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# STRUCTURAL AND PHYSICAL PROPERTIES OF SRAPHITE-ZIRCONIUM DIBORIDE-SILICON CARBIDE COMPOSITES

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UNION CARBIDE CORPORATION CARBON PRODUCTS DIVISION

TECHNICAL REPORT AFML-TR-70-50

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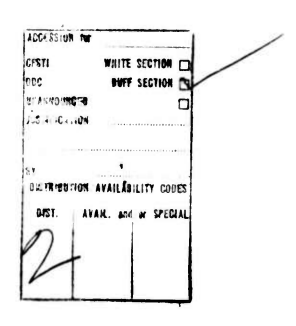
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# STRUCTURAL AND PHYSICAL PROPERTIES OF GRAPHITE-ZIRCONIUM DIBORIDE-SILICON CARBIDE COMPOSITES

Union Carbide Corporation Carbon Products Division

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#### FOREWORD

The work reported herein was performed under the sponsor-ship of the Advanced Research Projects Agency, Department of Defense, through a contract with the Air Force Materials Laboratory, MAN, AFSC Wright-Patterson Air Force Base, Ohio, Contract No. AF 33(615)-3110, ARPA Order No. 719, Program Code 6D10. Mr. H. S. Schwartz, MAN is the Air Force Program Manager.

The Contractor is Union Carbide Corporation, Carbon Products Division, Parma Technical Center, P.O. Box 6116, Cleveland, Ohio 44101. The Program Supervisor is Dr. Herbert F. Volk. Major contributors to the work reported herein were Dr. George B. Spence, Dr. Tu-Lung Weng, Mr. Owen L. Blakslee, Mr. Robert G. Fenish, and Mr. Charles W. Nezbeda.

The work reported herein was conducted during the period from May 1965 to June 1967, and the report was submitted for publication in May 1970.

This technical report has been reviewed and is approved.

A. M. Lovelace

Director

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Air Force Materials Laboratory

#### ABSTRACT

Three series of graphite based, refractory composites compositionally related to commercial Grade JTA, but containing 30, 50, and 70 weight percent additives (ZrB2 and Si), were prepared. Within each series, the porosity was varied from approximately 2 to 17 percent. Property measurements were carried out as a function of composition and porosity and along the two principal symmetry axes, i.e., normal and parallel to the molding direction. Property measurements carried out at room temperature consisted of tension, compression, flexural, torsion, ultrasonic, and resonant bar tests. The specific heat, thermal conductivity, coefficient of thermal expansion, and effect of high temperature annealing were also determined. Stress-strain relations and fracture strength under multiaxial stress were determined for the material with 50 percent additive. A generalized stress-strain relation for a transversely isotropic, nonlinear material has been obtained without assuming volume constancy. Various theories of fracture for anisotropic materials were compared with the fracture strength of JT materials. The comprehensive property data compilations in this report should provide designers with sufficient information for structural applications of JT materials.

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#### SECTION I

#### INTRODUCTION

The information presented in this report was generated as part of a much larger program, "Integrated Research on Carbon Composite Materials," which is designed to fulfill three different, but clearly interdependent needs of the Department of a material need, a structural design capability need, and a need for more scientists and engineers trained in applied materials problems and advanced design methods. The Carbon Products Division of Union Carbide Corporation, Case Western Reserve University, and Bell Aerospace Company have formed an Association through which a joint effort is made to meet these needs. The Association has formulated a broad program which includes the development of new materials, generation of advanced analyses and design methods, and education of graduate students. The graphite based refractory compostes (designated by Union Carbide as JT series) which are the subject of this report are excellent candidates to satisfy the needs of weapons and aerospace systems for structural materials exhibiting high temperature oxidation resistance, good thermal shock resistance, refractoriness, and good machinability. These composites, upon exposure to oxygen at elevated temperatures, enclose themselves with a protective oxide coating. In this fashion, they represent an improvement over coated systems by providing oxidation protection Should processes such as erosion, chemical reaction, or vaporization result in local or general loss of the protective layer. Based on this mode of behavior, this class of composites is a candidate material for use above 3000°F in an oxidizing envirorment. Previous Air Force Materials Laboratory reports containing information on the JT series composites are listed as Reference 1-5 inclusive. Commercial Grade JTA is but one example of this series of composites; the properties of these materials can be varied over a wide range through appropriate changes in composition and/or porosity. The objective of this work was to provide designers and structural engineers with sufficient characterization of the JT-series composites to permit proper materials selection and structural optimization.

The results obtained under the Association's program have been reported in four Annual(6-11)Reports. However, the overall program is primarily concerned with graphite fiber reinforced composites; the results on particulate composites are scattered throughout a number of these reports and are, therefore, not conveniently accessible. Moreover, some property measurements (1) on the JT-series composites have been superseded by more accurate work and/or by data obtained on billets fabricated by a only in Quarterly Progress Reports which are not as readily available

as the Annual Reports. For these reasons, it was decided to publish the present Topical Report, which in a convenient format provides the designer with sufficient details for the structural application of these materials.

#### SECTION II

#### SUMMARY

The fabrication procedures and properties of a series of graphite-based, particulate refractory composites containing ZrB<sub>2</sub> and SiC were investigated. The hot-pressing parameters (particularly temperature and pressure conditions) required to produce dense cylindrical billets up to 3-inch diameter by 3-1/2-inch high were determined. A float mold technique was found to produce the most uniform billets, and this method was used to produce twenty-seven 3 x 3.5-inch specimens for the property evaluation work.

The elements Zr, B, and Si (the "metallic additives") were always maintained in the same proportion as in the commercial composite Grade JTA\*, i.e., 42.4 parts of ZrB2 to 9.5 parts of Si. Three series of composites containing 30, 50, and 70 weight percent metallic additives were prepared; within each series, the porosities of the billets were intentionally varied from approximately 2 to 17 percent. Thus, the physical properties were determined as a function of composition and of porosity.

Since the JT-series materials have the symmetry characteristics of transverse isotropy (the symmetry axis is parallel to the direction of molding pressure), it was necessary to measure all physical properties in two directions: parallel and normal to the symmetry axis. Mechanical properties were determined at room temperature by both static and sonic techniques. The static measurements included tension, compression, flexural, and torsion tests; the sonic tests consisted of ultrasonic and resonant bar tests. In addition to the ambient temperature measurements, the thermal properties of the JT-series materials were also determined, again as a function of both composition and of porosity. The specific heat, thermal diffusivity, thermal conductivity, and coefficient of thermal expansion were determined. The effects of annealing JT-materials at various temperatures were also studied. The uniaxial stress-strain relations for JT-materials are nonlinear and nonconservative, and brittle fracture is preceded by plastic flow. Strengths and moduli increase with increasing metallic additive content and decreasing porosity. The physical property data, both mechanical and thermal, provide sufficient information to optimize the material and geometry of a structural component for specific applications.

<sup>\*</sup>A product of Union Carbide Corporation, Carbon Products Division.

Stress-strain relations and fracture strength of JT-50 (50 percent metallic additive) for all four biaxial stress quadrants were determined on thin-walled cylindrical specimens subjected to combinations of internal pressure, external pressure, and axial load. The stress-strain relations of this material under combined stresses are also nonlinear and nonconservative. A generalized stress-strain relation for a transversely isotropic, nonlinear material has been obtained without assuming volume constancy. The correlation between effective stress and effective strain for the uniaxial and biaxial stress tests is satisfactory for engineering applications.

Various theories of fracture for anisotropic materials were compared with the fracture strength data of JT-50 materials. An empirical fracture criterion has been formulated from the fracture strength data, and the fracture envelopes for JT-50 series materials as function of composition and porosity have been calculated with these fracture criteria. Significant influences of anisotropy and stress state on the modes of fracture were observed for JT-50 material. Catastrophic fracture resulted when the specimen was subjected to a large compressive stress, regardless of whether the other stress was tensile or compressive.

#### SECTION III

#### FABRICATION OF JT-SERIES MATERIALS

## 1. Description of JT-Series Materials

The JT-series materials are graphite-base refractory composites similar to the commercial material having the grade designation JTA produced by the Carbon Products Division of Union Carbide Corporation (JTA is one member of the series). On a weight basis, Grade JTA is composed of 48.1 percent C, 42.4 percent ZrB<sub>2</sub>, and 9.5 percent Si. The minimum density limit for Grade JTA is 3.0 g/cm<sup>3</sup>. During manufacture, silicon combines with part of the carbon to form silicon carbide.

In this work, the elements Zr, B, and Si are called the metallic additives; these elements are always present in the same proportions as in JTA: 42.4 parts of ZrB<sub>2</sub> to 9.5 parts of Si. Let

- c = mass fraction of metallic additive
  in the fabricated product, and
- $\rho$  = density;

Grade JTA may, then, be specified by composition c = 51.9 percent and density  $\rho > 3.0$  g/cm<sup>3</sup>.

The JT-series materials are a class of materials produced by varying the amount of metallic additives and the density. In order to vary the density at a fixed composition, variations had to be made in processing temperatures or pressures; otherwise, the manufacturing conditions are kept as close as possible to those for grade JTA.( $^1$ ) Therefore, to a good approximation, each member of the JT series can be uniquely identified by two values: the compositional variable, c, and a value that is equivalent to a processing variable, the density,  $\rho$ . To the same approximation, the physical properties of the JT-series materials may be considered to be functions of only two material variables, c and  $\rho$ .

# 2. <u>Material Symmetry, Coordinate Orientation, and</u> Billet Notation

All of the JT-series materials were molded in cylindrical billets. This process yields material with the symmetry characteristic of transverse isotropy, i.e., all physical properties are invariant with respect to arbitrary rotations about the symmetry axis, which is parallel to the direction of molding and perpendicular to the molding ram face.

Throughout this study, physical properties of the JT-series materials are specified with respect to a Cartesian co-ordinate system oriented with the x<sub>1</sub> and x<sub>2</sub> axes in the plane of transverse isotropy and the x<sub>3</sub> axis parallel to the axis of rotational symmetry. These symmetry oriented coordinates should not be confused with other types and orientations of coordinate systems used in the stress analysis problems. In JT material, as in other molded material, the elongated graphite particles or grains tend to be aligned with their two larger dimensions parallel to the plane of transverse isotropy, a situation which has led to properties (Young's modulus, coefficient of thermal expansion, etc.) in directions parallel to the plane of isotropy being called "with-grain" properties and properties in the direction parallel to the symmetry axis being called "against-grain" or "across-grain" properties.

The billets of JT material fabricated especially for this program are identified by the letters JT, followed by the nominal composition, and, finally, the sequential fabrication number. Thus, JT30-4 designates that the billet is the fourth fabricated and that the composition is approximately 30 percent metallic additive.

Commercial JTA billets from the standard production material which were also used in this program are identified by the letters JTA followed by the sequential order number. Thus, JTA-9 designates the ninth piece of commercial JTA used for this program.

# 3. Calculation of Theoretical Maximum Density and Porosity

Because the density varies with the composition as well as with porosity in the material, the composition c and porosity p were chosen as the material variables, rather than composition and density. Part of the porosity is in closed pores, the volume of which is not easily measured. Therefore, the porosity was calculated from the measured bulk density  $\rho$  and the calculated theoretical maximum density  $\rho_{\rm m}$ :

$$p = 1 - \rho/\rho_{\rm m} \tag{1}$$

The theoretical maximum density can be calculated from the mass fraction and X-ray density of each phase in the composite material. Based on the following assumptions:

mass ratio of  $ZrB_2$  to Si is 42.4/9.5, all Si is in the form of  $\alpha$ - and  $\beta$ -SiC, and X-ray density of pure C is 2.267 g/cm<sup>3</sup>, of  $ZrB_2$  is 6.10 g/cm<sup>3</sup>, and of SiC is 3.216 g/cm<sup>3</sup>,

the calculated theoretical maximum density is given by

$$\rho_{\rm m} = \frac{2.267}{1 - 0.5905 \, \rm c} \tag{2}$$

where c (0 < c < 1) is, as above, the mass fraction of Zr,B, and Si the fabricated composite. In practice, there are uncertainties of perhaps one percent in the mass fraction c and  $ZrB_2/Si$  ratio and in the density of  $ZrB_2$ . At low porosity levels of less than five percent, the corresponding uncertainty in the calculated porosity is several tens of percent. The possible error in the calculated porosity will not affect the stress analysis and structural synthesis work provided that Equations (1) and (2) are used consistently in calculating the porosity.

## 4. Fabrication of Initial Billets

The initial series of billets to be fabricated was selected to provide the initial trend of property data with composition at a fixed porosity and the trend with porosity at a fixed composition. Billets three inches in diameter by 3-1/2 inches in length were pressed on production-size equipment at a maximum observed outside-surface mold temperature of 2130°C and at pressures ranging between 3,300 and 5,500 lb/in.². Two attempts to fabricate composites with ten percent additive were unsuccessful due to mold failures. Two other billets were of smaller size. Table I gives the composition, average density and porosity of the good billets and of a commercial JTA billet. All of these billets were cut into specimens for physical property testing. The test results are given in Section IV.

TABLE I
COMPOSITION, DENSITY, AND POROSITY OF
INITIAL JT-SERIES BILLETS

Billet Number	Composition Percent Metallic Additive	Average Density g/cm <sup>3</sup>	Average Porosity Percent
JT-30-5	31.5	2.31	17.2
JT-30-6	31.5	2.38	14.7
JT-30-4	31.5	2.58	7.4
JT-40-9	41.7	2.81	•
JTA-9	51.9	3.05	6.7 6.7
JT-70-3	71.4	3.66	6.7
JT-70-8	71,4	3.66	6.7

# 5. Process Studies on 1- and 1.5-Inch Diameter Billets

A basic premise of the entire JT-series program was that the physical properties of the materials depend on only two material variables, composition and density, and not on, say, the individual values of processing temperature and pressure used to attain a certain density. When it was found that the physical properties of commercial JTA did not fit smoothly with those of the initial experimental billets, a limited study of processing conditions was initiated to determine conditions which would yield material more comparable with commercial JTA on laboratory size equipment.

- a. Method of Fabrication. Experimental billets 1.0 and 1.5 inches in diameter with the same composition as JTA were hot-pressed by a floating mold technique. The mold was heated in a 4-inch diameter tube furnace. Resistance heating, rather than the induction heating normally employed for manufacturing JTA, was used to obtain closer temperature uniformity. During hot pressing, an initial pressure of 2500 lb/in.² was applied and maintained until the mold temperature reached 1800°C; at that time, the selected maximum pressure was applied. All samples were held for one hour at the hot-pressing temperature. They were then cooled under maximum pressure to a temperature of 1700°C, at which time the pressure was released. These experimental billets are designated by the letter E followed by a sequential fabrication number.
- b. Temperature Series. Effects of pressing temperature on the density and microstructure were examined on samples hot pressed at temperatures ranging from 2210° to 2280°C at a pressure of 2900 lb/in.². The solidus temperature for JT composites was determined to be 2260° ±10°C by incipient melting experiments. Examination of the run-out material by metallography and X-ray diffraction disclosed a eutectic between ZrB2 and SiC. Processing and density data are given in Table II.

Figure 1 illustrates the effect of pressing temperature on density and microstructure. Density increases with increasing pressing temperature to an extrapolated temperature of 2258°C and then decreases at higher temperatures. This decrease in density is caused by some run-out of the ZrB2, SiC liquid eutectic. The 250 X magnification photomicrographs adjacent to the data points on Figure 1 show the representative structure of the material at different pressing temperatures. As the temperature is raised from 2210° to 2250°C, the porosity of the material decreases; the general physical structure and distribution of the ZrB2-SiC-C phases remain essentially unchanged. As the temperature is raised above 2260°C (the solidus temperature), some of the dense metallic phases are pressed out, causing a change in composition of the

billet. Both the porosity and the density of the billet decrease, and the structure shows the presence of an increased amount of a well-dispersed metallic phase.

TABLE II

PROCESSING CONDITIONS AND DENSITY OF
EXPERIMENTAL JT-SERIES BILLETS

Specimen Number	Processing Temperature T±10°C	Processing Pressure lb/in. <sup>2</sup>	Density g/cm <sup>3</sup>	Diameter in.
E-2#	2210	2900	2.84	1.5
E-3#	2230	2900	2.98	1.5
E-4#	2250	2900	3.11	1.5
E-5#	2270	2900	3.11	1.5
E-6#	2280	2900	3.06	
E-17	2240	2900	2.86	1.5
E-18	2260	2900	3.00	1.5
E-20	2265	2900	3.03	1.5
E-16	2290	2900	2.97	1.5
E-13	2150	3500	2.78	1.0
E-14	2150	4500	2.89	1.0
E-12	2150	5500	2.97	1.0
E-10 E-9 E-8 E-7	2210 2230 2230 2230	5500 4500 5000 5500	3.11 3.09 3.11 3.13	1.0 1.0 1.0
E-19	2260	5 <b>50</b> 0	3.15	1.0

\*Specimens E-2 through E-6 were inadvertently made with 53.7 instead of 51.9 percent metallic additive

An X-ray analysis was made on the liquid run-out from a specimen taken to 2400°C, well above the solidus temperature; the analysis showed only the presence of ZrB2, SiC, and graphite.

c. Pressure Series. The effect of molding pressure on density was examined on samples molded at pressures ranging from 3500 to 5500 lb/in. 2 at temperatures of 2150° ±10 and 2230° ±10°C.

