Introduction

As Entegris has continued to grow its market share of manufactured graphite material and to expand their usage into increasingly complex areas, the need to be more technically versed in both Entegris’ POCO® graphite and other manufactured graphite properties and how they are tested has also grown. It is with this thought in mind that this primer on properties and characteristics of graphite was developed.

POCO, now owned by Entegris, has been a supplier to the semiconductor industry for more than 35 years. Products include APCVD wafer carriers, E-Beam crucibles, heaters (small and large), ion implanter parts, LTO injector tubes, MOCVD susceptors, PECVD wafer trays and disk boats, plasma etch electrodes, quartz replacement parts, sealing plates, bonding fixtures and sputtering targets. Entegris has a wide range of materials, as well as post-processing and machining capabilities to meet the demanding requirements of semiconductor processing.

With an accumulation of experience and data in testing POCO graphites as well as other manufactured graphites, it is now possible to describe the properties and testing techniques and their interrelationship. The very discussion of these issues will bring forth many of the reasons why POCO graphites with their unique properties are superior to other manufactured graphites.

The purpose of this primer is to introduce the reader to graphite properties and to describe testing techniques, which enable true comparisons between a multitude of manufactured graphites. POCO graphites come in many grades, each designed for a specific range of applications. In the semiconductor industry, these are represented by grades such as SFG, CZR, TRA, DFP, HPD, PLS, SCF and ZEE. These are the materials upon which POCO facility has built its reputation as a manufacturer of the best graphites in the world.

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Structure

Definition: Carbon, the Element

Carbon is the sixth element on the periodic table and can be found in abundance in the sun, stars, comets and atmospheres of most planets.

Carbon is a Group 14 element (on older periodic tables, Group IV A) along with silicon, germanium, tin and lead. Carbon is distributed very widely in nature (Figure 1-1).

In 1961, the International Union of Pure and Applied Chemistry (IUPAC) adopted the isotope ¹²C as the basis for atomic weights. Carbon-14, ¹⁴C, an isotope with a half-life of 5730 years, is used to date such materials as wood, archeological specimens, etc. Carbon-13, ¹³C, is particularly useful for isotopic labeling studies since it is not radioactive, but has a spin I = ½ nucleus and therefore a good NMR nucleus.

Carbon has four electrons in its valence shell (outer shell). The electron configuration in carbon is 1s² 2s² 2p². Since this energy shell can hold eight electrons, each carbon atom can share electrons with up to four different atoms. This electronic configuration gives carbon its unique set of properties (Table 1-1). Carbon can combine with other elements as well as with itself. This allows carbon to form many different compounds of varying size and shape.

Carbon is present as carbon dioxide in the atmosphere and dissolved in all natural waters. It is a component of rocks as carbonates of calcium (limestone), magnesium and iron. Coal, petroleum and natural gas are chiefly hydrocarbons. Carbon is unique among the elements in the vast number of varieties of compounds it can form. Organic chemistry is the study of carbon and its compounds.

Forms of Carbon

Carbon is found free in nature in three allotropic forms: amorphous carbon, graphite and diamond. More recently, a fourth form of carbon, buckminsterfullerene, C₆₀, has been discovered. This new form of carbon is the subject of great interest in research laboratories today. Within the past few years, this research has centered on graphene and its derivatives, which have the potential to bring about a fundamental change in the semiconductor/electronic industry.

Carbon alone forms the familiar substances graphite and diamond. Both are made only of carbon atoms. Graphite is very soft and slippery, while diamond is one of the hardest substances known to man. Carbon, as microscopic diamonds, is found in some meteorites. Natural diamonds are found in ancient volcanic “pipes” such as found in South Africa. If both graphite and diamond are made only of carbon atoms, what gives them different properties? The answer lies in the way the carbon atoms form bonds with each other.

### Table 1-1. Properties of the Element Carbon

<table>
<thead>
<tr>
<th>Property</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Carbon</td>
</tr>
<tr>
<td>Symbol</td>
<td>C</td>
</tr>
<tr>
<td>Atomic number</td>
<td>6</td>
</tr>
<tr>
<td>Atomic mass</td>
<td>12.0107 amu</td>
</tr>
<tr>
<td>Melting point</td>
<td>3500.0°C</td>
</tr>
<tr>
<td></td>
<td>3773.15 K</td>
</tr>
<tr>
<td></td>
<td>6332.0°F</td>
</tr>
<tr>
<td>Boiling point</td>
<td>4827.0°C</td>
</tr>
<tr>
<td></td>
<td>5100.15 K</td>
</tr>
<tr>
<td></td>
<td>8720.6°F</td>
</tr>
<tr>
<td>Number of protons/electrons</td>
<td>6</td>
</tr>
<tr>
<td>Number of neutrons</td>
<td>6, 7, 8</td>
</tr>
<tr>
<td>Classification</td>
<td>Non-metal</td>
</tr>
<tr>
<td>Crystal structure</td>
<td>Hexagonal</td>
</tr>
<tr>
<td></td>
<td>Cubic</td>
</tr>
<tr>
<td>Density @ 293 K</td>
<td>Graphite – 2.26 g/cm³</td>
</tr>
<tr>
<td></td>
<td>Diamond – 3.53 g/cm³</td>
</tr>
<tr>
<td>Color</td>
<td>Black, gray</td>
</tr>
</tbody>
</table>

The history of manufactured graphite began at the end of the 19th century with a surge in carbon manufacturing technologies. The use of the electrical resistance furnace to manufacture synthetic graphite led to the development of manufactured forms of carbon in the early part of the 20th century and more recently, to a wide variety of high-performance materials such as carbon fibers and nanotubes (Figure 1-2).
The forces within and between crystallites determine the extreme difference in properties between these two forms. In diamond, the crystal structure is face-centered cubic (Figure 1-3). The interatomic distance is 1.54 Å with each atom covalently bonded to four other carbons in the form of a tetrahedron. This interatomic distance is close to that found in aliphatic hydrocarbons, which is in distinction to the smaller 1.42 Å carbon-carbon distance found in graphite and aromatic hydrocarbons (1.39 Å in benzene). This three-dimensional isotropic structure accounts for the extreme hardness of diamond.
Thermodynamically, graphite at atmospheric pressure is the more stable form of carbon. Diamond is transformed to graphite above 1500°C (Figure 1-4).

The structure of graphite consists of a succession of layers parallel to the basal plane of hexagonally linked carbon atoms. The ideal graphite structure is shown in Figure 1-5.

In this stable hexagonal lattice, the interatomic distance within a layer plane, $a$, is 1.42 Å and the interlayer distance, $d$, between planes is 3.35 Å. Crystal density is 2.266 g/cm$^3$ as compared with 3.53 g/cm$^3$ for diamond. In the graphite structure ($sp^2$ hybridization), only three of the four valence electrons of carbon form regular covalent bonds ($\sigma$-bonds) with adjacent carbon atoms. The fourth or $\pi$ electron resonates between the valence bond structures. Strong chemical bonding forces exist within the layer planes, yet the bonding energy between planes is only about two percent of that within the planes (150–170 kcal/[gram atom] vs. 1.3–4 kcal/[gram atom]). These weaker bonds between the planes are most often explained to be the result of van der Waals forces.

However, Spain identifies the $\pi$ orbital, which has a $p_z$ configuration, and not van der Waals forces as the correct source of bonding between the adjacent layers. In general, the $\pi$ bands overlap by ~40 meV to form the three-dimensional graphite network where the layer planes are stacked in the ABAB sequence illustrated in Figure 1-5. Spain concludes in his discussions on electronic structure and transport properties of graphite that the overlap of $\pi$ orbitals on adjacent atoms in a given plane also provides the electron bond network responsible for the high mobility (electronic) of graphite. This would appear more correct, since van der Waals forces are the result of dipole moments, which would not account for the high mobility.

Consequently, weak forces between layer planes account for (a) the tendency of graphite to fracture along planes, (b) the formation of interstitial compounds and (c) the lubricating, compressive and many other properties of graphite.

As previously mentioned for the hexagonal graphite structure, the stacking order of planes is ABAB, so that the atoms in alternate planes are congruent (Figure 1-5). Studies have shown that natural graphite contains 17 to 22 percent of a rhombohedral structure with a stacking sequence of ABCABC. In “artificial” or “synthetic” graphite, in the as-formed state, only a few percent at best could be found. However, deformation processes such as grinding substantially increase the percent of rhombohedral structure found in the otherwise hexagonal structure.

Amorphous carbon is also referred to as nongraphitic carbon. When examined by X-ray diffraction, these materials show only diffuse maxima at the normal scattering angles. This has been attributed to a random translation and rotation of the layers within the layer planes. This disorder has been called turbostratic. Some of these nongraphitic carbons will become graphitic, upon heating to 1700–3000°C. Some will remain nongraphitic above 3000°C.

Thus far, the discussion has centered on the crystal structure of graphites. On a more macroscopic level, the structure as routinely examined on a light microscope at magnifications of 100, 200 and 500 times reveals the porosity, particle or grain size and the general microstructure as it is commonly referred to. Photomicrographs of POCO DFP-1 graphite compared to a conventional graphite demonstrate some significant differences when viewed at 100× magnification (Figure 1-6) and at 500× magnification (Figure 1-7). It can be seen from these photos that vast differences do exist in graphite microstructure. These differences are directly related to raw material and processing parameters.

As seen in the photos, the dark or black regions represent the porosity while the lighter regions represent the graphite matrix. It is this matrix, composed of smaller particles bound together either chemically or mechanically, which is comprised of the crystals stacked layer upon layer. This is more easily seen in scanning electron micrographs (SEM).
The fourth form of carbon, buckminsterfullerene, formula \( C_{60} \), whose framework is reminiscent of the seams in an Association Football ("soccer") ball (Figure 1-8), is the subject of considerable interest at present and was only discovered a few years ago in work involving Harry Kroto, a Sheffield University graduate.

**Test Methods**

The structure of graphite has been determined through such methods as X-ray diffraction, transmission electron microscopy, neutron diffraction and convergent beam electron diffraction. These methods are highly sophisticated and generally require very expensive equipment with a highly skilled operator. This is normally beyond the scope of typical industrial laboratories. Since this type of testing or analysis is more research-oriented, no standard methods will be presented.

However, several books have been published on the structure of graphite and the reader is encouraged to review the bibliography in the appendix.

**Entegris’ POCO Graphites vs. Conventional Graphites**

With regard to crystalline structure, POCO graphite has a typical hexagonal structure. The layer spacings may vary, as they are a function of raw material and process conditions which vary from manufacturer to manufacturer. It is reasonable to assume that a certain degree of rhombohedral structure exists also in machined artifacts due to the machining-induced deformation mentioned previously. No testing has been done to confirm this.
POCO graphites are also highly isotropic with respect to their structure and properties. The isotropy factor is between 0.97 and 1.03 with 1.00 being perfect. A factor of 1.00 means the properties are identical no matter which direction they are measured in. Many conventional graphites are anisotropic. This means the properties vary depending on which direction you test them in. The high degree of isotropy makes POCO graphites useful in many applications where an anisotropic material would fail. It also allows for maximum utilization of material, as machining orientation is of no importance.

