately 2 μ s, it can be seen from Figure 31 that for the miniaturized Kolsky bar apparatus discussed here the time is approximately 4 μ s.



Figure 29. Synchronized strain pulses



Figure 30. Stress equilibrium comparison graph



Figure 31. Stress equilibrium verification graph

The other condition that must be checked to determine the validity of the test results is the duration of the test at which the specimen is under a constant strain rate. The strain rate as a function of time is given by Eq. (4) and is shown graphically for the example case in Figure 32. Clearly, from Figure 32 it is seen that the strain rate is not constant throughout the duration of the test, but an average strain rate after 4 μ s is used as the strain rate for the experiment and is calculated to be approximately 14470 s⁻¹ for the present example. The accumulated strain in the specimen as a function of time, shown in Figure 33 is found by integrating Eq. (4) at each time increment and adding the individual strain increments. The value of strain at 4 μ s is approximately 2%. Hence, the stress response corresponding to strains less than 2% cannot be accurately determined for this particular experiment due to the ring-up period. When the stress as a function of time, calculated from Eq. (5) is superimposed with the strain as a function of time, the engineering stress-strain curve is produced as shown in Figure 34. Again, a line is shown on the graph denoting the point before which the data cannot be validated. From Figure 34 the typical mechanical material properties can be estimated such as yield stress, ultimate stress, and the hardening/softening behavior.



Figure 32. Strain rate versus time

In summary, the data analysis procedure consists of processing the raw voltage pulse data recorded from the oscilloscope by smoothing and scaling it according to the calibration constants discussed in Chapter 3. Next the voltage is converted into strain through Eq. (14) and the incident, reflected, and transmitted pulses are synchronized from which the strain rate and stress as functions of time are calculated. The portion of the data that can be validated is determined by investigating the stress equilibrium and strain rate characteristics of the data. Finally, the true

stress-strain response is created from which the mechanical behavior of the specimen at that particular strain rate can be determined. This process is repeated for experiments at a range of strain rates to characterize the relationship between the mechanical behavior of the specimen material and strain rate.



Figure 33. Specimen strain versus time



Figure 34. Specimen stress strain curve

5.2.2 Experimental Results

Data was collected from 12 Kolsky bar experiments and analyzed using the methods presented in Section 5.2.1. For each test the stress equilibrium condition was verified and the average strain rate was computed. The specimens were recovered and measured after each experiment to verify the validity of the constant volume deformation assumption. The difference between the initial and final volumes for each of the experiments was within 10%. A comparison between a sample before and after a Kolsky bar experiment is shown in Figure 35. None of the samples tested fractured or showed visible signs of material failure.



Figure 35. Ultem 1000 specimen before and after Kolsky bar experiment

The dynamic stress strain curves for only a few of the experiments are presented here to illustrate some key rate dependent characteristics observed for Ultem 1000. The stress strain response of Ultem 1000 at the lowest $(8.7 \times 10^3 \text{ s}^{-1})$ and highest $(1.5 \times 10^4 \text{ s}^{-1})$ strain rates performed on the miniaturized Kolsky bar device are illustrated in Figure 36. The response for both the quasi-static and medium strain rate test is included in this figure for comparison. The yield stress at the strain rates of $8.7 \times 10^3 \text{ s}^{-1}$ and $1.5 \times 10^4 \text{ s}^{-1}$ were 290 MPa and 308 MPa, respectively. The yield stress for the intermediate strain rates showed a similar pattern of increasing with strain rate.



Figure 36. Stress-strain curves of Ultem 1000 at various strain rates

To illustrate the relationship between the strain rate and yield stress of Ultem 1000, the yield stress for a wide range of strain rates is plotted against the corresponding strain rate as shown in Figure 37. Clearly, the yield stress of Ultem 1000 exhibits a distinct bilinear dependence on $\log(\dot{\epsilon})$. It appears that the transition occurs at approximately 1.2×10^3 s⁻¹.



Figure 37. Ultem 1000 yield stress versus strain rate

Such a bilinear relationship has been observed in previous works for other amorphous polymers (PC, PMMA) over similar ranges of strain rates (See Section 2.2.2). It can therefore be reasonably concluded that the rate dependence of Ultern 1000 is dually affected by simultaneous activation processes which can be modeled with the Ree-Eyring Eq. (7). The experimental data shown in Figure 37 was fit to the Ree-Eyring model and the results are provided in Figure 38. The coefficients determined for the best fit are shown in the inset of Figure 38. The temperature term in the model ($\theta = 295$ °K) was set to the temperature at which the experiments were conducted (room temperature).