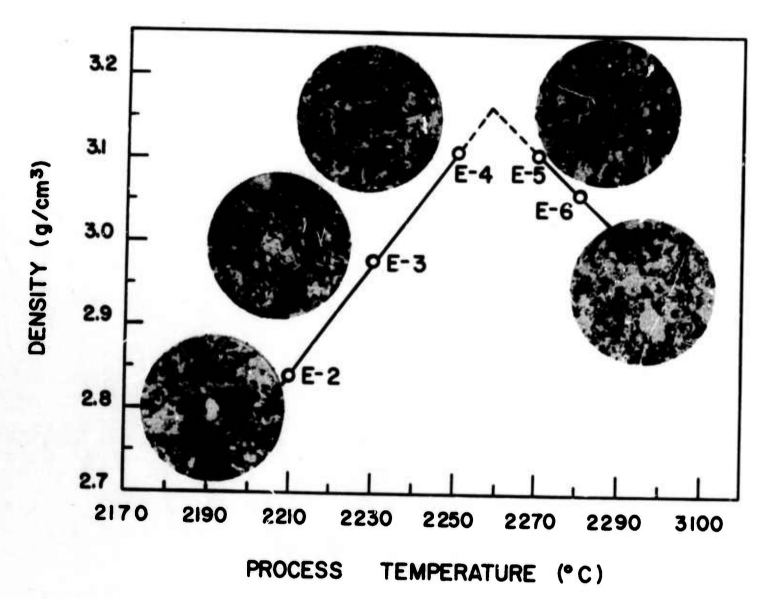


Figure 1. Effect of Processing Temperature on Density and Microstructure of JT Material of 53.7 Percent Composition. White Phase is ZrB2, Light Gray Phase is SiC, and Dark Gray and Black Phases are Graphite and Porosity.

N-9556

Figure 2 shows the dependence of density on molding pressure; the figure also shows the density change with temperature for four samples pressed at  $5500 \, \text{lb/in.}^2$ . These data indicate that, to achieve densities of 95 or higher percent of theoretical maximum density (.95 x 3.27 = 3.11 g/cm<sup>3</sup>), temperatures in excess of 2200°C and pressures in excess of 5000 lb/in.<sup>2</sup> are necessary.

d. <u>Ultrasonic Elastic Stiffness Constants</u>. The elastic stiffness constants c<sub>ij</sub> have been determined for the experimental billets from measurements of the velocity of ultrasonic pulses at

one megacycle per second frequency. Figure 3 shows the stiffness constants c<sub>11</sub>, c<sub>33</sub>, and c<sub>44</sub> and the density for a series of billets processed at successively higher temperatures. At a given density, the value of each constant is 15 to 20 percent higher if the processing temperature is above rather than below the solidus temperature. The consistently high values of the elastic constants of the initial billets (fabricated as described in Section III-4) are probably due to the maximum billet temperatures having been higher than the solidus temperature, even though the outside mold temperature was only 2130°C.

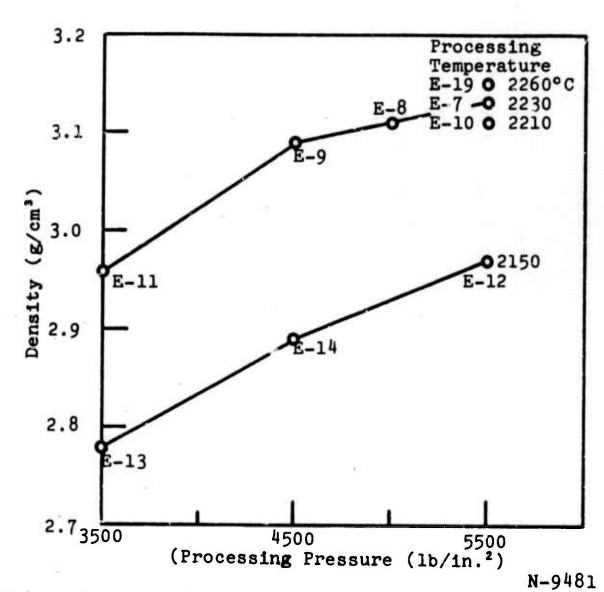


Figure 2. Effect of Processing Pressure and Temperature on Density of JT Material of 51.9 Percent Composition.

Figure 4 chows the stiffness constants c33 and c44 versus density for all the 51.9 percent composition billets fabricated at various pressures and various temperatures less than the colidus temperature (see Table II for the processing conditions). It

appears that, within the reproducibility of the experiments, the elastic constants at fixed composition depend only on the density and not on the individual processing temperatures and pressures used to achieve that density, provided that the temperature is less than the solidus temperature. This result strengthens the original premise that the JT-materials program can be based entirely on composition and density as material variables.

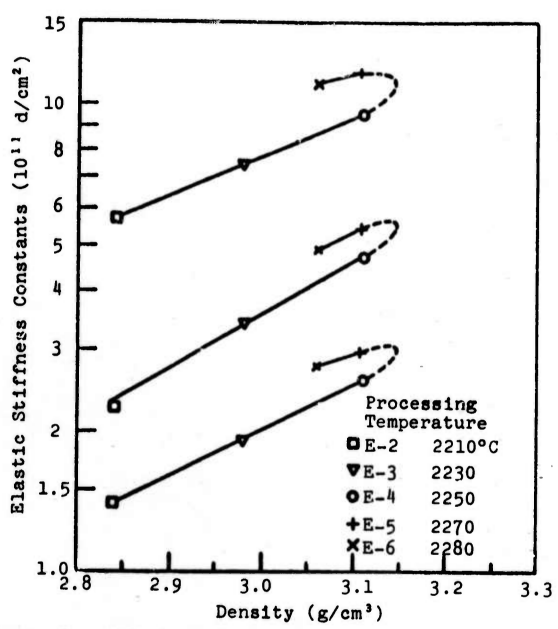


Figure 3. Effect of Processing Temperature on Density and Ultrasonic Elastic Stiffness Constants of JT Material of 53.7 Percent Composition.

N - 9479

# 6. Process Studies on 3-Inch Diameter Billets

A 40-ton press with an induction furnace capable of attaining temperatures higher than 3000°C was employed for the process studies. Figure 5 shows an overall view of the press, including the induction coil and furnace assembly.

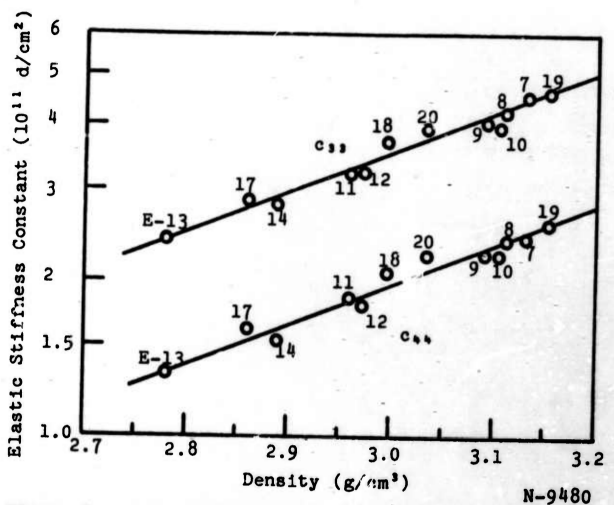


Figure 4. Ultrasonic Elastic Stiffness Constants Versus Density for JT Materials of 51.9 Percent Composition

In induction heating at high heating rates, there is a difference between the temperature of the outside surface of the mold (which acts as the susceptor) and the temperature of the billet. During normal fabrication, only the outside mold temperature is measured. Therefore, it was necessary to correlate the surface temperature of the mold to the interior temperature of the billet. This correlation was established by simultaneous temperature measurements on the mold surface and at the center of a specially prepared, prepressed billet of 51.9 percent composition. The maximum temperature difference was 140°C.

A three-inch diameter billet of JT material of 51.9 percent composition was hot pressed to 92.5 percent of theoretical density by single ram pressing. The billet was sectioned, and a density profile consisting of 15 specimens (1/2 x 1/2 x 1/2 inch) was obtained. The density variation within the billet was 3.0 percent; densities ranged from 3.01 to 3.10 g/cm³. A three-inch diameter billet of 31.5 percent composition was hot pressed by float molding. The density profile of the sectioned billet resulted in a 1.2 percent density variation; densities varied from 2.47 to 2.50 g/cm³. Thus, float molding reduced the density variation in a three-inch diameter billet from three percent to one percent.

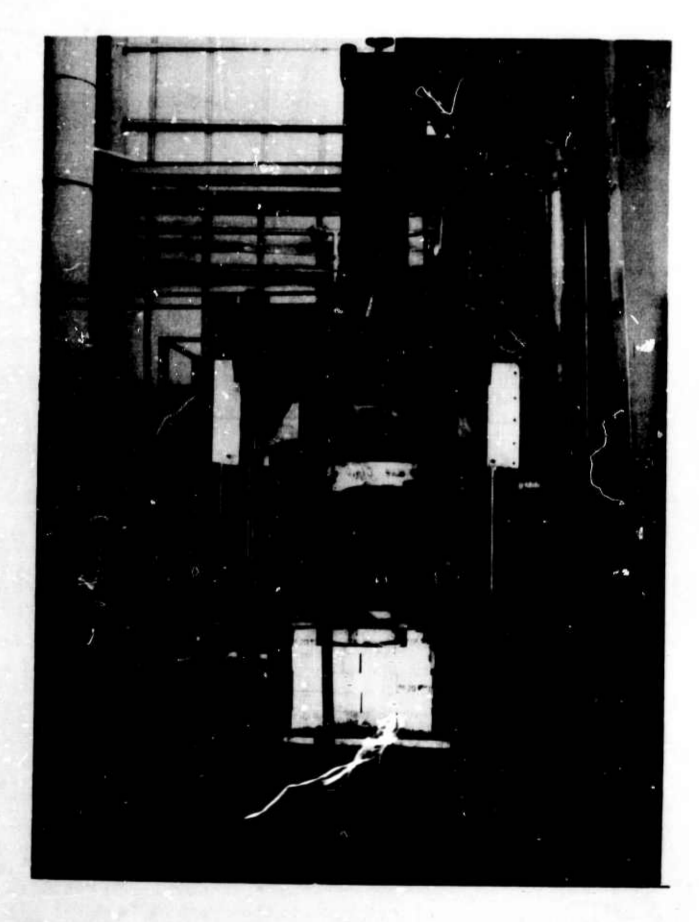


Figure 5. Forty-Ton Press and Induction Furnace Used for the Fabrication of the JT-Series Materials.

N-9557

Based on these results, additional billets of JT material were fabricated employing the float molding technique.

# 7. Fabrication of JT-Series Billets for Physical Property Evaluation

A series of JT materials was fabricated for mechanical and thermal properties evaluation. The float mold technique and carefully controlled processing conditions were employed to hot press billets three inches in diameter and approximately 3.5 inches in length. Nominal compositions of 30, 50, and 70 percent metallic additive were used; at each composition the porosity was varied from a low of two-to-six percent to a high of 17 percent. Table III lists the composition, density, and porosity of the 27 billets fabricated.

TABLE III

COMPOSITION, DENSITY, AND POROSITY OF JT-BILLETS
USED FOR PROPERTY EVALUATION

Billet Number	Composition Percent Additive	Density g/cm <sup>3</sup>	Porosity Percent
JT-30-16 JT-30-48 JT-30-40 JT-30-38 JT-30-35 JT-30-35	31.5 31.5 31.5 31.5 31.5 31.5	2.61 2.57 2.54 2.48 2.45 2.32 2.30	6.2 7.9 8.8 11.1 11.9 16.8 17.6
JT-50-46 JT-50-22 JT-50-23 JT-50-43 JT-50-44 JT-50-45 JT-50-28 JT-50-28 JT-50-27 JT-50-27	51.9 51.9 51.9 51.9 51.9 51.9 51.9 51.9	3.14 3.08 3.08 3.07 3.05 3.04 3.02 2.95 2.94 2.82 2.71	4.0 4.9 5.7 5.7 6.1 6.7 7.0 7.7 9.8 10.1 13.9 17.3
JT-70-19 JT-70-47 JT-70-20 JT-70-18 JT-70-33 JT-70-34 JT-70-37 JT-70-17	71.4 71.4 71.4 71.4 71.4 71.4 71.4	3.86 3.84 3.83 3.82 3.57 3.45 3.42 3.26	1.6 2.0 2.2 2.6 8.9 11.9 12.9

#### SECTION IV

### PHYSICAL PROPERTIES OF JT-SERIES MATERIALS

The objective of this work was to obtain physical property data which may be used as input information to stress analysis and structural synthesis calculations and for engineering applications of JT-series materials. The physical properties of interest are the elastic moduli, Poisson's ratios, stress-strain relations, fracture strengths, thermal conductivities, specific heats, and coefficients of thermal expansion.

#### 1. Methods of Measurement

- a. <u>Ultrasonic</u>. Elastic constants were determined from the velocities of propagation of longitudinal and transverse ultrasonic pulses at one megacycle per second frequency. Due to the high attenuation of the JT-series material, a through-transmission method was used with a path length of approximately 1.25 inches and with directions of propagation parallel, perpendicular, and at 45 degrees to the symmetry axis of the material.
- b. Resonant Bar. Elastic constants were determined from the frequencies of longitudinal, flexural, and torsional vibrations of bars of dimensions 1/4 x 1/4 x 3 inches. In addition, measurements were made on the static tension and flexural specimens to detect defects and unusual characteristics of the specimens prior to static testing and to provide a check of the moduli determined from the initial slope of the stress-strain curves.
- c. Tension and Compression Tests. Both longitudinal and transverse strains were measured with pairs of strain gages mounted on dog bones in tension and on rectangular prisms in compression. The dog-bone blank dimensions were  $1/4 \times 5/8 \times 2-3/4$  inches, and the gauge-section dimensions were  $1/4 \times 1/4 \times 1-1/2$  inches. The size of the compression sample was  $1/4 \times 1/4 \times 1$  inch.
- d. Flexural Tests. Four-point loading on 1/4 x 1/4 x 3 inch bars was used for flexural testing. Transverse and longitudinal tensile and compressive strains were measured at the top and bottom surfaces with strain gages. Because the load versus strain relations are nonlinear and slightly different in tension and compression, the data obtained from flexural tests were converted to uniaxial stress-strain relations by means of Naidai's graphical method. (12) The effect of frictional forces at the loading and supporting edges was taken into account in the calculation of the true stresses.

- e. Thermal Diffusivity. Thermal diffusivities were measured from ambient to approximately 600°C by a flash method. (13) The sample size was 5/8 x 5/8 x 0.04 inch.
- f. Specific Heat. Specific heats were calculated by multiplying the weight percent of each constituent phase by the specific heat per unit mass of that phase [carbon, (14) zirconium diboride, (15) and silicon carbide, (16) assuming complete conversion of Si into SiC]. The calculations were made at 100-degree intervals from room temperature to 700°C. These calculations have been checked from room temperature to 400°C by measurements made with a Perkin-Elmer Differential Scanning Calorimeter.
- of thermal expansion from room temperature to 700°C were measured by the Newton's rings method on 1/2-inch cubes. Measurements in the temperature range from 20° to 2000°C have been made on bars 2-1/2 inches in length heated in a tube furnace. Telescopes equipped with micrometer eyepieces were used to measure the elongation.

#### 2. Mechanical Properties

Room temperature mechanical property measurements on the billets listed in Table III were made by using ultrasonic and sonic resonant bar tests and by static tensile, flexural, and compressive tests. For each material composition and for each property, a graph of the property versus density and porosity was prepared on which the data points from the various test methods were plotted. Representative examples of these graphs for JT-50 material are shown in Figures 6, 7, and 8. A curve was drawn through the points, and property values were obtained from the curve at porosities of 5, 10, and 15 percent. These values are listed in Table IV.

a. Moduli. In most cases, static values of the Young's moduli are approximately 10 percent less than the sonic values. This discrepancy is due to the uncertainty in determining the initial slopes of the static stress-strain curves and, possibly, to other systematic errors. The sonic modulus results are considered to be more accurate and should be used at near-zero stress levels; the static stress-strain curves must, of course, be used at higher stress levels.

