AD-757 765

GLASSY CARBONS
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Prepared for :

Advanced Research Projects Agency January 1973

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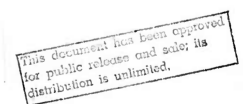
U. S. DEPARTMENT OF COMMERCE 5285 Port Royal Road, Springfield Va. 22151 Semi-Annual Progress Report for the Period June 1, 1972 to December 31, 1972

Glassy Carbons

January 1973

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#### GLASSY CARBONS

Semi-Annual Progress Report for the Period June 1, 1972 to December 31, 1972

January 1973

ARPA Order Number: 1824 Program Code Number: 1D10

Contractor: The Regents of The University of Michigan

Effective Date of Contract: 1 June 1972 Contract Expiration Date: 31 May 1973

Amount of Contract: \$162.580

Contract Number: DAHC15-71-C-0283

Principle Investigator: Professor Edward E. Hucke

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Engineering

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#### SUMMARY

A large number of glassy carbon samples was produced by controlled pyrolysis of furfural alcohol resins, examined for structural differences, and evaluated for physical and mechanical properties. It is concluded that glassy carbon is a family of materials whose properties can be varied over a significant range, depending both on the small scale (2-50Å) carbon structure, and the amount and distribution of porosity which may be present in substantial amounts.

The combination of availability, chemical stability, low density and high strength make glassy carbons look promising for mechanical applications, particularly at very high temperatures. In terms of strength per unit weight, glassy carbons compare favorably with the best available materials even at room temperature. Very fine pored (<100Å) glassy carbons have been produced in section thicknesses of 2 inches in processing times of six days, which significantly increases the range of potential application.

The carbon's fine structure determined by electron microscopy, electron diffraction and X-ray diffraction is not homogeneous on a size scale below 100 Angstroms. The material is paracrystalline with the crystallite size ranging from 10-100Å depending on processing, and with some very crystalline features occasionally existing in sizes up to 500Å.

A thermodynamic method has given results yielding a direct measure of the degree of order in terms of the configurational entropy and enthalpy relative to crystalline graphite.

Helium, xylene, mercury intrustion, small angle X-ray scattering, surface area analysis, and scanning electron microscopy show the pore structure of glassy carbons may be either isolated or interconnected, and in a size range from 5Å to 50 microns. Mechanical strength has an inverse relation to the pore size. The pore structure allows a decrease in density, as well as the opportunity to vary mechanical properties and provide chemical filtering and absorption appliances with substantial strength.

#### GLASSY CARBONS

### I. Introduction

This report covers work carried out during the period June 1972 to December 1972. Results of the previous contract periods are summarized in two previous semi-annual reports. 1,2 Since various property evaluations are being carried out simultaneously, the data tables included in this report are cumulative and have been revised to reflect additional samples as well as corrected in certain instances where samples were either improperly identified or errors in data reduction occurred. Complete data tables are given in Appendix A.

Glassy carbons have already been shown to have an unusual set of properties, particularly with respect to inertness and high strength to weight ratio. These attributes, together with the ready availability and high temperature capability of carbon naturally lead to potential applications in a wide variety of extreme conditions, such as reentry shields. Aside from mechanical applications, the unusual inertness in the human body has lead to a wide variety of biomedical applications, while its electrical properties have shown experimental promise as a semi-conductor switch, and its molecular sieve properties suggest applications in chemical separations.

Glassy carbon is still a new material and at present is

costly and available only in thin sections. The high cost and section limitation both stem from the need to use very long pyrolysis cycles in order to obtain material without cracks.

A major objective of this research has been to achieve larger section bodies with rapid processing times.

While all of the glassy carbons made from a variety of polymers and gaseous precursors are hard, strong and light, many subtle variations exist and considerable tailoring of properties is possible. A major difficulty in comparing the various carbon materials is that no simple criteria are available to distinguish one from the other. In general, many of the physical and mechanical properties show significant differences and one can give a comparison of one material with another only by specifying a complete property set. All of the properties derive ultimately from the structure and therefore it is important to be able to elucidate, control, and specify the structure of the material. The lack of a well-defined crystalline structure severely complicates matters since not only are the usual microstructural features (.1 to 100 microns) important, but also variations in the ultrastructure (3 to 100 Angstroms) are encountered. Even a seemingly straight-forward property such as real density becomes elusive because it can vary on size scale of the order of 50A.

With an understanding of structural variations possible, extension in the range of properties achievable can be made.

From data previously available in the literature and the early

results of this program<sup>2</sup>, it has become obvious that "glassy carbon" is not a single material, since even though it contains essentially only carbon atoms, its structure can be varied at all size levels. It seems more appropriate to think of glassy carbon as a material that may have short range atomic coordination with a variable state of crystallinity; but in addition, where rather large fractions of thermally stable voids can be arranged at size levels from 5 Angstroms to 50 microns. This accounts for the extremely wide variation in properties that can be achieved. The incorporation of voids into materials is not unique. However, it is unique to have up to 30% void remain stable in material at temperatures close to sublimation when the pore size is well below 100 Angstroms.

Glassy carbons possess some crystallinity on a scale of the order of 100 Angstroms or less, but the perfection of the crystallites is very poor by usual standards; and while the crystalline perfection may improve after heating to very high temperature, a well-defined graphitic structure is not achieved without unusual methods. The structure is better described as para-crystalline in the sense used to describe certain polymers. In fact, the glassy carbon structures are best considered polymers with vanishing contents of other than carbon atoms. The carbon structures are related to the polymer structures from which they evolve during pyrolysis. This observation follows from the differences obtainable in structure and properties of carbons after heating to 3000°C by merely changing the thermal history

of the polymer in a temperature range below 100°C where polymer structure is being formed. Thus, even though most of the atoms present in the precursor polymer are later removed, the resulting carbon inherits part of the structure.

The present study is largely concerned with structures obtainable together with the simultaneous measurement of certain key physical and mechanical properties. It is desired to achieve in different samples as wide a variation in structure and properties as possible.

The structure is being investigated from the viewpoints of the solid structure and the pore structure, which of course are related. Since significant differences can be induced in both over a size range of more than four orders of magnitude, no single technique is sufficient. Instead, combined techniques must be used. Each is discussed in detail in the following sections of the report. Representative property measurements are being carried out in conjunction with the structural examinations.

Due to the large number of samples processed, complete structural and property evaluations are not attempted on every sample. Apparent density, real density, wide angle X-ray diffraction, scanning electron microscopy, and strength measurements are made on every suitable sample. Small angle X-ray scattering, selected area electron diffraction, transmission electron microscopy, thermodynamic analysis, surface area analysis, mercury porosimetry, electrical resistivity, hardness, modulus of elasticity, and internal friction are carried out on a selected

small number of samples.

#### Materials Preparation II.

During this report period over 500 samples of glassy carbon were prepared with over 95 different processing conditions. The work has continued to concentrate on furfural alcohol and a furfural alcohol resin, Durez 16470\* as the carbon yielding material, with para-toluene sulfonic acid (PTSA) as the polymerization catalyst. However, a limited number of samples has been prepared with other type resins (Varcum 8051 and 4048)\*\* and other catalysts made experimentally by the Quaker Oats Company. While glassy carbons are obtained in all cases, the yield of good material and response varies for each system. Thus far, the best results have been obtained with the Durez-PTSA system, although this may be only due to the greater experience with this system.

Catalyst levels from 2 to 20% by weight based on the monomer or resin content have been explored. Temperatures for addition of the catalyst and subsequent curing time and temperatures have been varied from several minutes to several weeks with temperatures from -11°C to 150°C. In all cases where a particularly interesting set of properties has been found, a duplicate set of specimens has been made to check reproducibility. In general, reproducibility is fair, but only if meticulous care is exercised to exactly reproduce <u>all</u> conditions. Standard sample cylinders

<sup>\*</sup> Hooker Chemical Company, North Tonowanda, New York

<sup>\*\*</sup>Reichhold Chemical Company, White Plains, New York

approximately 3 cm in diameter and 20 cm in length have been adopted. Samples up to 15 cm in diameter have been produced from coarser pored material, while a few samples of 5 cm have been made from very fine pored (<100Å) material. The latter sample is far thicker than any other similar material previously reported even though its pyrolysis time of 6 days is much shorter than usual commercial practice.

The yield of crack free material has continued to improve due mainly to increased care in catalyst addition and prevention of trapped bubbles prior to and during casting. The presence of bubbles has been found to cause cracking in the subsequent pyrolysis of the samples. In most cases, it has been possible to perform a reduced pressure de-bubbling operation after addition of the catalyst.

In order to yield good material, it has been necessary to perform the hardening of the resin systems with a minimum temperature gradient throughout the piece, which often requires cooling after addition of the catalyst. The length of time curing, curing temperature and catalyst level have been varied over wide ranges.

The single-most important factor in obtaining crack free materials in pyrolysis is obtaining a uniform slow heating rate. While heating rates have been varied from 5 to 50°C per hour in the range of room temperature to 1000°C, the yield of crack free material is much greater at the slower rates. Rates up to 200°C per hour appear to cause no difficulty in heating material previously pyrolyzed to at least 700°C. Both flowing nitrogen and

reduced pressure pyrolysis have been employed. Thus far, flowing nitrogen appears to yield less difficulty with respect to cracking.

On selected samples, carbon recovery and shrinkage data have been gathered. In both cases the data are comparable with those of the literature, namely a carbon recovery based on the total resin weight of about 55% and linear shrinkage of about 29%. However, differences in both shrinkage and recovery of about 10% can be noticed with varying of curing and pyrolysis schedule.

Inhomogeneities of about lµ in diameter in the starting resin were noted in the previous report<sup>2</sup>. These regions carry through the pyrolysis and cause non-uniformity in the final structure. At present no completely satisfactory method has been found for eliminating this problem, although filtering the starting resin has made an improvement.

Rather subtle differences exist in the cured resins which result in different properties after pyrolysis. Thus far no satisfactory means for quantitatively studying the resin have been developed. Exploratory work using scanning calorimetry has been initiated in an effort to see if the differing final results can be detected from the thermal changes occuring in the polymer stages.

The single-most important processing variable is the maximum pyrolysis temperature (HTT) which has long been known to affect the structure and properties. However, all the other processing variables have some influence even though at this time the data show no clear-cut correlations. Part of the difficulty

in finding a correlation is undoubtedly due to the fact that the exothermic polymerization causes heating of parts of the sample above the temperatures of intended control. It is also probable that many of the effects noted in mechanical properties may be due to the existence of very fine cracks or possibly residual macroscopic stresses in specimens that appear to be macroscopically sound. In this case, a particular processing change may appear to cause a large effect on the properties, but may only aid in yielding a more flaw free sample. However, it is difficult for example, to understand how this explanation can be applied to the observed differences of 20% in real density (-325 mesh powder) with different curing schedules.

## III. Structural Studies

while the eventual goal of this study is to correlate structure at various stages of the process with properties, the results thus far are largely drawn from samples taken at only two processing stages. The first stage corresponds to a maximum temperature exposure in the range 650-1000°C, while the second stage is that formed at about 2000°C. More recently, samples have been examined at HTT between 1000°C and 2300°C, since for given processing conditions maximum values of mechanical properties were noted in this range.

The techniques chosen for structural examination fall into two broad categories with respect to the information yielded. The first yields information predominantly about the state of the

solid making up of the structure, while the second deals mainly with the void structure. In the first category this study is employing bright field and dark field transmission electron microscopy, electron diffraction, wide angle X-ray diffraction and scanning and electron microscopy. The last two methods also yield information on pore structure.

Additional techniques used to establish the pore structure are small angle X-ray scattering, helium pycnometry, mercury porosimetry, and surface adsorption.

An attempt is also being made to gain structural information through a precise measurement of the thermodynamics of the equilibrium

# Cgraphite = Cglassy

Since the effort required to carry out the above techniques varies substantially, a complete set of data will be collected on only a limited number of samples after screening with more routine tests.

### A. Solid Structure

### X-ray Studies

Several useful measures of the state of non-graphitizing carbons can be derived from wide angle X-ray diffraction. The usual data for d(002),  $L_{\rm c}$ , d(10), and  $L_{\rm a}$  have been measured for a sample from each processing condition with the results reported in Table 1 of the appendix. The procedures are the same as previously reported<sup>2</sup>. The "crystallite size" parameters,  $L_{\rm c}$ 

and L<sub>a</sub> are in reality only line broadening parameters and can not be literally interpreted as the size of graphite crystals, since the analysis considers only size broadening of highly crystalline material and ignores strain effects and the possibility of a distribution of layer spacings. These parameters are useful for comparing different samples of glassy carbon and correspond roughly to a paracrystalline size determined by electron diffraction.

The results obtained to date are similar to others in that they show an increasing  $L_{\rm c}$  and  $L_{\rm a}$ , and decreasing d(002) as the pyrolysis temperature is raised. These results are consistent with the idea that the structure is based on imperfect layers from 15 to 50 Angstroms thick with a layer spacing significantly larger than graphite (3.35Å).

As previously reported, a large number of samples, but not all, show multiple reflections within the (002) line giving the impression of a multiple phase material. These samples have been designated as 2P, 3P or NVS in Table 1 where other than a smooth broad 002 peak exists. These results give strong support for the behavior postulated by Houska and Warren<sup>3</sup> for carbon blacks where they concluded nearest-neighbor pairs of layers assume the ordered graphite relation independently. Layer spacings of 3.35Å and 3.44Å exist corresponding to ordered and disordered pairs, respectively. In the present study the 2P or 3P samples show very good agreement with these values. There is a strong tendency toward multiple reflections with higher pyrolysis tem-

perature, and some correlation with longer curing times and higher catalyst concentration. However, there are many reversals and no satisfactory reason can now be offered to explain why this behavior is sometimes present and sometimes not. It is possible that sharp lines are always present in the (002) but that their intensity is too low to allow resolution in all but exceptional cases. The electron diffraction results confirm this finding.

The only clear-cut correlation between the X-ray data and any processing variable or measured property exists with HTT. While additional correlations may be possible, it appears at present that the X-ray data are not very sensitive indicators of the structural differences that influence the measured properties. The table below indicates the relation of  $L_{\rm C}$  to pyrolysis temperature.

HTT °C	Lc, Angstroms
700	15-18
1300-1450	18-26
1500-1700	19-29
2000-2300	25-110

However, for a given HTT, a variation outside the experimental uncertainty exists for samples with no other apparent processing differences.

# Electron Microscopy and Diffraction

Transmission electron microscopy (TEM) is being used to examine the microstructure of selected glassy carbon samples.

Samples for direct TEM and selected area electron diffraction (SAED) are prepared essentially the same way reported earlier<sup>1,2</sup>, except that now ultrasonic treatment of powder is done for 5-10 minutes. This results in smaller particles which result in better micrographs. However, further improvements in the method of sample preparation are still being explored.

Bright field electron micrograph obtained at magnifications of 36400 to 66300 revealed similar structural features to those reported previously. However, in many samples cylindrical features and evidence for oriented texture have been seen. In general, irregularly shaped platelets 150-500Å in diameter were observed. The granular texture is more distinct in samples heat treated at higher temperatures in agreement with earlier findings. There is considerable variation in the size of platelets and granular texture within the same sample. The characteristic platelet and granular texture diameters are listed in Table 2.

Figure la, taken from sample 321-31D(2300) shows granular texture very clearly but the platelets are not very distinct. Similar features are also visible in certain parts of Fig. 2 taken from sample 317-48(2000). The granular texture size recorded in Table 2 is generally more than 'Lc', the crystallite height, and less than 'La', the crystallite diameter, as measured by X-ray broadening. This suggests that the texture represents domains of paracrystalline material. The variation of size of the granular texture is in agreement with the fact that the Lc and the La obtained from X-ray line broadening are only average values of

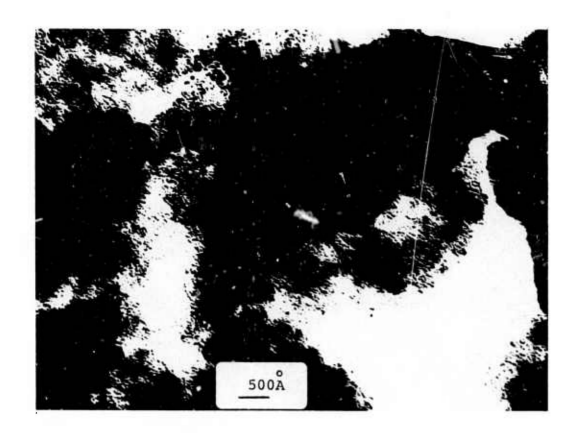


Figure la. Sample 321-31D(2300). Typical granular texture characteristic of bulk of the material. The platelets are not very distinct.

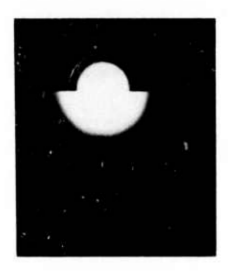


Figure 1b. SAED pattern from upper left corner of the above. Both (002) and (100) contains small spots.



Figure 2. Sample 317-48(2000). Typical platelet and granular features are shown. In addition, cylindrical features are also shown by arrows.



Figure 3a. SAED from 317-49(2000). Streaking can be seen in (100) and (110) halos indicating oriented texture (pattern was exposed thrice to bring out different rings and streaks).

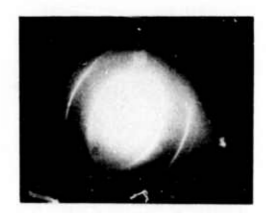


Figure 3b. SAED from 321-31C(2300). Elliptical diffraction pattern characteristic of oriented texture. (002) planes are inclined at an angle of 20 degrees from the beam.



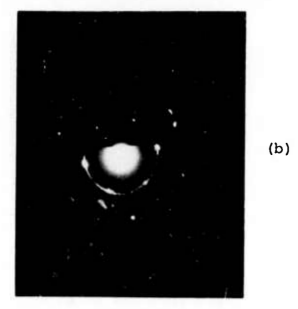


Figure 4. 321-31C(2300). (a) Very small granular texture can be seen on the upper right corner. SAED from this was typical of amorphous material. The region on the left side of the figure is more transparent and smooth. SAED of this region revealed that it is mainly graphite as shown in (b).