Figure 38. Ree-Erying model fit to experimental data

Many studies concerning the rate dependence of amorphous polymers limit their discussion to the effects on yield stress and devote little attention to the behavior of the material after this point. The present study however seeks to establish a relationship between the post yield behavior of the material and the strain rate. Admittedly, the development of such a relationship would require the consideration of the complex interaction between temperature rise and polymer chain rearrangements. However, a simplification of the post yield behavior facilitates the observation of some rate dependent trends which would otherwise be require further testing at a range of temperatures. Hence, as a first step in quantifying this relationship, the stress-strain curves collected from the Kolsky bar experiments were modeled with the power law Ramberg-Osgood equation which takes the form

$$\sigma = K \varepsilon_p^n \tag{15}$$

where ε_p is the plastic strain and *K* and *n* are fitting parameters that describe the yield point and hardening behavior, respectively. A comparison between the experimental stress-strain data and the model is illustrated in Figure 39. The plastic portion of the strain (ε_p) is separated from the total strain (ε) by subtracting the elastic strain, i.e.,

$$\varepsilon_p = \varepsilon - \frac{\sigma}{E} \tag{16}$$

Here, E is the slope of the initial linear portion of the stress strain response. It is appropriate to limit the scope of the mathematical modeling to only the plastic strains as defined by Eq. (16) since the response in the elastic strain range cannot be accurately determined for the Kolsky bar experiments discussed here (See Section 5.2.1).



Figure 39. Ramberg-Osgood model fit of experimental data

The stress-strain curves for various strain rates between 8.7×10^3 s⁻¹ and 1.5×10^4 s⁻¹ were fit to Eq. (15) and the results are provided in Figure 40. An observation readily made from Figure 40 is that the slope of the hardening portion of each of the curves is nearly a constant for each of the strain rates. This relationship can be easily seen by plotting the hardening coefficient, *n*, against the strain rate and observing a nearly linear correlation as is the case in Figure 41. The rate dependence of the Ramberg-Osgood coefficient *K* was found to increase linearly with strain rate, as is shown in Figure 42.



Figure 40. Ramberg-Osgood model for various strain rates



Figure 41. Rate dependence of hardening coefficient, n



Figure 42. Rate dependence of Ramberg-Osgood coefficient, K

6. DISCUSSION

Overall, the experimental results for both the quasi-static and high strain rate tests were consistent with trends observed previously for other amorphous polymers. This fact provides a level of confidence in the operation of the Kolsky bar device and in the data analysis techniques. The quasi-static stress strain behavior of Ultem 1000 exhibits the typical response for an amorphous polymer which has been well defined in terms of polymer chain sliding and rearranging. At this low strain rate, the effects of temperature rises due to plastic deformation can be ignored.

Regarding the high strain rate experiments, perhaps the most noticeable trend is the bilinear relationship between the yield stress and the $log(\dot{\varepsilon})$. For Ultem 1000, the transition between the two linear regions is approximately 10^3 s^{-1} . This transition point is similar to that of PC and PMMA determined by Walley and others [Walley, 1991]. Determining a more precise value for this transition point would require additional experiments in the range of 10^1 to 10^3 s^{-1} which are out of the operating range of both standard electromechanical test frames and Kolsky bar devices. Determining the cause of this transition is of considerable interest, especially if a molecular based constitutive model is to be developed. Past investigators have explained this bilinear relationship in terms of two activation processes concurrently taking place and the second process becoming dominant after the transition point. In order to fully understand the mechanisms causing this transition, the effects of local temperature rises must be considered. The effect of temperature on the stress-strain response can be investigated in two ways. One method would be to measure any thermal changes during an experiment and try to correlate this to an activation process. A secondary method to investigate thermal effects would be to

conducted experiments at a range of temperatures for each level of strain rate. The data from these experiments would allow the effects of temperature and strain rate to be decoupled, thus providing a better understanding of the molecular mechanisms causing the transition to occur. Conducting these experiments would require physical modifications to the Kolsky bar device and are saved for future investigations (See section 7.2).

A point to consider when discussing the apparent transition in yield stress rate dependence is the fact that the data for each of the two linear regions was acquired on different testing devices i.e., an MTS testing frame for the low and medium strain rates and the Kolsky bar device for the high strain rates. Although the specimen size employed for the two testing methods was the same, other factors such as differences in frictional forces and data acquisition methods could contribute to the apparent transition in the rate dependence. Therefore, care should be taken when conducting the experiments to minimize the variables between the two test methods that could affect the stress-strain response.