Typical longitudinal stress-strain curves for tension and compression of samples of JT-30, JT-50, and JT-70 material in the with-grain and against-grain directions are shown in Figures 9 and 10. Each curve is a composite of tension and compression data and represents the results of measurements on six samples (three tension and three compression). The range marks on each curve indicate both high and low stress values for the samples, and the

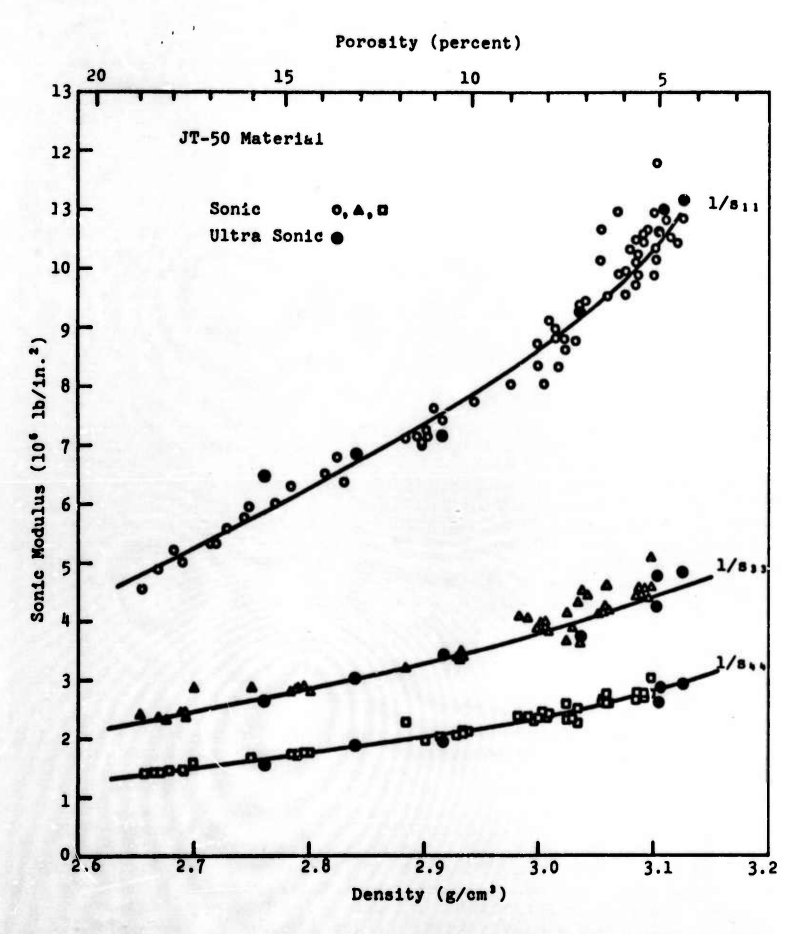


Figure 6. Variation of Sonic Moduli with Density and Porosity JT-50 Material.

N-22146

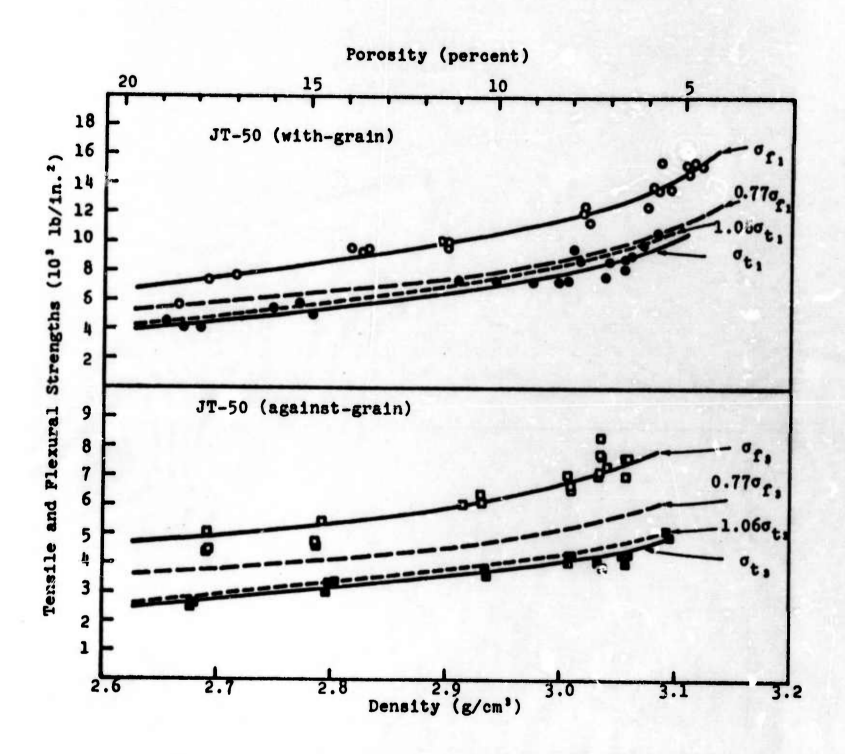


Figure 7. Variation in Tensile and Flexural Strengths with Density and Porosity for JT-50 Materials.

N-22145

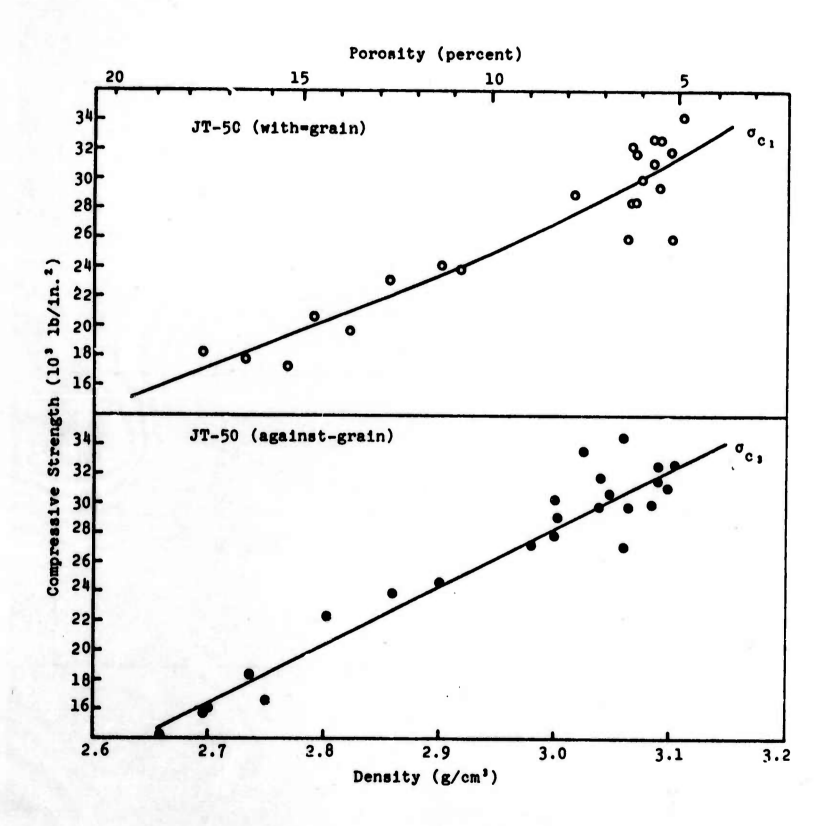


Figure 8. Variation in Compressive Strength with Density and Porcsity for JT-50 Material.

N-22147

TABLE IV SUMMARY OF MECHANICAL PROPERTY DATA FOR JT-SERIES COMPOSITES

Comp	Composition (%)		30			50			7.0	
Por	Porosities (%) Densities (g/cm³)	2.65	2.51	2.37	3.11	10	2.78	3.73	3.53	15
Young's	1/s11, sonic 1/s11, static	5.67	4.46	3.34	10.41	7.87	6.10	22.00	16.80	14.00
Modulus	1/833, sonic	2.26	1.86	1.51	4.42	3.50	2.77	11.90	9.04	13.75
	1/s33, static	2.04	1.70	1.40	3.81	3.17	2.53	11.70	8.93	7.11
Shear	1/s**, sonic	1.42	1.15	0.91	2.84	2.13	1.69	98 9	5.28	4.32
Poisson's Ratic	-812/811 -813/833	.095 .145	.089	.085	.110	.104	.099	.136	.136	.136
Tensile Strength	۵ و به	6.31	5.33	4.51	11.13	7.58	5.67	13.50	9.15	7.00
Compressive $\sigma_{c_1}$ Strength $\sigma_{c_3}$	ve d <sub>c1</sub>	17.6	15.0	12.5	32.5	25.1 26.1	19.7	70.0	56.0	41.6
Flexural Strength	of,	8.22	6.46	4.85	11.40	8.20	4.07	16.17	10.70	5.47
Units = Moduli -	duli - 106 psi; strength -		103 ps1							

curves were drawn through their midpoints. The general shapes of the curves in tension and compression are similar with compressive strengths and compressive strains being consistently higher in each of the two orientations. The tensile and compressive strengths are in each case an increasing function of the amount of metallic additive.

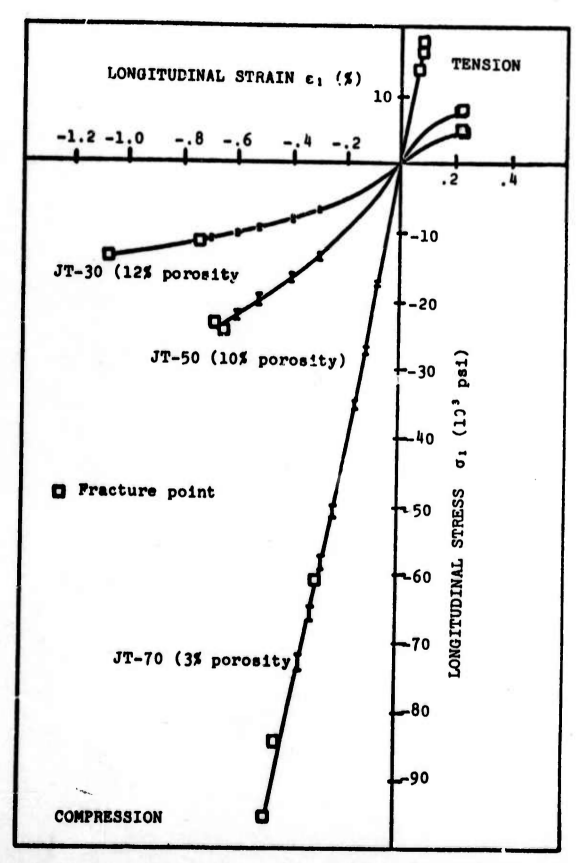


Figure 9. Stress-Strain Curves for With-Grain Samples of JT-30, JT-50, and JT-70 Material in Tension and Compression.

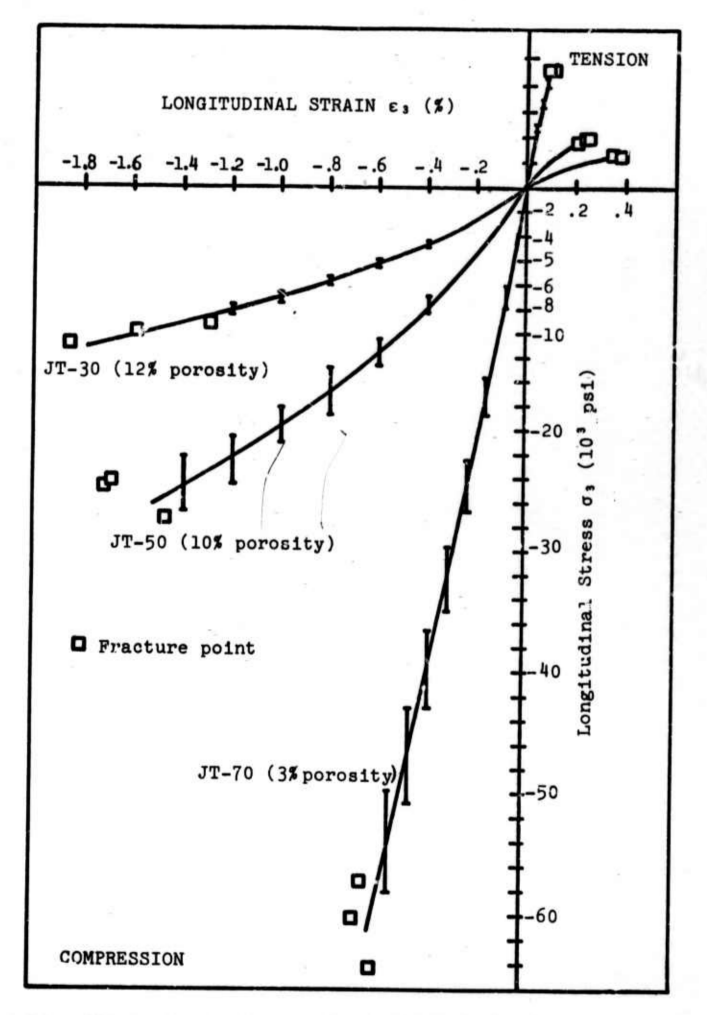


Figure 10. Stress-Strain Curves for Against-Grain Samples of JT-30, JT-50, and JT-70 Material in Tension and Compression.

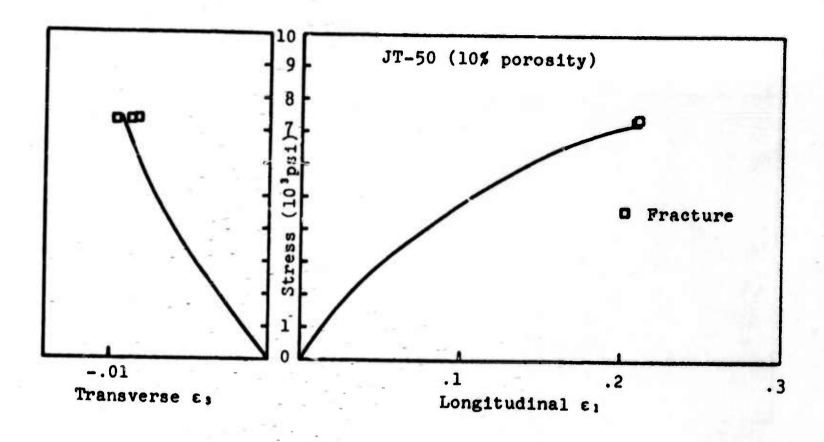
N-22150

Typical longitudinal and transverse stress-strain curves for tension and compression on samples of JT-50 material in the with-grain and against-grain directions are shown in Figures 11 and 12. The stresses are longitudinal in every case. The transverse curves have slightly different shapes not only in tension and compression, but also with orientation. In the tension curves, the transverse strains are negative indicating that when a longitudinal tensile stress is applied a contraction takes place in the transverse direction. In the compression curves the transverse strains are positive indicating that when a longitudinal compressive stress is applied an expansion takes place in the transverse direction. The transverse strains for the with-grain orientation are considerably larger than the transverse strains for the againstgrain orientation in both tension and compression. The transverse strain curves are all nonlinear, and, with the exception of one curve, have curvatures opposite to that of the longitudinal curves.

The variation of the with-grain and across-grain Young's moduli, with composition and porosity, determined by the sonic resonant bar test is shown in Figure 13. The moduli increase with increasing metallic additive composition and decreasing porosity. The variation of Poisson's ratios with composition is negligible, and the Poisson's ratios decrease slightly with increasing porosity. The range of Poisson's ratios are:  $-s_{12}/s_{11} = 0.08$  to 0.11, and  $-s_{13}/s_{33} = 0.12$  to 0.15, and  $-s_{13}/s_{11} = 0.28$  to 0.37.

The shear moduli follow the same general variation with composition and porosity as do the Young's moduli; i.e., the shear moduli increase with increasing composition and decreasing porosity. This variation of the shear moduli, determined by the sonic resonant bar test, is illustrated in Figure 14.

Strengths. The flexural test data (see results for JT 50 material, Figure 7) have been corrected by a factor of 0.77 for nonlinearity in the shear-strain curves and for friction between the specimens and the fixture. The tensile strengths, computed from load divided by cross-sectional area, have been increased by a factor of 1.06 (see results for JT 50 material, Figure 7) to allow for stray bending moments in the gauge section. Even after these corrections are made, the tensile strength is consistently less than the flexural strength by approximately 10 to 20 percent. Two possible causes of this difference are a statistical dependence of strength on test volume and stress concentrations at the fillet of the dog-bone shaped tensile specimen. The variations of flexural, tensile, and compressive strengths with composition and porosity are shown in Figures 15, 16, and 17 for both the with-grain and across-grain properties. The strengths increase with increasing composition and decreasing porosity; the variation in the compressive strength is particularly large.



