Natural Graphite-CVD Carbon Composites


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Graphite-carbon composites have been fabricated using a natural graphite powder which was molded at low pressures and room temperature and infiltrated by a CVD technique at 700°C using propylene as a source of carbon. The rate of removal of open porosity by carbon infiltration is proportional to open porosity up to an infiltration time of at least 24 hr. Flexural strength, Young's modulus and electrical conductivity of the composites progressively increase with increasing apparent density (d), being approximated by expressions of the type Y = ad^n over most of the infiltration range. Knoop hardness is increased by infiltration but remains much less than that found for cellulose carbon-CVD composites of comparable infiltration. Infiltration has interesting effects on CTE parallel and perpendicular to the original molding direction and CTE ratios. Heat treatment of infiltrated composites up to 2400°C results in density decreases and some degradation in mechanical properties.

1. INTRODUCTION

As has been shown previously (1, 2), natural graphite powders can be molded at room temperature into artifacts. Variables like molding pressure, subsequent HTT, and particle size distribution of the graphite powder affect the density and pore size distribution of the artifacts produced. However, compaction at pressures up to 9,000 kg/cm² does not decrease the surface area of the graphite, as measured by N₂ adsorption at 77 K. This means that compaction has not created apertures (slits) less than about 5.2A in thickness (3). Increasing molding pressure (increasing density) results in increasing crushing strengths and decreasing electrical resistivities for the artifacts (1, 2). However, reactivity of artifacts to CO₂ at 800°C is unaffected by previous molding pressures (1).

Recently we have reported on the preparation of composites by the infiltration of porous cellulose carbon artifacts with CVD carbon from propylene (4). The cellulose carbon artifacts (substrates) prepared by molding at room temperature, curing at 250°C, and carbonizing at 800°C consisted of a disordered carbon structure and exhibited isotropic properties. A reasonably high transverse breaking strength (100 kg/cm²) for the uninfiltreted artifact, despite its low density (0.7 g/cc), suggested that some particle consolidation had occurred during the carbonization step. This consolidation could be the result of cross linking reactions occurring during curing of the cellulose.

It was, therefore, of practical and scientific interest to study the preparation of composites from the CVD infiltration of molded natural graphite artifacts. Practically, such composites may have application where anisotropic properties are desired. Scientifically, it is of interest to compare rates of CVD over substrates as markedly different in structure as cellulose carbon and natural graphite.

2. EXPERIMENTAL

2.1 Preparation of uninfiltreted artifacts

Natural graphite powder (SP-1 grade from the Carbon Products Division of Union Carbide Corporation) was used as the starting material. It had a total impurity content of less than 6 ppm and a specific BET surface area of 1.8 m²/g, as measured by N₂ adsorption at 77 K. From quantitative x-ray diffraction studies on the (004) diffraction peak, the interlayer spacing of the material was 3.3541Å at 15°C. Microscopic investigation revealed the powder to be composed of flake-like particles having a mean diameter of 30µ on a number basis. The particle thickness is estimated at 0.5µm (5).

In order to produce artifacts with a large open porosity, the graphite powder was molded into bars at a low pressure, that is 370 kg/cm². The bars had dimensions of 10.9 cm long, 1.2 cm wide, and 0.35 cm thick, with the thickness dimension parallel to the molding direction. The bars had an apparent density of 1.65 g/cc or an open porosity (based on a helium density for the graphite powder of 2.25 g/cc) of 36.2%. This is to be compared with an open porosity of 62% in the unin-
filtrated cellulose carbon artifacts used in the previous CVD studies (4). Molding resulted in some preferred orientation of the graphite particles, with their long dimension perpendicular to the pressing direction. That is, the (002) peak intensity ratio for two mutually perpendicular directions (I_{002}/I_{T_{002}}), see reference 6, had a value of 11.3.

2.2 Infiltration conditions
A 50% propylene-50% helium mixture (by volume) was used at atmospheric pressure and 700°C for infiltration, as previously described (4). This temperature was selected previously since infiltration for 72 hr did not result in noticeable external surface deposition. The gas mixture was passed in laminar flow through the reactor, it having a residence time of 4 min in the 46 cm hot zone, as previously described (4). Up to four samples could be infiltrated at one time.

2.3 Measurement of sample properties
All properties were measured using approaches similar to those described previously to characterize the cellulose-CVD composites (4).