An observation that was made in the present work that is typically not discussed in polymer rate dependence studies is the post yield behavior of the material. The stress-strain response for the quasi-static and medium strain rate tests, shown in Figure 36, exhibit the typical strain softening followed by a steep strain hardening region. In contrast, the stress-strain response of the high strain rate tests, also shown in Figure 36, do not exhibit a clear strain softening region. Additionally, the hardening behavior, although still observed, is not as sharp as in quasi-static case. It is difficult to postulate the causes responsible for the apparent lack of a strain softening region because of the noise observed in the high strain rate stress-strain curves. Before a determination can be made whether this observation is representative of an actual

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material response or a mere artifact of the available data, more experiments need to be conducted with improvements to the Kolsky bar device to reduce noise. The differences in the hardening behavior between the quasi-static and high strain rate experiments could be a result of temperature rises in the material at the high strain rates. If it is assumed that the same molecular mechanisms that cause strain hardening in quasi-static case (polymer chain realignment) are also at work in the high strain rate cases, then the rise in temperature could have a softening effect on the material in conjunction with the hardening process due to the polymer chain realignment. This dual effect of polymer chain rearrangement with the introduction of heat generation could explain why hardening is still observed, but to a lesser degree. However, to be sure, more experiments should be conducted where the temperature rises are measured directly.

7. CONCLUSION AND FUTURE WORK

7.1 Conclusions

The achievements of the present work can be summarized into two categories: (1) successful Kolsky bar apparatus design, construction, and operation and (2) material characterization of Ultem 1000 over a wide range of strain rates. The provided methodologies for designing and fabricating a miniaturized Kolsky bar device as well as the provided data processing procedures and codes can serve as a guide to those seeking to build and employ these devices. A method for fabricating the test specimens used for these experiments was presented along with the engineering drawings for the custom polishing jig used. Additionally, the design for an economical amplification circuit board capable of acquiring the signal pulses from Kolsky bar experiments was provided.

The trends observed from the high strain rate testing of Ultem 1000 were consistent with those for other amorphous polymers tested under similar conditions. Specifically, the relationship between the yield stress and the $log(\dot{\epsilon})$ was bilinear. This correlation was modeled with the Ree-Eyring equation. Additionally, the differences between the post yield behavior of the quasi-static and high strain rate experiments was observed and possible explanations were provided.

7.2 Future Work

The Kolsky bar device in its present configuration is capable of testing a variety of polymeric materials at room temperature from strain rates of approximately 10^3 s⁻¹ to 10^5 s⁻¹.
From these experiments, a number of rate dependent characteristics can be observed. However, it is often desirable to characterize a material at both a wide range of strain rates and temperatures. As such, future work on the Kolsky bar device includes the addition of a thermal chamber capable of heating or cooling a sample just prior to testing. For room temperature experiments, the chamber will serve as a thermal barrier around the specimen so the temperature rise due to rapid plastic deformation can be measured. Such data will be helpful when correlating the macroscopic response of the material to molecular level activation processes. Additionally, pulse shaping techniques will be used in subsequent experiments to more precisely control the strain rate over the duration of an experiment.

Another important experimental characteristic that will be investigated in the future is the effects of the specimen fabrication on the results. Specifically, the geometric tolerances on the specimen end flatness and parallelism and the diametrical tolerances could have a significant effect on the accuracy of the stress-strain curves produced from Kolsky bar experiments. The effects of these tolerances on the stress-strain results can be investigated through a parametric finite element analysis. For example, the ends of the specimens can be modeled such that they are not parallel to each other or perpendicular to the specimen axis. The amount of angular offset can be incrementally varied and the numerical stress-strain results can be compared for various magnitudes of offset until an acceptable tolerance band is established. A similar procedure could be conducted with diametrical tolerances on the specimen. Once a geometric tolerance band that yields repeatable and accurate stress-strain results has been established, the specimen machining processes can be optimize to fabricate specimens which continually fall within the specified tolerances. The current specimen fabrication process, which involves

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considerable manual work, is time consuming and does not easily produce uniform specimens. As such, an improved manufacturing process involving micro-milling is being developed which is capable of fabricating specimens with a resolution of 50 micrometers. This fabrication process will facilitate the rapid production of hundreds of nearly identical specimens with exceptional geometric tolerances.