Strain (percent)

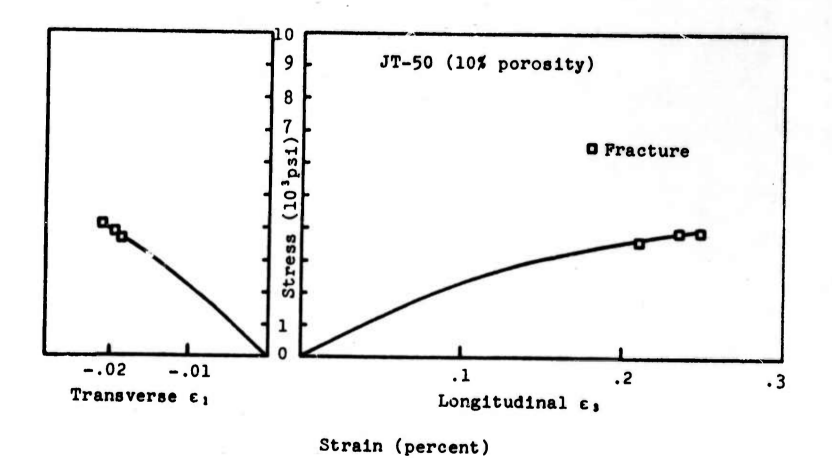
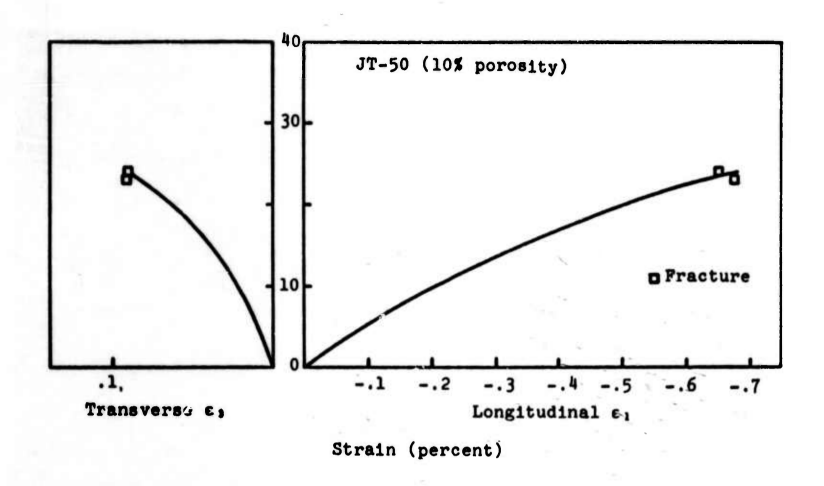


Figure 11. Stress-Strain Curves for With- and Against-Grain Samples of JT-50 Material in Tension Showing Longitudinal and Transverse Strains.



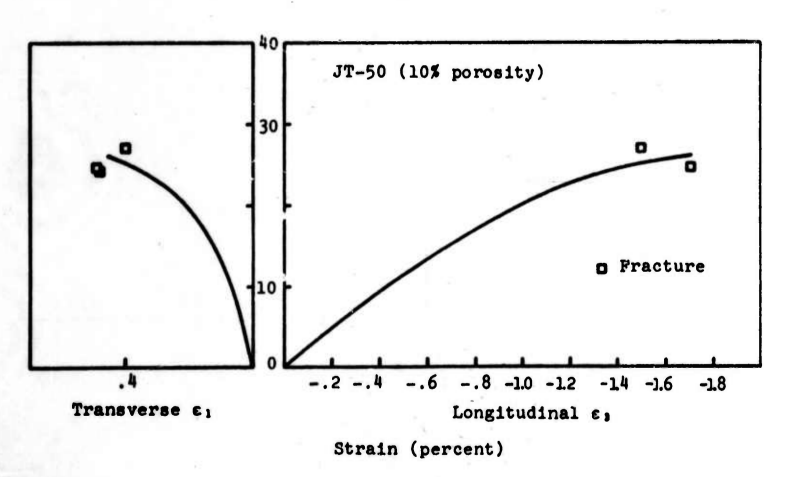
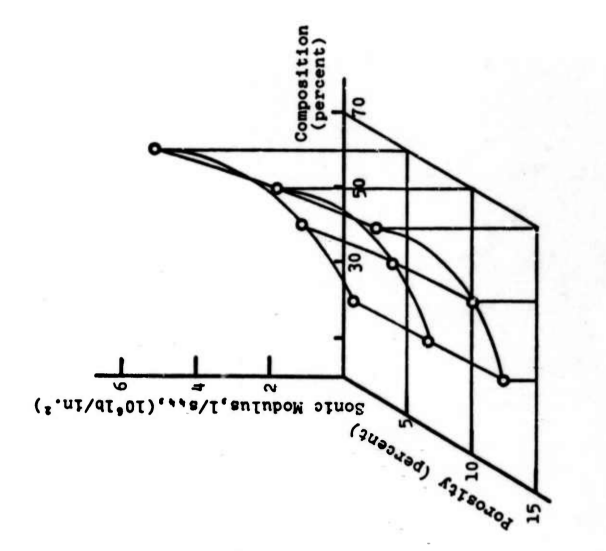


Figure 12. Stress-Strain Curves for With- and Against-Grain Samples of JT-50 Material in Compression Showing Longitudinal and Transverse Strains.



N-17224 Variation of Shear Moduli versus Composition and Porosity. Figure 14.

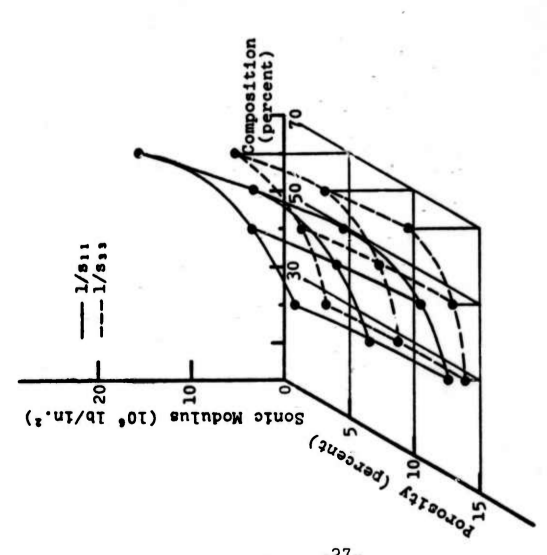
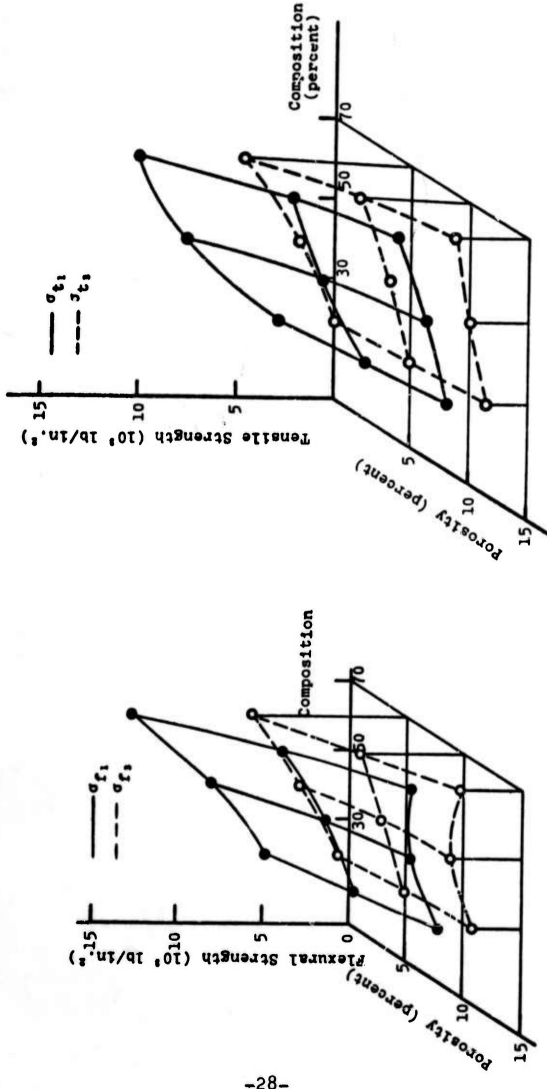


Figure 13. Variation of Young's Mcduli versus Composition and Porosity.

N-10155



N-17225 Variation of Flexural With-Grain and Across-Grain Strengths versus Composition and Porosity. 15. Figure

Across-Grain Tensile Strengths versus Composition and Porosity. Figure 16.

-28-

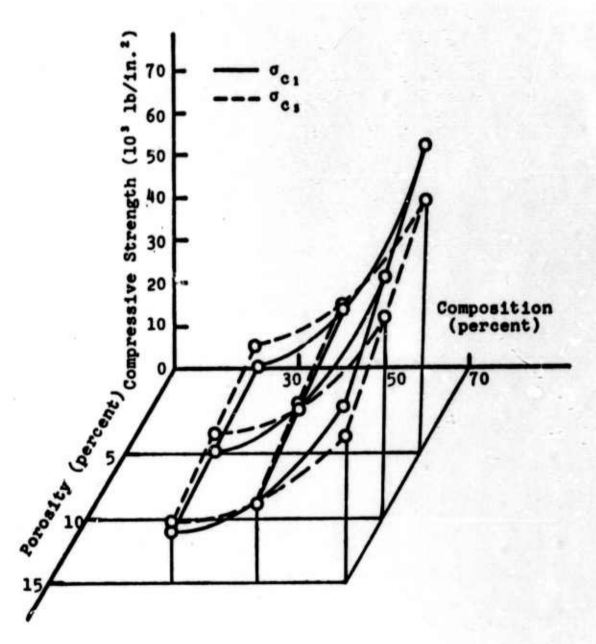


Figure 17. Variation of With-Grain and Across-Grain Compressive Strengths versus Composition and Porosity.

N-17327

c. Hoop Tensile Tests. A fixture, similar to one originally designed by the Stanford Research Institute, (17) was constructed for hoop tensile stress testing by pressurizing a short, open-end, thin-walled, hollow cylinder. The test fixture, containing a test specimen and a Neoprene bag for pressurization, is shown in Figure 18. By means of the hydraulic press, the upper and lower plates of the test fixture are pressed against the spacers which slide freely over the guiding pins. Because the length of the specimen is slightly shorter than the length of the spacers, there is a small clearance between the upper plate and the specimen; thus, the specimen is free from end loads. When the Neoprene bag is pressurized, the specimen is under pure hoop tensile stress.

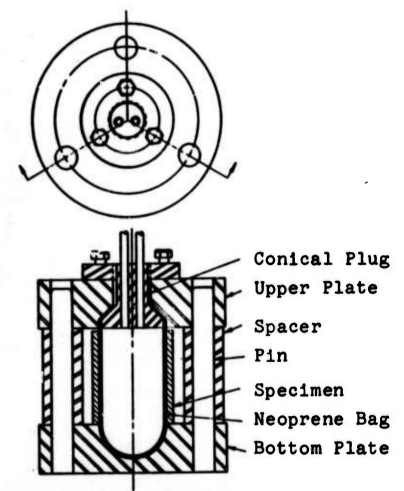


Figure 18. Specimen and Test Fixture for Hoop Tensile Stress Testing.
N-9469

Talc was applied between the specimen and the Neoprene bag to minimize friction. The inside and outside diameters of the test cylinders are, respectively, 0.885 and 0.985 inch; the length can be 0.5, 1.0, or 2.0 inches.

In order to check the test fixture, an epoxy cylinder two inches long and aluminum cylinders having lengths of one and two inches were tested to examine the effect of the specimen length and pressure loading on uniformity of strain distribution.

The strain distribution in the epoxy cylinder was investigated with a reflection polariscope, and the variation of fringe pattern was estimated to be less than 0.1 fringe. This small variation is believed to be due to variations in the properties of the epoxy cylinder itself, rather than to variations in loading stresses. Strain distributions in the 1.0 and 2.0 inch aluminum cylinders were investigated with strain gages and were also found to be uniform; the experimental and theoretical values of the strains are in good agreement.

Grade ATJ graphite cylinders with lengths of one and two inches were tested. The variation of the hoop strains measured with strain gages at four locations on the two-inch long ATJ graphite cylinder was two percent. Tensile strengths and strains obtained from these ATJ graphite cylinders are in good agreement with those obtained by the uniaxial tensile test. The tensile fracture strength and strain for a JTA-9 cylinder were 7500 lb/in. and ll80  $\mu$ in./in., respectively.

d. <u>Torsion Tests</u>. Four solid torsion specimens of JT-50 material were tested in the apparatus shown in Figure 19.

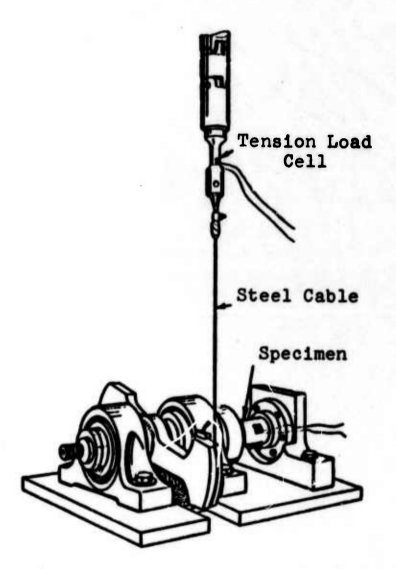


Figure 19. Torsion Testing Machine. N-10143

The overall length of the specimens was approximately 3.5 inches; the gage section was 0.75-inch in diameter and 2.0 inches in length. The cylinder axis was oriented parallel to the symmetry axis of the material. Shear strain and longitudinal strain were measured with pairs of strain gages. The shear-stress versus shear-strain curve and the shear-stress versus longitudinal-strain curve

are shown in Figures 20 and 21, respectively, for one of the specimens. The stress-strain behavior for these specimens is similar to their uniaxial stress-strain behavior: nonlinear and nonconservative.

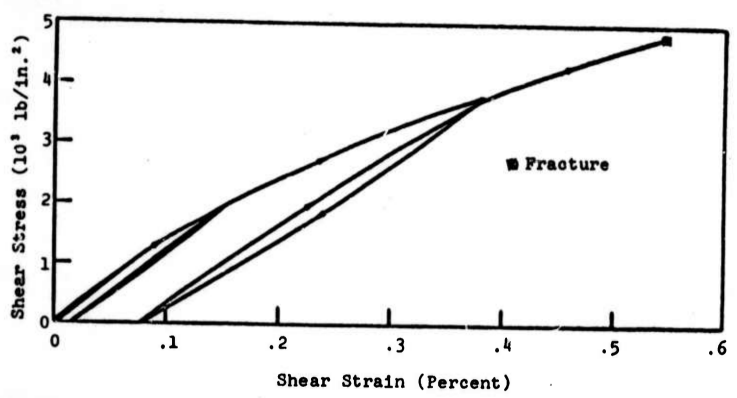


Figure 20. Shear-Stress versus Shear-Strain Curve for JT-50 Material.
N-11269

Since the torque versus shear-strain curve for JT-series composites is nonlinear and nonconservative, Naidai's graphical method(1'6) was employed to obtain the shear-stress versus shear-strain relationship from the torque versus shear-strain curve. The shear properties for these specimens are also given in Table V. The shear-strain for specimen I was not properly recorded, and, therefore, cannot be used for construction of the shear stress as a function of shear-strain curve. The nonlinearity correction on shear strength is approximately nine percent.

The shear strength tends to increase and the shear strain to decrease with increasing density. The value of the shear strength is approximately 10 to 20 percent greater than the tensile strength in the across-grain direction. The specimens exhibited a small longitudinal elongation. Length changes in plastically twisted circular metal specimens have been observed by Swift. (19) The inherent characteristics of the torsion test and the anisotropy of the materials are believed to be the source of this phenomenon. (18,19,20) The theory of anisotropic plastic flow developed by Hill(20) also supports the explanation of the length change due to the anisotropy of the materials.