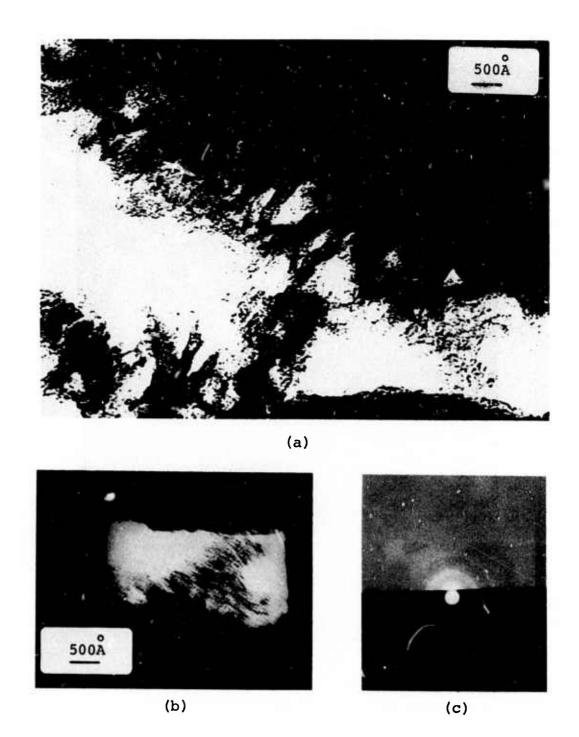


Figure 5. (a) Sample 317-48(2000). Typical granular texture is clearly visible. However, platelets are not present at all. (b) Selected area of (a) from which the SAED of (c) was taken.

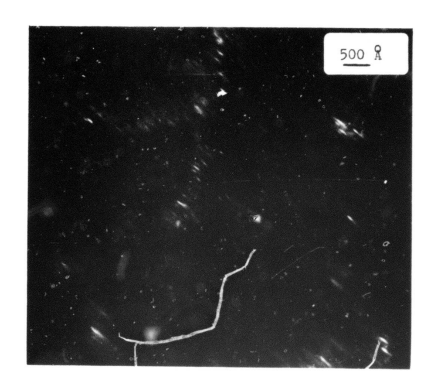


Figure 6. Sample (321-31D(2300). Dark field micrograph taken from (002) halo of SAED shown in Fig. 3b.

crystallite size and considerable variation might be present.

In some regions granular texture is extremely small and thus not very distinct. Micrograph of one such region is shown in Fig. 4a. These regions are generally highly crystalline as indicated by their corresponding SAED patterns. Fig. 4b is an example which was taken from the left side of Fig. 4a. The (002) ring of this pattern is very weak in intensity and spotty, suggesting that the basal planes are perpendicular to electron beam in most of the region. The d(002) of the region is 3.35 indicating presence of graphite crystals in the region.

Another exceptional feature reported earlier for 318-29 (2000) and  $311-19(750)^2$  has also been observed in the samples 317-48(2000), 317-49(2000) and 321-31C(2000). In these samples long cylindrical features are clearly visible as can be seen in Fig. 2 (indicated by arrows).

Selected area electron diffraction patterns, in most of the cases, slowed three diffraction rings corresponding to (002), (100) and (110) reflection (see Fig. 1b and 5c). However, in some cases many additional rings corresponding to higher reflections were also present. The sharpness of the diffraction pattern varied from area to area within the sample. This indicates that a particular sample is non-homogeneous on a microscopic level as would be expected for paracrystalline material.

In addition to the diffraction rings, some of the diffraction patterns contained diffraction spots on the rings (see Fig. 1b and 4b). In some cases diffraction spots were slightly

displaced from the rings corresponding to smaller d(002). The presence of spots on the rings suggests that larger crystals are present in that particular area; while spots corresponding to smaller d(002) suggest the presence of more perfect crystals. These crystals have d(002) greater than 3.35Å which indicates that they might correspond to the second phase seen by X-ray diffraction. Figure 4b shows the diffraction pattern from 321-31C(2300). The area from which this is taken is very graphitic as suggested by the d(002) which is 3.35Å and six spots on the (100) ring. The (002) ring consists of very small spots of low intensity while the (100) ring is very sharp as would be expected if most of the crystals were oriented with basal planes perpendicular to the beam.

Figure 3a shows SAED pattern from 317-49(2000). The (100) and (110) rings contain "streaking", i.e., tangential straight lines on these rings. The streaking is caused by the crystals oriented with their (002) planes parallel to the beam. "', Figure 3b shows elliptical diffraction patterns from (100) and (110) reflections. However, the (002) reflection gave a circular halo but cannot be seen in the figure due to its very low intensity. The distribution of reflections along ellipses is characteristic of an oriented material struck by a beam at an angle to some well-developed plane. In the above case, the (002) planes are tilted at an angle of about 20 degrees from the beam.

Electron diffraction spots on the SAED patterns have also been observed in 312-31(2000) which was originally identified to

be single phase according to X-ray diffraction study. This suggests that the so-called single phase type samples may contain other phases, though in minute amounts. However, no spots were detected in sample 321-31D(2300) which was also identified as a single phase type from X-ray study. In general, 3-phase type samples show more diffraction spots and more crystalline areas compared to other types. This was seen clearly in samples 317-33 (2000), 317-48(2000), 317-49(2000), 318-12(2000), and 321-31C (2300), which were 3-phase type, and 312-31(2000) and 321-31D (2300), which were single phase type.

Ultrasonic treatment of samples before depositing them on the microgrid has resulted in better dark field micrographs from (002) reflections and made it possible to get some micrographs from the (100) reflections as well. However, dark field micrographs from the (100) ring is usually extremely poor in intensity requiring high exposure times.

Dark field micrographs, obtained from (002) reflection, reveal as bright spots the regions or crystallites giving rise to the reflection. Accordingly, the diameter of these spots is equivalent to the crystallite height, Lc. Similarly, dark field micrographs from (100) reflection represents the crystallites giving that reflection. Thus, the diameter of the spots are equivalent to 'La'. Table 2 lists the diameter obtained from these micrographs for different samples. It can be seen that the diameter from (002) dark field is in fairly good agreement with the X-ray 'Lc'. But the diameter obtained from (100) dark field

is generally very much higher than La. A possible reason for this is drifting of the sample caused by long exposure times of up to 1 minute needed to record these micrographs. Figure 6 shows the dark field micrograph obtained from the (002) reflection from the sample 321-31D(2300).

Table 3 shows that average d(002) and d(10) spacings from SAED agree fairly well with the values obtained from X-ray diffraction. The most common error in measuring these values was perhaps varying contraction of films and thus a varying camera constant for every sample. The (100) ring of graphite powder was used to calibrate the microscope for every run. Yet a further check was made with the (100) reflection from the sample itself since ideally d(100) from the sample should always yield a value close to 2.13Å.

Often additional diffracting maxima were observed, which sometimes are not indexable. Further detailed investigation is being done to find if they are not caused by impurities in the sample. However, they may be due to one or more of the crystalline forms reported by Whittaker.<sup>6,7</sup>

### Thermodynamics

Work has progressed in the program for direct determination of the thermodynamic properties of the following equilibria:

$$C_{graphite} = C_{glassy}$$
 (1)

$$C_{\text{graphite}} + CO_2(g) = 2CO(g)$$
 (2)

$$CO(g) + \frac{1}{2}O_2(g) = CO_2(g)$$
 (3)

$$CO_2(g) + H_2(g) = CO(g) + H_2O(g)$$
 (4)

Results are available from at least one run on each of the above equilibria. The primary interest lies in precise measurements of reaction 1 for various types of glassy carbon, while the others are of interest as supplements to already available data.

It has been previously noted 8,2, that values of the free energy for reaction 1 versus temperature can be used as a direct structural measure of the disorder of a particular carbon structure relative to graphite. Available data for the heat capacities of graphite and various glassy carbons 9,10,11 show that any vibrational contributions to the free energy in the range of temperature of interest will be 10 cal/mole or less, which is completely negligible. Therefore, the measured differences can be considered wholly configurational and separated into enthalpy and entropy terms. Disorder such as mixed bonding or strained bonds will cause changes in the configurational enthalpy and a definite increase in the configurational entropy, which can be directly interpreted as a numerical disorder parameter. Such techniques have already been applied in the case of polymers to determine degree of crystallinity 12, where configurational (residual) entropies up to 5 cal/  $mole^{-o}K$  are found. Configurational enthalpies of  $\sim 1500$  cal/mole have been previously measured 13 between glassy carbon and graphite by heat of combustion. Using another reaction, rather large energy differences have also been measured 14 between graphite in

normal and rhombohedral stacking. In addition, Gordon <sup>15</sup> has pointed out from an examination of the discrepancy between experimental determinations of the equilibrium constant for the producer gas reaction and values derived from heat of combustion and spectroscopic data that a residual entropy of .5 cal/mole- $^{\circ}$ K may exist in near perfect graphite. The proposed cell method should be considerably more accurate than the previous methods and yields both the  $\Delta H$  and  $\Delta S$ .

A rather extensive set of check runs has been made in order to insure precision and otherwise check experimental procedures. A series of runs previously reported<sup>2</sup> was carried out on oxides (NiO, FeO, CoO, NbO<sub>2</sub>, Nb<sub>2</sub>O<sub>5</sub>, CO<sub>2</sub>, H<sub>2</sub>O) with known thermodynamic properties in order to verify cell configurations, electrical and temperature measurement techniques, gas train functioning, thermal diffusion effects, and solid electrolyte composition. In all cases, extremely gratifying confirmations were found.

Measurements have been carried out with two entirely different cell techniques; however, at this time, not for the same sample. The first type uses a molten salt, CaCl<sub>2</sub> with CaC<sub>2</sub> dissolved, and involves reversible transfer of carbon directly from one electrode to the other. While the method is straight-forward, the useable temperature range and other experimental difficulties cause limitations. Also, the details of the charge carrying species are less certain since only one previous study<sup>16</sup> has been completed. However, Fig. 7 shows

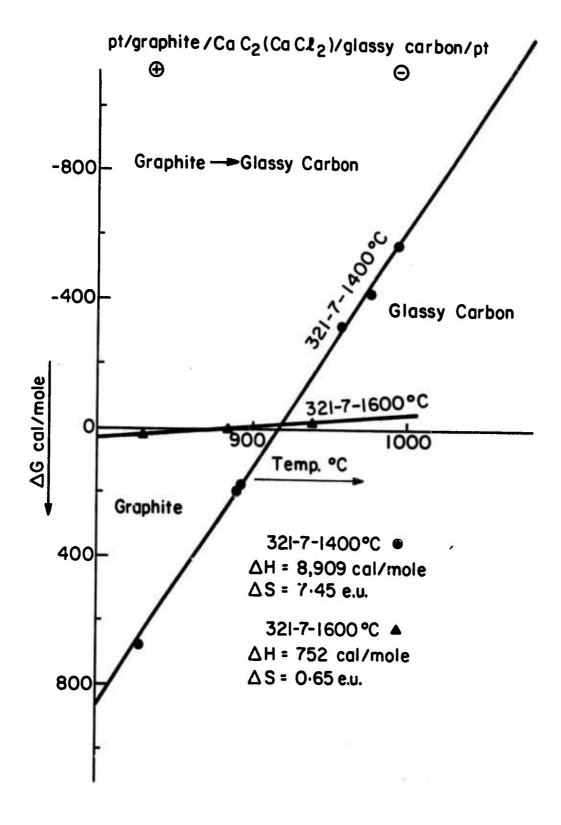


Figure 7.

the results obtained for two portions of a glassy carbon sample differing only in the HTT, i.e., 1400°C vs. 1600°C. The data, as expected, fall on two straight lines which yield the results below

Sample	ΔH, Cal/mole	ΔS, Cal/mole-°K
321-7 (1400°C)	8909	7.45
321-7(1600°C)	752	.65

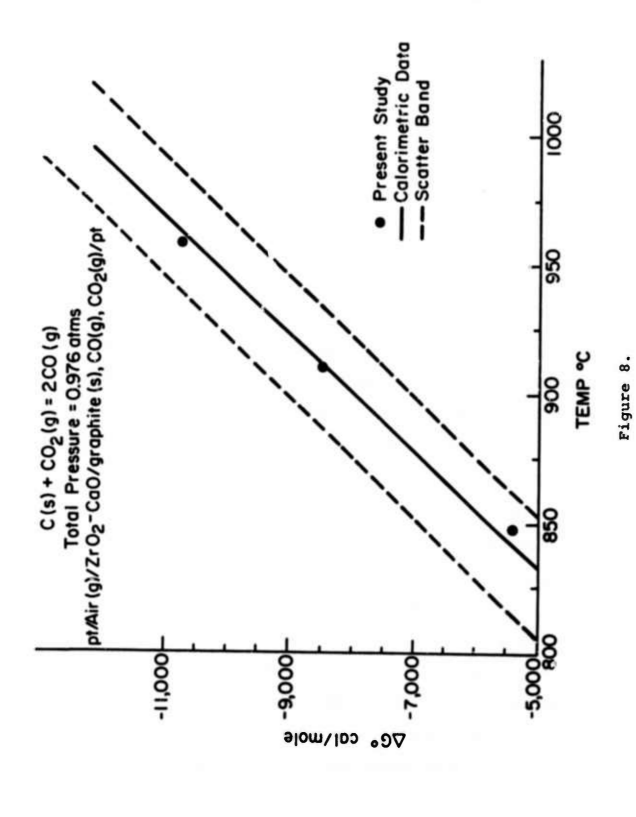
The results, especially for the 1400°C sample are remarkedly high but are quite consistent with the expected behavior of less disorder after higher treatment temperatures. While this cell technique appears feasible, it will be held as an alternative and check procedure since the solid electrolyte technique seems more versatile.

The solid electrolyte cell involves measuring the difference in  $O_2$  pressure existing of opposite sides of the electrolyte where this pressure is additionally constrained to be in equilibrium with CO and  $CO_2$  and solid carbon. The details of analysis have been presented elsewhere.  $^8$ ,  $^2$  If a difference in  $^2$ O<sub>2</sub> exists between the two sides at equilibrium with the same temperature and total pressure after starting with either CO or  $CO_2$  or  $O_2$ , then it can only be due to a difference in the activity of the two carbons. While the equilibrium value may be the same regardless of the starting gas, the rate may be quite different. In order to check the functioning of the cell, several runs were made with Pt only on both sides and then with graphite on both

sides in order to verify that no stray EMF was involved. The cell voltages measured were always less than .1 millivolts. The graphite used in all the runs to date has been a common electrographite (UCC-grade CS) and will be replaced by a carefully selected graphite standard.

The cell was next run with air on one side so that the measured EMF is a direct measure of reaction 2, where the activity of the solid carbon (graphite) is unity by definition. The results are shown in Fig. 8. The agreement is extremely good, since the experimental points lie very close to the best known line and well within the scatter band for the previously available data. Even more refined measurements over a wider temperature interval will be made on the graphite standard. The cell was well-behaved with equilibrium established within several hours. Equilibrium could be approached from both directions and the same value obtained after thermal cycling. The cell voltages have been measured with a high impedance Keithley differential voltmeter, a Keithley electrometer, and a calibrated L&N potentiometer, purposely set to polarize the cell in opposite directions. These experiments show the cell EMF to be reproducible, sensitive, accurate, reversible, and sign consistent.

The next series utilized graphite in one compartment and various glass carbon solid discs in the other. The glass carbons are weighted for total carbon loss and both sides are exposed to pure CO<sub>2</sub> after evacuation. Companion runs on the same samples starting with O<sub>2</sub> both before and after CO<sub>2</sub> exposure are planned.



The results for portions of a glassy carbon pyrolyzed to 700°C and then vacuum heated for 1 hour to 1060°C, 1243°C, and 1800°C are given in Fig. 9. Again, as expected, the data fall on good straight lines giving the following values for  $\Delta H$  and  $\Delta S$  which are shown in a provisional plot in Fig. 10.

Specimen	ΔH, Cal/mole	ΔS, Cal/mole-°K
321-13(1060°C-Vac)	9029	8.66
321-13(1243°C-Vac)	4934	4.70
321-13(1800°C-Vac)	3689	4.60

The rate of approach to equilibrium at the higher temperatures was rapid, but took 2 to 3 days at 650°C, the lower limit for the electrolyte. Future runs call for use of crushed samples to hasten equilibrium. The EMF's of up to 40 millivolts were stable, sensitive, reversible and could be approached from heating or cooling. There can be no doubt that replacing graphite with glassy carbon caused a dramatic change in cell voltage.

While the data are yet too few to draw sweeping conclusions, they do show the expected trends. Again, the magnitudes are embarrassingly high. It is interesting to note that the configurational enthalpy drops steadily with HTT while the order increases rapidly from 1060°C to 1200°C, but very little from 1200°C to 1800°C. Further conclusions must await data for more samples including carbons of many different types and measurements over a larger range of temperature.

It is interesting to note that the above data can be

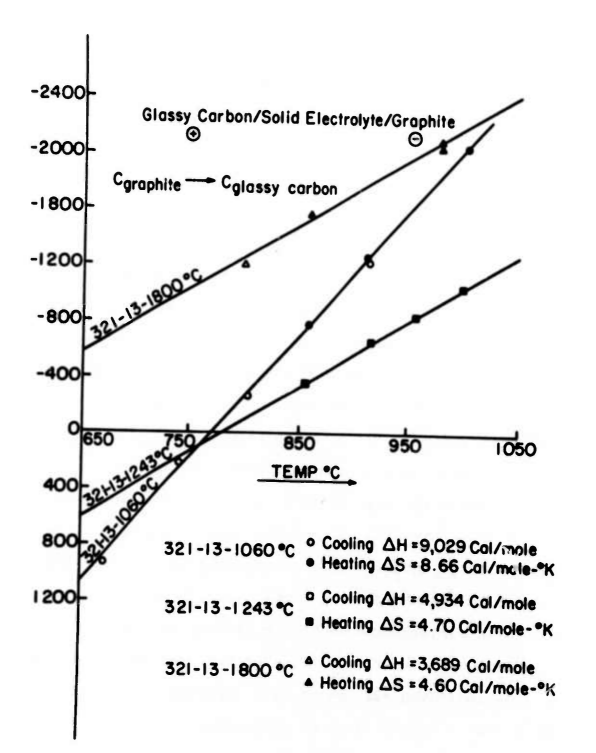
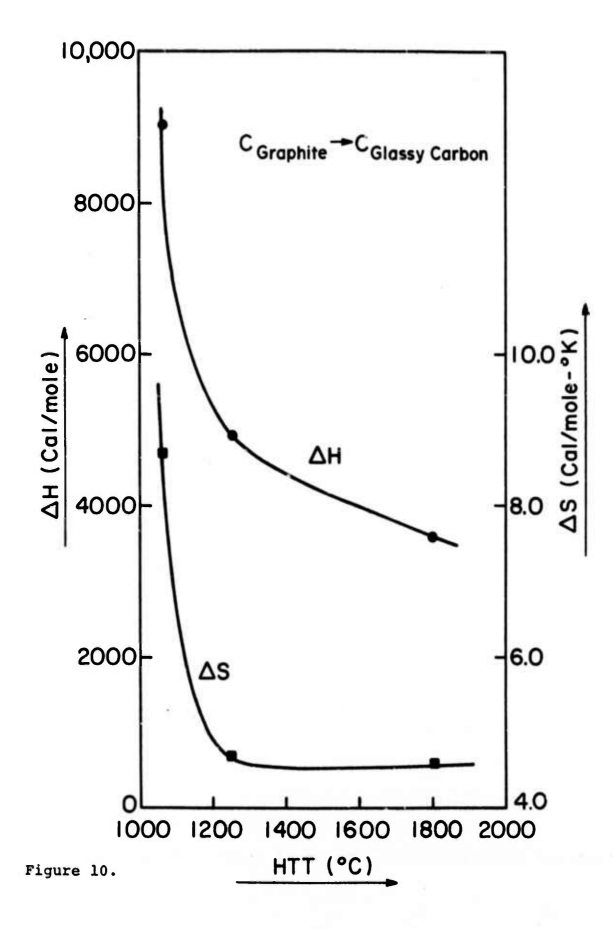


Figure 9.



used to directly calculate the ratio of carbon sites occupied by 0 to those free, if one uses data available for reaction 3 and for the oxygen exchange reaction<sup>17</sup>,

$$C_0 + CO(g) = CO_2(g) + C_f$$
 (5)

Example calculations are given below.