Temperature Effects
There are two general types of carbon, those considered to be “graphitizing” carbons and those that are “nongraphitizing.” The most significant difference is found in the apparent layer size and apparent stack height. For equal layer sizes, the apparent stack height, i.e., average number of layers per stack, is less for nongraphitizing carbons than graphitizing carbons. The layer stacking is more perfect in graphitizing carbons than nongraphitizing. These apparent sizes and heights are important in the first stages of carbonization.

Density Effects
Isotropy is independent of density. A high- or low-density material can be isotropic or anisotropic. The general crystal “structure” is also independent in that the greatest effects on density are due to process parameters. The same crystal “structure” can exist independent of the density of the bulk piece.

Apparent Density
Definition
The density of a substance is the amount of material, or mass, per unit volume. Density is ordinarily expressed in grams per cubic centimeter or pounds per cubic foot (1 g/cm$^3$ = 62.4 lb/ft$^3$). To determine the density of a specimen, one would first calculate its volume from the physical dimensions (for a rectangular solid, the volume is equal to the product of the length, width and thickness). Next, the mass would be determined by weighing the specimen. The density is determined by dividing the mass by the calculated volume.

If the specimen were completely homogeneous, with no flaws or voids, this method of determining density would yield the theoretical value. Graphite materials are, however, porous; hence, the term apparent density.

In general, the differences in density of POCO graphites reflect what some of the other physical properties will be. The higher-density graphite will, generally, be stronger with a higher hardness value plus improvement in many other properties and characteristics.

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3 See footnote 1.
The mathematical expression for the determination of density is:

\[ D = \frac{W}{V} \]

Where: \( D \) = Density in g/cm\(^3\)  
\( W \) = Weight of specimen in grams  
\( V \) = Volume of specimen in cm\(^3\)

Or, if the weight is expressed in pounds and the volume is expressed in cubic feet, then the density would be in units of pounds per cubic foot.

**Sample Calculation:**

A graphite specimen has a length \( (l) \) of 1.000 inches, a width \( (w) \) and thickness \( (t) \) of 0.500 inches, and a weight of 7.500 grams. Calculate the apparent density \( (D) \).

\[ V = l \cdot w \cdot t = 1.000 \text{ in} \cdot 0.5000 \text{ in} \cdot 0.5000 \text{ in} \]

\[ V = 0.25 \text{ in}^3 \]

To convert to cm\(^3\), multiply by 16.387 (1 inch = 2.54 cm, Appendix A).

\[ D = \frac{W}{V} = \frac{7.500 \text{ g}}{4.097 \text{ cm}^3} = 1.831 \text{ g/cm}^3 \]

**Test Methods**

The standard method commonly used to determine the apparent density of graphite is described in ASTM® Standard C559 and Research & Development – Analytical Services Laboratory Instruction (TDI) 4.1.1.1 (Appendix B).

For premium graphites, such as the POCO grades, an alternate method of determining apparent density is the “water method.” This is a method that can be used on objects of irregular shape where the volume would be difficult to calculate. Even though the graphite is porous, the intrusion of water into the porosity is slow and the accuracy with this method is ±1 percent if the submerged weight is taken quickly.

The general steps in the “water method” are as follows:

1. Support the piece of graphite by a thin wire/thread and weigh the piece of graphite in air.
2. Submerge the piece in a container of water in such a way that the submerged weight can be determined.
3. Calculate the density by the following formula:

\[ D = \frac{W}{W_1 - W_2} \times D_1 \]

Where: \( D \) = Density in g/cm\(^3\)  
\( W_1 \) = Weight in air (in grams)  
\( W_2 \) = Weight in water (in grams)  
\( D_1 \) = Density of water

**POCO Graphites vs. Conventional Graphites**

POCO graphites are manufactured in a variety of grades covering the density range from 1.30 g/cm\(^3\) to 1.88 g/cm\(^3\). The density is a particularly important characteristic of graphite because, in addition to its inherent significance, it has a direct influence on other properties. Generally, the physical and mechanical properties improve as the density is increased; details will be presented in later sections. Commercial, polycrystalline graphites seldom exceed 80 percent of the theoretical density figure (2.26 g/cm\(^3\)) due to voids and pores. Single crystal and pyrolytic graphites, because of their highly ordered structures and absence of pores, have densities closely approaching the theoretical value. In comparison to most other materials of construction, graphite has a low density (Figure 2-1). This is a decided advantage for the large majority of applications.
Temperature Effect
The apparent density will be influenced by temperature during the graphitization process. Generally, the higher the graphitization temperature, the higher the density will become. There are other factors which may contribute to this also, but there is an appreciable density increase as you go from 2000°C to 3000°C.

Porosity

Definition
The standard definition for porosity, as found in ASTM Standard C709 which has definitions of terms relating to manufactured carbon and graphite, is “the percentage of the total volume of a material occupied by both open and closed pores.” When one calculates the apparent density of a material, the pore volume is included in the calculation. This results in typical maximum densities for nonimpregnated manufactured graphites of 1.90 g/cm³. The theoretical density of graphite is 2.26 g/cm³. This means that in the very best case, about 16 percent of the volume of a bulk piece of graphite is open or closed pores. This porosity plays an important role in many ways, as will be discussed later.

The characteristics of the porosity of POCO fine-grained graphites have been studied extensively.

Test Methods
There is no recognized ASTM standard for measuring the porosity of manufactured graphites at this time. A number of techniques may be employed for the purpose and are widely in use today. It is important to state the method by which porosity data is determined because each method imparts its own bias.

One of the more widely used methods is mercury porosimetry. Two other methods in use are gas absorption by the BET technique and direct image analysis of the microstructure. The latter is gaining increased acceptance as a means of measuring more accurately the real pore structure. The advent of computer and video equipment has pushed this technique to the forefront of the porosimetry field. There are, nonetheless, limitations to this method also.

The mercury porosimetry technique is the method used for the data reported in POCO graphite literature. It involves basically pushing, under increasing pressure, mercury into the pores and as a function of pressure and volume filled, the pore size and pore volume can be determined. There are certain disadvantages of this method, such as:

1. The pores are not usually circular in cross-section and so the results can only be comparative.
2. The presence of “ink-bottle” pores or some other shape with constricted “necks” opening into large void volumes. The pore radius calculated by the Washburn equation is not truly indicative of the true pore radius and capillaries are classified at too small a radius.
3. The effect of compressibility of mercury with increasing pressure. This should be corrected for by carrying out a blank run.
4. The compressibility of the material under test. This is a problem of particular importance for materials which have pores that are not connected to the surface, e.g., cork. Additionally, pore walls may break under the pressures used if the material under test is relatively weak. This could cause a bias in the data.
5. The assumption of a constant value for the surface tension of mercury.
6. The assumption of a constant value for the angle of contact of mercury.
Entegris has carried out extensive analysis via mercury porosimetry to determine fundamental porosity parameters such as pore size and distribution, pore volume and surface area. Using these pore characteristics, two basic linear correlations were observed: closed porosity versus graphite apparent density (Figure 3-1) and average pore diameter versus graphite apparent density (Figure 3-2).

The mercury porosimetry measurements were made on a Micromeritics Mercury Porosimeter, Model 915-2. A surface tension constant of 480 dynes/cm and a wetting contact angle of 140 degrees were assumed and used in the Washburn equation where \( P \) is pressure in psi, \( r \) is the pore radius in cm, \( \gamma \) is the surface tension and \( \theta \) is the contact angle. A pressure and penetration volume reading were derived from the mercury porosimetry apparatus.

Penetration volume and pressure data were used to generate a printout of pressure, volume, pore size and percent porosity relationships. Graphical plots were generated to summarize pore size distribution information; percent porosity was plotted as a function of pore diameter.

\[
Pr = -2\gamma \cos \theta
\]

The surface area was also determined for each sample as described in the relationship where \( S \) is the surface area in square meters per gram. In addition, the total closed porosity was determined from the theoretical porosity and the observed (open) porosity. Percent open and closed porosity were expressed as theoretical porosity.

\[
S = 0.0225 \left( \frac{V_{max}}{PdV} \right)
\]

The above porosity parameters are displayed in graphical form as closed porosity versus apparent density and average pore diameter versus apparent density and average pore diameter versus apparent density in Figures 3-1 and 3-2, respectively. Linear regression analysis was performed to determine the best fit equation (dotted lines represent limits at the 95 percent confidence level).

The results of these porosimetry studies indicate a linear relationship between average pore diameter and graphite apparent density. That is, as the graphite apparent density increases the average size of the pores increases. Previous in-house photomicrograph studies confirm this observation. Closed porosity was also found to increase with graphite apparent density. As can be predicted on the basis of the above surface area equation and the above observations, surface area varies inversely with graphite apparent density.

A greater amount of surface area is observed in the lower-density graphite than in the higher-density product. At first glance, these observations are surprising and may even seem contradictory. Why should closed porosity increase when a concomitant increase in the pore diameter is also observed?

Although cause-and-effect relationships are difficult to establish, graphite porosity, pore size and surface area are all physically related to density. Their relationship to density, whether direct or inverse, has implications on the structural properties of POCO graphite. The following physical model is advanced to rationalize the above relationships.

One notes that as graphite density increases, closed porosity and average pore size also increase while graphite surface area decreases. As the graphite structure increases in density, the smaller pores can be imagined to become more and more occluded until they are isolated from the rest of the pore system. As this process occurs, the smaller-diameter pores are systematically eliminated until only the larger, less complex pores remain. This also creates a larger amount of closed porosity. Thus, not only does the average pore diameter increase as a result of the elimination of small open pores, but pore surface area is reduced, since only pores with less branched structures remain.

Mercury porosimetry data on fine-grained graphites reveals a definite relationship between closed porosity and apparent density with the closed porosity increasing as the apparent density increases. The pore size also increases as apparent density increases.

**POCO Graphite vs. Conventional Graphite**

The pore volume will be the same for all graphite with the same apparent density, but that is where the similarity ends. The pore diameter of POCO graphite ranges from 0.2 microns nominal to 0.8 microns nominal for our 1 and 5 micron grades and densities. The open porosity ranges from 75 percent open to 95 percent open and the pores are generally spherical in shape. The smaller pore size materials have a very large surface area associated with them. Conventional graphites will, at best, have typical pore sizes only down to several microns in size. The distribution of pore sizes is very narrow for POCO graphites, while they are generally broader for conventional graphites (Figure 3-3).

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The fineness of the porosity allows Entegris materials to be modified to create truly impermeable graphite. With various post-processing techniques, Entegris can seal, fill or close the porosity, depending on the end application. The high degree of open porosity also allows Entegris to purify the material to less than 5 parts-per-million (ppm) total impurities, by ash analysis. Figure 3-4 shows helium flow data on various grades of POCO graphite. It clearly shows a wide variety of capabilities for POCO graphites. The helium flow test for checking the permeability, like the mercury porosimetry test, has its limits or bias and should be clearly identified when using data generated by it. Permeability is simply the rate of flow of a medium such as a gas or liquid through a material while under a pressure gradient. The pores of POCO graphite are not only uniformly distributed, but are well interconnected so flow can take place through them. There are applications (such as filters) where this is important.

