TUNABLE STOICHIOMETRY OF BC\textsubscript{x}N\textsubscript{y} THIN FILMS THROUGH MULTITARGET PULSED LASER DEPOSITION MONITORED VIA IN SITU ELLIPSOMETRY (POSTPRINT)

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AFRL/RXAN

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Interim Report

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Tunable stoichiometry of $\text{BC}_x\text{N}_y$ thin films through multitarget pulsed laser deposition monitored via in situ ellipsometry

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Abstract. Pulsed laser deposition is an energetic deposition technique in which thin films are deposited when a laser pulse at 248-nm wavelength strikes a target and material is subsequently deposited onto a substrate with ideally the same stoichiometry. By synchronizing a high-speed mirror system with the pulsing of the laser, and using two separate targets, thin films having tunable stoichiometry have been deposited. Depositions were performed in a high vacuum environment to obtain as much kinetic energy as possible during growth. Typically, some 150 pulses at 300 mJ/pulse were required to deposit 1 nm. Island growth must occur on a per pulse basis since over 100 pulses are required to deposit a 1 nm film thickness. Films were deposited to ∼100-nm thickness, and in situ ellipsometry data were modeled to calculate thickness, $n$, and $k$. X-ray photoelectron spectroscopy (XPS), Raman spectroscopy, and atomic force microscopy (AFM) were all performed on each of the films. XPS demonstrated change in film composition with change in laser pulse ratio; ellipsometry displayed thickness from the model generated as well as the optical properties from 370 to 1690 nm. AFM thickness measurements were in agreement with independently modeled ellipsometry thickness values. © 2014 Society of Photo-Optical Instrumentation Engineers (SPIE) [DOI: 10.1117/1.JNP.8.083999]

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1 Introduction

Nanotechnology applications require material feature sizes or structures between 1 and 100 nm. Thermal materials, optical materials, metamaterials, and many others are all examples of areas which would greatly benefit from films having improved properties of interest such as thermal conductivity, resistivity, refractive index, or chemical stability, as well as required nano-sized features. A current significant problem is being able to obtain material samples having precise material composition uniformity on the nanometer scale. The novelty of this paper is that by synchronizing a pulsed laser with two different pulsed laser deposition (PLD) target materials and a programmable galvanometer, the stoichiometry of a film can be tuned at the subnanometer scale resulting in thin film growth having either constant or functionally graded composition. If a particular PLD target requires approximately 130 laser pulses to generate 1-nm thickness at a substrate, then clearly on a per pulse basis island growth occurs. Island growth results from a few laser pulses on a target, and by repeatedly applying a fixed number of pulses onto each PLD target an amorphous thin film can be generated at room temperature having a desired composition that is very well mixed at the subnanometer level throughout the thin film.
Each different composition results in different material properties and is expected to be defined by the rule of mixtures. PLD is an efficient vacuum deposition technique for depositing films from a target material, with compressive stresses ultimately defining the maximum thickness for each different film of interest. To illustrate this novel growth technique, $\text{BC}_x\text{N}_y$ was chosen as a materials system with PLD targets of BN and C used as basic components. Three amorphous thin films 100 nm or less in thickness were deposited using this novel technique to illustrate the resulting tunable amorphous materials properties, as well as films of pure BN and C.

$\text{BC}_x\text{N}_y$ thin films are of interest due to their potential to be more chemically stable and oxidation resistant at higher temperatures as compared with pure carbon based materials. BN has a wide band gap, $>5.0$ eV for different phases such as hexagonal or cubic, making it a good insulator and when in the cubic form is not as hard as diamond, but is more chemically stable. $\text{BC}_x\text{N}_y$ is also more chemically stable than carbon with both $\text{BC}_4\text{N}$ nanotubes stable to decomposition in air up to 900°C, as well as hexagonal BN (h-BN) to 1000°C. Oxidation resistance testing was beyond the scope of this effort, but motivated the choice of materials. For future efforts, the films will need to be deposited at high temperatures to obtain crystalline bonding, instead of strictly amorphous thin films that result from room temperature deposition. Novel photonic applications involving high temperature thermal management and index matching could be well suited to use $\text{BC}_x\text{N}_y$, such as optical metamaterials for high temperature application.