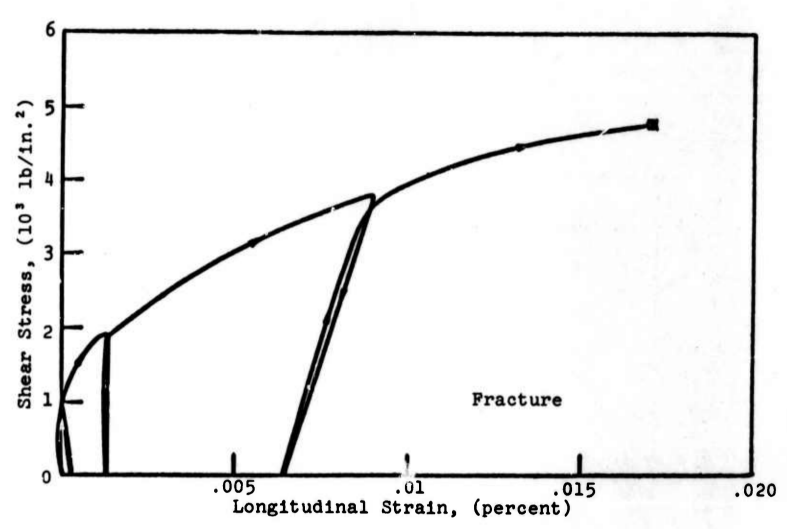


Figure 21. Shear-Stress versus Longitudinal-Strain Curve for JT-50 Composite Material.

N-11270

TABLE V
SHEAR PROPERTIES FOR JT-50 COMPOSITE MATERIAL

	Density	Shear 1/s, (10	Modulus black blac	Shear St (1b/1		Shear Strain	Long. Strain
Specimen	(g/cm <sup>3</sup> )	sonic	static	uncorrected	corrected	(percent)	(percent)
1	2.934	2.11		5750			0.01
2	2.931	2.11	1.98	5160	4740	0.55	0.03
3	3.014	2.27	2.43	6160	5630	0.44	0.03
4	3.012	2.28	2.43	6690	6080	0.48	0.04

#### 3. Thermal Properties

a. Specific Heat. The dependence of specific heat, cp, on temperature is given in Figure 22 for the nominal compositions at 30, 50, and 70 percent. The curves were calculated from the specific heats of the constituent phases. The measured points are in very good agreement with the calculated values.

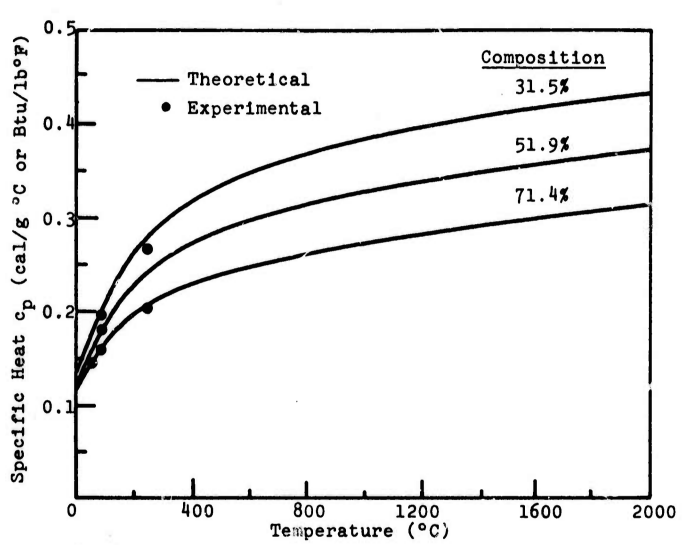


Figure 22. Specific Heat versus Temperature for JT-Material. N-18117

b. Thermal Diffusivity and Conductivity. The thermal diffusivity,  $\kappa_i$ , has been measured from room temperature to 600°C. The thermal conductivity  $k_i$ , has been calculated from the thermal diffusivity by means of the relation

$$k_{i} = \rho c_{p} \kappa_{i}, \qquad (3)$$

where  $\rho$  is the density. The thermal diffusivity and thermal conductivity results for the nominal 30, 50, and 70 percent composition and 5, 10, and 15 percent porosities are reported in Tables VI, VII, and VIII.

TABLE VI

THERMAL DIFFUSIVITY AND THERMAL CONDUCTIVITY OF JT-30 MATERIAL

.352       .179       .194       .097       .179       .091       .165         .282       .144       .195       .098       .180       .091       .164         .236       .119       .187       .093       .171       .086       .155         .205       .100       .178       .086       .161       .078       .146         .181       .087       .168       .079       .129       .129         .164       .075       .160       .073       .144       .066       .129	.186 .362 .183 .352 .179 .194 .097 .179 .091 .165 .151 .292 .148 .282 .144 .195 .098 .180 .091 .164 .157 .295 .149 .282 .144 .195 .098 .180 .091 .164 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.186 .362 .183 .352 .179 .194 .097 .179 .091 .165 .151 .292 .148 .282 .144 .195 .098 .160 .091 .164 .127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	Temperature 5% Po	The The Strain King King King King King King King Ki	Cm <sup>2</sup> (cm <sup>2</sup>   10% Po	Thermal Diffusivity (cm²/sec)  Y 10% Porosity K1	l N ×	Porosity	T 5% Porosity	The	cal/sec	os Ca	ty 15% Porosity	rosity
.186 .362 .183 .352 .179 .194 .097 .179 .091 .165 .151 .292 .148 .282 .144 .195 .098 .160 .091 .164 .151 .292 .148 .282 .144 .195 .098 .160 .091 .164 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.186       .362       .183       .352       .179       .194       .097       .179       .091       .165         .151       .292       .148       .282       .144       .195       .098       .160       .091       .164         .127       .245       .123       .236       .119       .187       .093       .171       .086       .151       .078       .146         .108       .215       .104       .205       .100       .178       .086       .161       .078       .146         .094       .191       .090       .181       .087       .168       .079       .152       .072       .136         .083       .174       .079       .164       .075       .160       .073       .144       .066       .129	.186 .362 .183 .352 .179 .194 .097 .179 .091 .165 .151 .292 .148 .282 .144 .195 .098 .160 .091 .164 .127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129		10				2	1 4	A.3	K.1	K3	K <sub>1</sub>	k 3
.151 .292 .148 .282 .144 .195 .098 .160 .091 .164 .157 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.151 .292 .148 .282 .144 .195 .098 .180 .091 .164 .127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.151 .292 .148 .282 .144 .195 .098 .180 .091 .164 .165 .187 .093 .171 .086 .155 .164 .195 .098 .180 .091 .164 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.371	.186	.362	.183	.352	179	194	100	170	.00	1,75	000
.127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .155 .108 .215 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .155 .108 .215 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.127 .245 .123 .236 .119 .187 .093 .171 .086 .155 .108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.301	.151	.292	.148	.282	771	105	800	617.	160.	COT.	900.
.108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.108 .215 .104 .205 .100 .178 .086 .161 .078 .146 .094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.254	.127	.245	.123	.236	110	187	260.	121	160.	. To4	.084
.094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.094 .191 .090 .181 .087 .168 .079 .152 .078 .146 .084 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.094 .191 .090 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129 TABLE VII	.224	.108	.215	104	205	90.	101.	660.	7)7.	990.	.155	.078
.083 .174 .079 .181 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.083 .174 .079 .164 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.083 .174 .079 .164 .087 .168 .079 .152 .072 .136 .083 .174 .079 .164 .075 .160 .073 .144 .066 .129	200	100	101	000		007.	0/1.	000.	.101	.078	.146	170.
223 .174 .079 .164 .075 .160 .073 .144 .066 .129	.083 .174 .079 .164 .075 .160 .073 .144 .066 .129	.083 .174 .079 .164 .075 .160 .073 .144 .066 .129 TABLE VII	001	160.	161.	060.	181	.087	.168	.079	.152	.072	.136	.065
			.183	.083	.174	.079	164	.075	.160	.073	144	990.	129	.059

L CONDUCTIVITY OF JT-50 MATERIAL	TUTILITIES AC -
LTY OF	
INERMAL C	
DATE .	
TTATOO TATA	

	15% Porosity	K <sub>3</sub>	100	****	0.092	0 086	000	000.0	0.073	0.068
ity	1. "	ž	ואר ס	107.0	0.164	0.157	0110	0.1.0	0.140	0.135
nduct1v	10% Porosity	K 3	אשר ט טטר ט	001.0	650.0	0.094	200	00.0		0.00
Thermal Conductivity	10% Po	K	0.1.2	1	0.374	0.175	0 160	001.0	761.0	0.147
£	5% Porosity	K3	0.200 0.117 0.132		0.117	0.107	0.100	100	160.0	0.084
	5% Po	2			0.204	0.198	0.190	0 182	707.0	0.087 0.177
	5% Porosity	7	0.191		0.153	0.128	0.111	0.097	100	0.087
ıty	15% Pc	1	0.331	470 0	+/2.0	0.235	0.207	0.186		0.172
lffusiv /sec)	Porosity		0.202	אאר ט	101.0	0.135	0.118	0.103		0.091
Thermal Diffusivity (cm <sup>2</sup> /sec)	10% Po	•	0.346	0 280	603.0	0.250	0.222	0.200	781 0	007.0
T	Porosity		0.212	0.175		0.143	0.125	0.108	900	0.030
	5% Po		0.361	0.304		0.204	0.237	0.215	1000	1010
	Temperature (°C)		100	200		300	00#	200	600	

There is a large uncertainty in the last significant figure.

THERMAL DIFFUSIVITY AND THERMAL CONDUCTIVITY OF JT-70 MATERIAL TABLE VIII

		Ē	ermal D	Thermal Diffusivity	lty.			The	ermel Co	Thermal Conductivity	lty	
Temperature (°C)	5% Por	Porosity	10% Po	ros	15% Po	Porosity	5% Por	Porosity	10% Po	Porosity	15% Po	5% Porosity
		2	1				14	h3	W.1	k3	K1	K.3
100	.324	.215	.307	.204	.292	.192	.193	.128	.174	.115	.109	.072
200	.274	.178	.257	.169	.241	.160	.192	.125	.171	.112	901.	.070
300	.238	.154	224	941.	.211	.139	.183	.118	.163	901.	.101	190°
4:00	.214	.140	. 500	.133	186	.125	.175	.114	.155	.103	.095	190.
200	.196	.131	.183	.124	.170	.115	.167	.111	.147	.100	060.	.061
009	921.	.111	991.	.108	.156	.105	.155	860.	.139	060.	.086	.058

expansion data showed somewhat greater scatter than did the other property data. This scatter masked any dependence of the thermal expansion on porosity that might exist. The thermal expansion does depend slightly on composition, but the dependence is opposite for the with-grain and across-grain directions. The coefficients of thermal expansion  $\alpha_1$  and  $\alpha_3$  are almost equal for the 70 percent composition. The trends of  $\alpha_1$  and  $\alpha_3$  with temperature up to 1500°C are shown in Figure 23 for the nominal compositions of 30, and 70 percent.

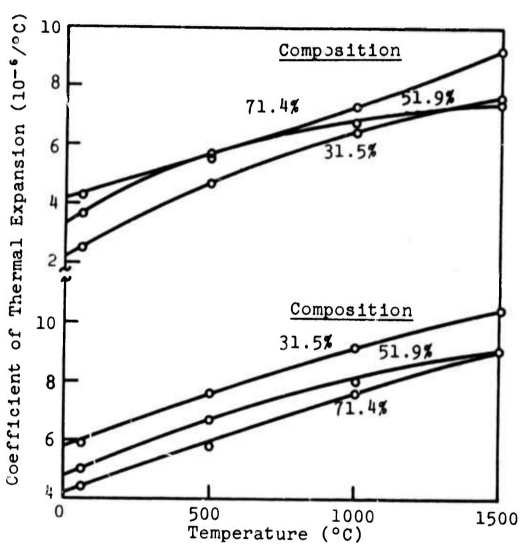


Figure 23. Coefficient of Thermal Expansion versus Temperature for JT-Material.
N-17328

# 4. Effects of Annealing on Thermal Expansion and Mechanical Properties

Irreversible dimensional changes occur during the initial heating of the JT-material above approximately 1600°C. In the across-grain direction, the material expands between 1600° and 2000°C by amounts of the same order as the intrinsic thermal expansion up to 1600°C. In the with-grain direction, the irreversible

dimensional changes are much smaller and are sometimes positive and sometimes negative. During the fabrication process, the JT-billets are heated to temperatures greater than 2000°C and cooled under pressure; stress relief apparently occurs when the billets are reheated. When the same specimens are cycled a second time, irreversible changes still occur on a reduced scale.

Two simple experiments have been performed to study the dimensional changes that occur during reheating. Specimens of JT-30, 50, and 70 material were machined to precise dimensions for the first experiment. These specimens were repeatedly cycled in an induction furnace (argon atmosphere) from room temperature to various higher temperatures. After each temperature cycle, the specimen dimensions were checked with a micrometer. Table IX summarizes the results of this experiment, and Figure 24 shows a representative plot of the data for one of the specimens (JT-50-26).

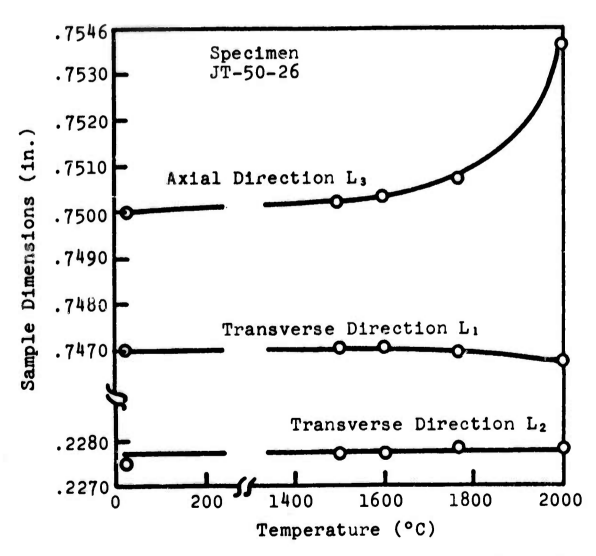


Figure 24. Room Temperature Dimensions of JT-50 Material After Repeated Anneals to Each Higher Temperature.

N-11413

In Table IX and Figure 24,  $L_3$  is in the across-grain direction and  $L_1$  and  $L_2$  are in the with-grain directions. In this experiment, several minutes were required before temperature equilibrium was achieved (the hold-time was approximately 30 seconds), whereas several hours at high temperatures were required for the coefficient of thermal expansion tests. Since approximately the same amount of growth occurs in both instances, the growth is apparently independent of these heating rates.

Several additional specimens were machined for the second experiment. These specimens were repeatedly cycled between room temperature and 2000°C, and specimen dimensions were measured after each cycle. Table X summarizes the results: after two temperature cycles, no further growth occurs in either the across-grain or with-grain directions.

The specimens should be pre-annealed or cycled several times to 2000°C before measuring the coefficient of thermal expansion at temperatures greater than 1600°C to insure that no growth will occur during measurement. This procedure should insure that only the intrinsic coefficient of thermal expansion of the material is measured above 1600°C.

A third experiment was run to determine the effect that annealing may have on mechanical and thermal properties. Two-thirds of the billet JT-30-38 was annealed at 2100°C for several hours and one-third was left as received. Specimens for mechanical and thermal tests were cut from both parts.

The solid curves in Figure 25 are the best fits through all the sonic and ultrasonic moduli data taken on JT materials. The data points plotted are for samples cut from the annealed and as-received portions of billet JT-30-38. The agreement between the plotted points and the solid curves is quite good; for the JT-30 material, no appreciable change in the moduli is apparent after the material has been annealed.

Figures 26 and 27 show similar best-fit curves and points for annealed and as-received JT-30-38 material for the flexural and compressive strength data, respectively. Again, the agreement between the plotted points and solid curves is quite good. In addition, the flexural and compressive strength data for billet JT-30-38 fall well within the experimental scatter of the data points used to determine the solid curves. Therefore, no significant change in flexural or compressive strength of JT-30 material after annealing was observed.

Thermal diffusivity data and best-fit curves are shown in Figure 28 for with-grain specimens of billet JT-30-38. These data indicate that the with-grain thermal diffusivity at every temperature increases after the material has been annealed at 2100°C.

DIMENSIONS OF JT-MATERIAL AFTER CYCLING TO VARIOUS TEMPERATURES\* TABLE IX

		A 0 0 0 0 0	7	18	lst Cycle	<b>.</b>	Ŕ.	2nd Cycle	• •	¥.	3rd Cycle	ø c	at t	4th Cycle	•
		TECON.	Į.	11	71	13	3 5	1000	i	3	L2	2	1	L2	15
JT-30-16	.7500	.7500 .2275 .7535	.7535	.7505 .2278	.2278	.7535	.7505	.2277	.7534	.7504	7504 .2277	.7538	.7502	.2277	.7581
JT-30-35	.7535	.2275	.7505	.7536	.2276	.7504	.7538	.2275	.7502	.7539	.2276	.7506	.7538	.2275	.7559
JT-50-21	.753+	.2275	.7505	.7536	.2274	.7504	.7536	.2273	.7504	.7536	.2274	.7507	.7532	.2273	.7582
JT-50-26	.7470	.2275	.750+	.7470	.2277	.7502	.7470	.2277	.7503	.7469	.2278	.7507	7467	.2278	.7535
JT-70-18	.7480	.227+	.7505	7477.	.2274	.7503	.7478	.2273	.7502	.7478	.2273	.7503	.7476	.2272	.7516
JT-70-34	.7470	. 2265	.7505	.7469	.2267	.7505	.7468	.2267	.7505	.7468	. 2267	.7508	.7467	.2266	.7531

dimensions measured in inches at room temperature.