Mommowahuwa	Graphite $\left(\frac{Co}{C_c}\right)$	Glassy Carbon 321-13(1060°C, Vac) $\left\{\frac{Co}{C}\right\}$
Temperature		(°f)
800°C	0.0128	0.0143
1000°C	0.00458	0.01

It can be seen that the surface of glassy carbon is much better covered with oxygen than is graphite at 1000°C.

Work has progressed on a structural model for carbons allowing calculation of the entropy of disorder. At present the results are encouraging but not complete enough for presentation.

#### B. Pore Structure

The details of pore structure are being investigated with a variety of techniques, including small angle X-ray scattering, scanning electron microscopy, pycnometry, surface adsorption, and Hg porosimetry.

# Small Angle X-ray Scattering

Microporosity is one of the main characteristics of the non-graphitizing carbons. The small angle X-ray scattering

permits the evaluation of general structural parameters if recent developments in theory are used to analyze the scattering. Perret and Ruland's 18 work on the non-graphitizing carbons gives an outstanding example. The determination of the pore size and the pore size distribution can be made using Guinier's analysis which has been performed on several samples.

For the evaluation of  $R_{\mathbf{G}}$ , the Guinier radius, the approximation

$$I(S) = n^2 \exp(-\frac{4}{3} \pi^2 R_G^2 s^2)$$

is considered to be valid. Here I(s) is the observed X-ray intensity and s is  $2 \sin\theta/\lambda$ , where  $\theta$  and  $\lambda$  are the Bragg angle and wave length, respectively. The R<sub>G</sub> can be obtained from a plot of I(s) versus s<sup>2</sup>, which is a straight line if the distribution of particle size is narrow. Pore size can be calculated from R<sub>G</sub> if the shape of the pores is known. If the pores are spherical then

$$a = \sqrt{\frac{5}{3}} R_G$$

where a is the radius of the pore.

The extent to which the log I(s) versus s<sup>2</sup> plot follows a straight line gives information about the pore size distribution. In the case of a wide distribution of pore size, the slope of the I(s) versus s<sup>2</sup> curve varies from point to point.

## Experimental Procedure

Copper  $K\alpha$  radiation was employed with pinhole collimation. The intensity was collected on the photographic films. The

density of the films was measured using a Joyce-Loebl microdensitometer. The samples in all the cases were a few millimeters thick.

#### Results

Figs. 11 and 12 show results for some of the samples studied. As can be seen, most of the samples follow Guinier's law, i.e., the log I versus s<sup>2</sup> plot is a straight line.

Table 4 gives our values of "a" along with the reported values for some commercial samples. The agreement is not very good. As in our case, the values are almost twice the reported values. However, all three samples follow Guinier's law fairly well up to  $S^2 = 2 \times 10^{-4} \mbox{Å}^{-2}$  and very little polydispersity is seen which agrees very well with reported results.

Table 5 shows the "a" values for some of our samples. It has been found that samples 317-45(2000), 318-11(2000), 318-42 (2000) are not monodisperse. In these cases the pore size has been determined from the average value of the slope.

The pore size is smaller for the samples heat treated at 700°C than the samples heat treated at 2000°C. This is in agreement with the work of Perret and Ruland on the non-graphitizing carbons. 18

There seems to be a strong correlation between the appearance of multiple reflections in the (002) line and the appearance of polydispersity in small angle scattering. A further investigation of this point is in order.

Currently, the unit is being adjusted for the slit colli-

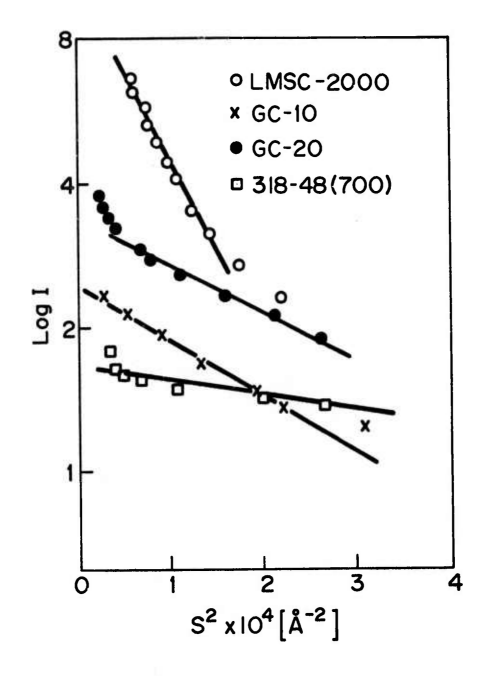


Figure 11. Guinier plots for four different samples. Intensity values are in arbitrary units.

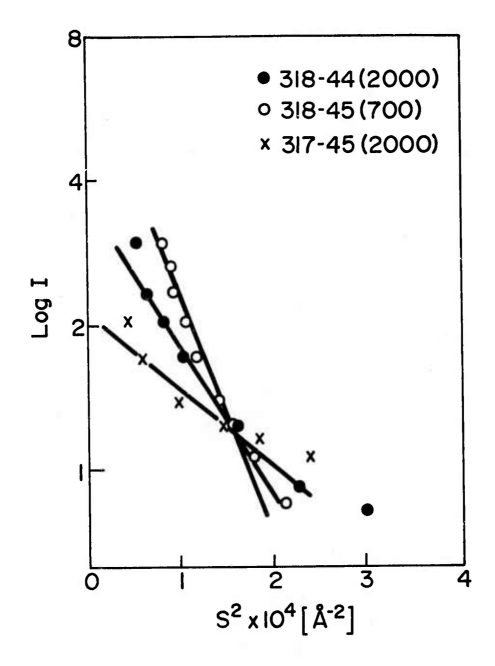


Figure 12. Guinier plots for three different samples. Intensity values are in arbitrary units.

mation so that a counter can be used to determine the intensity and reduce the experimental time per sample.

# Electron Scanning Microscopy

Electron scanning microscopy is employed on a fracture surface of each sample to obtain additional information on the pore structure. For the coarser materials (>200Å) the pore structure is easily discernible with this method. However, for the finer materials, the size of the pores is equal to or finer than the 100Å resolving power of the instrument, rendering the determination of quantitative information impossible. Also, the SEM pictures show a surface roughness of the same order of size as the pore structure which makes pore size assessment difficult.

In many instances, it has been possible to compare the average pore size as measured with SEM to that obtained for Hg porosimetry. The agreement is satisfactory though not precise. For the finer pored materials, lack of resolving power and surface roughness preclude even a rough quantitative check. However, from the observations made on coarser materials, it may be concluded that the Hg intrusion pore diameter is approximately the same as actually seen in the microscope.

The microstructures examined show the same general features as presented previously<sup>1</sup>,<sup>2</sup>, although some improvement in presence of foreign "agglomerate" particles has apparently resulted from filtering the starting resin.

## Pycnometry

As a routine characterization of the pore volume open to He, the real density was measured in a commercial pycnometer made by Micrometrics Inc. of Norcross, Georgia. Powder samples, -325 mesh, are used routinely. Data for samples thus far measured is included in Table 8. Considerable difficulty has been encountered in the He density measurements for carbons with HTT in the range of 700°C. In this range the carbons are known to have a very large micropore volume which has been verified by our surface area determinations. Data for some of these samples yielded an apparent negative density, probably due to a rather large absorption of He in the micropores of the carbon. While in these cases it is obvious that an error exists, it may be that much of the variation in He density of other samples is caused by smaller, but variable, amounts of adsorption. 19 Many of the 700°C samples, however, behaved normally. In general, as expected with these samples, the 700°C density was higher than that at 2000°C as can be seen for example in Table 6. As a result, densities are now being measured by Xylene immersion. In most cases the Xylene density is quite reproducible, and in most cases, slightly lower than the He density as expected.

In addition to real density, the apparent density of the bulk samples is determined geometrically. Table 2 includes the results obtained to date.

#### Surface Area

Additional surface area data for samples 321-9 have been gathered since those in Table 6 reported previously2. The high surface area (~500 m²/gm) of glassy carbons treated to 700°C was again confirmed, as was the dramatic drop after pyrolysis to 2000°C. Sample 321-9 is characterized by relatively high apparent density and very fine pore size as seen in the Hg porosimetry data of Table 7. However, the open porosity of 57 to 73A can not be the porosity accounting for the very high surface area of the 700°C sample as can be seen by comparing the Hg data for the 700°C and 2000°C samples. While the open pore system is about the same size, the 700°C sample has slightly less intrusion pore volume, but almost 50 times the open area for nitrogen adsorption. Similarly, for sample 318-22, the slightly higher Hg intrusion pore volume gives no significantly larger nitrogen surface area. The large surface area is probably associated with the smaller pore system (5-20A) disclosed by small angle scattering and molecular probe techniques.

### Mercury Porosimetry

Mercury porosimetry was used to determine pore size distribution, interconnected pore volume, density and median pore diameter values. Data obtained to date are shown in Table 7.

In most cases, a rather sharp pore size distribution was observed, however, a few samples yielded a wide distribution. The porosimetry data indicate that it is possible to produce material with a rather large open pore volume (up to 65%) with a pore size

ranging from 30 Angstroms to 50 microns. In addition, there is a pore structure at a smaller level of 5-15Å which is open in low temperature carbons to the macropore system. This allows rather exciting possibilities in providing structurally strong, inert elements that could serve both as filters and absorbers.

Higher processing temperature (HTT) closes the very fine pores and causes them to grow according to surface area and X-ray scattering results. The macropore system, even though it is very fine  $(\sim 30\text{\AA})$  in some cases, changes relatively little up to  $2000^{\circ}\text{C}$ . In most cases, noted in Table 7, the pore size decreases slightly while the pore volume remains about constant.

# IV. Property Evaluation

Because the glassy carbons under investigation were produced under a wide range of processing variables, a large degree of variation in structure and physical properties was observed. Based on previous information, some of which was included in a previous report<sup>2</sup>, several tests were chosen as a means of obtaining preliminary mechanical property data. The mechanical properties investigated thus far include hardness (DPH), compressive strength, ultimate tensile strength (Dimetral compression) and sonic modulus of elasticity. In addition, internal friction and electrical resistiveity were measured on selected samples.

#### Hardness

Hardness measurements of these materials have posed constant problems, despite the appeal of providing a quick mechanical measurement. At present no test has been found completely satisfactory since the various carbons vary so widely in pore volume and pore size. The indentation in many of the materials occurs largely by densification under the indentor. In most cases there is a large recovery. Attempts were made during this report period to develop a Rockwell type test which measures depth of indentation instead of indentation stress. Satisfactory testing could be done without cracking at rather large loads using a 1/8" diameter ball. However, the results on widely differing carbons failed to correlate with strength, Vickers hardness, modulus, or scratch hardness. Further hardness testing awaits development of a more relevant test.

#### Compressive and Ultimate Tensile Strength

Further results are reported in Table 8 and presented on "reduced" basis in Table 9. Values were revised where additional data warranted change.

To show the relevance of the diametral disc rupture test, tensile bars 2 inches long by 3/8 inch diameter were ground, threaded gripping ends epoxied in place, and tested in uniaxial tension on an Instron. In this way, a large sample volume is under stress. The results are given in Table 10 showing a very gratifying agreement. The tensile data surpassed the disc results in half the cases. A similar comparison is planned with

three point bending tests on very small samples, which normally gives unrealistically high values.

The tensile disc test often gives a stepped stress-strain curve which raised the possibility that the pores may be functioning as crack stoppers. This observation can not be definitely confirmed at this time since there is a possibility of compressive failure occurring at the disc edges due to stress concentration before the tensile strength has been exceeded in the center of the sample. However, Tingey has noted definite acoustic emissions from these materials prior to catastrophic fracture indicating that the steps may indeed correspond to cracking between pores.

Various correlations have been attempted between strength properties and processing variables, apparent density, real density and pore size without great success. The raw data show a broad correlation with apparent density; with higher strength occurring at higher density. Also, a fair correlation (Figs. 13 and 14) exists with intrusion pore diameter, higher strengths with smaller pores. No other correlations are evident at this time. Since strength is controlled by fracture path, it would be determined by the maximum path within the carbon, i.e., distance between pores. For similar structures and constant densities (real and apparent), the mean free path would vary directly with the average pore size. Unfortunately, these conditions are not met between different samples and further, the only measure of the pore size over an adequate range is the mean

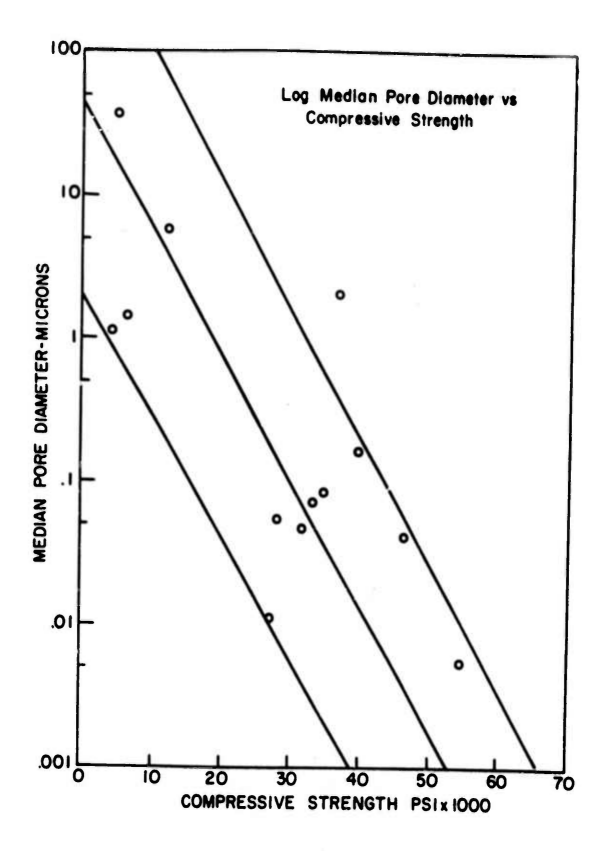


Figure 13.

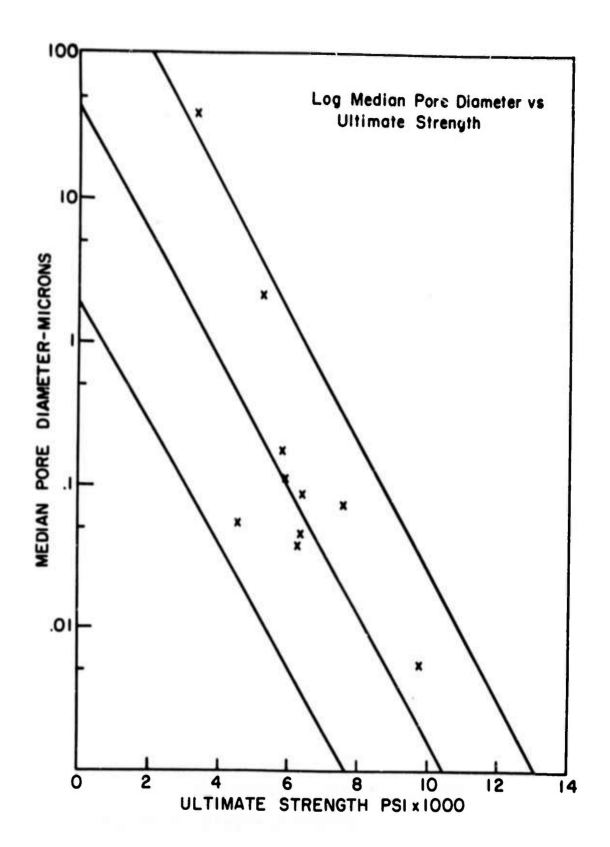


Figure 14.

intrustion pore size. Attempts to use SEM measurements are not very fruitful since the interesting range occurs at sizes where the path length is not easily measured.

The reduced data shown in Table 9 show less range than the raw data. As one method of data reduction, the various mechanical properties are divided by density to yield value of specific strength. On a specific basis these carbon materials compare very favorably with the best available structural materials. Considering the high temperature capability of carbon, the comparison becomes even more favorable.

In addition, the area dependent properties have been normalized by either multiplying or dividing by the area fraction carbon, which is proportional to the volume fraction unavailable to He (i.e.,  $\rho_{\rm a}/\rho_{\rm He})$ , assuming all pores open to He. It can be seen that this normalization considerably reduces the range of values exhibited by the various carbons. However, since the compressive and tensile strengths are certainly dependent on macrostructure as well as fine scale binding of carbon atoms, some range of reduced properties is to be expected.

#### Sonic Modulus and Internal Friction

A conversion factor error was noted in the modulus data presented in the last report<sup>2</sup>. All data subject to this error have been recalculated reducing the previous results by a factor of .75. However, the extremely wide range possible in porous glassy carbons is still present, i.e., from .35×10<sup>6</sup> psi to 5.8×10<sup>6</sup> psi. When the data are reduced to reflect the volume

fraction carbon, the range goes from .32×10<sup>6</sup> psi to 10.06×10<sup>6</sup> psi, with most samples falling around 2.5×10<sup>6</sup> psi. This value is reasonably close to that reported for other more dense glassy carbons. However, the existence of such a wide range is quite interesting since the modulus should not be very dependent upon gross microstructure, but should be determined by shorter range carbon-carbon bonding.

In order to determine whether the sonic modulus, which involves very small strains, well approximates the mechanical modulus, test bars were fitted with strain gages. Prior calculation had already justified that the difference between adiabatic and isothermal moduli for these materials should be negligible. Table 11 shows a comparison of results, which indeed shows that the sonic modulus is a good measure of the modulus at higher strains.