3. RESULTS AND DISCUSSION

3.1 Kinetics of the CVD process
As found previously (4), the rate of decrease in open porosity upon infiltration of the artifacts was proportional to the fraction of open porosity (δ) present in the sample at time (t) over most of the run. That is, 

\[-d\delta/dt = k\delta\]

Up to 24 hr infiltration time, the first order plot of log(100 δ₀ / δ) vs time was linear, having a slope of -0.0137 hr⁻¹. This is to be compared with a slope of -0.00877 hr⁻¹ found for the first order plot for infiltration of the cellulose carbon artifact.

Figure 1 presents results for the weight of carbon pick-up versus time on the natural graphite and cellulose artifacts. Results are normalized by dividing the weight of carbon deposited by the initial weight of the artifact used. Interestingly, up to about 24 hr infiltration time, the rate of carbon pick-up on the natural graphite is greater than that on the cellulose carbon, despite the fact that their initial total surface areas, as measured by N₂ adsorption at 77K, are 1.8 and 480 m²/g, respectively. Studies in progress in this laboratory are showing, however, that it is not the total surface area (TSA) but the active surface area (ASA) which determines the rate of propylene cracking. The active area is located at the prismatic faces of carbon crystallites and around point defects within the basal plane. On the basis of our recent studies in this laboratory on the cracking of propylene over activated Graphon, the disappearance of propylene is first order in propylene pressure with a rate constant at 700°C of 8.3 x 10⁻⁵ cm/sec. To calculate this rate constant, the ASA of the Graphon was measured by oxygen chemisorption at 300°C (7). Using this value and the rate expression

\[\text{Rate} = (k_{111})(C_{C_{2}H_{4}})(ASA)\]

where \(k_{111}\) is the rate constant and \(C_{C_{2}H_{4}}\) is the concentration of propylene in the reactor over the graphite artifacts, it is possible to predict an initial rate for carbon deposition over the SP-1 graphite. If the ASA of SP-1 graphite is taken as 3.7% of the TSA (5) and if 50% of the carbon in the propylene molecule goes to deposited carbon and 50% to methane and
ethylene (as is roughly found), the initial rate of carbon deposition is predicted to be $2.3 \times 10^{-3}$ g/hr. This agrees reasonably well with the estimated initial slope for the plot of carbon deposition over SP-1 graphite in Figure 1, that is $1.6 \times 10^{-2}$ g/hr.

The possible role which diffusional resistance plays in limiting the rate of carbon deposition can be estimated from the dimensionless criteria, $\phi \eta$. That is

$$\phi \eta = \left( \frac{R}{C_{\text{eff}} D_{\text{eff}}} \right) \frac{dW}{dt}$$

where $R$ is one-half the sample dimension in the direction where diffusion should be most rapid (taken as one-half the width dimension in this case), $D_{\text{eff}}$ is the effective diffusion coefficient, and $dW/dt$ is the rate of propylene cracking per unit external area. $D_{\text{eff}}$ is calculated for the counter-diffusion of propylene in hydrogen and is corrected for tortuosity (cos 45°) and sample porosity (0.362). At 700°C, $\phi \eta$ is estimated to equal $1.2 \times 10^{-4}$, which is significantly less than 0.1. The significance of $\phi \eta < 0.1$ is that under these circumstances the rate of a heterogeneous gas-solid reaction is chemically controlled (8). Therefore, we conclude that diffusional resistance to the initial rate of carbon infiltration of the SP-1 graphite artifacts was negligible.

The fact that the rate of carbon infiltration in the SP-1 graphite was greater than that in the cellulose carbon might be explained by three possibilities: i) SP-1 graphite has a higher inorganic impurity content which might catalyze the reaction, ii) SP-1 graphite has a higher ASA, and iii) infiltration of the cellulose artifact was diffusion limited. The first possibility is ruled out because of the very high purity of SP-1 graphite and close agreement in deposition rate with that predicted from studies using Graphon. Because of its smaller crystallite size and more defective structure, the ASA of cellulose carbon is expected to be higher than that of SP-1 graphite, suggesting that possibility (ii) should be ruled out. It is concluded that infiltration into the cellulose carbon artifacts was diffusion limited. The as-formed cellulose artifact exhibited molecular sieve properties in the uptake of $N_2$ and $CO_2$ (4); and, therefore, it is concluded that much of the internal surface area was located in voids between about 4.9 and 5.2A in size. Indeed, infiltration for only 3 hr sharply reduced the internal $N_2$ surface area in the cellulose artifact from 480 to 4 m$^2$g$^{-1}$ (4).