Another feature of the small pore size is that some liquids will not generally penetrate the pores readily. For instance, it has been determined that after soaking in water for seven days, a sample of DFP-1 picked up less than one percent by weight of the water (Figure 3-5). This could be an advantage in some applications. However, in other applications, such as wanting to infiltrate a liquid such as molten copper, very high pressures are required at high temperatures to accomplish it. This is a decided disadvantage.
Temperature Effects
As temperature increases, pores will expand along with the rest of the matrix to the point of opening what were previously closed pores. As graphitization temperature increases, the pores will generally be found to be slightly smaller in size. It is uncertain what would happen to an impermeable graphite at room temperature if it were raised to very high temperatures. Theoretically, if it is surface sealed, it may retain its impermeability. If it is densified, additional pores may open and thus render it permeable once more.

Density Effects
Mercury porosimetry data on fine-grained graphites reveals a definite relationship between closed porosity and apparent density with the closed porosity increasing as the apparent density increases. The pore size also increases as apparent density increases. The relationship is linear, with closed porosity increasing as density increases and pore size also increasing as density increases. If post-processing densification techniques were employed to raise the density, the pore size and pore volume would decrease.

Hardness

Definition
Although the term “hardness” has comparative significance in an engineering sense, it is generally not considered to be a fundamental property of matter. The index of hardness is generally a manifestation of several related properties. In graphite, for instance, the particle size and porosity, and apparent density have an influence on the hardness value. The hardness value can be changed by the graphitization temperature as well, which generally relates back to a strength characteristic such as shear strength at the crystallographic level. Different hardness testers are influenced by different properties and as a consequence cannot be correlated very well. Comparatively soft materials may be hard in the sense that they can resist abrasion stresses, whereas harder materials in the sense of indentation hardness may fail completely under the same circumstances. It should be obvious at this point that the term “hardness” is relative and hardness data must be interpreted in relation to the type of hardness tests used.

An appropriate definition of hardness then would be the resistance of a material to permanent deformation of its surface. The deformation may be from scratching, mechanical wear, indentation or in a broader sense, cutting. Cutting would clearly include machinability as an index of hardness. This is much less precise than conventional hardness testers, but can be an indicator of relative usefulness.

Test Methods
There are two standard test methods for hardness of graphite: ASTM Standard C886, the Scleroscope Hardness method, and C748, the Rockwell Hardness method. The Shore Scleroscope measures hardness in terms of the elasticity of the material. A diamond-tipped hammer in a graduated glass tube is allowed to fall from a known height onto the specimen to be tested, and the hardness number depends on the height to which the hammer rebounds; the harder the material, the higher the rebound (see also Mohs hardness). Brinell hardness is determined by forcing a hardened steel or carbide ball of known diameter under a known load into a surface and measuring the diameter of the indentation with a microscope. The Brinell hardness number is obtained by dividing the load, in kilograms, by the spherical area of the indentation in square millimeters; this area is a function of the ball diameter and the depth of the indentation.

The Rockwell hardness tester utilizes either a steel ball or a conical diamond known as a bradle, and indicates hardness by determining the depth of penetration of the indenter under a known load. This depth is relative to the position under a minor initial load; the corresponding hardness number is indicated on a dial. For hardened steel, Rockwell testers with bradle indenters are particularly suitable; they are widely used in metalworking plants.

The Vickers hardness tester uses a square-based diamond pyramid indenter, and the hardness number is equal to the load divided by the product of the lengths of the diagonals of the square impression. Vickers hardness is the most accurate for very hard materials and can be used on thin sheets.

The Scleroscope hardness test method is based on the rebound height of a diamond-tipped hammer off the sample’s surface after it falls a fixed distance. The scale is unitless, with the degree of hardness directly related to the height of the rebound. The higher the rebound, the “harder” the material. There are many factors that affect the reproducibility and accuracy of the data. Studies at Entegris indicate that sample size, i.e., mass, has a significant bearing on the results as well. The results of a large billet will generally be higher than a small test sample.

The Rockwell hardness test method is based on the differential—the depth of indentation—produced on
Compressive Strength

Definition

The easiest strength characteristic of a material to understand and measure is the compressive strength. When a material is positioned between two flat, parallel platens and a continually increasing compressive force is applied, either of two things can occur:

1. If the material is ductile, such as copper or iron, atomic or molecular bonds can be re-formed easily; therefore, when crystalline planes begin to slip across each other, the atoms will readily re-form bonds with other atoms. Consequently, it is quite possible to flatten a very ductile material, such as gold, into a very thin sheet (if high enough compressive load is applied).

2. If the material is brittle, such as graphite or many ceramic materials, atomic or molecular bonds cannot be re-formed easily; therefore, when crystalline planes begin to slip, catastrophic failure occurs and the material fractures.
The compressive strength of a brittle material is expressed as the maximum force per unit area that can be withstood before failure occurs. It is usually expressed in pounds per square inch (lb/in² or psi) or in megapascal (MPa) in metric units. The mathematical expression for the determination of compressive strength is:

\[ \text{C.S.} = \frac{L}{A} \]

Where:  
- \( \text{C.S.} \) = Compressive strength  
- \( L \) = Load required to cause failure  
- \( A \) = Cross-sectional area of specimen

**Sample Calculation:**

A graphite specimen with a cross-sectional area of 0.25 square inches (in²) fails when a load of 3500 pounds is applied; calculate the compressive strength.

\[
\text{C.S.} = \frac{L}{A} = \frac{3500 \text{ lbs}}{0.25 \text{ in}^2} = 14,000 \text{ psi}
\]

To convert to MPa, multiply by 0.0068948.

Compressive strength = 97 MPa

**NOTE:** 1 psi = 0.0068948 MPa

Other units and the appropriate conversion factors are given in Appendix A.

**Test Method**

The standard method of measuring the compressive strength of a graphite specimen is described in ASTM Standard C695. Entegris’ method differs in that the specimen used is rectangular rather than a right cylinder. Cushion pads are not used either (TDI 4.1.1.14 in Appendix B and Figure 5-1).

**POCO Graphites vs. Conventional Graphites**

POCO graphites have high compressive strengths compared to conventional materials. The compressive strengths of Entegris materials range from 69 MPa (10,000 psi) to over 207 MPa (30,000 psi) depending on the grade selected. These values are two to three times higher than most other graphites. The final heat-treating temperature used in the manufacture of a carbon/graphite material has a marked effect on its compressive strength; the lower the final temperature, the higher the compressive strength.

**Temperature Effects**

As the temperature of a piece of graphite is increased, its compressive strength increases, up to about 2500°C. This is particularly important in such applications as hot pressing dies, where the material is subjected to both high temperature and high stress levels. Depending on the grade and type of graphite, the increase in strength will be from 15 to 50 percent higher at 2500°C than it is at 25°C.

The graphitization temperature also has a marked effect on the compressive strength. The lower the graphitization temperature, the less graphitic the material is and the higher the compressive strength will be. This is readily seen when comparing carbon-based material to graphite-based material for areas such as mechanical application.

**Density Effects**

As with many other properties, the compressive strength of graphites changes with apparent density; the higher density having the highest strength (Figure 5-2).
Flexural Strength

Definition

Flexural strength, tensile strength and compressive strength are the three common tests performed to measure the strength of materials. In flexural strength testing, a steadily increasing bending movement is applied to a long bar until the material eventually ruptures. If the material is ductile (like copper), it will bend prior to breaking. However, if the material is brittle (such as chalk or graphite), it will bend very little before it fails catastrophically.

Flexural strength can be defined as the maximum stress in bending that can be withstood by the outer fibers of a specimen before rupturing (Figure 6-1).

\[
F.S. = \frac{PL}{W(T)^2}
\]

Where:
- \( F.S. \) = Flexural strength in pounds per square inch (psi)
- \( P \) = Load in pounds at failure
- \( L \) = Length between outer support roller pins in inches
- \( W \) = Width of specimen in inches
- \( T \) = Thickness of specimen in inches

Sample Calculation:

Assume a graphite specimen 4.0 inches long by 0.75 inches wide by 0.50 inches thick is flexure tested on a fixture with support roller pins 3.0 inches apart. The specimen fails under a 500 pound load; calculate its flexural strength.

\[
F.S. = \frac{PL}{W(T)^2} = \frac{500 \times 3.0 \text{ lbs}}{0.75 \times (0.50)^2} = \frac{1500}{0.1875 \text{ in}^2} \text{ psi} = 8000 \text{ psi}
\]

To convert to MPa, multiply by 0.0068948.

Flexural strength = 55 MPa

Test Method

The details of the procedure commonly used for testing the flexural strength of graphite are found in ASTM Standard C651. Entegris’ method (TDI 4.1.1.13 in Appendix B) differs in that the specimen geometry has a 1:1 ratio in thickness and width rather than the 2:1 width to thickness. The fixture used is not exactly as described either, but with a surface finish of less than 32 micro inches Ra on the sample, the frictional component is minimized, and results are comparable to those obtained with the fixture recommended by ASTM. Regardless of the procedure followed, steps must be taken to avoid factors that bias the results, such as improper sample alignment, rough and/or non-parallel surfaces. It is also important to know whether the reported strength values were obtained from three-point or four-point loading.

For example, a test specimen typically has a uniform rectangular cross-section but the load may be applied in three or four point as illustrated in Figure 6-2. Note the shaded areas indicating the stress distribution. In 3-point loading, the peak stress occurs on a single line at the surface of the test bar and opposite the point of loading. The stress decreases linearly along the length of the bar, and into the thickness of the bar, until reaching zero at the bottom supports.

Unlike the 3-point bend tests, where the peak stress occurs on a single line opposite the point of loading, 4-point loading distributes the peak stress over an area that is determined by the width of the sample and the span of the top loading supports, respectively. Observe how the tensile stress distribution decreases linearly from the area of peak stress on the tensile face, and into the thickness of the bar, until reaching zero at the bottom span supports. The increased area and volume under peak tensile stress, or near the peak tensile stress, in 4-point loading increases the probability of encountering a larger flaw.
The probability of detecting the largest flaw in the specimen during 3-point loading is minimized since the largest flaw must be at the surface and along the line of peak stress. Consequently, the specimen fractures at either a smaller flaw or within a region of lower stress, thus yielding artificially higher strength values in comparison with 4-point loading results. The strength limit of the material, or even the local stress and flaw size that caused fracture, is not revealed in 3-point loading. It only indicates the peak stress on the tensile surface at the time of fracture for a given material.

POCO Graphites vs. Conventional Graphites

In a brittle material such as graphite, the flexural strength is particularly sensitive to flaws or defects in the material. If a flaw is present within span L, i.e., between the outer support pins of the flexural test specimen, then the load P required to break the sample will be reduced. When failure occurs, it is catastrophic. The sample breaks suddenly and, frequently, small chips and flakes of material break away at the point of failure.

POCO graphites have flexural strengths covering the range from 34 MPa (5000 psi) to over 124 MPa (18,000 psi). These values are quite high when compared to conventional graphites, which may range from less than 7 MPa (1000 psi) to around 41 MPa (6000 psi). The fine particle size and homogeneous structure of POCO graphites contribute to their high flexural strengths. Another important characteristic of this property in POCO graphites is that the flexural strength is the same for samples cut from any direction or orientation of the parent block of material. This characteristic is called isotropy; POCO graphites are said to be isotropic, whereas most other graphites are anisotropic. In conventional graphites that are molded or extruded, the ratio of flexural strengths between the “against grain” and “with grain” directions may range from 0.3 to 0.5.

Temperature Effects

One of the unusual properties of graphite is that it gets stronger as it gets hotter (up to about 2500°C). This is contrary to most materials, which lose strength as the temperature increases. The flexural strength of graphites will increase by 20 to 50 percent when the test temperature is increased from 25°C to 2500°C.