DIMENSIONS OF JT-MATERIAL AFTER CYCLING TO 2000°C\* TABLE X

		As Received	Pe	ă Fi	1st Cycle to 2000°C	<b>0</b> C)	ά iλ	2nd Cycle to 2000°C	<b>4</b> C)	# #	3rd Cycle	di Ci	
	r.	L.	1	51	L	1	i	r.	į.	13	L2	Ls	
JT-30-16	.7500	.0418	.7501	.7501	.0418	.7527	.7501	.0417	.7530	.7500	7140.	.7530	
JT-50-21	.7501	4140.	.7504	.7499	.0415	.7541	7499	.0414	.7546	.7498	.0414	.7546	
JT-70-18	.7502	.0419	.7500	.7499	.0419	.7507	.7498	.0450	.7507	.7498	.0420	.7507	
1 dimen	stons n	leasured	*All dimensions measured in inches at room temperature.	at ro	om temp	erature.							,

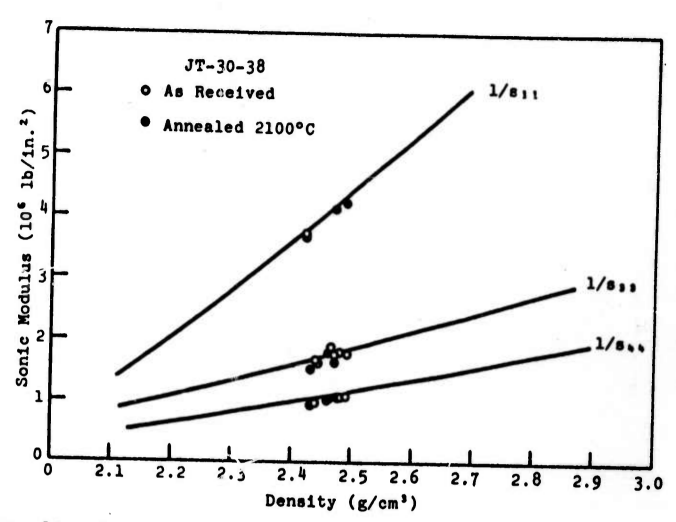


Figure 25. Comparison of Sonic Moduli Between Samples of JT-30-38 and Best Fit Through All the JT-30 Sonic Data.

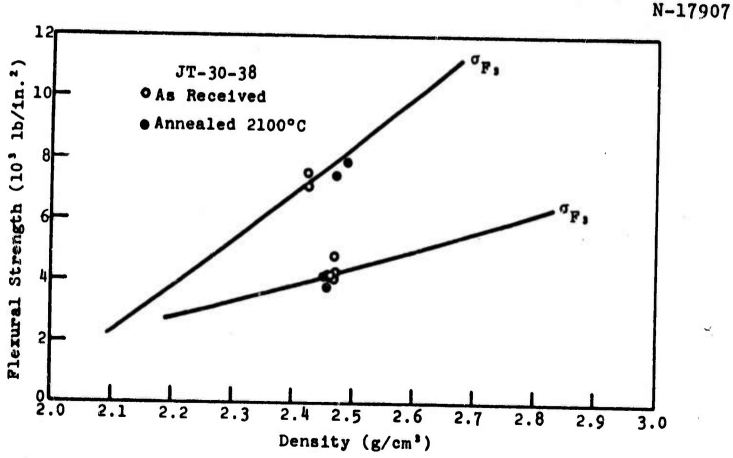


Figure 26. Comparison of Flexural Strengths Between Samples of JT-30-38 and Best Fit Through all the JT-30 Flexural Strength Data.

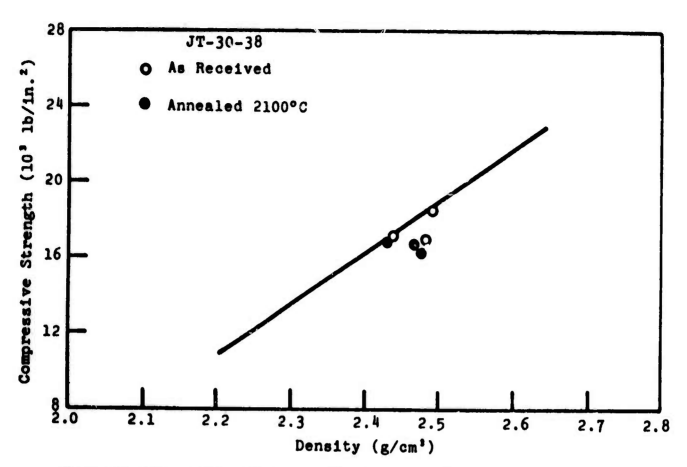


Figure 27. Comparison of Compressive Strength Between Samples of JT-30-38 and Best Fit Through All the JT-30 Compressive Strength Data.

N-17904

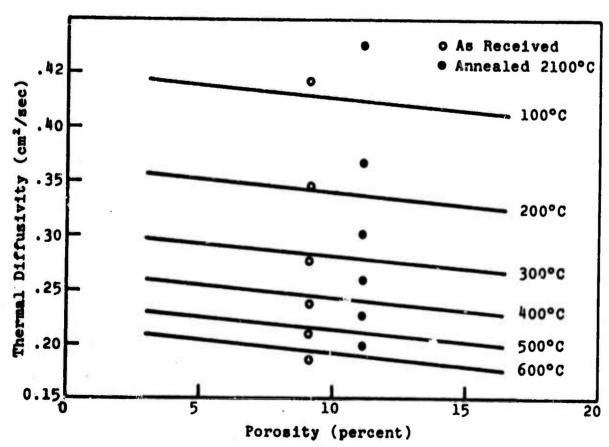


Figure 28. Comparison of With-Grain Thermal Diffusivity Between Samples of JT-30-38 and all the JT-30 Diffusivity Data.

N-17905

The increase is approximately 11 percent at 100°C and approximately five percent at 600°C. Material was available for the annealing of only one across-grain specimen. Comparison of the thermal diffusivity of this specimen with that of the best-fit curves suggests a slight increase in diffusivity after annealing of approximately five percent. Although the effect of annealing on the thermal diffusivity and mechanical properties was studied on only 30 percent additive material, there is no reason to expect that the results would be different for other composites.

#### SECTION V

## MULTIAXIAL STRESS TESTING AND FAILURE CRITERIA

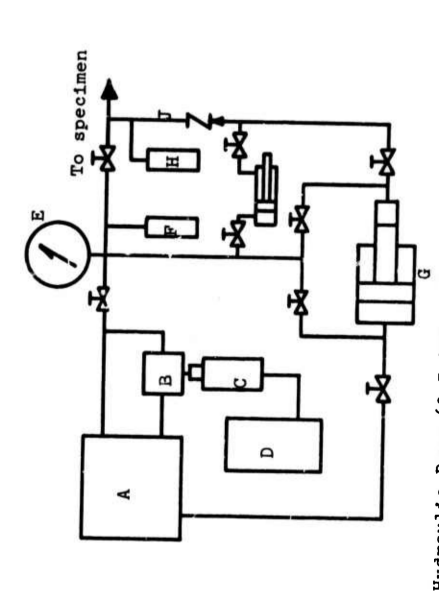
The objective of this work was to extend and complement the physical property data presented in Section IV by determining the stress-strain relations and fracture strength under various states of combined stresses. For this work, thin-walled cylindrical specimens were subjected to combinations of internal, external, and axial pressures.

# 1. Multiaxial Stress Testing Apparatus

Since the JT materials are hard and abrasive, test specimens were first machined to rough dimensions from the billets by using diamond core drills and then ground to the specified dimensions with a diamond wheel. The longitudinal axis of the test specimens was oriented parallel to the symmetry axis of the material. The test specimens had the following dimensions: inside diameter, 0.885 inch; outside diameter, 0.985 inch; and length, 3.50 inches. In general, the variation of the wall-thickness throughout the test specimens was less than 0.002 inch. A thin layer of synthetic rubber coating was applied on strain gages and specimen surfaces to prevent the hydraulic oil from damaging the strain gages and penetrating pores.

The schematic diagram of the hydraulic system for the multiaxial stress test apparatus is shown in Figure 29. The basic hydraulic power unit can supply a continuous pressure to 5000 psi. The pressure from the power unit can be further increased to 7500 psi by means of a precision hydraulic cylinder and to 50,000 psi by means of a pressure intensifier. The pressure from the power unit can be increased or decreased at prescribed rates by replacing the screw for pressure adjustment in the pressure relief valve with a threaded plunger which rotates and moves axially in a linear actuator. The rate and direction of the axial motion of the threaded plunger are remotely controlled.

The pressure chamber, specially designed for combined stress tests, is shown in Figure 30. The specimen assembly, which consisted of a test specimen and two steel end-caps, is shown inside of the pressure chamber. The interior surfaces of the steel end-caps were slightly tapered, and the steel caps were attached to the test specimen with epoxy cement. Since the modulus of epoxy cement is considerably lower than that of the test specimen, the test specimen can deform without serious end-constraints. The variation of strain due to end-constraints was approximately five percent. The strain gage wires were led out from the pressure chamber by means of connectors normally used for thermocouple wires.



Intensifier Transducer Transducer Hydraulic Cylinder Pressure Pressure Pressure H. I. Control Box for Linear Actuator Pressure Gage (0-5,000 ps1) Hydraulic Pump (0-5,000 ps1) Pressure Relief Valve Linear Actuator A. Hydraull B. Pressure C. Linear A D. Control E. Pressure Figure 29.

Schematic Diagram of Hydraulic System for Multiaxial Stress Test Apparatus.

Pressure Check Valve

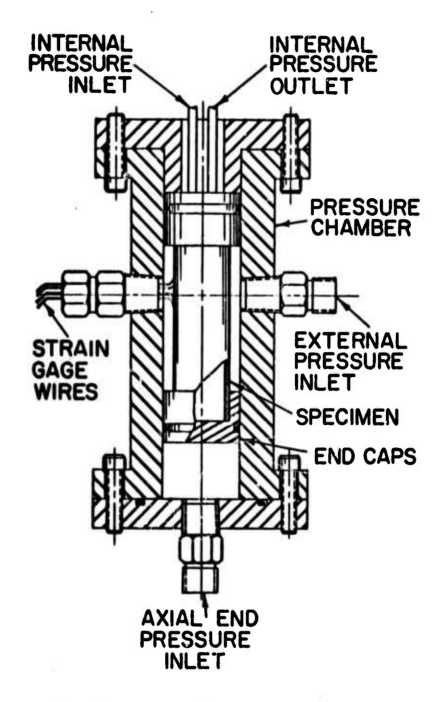


Figure 30. Pressure Chamber Assembly for Combined Stress Tests.
N-9978

Four foil strain gages, two longitudinal and two tangential, were mounted at diametrically opposite locations on the outside surfaces of most of the test specimens. The output of the longitudinal strain gages was measured individually so that any bending moment could be determined and taken into consideration in the calculation of stresses. The corrections for transverse sensitivity of the strain gages were made on the measured strains. The hydraulic power unit could supply a continuous pressure to 50,000 psi. The application of two pressures was usually necessary to obtain a

desired state of combined stresses. For most cases, the ratio of the two pressures was maintained constant to provide the so-called radial or constant stress-ratio loading condition. Pressure-strain relations and pressure-time relations were plotted by X-Y recorders.

# 2. Combined Stress Test Results for Nominal 50 Percent Additive JT-Series Composite Material

The tangential stress,  $\sigma_1$ , and radial stress,  $\sigma_2$ , were calculated by Lame's equations, and the longitudinal stress was calculated by the equation

$$\sigma_3 = \frac{a^2 P_1 + (d^2 - b^2) P_0 - d^2 P_e}{b^2 - a^2}, \quad (4)$$

where a is the inside radius of test specimen, b is the outside radius of test specimen, d is the inside radius of the pressure test chamber, F<sub>1</sub> is the internal pressure, P<sub>0</sub> is the external pressure, and P<sub>0</sub> is the axial end pressure. The radial stress, which is equal to the oil pressure at the surface of the test specimen, is small compared with the tangential and longitudinal stresses and could be neglected in most cases.

The longitudinal and tangential stress-strain curves, obtained by constant stress-ratio loading in various states of combined stresses are shown in Figure 31 (a) and 31 (b). Compressive stress and strain are indicated with a negative sign. values of stress and strain at the outside surface of the test specimen are used in Figure 31 (b). The curves designated as AT and AC in Figure 31 (a) are, respectively, uniaxial tensile and compressive stress-strain curves in the against-grain direction. The curves designated as WT and WC in Figure 31 (b) are, respectively, uniaxial tensile and compressive stress-strain curves in the with-grain direction. The material deforms considerably more in the against-grain (longitudinal) direction than in the withgrain (tangential) direction under a given compressive stress. scatter was present in the biaxial stress-strain data, particularly in the state of biaxial compression; this scatter was attributed to the variation of porosity of the test specimens and to errors in strain measurements. Strain measured by a strain gage depends slightly on the surface condition (size and distribution of pores) on which the strain gage is placed. The longitudinal and tangential stress-strain relations for a JT-50 composite specimen subjected to internal pressure are shown in Figure 32. The longitudinal strain and the tangential strain at a given internal pressure are approximately the same magnitude because of the highly anisotropic properties of the material. The nonconservative response under cyclic internal pressure loading is similar to that obtained for uniaxial

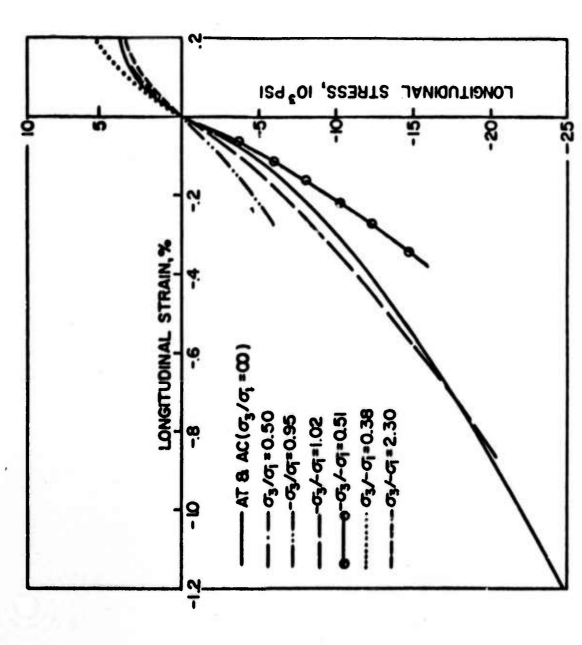


Figure 31 (a). Longitudinal Stress-Strain Curves Under Combined Stresses for the JT-50 Composite Material. N-18224

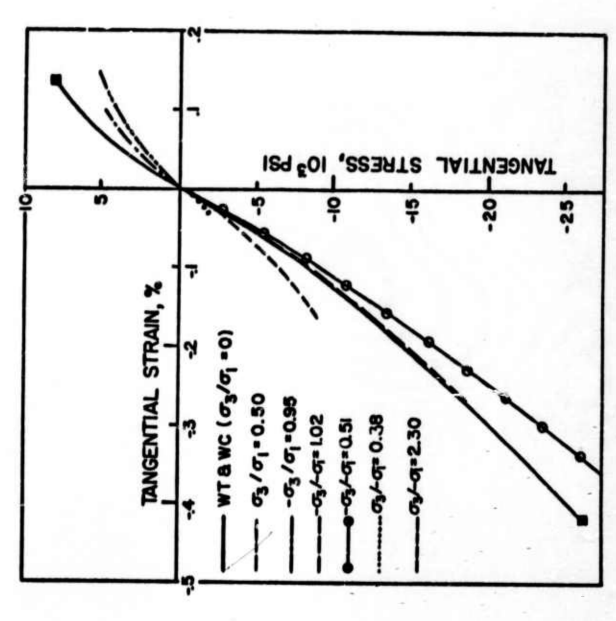


Figure 31 (b). Tangential Stress-Strain Curves Under Combined Stresses for the JT-50 Composite Material.