The modulus data versus HTT are shown in Table 12. In all samples the modulus rises from about 10<sup>4</sup> psi after low temperature curing to about 10<sup>6</sup> psi after 700°C HTT and then passes through a broad maximum (√3×10<sup>6</sup> psi) between 900-1500°C HTT. The HTT for this maximum for a single sample series will be investigated including some temperatures below 700°C.

The rather remarkable increase in modulus between the cured and pyrolyzed states illustrated below warrants further study. The evolution of the stiffness with curing may be useful in sensing changes in "crosslinking" in a given sample and between

samples if a suitable method can be devised for normalizing the data with respect to structural changes.

Sonic Modulus, psi
Sample Treatment Temperature °C

Sample	70°C	95°C	700°C
318-35		1.52×10 <sup>4</sup>	.76×10 <sup>6</sup>
318-36		2.47×10 <sup>4</sup>	1.2×10 <sup>6</sup>
318-37		1.56×10 <sup>4</sup>	.69×10 <sup>6</sup>
318-39	.74×10 <sup>4</sup>		.9×10 <sup>6</sup>
318-45	5.5×10 <sup>4</sup>		1.09×106
318-46	6.2×10 <sup>4</sup>	MAL quan	89×106

Internal friction measurements have been discontinued for the present. While remarkable differences in behavior of the various samples have been noted, no straightforward method of correlating these changes to structure has been developed. The anomalous decay curves found in some samples earlier<sup>2</sup> are believed to be associated with mixed frequency excitation due to the specimen claiming device rather than related to porosity as first thought.

## Resistivity

Refinements of the measuring set-up have allowed more accurate measurements, particularly at low resistivity. Cumulative data is presented in Table 13. Resistivity will be continued on a limited basis, especially as a parallel check of sonic modulus data for the low temperature development of crosslinking.

It is interesting to note that the resistivity of

nearly all samples on a reduced basis falls between  $10^{-3}$ - $10^{-4}$   $\Omega$ -cm which is the same as for other glassy carbon materials pyrolyzed to 2000°C.

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APPENDIX

# SUMMARY OF X-RAY DATA (All values in Angstroms)

# Symbols Used in the Tables

## Experimental Condition

All the samples were run in a Phillips-Norelco Diffractometer using  $CuK\alpha$  radiation under the following conditions:

Tube Voltage: 45KV
Tube Current: 14mA

Proportional Counter Voltage: 1.622KV Proportional Counter Time Constant: 4 sec.

Chart Speed: 1/2 inch/min. Scan Speed: 1.2 degree (20)/min.

Slits: 1°/006"/1° at Primary/Scattering/Secondary

Sample used of thickness of 3mm in all cases except where otherwise designated. The value of d(10) refers to the unresolved (100) and (101) peak.

# (002) Peak Type

S: "Smooth" (or single phase) Peak

NVS: "Not Very Smooth" Peak

2P: "2 Phase" Peak 3P: "3 Phase" Peak

Sample Designation	Temp.	(002) Peak Type	d (002)	Lc	d(10)	La
Graphite, solid 3mm & 1mm thick		S	3.37	Very High	2.13	Very High
Graphite		S	3.37	Very	2.13	Very
Graphite, natural (Reported)		S	3.35	High Very High	2.13	High Very
Graphite, synthetic (Reported)		S	3.37	Very High	2.13	High Very High

	Потт	(002)				
Sample Designation	Temp.	Peak Type	d(002)	Lc	d(10)	La
Commercial Samples						
Lockheed, solid Lockheed, reported	2000	S	3.53	21.2	2.09	98.0
Beckwith, solid	2000	•	3.56	19.0		
Beckwith, reported	2000 2000	S	3.55	23.2	2.09	112.0
Tokai, solid	1000		3.54	15.0		50.0
Tokai, reported	2000	S	3.70	14.2	2.07	43.7
Atomergic Chemicals						
Co., V-25, solid	2500	S	3.52	30.2	2.09	66.0
Atomergic Chemicals	2500					
Co., V-25, report			• .			
Atomergic Chemicals	1000	MIC	2 44	45.0		
Co., V-10	1000	NVS	3.44	45.0	2.10	70.0
Atomergic Chemicals	1000					
Co., V-10, report						
200, 10poz 0						
211.0			-			
311-9	2000	S	3.54	28.0	2.10	57.0
311-19	700	S	3.63	18.0		
311-20	2000	S	3.53	27.2	2.10	46.0
311-21	2000	S	3.52	27.2	2.10	54.0
311-22	2000	2P	2.49	29.0	2.12	>125.0
211-25	=1.		3.45			
311-25	700	S	3.70	21.0		
311-30A	2000	S	3.51	23.4	2.10	54.0
311-31	2000	S	3.50	25.0	2.10	51.0
312-8	2000	S	3.52	27.0	2.10	42.0
312-9	2000	2P	3.52	27.0	2.10	57.0
212 10			3.45			
312-10	700	S	3.65	16.2		
312-10 312-14	2000	S	3.49	35.0	2.11	53.0
	2000	S	3.51	29.0	2.09	61.0
312-14A	2000	3 <b>P</b>	3.46	32.0	2.12	>150.0
			3.43			
312-15	2222		3.36			
312-16	2000	S	3.52	37.1	2.11	51.0
312-16	2000	3P	3.47	32.1	2.11	51.0
			3.43			
312-21	2000		3.36			
312-21	2000	S	3.51	30.8	2.10	57.0
312-28	2000	S	3.51	27.8	2.09	48.0
312-28	2000	S	3.51	27.2	2.10	51.0
	2000	S	3.51	34.0	2.10	57.0
312-31, solid 312-32	2000	S	3.57	27.6	2.10	56.0
312-32	2000	S	3.51	30.8	2.10	54.0
312-33	2000	NVS	3.48	30.8	2.10	52.8

	Temp.	(002) Peak				
Sample Designation	(°C)	Type	d(002)	Lc	d(10)	La
312-34	2000	2P	3.46 3.44	33.0	2.11	46.2
312-39	2000	S	3.49	29.8	2.10	53.0
312-40	2000	2P	3.48 3.45	30.1	2.09	54.0
312-43	2000	2P	3.48 3.43	33.2	2.11	51.0
312-44	2000	2P	3.48	42.0	2.11	51.0
312-48	2000	3P	3.46 3.43 3.37	30.4	2.10	37.0
312-49	2000	S	3.52	31.1	2.11	63.0
315-1	2000	S	3.53	27.2	2.11	61.0
315-2	2000	2P	3.49 3.43	29.0	2.10	48.0 54.0
315-3	700	S	3.71	16.2		
315-5	2000	2P	3.49 3.44	29.0	2.11	54.0
315-8	2000	2P	3.49	28.2	2.10	56.0
315-9	2000	2P	3.47	33.0	2.11	47.0
315-14	2000	S	3.53	26.3	2.10	54.0
315-18	2000	3P	3.40 3.382 3.351	45.0		
315-20	680	S	3.70	16.3		
315-20A	2000	2F	3.53 3.43	28.0	2.10	57.0
315-21C	2000	2P	3.52 3.43	27.8	2.10	57.0
315-22	665	S	3.67	16.4		
315-22	2000	S	3.52	28.0	2.10	51.0
315-24A	2000	2P	3.47 3.38	20.0		
315-25A	2000	S	3.52	26.5	2.09	48.0
315-26B	2000	3P	3.50 3.41 3.37	26.5	2.10	46.0
315-26C	680	S	3.69	17.1		
315-28	2000	2P	3.52 3.43	26.0	2.10	46.0
315-28B	600	S	3.70	16.8		
315-30	2000	2P	3.56 3.43	24.0	2.09	48.0
315-31	680	S	3.70	18.2		
315-34	680	S	3.69	15.4		

		(002)				
Sample Designation	Temp.	Peak	4/002\	_	W 4 W - A	
315-36		Туре	d(002)	Lc	d(10)	La
313-36	2000	3P	3.52	24.3	2.10	48.5
			3.43 3.37			
315-37	680	S	3.63	17.5		
315-37	2000	S	3.50	26.3	2.098	42.0
315-38	680	S	3.63	18.8	2.098	42.0
315-38	2000	2P	3.49	27.1	2.097	46.0
47.61.42			3.43		2.007	40.0
315-39	2000	2P	3.53	27.2	2.098	57.0
315-39	600		3.43			
315-40	680	S	3.63	20.0		
315-41	2000	S	3.54	25.6	2.097	51.0
315-42	2000 2000	NVS	3.49	23.6	2.098	51.0
315-43	2000	S	3.56	27.2	2.098	46.0
315-43	700	NVS S	3.52	24.3	2.098	51.0
315-44	2000	2P	3.67 3.55	17.4	2 10	40.0
	2000	2.5	3.45	23.1	2.10	40.2
315-45	2000	s	3.49	27.2	2.10	46.6
315-46A	2000	2P	3.55	23.1	2.098	57.0
			3.43	20.1	2.030	37.0
316-6	2000	NVS	3.50	27.0	2.11	57.0
316-7, Run I	2000	S	3.49	28.0	2.10	45.0
316-7, Run 2	2000	S	3.52	27.0	2.10	53.0
316-15 316-28	2000	2P	3.40	32.0		
316-28	2000	S	3.50	27.2	2.10	51.0
310-32	2000	2P	3.42	53.0		·
317-1	700	C	3.40			
317-1	2000	S	3.71	20.0		
317-2	700	S	3.46 3.68	45.0	2.11	63.0
317-2	2000	NVS	3.48	15.7 24.6	2.09	47.0
317-6	700	S	3.71	13.0	2.09	47.0
317-6	2000	NVS	3.55	22.0	2.10	55.0
317-7	700	S	3.68	16.0		
317-7	2000	NVS	3.46	27.5	2.10	50.0
317-8	700	S	3.71	11.5		
317-8	2000	2P	3.56	20.0	2.10	44.0
317-10	2000	\***	3.46	200		
317-11	2000 700	NVS	3.48	26.0	2.10	68.0
317-13	700	S S	3.71	16.3		
317-13	2000	NVS	3.72	15.0		
317-14	700	S	3.47 3.71	24.0 15.7	2.08	66.0
317-14	2000	NVS	3.45	27.0	2.09	46.0
317-15	700	S	3.71	15.3		46.0
317-15	2000	NVS	3.47	26.0	2.09	54.0
317-16	2000	S	3.54	24.0	2.09	53.0

	Mamm	(002)				
Sample Designation	Temp.	Peak Type	d(002)	Lc	d(10)	La
317-18	2000	S	3.59	21.0	2.09	58.0
317-19	700	S	3.68	16.5		
317-19	2000	NVS	3.49	30.0	2.09	52.0
317-20	700	S	3.66	17.5		
317-20	2000	S	3.50	25.6	2.09	48.0
317-24, Run 1	2000	NVS	3.52	24.0	2.09	45.0
317-24, Run 2, solid	2000	NVS	3.49	21.0	2.09	50.0
317-25	2000	S	3.53	20.0	2.09	50.0
317-26, Run 1	2000	NVS	3.48	26.0	2.09	48.0
317-26, Run 2, solid	2000	2P	3.46	24.2	2.10	51.0
			3.43			
317-28	2000	NVS	3.46	25.0	2.09	52.0
317-29	700	2P	3.43	21.5		
			3.42			
317-29, Run 1	2000	NVS	3.43	65.0		
317-29, Run 2	2000	NVS	3.426	75.0		
317-30	2000	2P	3.44	29.5		
			3.40			
317-31A	2000	S	3.58	22.0	2.10	46.0
317-32	700	S				
317-32	2000	2P	3.51	23.6	2.08	49.0
			3.48			
317-33	700	S	3.68	17.0		
317-33	2000	S	3.414	92.0	2.10	49.0
317-34	700	S	3.68	17.0		
317-34	2000	3P	3.44	30.0	2.09	50.0
			3.42			
		-	3.36			
317-35	700	S	3.71	16.0		
317-35	2000	3 <b>P</b>	3.50	26.5	2.10	62.0
			3.43			
		161	3.36			
317-37	700	S	3.68	15.6		
317-37	2000	NVS	3.43	43.0	2.10	63.0
317-38	700	S	3.68	15.6		
317-38	2000	3P	3.54	25.0	2.09	51.0
			3.43			
217 20 5 1	2222	2.5	3.37	00.0	0.10	F0 0
317-39, Run 1	2000	3P	3.45	28.0	2.10	52.0
			3.43			
217 20 20- 2	2000	25	3.36	24.2	2 00	AE O
317-39, Run 2,	2000	3P	3.52	24.2	2.09	45.0
solid, lmm thick			3.42			
217 20 5 2	2000	25	3.37	22 5	2 00	40.0
317-39, Run 3,	2000	3P	3.52	23.5	2.09	49.0
solid, lmm thick			3.41			
			3.37			

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	Temp.	(002) Peak				
Sample Designation	(°C)	Type	d(002)	Lc	d(10)	La
317-40	2000	3P	3.49 3.42 3.36	24.8	2.10	48.0
317-41A	2000	S	3.53	26.5	2.10	40.0
317-41B	2000	S	3.53	28.0	2.10	48.0
317-42	2000	3P	3.49	26.0	2.10	54.0 42.6
			3.42	20.0	2.10	42.0
217.42			3.36			
317-43	2000	3P	3.45	26.0	2.09	56.0
			3.42			
317-44	2000		3.35			
217-44	2000	3 <b>P</b>	3.48	30.0	2.09	55.0
			3.42			
317-45, solid,	700		3.36			
lmm thick	700	S	3.75	12.9		
317-45, Run 1	2000	3P	2.40		4	
	2000	JF	3.48	25.0	2.09	60.0
			3.40 3.35			
317-45, Run 2	2000	3P	3.46	24.0	2 00	40.0
		<b>J.</b>	3.42	24.0	2.09	42.0
			3.35			
317-46	2000	3P	3.43	31.5	2.10	75 0
			3.42	31.3	2.10	75.0
			3.36			
317-47	2000	3P	3.50	27.0	2.10	60.0
			3.42			00.0
217 40 5			3.36			
317-48, Run 1	700	S	3.71	16.2		
317-48, Run 2 317-48, Run 1	700	S	3.87	17.4		
317-48, Run 1	2000	3P	3.45	40.0	2.10	60.0
			3.43			
317-48, Run 2	2000	25	3.37			
solid, lmm thick	2000	3P	3.46	34.0	2.10	59.0
John Child			3.43			
317-49	700	S	3.37	15 7		
317-49, Run 1	2000	3P	3.71 3.49	15.7 29.0	2 00	
		31	3.41	29.0	2.09	62.0
			3.35			
317-49, Run 2	2000	3P	3.46	33.0	2.09	60 0
solid, 1mm thick			3.44	33.0	2.09	60.0
			3.37			
317-50	700	S	3.67	15.6		
318-1	2000	S	3.55	28.0	2.10	54.0
318-2	2000	S	3.51	27.0	2.10	55.0
318-3, Run 1	700	S	3.70	16.7		
318-3, Run 2	700	S	3.69	16.7		

		(002)				
Sample Designation	Temp.	Peak	4(002)	T = 1	a (10)	•
		Type	d(002)	Lc		La
318-3	2000	2P	3.46	26.0	2.11	51.0
210 4	700	C	3.41	16.0		
318-4 318-6A	700 2000	S S	3.66 3.50	16.8	2 00	
318-7, Run 1	2000	S	3.50	31.0 28.0	2.09 2.10	59.0 65.0
318-7, Run 2, solid	2000	S	3.49	28.0	2.10	45.0
318-8, Run 1	2000	S	3.45	39.0	2.10	63.0
318-8, Run 2	2000	s	3.45	43.5	2.10	77.0
solid, 2mm thick						
318-9	2000	2P	3.48	32.5	2.11	57.0
			3.46			
318-10	520	S	3.74	15.2		
318-10	2000	S	3.49	33.8		50.0
318-11, Run 1	2000	NVS	3.42	77.0	2.10	38.0
318-11, Run 2	2000	NVS	3.43	78.0	2.11	40.0
318-12	2000	3P	3.49	31.4	2.09	59.0
			3.43			
318-13	2000	NVS	3.36 3.42	44.0	2.10	E0 0
318-14	700	S	3.65	16.0	2.10	58.0
318-14	2000	3P	3.48	30.5		60.0
310 14	2000	31	3.43	30.3	2.10	00.0
			3.36			
318-15, Run 1	700	S	3.75	16.0		
318-15, Run 2	700	S	3.75	15.1		
318-15	2000	3 <b>P</b>	3.45	30.2	2.10	60.0
			3.42			
			<b>3.37</b>			
318-16	700	S	3.72	15.7		
318-16	2000	2P	3.43	39.0	2.09	49.0
210 17	700	G	3.41	16 7		
318-17 318-17	700 2000	S NVS	3.68 3.45	16.7 42.0	2.11	59.0
318-18, Run 1	700	_	3.68	16.4	2.11	39.0
318-18, Run 2	700	S	3.71	16.3		
318-18	2000	S	3.55	25.6	2.10	44.0
318-19	2000	S	3.52	26.0	2.09	59.0
318-20	700	S	3.67	16.0		
318-20	2000	S	3.53	21.0	2.09	48.0
318-21, Run 1	700	S	3.78	14.0	~-	
318-21, Run 2	700	S	3.75	15.4		
318-21	2000	S	3.55	23.6	2.10	55.0
318-22	700	S	3.70	15.4		
318-22, Run 1	2000	NVS	3.44	65.0	2.10	55.0
318-22, Run 2	2000	NVS	3.44	64.0	2.11	54.0
318-23 318-23	700 2000	S S	3.74 3.63	16.0 63.0	2.10	73.0
318-24	700	S	3.64	16.7		73.0
	. 5 0	_	2.03	2007		