The end goal of many material characterization studies is the implementation of the experimental results into a numerical simulation. This can be accomplished by either determining the fitting constants for an existing constitutive model or by developing a novel constitutive model. Future studies will attempt to fit the experimental data for Ultern 1000 into existing constitutive models for amorphous polymers. Such a constitutive model can then be employed in an explicit numerical simulation code, such as LS-DYNA, to verify the experimental results as well as determine the response of Ultern 1000 under novel combinations of temperature and load.

APPENDIX A: TEST HARDWARE



Figure 43. Launch tube drawing (dimensions in inches)



Figure 44. Launch tube alignment block drawing (dimensions in inches)



Figure 45. LH1 lens holder drawing (Thor Labs)



Figure 46. TR3 lens mount post drawing (Thor Labs)



Figure 47. PH2 post holder (Thor Labs)



Figure 48. Striker bar bushing drawing (dimensions are in inches)

APPENDIX B: STRIKER BAR VELOCITY LABVIEW VI







Figure 49. LabVIEW VI block diagram (a) and front panel (b)

APPENDIX C: DATA PROCESSING CODE

close all clc clear all

tic

% Define variables	
file_name='ultem_data';	% File name
CH1_gain=1208.5;	% Gain for CH1
CH2_gain=1203.8;	% Gain for CH2
CH1_intcp=0.0785;	% Gain intercept for CH1 (V)
CH2_intcp=-0.1742;	% Gain intercept for CH2 (V)
gauge_factor=2.08;	% Strain gauge factor
ex_volt=3.288;	% Bridge excitation voltage (V)
Cb=4943;	% Bar wave speed (m/s)
span=101;	% Smoothing span value (must be odd)
run=1;	% Refers to the sheet number in Excel

% Read record length and time increment N=xlsread(file_name,run,'B1:B1'); inc=xlsread(file_name,run,'B2:B2');

for run=1:13 % Loop for all sheets

% Read time, incident pulse, and transmission pulse vectors CH1=xlsread(file_name,run,'B14:B16397'); CH2=xlsread(file_name,run,'C14:C16397');

% Reduce magnitude of pulses using gain equation CH1_gain=(CH1-CH1_intcp)/CH1_gain; CH2_gain=(CH2-CH2_intcp)/CH2_gain;

% Determine pulse offset zeros CH1_zero=mean(CH1_gain(1:500)); CH2_zero=mean(CH2_gain(1:500));

% Offset pulses to start at zero CH1_adj=CH1_gain-CH1_zero; CH2_adj=CH2_gain-CH2_zero;

% Convert pulses from voltage to strain CH1_strain=(4*CH1_adj)/(gauge_factor*ex_volt); CH2_strain=(4*CH2_adj)/(gauge_factor*ex_volt); % Smooth data CH1_smooth=smooth(CH1_strain,span,'rlowess'); CH2_smooth=smooth(CH2_strain,span,'rlowess');

% Build matrix of values to write back into excel workbook output(:,1)=CH1_gain; output(:,2)=CH2_gain; output(:,3)=CH1_adj; output(:,4)=CH2_adj; output(:,5)=CH1_strain; output(:,6)=CH2_strain; output(:,7)=CH1_smooth; output(:,8)=CH2_smooth;

% Write output matrix back into excel workbook xlswrite(file_name,output,run,'D14:K16397');

end

toc

APPENDIX D: SPECIMENS AND TEST DATA

Specimen #	Average Strain Rate	Yield Stress	Total Strain
-	s ⁻¹	MPa	%
100	14277	289	32.5
101	12744	285	28.3
102	12787	302	28.7
103	12447	292	37.0
104	13235	293	35.6
107	8991	289	18.2
108	8677	308	12.6
109	10291	302	19.7
110	9441	309	20.3
111	13682	311	30.7
112	14509	311	25.2
113	14951	289	37.4

Table 4. Summary of Kolsky bar experimental data



Figure 50. True stress-strain curve for specimen 100



Figure 51. True stress-strain curve for specimen 101



Figure 52. True stress-strain curve for specimen 102



Figure 53. True stress-strain curve for specimen 103



Figure 54. True stress-strain curve for specimen 104



Figure 55. True stress-strain curve for specimen 107



Figure 56. True stress-strain curve for specimen 108



Figure 57. True stress-strain curve for specimen 109



Figure 58. True stress-strain curve for specimen 110



Figure 59. True stress-strain curve for specimen 111



Figure 60. True stress-strain curve for specimen 112



Figure 61. True stress-strain curve for specimen 113

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