Density Effects

Flexural strength, like many other physical properties of graphite, increases with increasing density. For example, nominal flexural strength of 5 micron graphite (DFP-1, TRA-1 and CZR-1) ranges from 6000–17,000 psi for densities of 1.60 (g/cm$^2$) and 1.88 (g/cm$^2$), respectively (Figure 6-3). One micron or SFG exhibits the highest flexural strength indicating 18,000 psi.

![Figure 6-3. Nominal flexural strength vs. apparent density of DFP-1, TRA-1 and CZR-1 graphites](image-url)
Tensile Strength

Definition

The tensile strength of a material can be defined as its strength when a pulling force is applied along the length of a sample. If a cylindrical bar of uniform cross-section is subjected to a steadily increasing tensile (“pulling apart”) force along its axis, the material will eventually rupture and tear apart when a large enough force is applied. It is extremely difficult to correctly determine the tensile strength of a brittle material, such as graphite. The reason for this is that when brittle materials fail in tension, a crack originates at some point and is rapidly propagated across the specimen, causing catastrophic failure. Any imperfection in the material, such as voids, or surface scratches, will cause stress concentrations at that point. Since the stress at this point is higher than the overall average throughout the specimen, a crack will begin to propagate and premature fracture will occur. To alleviate this situation, carefully machined, highly polished specimens are used. Also, of no lesser importance is the alignment of the gripping apparatus on the specimen. Even small misalignments could cause premature fracture, resulting in an erroneous (low) value for the tensile strength.

Tensile strength, like compressive strength, is expressed in pounds per square inch and is calculated in the same manner as compressive strength, i.e., the applied force at failure is divided by the cross-sectional area of the sample.

Sample Calculation:

If a rod with a cross-sectional area of 0.25 in² breaks at a load of 2000 pounds, then the tensile strength is 8000 psi.

\[
\text{Tensile strength} = \frac{\text{Load}}{\text{Area}} = \frac{2000 \text{ lbs}}{0.25 \text{ in}^2} = 8000 \text{ psi}
\]

To convert to MPa, multiply by 0.0068948.

\[
\text{Tensile strength} = 55 \text{ MPa}
\]

Test Method

The details of the commonly used method for tensile strength testing are shown in ASTM Standard C565. Other, more sophisticated testing equipment has been developed for this test; one of the better ones uses hemispherical air bearings instead of chain connectors to eliminate misalignment of the sample.

POCO Graphites vs. Conventional Graphites

The tensile strength of a brittle material, such as graphite, is very sensitive to defects or imperfections in the material. The fine structure and uniformity of POCO graphites result in higher tensile strength for Entegris materials as compared to conventional graphites. Typical tensile strength for conventional graphites ranges from 14 MPa (2000 psi) to 34 MPa (5000 psi), as compared to 34 MPa (5000 psi) to 69 MPa (10,000 psi) for Entegris materials (Table 7-1).

<table>
<thead>
<tr>
<th>Material</th>
<th>Ultimate Tensile Strengths (10^3 psi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural rubber</td>
<td>&gt;4</td>
</tr>
<tr>
<td>ABS/PVC plastic</td>
<td>3–6</td>
</tr>
<tr>
<td>Graphite (polycrystalline)</td>
<td>2–10</td>
</tr>
<tr>
<td>Aluminum (wrought)</td>
<td>13–98</td>
</tr>
<tr>
<td>Silicon carbide</td>
<td>3–20</td>
</tr>
<tr>
<td>Alumina (ceramic)</td>
<td>20–30</td>
</tr>
<tr>
<td>Steel (wrought)</td>
<td>90–290</td>
</tr>
<tr>
<td>Brass</td>
<td>34–129</td>
</tr>
<tr>
<td>Copper</td>
<td>29–76</td>
</tr>
<tr>
<td>Nickel</td>
<td>50–290</td>
</tr>
<tr>
<td>Gold</td>
<td>19–32</td>
</tr>
<tr>
<td>Platinum</td>
<td>18–30</td>
</tr>
</tbody>
</table>

\[F.S. = \frac{PL}{W(T)} = 8000 \text{ psi}\]

\[ER = \frac{AR}{P(L)}\]
Temperature Effects
As the testing temperature of graphite is increased, its tensile strength increases; this is in sharp contrast to the behavior of metals, which show a decrease in strength as temperature increases. In an inert atmosphere (to avoid oxidation), the tensile strength of POCO DFP-1 graphite is almost 138 MPa (20,000 psi) at 5000°F (2760°C), as compared to 63 MPa (9150 psi) at room temperature (Figure 7-1). This dramatic change is characteristic of graphite materials, even though most grades do not show the 100 percent increase that POCO graphites exhibit.

The final heat-treat temperature of the graphite has a marked effect on its room temperature tensile strength: the lower the heat-treat temperature, the higher the strength. This is similar to the effect seen on other strength characteristics where the less graphitic materials are harder and stronger. The relationship between graphitization temperature and tensile strength is shown in Figure 7-2.

Density Effects
The tensile strength of graphite has a strong correlation with density: as the density increases, the tensile strength increases. This is typical of the other strength characteristics of graphites. Figure 7-3 shows the general relationship between density and tensile strength for the DFP-1, TRA-1 and CZR-1 graphites. The nominal values of tensile strength range from 34 MPa (5000 psi) at 1.60 g/cm³ to 55 MPa (8000 psi) at 1.84 g/cm³.

---

Figure 7-1. Ultimate tensile strength of DFP-1

Figure 7-2. Graphitization temperature effect on tensile strength

Figure 7-3. Nominal tensile strength vs. apparent density of POCO DFP-1, TRA-1 and CZR-1 graphites
Modulus of Elasticity

Definition

The elasticity of a material is related to a uniformly increasing separation between the atoms of that material. As a consequence, the elasticity is directly related to the bonding between atoms and the respective energies associated therewith. This can be readily demonstrated by showing the general relationship between modulus of elasticity (MOE) and melting points of various materials. The higher the melting point (i.e., the energy required to disrupt the atom-to-atom bonds), the higher the modulus of elasticity (Figure 8-1). The presence of a second phase of differing modulus results in most of the stress being carried by the higher modulus phase. Porosity, which is uniformly distributed and continuous, constitutes a second phase. The effect on the modulus in materials of less than 50 percent pore volume can be represented by the relationship:

\[ E_o (1 - 1.9P + 0.9P^2) \]

Where: \( P \) = Porosity (volume fraction)  
\( E_o \) = Original modulus of elasticity

Another way of expressing the modulus of elasticity is by referring to Hooke’s law, which applies to materials below their elastic limit. Basically, Hooke’s law says that the average stress is proportional to the average strain.

\[ \frac{T}{e} = E = \text{Constant} \]

Where: \( T \) = Stress 
\( e \) = Strain 
\( E \) = Modulus of elasticity, or Young’s modulus

Stress is the load per unit area and strain is the ratio of change in length to the original length, i.e.,

\[ T = \frac{P}{A} \quad e = \frac{\delta L}{L_o} \]

Where: \( P \) = Load 
\( A \) = Area  
\( \delta L \) = Change in length (\( L - L_o \)) 
\( L_o \) = Original length

Therefore:

\[ E = \frac{P}{A} \frac{\delta L}{L_o} = \frac{PL_o}{\delta L} \]

Sample Calculation:

If a tensile sample with a diameter of 0.220 inch and a gage length of 1.000 inch breaks at 500 pounds with a gage length increase of 0.008 inch, the tensile MOE would be \( 1.6 \times 10^6 \) psi.

\[ E = \frac{PL_o}{\delta L} = \frac{(500 \text{ lbs})(1)}{(0.008 \text{ in})(0.038 \text{ in}^2)} = 1.6 \times 10^6 \text{ psi} \]

The modulus of elasticity is usually expressed in millions of pounds per square inch (10^6 psi), or in N/mm² in metric units.

Test Method

The standard method of measuring modulus of elasticity for graphite when not determined from samples tested in tension is described in ASTM Standards C747 and C769. The methods are approximations derived from other properties. A more accurate, but also more difficult, means is to attach proper strain measuring gages to tensile strength samples and measure strain along with stress during a tensile test. Then, employing Hooke’s law, the modulus can be calculated.

POCO Graphites vs. Conventional Graphites

The modulus of elasticity varies for the many different grades of graphite available; therefore, a significant difference in the modulus for POCO materials compared to conventional graphite is not seen (Table 8-1). The most notable difference, however, is the isotropy of POCO which assures a modulus of the same value in any direction as compared to many conventional graphites being anisotropic.
### Table 8-1. Modulus of Elasticity of Various Graphites

<table>
<thead>
<tr>
<th>Material</th>
<th>Apparent Density (g/cm³)</th>
<th>MOE (10⁶ psi)</th>
</tr>
</thead>
<tbody>
<tr>
<td>POCO CZR</td>
<td>1.65</td>
<td>1.3</td>
</tr>
<tr>
<td>TRA</td>
<td>1.72</td>
<td>1.5</td>
</tr>
<tr>
<td>DFP</td>
<td>1.77</td>
<td>1.6</td>
</tr>
<tr>
<td>HPD</td>
<td>1.76</td>
<td>1.7</td>
</tr>
<tr>
<td>Mersen®</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2161</td>
<td>1.66</td>
<td>1.3</td>
</tr>
<tr>
<td>2020</td>
<td>1.77</td>
<td>1.3</td>
</tr>
<tr>
<td>2080</td>
<td>1.87</td>
<td>1.8</td>
</tr>
<tr>
<td>Morgan P3W</td>
<td>1.60</td>
<td>1.4</td>
</tr>
<tr>
<td>L-56</td>
<td>1.63</td>
<td>0.5</td>
</tr>
<tr>
<td>P5</td>
<td>1.72</td>
<td>2.4</td>
</tr>
<tr>
<td>P03 1</td>
<td>0.82</td>
<td>1.8</td>
</tr>
<tr>
<td>SGL Group, The Carbon Company®</td>
<td></td>
<td></td>
</tr>
<tr>
<td>H-440</td>
<td>1.75</td>
<td>1.5</td>
</tr>
<tr>
<td>HLM</td>
<td>1.75</td>
<td>1.8</td>
</tr>
<tr>
<td>H-463</td>
<td>1.75</td>
<td>1.8</td>
</tr>
<tr>
<td>H478</td>
<td>1.88</td>
<td>2.2</td>
</tr>
<tr>
<td>Toyo Tanso USA®</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SEM5</td>
<td>1.80</td>
<td>2.4</td>
</tr>
<tr>
<td>SEM3</td>
<td>1.85</td>
<td>1.4</td>
</tr>
<tr>
<td>UCAR®</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AGSR</td>
<td>1.58</td>
<td>1.6</td>
</tr>
<tr>
<td>CBN</td>
<td>1.67</td>
<td>1.8</td>
</tr>
<tr>
<td>CS</td>
<td>1.70</td>
<td>2.0 WG*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.1 AG*</td>
</tr>
</tbody>
</table>

* Shows typical anisotropy effects

#### Temperature Effects

As with the other strength properties of graphite, as temperature increases, the modulus of elasticity also increases, to a point beyond which it begins to drop rapidly. This is typically around 2500°C. The increase can be as much as 25 percent higher before the drop begins. The final graphitization temperature also has an effect, but is relatively small.