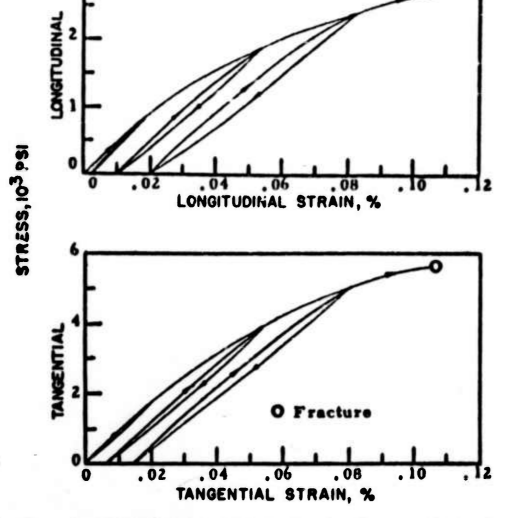


Figure 32. Longitudinal and Outside Tangential Stress-Strain Curves for JT-50 Composite Cylinder Under Internal Pressure.

N-9607

The biaxial fracture strength of the JT-50 composite is shown in Figure 33, which is a two-dimensional representation of the longitudinal stress,  $\sigma_3$ , versus the tangential stress,  $\sigma_1$ , at fracture. The radial stress,  $\sigma_2$ , is considered to have no effect in Figure 33 because of its small magnitude. The scatter in the fracture strength data in the state of biaxial compression is believed to be due primarily to the variation of percent porosity of the test specimen and premature failure initiated by localized defects. An empirical fracture criterion for the JT-50 composite is assumed to be in the form of a quadratic equation for an ellipse, and the constants in the quadratic equation have been determined from the fracture strength data by means of the least squares, best fit technique. The fracture envelope predicted by the empirical fracture criterion is shown in Figure 33, and the empirical fracture criterion\* is given as

$$(12100)^2 = \sigma_1^2 - 0.83_1\sigma_3^2 + 1.04\sigma_3^2 + 15515\sigma_1 + 20208\sigma_3.$$
 (5)

The empirical fracture criterion was obtained by Dr. C. C. Chamis of Case Western Reserve University.

The agreement between the empirical fracture criterion and fracture strength data is not completely satisfactory, indicating that the assumption of a continuous ellipse for the fracture envelope may not be accurate.

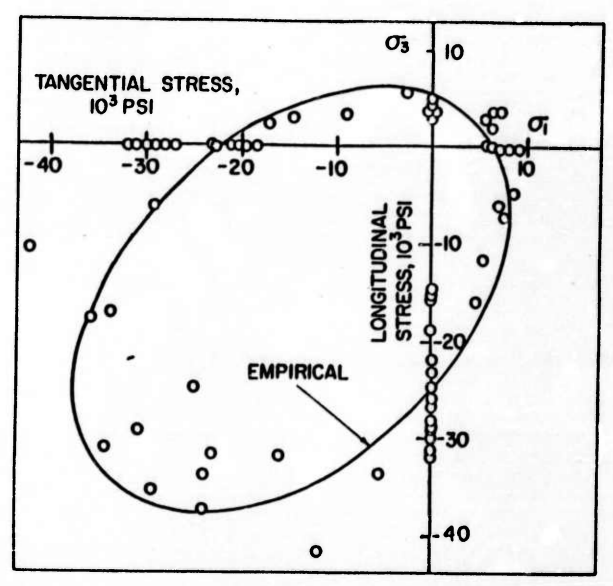


Figure 33. Biaxial Fracture Strength for the JT-50 Composite Material.

N-18505

# 3. Failure Criteria for Nominal 30 Percent and 70 Percent Additive JT-Series Composite Material

A limited number of JT-30 and JT-70 composite specimens were tested under biaxial stresses so that the approximate shapes of the fracture strength surfaces for these composites and the effects of the composition on biaxial fracture strength could be determined. Two specimens from each composite were tested; the results are given in Table XI. The biaxial fracture strength for the JT-30-49-2 specimen appeared to be lower than the estimated value. The strains measured on the outside surface of the specimen are also given in Table XI.

TABLE XI
BIAXIAL FRACTURE STRENGTH AND STRAIN OF
JT-30 AND JT-70 COMPOSITES

Specimen	Density (g/cm³)	Strength*(103 Inside Hoop our	lb/in.²) Axial σus	Strain* Hoop  Eul	(percent) Axial <sub>Eus</sub>
JT-30-48-1 JT-30-49-2 JT-70-47-1 JT-70-47-2		4.4 -13.8 16.5 17.2	2.0 -11.1 -16.6 7.7	0.15 -1.15 0.09 0.06	0.17 -0.24 -0.15 0.06
*Negative	values in	ndicate compre	ssion.		

# 4. Comparison of Theories of Fracture with the Experimental Data

A number of theories of fracture have been developed for the determination of allowable design stresses for the complicated stress conditions which occur in actual situations. By using such theories of fracture, designers are able to predict when failure will occur under combined stresses from the known fracture strength data in simple tension, compression, and shear loading or from the shape and size of cracks in the material. Maximum stress theory and Mohr's theory are, perhaps, the earliest theories of fracture which take into account different strengths in tension and compres-Several theories of fracture have been proposed more recently for materials having various types of anisotropy. (2 1-28) theories are generally in the form of a quadratic equation for an ellipse in terms of stresses, and the constants in the equation are related to uniaxial tensile and compressive strengths and shear strength in different ways. The fracture envelopes predicted by these theories of fracture were compared with the experimental data for the JT-50 composite material; most theories of fracture do not predict satisfactorily the experimental data. Among the theories which predict a fracture envelope in the form of an ellipse, the theories proposed by Stassi(23) and Hoffman(27) gave the best agreement with the experimental data.

The correlations of the experimental data with the empirical fracture criterion shown in Figure 33 and with the theories of fracture which predict a continuous ellipse for the fracture envelopes are not entirely satisfactory, since these theories do not consider the different fracture mechanisms operative under different states of biaxial stresses for this composite. Consequently, the biaxial fracture envelope of this material may be taken to be

composed of four separate fracture envelopes which are consistent with the fracture mechanism operative for each state of braxial stresses.

Norris, et al.(29)\* formulated a theory of fracture for orthotropic materials based on the assumptions that the orthotropic material is composed of walls of an isotropic material surrounding voids in the shape of equal rectangular prisms and that fracture under combined stresses is governed by the distortionenergy criterion. Norris's fracture criterion can be written in the generalized form

$$\frac{\sigma_1^2}{\sigma_{\bar{1}_1}^2} - \frac{\sigma_1\sigma_3}{\sigma_{\bar{1}_1}\sigma_{\bar{j}_3}} + \frac{\sigma_3^2}{\sigma_{\bar{j}_3}^2} = 1, \qquad (5)$$

where  $\sigma_{i_1}$  and  $\sigma_{j_3}$  are the uniaxial fracture strengths in the symmetry directions. The values of  $\sigma_{i_1}$  and  $\sigma_{j_3}$  can be either tensile strengths  $\sigma_{t_1}$  and  $\sigma_{t_3}$  or compressive strengths  $\sigma_{c_1}$  and  $\sigma_{c_3}$ , depending on whether the principal stresses  $\sigma_{i}$  and  $\sigma_{i}$  are tensile or compressive stresses. Four fracture envelopes, each of which is applicable to only a particular stress state, are obtained when the appropriate uniaxial fracture strengths for each stress state are substituted in Equation (6). Equation (6) can be obtained in a different manner through the generalization of the distortion-energy criterion to include the different strengths in tension and compression and the transverse isotropy.

Norris's theory has been applied to the fracture data for the JT-50 composite material. Good agreement between predicted and measured strengths was obtained, except in the biaxlal-tension stress state for which the predicted strength was somewhat high. The fracture envelope in biaxial tension may be best represented by the modified maximum stress theory given by

$$\frac{c_1}{\sigma_{t_1}} = 1 \text{ or } \frac{\sigma_3}{\sigma_{t_3}} = 1. \tag{7}$$

Therefore, the fracture strength of JT-series composites can be predicted satisfactorily by Equation (7) for the state of biaxial tension and by Equation (6) for the states of tension-compression and biaxial compression.

<sup>\*</sup>A series of reports pertaining to this subject have been published by C. B. Norris and his associates at the Forest Products Laboratory.

The fracture envelopes for the JT-50 composite material at five, ten, and fifteen percent porosity have been calculated with Equations (6) and (7) and the uniaxial strength data given in Table IV and are shown in Figure 34. Most of the fracture strength data fall between the fracture envelopes of ten and five percent porosity, since the porosities of the JT-50 composite billets fabricated for this program were generally slightly smaller than eight percent. The calculated fracture envelopes for JT-30, JT-50, and JT-70 composites at ten percent porosity are shown in Figure 35.

Figure 34. Biax al Fracture Strength as a Function of Porosity, Composition = 50 Percent.

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POROSITY=5%

### 5. Correlation of Scress and Strain Under Combined Stress

JT-50 Composites. In the designing of machine and structural components, the designers must be able to predict the plastic stress-strain relations under various states of combined stresses from the uniaxial stress-strain relations. These predictions have been made with some degree of success for ductile materials by correlating certain invariant characteristics of the states of stress and strain which are often defined as the effective stress and effective strain. The relationship between the effective stress and

effective strain is commonly called the generalized plastic stressstrain relation. If a generalized stress-strain relation can be established for the JT-series composites, the known solutions in plasticity theory may be applied to these composites.

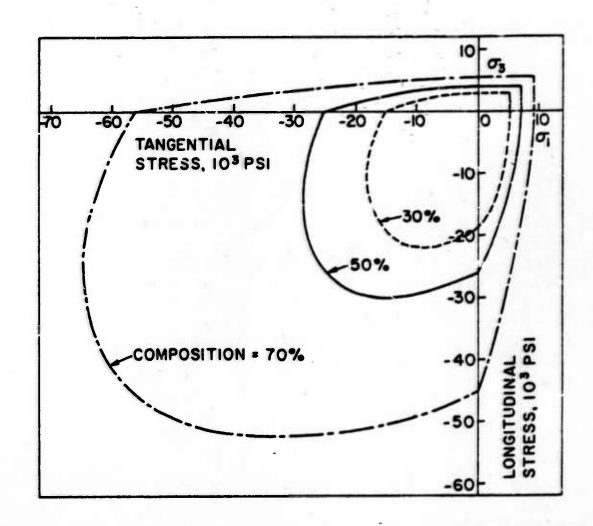


Figure 35. Biaxial Fracture Strength as a Function of Composition, Porosity = 10 percent.

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The effective stress,  $\bar{\sigma}$ , and the effective strain,  $\bar{\epsilon}$ , defined by the deformation-type theory of plasticity for ductile isotropic material are

$$\bar{\sigma} = \frac{1}{\sqrt{2}} \left[ (\sigma_1 - \sigma_2)^2 + (\sigma_2 - \sigma_3)^2 + (\sigma_3 - \sigma_1)^2 \right]^{\frac{1}{2}}$$
 (8)

and

$$\overline{\varepsilon} = \frac{\sqrt{2}}{3} \left[ (\varepsilon_1 - \varepsilon_2)^2 + (\varepsilon_2 - \varepsilon_3)^2 + (\varepsilon_3 - \varepsilon_1)^2 \right]^{\frac{1}{2}}, \quad (9)$$

where  $\sigma_1$ ,  $\sigma_2$ , and  $\sigma_3$  are principal stresses and  $\varepsilon_1$ ,  $\varepsilon_2$ , and  $\varepsilon_3$  are principal strains. Hill( $^{30}$ ) suggested a theory of plastic flow for anisotropic materials by postulating the effective stress to be given by

$$\vec{\sigma} = [F(\sigma_1 - \sigma_2)^2 + G(\sigma_2 - \sigma_3)^2 + H(\sigma_3 - \sigma_1)^2]^{\frac{1}{2}},$$
 (10)

where F, G, and H are parameters characteristic of the current state of anisotropy. Similar stress-strain relations for aniso-tropic materials were proposed by Jackson, Smith, and Lankford, (31) Dorn, (32) and Fisher(33). More recently, Hsu(34) proposed a yield criterion and plastic stress-strain relations for anisotropic materials exhibiting the Bauschinger effect. Hu, (35) starting with Hill's effective stress, obtained an expression for the incremental effective strain for materials which obey the law of constancy of plastic volume. The assumption of plastic volume constancy is approximately correct for many ductile materials and is often used in the theory of plasticity because the assumption leads to greatly simplified solutions to many problems. The assumption of volume constancy cannot be used in the study of deformation and fracture of the JT-..eries composites since significant volume changes under uniaxial stresses have been observed for these materials. The volume changes under stresses are believed to be due largely to changes in the size of the macroscopic pores and, to a lesser extent, to volume changes in the graphite particles (the particles contain microscopic porosity).

b. A Generalized Stress-Strain Relation. A generalized form of the plastic stress-strain relation for anisotropic materials can be written as

$$\varepsilon_{i} = \sum_{j} \alpha_{i,j} \sigma_{j} \phi \quad (i,j, = 1,2, --,6),$$
(11)

where the  $\alpha_{ij}$  are the constants of anisotropy and  $\phi$  is the proportionality function defined by

$$\phi = \bar{\epsilon}/\bar{\sigma}; \tag{12}$$

the expressions for  $\bar{\epsilon}$  and  $\bar{\sigma}$  are given later. The proportionality function,  $\phi$ , is a monotonically increasing function of  $\bar{\sigma}$  and describes the degree of nonlinearity in the stress-strain behavior. The constants of anisotropy,  $\alpha_{ij}$ , are assumed to be independent of deformation. The use of the total strain components is permissible if the stress-ratio is constant during deformation, the stress increases monotonically, and the material exhibits little elastic behavior. These assumptions are justified for the JT-50 composite and the present test program. There are 36 constants of anisotropy

in Equation (11). Because of the symmetric property  $\alpha_{ij} = \alpha_{ji}$  and the transversely isotropic property of the material, the number of independent  $\alpha_{ij}$  is reduced to 5, namely,  $\alpha_{11}$ ,  $\alpha_{33}$ ,  $\alpha_{12}$ ,  $\alpha_{13}$ , and  $\alpha_{44}$ . Since the material symmetry axes coincide with the principal axes of the test specimens, the stress distribution is axially symmetric and the shear stress components vanish; hence,  $\alpha_{44}$  does not appear in Equation (11). Therefore, the plastic stress-strain relations of interest become

$$\varepsilon_1 = (\alpha_{11}\sigma_1 + \alpha_{12}\sigma_2 + \alpha_{13}\sigma_3) \phi, \qquad (13a)$$

$$\varepsilon_2 = (\alpha_{12}\sigma_1 + \alpha_{11}\sigma_2 + \alpha_{13}\sigma_3) \phi$$
, and (13b)

$$\varepsilon_3 = (\alpha_{13}\sigma_1 + \alpha_{13}\sigma_2 + \alpha_{33}\sigma_3) \phi.$$
 (13c)

The stress components, in terms of the strain components, can be obtained for Equations (13) and are given by

$$\sigma_1 = (k_{11}\varepsilon_1 + k_{12}\varepsilon_2 + k_{13}\varepsilon_3)/\phi, \qquad (14a)$$

$$\sigma_2 = (k_{12}\varepsilon_1 + k_{11}\varepsilon_2 + k_{13}\varepsilon_3)/\phi$$
, and (14b)

$$\sigma_3 = (k_{13}\varepsilon_1 + k_{13}\varepsilon_2 + k_{33}\varepsilon_3)/\phi, \qquad (14c)$$

where

$$k_{11} = (\alpha_{11}\alpha_{33} - \alpha_{13}^2)/\Delta,$$

$$k_{12} = (\alpha_{13}^2 - \alpha_{12}\alpha_{33})/\Delta,$$

$$k_{13} = (\alpha_{12} - \alpha_{11}) \alpha_{13}/\Delta,$$
 (15)

$$k_{33} = (\alpha_{11}^2 - \alpha_{12}^2)/\Delta$$
, and

$$\Delta = (\alpha_{11}^2 - \alpha_{12}^2) \alpha_{33} + 2 (\alpha_{12} - \alpha_{11}) \alpha_{13}^2.$$

For a deformed body, the total work per unit volume is

$$w = \overline{\sigma}\overline{\epsilon} = \sigma_1 \epsilon_1 + \sigma_2 \epsilon_2 + \sigma_3 \epsilon_3. \tag{16}$$

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The expression for the effective stress for the present case can be obtained by the substitution of Equations (12) and (13) into Equation (16):

$$\bar{\sigma} = \left[\alpha_{11} \left(\sigma_1^2 + \sigma_2^2\right) + \alpha_{33}\sigma_3^2 + 2\alpha_{12}\sigma_1\sigma_2 + 2\alpha_{13}\left(\sigma_2\sigma_3 + \sigma_3\sigma_1\right)\right]^{\frac{1}{2}}$$

(17)

The corresponding expression for the effective strain can be derived from Equations (12), (14), and (16):

 $\overline{\varepsilon} = \left[k_{11} \left(\varepsilon_1^2 + \varepsilon_2^2\right) + k_{32}\varepsilon_3^2 + 2 k_{12}\varepsilon_1\varepsilon_2 + 2 k_{13} \left(\varepsilon_2\varepsilon_3 + \varepsilon_3\varepsilon_1\right)\right]^{\frac{1}{2}}$ 

(18)

to represent the stress-strain curves at the origin (Hooke's Law for transverse isotropy), the proportionality function  $\phi$  is defined to have the value  $10^{-6}$  in.  $^2/\text{lb}$  in the limit of zero stress and each anisotropy constant  $\alpha_{i,j}$  is chosen to be a dimensionless quantity having the numerical value of  $10^6$  times the value of the corresponding elastic compliance constant  $s_{i,j}$ ; the values of the  $s_{i,j}$  can be calculated from the data given in Table IV. The value of  $\phi$  for a given value of  $\bar{\sigma}$  can be computed from the average effective stress-effective strain curve which was determined from the uniaxial tensile and compressive stress tests. For the biaxial tests, the values of  $\bar{\sigma}$  can be calculated with Equation (17) and the values of  $\bar{\varepsilon}$  can be computed with Equation (18) by using the experimental values of  $\varepsilon_1$  and  $\varepsilon_3$  and the values of  $\varepsilon_2$  calculated with Equation (13b).

