Role of nanocrystalline domain size on the electrochemical double-layer capacitance of high edge density carbon nanostructures

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Abstract
Nanostructured carbon materials, especially activated carbon, carbon nanotubes, and graphene, have been widely studied for supercapacitor applications. To maximize the efficacy of these materials for electrochemical energy storage, a detailed understanding of the relationship between the nanostructure of these materials and their performance as supercapacitors is required. A fundamental structural parameter obtained from the Raman spectra of these materials, the in-plane correlation length or nanocrystalline domain size, is found to correlate with the electrochemical capacitance, regardless of other morphological features. This correlation for a common nanostructural characteristic is believed to be the first result of its kind to span several distinct nanostructured carbon morphologies, including graphene–carbon nanotubes hybrid materials, and may allow more effective nanoscale engineering of supercapacitor electrode materials.

Introduction
Nanostructured carbon materials have been widely studied for applications in electrochemical energy storage, including activated carbon,[1–3] carbon nanotubes (CNTs),[4–7] and graphene.[8–11] Applications of such devices include hybrid structures in automotive energy storage,[12] flexible electronics,[13–15] and neural stimulation electrodes.[16] In order to optimize the performance of such devices, a clear understanding of the relationship between the physical properties of nanostructured carbon materials and electrochemical capacitance is required. As a nondestructive process, determination of nanocrystalline domain size using the Raman I_D/I_G ratio,[17–20] referred to interchangeably as graphitic cluster size or in-plane correlation length,[21] provides a facile and inexpensive method to engineer carbon nanostructures for energy storage applications. In this study, we present a relationship between this nanocrystalline domain size and the specific capacitance of various carbon nanostructures grown by plasma-enhanced chemical vapor deposition (PECVD) and measured by Raman spectroscopy. Previous works[22–26] have suggested a relationship between graphitic edge planes and specific capacitance for graphite, glassy carbon, highly ordered pyrolytic graphite, carbon nanofibers, multi-walled carbon nanotubes, and doped graphene. This work is believed to be the first to quantify the relationship between structural defects, largely contributed by few-layered graphene (FLG) edge planes, and the specific capacitance in terms of nanocrystalline domain size for graphene–CNT hybrid materials. It is also believed to be the first study of its kind across materials that span such a large range of domain sizes.

Materials and methods
The details of the PECVD growth system are described elsewhere,[27] as is the deposition process for growth of the graphenated carbon nanotubes (g-CNTs) used in this study.[28] In brief, nanostructures were grown in a 915 MHz microwave PECVD reactor using 50 Å Fe catalyst deposited on silicon. Prior to deposition, the substrates were heated to the desired deposition temperature in 100 sccm NH_3, followed by striking and stabilizing the plasma at 21 Torr and 2.15 kW magnetron input power. The substrates were pretreated for several minutes in the ammonia plasma to dewet the Fe catalyst film to form nanoparticles. The gas flow was subsequently changed to 150 sccm CH_4 and 50 sccm NH_3 for the desired deposition time. The graphene and CNT formed simultaneously during the growth process. Possible mechanisms for this formation have been discussed elsewhere.[28,29] In contrast to the high-temperature conditions of g-CNT growth, which results in a high foliate density but poor nanotube alignment, vertically aligned graphenated carbon nanotubes (VA g-CNTs) were achieved by reducing deposition temperature to 900 °C and extending deposition time to 30 min. Hierarchical structures of few-layered graphene and carbon nanotubes were created by further extending the deposition time, hereafter referred to as carbon nanosheets on carbon nanotubes or CNS-on-CNTs.
Raman spectroscopy was carried out using a Horiba Jobin Yvon LabRam ARAMIS spectrometer operating with a 633 nm HeNe laser. The $D$ and $G$ Raman modes were deconvolved as Lorentzian peaks and the $D'$ was modeled as a Gaussian peak, as suggested by Ferrari and Robertson [21] and Mennella et al. [30]. The ratio of the $D$ and $G$ band intensities were calculated from the intensities of the isolated peaks. The Raman spectra of the nanostructured materials and deconvolution of the peaks are shown in Fig. 1. Scanning electron micrographs of the various nanostructures examined in this study are shown in Fig. 2. Carbon nanostructures were imaged with a FEI XL30 SEM-FEG scanning electron microscope.

The electrochemical cell and sample preparation have been described in detail elsewhere. [31] A three-terminal cell (K0235 by Princeton Applied Research) was used with the nanostructure under study as the working electrode, a Pt mesh as the counter-electrode, and an Ag wire in 1 M tetrabutylammonium and 0.01 M AgNO3 in acetonitrile as the reference electrode. The electrolyte used was 1 M LiClO4 in acetonitrile. To perform electrochemical measurements, the nanostructured electrode was mounted on a piece of sheet metal using copper tape, and electrical contact was made by painting conductive silver epoxy on the nanostructure side. The nominal active area of the electrode was 1.43 cm2, defined by a PTFE gasket.