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	-	(002)				
Sample Designation	Temp.	Peak Type	d(002)	Lc	d(10)	La
318-24	2000	S	3.44	45.0	2.10	68.0
318-26, Run 1	700	S	3.69	15.7		
318-26, Run 2	700	S	3.75	16.1		
318-26, Run 3	700	s	3.69	16.7		
318-27	2000	2P	3.45	35.4	2.10	47.0
			3.41			
318-28	700	s	3.75	18.0		
318-28	2000	2P	3.47	27.0		
			3.42			
318-29, Run 1	2000	NVS	3.45	30.0	2.08	62.0
318-29, Run 2,	2000	2P	3.50	30.5	2.10	65.0
solid, 1mm thick			3.44			
318-29, Run 3	2000	2P	3.52	31.0	2.10	60.0
<b></b>		<u></u> -	3.42			
318-30	700	s	3.64	15.2		
318-30, Run 1	2000	2 <b>P</b>	3.48	34.1	2.11	69.0
320 00, 11411 -			3.43	• • • •		
318-30, Run 2	2000	3P	3.45	31.0	2.11	63.0
323 33, 11411 2		-	3.41			55.5
			3.36			
318-31, Run 1	2000	2P	3.45	35.5	2.10	64.0
Sio Si, Kan i	2000		3.43	33.3	2.10	04.0
318-31, Run 2	2000	3P	3.47	31.0	2.11	63.0
310 317 Man 2	2000	<b>J.</b>	3.41	31.0	3.11	03.0
			3.36			
318-32, Run 1	700	S	3.64	15.7		
318-32, Run 2	700	S	3.63	16.0		
318-32 Kan 2	2000	S	3.44	47.0	2.10	65.0
318-33	700	S	3.66	16.7		
318-33	2000	NVS	3.46	28.0	2.11	64.0
318-34	700	S	3.63	16.5		
318-34	2000	3P	3.49	37.0	2.10	59.0
310 34	2000	<b>J</b> -	3.43	37.0	2020	33.0
			3.36			
318-35	700	S	3.71	15.3		
318-35	2000	3P	3.50	34.0	2.11	67.0
310 33	2000	0.2	3.44			
			3.37			
318-36	700	S	3.68	17.0		
318-36	2000	2P	3.51	28.0	2.10	49.0
318-30	2000	21	3.44	20.0	2.10	43.0
318-37	700	S	3.71	16.1		
318-37	2000	3P	3.46	33.6	2.10	52.0
310-37	2000	32	3.43	33.0	2120	5215
			3.376			
318-38	700	S	3.71	15.6		
318-38	2000	3P	3.47	28.0	2.10	49.0
J_0- J0	2000	<b>J</b>	3.43	20.0	2110	
			3.37			
			3.37			

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	Temp.	(002) Peak				
Sample Designation	(°C)	Type	d(002)	Lc	d(10)	La
318-39, Run 1	700	S	3.71	17.0		
318-39, Run 2	700	S	3.65	17.2		
solid, lmm thick	0000	_	. =-			
318-39 318-40	2000	S	3.51	26.1	2.09	60.0
318-40	700	S	3.71	15.0		
310-40	2000	2P	3.52	28.0	2.11	54.0
318-41	700	S	3.45 3.71	14.0		
318-41	2000	S	3.50	14.8	2.00	 
318-43, Run 1	700	S	3.69	28.0 17.0	2.09	57.0
318-43, Run 2	700	s	3.71	13.8		
solid, 1mm thick		_	3.71	13.0		
318-43, solid	2000	S	3.44	31.0	2.12	58.0
318-44	700	S	3.72	15.6		
318-44	2000	S	3.55	27.2	2.10	44.0
318-45	700	S	3.71	15.7		
318-45	2000	S	3.56	25.4	2.10	46.0
318-46	700	S	3.71	15.9		
318-46, solid	2000	S	3.53	26.2	2.11	51.0
lmm thick						
318-47	700	S	3.71	15.0		
318-47	2000	NVS	3.49	29.0	2.10	48.0
318-48, Run 1 318-48, Run 2	2000	S	3.53	26.8	2.10	54.0
318-50, Run 1	2000 700	S	3.52	29.2	2.10	42.0
318-50, Run 2	700	S S	3.71	14.3		
318-50	2000	S	3.71	15.5		
318-51	2000	S	3.53 3.56	26.0 27.2	2.10	46.0
318-52	2000	S	3.53	26.5	2.10 2.10	56.0
318-53, Run 1	2000	S	3.52	26.5	2.10	54.0 54.0
318-53, Run 2	2000	S	3.54	30.0	2.10	60.0
318-54	700	S	3.66	17.0		
318-55	700	S	3.71	15.2		
318-56	2000	S	3.54	27.0	2.10	54.0
318-58	700	S	3.71	18.0		
318-58	2000	NVS	3.51	28.2	2.10	51.0
318-59	700	S	3.68	16.7		
318-59	2000	S	3.51	26.0		
318-60	700	S	3.70	15.7		
318-60	2000	2P	3.47	32.0		
318-61	700	C	3.44	10.6		
318-61	700 2000	S	3.71	18.6	2.00	
318-62	700	S S	3.52 3.70	23.3	2.09	55.0
318-62	2000	S	3.70	15.3 22.5	2.10	51 0
321-1	700	S	3.66	5.0	2.10	51.0
321-2	700	2 <b>P</b>	3.63	17.4		
			3.57			

	Temp.	(002) Peak				
Sample Designation	(°C)	Type	đ (002)	Lc	d(10)	La
321-2	2000	3P	3.54	22.8	2.10	51.5
			3.43			
201 2	<b>500</b>	-	3.38			
321-3 321-3	700	S	3.64	17.4		
321-4	2000 700	S S	3.53	24.3	2.10	51.0
321-4	2000	- -	3.64	17.2		
321-5	700	S	3.63	15.4		
321-5	2000	S	3.49	26.4	2.09	53.7
321-6	700	s	3.64	17.0	2.03	33.7
321-6	2000	S	3.54	27.7	2.10	48.0
321-7	700	S	3.69	18.0		
321-7	2000	-				
321-8	700	S	3.69	17.5		
321-8	2000	-				
321-9	700	S	3.67	17.4		
321-9	2000	_				
321-10 321-10	700	S	3.67	17.0		
321-10	2000 700	-	2 73	17.0		
321-11	2000	S 2P	3.71 3.54	17.0	2.16	
321 11	2000	2 <b>F</b>	3.46	27.2	2.10	65.0
321-12	700	S	3.63	16.8		
321-12	2000	S	3.53	26.4	2.094	56.0
321-13	700	S	3.66	17.0	2.034	30.0
321-16A	2000	3P	3.50	30.8	2.10	57.0
			3.43			
			3 <b>.36</b>			
321-16B	2000	S	3.50	28.8	2.10	44.0
321-16C	700	S	3.63	15.2		
321-17	700	S	3.60	18.7		
321-17B	2000	S	3.49	28.8	2.10	44.0
321-18A	2000	3P	3.50	29.8	2.10	49.0
			3.43 3.37			
321-18B	700	S	3.63	15.2		
321-19A	2000	2P	3.54	25.0	2.09	46.0
			3.426	23.0	2.05	40.0
321-19A	700	s	3.63	17.1		
321-19B	2000	NVS	3.43	39.0	2.10	53.8
3 <b>21-20A</b>	700	S	3.63	17.5		
321-20A	2000	3P	3.52	28.0	2.10	61.0
			3.42			
201 005			3.36			72-01
321-20B	2000	3P	3.53	37.0	2.10	46.0
			3.426			
321-21A	700	s	3.37	10 0		
JZI-ZIR	700	3	3.63	18.0		

	Temp.	(002) Peak				
Sample Designation	(°C)	Type	d(002)	Lc	d(10)	La
321-21A	2000	NVS	3.43	41.6	2.10	51.0
321-21B	700	S	3.64	18.4		
321-21B 321-22A	2000 2000	S 2P	3.52 3.52	27.0 27.6	2.10 2.10	57.0 53.0
321-228	2000	2.5	3.43	27.0	2.10	53.0
321-22B	700	S	3.70	16.1		
321-23	2000	S	3.49	33.0	2.10	54.0
321-23A	700	S	3.63	18.4		
321-23B 321-23B	700 2000	S 3P	3.36 3.52	16.5 29.6	2.10	51.0
321 235	2000	Jr	3.43	29.0	2.10	31.0
			3.36			
321-24	700	S	3.70	15.8		
321-24A	700	S	3.63	16.2		
321-24B 321-24B	700 200 <b>0</b>	S	3.63	16.5	2.00	 
321-25	700	S S	3.43 3.63	35.4 18.4	2.09	57.0
321-25	2000	NVS	3.47	30.8	2.10	60.8
321-25A	700	S	3.60	21.0		
321-25A	2000	NVS	3.43	40.2	2.10	60.5
321-26	700	S	3.67	15.0		
321-26 321-26A	2000 700	S S	3.52	27.2	2.094	48.5
321-26A 321-26A	2000	S	3.63 3.52	18.1 27.2	2.094	54.0
321-27	2000	S	3.52	29.8	2.10	51.0
321-29	700	S	3.63	16.8		
321-29	2000	3P	3.49	24.8	2.094	58.0
			3.40			
321-30	700	S	3.35 3.63	19.6		
321-30	2300	3P	3.44	49.0	2.10	60.5
			3.41		2.10	00.5
			3.37			
321-31A	2300	NVS	3.40	90.0	2.11	58.0
321-31B	2300	3P	3.49 3.43	37.0	2.11	69.0
			3.43			
321-31C	700	S	3.60	18.8		
321-31C	2300	3P	3.49	34.5	2.11	69.0
			3.426			
321-31D	700	0	3.37	17 4		
321-31D 321-31D	700 2300	S S	3.63 3.47	17.4 37.2	2.11	69.0
321-31E	700	S	3.63	17.4	2.11	
321-31E	2300	NVS	3.426	61.6	2.11	69.0
321-31F	700	S	3.60	18.5		
321-31F	2300	S	3.45	44.0	2.10	69.0
321-31G	2300	2P	3.47 3.43	35.0	2.10	56.0
			3.43			

No.

I

	Temp.	(002) Peak				
Sample Designation	(°C)	Type	d(002)	Lc	d(10)	La
3 <b>21-31</b> I	2300	2P	3.49 3.38	40.0	2.11	69.0
321-32	700	S	3.60	18.9		
321-34A	2300	ŝ	3.426	51.4	2.10	42.0
321-36A	2300	S	3.47	53.6	2.11	42.0
3 <b>21-36</b> B	700	Š	3.63	18.2	2.11	
321-36C	2300	S	3.43	70.0	2.10	46.0
3 <b>21-</b> 37	2000	S	3.52	30.8	2.10	54.0
3 <b>21-37A</b>	700	S	3.60	18.1		J4.0
321-37B	2000	3P	3.53	27.2	2.10	54.0
			3.44			34.0
321-38B	700	~	3.36	_		
321-36B 321-39	700	S	3.63	17.7		
321 39	2000	2P	3.48	34.1	2.10	57.0
321-39B	700	C	3.426			
321-41C	2000	S 3P	3.63	16.4		
227 470	2000	3P	3.49	32.6	2.10	62.5
			3.43 3.36			
321-42A	700	S	3.63	18.8		
321-42A	2000	29	3.49	35.5	2.10	
0		2.	3.43	33.3	2.10	60.5
321-42B	700	S	3.63	16.7		
321-42B	2000	2P	3.50	25.3	2.10	64.4
			3.43	23.3	2.10	04.4
3 <b>21-43</b> B	700	S	3.63	19.3		
3 <b>21-4</b> 3B	2000	2P	3.49	35.0	2.098	59.4
			3.42			3311
321-43B <sub>1</sub>	2000	2P	3.50	36.8	2.11	59.5
227 425			3.426			
321-43B <sub>2</sub>	2000	2P	3.50	33.0	2.10	57.0
207 442		_	3.43			
321-44A	2000	3P	3.50	30.8	2.10	51.0
			3.43			
321-44B	2000	25	3.36			
321-44B	2000	3P	3.50	29.0	2.10	54.0
			3.43			
321-45A	2200	C	3.36	22.0	2.10	
321-45B	2200	S S	3.49 3.47	33.0	2.10	57.0
321-46A	2000	2P	3.49	45.5	2.10	40.5
	2000	45	3.43	31.7	2.10	60.4
321-46B	2000	3 <b>P</b>	3.46	30.8	2.10	51.0
			3.43		- · <b>- ·</b>	
			3.36			
321-46C	2000	3P	3.47	35.6	2.10	51.0
			3.43			
			3.36			

	Temp.	(002) Peak				
Sample Designation	(°C)	Туре	d(002)	Lc	d(10)	La
321-46D	2000	3P	3.50	31.7	2.10	64.4
			3.43			
321-48A	2000	c	3.36	27.0		14
321-48B	2000	S S	3.50 3.50	31.8 33.0	2.10 2.10	57.0
321-48C	2000	S	3.50	33.0	2.10	64.4 60.4
321-49A	2000	S	3.49	33.0	2.10	60.4
321-49B	2000	NVS	3.426	51.2	2.10	54.0
321-51	2000	NVS	3.43	91.5	2.11	69.0
321-51A 321-52	2000	NVS	3.43	107.5	2.11	68.0
321-32 322-1A	2000 2000	NVS	3.43	91.2	2.11	60.5
322 IA	2000	3 <b>P</b>	3.44	26.4	2.085	57.3
			3.37 3.33			
322-1B	2000	S	3.40	41.5	2.085	
322-2B	1600	S	3.50	23.0	2.085	37.3
322-3B	1600	S	3.53	22.0	2.085	61.0
322-10A	2000	2P	3.46	33.0	2.085	
322-10B	2000		3.42			
322-10B	2000	2P	3.50	30.8	2.10	69.0
322-10C	2000	2P	3.43 3.49	22.4	2 225	
311,233	2000	2.5	3.49	33.4	2.085	
322-10D	2000	2P	3.50	28.6	2.10	60.0
			3.43	20.0	2.10	00.0
322-11A	1670	S	3.55	22.0	2.085	51.2
322-12A	1600	S	3.56	23.0	2.085	40.5
322-12B	1670	S	3.56	22.0	2.10	57.0
322-13B 322-14A	1670	S	3.53	31.8	2.085	50.2
322-14B	1670 1670	S	3.50	23.0	2.085	
322-15B	1670	s s	3.56 3.56	21.0	2.09	46.0
322-16A	1670	S	3.56	25.0 23.5	2.085 2.085	48.4
322-16B	1670	S	3.56	22.0	2.085	44.0
322-17A	1670	S	3.56	22.2	2.085	88.0
322-17B	1670	S	3.56	22.0	2.085	51.4
322-18B	1670	S	3.50	28.8	2.085	61.5
322-19B	1670	S	3.56	20.6	2.085	48.5
322-20 322-21B	1670	S	3.50	21.4	2.085	30.2
322-21B 322-25A	1670 1410	S	3.56	17.0	2.085	69.5
322-25B	1410	s s	3.56 3.56	18.5	2.085	46.1
322-26A	1410	S	3.50	19.2 25.6	2.07	121.0
322-26B	1410	S	3.50	19.2	2.085 2.085	51.0 40.2
322-27A	1410	s	3.59	20.0	2.085	57.3
322-27B	1410	S	3.56	19.3	2.085	
322-28A	1410	S	3.56	19.4	2.08	42.0
322-28B	1410	S	3.56	17.8	2.085	53.2

Sample Designation	Temp.	(002) Peak Type	đ (002)	Lc	d(10)	La
322-29	1410	S	3.56	20.1	2.085	40.2
322-29B	1410	S	3.56	21.0	2.085	53.2
322-31A	1410	S	3.56	23.0	2.085	40.4
322-36	1543	S	3.56	19.2	2.085	53.0
322-37	1543	S	3.56	23.0	2.085	
322-40	1440	S	3.53	23.6	2.085	66.0
322-41	1440	S	3.56	20.4	2.085	61.0
322-42A	1440	S	3.56	21.4	2.085	37.1
322-42B	1440	S	3.56	21.4	2.085	65.0
322-46	1440	S	3.53	19.6	2.08	
322-47A	1440	S	3.50	21.5	2.085	74.0

Sizes of the Structural Features Observed in TABLE 2 Bright and Dark Field Electron Micrographs Compared to Crystallite Sizes Obtained from X-ray Analysis

Sample #	Platelet Dia. A	Granu- lation* <u>Dia. A</u>	Dark F Dia. (002)		X-ray <u>Lc</u>	(Å) <u>La</u>	(002) Peak Type
311-19 (2000)	150-500	30-40	20-40				
311-19 (750)×	150-350	20-30			14	19	S
312-31 (2000)	200-500	20-45	20-45		27.6	56	S
312-31 (2000)	150	35	30	>100	28	56	S
317-24 (2000)	250	42	60†		24	45	NVS
317-29 (2000)	>250	60	30-70+		65-75		NVS
317-33 (2000)	250-500	35			92	49	S
317-45 (2000)	>500	30			25	60	3P
317-48 (2000) ×	250	55			34	59	3P
317-49 (2000)×	250-500	45	40†		33	60	3P
318-12 (2000)	250-500	60	50	110†	31	59	3P
318-22 (2000)	>500	40-60	35	<b></b>	65	55	NVS
318-22(700)	250				15.7		S
318-23 (2000)	250	50	50		63	73	S
318-23(700)					16	<del>-</del> -	S
318-29 (2000) ×	>500	30-40	60		31	63	2P
321-31C(2000)×	250	35	60	80	35	69	3P
321-31D(2300)	250	40	35	80	33 37	69	S
		7.5		30	51	כט	3

<sup>\*</sup>Diameter corresponds to distances between nearest neighbor. \*\*Diameter of diffracting regions obtained from (002) or (100) diffraction halos.

†Some of the crystallites giving rise to halos or spots are

very large in size, i.e., up to 500Å.

\*A second structural feature was observed in the bright field micrographs of these samples. This new feature appeared to be long regular cylinders 500Å in diameter by about  $1\mu$  long. Regular striations along the length were spaced 45Å apart.

TABLE 3 Electron Diffraction Results Compared to X-ray Diffraction Results for d(002) and d(10) Spacings (A)

Sample #	X-r d(002)	d (10)	Elec Diffra d(002)	tron ction <u>d(10)</u>	(002) <u>Peak Type</u>
Graphite 311-19(2000) 311-19(750) 312-31(2000) 317-24(2000) 317-29(2000)	3.35 3.56 3.70 3.54 3.57 3.50 3.43	2.13 2.17 2.19 2.12 2.10 2.10	3.37 3.45  3.53 3.53† 3.53† 3.35†	3.12 2.09 2.07 2.16 2.12 2.10+	 s s s nvs
317-33 (2000) 317-45 (2000) 317-48 (2000)	3.414 3.35 3.48 3.46	2.10 2.09 2.10	3.45 3.35† 3.50 3.48†	2.12 2.10 2.10 2.12	NVS S 3P 3P
317-49 (2000) 318-12 (2000) 318-22 (2000) 318-22 (700)	3.48 3.49 3.44 3.70	2.09 2.09 2.10	3.48† 3.47† 3.37† 3.50 3.42†	2.10 2.11 2.07 2.11	3P 3P NVS S
318-23(2000) 318-23(700) 318-29(2000) 321-31C(2300) 321-31D(2300)	3.43 3.74 3.45 3.43 3.47	2.10  2.08 2.11 2.11	3.50+  3.45 3.56+ 3.50*	2.10 2.07 2.12 2.12 2.125	S S 2P 3P S

<sup>\*</sup>In this sample no spots were seen on any diffraction halo.
†In addition to Debye-Scherrer rings, a number of sharp
diffracting spots were observed on or close to the ring.