#### Density Effects

The density relationship is evident with about a 15 to 20 percent increase in modulus detected as the density increases from 1.65 g/cm³ to 1.77 g/cm³. This follows the same pattern as other strength properties for graphite.

#### Electrical Resistivity

**Definition**

The electrical resistivity is that property of a material which determines its resistance to the flow of an electrical current and is an intrinsic property. The electrical resistance of a substance is directly proportional to the length of the current path, i.e., as the current path increases, the resistance increases. It is also inversely proportional to its cross-sectional area, i.e., as the area increases, the resistance decreases.

The mathematical expression for the determination of electrical resistivity is:

\[
ER = \frac{AR}{L}
\]

Where:
- \(ER\) = Electrical resistivity at room temperature
- \(A\) = Cross-sectional area (square inches)
- \(R\) = Electrical resistance of the material (ohms)
- \(L\) = Distance between potential contacts (inches)

**Sample Calculation:**

For a graphite sample with a cross-sectional area of 0.25 in², distance between the potential contacts of 2.0 inches and an electrical resistance reading of 0.00425 Ω (ohms); calculate the electrical resistivity.

\[
ER = \frac{0.25 \text{ in}^2 \times 0.00425 \Omega}{2.0 \text{ in}} = \frac{0.10625 \Omega \cdot \text{in}}{2.0 \text{ in}} = 0.053125 \Omega \cdot \text{in}
\]

\[ER = 0.000531 \Omega \cdot \text{in}\]

\[ER = 531 \mu\Omega \cdot \text{in}\]

To convert to \(\mu\Omega\cdot\text{cm}\), multiply by 2.54

Electrical resistivity = 1349 \(\mu\Omega\cdot\text{cm}\)

**Test Method**

The standard method of measuring the electrical resistivity of a graphite sample is described in ASTM Standard C611. At Entegris only two resistance readings are taken, using a special sample holder with eight electrical contacts which takes the equivalent of four readings at once. Entegris uses test method TDI 4.1.1.2 (Appendix B).
POCO Graphite vs. Conventional Graphites

POCO graphites have electrical resistivity values that fall into the range of most conventional graphites. POCO graphites have a range from around 450 μΩ-in to 1050 μΩ-in. This may be compared to copper which has a range of 1.1 to 1.5 μΩ-in or tool steels which range from 7.1 to 7.5 μΩ-in. See Table 9-1 for other common material resistivities.

**TABLE 9–1. TYPICAL ELECTRICAL RESISTIVITY OF VARIOUS MATERIALS**

<table>
<thead>
<tr>
<th>Material**</th>
<th>Electrical Resistivity Range (μΩ-in)</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>POCO graphites</td>
<td>450–1000 Polycrystalline graphite</td>
<td></td>
</tr>
<tr>
<td>Toyo Carbon graphites</td>
<td>300–600 Polycrystalline graphite</td>
<td></td>
</tr>
<tr>
<td>Mersen graphites</td>
<td>550–1200 Polycrystalline graphite</td>
<td></td>
</tr>
<tr>
<td>Toyo Tanso USA graphites</td>
<td>400–500 Polycrystalline graphite</td>
<td></td>
</tr>
<tr>
<td>Copper</td>
<td>1.1–1.5 Pure, wrought</td>
<td></td>
</tr>
<tr>
<td>Gold</td>
<td>0.9 Pure</td>
<td></td>
</tr>
<tr>
<td>Silver</td>
<td>0.6 Pure</td>
<td></td>
</tr>
<tr>
<td>Tungsten</td>
<td>2.2 Pure</td>
<td></td>
</tr>
<tr>
<td>Carbon steels</td>
<td>7.1–7.5 Hardening grades, wrought</td>
<td></td>
</tr>
<tr>
<td>Stainless steel</td>
<td>15.7–28.3 400 series, wrought</td>
<td></td>
</tr>
<tr>
<td>Cobalt base superalloys</td>
<td>36.6–74.3 Wrought</td>
<td></td>
</tr>
<tr>
<td>Nickel base superalloys</td>
<td>3.0–52.4 Wrought and cast</td>
<td></td>
</tr>
<tr>
<td>Silicon</td>
<td>6 × 10⁶ Semiconductor</td>
<td></td>
</tr>
<tr>
<td>Silicon carbide</td>
<td>4 × 10⁶ Semiconductor</td>
<td></td>
</tr>
<tr>
<td>Silicon nitride</td>
<td>4 × 10²⁰ Insulator</td>
<td></td>
</tr>
<tr>
<td>Alumina</td>
<td>&gt;4 × 10³¹ Insulator</td>
<td></td>
</tr>
</tbody>
</table>

* Measured at room temperature in accordance with ASTM Standard C611 and TDI 4.1.1.2.

** Select samples from each manufacturer but not all-inclusive

Temperature Effects

Electrical resistivity varies with temperature (Figure 9-1). As the temperature increases from room temperature to about 700°C, the electrical resistivity decreases. From that point, however, as the temperature increases, the resistivity also increases.

Density Effects

Density is a particularly important characteristic of graphite because, in addition to its inherent significance, it has a large and direct influence on other properties. As the density of POCO graphite increases, its electrical resistivity decreases (Figure 9-2).
Thermal Expansion

Definition

The physical dimensions of a body are determined by the number and spacing of its atoms. At temperatures above 0 K, the atoms are always in constant vibration about their positions in the lattice. If energy is added to the material (by heating it), the atoms vibrate more vigorously, causing the macroscopic dimensions of the material to increase.

The coefficient of thermal expansion (CTE) is defined as the change in length of a substance per unit length for a specific change in temperature. If this thermal expansion is hindered in any way, internal stresses will occur.

When two different materials are to be joined permanently, as in the electrical connection to an incandescent lamp or a vacuum tube, it is important that they have nearly the same values of CTE if there is to be no danger of failure from cracking. The same precaution applies to coatings or cladding of one material to another.

The CTE is normally expressed as inch per inch per °C or as inch per inch per °F, with the temperature range over which the CTE is applicable being specified. This is done because, for some materials, the CTE will change with temperature.

The mathematical expression for the determination of the coefficient of thermal expansion is:

\[
CTE = \frac{\delta L}{L(T_2 - T_1)}
\]

Where: CTE = Coefficient of thermal expansion  
\( \delta L \) = Change in sample length from the lower temperature to the upper temperature  
\( L \) = Sample length at the lower temperature  
\( T_1 \) = Lower temperature  
\( T_2 \) = Upper temperature seen by sample

Sample Calculation:

A two-inch graphite sample is heated from room temperature to 1000°C.

The length uniformly increases until it is 2.0164 inches in length at 1000°C; calculate the CTE.

\[
CTE = \frac{2.0164 \text{ in} - 2.0000 \text{ in}}{2.0000 \text{ in} (1000°C - 23°C)}
\]

\[
CTE = 8.39 \times 10^{-6} \ (\text{in/in})/°C
\]

To convert to (in/in)/°F; divide by 1.8

\[
CTE = 4.65 \times 10^{-6} \ (\text{in/in})/°F
\]

Test Method

The standard method of measuring the coefficient of thermal expansion of a graphite sample is similar to ASTM Standard E228, but there is no specific method for graphite. POCO has a horizontal silica (orton) dilatometer and follows TDI 4.1.1.5 (Appendix B).

POCO Graphites vs. Conventional Graphites

POCO graphites have high coefficients of thermal expansion compared to conventional materials (Table 10-1). The coefficient of thermal expansion of POCO materials range from 7.0 \times 10^{-6} to 9.0 \times 10^{-6} (in/in)/°C, depending on the grade selected. These values are two to four times higher than most other graphites.

<table>
<thead>
<tr>
<th>Material</th>
<th>High</th>
<th>Low</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminum alloys (68–212°F)</td>
<td>24.1</td>
<td>22.3</td>
</tr>
<tr>
<td>300 Stainless steels (32–212°F)</td>
<td>18.7</td>
<td>14.9</td>
</tr>
<tr>
<td>Copper (68–572°F)</td>
<td>17.6</td>
<td>16.7</td>
</tr>
<tr>
<td>Nickel base superalloys (70–200°F)</td>
<td>17.8</td>
<td>10.6</td>
</tr>
<tr>
<td>Cobalt base superalloys (70–1800°F)</td>
<td>17.1</td>
<td>16.2</td>
</tr>
<tr>
<td>Boron nitride (70–1800°F)</td>
<td>7.5</td>
<td>–</td>
</tr>
<tr>
<td>Titanium carbide (77–1472°F)</td>
<td>7.4</td>
<td>6.7</td>
</tr>
<tr>
<td>Tungsten carbide</td>
<td>7.4</td>
<td>4.5</td>
</tr>
<tr>
<td>Silicon carbide (0–2550°F)</td>
<td>4.3</td>
<td>3.9</td>
</tr>
<tr>
<td>Silicon nitride (70–1800°F)</td>
<td>2.5</td>
<td>–</td>
</tr>
<tr>
<td>POCO graphite (70–1832°F)</td>
<td>8.8</td>
<td>7.0</td>
</tr>
</tbody>
</table>
Temperature Effects

As the temperature of a piece of graphite is increased, its coefficient of thermal expansion will become greater (Figure 10-1). There is limited data to describe its expansion characteristics above 1000°C, but the data available indicates a linear increase in expansion as high as 2500°C.

Density Effects

As with many other properties, the coefficient of thermal expansion changes with apparent density; as the density of the material increases, so will the coefficient of thermal expansion.

Since the higher-density material has less porosity, there is less distance for the crystals to move unobstructed as the temperature is increased; therefore, higher density means greater expansion (Figure 10-2).

Thermal Conductivity

Definition

The thermal conductivity of a material is a measure of its ability to conduct heat. A high thermal conductivity denotes a good heat conductor, while a low thermal conductivity indicates a good thermal insulator (Table 11-1).