![Figure 1. Raman spectra and Lorentzian deconvolution of the first-order Raman-active modes in various carbon nanostructures grown by MPECVD. The peaks shown are the D, G, and D' in order of increasing wavenumber.](https://www.cambridge.org/core)
Results and discussion

Relationship between nanocrystalline domain size and nanostructure for high edge density carbon nanostructures

The classical relationship between the $I_D/I_G$ ratio and nanocrystalline domain size was developed by Tuinstra and Koenig\textsuperscript{[17]} as $L_a = C (I_D/I_G)^{-1}$, where $L_a$ is the in-plane correlation length and $C = 44$ Å for an excitation wavelength of $\lambda_L = 514.5$ nm. Matthews et al.\textsuperscript{[32]} appended this expression to include the wavelength dependence of the constant term, where $C(\lambda) = -126$ Å + 0.033$\lambda_L$ for visible wavelengths. Using this expression, the nanocrystalline domain size of each material was calculated (Fig. 2). A clear correlation is observed between the nanostructure observed in the SEM and the nanocrystalline domain size as estimated from the Raman spectra. The CNS nanostructure in Fig. 2(a) contains the highest density and the thinnest nanosheets of all the samples studied, indicating that the crystalline domain size is expected to be the smallest. TEM reported elsewhere\textsuperscript{[29]} has shown that such structures terminate in few-layered graphene, down to two or three layers. The CNS structure in Fig. 2(b) forms at the top of a CNT forest and the thickness of the nanosheets increases dramatically, while the density decreases from the samples in Fig. 2(a), indicating a larger spacing between the edges. As one observes the nanostructure evolution from Figs. 2(c) to 2(e) a decreasing density of graphene edges is observed on the g-CNT sidewalls until no foliates remain and the nanostructure is a standard vertically aligned CNT morphology. Samples shown in Figs. 2(d) and 2(e) are interesting in that both the SEM and the Raman-derived crystalline domain sizes show very minor differences. It is intriguing that the nanocrystalline domain sizes are so similar for these samples, which could imply that FLG may nucleate at the sites of previously existing defects in support of the proposed nucleation and growth model for graphene foliates.\textsuperscript{[29,33]}

Interpretation of the nanocrystalline domain size calculation

The data in Fig. 2 and the discussion above illustrate that the observed edge density follows trends in the Raman-derived crystalline domain size. Although the Tuinstra and Koenig expression originally attempted to link the $D$ peak intensity to phonon confinement, it has been discovered more recently that double resonance is the activation mechanism.\textsuperscript{[34]} Therefore, one may consider $L_a$ as an average interdefect spacing with the expectation that a larger defect density gives rise to a higher $D$ peak intensity, and thus a smaller $L_a$.\textsuperscript{[35]} Comparing this Raman expression with crystallite size estimates by x-ray
diffraction, the larger crystallites are given less weight in this case and the above expression underestimates $L_a$ due to the dominant effect of small crystallites.\cite{36–38} Bearing in mind the limitations of the above formulation of crystalline domain size, the trends detailed in this letter are nevertheless tooted in the existence and spatial frequency of defects in the materials. Thus, $L_a$ can be equally referred to as the mean spacing between defects, including the edge defects that are prominent in the materials of the current study.

**Relationship between nanocrystalline domain size and specific capacitance**

Data from a previously reported design of experiments (DOE) study of MPECVD growth\cite{29} was used to compare the $I_D/I_G$ ratio and specific capacitance of a variety of carbon nanostructures from CNTs to highly defective amorphous carbon (a-C), thus extending the range of nanostructured carbons beyond those depicted in Fig. 2. Briefly, the DOE process employed statistical methods to populate the process space of growth in the PECVD reactor, analyzing catalyst thickness, pretreatment time, process gas ratio, deposition temperature, and deposition time as factors and presence of CNTs, presence of CNSs, CNT diameter, $I_D/I_G$ ratio, and capacitance as responses.

A monotonically increasing trend was discovered between $I_D/I_G$ and capacitance, which prompted a comparison with $L_a$. As can be seen in Fig. 3, a linear relationship exists between capacitance as a function of nominal surface area and the nanocrystalline domain size calculated from the Raman data.

**Fig. 4.** CV curves (100 mV/s) of structures from the plot in Fig. 3. SEM micrographs are representative images of the nanostructures grown for the capacitance data set, including CNTs, g-CNTs, and a-C.

**Fig. 3.** The relationship between nanocrystalline domain size and specific capacitance. A linear fit converges with an $R^2$ value of 0.89 for various carbon thin films deposited according to the DOE procedure.\cite{29}

By roughly estimating and comparing the surface area of CNTs and CNSs, it is possible to eliminate the possibility that the trend in Fig. 3 arises primarily due to differences in the surface area. CNSs grown by PECVD can be modeled as semi-circular two-dimensional objects with a radius of $\sim$700 nm and inter-sheet spacing of $\sim$500 nm based on SEM images. For a substrate of 1 cm$^2$ nominal area, this corresponds to a surface area on the order of $10^{-4}$ m$^2$. Similarly for CNTs on a 1 cm$^2$ substrate, assuming a CNT diameter of $\sim$50 nm, height $\sim$10 μm, and density $1.5 \times 10^{10}$ CNTs/cm$^2$,\cite{31} the surface area is on the order of $10^{-2}$ m$^2$. Thus, CNTs have approximately two orders of magnitude higher surface area compared with CNSs under these experimental conditions. Since CNSs possess a smaller $L_a$ and higher specific capacitance than CNTs, it can therefore be concluded that defect density, which influences electronic structure and adsorption, more strongly
“foliates” of FLG grown directly on substrates or simultaneously with CNTs. A linear trend emerged between the measured specific capacitance and \( I_0 \), yielding a novel perspective for engineering carbon nanomaterials for energy storage applications.

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