TABLE 4

Sample	a i	n A	Dispersity
Emm San Sa	Our value	Reported Value	Dispersity
LMSC-20	42.4	17.1	Monodisperse
GC-10	14.0	9.0	11
GC-20	12.9	11.5	11

TABLE 5

Sample	a in A	Dispersity
312-31 (2000)	40.8	Polydisperse
317-45 (2000)	21.2	Wide distribution of pores
318-11 (2000)	20.4	Bidisperse
,	44.4	
318-41B (700)	18.7	Monodisperse
318-42 (2000)	41.0	Polydisperse
318-44 (2000)	30.4	Monodisperse
318-45 (700)	28.0	Monodisperse
318-45 (2000)	35.3	Monodisperse
318-46 (2000)	20.4	Monodisperse
318-48 (700)	9.5	Monodisperse
318-48 (2000)	25.4	Monodisperse

TABLE 6

Sample #	Temp.°C	He Density (gm/cm <sup>3</sup> )	Surface Area Knudsen Flow (m²/gm)	Specific Surface Area (m²/gm)
311-32	2000	1.41	3.0	26.4
317-9	700	1.83		506.0
317-9	2000	1.70	12.5	59.9
317-12	700	1.80	9.1	510.0
317-12	2000	1.72		109.0
318-22	700	1.79		459.0
318-22	2000	1.51		49.6
321-9	700	1.46		541.2
321-9	2000	1.36		12.7

TABLE 7

Sample #	Temp.°C	<sup>ρ</sup> He real¹ (g/cc)	<sup>ρ</sup> Hg real <sup>2</sup> (g/cc)	<sup>ρ</sup> Hg app. (g/cc)	MPD (µ)	IPV (cc/g)
GC No.1		1.47	1.482	1.424	.003	.0273
302-5	2320		1.509	.647	2.97	.8828
302-12	2320		1.501	.559	3.62	1.1224
305-6	2000	1.94	1.802	.636	2.54	1.0151
6.62 Mo						
305-12	2000	1.55	1.562	.557	4.19	1.1560
305-18	2000	1.77	1.718	.606	2.49	1.0678
.4 Mo						
308P-2 #2		1.586	1.505	1.034	.009	.3030
308P-3 #3		1.611	1.486	1.077	.008	.2559
310-1	1000	1.27	1.446	.814	.023	.5411
310-3	1000		1.424	.805	.020	.5454
310-17A	2000	1.50	1.175	.639	.119	.7130
310-18	1000	1.48	1.452	.687	.039	.7666
310-18	2000	1.15	1.366	.648	.044	.8110
310-20	2000	1.09	1.458	1.029	.009	.2855
310-29	2000	1.89	1.533	.944	.014	.3959
311-21	2000	1.59	1.339	.731	.038	.6221
311-22	2000	1.00	.847	.484	.154	.8809
312-19A	730	1.20	1.481	.879	.629	.4626
312-29	728	1.52	1.441	1.038	.014	.2709
312-31	2000	1.41	1.490	.923	.025	.4118
312-45	2000	1.26	1.302	1.214	.005	.0540
312-48	2000	1.53	1.392	.861	.121	.4425
312-49	2000	1.34	1.404	1.031	.011	.2579
315-1	2000	1.50	1.431	.962	47.0	.3412
317-5	2000	1.42	1.313	.873	.071	.4039
317-18	2000	1.50	1.255	.953	39.1	.281
318-22	700	1.79	1.426	.771	.057	.5958
318-22	2000	1.51	1.576	.937	.054	.4334
318-45	2000		1.20	.78	.0078	.021
321-9	700	1.46	1.24	.98	.0073	.205
321-9	2000	1.26	1.4	1.2	.0057	.016
321-17	2000	1.43	1.17	. 59	2.15	.299
321-18	2000	1.67	1.16	.87	.175	.247
321-19	2000	1.80	1.56	.98	.049	.379
321-20	2000	1.60	1.63	.70	.088	.345
321-21	2000	1.79	1.30	.85	.041	.377
321-25	2000		2.20	1.14	.011	.088

<sup>\*</sup>Glassy Carbon No. 1 - Le Carbone, p. 6927.

<sup>1</sup>Real density as determined by He pycnometry

<sup>2</sup>Real density as determined by Hg

Sample #	Temp.°C	<sup>ρ</sup> He real <sup>1</sup> (g/cc)	<sup>ρ</sup> Hg real <sup>2</sup> (g/cc)	<sup>ρ</sup> Hg app. (g/cc)	MPD (μ)	IPV (cc/g)
321-31	2000	1.41	1.49	1.34	.0195	.075
322-14A	1300				2.2	.826
322-14A	1412	1.74			1.7	.494
322-14B	1300				2.3	.501
322-14B	1412				1.5	.496
322-17A	1300				4.5	.604
322-17A	1412				4.4	.271
322-17B	1300				2.5	.382
322-17B	1412				2.0	.534
322-19A	1300				1.0	.461
322-19A	1412	1.9			.08	.472
322 <b>-19</b> B	1300				.65	.466
322-19B	1412				.95	.468
322-20	1300				1.8	.432
322-20	1412				1.5	.666
322-21A	1300				18.	.503
322-21A	1412				3.5	.420
322-21B	1412				10.	.400
322-21D	1300				8.	.780
322-22A	1300				1.1	.308
322-22A	1412				1.2	.443
322-22B	1300				1.4	.457
322-22B	1412	1.49X*			1.2	.440
322-23A	1300				1.5	.443
322-23A	1412				1.2	.458
322-23B	1300	2.03X			.32	.453
322-23B	1412				.35	.458
322-24A	1300	1.74X			1.3	.395
322-24A	1412				1.3	.563
322-24B	1300				1.9	.620
322-24B	1412	1.59X			1.4	.888
322-32	1350	1.60X			1.4	.571
322-35	1350	1.43X			6.0	.472
322-41	1440	1.59X			.07	.669
322-45	1500	1.72X			4.2	.421
322-46	1500				1.4	.550
322-47A	1500				1.3	.652
322-48	1605				6.0	.634
322-49	1400				7.0	.841
322-49	1400				7.0	.607
322-49	1600				7.0	.595
322-50	1600				6.0	.545
322-50	1400				6.0	.679

<sup>\*</sup>X indicates Xylene

Ult. Str. psi (x10-3)		1.01	7.04	1.23	4.85	1	2 83		0/./	<u> </u>	5.13	!	1,11	1 1		! !	!	<b>!</b>	+	5,96	1	0.36	) )     	;	4.7	נא כ		!	<u> </u>	<b>¦</b>	!	09.9	7.13
Compr. Str. psi (x10-3)		5.18	7.2	6.85	50.0	36.0	1 73	) • I	1 1	39.7	29.5	<b>!</b>	27.3		H		!	!	!	-	1	1.47	1	}	47.7	29.3			!	ļ.	<b>!</b>	!	46.8
Sonic Mod. psi (x10-6)			!	0.35	<b>!</b>	!	-	¦		!	!	!	!!	+	!	ļ		!	¦	!	1	1.27	1	1	-	;	į	1	4. 6	L.37	1.52	-	1.54
Int. Frict.		! !		I.43	!	!	;	<b>!</b>	ļ		!	! !	!	;	;			!	¦	1	<b>!</b>	1	!	!	<b>¦</b>	1	¦	0 0 0		1.03	0.54	!	0.26
Hard- ness (DPH)		<b>I</b>	! !	!	!	<b>¦</b>	!	!	!	ij		90		86	135	176	201	707	702	!	-	ł	!	<u> </u>	1	!	!	!			:	!	;
Resistivity N-cm(×10)	ł	1	1	467.	;	•	;	-	-	!		1	!	-	!	!	1		!	<b>;</b>	;	.349	!	!	<b>!</b>	1	1	.180	275		. 203	!	.147
ρ <sub>He</sub> (9/cc)	¦	!				L - 44	L.27	<b>¦</b>	1.52	1.47	פני		F. 50	8T•T	!	1.26	;	i	, ,	7 · · ·	L • 4	1.52	χ. Υ.	T.00	P.0	1 ;	1.6	1.6	1.60	1.60	) 	, L	70.1
papp.	.57	09		10			//-	.15	.07	90	. !	00	1		<u> </u>		!		9	0 0		0/.		90	000	٥,٠	. 84	. 84	.77	88	96.0	) [	T 6 .
Temp.°C	O	0	0		· C	0	2	<u> </u>	$\bar{}$	ā	$\overline{}$	_		5 :	~	=	~	=	<u> </u>	_	, ,	ى ر	2000	, C	) C	<b>)</b> (	<b>-</b>	0	0	0	0		•
Sample #	10-3	11-3	11-3	12	12-1	. ה ה	7 (	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	7-71	12 - 3	12-3	12 - 3	2-4	1 (	C#17-	1-4	.2-4	2-4	2-4	5-1	ď	י ע	315-4	5	7 .	יה הו		5-20	5-20	5 - 20	5	5-21	1

August

Ult. Str. psi (x10-3)	7.62	1	1	4.61	4.78	7.38	6.63	4.24	1	4.39	1	1	5.15	ļ	9.9		- 1	2.95		;	2.51	!	2.97	5.59	4.41	!	į	2.0	2.02	1	!	!	!
Compr. Str. psi (x10-3)	2.70	1	!	24.3		35.5	30.5	25.6	36.6	37.3	1	1	35.1	!	•	45.0	!	21.0	16.4	;	14.2	}	2.40	35.9	28.5	!	15.0	18.6	16.0	!	50.0	!	!
Sonic Mod. psi (x10-6)	;	!	!	!	;	1.55	1	1.20	1.38	;	!	!	1.44	1.65	1.60		1.26	0.87	.2	!	0.61	-	0.93	1.78	1.76	<u>}</u>	;	1.35	1.28	1.65	!	1.43	;
Int. Frict.	ļ	!	!	1	!	2.38	!	!	!	!	!	!	0.33	0.47	0.42	1	1.50	0.31	•	ļ	0.31	!	!	0.47	1.28	!	!	1.18	96.0	0.35	!	0.32	;
Hard- ness (DPH)	1	!	!		!	!	!	!	}	!	!	!	!	1		1	!	1	1	1	1	!	!	!	1	!	-	!	!	!	!	-	!
Resistivity -cm(×10)	1	:	1	1	;	.317	!	.057	.149	1	;	;	.119	.237	. 229	-	.195	. 294	.137	1	.262	!	.237	.188	.029	!	.220	.038	.157	.249	!	.214	1
P <sub>He</sub>		1.63	1.78	!	!	1.41	!	1.45	1.45	1.46	1.49	1.48	1.46	1.48	1.46	1.43	1.50	1.58	1.57	1.89	1.61	1.51	1.51	1.64	1.64	1.33	1.67	1.67	1.67	1.83	1.90	1.78	1.66
papp.	1.01	Ġ,	٦.	Φ,	Φ,	ω.	Φ,	0.80	Φ.	٠ ص	ο.	Φ.	Φ.	٠ و	σ.	σ.		9	9	œ	N.	. 7	7	9	0	φ	9	7	7	φ	0	7	ω
Temp. °C	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000	2000
Sample #	315-21D	15-2	15-2	15-2	15-2	15-2	15-2	15-2	15-2	15-2	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-3	15-4	15-4	15-4	15-4	15-	12-4	315-44	L5-4

Ult. Str. psi ×10-3)	H	1	!	2.23	7.5	2.97	7.50	1	1	2.29	5.77	00.9	5.82		4.69	4.09	3.30	8.75	;	1.90	4.37	2.85	0.92	-	;	2.48	-	-	5.05	5.60	4.61	1	06.9
Compr. Str. psi (x10-3)	ł	1	¦	2.5	56.5	23.7	33.1	-	!	40.5	32.3	43.7	-	-	27.4	33.6	5.1	28.2	-	7.6	49.1	34.1	4.7	37.3	<b>!</b>	16.4	1	1	44.2	40.2	24.0	!	40.6
Sonic Mod. psi (x10-6)	5.8	;	I		1	0.91	!	!	!	1.82	1	1.45	1	1	!	1	0.86	¦	;	1	1.45	1	1.51	1		0.86	;	ļ	1.64	;	2.07	!	1.61
Int. Frict. (×10³)	0.28	!	-	1	-	1	!	!	!	!	!	0.75	!	!	!	1	0.39	1	!	!	92.0	1	99.0	1	}	!	1	1	1	;	1	1	0.31
Hard- ness (DPH)	1	240	105	28	!		28	!	1	!	!	!	!	!	!	!	;	1	!	!	!	!	14	!	1	!	!	<b>¦</b>	1	73	!	!	80
Resistivity $\Omega$ -cm(×10)	.039	<b>!</b>	;	!	;	.088	<b>!</b>	;	;	<b>!</b>	<b>!</b>	600.	!	!	;	!	.088	1	1	:	.187	;	.195	:	1	.122	;	1	.224	1	.321	;	. 225
ρ <sub>He</sub> (g/cc)	1.39	!	!	1.55	1.67	1.74	1.42	1.88	1.82	1.64	1.76	1.42	1.60	1.88	1.49	1.46	1.50	1.68	1.51	!	1.57	1.69	1.48	<b>¦</b>	1.7	1.65	1.68	1.45	1.72	1.46	1.56	1.40	1.34
Papp.	.7	.09	•	. 89	.2	.7	0.78		.7	6	9	. 7	φ.	φ.	φ.	σ.	.7	4	0.	ω.	-	œ		!	9	.7			φ.	0	0.65	9	9
Temp.°C	00	00	00	00	00	00	2000	00	00	00	00	00	00	00	00	00	00	00	00	9	00	00	00	00	9	90	00	90	00	90	00	0	00
Sample #	15-	15-4	15-4	15-4	17-	17-	317-5	17-	17-	17-	17-	17-1	17-1	1.7-1	17-1	17-1	17-1	17-1	17-2	17-2	17-2	17-2	17-	17-2	17-2	17-2	17-3	17-3	17-3	17-3	17-3	17-3	17-3

Ult. Str. psi (×10 <sup>-3</sup> )	4.20	3.40	2.35	6	3.90	5.30	•		4.95	5.62		3.0	2.56	1	4.77	1	;	79-1	5.62		5.19	1	5.09	1	1	0.83	1	4.02	4.35	1	5.28
Compr. Str. psi (x10-3)	37.6	22.8		7.4	27.0	39°8	27.3	32.3	34.6	27.5		11.1	;	!	34.5	ı	!				27.4	!	1	!	!	4.74		28.2	33.2	1	!
Sonic Mod. psi (×10 <sup>-6</sup> )	1.01	1.25	1	!	1	1.54	1.35	)     	1.27	1	1.23	0.89	0.71	¦	0.80	1	1	0.65			ļ 3	1	1.48	1	;	0.41	1	1.43	1.18	!	!
Int. Frict. (×10³)	1.27	; ;	1		!	1 1	1.68	1	1	1	1	1.31	1		{	}	1	!	;		+	1	!	1	!	0.41	1	}	0.18	!	!
Hard- ness (DPH)	62		;	!	1 1	ا کے ا	52	1	71	49	!	1	!	1	1	!	!		09	}	26	!	51	19	71	31	40	47	53	46	ł
Resistivity N-cm(×10)	.268	) ω	1	!	1 °	.135	.007	;	.112	;	1	.34	.169	<b>¦</b>	907.0	;	1	.165	!	1	1	!	!	:	:	.285	:	.270	.189	;	+
θH <sub>σ</sub>	1.34	1.47	!	•	1.48	1.40	1.51	1.40	1.48	1.39	1.46	1.51	1.51	1.45	!	1.37	1.45	1	1.49	<b>¦</b>	1.50	1.48	1.58	1.50	!	1.47	1.51	1.48	1.46	1.48	:
Papp.	0.92	ω.	9	י נכ	ન ∘	0	α	ထ	φ.	٥	Τ.	00		0	φ.		7		ο.	ο.	96.0	ο.	σ.	0	0		6	6		!	•
Temp.°C	2000	ŏ	56	58	56	50	$\approx$	ŏ	ŏ.,	$\ddot{c}$	<u> </u>	2	ָ כ	$\sim$	io i	9	8	20	9	9	9	9	$\circ$	0	9	9	$\mathbf{o}$	0	$\circ$	$\geq$	9
Sample #	317-38 317-39	17-4	17-4	7 <del>7 - 7 - 7 - 7 - 7 - 7 - 7 - 7 - 7 - 7</del>	1 / - 4 7 / 7	17	17-4	17	17-	-	- 1	2	8	8	φ.	φ.	8	8	<u>φ</u>	φ.	ᅇ.	ᇷ	ᅘ	ည်း	φ.	φ.	φ,	۳ ا	p o	p d	Ď

Ult. Str. psi ×10-3)				1 ,	4.3/	l		4	7. 1		<b>1</b>	1 7	2.0	28.0	0.19			4.85	4.57	3.34	3.90		7.95	!	<b>!</b>	!	¦	7,35	, (	D . C . 7	- 41	F • 4
Compr. Str. psi x10-3)	i				C.67		<b> </b>	20	0 1	¦ ¦		200	2.63	6.12	٦. ١٠٥	19.7	32.2	28.7	26.4*	18.6*	25.7*	İ,	34.9*	!		!	<b>¦</b>	42.3*	, v	)   •   )	17.3	)
Sonic Moa. psi x10-6)	1	ļ	¦	סכר	C7.T	<b> </b>	7 49			1	1 33	) 	כח	L.02	0.135	[   (	1.3/	'	1.48	1.35	2.17	G	66.7	!	!	ı	•	2.34	i		0.83	)
Int. Frict.	;	;	¦	ŀ		!	}	¦	!	• ¦		¦		רף כ	T # - 7	1 1	0.73	!	!	!		!	!	<b>!</b>	<u> </u>	!	<b>!</b>	į	!	!	-	
Hard- ness (DPH)	<b>!</b>	65	יי פיי	44	36	) T.	19	-	7.0	)  -  -	¦	26	9 9	5 6	4 C	ט ת ט ג	/ 0	!		/ 9		!		0 0	907r	707	!	26	!	!	1	
Resistivity $\Omega-cm(\times10)$	!	1	<b>!</b>	237		!	!	1	1	1	.177	194	 	216	017.		T 0 1 1	0.	T. L.	7°-	71.1	น เ	. i				.39	.41	!	;	;	
(b) (b) (b) (b) (c) (c) (d) (d) (d) (d) (d) (d) (d) (d) (d) (d	1.41	1.50	1.37	1.45	1.48	1.49	1.29	!	1.59	1.38	1.45	1.45	1.49	1.31	7.57		1.45	7.7	7 - V	- T	4. t	7. T	7.44	· • I	ļ		!	!	1.43	!	1.43	
$\rho_{\text{app.}}$	0.77	ı	!	Ψ,		0.91	9	9	9	ω.	Φ,	9	0	ഹ	α	0		α	9 0	0	1	C		2	60.	, c	7.	0	0		0.88	
Temp.°C	2000	2000	2000	2000	7	2000	ō	ō	5	5	ŏ	ŏ	ŏ	$\sim$	$\sim$	$\sim$	$\simeq$		: <b>:</b> :	ς ς	? >		$\sim$	0	9	2	2 9	⊇ ′	0	Ō	0	
Sample #	318-19	18-	18-	18-	18-	18-	18-24	18-	18-	18-	 18-1	L8-1	-87	8-	8	8	8	& -	00	α	2 0	, en	8-4	8-4	8-4	2-4	ר יי ס כ	χ. 4	20 1	8-5	8-5	

\*Head Speed .05 in/min., all others .02 in/min.