In situ spectroscopic ellipsometry was employed in order to characterize optical properties from 370 to 1690 nm. Ellipsometric analysis measures the change in the polarization of light after subsequent reflection through a medium. By analyzing the phase difference ($\Delta$) and magnitude ($\Psi$) of the refracted, elliptically polarized light, ellipsometry can provide information pertaining to several optical and physical properties of the film during growth. Due to the different optical behaviors of various classes of materials, raw ellipsometry data must be fit using an appropriate model. Successful fitting of experimental data can lead to an accurate estimation of the material’s refractive index ($n$), extinction coefficient ($k$), and thickness. The final thickness calculated from the models generated from ellipsometry data was verified with atomic force microscopy (AFM), X-ray photoelectron spectroscopy (XPS) and Raman spectroscopy were used to measure and compare the composition of the films. In this paper, four different desired thin film compositions were deposited and compared: (a) 100% BN, (b) 80% BN–20% C, (c) 67% BN–33% C, and (d) 50% BN–50% C. Addition of C to the film increased the extinction coefficient and caused the film to be much more opaque.

2 Experimental Setup

As shown in Fig. 1, a Lambda Physik LPX 220 excimer laser was used to ablate material from two different 50.8 mm rotating target materials of 6.3-mm thick. The target to substrate distance was 125 mm. The laser was pulsed in a synchronous fashion at $\sim$16 Hz by computer software which also constrained the laser beam path using a Cambridge Technology galvanometer system (two axes servo mirror system with capacitive coupled position feedback for high speed) to a uniform elliptical pattern on each target, and pulsed the required number of pulses sequentially.

![Fig. 1 Pulsed laser deposition (PLD) process utilizing multitarget galvanometer: PLD A–carbon and PLD B–BN.](image-url)
on each target. For each of four depositions 13,000 pulses at 300 mJ/pulse were used with different ratios of BN to C: 100% BN–0% C, 80% BN–20% C, 67% BN–33% C, and 50% BN–50% C. These ratios were achieved by placing the following sequence of pulses of BN and then C in a repeated fashion: 100–0, 16–4, 16–8, and 8–8. Each trigger event of the excimer laser generated a 20 ns output pulse at 300 mJ of 248 nm light having ∼60 J/cm² that created a plume of ablated material which was visible for <20 μs. Each laser pulse is 24 mm × 8 mm as measured at the laser, before focusing. The laser output was focused to strike the center of the target with a focused spot size of approximately 1.2 mm × 0.4 mm. The depositions were performed in high vacuum (HV) to be as energetic as possible, with initial base pressures ∼9.3 × 10⁻⁵ Pa.

Ellipsometry data were collected in situ using a Woollam M2000, and ellipsometry modeling was performed by using a multilayer model which consisted of a substrate model for silicon, a SiO₂ layer for the oxidation layer that is always initially present on silicon, and a second modeled layer representative of the data collected in situ during the thin film deposition. After a model is generated matching each of the three modeled layers, the film thickness can be calculated.

Initial alignment verification and data collection were performed for each sample and the native silicon oxide thickness was calculated to be ∼2.25 nm for each wafer. A model generated parameters defined to constantly fit the data generated during deposition, and the primary parameter of interest for modeling was the real-time thickness. The thickness measured for 100% BN thin film by ellipsometry was 100 nm at completion. Samples were analyzed with a Renishaw inVia Raman microscope, Staib XPS, and Asylum Research MFP3D AFM.

### 3 Results and Discussion

Ellipsometry data were collected for each of four deposition conditions and used to calculate n and k at five exemplary wavelengths as well as the final film thickness. Table 1 shows the numerical n and k results at one of the five wavelengths, 632 nm, for each of the experimental conditions. The ellipsometer model used contains three layers: the first layer is an infinitely thick silicon substrate, the second layer is the native silicon oxide layer, and the third layer is a dynamic model of the grown thin film. The silicon substrate and native oxide were measured prior to deposition process and were assumed to remain unchanged. The parameters shown in Table 1 were calculated using a Tauc-Lorentz model, due to its ability to accurately characterize the optical behavior of amorphous materials, including carbon and various nitrides.

Throughout the deposition process, raw psi (Ψ) and del (Δ) data were analyzed within the J.A. Woollam CompleteEase software package using a Cauchy model because of its simplicity and small mean squared error. Numerical components of the model were continuously optimized to approximate n and k across the wavelength range of the device. The thickness of the growth layer was both calculated and displayed every 8 s.