<table>
<thead>
<tr>
<th>Material</th>
<th>Thermal Conductivity Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Natural rubber</td>
<td>0.082</td>
</tr>
<tr>
<td>Nickel</td>
<td>6–50</td>
</tr>
<tr>
<td>Steel (wrought)</td>
<td>8–21</td>
</tr>
<tr>
<td>Silicon carbide</td>
<td>9–25</td>
</tr>
<tr>
<td>Tungsten carbide</td>
<td>16–51</td>
</tr>
<tr>
<td>Brass</td>
<td>15–135</td>
</tr>
<tr>
<td>Iron (cast)</td>
<td>25–30</td>
</tr>
<tr>
<td>POCO graphite</td>
<td>40–70</td>
</tr>
<tr>
<td>Platinum</td>
<td>42</td>
</tr>
<tr>
<td>Conventional graphite</td>
<td>65–95</td>
</tr>
<tr>
<td>Aluminum</td>
<td>67–135</td>
</tr>
<tr>
<td>Tungsten</td>
<td>97</td>
</tr>
<tr>
<td>Copper</td>
<td>112–226</td>
</tr>
<tr>
<td>Gold</td>
<td>172</td>
</tr>
<tr>
<td>Silver</td>
<td>242</td>
</tr>
</tbody>
</table>

The mathematical expression for thermal conductivity ($K$) is:

$$K = \frac{Q\partial X}{A\partial T}$$

Where:
- $Q$ = Rate of heat flow through a slab in Btu per hour
- $A$ = Cross-sectional area of the slab in feet$^2$
- $\partial X$ = Thickness of slab in feet
- $\partial T$ = Temperature drop across the slab (°F)

Therefore, $K$ is expressed as:

- Btu-ft/hr/ft$^2$ °F
- cal/(meter °C sec)
- W/m-K
Sample Calculation:
Assume a flat plate of graphite 0.1 foot thick has a surface area of one square foot. Heat is flowing through this plate at a rate of 1000 Btu per hour; the hotter surface is at a temperature of 1036°F, while the cooler surface is at 1018°F; calculate the thermal conductivity.

\[
K = \frac{Q \alpha X}{A \delta T} = \frac{(1000 \text{ Btu})(0.1 \text{ ft})}{(\text{hr})(1 \text{ ft}^2)(1036^\circ F - 1018^\circ F)}
\]

\[
K = \frac{(1000 \text{ Btu})(0.1 \text{ ft})}{(\text{hr})(1 \text{ ft}^2)(18^\circ F)} = 55.5 \frac{\text{Btu-ft}}{\text{hr/ft}^2 \text{ °F}}
\]

To convert to metric system of units, multiply by 0.413 to obtain:

\[
K = 22.9 \text{ cal/} (\text{meter °C sec})
\]

To convert to SI system of units, multiply by 1.73 to obtain:

\[
K = 96.0 \text{ W/m-K}
\]

Test Method
Historically, two different techniques were employed for determining thermal conductivity over the temperature range of 110–3300 K. They are:

1. A comparative rod apparatus from 110 K to 1250 K
2. A radial inflow apparatus from 1250 K to 3300 K

There is no standard method associated with graphite for measurement of thermal conductivity.

However, ASTM Standard C714-72 is routinely used for determining thermal diffusivity and the corresponding results are used to calculate thermal conductivity. POCO graphite deviates from ASTM Standard C714-72 in that this standard requires the use of a flash lamp for sample heating and thermocouples for monitoring temperature. A similar technique employed by Entegris utilizes a Netzsch® LFA-427 (Laser Flash Apparatus), which has a high-intensity laser to heat the sample surface and the resultant temperature change is monitored with an infrared detector (Figure 11-1).

Thermal diffusivity measurements involve quantifying the rate of diffusion of heat through a solid where local temperature and temperature gradients vary with time. This is a decided advantage in that thermal diffusivity measurements are nonsteady-state, due to the transient nature of diffusion, thus eliminating the need for determining heat flux and maintaining steady-state conditions. Nonetheless, close control of local time-temperature relationships in the system is required. Thermal diffusivity methods are also preferred over heat flux measurements due to the ease of sample preparation. With less material requirements and fast measurements, one typically obtains accuracy within ±5% when implementing careful measurement techniques. Thermal conductivity is calculated from the relationship:

\[
K = \alpha C_P \rho
\]

Where:
- \( K \) = Thermal conductivity (W/cm K)
- \( \alpha \) = Thermal diffusivity (cm²/sec)
- \( C_P \) = Heat capacity at constant pressure (J/g K)
- \( \rho \) = Apparent density (g/cm³)

POCO Graphite vs. Conventional Graphites
From studies on POCO graphite and experimental work on other graphites, it is believed that the conductivity of most graphites is mainly governed by Umklapp phonon-phonon interactions.

For POCO graphite and most other graphites, thermal conductivity begins to decrease around 27°C (70°F) and continues to decrease as the temperature increases to 3227°C (5800°F). Thermal diffusivity and thermal conductivity results for POCO graphite grades are indicated in Figures 11-2 and 11-3, respectively, as a function of temperature from 23–1650°C.

It has been reported that above 2727°C (4900°F), electrons contribute about 20 percent to the total thermal conductivity of these graphites. For more highly graphitized graphites and other feedstocks, the results can be different.
Density Effects

The density of graphite is significant in its effect, as on the other properties. As the density of POCO graphite increases, its thermal conductivity also increases.

Thermal Shock

Definition

The ability of a material to be subjected to sudden thermal gradients without weakening or fracturing is referred to as its thermal shock resistance. The excellent resistance of graphite to thermal shock is due to a unique set of properties. High strength, low modulus of elasticity and low coefficient of thermal expansion are the properties that are most important. This theoretical relationship can be expressed as:

$$ R = \frac{\sigma_f (1-\Omega) K}{\alpha E} $$

Where: $R$ = Thermal shock resistance  
$\sigma_f$ = Fracture stress  
$\Omega$ = Poisson’s ratio  
$\alpha$ = Coefficient of thermal expansion  
$E$ = Modulus of elasticity

This relationship applies only when the quench is so fast that the surface temperature reaches final value before the average temperature changes.

For conditions when the heating rate is not very high, a second relationship can be established which includes thermal conductivity. This is expressed as:

$$ R' = \frac{\alpha_f (1-\Omega) K}{\alpha E} $$

Where: $R'$ = Thermal shock resistance  
$\sigma_f$ = Fracture stress  
$\Omega$ = Poisson’s ratio  
$K$ = Thermal conductivity  
$\alpha$ = Coefficient of thermal expansion  
$E$ = Modulus of elasticity

Another way to express this would be:

$$ R = \frac{KT_S}{\alpha E_T} $$

Where: $R$ = Thermal shock resistance  
$K$ = Thermal conductivity  
$T_S$ = Tensile strength  
$\alpha$ = Coefficient of thermal expansion  
$E_T$ = Tensile modulus of elasticity

This expression may be a simpler form to use and consequently will be used in the sample calculation.

---

In either case, the typical properties of carbon and graphite generally yield the best overall thermal shock resistance of all the temperature-resistant, non-metallic materials available. This is useful for rocket nozzle and reentry nose-cone applications, among others.

**Sample Calculation:**

A graphite sample with a thermal conductivity of 0.29 (cal cm)/(cm² sec °C) and a tensile strength of 10,000 psi (703 kg/cm²) has a modulus of elasticity of $1.6 \times 10^6$ psi ($0.112 \times 10^6$ kg/cm²) and a CTE of $8.4 \times 10^{-6}$ (in/in)/(°F); calculate its thermal shock resistance.

\[
R = \frac{KT_s}{\alpha E_T} = \frac{0.29 \text{ cal} \times \text{cm}^2}{\text{cm}^2 \times \text{sec} \times \degree \text{C}} \times \frac{703 \text{ kg}}{0.112 \times 10^{-6} \text{ kg/cm}^2} = 216.7 \text{ cal/cm} \times \text{sec}
\]

**Test Method**

There are no standard test methods for accurately measuring the thermal shock resistance of materials. A number of practical tests are being used for specific applications. One such testing device is described by Sato. Use of the relationships previously described are generally accepted for prediction of the thermal shock resistance of a material, but many other considerations must be made such as sample size, stress distribution in material (i.e., geometry and stress duration).

**POCO Graphite vs. Conventional Graphites**

Some early studies reported that coarse-particle graphite had better thermal shock resistance than finer-particle graphite despite the higher strength of finer-particle graphite. The study further indicated a correlation to the binder. If the binder were decreased, the thermal shock resistance also dropped, suggesting the thermal stresses were being absorbed by the binder material. Other studies have shown the precursor material, e.g., tar coke versus oil coke, to have effects on the thermal shock resistance. The oil coke had resistance values about seven times higher than the tar coke.

Generally, due to small particle size and high CTE, POCO graphite may not fare as well as some other graphite in overall thermal shock resistance. Tests at Entegris have shown that samples of the DFP-1 grade, measuring one inch cubed, can survive a 1000°C to 10°C instantaneous change without fracturing in repeated tests. Other tests on POCO graphites have subjected the samples to 335°F to -335°F instantaneous changes without affecting the graphite. A comparison of some thermal shock resistance values for some materials is shown in Table 12-1.

<table>
<thead>
<tr>
<th>Material</th>
<th>Resistant to Thermal Shock</th>
</tr>
</thead>
<tbody>
<tr>
<td>POCO graphite</td>
<td>217</td>
</tr>
<tr>
<td>GLC</td>
<td>450</td>
</tr>
<tr>
<td>Mersen graphite</td>
<td>140</td>
</tr>
<tr>
<td>Pyrolytic graphite</td>
<td>4300</td>
</tr>
<tr>
<td>Titanium carbide</td>
<td>3.45</td>
</tr>
<tr>
<td>Toyo Tanso USA graphite</td>
<td>340</td>
</tr>
<tr>
<td>UCAR</td>
<td>329</td>
</tr>
</tbody>
</table>

While the differences are not as great between the various grades as the pyrolytic graphite shows between orientations, it should still be noted that processing variables, starting materials, etc., can influence the thermal shock resistance of graphite. For example, the Ringsdorff EK-87 grade has an average particle size of 20 microns, yet it has about 30 percent more calculated thermal shock resistance than the UCAR ATJ, which has an average particle size of 25 microns and nearly twice the calculated resistance of a POCO graphite with a particle size average of 4 microns.

**Temperature Effects**

Depending on the type of graphite used and its particular characteristics or properties, and considering shape, size, etc., higher temperature drops could be sustained without affecting the graphite. The temperature starting and ending points may have some bearing on the overall resistance to fracture, because the material will be thermally stressed differently at elevated temperatures. A drop from 1000°C to 500°C may respond differently than from 500°C to 0°C. Little data is available to clearly define what the relationship or limits are for the different types of graphite.

---

Density Effects

Since, generally, strength and thermal conductivity increase with density, higher thermal shock resistance should be found in higher-density ranges. However, with higher density, higher modulus and CTE are also usually found and these would tend to offset the gains in strength. The coarser-particle systems may be more effective than the finer-particle systems on an equal density basis. Many factors are involved, and well defined tests to measure these effects are not in common use.

Specific Heat

Definition

Heat capacity is the quantity of heat required to raise the temperature of a unit mass of material 1°, and the relationship between the two most frequently used units is as follows:

\[
\frac{1 \text{ Btu}}{\text{lb} \cdot \text{°F}} = \frac{1 \text{ cal}}{\text{g} \cdot \text{°C}}
\]

It has been found that the values of heat capacity for all types of natural and manufactured graphites are basically the same, except near absolute-zero temperatures. The differences found in measuring the same grade of graphite are as great as the differences in measuring different grades of graphite. Differences as much as nine percent have been found between natural graphite and manufactured graphite at low temperatures (120 K to 300 K), but at higher temperatures the differences for all types of graphite has been found to be less than the experimental error. Table 13-1 shows comparisons to other materials.

Heat Capacity Equation

The heat capacity at constant pressure \((C_P)\) can be expressed by polynomial functions of the absolute temperature \(T\), with \(T\) in K to give \(C_P\) in cal/(g°C).

Within the temperature interval 0 K to 300 K:

\[
C_P = (0.19210 \times 10^{-4})T - (0.41200 \times 10^{-5})T^2 - (0.10831 \times 10^{-7})T^3 - (0.10885 \times 10^{-10})T^4
\]

However, below 40 K, this equation has not been reliable in correlation with experimental data. The calculated values are in good agreement (<2.2 percent) with the experimental values for temperatures above 40 K.

For the temperature range 300 K to 3200 K:

\[
C_P = 0.44391 + (0.30795 \times 10^{-4})T - (0.61257 \times 10^{-7})T^2 + (0.10795 \times 10^{-9})T^3
\]

This expression yields calculated values within 1.5 percent of the experimental values for the entire temperature range.