Section of the second

Total Park

ult. Str.	$\frac{\text{psi}}{(\times 10^{-3})}$	5	1.60		C	•	9	6	6	00	Н	-	8.27		0.		- α	5.1		, r	6.04	6	S		5.26	:	5,67	. 0	77. 7	:	C	6.35
Compr. Str.	$\frac{\text{psi}}{(\times 10^{-3})}$	•	4	;	!	7		7	1	3	-	1	41.7*	ı	ω.	4	•	0	7	4	36.2	4	4	i	36.7		39.6	•	31.4		34.8	. N
Sonic Mod.	$\frac{\text{psi}}{(\times 10^{-6})}$	1.66	1		0.10	1		4.	-	;	!	!	2.33	- 1	1.48	!	2.99	•	1	1.22		1	1.55	- 1	1.81		1.73	1.83			!	1.65
Int.	Frict. (×10³)	1	-	1	!	!	1	1	1	!	1	;	1	;	1	!	;	!	!	1	!	<b>¦</b>	0.21	1	0.22	1	0.2		0.11		1	1
Hard-	(DPH)	1		-	-	1	54	1	69	1	78	!	81	-	105	120	66	95	132	;	131	1	}	!	!	1	!	115	87		!	;
	Resistivity A-cm(×10)	1.30	1	!	2.37	1	1.50	.403	;	1	.340	;	.31	ļ	.546	1	1.00	4	!	1.21	1.15	:	!	;	1	;	;	;	1	1	ľ	.2
0	(g/cc)	1.41	-	1.34	i	1.38		1.75	1.39	;	1.57	1.52	1.60	1.54	1.46	1.36	1.34	1.43	1	1.32	1.48	1.56	•	. 7	1.41	9	;		1.80			1.74
٥	(g/cc)	0.	ဏ	φ.	6	1.18	6.	0	6	1	6	6	0.	6	06.0	7	7	6	-	٠.	0.95	0.	ထ	0	0.	6	•	ထ	φ.	6.	.7	6
	Temp.°C	2000	ŏ	ŏ	ŏ	ŏ	ŏ	ŏ	5	$\ddot{\circ}$	0	$\tilde{c}$	$\frac{1}{2}$	$\sim$	$\sim$	2		0	0	2	2	9	9	9	9	9	9	0	0	0	0	0
	Sample #	318-52	18-5	18-5	18-5	18-5	18-6	18-6	9-87	7-17	7:	21-	7.7	-77	7.	5 – T :	21-1	21-1	21-1	21-1	77	1-1	21-1	7-7	1-1	1-1	1-1	1-1	1-1	:1-2	1-2	1-2

\*Head Speed .05 in/min., all others .02 in/min.

ult. Str.	psi (×10 <sup>-3</sup> )		77.9	!	<u> </u>	5.89	4.28	S	5.96	9	;	5	2.92		7.43	24.0	7.40	1.23	!	ï	٠.	4	4	0.22	4	3	1	;	0.57	0.42	0.37	0.22			!	;
Compr. Str.	$\frac{\text{psi}}{(\times 10^{-3})}$	-	4. I	! !		41.9	33.3	31.8	39.8	53.6	!		33,3		40.9	• 1	705	•		1 1	5.53	2.51	1.39	1.05	1.67	6.80	!	;	2.05		3	7	-	<b>¦</b>	;	!
Sonic Mod.	psi ×10-6)	0	0.00	י ר	7 1	٠,	0.91	.5		1.49	!	1.49	2.94	1.46	0.92		1.15	1		'	0.42	1	0.94	0.08	0.2	0.72	1	ŀ	0.35	7	9	7		;	!	!
11	Frict. (×10³)	ł	;	!	!	!	¦	!	;	1	<b>!</b>	;	;	!	-	!	;	ļ			!	!	!	<b>!</b>	!		!	!	!	!	!	;	!	;	<b> </b>	ŀ
Hard-	(DPH)	ł	†	!		1	¦	<b>¦</b>	1	!	!	1	1	!		-	;	¦	¦				!	!	!	<u> </u>	! !	1	1		!		!	1	!	!
	α-cm(×10)	.21	.25	. 29	70	* V	1 6	77.	•16	m	!	.27	.18	!	.38	;	. 29	-	ł		₹ 7 1 • I	-	*	<u>.</u>	T. C.		00.	(1)	. 29	87.	.36	• 46	!	;	;	<b>!</b>
o <sub>H</sub> O	(g/cc)	1.25	ŀ	;	1,33	) 			1.41	1.30	٠, ٢	T.59	!	1	1.21	1.80	1.66	1.43	1.44	1.50	)   	!	ן קפע	•	ואט	1.00	75.1	70.1	1.44X		1.55	1.44	18.1	1.56	1.78	1.84
d ann	(၁၁/6)	•	•	•			•	•	•	0.0	•	•			_	:	_	٠.	_	_	•	•	,			. 4	ין נ	, [	•	יו כ	•	9 0	, ע	0.98 2.98	٠, ١	0
	Temp.°C	2000	0	0	0			· ~	) (	) M	١ ۸	~ ~	<b>م</b> د		~ ~	~ .	$\sim$	$\overline{\sim}$	=	=	=	=	$\sim$		_	_	_	_	, C	, C	) C	) C	<b>)</b> (	$\supset$ $\circ$	<b>)</b> (	_
	Sample #	321-32C	21 - 32	21 - 32	21-32	21 - 32	21 - 32	21-33	21-22	21-34	7-10	17	7-10	מדר די	7C-T2	00111	95-T;	T-36	1-37	1-37	1-37	1-37	1-37	1-3	1 - 39	1-4	1-41	1-42	1-42	1-43	1-43	1-44	7 7 T	# V 	יים דיור	T 40

Ult.	psi (x10-3)		1	0.49	; ;	¦	!	i		0 9 0	•	•	i	! !	1	!	4.52	1	1.36	) 	1	i	0.78	!	!	!	¦	!	i	ŀ	!	   		<b>:</b>	!	-
Compr.	psi (×10 <sup>-3</sup> )		!	2.04	!	·	;	i	1	2,65	)	i		!	i	!	28.2	!	9.1	1	;	;	4.00	!	;	!	!	!	1	!	¦	!	!	!	!	;
	psi (×10-6)																																			
Int.	Frict. $(\times 10^3)$		i	;	1	;	!	;	;	;	!	;	ļ	ļ		!	<u> </u>	!	-	1	1	1	!	;	1	!	!	!		!	;	1	-	!	!	;
Hard-	ness (DPH)		1	1	¦	!	!	!	!	!	!	1	!	!		1	!	i i	!	!	!	i		;	!	!	1	!	!	!	<b>!</b>	-	-	<b>;</b>	!	!
	Resistivity $\Omega$ -cm(×10)																																			
Ö.	(g/cc)	1.40	0 V V		7.0	70.7	70.7	L.04	1.40	1.43	T.58	1.51	T.44	1.51	1.69	1.43	7 7 7			L.33	7.0	70.7	1.98	20.0	70.7	2.0.0	ا ر ب ب		7.4T	7.0	1.49X	1.9	1.43X	X.20.7	1.40X	T • / 4
9	(g/cc)	.83	82	200	٠ ۲	α	2 6	4 6	, , ,	0.0	0 5	100	200	91	00	02	0.3	66	) [	<del> </del>	.12	777	0.77	.87	7.8	78	98	2	200	7 0	7 0	היר	10	7 7	27	2
	Temp.°C																						1600													
	Sample #	1-46A	21-46B	21-46C	21-47A	21-47B	21-47C	21-48A	21-48B	21-48C	494-F	1-49B	מסער דר	7 6 7	7-20	1-50B	1-50C	1-51	1-51A	1-52	1-53	2-1A	2-1B	2-2A	2-3A	2-3B	2-5	5-6	2-10C	2-11A	2-11B	-12A	-12B	-13A	-14A	

ult. Str. psi (x10-3)		:	-	;	;	1	1	!	1	!	1.19	1.19	!	4.07		1.28	-	!	!	-	3.16	1	1	-	0.79	!	!	1.33	3.74	!	!	!	<b>{</b>	<b>!</b>
Compr. Str. psi (x10-3)		!	;	:	;	;	1	!	ł	;	4.1	5.2	1	;	!	-	-	1	1	-	18.4	1	!	1	5.8	1	<b>¦</b>	11.8	23.0	!	!	:	<b>;</b>	<b>;</b>
Sonic Mod. psi (x10-6)	1	0.57	1	•	7	2.17	;	0.33	0.77	0.53	-	-	1	}	1	!	1.56	1.48	2.04	1.58	!	9.	0.79	.2		9.	0	9.	1.29	4.	0.93	0.72	1.14	0.84
Int. Frict.		!	1	;	!	1	1	!	!		!	!	1	}	1	!	1	!	!	!	1	!	!	}	1	!	!	!	!	!	!	}	1	1
Hard- ness (DPH)		1	!	!	!	-	!	}	1	<b>!</b>	1	-	-	ł		1	1	1	!	1	!	1	!	!	!	!	!		!	!	!	1	1	1
Resistivity Q-cm(×10)		61.	-	.22	.37	60.	-	.28	.17	.22	<b>¦</b>	<b>!</b>	1	1	1	ľ	.11	.13	.11	80.	.10	.19	.18	.34	.24	.22	.13	.22	.10	. 20	.17	.18	.11	.17
ρ <sub>He</sub> (g/cc)		1.48X	1.89	1.81x	1.71x	1	1.9	-	!	!	1	1.49X	ł	2.03X	1.74X	1.59X	!	!	!	1.42X	1.44X	1.46X	1.64	1.58X	1.60X	1.78X	1.34X	1.43X	1.45X	1.46X	;	-	1.86X	1.59X
$\rho^{\text{app}}$ .		∞.	φ.	φ.		0.	φ.	0.89	φ.	φ.	φ.	φ.	φ.	φ.	φ.	φ.	φ.	φ.	0.	ω.	6.	9.	. 9	.7	.7	.7	φ	φ.	6	φ.		φ.	ω	
Temp.°C	- ;	9	67	67	67	67	67	1670	40	40	40	40	40	30	30	40	41	67	40	40	40	40	47	47	35	35	35	35	54	54	54	14	44	7
Sample #		22 - 1	22 - 1	22 - 1	22 - 1	22-1	22-1	322-19B	22-2	22-2	22 - 2	22 - 2	22 - 2	22 - 2	22 - 2	22 - 2	22-2	22 - 2	22 - 2	22 - 2	22-2	22 - 2	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22 - 3	22-4	22-4

ult. Str.	psi (×10-3)	}	1	1	1	3.11	1	!	!	!	!	!	!	!	1	!	!	!	!	1	ł	1	1	1		!	ţ	!	1	!	}	!	<b>!</b>	<b>¦</b>
Compr. Str.	psi (×10-3)	;	1	1	!	19.4	1	!	!	1	ļ		<b>¦</b>	1	-		ł	¦	!	-	-	;	;	;	1	1	!	1	!	!	<b>¦</b>	<b>!</b>	<b>!</b>	1
Sonic Mod.	psi (×10-6)	0.88	0.68	0.69	0.87	0.89	0.70	0.76	0.97	0.68	0.43	0.79	0.56	1	1	!	!	i	0.4	1.21	1.69	2.47	1.92	1.92	1.89	;	¦	1,32	<b>¦</b>	1,33	0.53	<b>!</b>	1	0.42
Int.	Frict. (x103)	;	!	;	;	ì		1	!	!	1	1	1		1	!	1	!	!	!	1	;	!	†	}	<u> </u>	<b>¦</b>	!	<b>!</b>	;	1	<b>¦</b>	;	<b>;</b>
Hard-	ness (DPH)	;		!	;	!	-	!	!	!	1	1		;	-	!	1	!	!	1	1	<b>!</b>	1	<b>!</b>	!	!	¦	1		!	!	<b>!</b>	;	<b>!</b>
	Resistivity A-cm(×10)	.14	.23	.20	.18	.27	.17	.19	.19	. 20	.29	.15	.15	;	ł	i	;	;	;	!	;	;	;	<b>!</b>	-	;	!	•	;	;	<b>!</b>	!	<b>!</b>	1
•	(g/cc)	;	!	!	1	!	!	!	!	1.72X	!	1.51X	!	1.56	1.48	1.33	1.63	1.85	1.50	1.62	1.56	1.49	1.60	1.55	1.61	1.45	1.37	1.64	1.65	1.84	1.52	1.25	1.48	1.50
•	papp.	7	0.73			ω.	7		ω	0.				.7	0.	σ.	0.	0.	7	0	0	٦.	0	7.	σ.	2	0	∞.	ω.	ω.	7	7.	7.	7
	Temp.°C	44	1440	44	44	44	44	44	Ž Ž	44	9	9	9	46	50	50	50	50	50	50	50	50	37	37	37	37	37	35	37	37	37	37	37	37
	Šample #	22-42A	322-42A4	22-42B	22-42B	22-42B	22-42B	22-42B	22-42B	22 - 4	22 - 4	22 - 4	22 - 5	22 - 5	22 - 5	22 - 5	22 - 5	22 - 5	22-6	22-6	22-6	22-6	22-6	22-6	22-6	22-6	22-6	2	22-67	22-6	22-	22-6	22-68	22-6

X

Ult. Str. psi (×10 <sup>-3</sup> )	ł	¦	<b>:</b>					I	ŀ	!
Compr. Str. psi (x10-3)	ł	ŀ	\	¦	ł	\ 		;	;	;
Sonic Mod. psi (x10-6)	0.42	38.0		זוק	30	) ) •	2.48	25.0		) 
Int. Frict. (×10³)	1	;	-	}	;	;	1	!	<b>¦</b>	;
Hard- ness (DPH)	ł	<b>¦</b>		!	<b>¦</b>	!	!	!	!	;
Resistivity N-cm(×10)	ŀ	}	ł	;	!	!	1	!	1	;
ρ <sub>He</sub> (g/cc)	1.56	1.56	1.47	1.51	1.50	1.36	1.61	1.49	1.49	1.52
ρ <sub>app</sub> .	0.76	0.69	1.01	0.81		0.98	1.13	0.78	0.76	0.95
Temp.°C	1370	1370	1370	1370	1370	1370	1370	1370	1370	1000
Sample #	322-69A	322-70	323-1	323-2	323-2A	323-3	323-3A	323-4	323-4A	323-8A

Physical Properties Correlated with Density
TABLE 9

$\rho\left(\frac{\rho_{\mathbf{a}}}{\rho_{\mathbf{He}}}\right)$		<u> </u>	!	<b>!</b>	<b>!</b>		<u> </u>		ı <u>1</u>	¦ ¦		-	010.		!	013	0 0	<b>O</b>	600		1	ļ		040			<b>-</b>	1 <	<b>D</b> r	.015	- I
$\sigma_{\mathrm{UTS}}\left(rac{ ho_{\mathrm{He}}}{ ho_{\mathrm{app}}}\right)$				! !	<b>!</b>	4 67	• 1	!	~	•	•		1.0	. !		ł	ľ	ļ	11.9	i	;	<b>¦</b>	8	. i	7 7			0 5	•	ı	6.5
$\sigma_{\rm cs}\left(rac{ ho_{ m He}}{ ho_{ m app.}} ight)$	ľ	<b>!</b>	   	ŀ	_	28.5	. !	56.4	47.7	40.9	i		71.2	1	1	!	-	ľ	78.2	1	!	!	56.9		v	. ~	7. 7.	7.00	r • 1	0	65.0
$\mathbf{E_{s}}\left(\frac{\rho_{\mathrm{He}}}{\rho_{\mathrm{app}}}\right)$	;	;	ł	<b>¦</b>	-	1	!	1	1	-	!	5 76	1	1	φ.	2.85	7	1	2.51	-	}	ļ	2.41		7	2.34	.	9	ע פ	2.56	
$\sigma_{\rm UTS}^{/\rho}{\rm app.}$	6	י רע י	6.99			0	188.0	-	8	33.	0	4	136.0	œ		+	!	91.	217.6	.60	45.	52.	32.	960	47.	!	27	178.8		01.	178.2
$\sigma_{cs}/\rho_{app}$ .	52.	'n	37.	298.	1000.0	24.	ł	30.	901.2	24.	1	φ.	1235.5	030.	-	<b>!</b>	!	;	28.		67.	 	20.	62.	888.9	224.	079.	13.	;	105.	1262.6
$E_s/\rho_{app}$ .	;	<b>¦</b>	19.2	1	!	!	;	2	:	1	}	50.3		l	48.8	· 0	Ф	ł	45.8	!	!	-	48.8	-			ł	•	6	48.8	;
Sample #	10-3	11 - 3	11 - 3	12 - 1	12.	12-1	12-2	12-2	12 - 3	12 - 3	12-4	15 - 2	[2-]	15-17	5-20	5-20	5-20	.5-21	5-2	$\frac{5-21}{2}$	5-25	5-25	5-25	5-26	5-26	5-26	5-2	5-3	5-31	315-31D	5–3

Specifical Services

Time Control of

$\rho\left(\frac{\rho_{\mathbf{a}}}{\rho_{\mathbf{He}}}\right)$		.009 .001 .011		010	000		.001	.004	.009 .016 .009
$\sigma_{\mathrm{UTS}}\left(rac{ ho_{\mathrm{He}}}{ ho_{\mathrm{app}}}\right)$		0 6 9 6 7 0 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6				V M 4 0		• • •	9.0 1.7 7.0 8.0
$\sigma_{cs}\left(\frac{\rho_{He}}{\rho_{app}}\right)$	1 10 6	43.1 50.3 61.3	10.0	91.3	47	56.1 73.8 61.1	.	911	. 65.5 36.6 85.4 57.4
$E_{s} \left( \frac{\rho_{He}}{\rho_{app.}} \right)$	2130	1.85 1.95 3.03	2.93	*   "	5 ' 6	1 1 1 6	1     1	2.99	1 8 9 4 1
<sup>0</sup> UTS <sup>/ p</sup> app. in(×10 <sup>-3</sup> )	36.	n	72.2 69.9		NNN	66 70 72 10	1	5000	89.9 32.8 92.8 113.0 152.0
ocs.	72 90 44	925.9 1038.8 824.7 612.0	671. 563.	1335.5 	297. 922.	11/5.0 1247.0 964.8 1536.6	74 74 25	6 4 4	10/6.4 167.4 615.0 1380.0 1092.0
Es/p	440.3 51.5 31.9	10-1-1:	48.7 45.9 52.7		35.6		33.0	52.9	53.7 32.3 51.3
O1 I		315-38A 315-39A 315-39B 315-41	1 4 4 4	7 <b>5</b> 5 5 6	# - 01.10	317-8 317-9 317-10	317-14 317-15 317-18	317-19 317-23 317-24 317-25	317-26 317-29 317-32 317-33