Figure 36 shows the effective stress-effective strain relations calculated for a number of test specimens, including those represented in Figures 31(a) and 31(b). The average effective stress-effective strain curve for the uniaxial stress tests is also plotted in Figure 36 (the dashed line). Some scatter exists in the correlation between the effective stress and the effective strain, but the correlation is satisfactory for engineering applications. An increase of scatter in the correlation with increasing compressive stresses is probably due to statistical variation and to difficulties in determining the values of strain under such unstable stress states. The scatter in the correlation for the combined stress test specimens is not any greater than that for the uniaxial stress test specimens. In fact, the effective stress-effective strain relations for the combined stress test specimens generally fall within a band formed by the effective stress-effective strain relations for uniaxial stress test specimens. Thus, the stress-strain relations under combined stresses can be predicted for the JT-50 composite material with the same degree of accuracy which is obtainable in the measurements of uniaxial stress-strain relations.

### 6. Microscopic Observation

a. Modes of Fracture. The influence of the high degree of anisotropy and the state of combined stresses on the modes of fracture can be observed from the photographs of the fractured specimens of the JT-50 composite material shown in Figure 37.

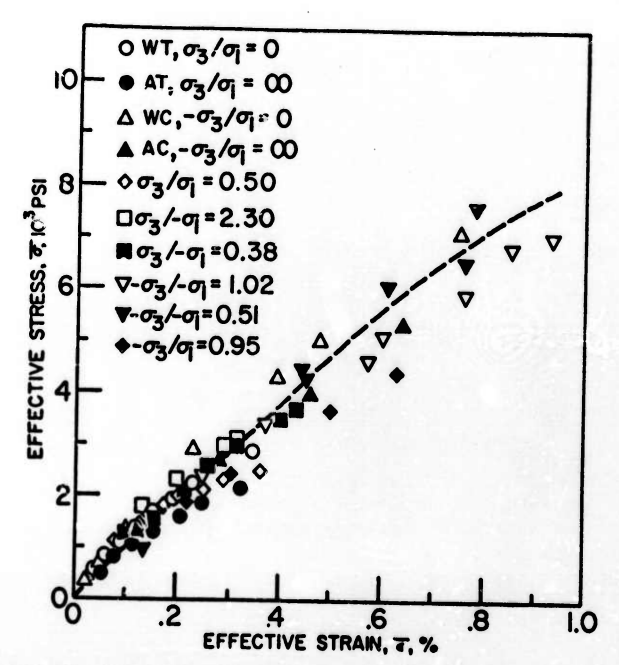


Figure 36. Effective Stress-Effective Strain Relation for the JT-50 Composite Material.

N-18397

Specimen (a) was fractured by an internal pressure, but it does not have the longitudinal crack which occurs normal to the direction of the largest tensile stress in an isotropic cylinder. The irregular fractured surface may be due to the fact that the longitudinal and tangential stresses, just prior to fracture, were approaching simultaneously the values of the uniaxial tensile strengths in the against-grain and with-grain directions of the composite, since the stress-ratio and the uniaxial tensile strength-ratio are both approximately 0.5.

The effect of a state of combined stresses on the mode of fracture is clearly illustrated by specimens (b) and (c). These specimens were fractured under an internal pressure and an axial

compressive load. The axial compressive load applied to specimen (c) was approximately twice that applied to specimen (b). The fracture of specimen (b) resulted from a barely visible longitudinal cleavage crack caused by the tangential tensile stress. The fracture of specimen (c) occurred as a result of catastrophic, random cracks. Catastrophic cracks such as those shown by specimen (c) have been observed in specimens subjected to a large compressive stress, regardless of whether the other stress was tensile or compressive.

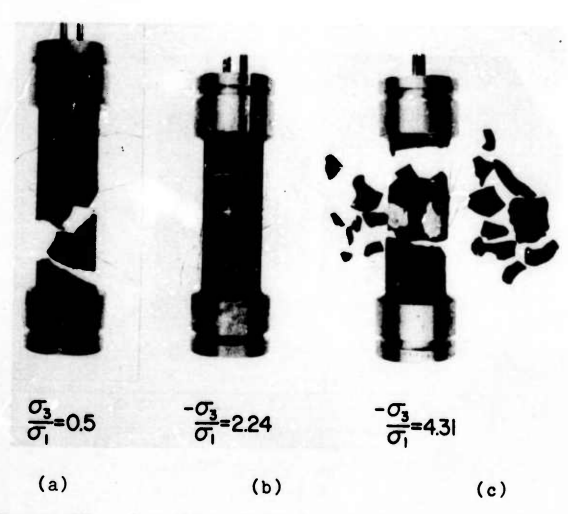


Figure 37. Specimens Fractured Under Combined Stresses.

N-20612

b. Factors Affecting Stresses. The degree of anisotropy in stress-strain properties varies with the orientation of graphite particles and the quantity of the metallic additives. The properties of the graphite particle itself are highly anisotropic, whereas the properties of the metallic additives, ZrB2 and Si, are relatively isotropic. Thus, the degree of anisotropy of the JT-series composites varies from the moderately anisotropic characteristics of graphite materials (no metallic additives) to the fairly isotropic characteristics of the ZrB2 and Si phases (all metallic additives).

Tensile strength of the graphite particles in the direction of the symmetry axis is very weak because of the low bonding forces between basal planes. This anisotropic behavior of the graphite particles and the internal stress concentrations caused by the notch effect of pores are responsible for the fact that tensile strengths are considerably lower than compressive strengths for the JT-series composites. If stresses are concentrated around the pores, fracture in those graphite particles which are oriented with their symmetry axes parallel to the direction of loading can be initiated under a relatively low tensile stress. The anisotropy in the strengths in the graphite particles, together with the preferred orientation of the particles, are largely responsible for the fact that the against-grain tensile strength of the JTseries composites is lower than the with-grain tensile strength. The compressive strength of the graphite particles in the direction parallel to the basal planes is weaker than that in the direction normal to the basal planes, probably because a condition of the low bonding forces between the basal planes leads to premature failure due to separation and buckling of the basal planes when a compressive load is applied along the basal planes. When the metallic additives are added to a graphite matrix, the compressive strength increases because the metallic additives can carry more load and also act as interlocking agents which prevent the basal planes of the graphite particles from separating and buckling prematurely. Therefore, the compressive strength in the with-grain direction of the JT-series composites increases more rapidly than that in the against-grain direction as the percent of metallic additives increases.

## SYMBOLS AND UNITS

Symbol	Unit	Description
1/811	10 <sup>6</sup> psi	Young's modulus (with-grain direction)
1/833	10 <sup>6</sup> psi	Young's modulus (against-grain direction)
1/844	106 psi	Shear modulus
ot 1	10 <sup>3</sup> psi	Tensile strength (with-grain direction)
ot,	10 <sup>3</sup> psi	Tensile strength (against-grain direction)
oc 1	10 <sup>9</sup> psi	Compressive strength (with-grain direction)
oc.	10 <sup>3</sup> ps1	Compressive strength (against- grain direction)
σ <sub>f1</sub>	10 <sup>3</sup> psi	Flexural strength (with-grain direction)
σ <sub>f</sub> ,	10 <sup>3</sup> psi	Flexural strength (against-grain direction)
$\sigma_1$	psi	Stress (with-grain direction)
σ	psi	Stress (against-grain direction)
ε1	percent	Strain (with-grain direction)
E 3	percent	Strain (against-grain direction
a <sub>1</sub>	10 <sup>-6</sup> /°C	Coefficient thermal expansion (with-grain direction)
a 3	10 <sup>-6</sup> /°C	Coefficient thermal expansion (against-grain direction)
K <sub>1</sub>	cal/sec cm-deg	Thermal conductivity (with-grain direction)
К,	cal/sec cm deg	Thermal conductivity (against-direction)
κ1	cm²/sec	Thermal diffusivity (with- grain direction)
K 3	cm²/sec	Thermal diffusivity (against-grain direction)
C <sub>p</sub>	cal/g°C	Specific heat

#### REFERENCES

- 1. Union Carbide Corporation, National Carbon Company, Research and Development on Advanced Graphite Materials Oxidation Resistant Graphite Base Composites, by K. J. Zeitsch and J. M. Criscione, WADD TR61-72, Vol. XXX, (April 1964), (AF 33(615)-6915).
- 2. Union Carbide Corporation, Carbon Products Division, <u>Tropoved</u>
  Graphite Materials for High Temperature Aerospace Use Development of Graphite Refractory Composites, by R. M. Bushong, et al, ML-TDR-64-125, Vol. II (Oct. 1964) (AF33(657)-11271).
- 3. IITRI, Properties of JTA Graphite, by Seymour A. Bortz, AFML-TR-65-178 (July 1968) (AF 33(615)-3028).
- 4. Bell Aerosystems Company, Selection Techniques for Brittle Materials--The Evaluation of TA Graphite Composites as a Structural Refractory Body, by Frank M. Anthony, et al, AFML-TR-67-78 (May 1957) (AF33(615)-2376).
- Union Carbide Corporation, Carbon Products Division, JTA Graphite Composite Manufacturing Methods, by Robert A. Reuter, AFML-TR-69-57 (March 1969) (F33615-67-C-1498).
- 6. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310 Part I (October 1966) (AF 33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).
- 7. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310, Part II (December 1967) (AF33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).
- 8. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310 Part III (January 1969) (AF 33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).
- 9. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310 Part IV Volume I (September 1969) (AF 33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).

#### REFERENCES (Cont'd.)

- 10. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310 Part IV Volume II (October 1969) (AF 33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).
- 11. Union Carbide Corporation, Carbon Products Division, in Association with Case Western Reserve University and Bell Aerosystems Company, a Textron Company, Integrated Research on Carbon Composite Materials, AFML-TR-66-310 Part IV Volume III (April 1970)(AF 33(615)-3110, Air Force Materials Laboratory, Wright-Patterson Air Force Base, Ohio).
- 12. A. Naidai, Theory of Flow and Fracture of Solids, Vol. I, (McGraw-Hill Book Company, New York, 1950), pp. 353-359.
- 13. W. J. Parker, R. J. Jenkins, C. P. Butler, and G. L. Abbott, "Flash Method of Determining Thermal Diffusivity, Heat Capacity, and Thermal Conductivity," J. Appl. Phys. 32, 1979-1984 (1961).
- Union Carbide Corporation, National Carbon Company, Research and Development on Advanced Graphite Materials Survey and Analytical Representation of the Measurements of the Specific Heat of Graphite, by G. B. Spence, WADD TR 61-72, Vol. XLI, (November 1963), (AF33(615)-6915).
- 15. Battelle Memorial Institute (Columbus), Refractory Ceramics of Interest in Aerospace Structural Applications—A Materials Selection Handbook, by J. R. Hague, J. F. Lynch, A Rudnick, F. C. Holden, and W. H. Duckworth, RTD-TDR-63-4102 (1963), p. 55, (AF33(657)-8326).
- 16. A. Goldsmith, T. E. Waterman, and H. J. Hirschhorn, <u>Handbook of Thermophysical Properties of Solid Materials</u>, Vol. III: <u>Ceramics</u>, (MacMillan Company, New York, 1961), p. 925.
- 17. Loc. cit., Reference 12, pp. 347-352.
- 18. H. W. Swift, "Length Changes in Metals Under Torsional Overstrain," Engineering, 163, 253-257 (1947).
- 19. R. Hill, The Mathematical Theory of Plasticity, (Oxford University Press, London, 1950) pp. 325-328.

#### REFERENCES (Cont'd.)

- 20. R. C. Grassi and I. Cornet, "Fracture of Gray-Cast-Iron Tubes Under Biaxial Stresses," J. of Appl. Mech. 16, Trans. ASME 71, 178-182 (1949).
- 21. H. W. Babel, "Biaxial-Fracture Strength of Brittle Materials," Technical Report AFML-TR-66-51, March 1966.
- 22. Joseph Marin, "Theories of Strength for Combined Stresses and Nonisotropic Materials," <u>Journal of Aeronautical Sciences</u>, Vol. 24, No. 4, April 1957, pp. 265-269.
- 23. F. Stassi-D'Alia, "Limiting Conditions of Yielding of Thick Walled-Cylinders and Spherical Shells," U. S. Army European Research Office (9851DU), Frankfort, Final Status Report Contract DA-91-591 EVC-1351 01-4263-61, November 24, 1959.
- 24. K. V. Zakharov, "Strength Criteria for Laminates," Plastmassy, Izd-vo Khimiya No. 8, 1961, pp. 59-62.
- John E. Griffith and William M. Baldwin, "Failure Theories for Generally Orthotropic Materials," <u>Development in Theoretical and Applied Mechanics</u>, Vol. I, Proc. of the First Southeastern Conference on Theoretical and Applied Mechanics, Plenum Press, New York, 1963.
- 26. I. I. Gol'denblat and V. A. Kopnov, "Strength Criteria of Anisotropic Materials," Izv. Akad, Nank SSSR, Mekhanika No. 6, 1965, pp. 77-83.
- 27. Oscar Hoffman, "The Brittle Strength of Orthotropic Materials, Journal of Composite Materials, Vol. I, No. 2, April 1967, pp. 200-205.
- 28. L. Fisher, "Optimization of Orthotropic Laminates," <u>Journal of Engineering for Industry</u>, Trans. ASME, Vol. 89, Ser. B, No. 3, August 1967, pp. 399-402.
- 29. Charles Brazer Norris, "Strength of Orthotropic Materials Subjected to Combined Stresses," FPL Report No. 1816, 3rd ed., Madison, Wisconsin, U. S. Forest Products Laboratory, 1962.
- 30. R. Hill, The Mathematical Theory of Plasticity, Oxford University Press, London, England, 1950, pp. 315-321.
- 31. L. R. Jackson, K. F. Smith, and W. T. Lankford, "Plastic Flow in Anisotropic Sheet Metal," Metals Technology, T. P. 2440, August 1948.

### REFERENCES, (Cont'd.)

- 32. J. E. Dorn, "Stress-Strain Relations for Anisotropic Plastic Flow," <u>Journal of Applied Physics</u>, Vol. 20, January 1949, pp. 15-20.
- 33. J. C. Fisher, "Anisotropic Plastic Flow," Trans. ASME, Vol. 71, 1949, pp. 349-356.
- 34. T. C. Hsu, "A Theory of the Yield Locus and Flow Rule of Anisotropic Materials," <u>Journal of Strain Analysis</u>, Vol. 1, No. 3, 1966.
- 35. L. W. Hu, "Studies on Plastic Flow of Anisotropic Metals,"

  Journal of Applied Mechanics, Vol. 23, ASME Paper No. 55-A-79,

  September 1956, pp. 444-450.

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Three series of graphite based, refractory composites compositionally related to commercial Grade JTA, but containing 30, 50, and 70 weight percent additives (ZrB2 and Si)2 were prepared. Within each series, the porosity was varied from approximately 2 to 17 percent. Property measurements were carried out as a function of composition and porosity and along the two principal symmetry axes, i.e., normal and parallel to the molding direction. Property measurements carried out at room temperature consisted of tension, compression, flexural, torsion, ultrasonic, and resonant bar tests. The specific heat, thermal conductivity, coefficient of thermal expansion, and effect of high temperature annealing were also determined. Stress-strain relations and fracture strength under multiaxial stress were determined for the material with 50 percent additive. A generalized stress-strain relation for a transversely isotropic, nonlinear material has been obtained without assuming volume constancy. Various theories of fracture for anisotropic materials were compared with the fracture strength of JT materials. The comprehensive property data compilations in this report should provide designers with sufficient information for structural applications of JT materials.

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