Table 13-2 represents the best fit of the values for typical manufactured graphite with proper consideration given to the accuracy of the individual measurements.

Sample Calculation:

Determine the heat capacity of a sample of graphite at 500°C using a differential scanning calorimeter (DSC) and a strip chart recorder to follow the calorimetric response for the temperature scans of 30°C. DSC scans are taken for the sample of graphite, a sapphire reference sample and the baseline for correction of the measured response. The heat capacity, \(C_P\) is then calculated for the graphite using the formula:

\[
C_P = \frac{W_S}{W_G} \times \frac{D_S}{D_G} \times C_P \text{ (sapphire)}
\]

Where: \(W_S\) = Weight of sapphire
\(W_G\) = Weight of graphite
\(D_S\) = Signal displacement of sapphire
\(D_G\) = Signal displacement of graphite

<table>
<thead>
<tr>
<th>Material</th>
<th>Heat Capacity Range cal/(g°C)</th>
<th>Heat Capacity Range J/(g K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gold</td>
<td>0.031</td>
<td>0.13</td>
</tr>
<tr>
<td>Platinum</td>
<td>0.031</td>
<td>0.13</td>
</tr>
<tr>
<td>Tungsten</td>
<td>0.034</td>
<td>0.14</td>
</tr>
<tr>
<td>Tungsten carbide</td>
<td>0.04</td>
<td>0.17</td>
</tr>
<tr>
<td>Silver</td>
<td>0.056</td>
<td>0.23</td>
</tr>
<tr>
<td>Brass</td>
<td>0.090</td>
<td>0.38</td>
</tr>
<tr>
<td>Nickel</td>
<td>0.091–0.14</td>
<td>0.38–0.59</td>
</tr>
<tr>
<td>Copper</td>
<td>0.092</td>
<td>0.38</td>
</tr>
<tr>
<td>Steel (wrought)</td>
<td>0.11</td>
<td>0.46</td>
</tr>
<tr>
<td>Iron (cast)</td>
<td>0.13</td>
<td>0.54</td>
</tr>
<tr>
<td>Graphite</td>
<td>0.17</td>
<td>0.72</td>
</tr>
<tr>
<td>Alumina</td>
<td>0.19</td>
<td>0.79</td>
</tr>
<tr>
<td>Aluminum</td>
<td>0.22–0.23</td>
<td>0.92–0.96</td>
</tr>
<tr>
<td>Silicon carbide</td>
<td>0.285–0.34</td>
<td>1.19–1.42</td>
</tr>
</tbody>
</table>
Let the weight of the sapphire reference = 0.203 g and a sample of DFP graphite, i.e., indicating a density of 1.82 g/cm$^3$ and measuring 1.5 mm × 6 mm diameter, weigh 0.077 g, respectively. $C_p$ of sapphire is 0.19 cal/(g°C). $C_p$ of graphite at 500°C can be calculated by knowing the displacement as determined from the strip chart recorder.

Assume the displacement (corrected) is 40.0 for the graphite and 52.5 for the sapphire reference.

Therefore:

$$C_p \text{ (graphite)} = \frac{0.023 \text{ g}}{0.077 \text{ g}} \times \frac{40}{52.5} \times 0.19 \frac{\text{cal}}{\text{g °C}}$$

$$C_p \text{ (graphite)} = 0.382 \frac{\text{cal}}{\text{g °C}} = 1.6 \frac{\text{J}}{\text{g K}}$$

**Test Method**

The previous example best illustrates how a DSC and a chart recorder may be used in determining heat capacity. However, modern technology has integrated the DSC with computer and software technology, thereby eliminating the need for calculating $C_p$ from a chart recorder. Entegris currently uses a Netzsch STA-449 Heat Flux DSC (Figure 13-1) for evaluating graphite and other related materials. Heat capacity calculations are made via software but the fundamental principles remain the same.

![Figure 13-1. Netzsch STA-449](image-url)

In DSC, a distinction must be made between the recorded baseline in the presence and absence of a sample. Thus, measurements are first taken with an empty sample crucible to determine the instrument baseline. A second run is made to verify the accuracy of the instrument by placing a sapphire reference sample in the sample crucible and taking measurements.
under the same experimental conditions. The DSC curve rises linearly resulting from the change in heat capacity of the sample as a function of temperature. Sapphire is routinely used as a reference material since it is stabilized, i.e., will not oxidize, and due to the fact that heat capacity for sapphire has been thoroughly investigated and well documented. Once the baseline and accuracy have been established, a third run is made on the sample of interest under the same experimental conditions.

There is no standard test method for determining specific heat of graphite, but the DSC method is commonly used for specific heat determination of other materials.

Another simple method which yields approximate specific heat data is an ice calorimeter. If a body of mass \(m\) and temperature \(t\) melts a mass \(m_1\) of ice, its temperature being reduced to 0°C, the specific heat of the substance is:

\[
C_p = \frac{m_1 \cdot C_i}{m - m_1}
\]

**POCO Graphite vs. Conventional Graphites**

POCO graphites have heat capacity values that are in the same range as conventional manufactured graphites. POCO graphites have a range from around 0.72 J/(g K) to 1.60 J/(g K) between 20° and 500°C (Figure 13-2).

**Temperature Effects**

The heat capacity of manufactured graphite increases with increasing temperature (Figure 13-3).

**Density Effects**

Heat capacity has no significant change as a function of density. It is intrinsic to the graphite crystal itself and apparently independent of density.

**Emissivity**

**Definition**

A measure of the heat radiating ability of a surface is referred to as emissivity. Total emissivity or spectral emissivity are ways to express it. The total emissivity is defined as the ratio of thermal energy emitted by a material per unit time per unit area to that emitted by a black body over the entire band of wavelengths when both are at the same temperature. A black body, i.e., ideal radiator, is one which absorbs all the radiant energy falling on it and, at a given temperature, radiates the maximum amount of energy possible over the entire spectrum of wavelengths.

The spectral emissivity is defined as the ratio of thermal energy emitted by a material per unit time per unit area to that emitted by a black-body reference where radiation from both are of the same wavelength and both are at the same temperature. The spectral emissivity of graphite has been measured at 6500 Å. Dull surfaced graphite and polished graphite have had typical values of 0.90 and 0.77, respectively. It has been determined that the emissivity of graphite does vary slightly with temperature. A temperature coefficient of \(1.9 \times 10^{-5}/\text{K}\) has been determined. Graphite emissivity has been found to be nearly constant in the wavelength range 2000 Å to 60,000 Å. Spectral emissivity values approaching 0.99 have been measured for graphite near sublimation temperatures of 3600 K.

---

For a specific temperature, at a specific wavelength, normal spectral emissivity can be expressed as:

\[ E_{n,\lambda} = \frac{W_{s,\lambda}}{W_{BB,\lambda}} \]

Where:
- \( W_{s,\lambda} \) = Normal spectral radiant energy emitted by a specimen material per unit time, unit area and unit solid angle.
- \( W_{BB,\lambda} \) = Normal spectral radiant energy emitted by a black-body reference per unit time, unit area, and unit solid angle.

Normal spectral emissivity written as a function of temperature is:

\[ \log E_{n,\lambda} = \frac{C}{2.303\lambda} \left[ \frac{1}{T} - \frac{1}{T_{A,\lambda}} \right] \]

Where:
- \( C = 14,380 \) micron – K
- \( \lambda = 0.65 \) micron
- \( T = \) Temperature of material (K)
- \( T_{A,\lambda} = \) Apparent temperature of material (K)

For a specific temperature the total normal emissivity can be expressed as:

\[ E_{n,t} = \frac{W_{s}}{W_{BB}} \]

Where:
- \( W_{s} \) = Normal total radiant energy emitted by a specimen material per unit time, unit area and unit solid angle.
- \( W_{BB} \) = Normal total radiant energy emitted by a black body reference per unit time, unit area and unit solid angle.

Total normal emissivity at elevated temperatures can be expressed as:

\[ E_{n,t} = \frac{T_{A,t}}{T_{4}} \]

Where:
- \( T = \) Temperature of material (K)
- \( T_{A,t} = \) Apparent temperature of material (K)

Emissivity data has usefulness in applications such as aerospace, heaters, pyrometry, etc., but published data is difficult to find.

**Test Method**

There are no standard methods for measuring emissivity of graphite, but one method used to measure the emissivity of graphite is described by Grenis and Levitt.\(^{13}\) A schematic of this equipment can be seen in Figure 14-1. The methods of measuring emissivity are wrought with difficulties and results may be biased due to equipment and/or assumptions made in running the tests. At best, the testing is at a research level and not a routine procedure.

**POCO Graphites vs. Conventional Graphites**

Very little information on the emissivity of graphite is available. The following information on total emittance of two grades of POCO graphite and pyrolytic graphite has been reported by Cezairliyan and Righini.\(^{14}\)

The equation for total emittance of POCO (TRA-1) graphite as a function of temperature, between 1800 to 2900 K is:

\[ E = 0.679 + (6.00 \times 10^{-5})T \]

The equation for POCO (DFP-2) graphite, between 1700 to 2900 K is:

\[ E = 0.794 + (2.28 \times 10^{-5})T \]

The equation for pyrolytic graphite, between 2300 to 3000 K is:

\[ E = 0.641 - (5.70 \times 10^{-5})T \]

Where: \( T \) is in K.

---


The measured values of the total emittance of the graphite specimens are shown in Table 14-1 and again graphically in Figure 14-2.

<table>
<thead>
<tr>
<th>Temperature (K)</th>
<th>POCO TRA-1</th>
<th>POCO DFP-1</th>
<th>Pyrolytic Graphite</th>
</tr>
</thead>
<tbody>
<tr>
<td>1700</td>
<td>—</td>
<td>0.833</td>
<td>—</td>
</tr>
<tr>
<td>1800</td>
<td>0.787</td>
<td>0.835</td>
<td>—</td>
</tr>
<tr>
<td>1900</td>
<td>0.793</td>
<td>0.837</td>
<td>—</td>
</tr>
<tr>
<td>2000</td>
<td>0.799</td>
<td>0.840</td>
<td>—</td>
</tr>
<tr>
<td>2100</td>
<td>0.805</td>
<td>0.842</td>
<td>—</td>
</tr>
<tr>
<td>2200</td>
<td>0.811</td>
<td>0.844</td>
<td>—</td>
</tr>
<tr>
<td>2300</td>
<td>0.817</td>
<td>0.846</td>
<td>0.510</td>
</tr>
<tr>
<td>2400</td>
<td>0.823</td>
<td>0.849</td>
<td>0.504</td>
</tr>
<tr>
<td>2500</td>
<td>0.829</td>
<td>0.851</td>
<td>0.499</td>
</tr>
<tr>
<td>2600</td>
<td>0.835</td>
<td>0.853</td>
<td>0.493</td>
</tr>
<tr>
<td>2700</td>
<td>0.841</td>
<td>0.856</td>
<td>0.487</td>
</tr>
<tr>
<td>2800</td>
<td>0.847</td>
<td>0.858</td>
<td>0.481</td>
</tr>
<tr>
<td>2900</td>
<td>0.853</td>
<td>0.860</td>
<td>0.476</td>
</tr>
<tr>
<td>3000</td>
<td>—</td>
<td>—</td>
<td>0.470</td>
</tr>
</tbody>
</table>

### Ash

#### Definition

ASTM Standard C709 defines ash of manufactured carbon and graphite as the “residual product following oxidation of the base carbon as determined by prescribed methods.” This ash or residual product generally results from natural contamination of the raw material used to produce graphite and may be introduced to a lesser extent during processing of the raw material or graphite products. Table 15-1 shows a general listing of typical elements and their level of contamination in graphite bulk product. As will be discussed later, there are means by which the contamination can be reduced or removed.