$\rho \left( \frac{\rho_{\mathbf{a}}}{\rho_{\mathbf{He}}} \right)$ $\Omega - \operatorname{cm} (\times 10)$	_	ı –	۱,	010.	<b>D</b> ,	⊢	<u> </u>	!	1	800.	1	.0004	1	900.	!	1	.018	0	:	<b>;</b> 1	ł	<b>!</b>	1	1	.015	-	Н		;	1		0	1 0	<b>O</b>
$\sigma_{\rm UTS}\left(\frac{\rho_{\rm He}}{\rho_{\rm app.}}\right)$	_	•		11.9	•	•	!		5.15	φ.	•	œ	•		•	!	5.7	•	!	!	8.7	İ	•	•	1.6	m.	•	•	1	1	!	.7	8 0.8	Ç.
$\sigma_{cs}\left(\frac{\rho_{He}}{\rho_{app.}}\right)$	7		֓֞֜֞֜֜֜֜֓֓֓֓֜֜֜֓֓֓֓֓֜֜֜֓֓֓֓֜֜֓֓֓֜֜֜֓֓֓֜֜֓֓֓֜֓֜	54.76	4	ω	†	!	5.	66.3	3	•	÷	63.2	6	<b>!</b>	20.9	<b>!</b>	!		28.2	ı	42.8	-	0.6		65.5	51.5	1	;	;	•	30.2	•
$\mathbf{E_s} \left( \frac{\rho_{\mathrm{He}}}{\rho_{\mathrm{app.}}} \right)$	d		4.	1.47	6	۲.	!	!	1	2.63	:	2.43	1	2.32	<b>!</b>	•	1.67	•	;	<b>¦</b>	<b>!</b>	<b>!</b>	!		.76	.2	9.		6	;	2.30		2.10	.32
σurs/ρapp. in(×10-³)	,		17.	9	29.	11.	0	00	96.5	6	76.	0	64.	215.0	60.	;	4.	•	148.9	•	3	1	150.2	•	29.9	•	•	•	;	129.0	1	•	149.62	•
σ <sub>cs</sub> /ρ <sub>app</sub> . in(×10 <sup>-3</sup> )	L	20	049.	158.	63.	36.	98.	28.	0.899	.99	462.	03.	017.	83.	85.	ļ	385.4	¦	ė	• 6	2	36.	2	<b>¦</b>	70.	31.	4	7.	1	715.0		0	563.3	0
Es/papp.	ć	· 0	ъ	31.3	е М	0	•	;	1,	49.1	-		1	43.5	-	6		4.	5	m "	-	1	;	5	14.6	2	4.	3	5	-	43.9	I	39.2	•
Sample #		1/-3	17 - 3	17 - 3	17 - 3	17-4	17-	17-4	17-4	17-42	17-4	17-4	17-4	17-	17-4	17-4	17-4	18-1	18-	18-7	18-	18-	18-9	18-1	18-1	18-1	18-1	18-2	18-2	18-2	18-28	18-2	318-30	18-3

P P P P P P P P P P P P P P P P P P P	Ω-cm(×10)	<b>¦</b>	!	;			.070	0	;	;	;	;	600.	1	;	1		.023		.021	2	.034		0	7	680.		1	!	1	!	!	;	;
	psi(x10-3)	1	;	1,13	7.4	•			<b>!</b>	!	;	2.3	3.55	2.7	ŀ	!	13.7		•	11.9	12.1	11.4	11.3	11.5	•	3.4	9.4	7.6	8.6	7.9		11.7	12.5	
	psi (×10-3)	36.5	l	•	42.9	6	•		;	1	•	28.2	•	7.9	1	6	5	39.2	0	9	į.	8	8	φ.	0	0			٥.		1		68.1	
$\mathbf{E_s}\left(\frac{\rho_{He}}{\rho_{app}}\right)$		1	<b>!</b>	;	4.	٦.	3.50	•	!	1	<b>!</b>	7	2.35	1		•	•	2.55	ł		4.	2.39			٣.	1.67	5	1	3	2.71	-	3,53	•	!
ours/papp.	in(×10 <sup>-3</sup> )		9	2	144.0	03.	$\vdash$	78.	1	200.0	20.	5	6	i.	δ.	98.6	22.	6	01.	09.	10.	15.	31.	38.	51.	2	.94	73.	48.	55.	46.	3	9	4
ocs/papp.	in(x10-3)	50.	18.	29.	831.0	73.	73.	78.		6	3.	46.	486.8	51.	1	44.	55.	620.0	195.	166.	61.	197.	283.	207.	184.	427.	32.	94.	792.	.680	726.	96.	055.	381.
Es/papp.	in (×10-6)	1	46.9	1	9	5	65.7	4.	7	5	!	•	46.2	1	•	50.6	2	0	!	<b>!</b>	59.5	5		0.99	4.	5.	о Ф	-	0	3	0	58.4	0	;
	Sample #	18-3	18 - 3	18 - 3	318-35	18 - 3	18 - 3	18-3	18-4	18-4	18 - 4	18 - 5	18 - 5	18-5	18-5	18-5	9-81	18-6	18 - 6	21-	12-	21-	21 - 9	21 - 1	21 - 1	21-1	21-1	21-1	21-16	31-17	21-18	$\sim$ 1	21-19	21-20

$\rho\left(\frac{\rho a}{\rho_{He}}\right)$ $\Omega - cm(\times 10)$	600.		1	, ,	1 0	200.	1 1	$\vdash$		1	600.	1	3 5	4 -	. כבס	ין ר	4 -	. C. C.	1 1		1	.015	01		1	.018		1	.011	7		
	12.1	, -i	7.2	•	7	· ·		11.1	1	4		2.2	5	•	•		•	∞ (	1.36	1	•	7.5	0	1	i	•	6.9	1				
$\sigma_{cs}\left(\frac{\rho_{He}}{\rho_{app.}}\right)$	9.98	60.	54.7	10	÷	œ	1	٠,	,		~	10	7	œ	ö	ė	e,	33.6	i.	1	0,	51.4		:    -		v	200	. !	7	78.3	,	
$\mathrm{E_{s}}\left(\frac{\rho_{\mathrm{He}}}{\rho_{\mathrm{app.}}}\right)$	0.0	4.21	:		2.5				•				•	•	1.6	1	7	1.56	0	ļ	۲.	1.36	7	7		1	2.12	4	1	מר כ	7.10	
σ <sub>UTS</sub> /ρ <sub>app</sub> .	17.	171.1		, ,	. o	4.	84.	71.	93.	12.	0,0	30.	•	• -	, 60	•	•	. ~		1 10	m	Ä	Q	ω	!	1	.99	23.	127.1	70.	0	
ocs/Papp.	344.	1288.9	ش ر	986.	າ ເ	. u	200	7	443	482	170	738	، م	313	> 0	2 0	ָר ע	2. 1	`	7 7 7 7	7	49	67	10	1	1	87.	963.	892.3	39.	673.	
$E_{\rm s}/\rho_{ m app}$ .	48.	53.6	5	!	·	.; ·	•	4.84	ο α	i	7	œ	!	0	46.0	9	!	· ·	-			, u	• v		4	LC.			43.8		46.5	
Sample #	4,5	1, 1	12	-2	12	[-2				<u> </u>	1-24	1-2	1-26	1-2	1 - 2	1-31	1 - 31	1-31	1-3]	1-3]	1-3	7-7	) (	) (   T	7	10	1	C - T 2	21-12 21-3	21-3	321-33B	

	$\rho\left(\frac{\rho_{\mathbf{a}}}{\rho_{\mathbf{He}}}\right)$	)	.016	;	1	.030	<b>¦</b>	_	.011	.020	;	.012	.013	.013	.012	.012	.018	.020	.015	.014	.011	1	.014	.012	.015	600.	;	.011	.014	.012	600.	.011	<b> </b>		1
	$\sigma_{\rm UTS}\left(\frac{\rho_{\rm He}}{\rho_{\rm app_*}}\right)$	1	13.0	!	1 .	•	m .	$\frac{11.2}{1}$	•		.50		2.5	•	1.07	96.	.75	ų.	•	•	6.3	•	;	1.6	-	1	!	!	:	1	;	1	ŀ	;	:
	$\sigma_{cs}\left(\frac{\rho_{He}}{\rho_{app_{\bullet}}}\right)$		54.1		ָרַ הַּ	0.10	1 0	78.7	12.6	•	•	ນ ເ ວ ດ			•	5.6	•	•		4.	39.5	3	ı	œ ش	ľ		:	!	:	<b>!</b>	<b>!</b>	1	;	:	<b>!</b>
•	$\mathrm{E}_{\mathrm{S}\left(egin{array}{c}  ho_{\mathrm{He}} \ \mathrm{sp.} \end{array} ight)}$	0			זר	7	7.0	0 / · T	2.12	31.7	4 <	* 4	•	ı	•	9 (	, (	57.	87.	•	2.1	•	, ì	90.	70.	₽.4 0	4. 1 00 (0	.58 8.5	4 30 .	9,	T.T	86.	.75	.45	<b>¦</b>
	σ <sub>UTS</sub> /ρ <sub>app</sub> . in(×10 <sup>-3</sup> )	0	84.5		155.5	61.6	187.7	45	2	ω,	ľ	42.8			•	13.57	•				200		5	T0.7				! !				!	<b>:</b>	<b>!</b> II	! !
	$^{\sigma}$ cs/ $^{\rho}$ app.	45.	963.5	1	1171.2	ł	08.	2	6	2	2	222.2		ω.	8	47.9				6	•	• } !	144.3		!	;	;	ł	;	ļ	;	ļ		ł	
	Es/papp. in(x10-6)	5.	85.1	7	ė.	ŀ	6	•	33.5	•		23.5	1	•	•	21.9				0		!	7.		0	œ			2	0				59.7	
	Sample #	321-34A	2	7	į,	<u> </u>	ַלֻ י	ή.	į.	Ė	Ä	4	4	4	4	4	4	뉴	뉴	4	$\dot{\mathbf{H}}$	$\dot{\boldsymbol{H}}$	Ċ	Ċ	'n	ζ.	4	4	4	7.	۲.	Ι.	١.	322-18A	

$\rho\left(\frac{\rho_{\mathbf{a}}}{\rho_{\mathbf{He}}}\right)$ $\Omega-\mathrm{cm}\left(\times10\right)$	1	:	¦	! <b>!</b>	:	1	;	;	-	;	.005	900	600	010	.016	.012	.010	600	. 013	900	.011	•	;	.005	800.	1	-	;	-	;	:	1
$\sigma_{\rm UTS}\left(\frac{\rho_{\rm He}}{\rho_{\rm app,}}\right)$	1	:	!	56 [	10.20	2.56	2.48	1	1	!	;	4.89	1	!	;	1.6	;	;	2.2	5.83	;	;	;	;	1	;	;	;	!	!	1	1
$\sigma_{\rm cs}\left(\frac{\rho_{\rm He}}{\rho_{\rm app.}}\right)$	1	:	!	2.7	: : !	ł	;	;	;	;	;	28.5	;	ł	ł	11.75	;	!	19.6	35.9	!	;	}	!	;	;	1	1	;	!	1	1
$\mathbf{E_{s}}\left(\frac{\rho_{He}}{\rho_{app.}}\right)$	1	!	. !	1	;	1	;	;	;	;	2.61	!	1.44	1.41	.51	.73	1.35	1.63	1.13	2.01	.84	;	1		1.69	;	}	;	;	;	!	1
συτς/ρ <sub>αpp</sub> .	1	: :	37 6	37.1	136.2	40.9	60.3		!	ł	ŀ	94.4	1	1	-	27.8	1	+	43.0	111.7	1	;	!	!	!	!	!	!	1	108.0	;	<b>!</b>
$\sigma_{cs}/\rho_{app}$ .	-	!!		162.3		;	1	1	;	-	;	549.6	-	;	;	203.9	;	1	$\infty$	6.989	;	;	;	;	;	!	1	!	1	673.6	!	!
Es/papp.	10.3		. !		¦	;	!	6	•	56.7	<b>;</b>	1		e,	œ	2	1.	3	21.9	œ	5	5	ω.	ė	•	;	'n	·	i	7		œ.
Sample #	322-19B	2-77	22-22	22-2	22-23	22-2	22-24	22-2	22-2	22-2	22-2	22-2	22-2	22-3	22-3	22-3	22-3	22 - 3	22-3	22 - 3	22 - 3	22-3	22-3	22-4	224	22-4	$\sim$ 1	22-42B	22-4	22-42B	2-4	2-42B

THE PARTY NAMED IN

1

p b	0-cm(×10)			110.	1 0	000	1	1 1				<b>:</b>			<b>!</b>	1 1			1						!
$ \sigma_{\rm UTS} \left( \frac{\rho_{\rm He}}{\rho} \right) $	psi (×10 <sup>-3</sup> )	;				1 ;		1	1					1 1	l	1	1	i	1		1 1				1
ocs PHe	psi (×10-3)	¦	1					ŀ	ł	1	1	1	ł		ł	ł	ł	;	;	6	1	ł	1		   
	psi (×10-6)								2.6	3.29	3.65	2.7	3.14		2 6	0	0	000	6	2.1	2.7	3.5		-	•
o. Turs/papp.	$in(\times 10^{-3})$	;	1	1	<b>¦</b>	}	4	;	1	H	}	ł	ł	ł	+	1	1	;	;	1	ł	1	;	ł	
ocs/papp.	$in(\times10^{-3})$								;	.	1	1	1	1	1	1	1	1	;	1	1	1	1	ł	
Es/papp.	in(×10-6)	5	20.8	9	27.8	19.7	14.1	•	45.6	•								5	15.3	39.4	6	61.0	6	20.1	,
	Sample #	322-42B6	322-45	322-48	322-49A	322-50	322-61	322-62	322-63	322-63A	322-64	322-64A	322-64B	322-67	322-67B	322-68	322-69	322-69A	322-70	323-2	323-2A	323-3A	323-4	323-4A	

TABLE 10 Comparison of Ultimate Strength
Determined by Direct Tension
vs. Disc Rupture

Sample #	Direct Tension	Disc Rupture
317-38	3780 psi	5100 psi
318-59	4600 psi	4153 psi
321-16B	5306 psi	4705 psi
321-36A	4081 psi*	2462 psi
321-50B	3780 psi	5228 psi
322-23B	3163 psi	4050 psi
322-24A	1367 psi	1279 psi
322-24B	1122 psi	1279 psi

\*Broke in Grip

TABLE 11 Comparison of Sonic Modulus and Mechanical Modulus

Sample #	HTT Temp.°C	Mechanical Modulus	Sonic Modulus
317-8	2000	$1.27 \times 10^{6}$	$1.82 \times 10^{6}$
317-26	2000	$0.38 \times 10^{6}$	$0.31 \times 10^{6}$
317-42	2000	$1.4 \times 10^{6}$	$1.5 \times 10^{6}$
317-46	2000	$1.59 \times 10^{6}$	$1.27 \times 10^{6}$
318-17	2000	$1.16 \times 10^{6}$	$1.18 \times 10^{6}$
318-52	2000	$2.0 \times 10^{6}$	$1.69 \times 10^{6}$
321-12	2000	$1.01 \times 10^{6}$	$1.22 \times 10^{6}$

TABLE 12

## Sonic Modulus vs. Pyrolysis Temperature psi(×10<sup>-6</sup>)

Sample #	700°C	800°C	900°C	1000°C	1577°C	1800°C	2000°C
318-59 #1L	1.02			2.16	1.95	1.77	1.46
318-59 #1L	1.06			2.07	2.10	1.82	1.76
318-60L	.93				1.82	1.77	1.70
321-11L	.75			2.37	1.68	1.72	1.58
321-11L 321-11CL	.73			1.75	1.75	1.72	1.66
321-11CL 321-12L	.66			1.49	2.03	1.23	1.18
321-12L 321-13 #1L	.80			1.84	1.77	1.73	1.75
	.82			1.84		1.79	1.72
321-13 #2L 321-13 #3L	.88			1.91		1.75	1.72
"	.81		1.69	1.92	1.91	1.84	1.69
321-15 #1 321-15 #2		1.53		1.96	2.01	1.86	1.71
321-15 #2 321-15L		1.69		2.06	1.875	1.86	1.79
		1.60		1.88	1.87	1.70	1.55
321-16A 321-16AL		1.57		1.97	1.87	1.67	1.58
321-16AL 321-16B		1.68		1.87	1.68	1.55	1.48
321-16B 321-17B	.80		1.80	2.12	1.96	1.91	1.81
321-17B 321-18A	.00	1.63		2.2	2.14	2.09	1.98
321-18B #1		1.73		1.48	1.86	1.81	1.73
321-18B #2		1.52		1.97	2.09	1.74	1.73
321-18BL		1.56		1.60	1.90	1.83	1.75
321-18BL 321-19A		1.53		2.03	1.96	1.89	1.83
321-19A 321-19B #1	.75		3.27	1.58	1.78	1.49	1.49
321-19B #1		1.32		1.69	1.61	1.61	1.55
321-19B #2 321-19BL		1.40		2.15	1.63	1.64	1.55
251-1300							

TABLE 13 Resistivity vs. Pyrolysis Temperature  $\Omega$ -cm(×10)

Sample #	800°C	900°C	1000°C	1577°C	1800°C	2000°C
318-59 #1L			.499	.102	.083	.110
318-59 #2L			.522	.099	.081	.060
318-60L			.523	.0108	.016	.170
321-11L			.643	.116	.120	.120
321-11CL			.471	.098	.090	.170
321-12L			.542	.148	.120	.06
321-13 #1L			.506	.120	.170	.100
321-13 #2L			.521	.106	.160	.110
321-13 #3L			.490	.111	.110	.110
321-15 #1		.143	.014	.0302	.015	.015
321-15 #2	.448		.029	.0319	.015	.015
321-15L	.324		.110	.115	.170	.110
321-16A	.279		.014	.029	.015	.015
321-16AL	.366		.170	.089	.120	.100
321-16B	.445		.030	.0328	.017	.017
321-17B		.083	.080	.0357	.018	.018
321-18A	.238		.047	.0333	.017	.017
321-18B #1	.611		.059	.0334	.016	.016
321-18B #2	.302		.03	.0325	.016	.016
321-18BL	.374		.160	.114	.110	.110
321-19A	.317		.031	.0332	.017	.017
321-19B #1		.171	.017	.0368	.018	.018
321-19B #2	.302		.03	.0325	.016	.016
321-19BL	.355		.150	.081	.110	.110