3.2 Effect of carbon infiltration on composite properties

As with the cellulose composites (4), changes in $\varepsilon$, Fig. 2 Dependence of flexural strength ( ), Young's flexural strength ( ), Young's modulus ( ), and electrical resistivity ( ) on apparent density of composites.

infiltration can be expressed by log-log plots, as seen in Figure 2. Experimental points for $\sigma$ and $E$ fit linear plots reasonably well, except at low levels of infiltration, where the artifacts are quite weak. Values of $\sigma$ also fall below the line following infiltration for 72 hr. This is attributed to noticeable build-up of deposited carbon at the exterior surface of the artifact, as previously discussed (4). Infiltration of the graphite artifacts produced much greater proportionate increases in $\sigma$ and $E$ than did infiltration of the cellulose artifacts, for similar extents of carbon pick-up. For the graphite artifact, following 72 hr of infiltration, $\sigma$ increased from about 4 to $500$ kg/cm$^2$; $E$ increased from about $2.5 \times 10^4$ to $4.0 \times 10^5$ kg/cm$^2$.

Conversely, infiltration of the graphite artifact produced a smaller decrease in $\rho$ than infiltration of the cellulose artifact. Following carbon deposition for 72 hr, $\rho$ decreased from 4.6 to 1.2 mohm-cm for the former and from 200 to 29 mohm-cm for the latter. The extent of the change in properties of these two artifacts upon infiltration is consistent with these facts.
First, the CVD carbon is less crystalline and more disordered than the natural graphite but more crystalline and less disordered than the cellulose carbon. Second, in the uninfilt rated cellulose artifact, some bonding has taken place between particles; it behaves like a consolidated body. By contrast, particles in the uninfilt rated graphite artifact are attracted by weak van der Waals forces; it behaves like an unconsolidated body.

Table 1 summarizes the effect of carbon infiltration on other important properties. Even though infiltration increases the hardness of the graphite artifact, its hardness is much less than that for the cellulose body. For example, infiltration for 72 hr only results in a Knoop hardness of about 45 kg/cm² compared to 186 kg/cm² for the uninfilt rated cellulose artifact. Wear resistance is poor, that is 1980 mg weight loss following 500 revolutions for the 72 hr infiltrated sample. By contrast, the infiltrated cellulose sample, underwent a weight loss of only 3.0 mg in 500 revolutions. This latter figure for wear resistance is superior to that found for stainless steel.

Effects of infiltration on the coefficient of thermal expansion (CTE) between 20 and 400°C are interesting. As expected because of crystallite alignment produced upon molding of the graphite artifacts, CTE parallel to the molding direction is larger than that perpendicular to the molding direction. However, since CTE (⊥) monotonically increases with increasing extent of infiltration whereas CTE (∥) goes through a pronounced maximum with increasing infiltration, the ratio of CTE’s (∥/⊥) goes through a pronounced minimum. Generally, the CTE of artifacts increases as the void volume existing between particles and crystallites (which can take up crystallite thermal expansion) decreases. However, in this case it is apparent that the deposition of sufficient CVD carbon, in effect, encapsulated individual graphite particles and retarded their normal expansion in the c-axis direction.

3.3 Effect of heat treatment on composite properties

As seen in Table 2, heat treatment (HT) decreased the densities of the infiltrated graphite artifacts. Artifact weight and planar dimensions did not change by more than 0.2% upon HT to 2400°C; on the other hand, sample thickness increased substantially more (up to 2.8%) and was responsible for most of the density decrease.

Heat treatment resulted in some degradation of mechanical properties. For example, for the 72 hr infiltrated sample fell from 500 to 400 kg/cm² following HT to 2400°C. Young’s modulus fell from 4.0×10⁶ kg/cm² to 3.0×10⁵ kg/cm². Such decreases are greater than would be expected on the basis of the decrease in artifact density as a result of HT and is attributed to some graphitization of the CVD carbon, as was previously reported (4).

As is seen in Table 3, HT drastically decreased the hardness of the artifacts. It also made wear resistance
too low to measure. HT also markedly lowered CTE's in both the parallel and perpendicular directions, as well as the maximum ratio of the CTE's found in these directions. These results can qualitatively be explained on the basis of crystallite growth in the CVD carbon and significant permanent expansion in the composites in the direction parallel to molding as a result of HT. Clearly there is no incentive to heat treat CVD infiltrated graphite artifacts on the basis of these results.

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REFERENCES