#### Test Method

The standard test method for ash content of graphite is ASTM Standard C561. This technique applies principally to nonpurified graphites of several hundred ppm, or more, of total impurities. The method is generally too crude for good reproducibility on high-purity samples.
One of the disadvantages of this method is the potential loss of low melting point impurities due to the temperature at which the test is run. Theoretically, at 950°C as a final test temperature, nothing should be lost that was not already volatilized during graphitization, but as a practical matter, it seems to occur. These are possibly impurities trapped in closed pores that did not successfully escape during the graphitization cycle. The test temperature could be lowered to preclude some of this from occurring, but then the time to complete the test is extended considerably as oxidation is a time-temperature dependent reaction.

It has also been found in previous studies that at higher temperatures (desirable for shorter test cycles) reactions of carbon and platinum, of which the crucibles are made, can occur and consequently bias the test results.

Entegris’ method, TDI 4.1.1.6 (Appendix B), differs slightly from the ASTM procedure, in that a solid sample is used rather than a powdered sample. The risk of contamination in powdering the sample is nonexistent if a solid sample is used. A standard test cycle for nonpurified material is 24 hours, whereas, a purified material usually takes 48–72 hours to complete the cycle.

Sample Calculation:
If a graphite sample weighed 80.0000 grams after drying and was ashed in a 35.0000 grams crucible where the total ash after testing was 0.4 milligrams, the total ash would be 5 ±1 parts per million.

\[
\text{Ash (ppm)} = \frac{C - A}{B - A} \times 1,000,000 = \frac{35.0004 - 35.000}{115.0000 - 35.000} \times 1,000,000 = 5 \pm 1 \text{ ppm}
\]

Where:  
A = Crucible weight  
B = Crucible plus dried sample weight  
C = Crucible plus ash weight

If ash % is required, multiply by \( \frac{C - A}{B - A} \times 100 \)  
Ash % = 0.0005

POCO Graphites vs. Conventional Graphites

When discussing nonpurified graphites, POCO graphites are fairly clean, i.e., contain relatively low levels of impurities. However, since graphitization temperature has a strong bearing on the impurities which remain, the ash levels on any product can vary accordingly. There are other process steps which influence the final ash level also.

In material which has been impregnated for densification, contaminants in the raw material used could increase the impurity content as well.

POCO graphites will typically range from 300–3000 ppm total impurities, depending on the grade and density. Many conventional graphites fall within this range also, but some will be in the range of 0.1 percent (1000 ppm) to several percent (20,000–30,000 ppm).

Purified POCO graphite will have less than 5 ppm total impurities. This is one of the best grades of purified graphite available anywhere. The unique nature of the POCO pore structure and relatively clean raw materials aids in allowing this ultra-high purity. Typical analysis of a purified POCO graphite is seen in Table 15-2. Purified graphites can be produced in several ways. Simply heat-treating to very high temperatures, usually in excess of 3000°C, will volatilize most heavy elements present. Halogen gases, such as chlorine or fluorine, will react with the impurities at high temperature and volatilize off as chloride or fluoride salts. Some elements such as boron are very stable and difficult to remove, especially since they can substitute for carbon atoms in the crystal structure.

Temperature Effects

The only real temperature effect on ash levels comes from the graphitization temperature or a subsequent thermal processing to remove impurities. As mentioned previously, the higher the final thermal treatment, the lower the ash level will usually be.

Density Effects

In POCO graphite, a slight trend has been noted with ash as related to density. Since POCO graphites have increasingly more open porosity as the density decreases, more opportunity for volatilization escape of the impurities during graphitization can occur. Consequently, slightly lower ash levels are typically found for the lower-density grades. This may be unique to POCO graphites. The difference is relatively small though, so it is mentioned only in passing, as a number of other factors may have more significant effects on the final ash level.
TABLE 15-2. TYPICAL PURITY ANALYSIS
PURIFIED POCO GRADES
TOTAL ASH RANGE 5 PPM OR LESS

<table>
<thead>
<tr>
<th>Element</th>
<th>Element Detected</th>
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</thead>
<tbody>
<tr>
<td>Silicon (Si)</td>
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<tr>
<td>Sulfur (S)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Vanadium (V)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Calcium (Ca)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Boron (B)</td>
<td>Yes</td>
<td>Trace</td>
</tr>
<tr>
<td>Aluminum (Al)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Magnesium (Mg)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Iron (Fe)</td>
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</tr>
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<td>Molybdenum (Mo)</td>
<td>Yes</td>
<td>Trace</td>
</tr>
<tr>
<td>Phosphorus (P)</td>
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<td>Trace</td>
</tr>
<tr>
<td>Nickel (Ni)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Titanium (Ti)</td>
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<td>—</td>
</tr>
<tr>
<td>Potassium (K)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Sodium (Na)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Copper (Cu)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Chromium (Cr)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Zinc (Zn)</td>
<td>No</td>
<td>—</td>
</tr>
<tr>
<td>Lithium (Li)</td>
<td>No</td>
<td>—</td>
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</tbody>
</table>

Oxidation

Definition

Oxidation is a chemical reaction that results in electrons being removed from an atom, which makes the atom more “reactive” so that it may join with one or more other atoms to form a compound. When talking about carbon and graphite, oxidation is normally thought of as the reaction of carbon atoms with oxygen to form carbon monoxide (CO) and carbon dioxide (CO₂). However, many other gases (besides oxygen) will react with carbon in an oxidation reaction (e.g., CO₂, H₂O, N₂O, etc.). The result of the oxidation reaction is a loss of carbon atoms from the carbon or graphite material, which obviously affects many of its properties and characteristics. If oxidation of a piece of carbon is allowed to continue long enough, all the carbon atoms will react to form a gas (or gases) which dissipates and leaves behind nothing except a small amount of ash, which is the oxides of the metallic impurities that were present in the carbon to begin with.

The oxidation of carbon by oxygen (as well as other gases) is highly temperature-dependent. No detectable reaction occurs at temperatures up to about 350°C. As the temperature is increased, the rate of reaction increases rapidly, according to the well-known Arrhenius expression:

\[ \ln K \propto \frac{E}{RT} \]

Where: 
- \( T \) = Temperature
- \( E \) = Activation energy
- \( K \) = Reaction rate
- \( R \) = Universal gas constant

Test Method

To measure the oxidation characteristics of a particular carbon material, a piece of known weight is exposed to the “oxidizing environment” of interest for some period of time, and then reweighed, to determine a weight loss (the oxidation weight loss). There is not a widely accepted standard test for measuring the oxidation characteristics of carbon materials (TDI 4.1.1.7 and 4.1.1.8 in Appendix B).

The oxidation characteristics of carbon are usually expressed in one of two different ways:

1. The percent weight loss in 24 hours at a given temperature, often referred to as oxidation resistance
2. The temperature at which a sample loses approximately one percent of its weight in a 24-hour period, which is called the oxidation threshold temperature of the material being tested

Sample Calculation:

If a sample originally weighed 10.0000 grams after being dried and 9.9900 grams after oxidation testing, the percent weight loss would be 0.1 percent.

\[ \%WT_L = \left( \frac{WT_O - WT_F}{WT_O} \right) \times 100 = \left( \frac{10.0000 - 9.9900}{10.0000} \right) \times 100 = 0.1\% \]

Where:
- \( WT_O \) = Initial sample weight (dried)
- \( WT_F \) = Sample weight after oxidation testing percent
- \( WT_L \) = Percent weight loss
POCO Graphites vs. Conventional Graphites

The oxidation characteristics of any graphite at the same temperature and atmospheric conditions are dependent on the amount of impurities in the material, the density of the material and the amount of surface area available to react with the oxidizing atmosphere. POCO graphite can be impregnated with a proprietary oxidation inhibitor that can substantially increase its oxidation resistance. Purification also reduces oxidation significantly by removing metallic impurities, which act as oxidation catalysts.

As the surface area increases, the oxidation rate increases, too. This is expected, as the surface exposure is important to the oxidation process. For purposes of standardization and to minimize the effect of variable surface area to volume (SA/V) ratios, the use of a sample with an SA/V ratio of 10:1 is used at Entegris for testing oxidation rates.

Temperature Effects

The differences in oxidation behavior of the various grades of graphite are widest at the lowest temperatures, tending to disappear as the temperature increases. The “oxidation threshold temperature,” defined as that at which a sample loses approximately one percent of its weight in 24 hours, is about 570°C for purified graphite, while nonpurified graphite is about 430°C (Figure 16-1).

The oxidation of graphite is highly temperature-dependent. At temperatures up to about 350°C, no detectable oxidation occurs. As the temperature is increased, the rate of reaction increases rapidly, according to the Arrhenius equation (Figure 16-2).

Density Effects

Since the oxidation of graphite is a surface reaction, it is known that the rate of oxidation is affected by the porosity of the material. As the percent porosity increases, the apparent density of the material will decrease. Therefore, as the apparent density decreases, the rate of oxidation will increase. However, the effects of impurities and their catalytic action can easily override the density effects. Purified data shows the density difference more clearly, but it is, nevertheless, relatively small when other factors are considered.

Figure 16-1. Oxidation threshold

Figure 16-2. Oxidation rate
## Appendix A

### Prefix Conversion Factors

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<thead>
<tr>
<th>Prefix</th>
<th>Symbol</th>
<th>Unit Multiplier</th>
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<td>$10^9$</td>
</tr>
<tr>
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<tr>
<td>kilo</td>
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<td>d</td>
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<tr>
<td>centi</td>
<td>c</td>
<td>$10^{-2}$</td>
</tr>
<tr>
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<td>m</td>
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<tr>
<td>micro</td>
<td>μ</td>
<td>$10^{-6}$</td>
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<td>n</td>
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<td>pico</td>
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### Density Conversion Factors

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<td>g/m$^3$</td>
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<td>lb/in$^3$</td>
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### Length Conversion Factors

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<td>inch</td>
<td>m</td>
<td>0.0254</td>
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<td>inch</td>
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<td>mm</td>
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</tr>
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<td>feet</td>
<td>in</td>
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</tr>
<tr>
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<td>m</td>
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<td>feet</td>
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### Strength Conversion Factors

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<td>kg/m$^2$</td>
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</tr>
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<td>lb/in$^2$</td>
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ELECTRICAL RESISTIVITY CONVERSION FACTORS

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</tr>
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<td>(\Omega)-cm</td>
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COEFFICIENT OF THERMAL EXPANSION

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<td>(in/in)/°C</td>
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<td>(in/in)/°C</td>
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Conversion constants for 1/°C to 1/°F and 1/°F to 1/°C are true for any dimensional unity, i.e., in/in, cm/cm, ft/ft, m/m.

THERMAL CONDUCTIVITY CONVERSION FACTORS

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</thead>
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<td>0.0173</td>
</tr>
</tbody>
</table>

1 calorie = 1 cal = 1 gram cal
1 calorie = 1 kilogram cal = 1 Kcal = 1000 cal
Appendix C Bibliography

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