#### 3.1 XPS Characterization

XPS characterization was performed on each sample using a Staib XPS system consisting of an ion pump, x-ray source, and energy analyzer. Samples were loaded into an antechamber with a base pressure of 6.7 × 10⁻⁶ Pa or better. The samples were then transferred into the analysis

<table>
<thead>
<tr>
<th>PLD A–carbon (%)</th>
<th>PLD B–BN (%)</th>
<th>Thickness (nm)</th>
<th>n₆₃₂</th>
<th>k₆₃₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>100</td>
<td>100</td>
<td>2.02</td>
<td>0.006</td>
</tr>
<tr>
<td>20</td>
<td>80</td>
<td>91</td>
<td>2.04</td>
<td>0.056</td>
</tr>
<tr>
<td>33</td>
<td>67</td>
<td>88</td>
<td>2.14</td>
<td>0.145</td>
</tr>
<tr>
<td>50</td>
<td>50</td>
<td>85</td>
<td>2.36</td>
<td>0.304</td>
</tr>
</tbody>
</table>

Table 1 Ellipsometer summary of film growth on Si wafer with ∼2.25 nm oxide layers.
chamber which had a base pressure of $4.0 \times 10^{-8}$ Pa. XPS data were collected using a pass energy of 20 eV. All spectra were background corrected using a Shirley background subtraction. Component curves were fitted to each spectrum using a Levenberg–Marquardt algorithm that minimizes $\chi^2$, Fig. 2. Each component curve consists of a convolution of Gaussian and Lorentzian functions. Photoemission intensities were corrected to account for Scofield cross sections, kinetic energy, and analyzer transmission function. XPS data were collected with a standard Mg Ka (1253.6 eV) excitation source. The energy scale was calibrated against Au 4f7/2 and Cu 2p3/2 at the binding energies of 84.0 and 932.6 eV, respectively. The XPS resolution was determined by the FWHM of the Au 4f7/2 line, which was found to be 1.2 eV.

Table 2 shows the binding energy ($E_B$) for boron and carbon (B 1s and C 1s). The normalized concentration of each element is equal to the measured intensity divided by the instrument sensitivity factor for that element. The resulting percentage of the normalized concentration for each element is then the atomic percent. Table 2 shows the surface characterization measurements for the 50% C-50% BN sample. Oxygen contamination, O 1s peak, is shown as being some 28%; this is expected due to the fact that the samples have been exposed to air and were exposed to an ion beam for a brief time before sampling. The same measurement and analysis techniques were used on each of the four samples. Table 2 shows that the ratio of B 1s to C 1s is approximately slightly <1.0 and the ratio of B 1s to N 1s is just under 1.7.

Figure 2 shows plots of the binding energies for boron and carbon 1s peaks. The spectra show cross-contamination that occurred during growth, and are obvious for the first growth of 100% BN, i.e., a laser pulse of carbon would place small amounts of carbon onto the BN target, and pulse onto the BN target would contaminate the carbon target. This issue has been largely resolved with the addition of a shutter over the sample as well as a shield between the two PLD targets which provides for the ability to operate the ellipsometer even while the shutter is closed with PLD operating for the purposes of precleaning the targets, and has been shown to be required.

Figure 2 shows the composition summary of B, C, and N for each of the four films deposited and the corresponding XPS composition. The nitrogen content is lower than boron content for each sample, which can be explained by the fact that ionization and atomization of the BN target during the PLD process in vacuum results in a loss of nitrogen. Since the pressure was constant, HV, it was also expected that the same boron to nitrogen ratio will essentially be maintained.

![Fig. 2](http://nanophotonics.spiedigitallibrary.org/) X-ray photoelectron spectroscopy (XPS) showing boron opposite of carbon for desired BN: C ratio. Both sets of stacked spectra consist of raw data (black), fitted components (blue), and sum curves (red).
regardless of BN:C ratio of growth, as shown in Fig. 3, and the B:N ratio is \( \sim 1.5 \) for XPS characterized samples.

The desired growth setpoints that were also characterized by XPS were BN: 100%, BN: 80%, BN: 67%, and BN: 50%, but as is shown in Fig. 3, the actual boron to carbon ratios ended up being 4.33, 2.96, 1.44, and 0.901, respectively. The ratio of boron to nitrogen ended up being 1.472, 1.493, 1.480, and 1.678, respectively. This represents ratios on the order of \( \sim B_4CN \), \( \sim B_3CN \), as well as two samples with a \( \sim BCN \) ratio near unity, i.e., one B:C sample rich in boron and the other rich in carbon.

As was previously mentioned, the four samples grown without mechanical shielding between targets and characterized by XPS show approximately a 4:1 ratio of boron to carbon for the first sample which should not have had any significant amount of carbon. This error can be from either XPS measurement technique due to air exposure, or due to contamination occurring during the deposition process. Contamination can occur by the fact that if the targets are not physically separated using mechanical shielding, then each laser pulse will have material both deposited on the substrate, but also onto the other laser target and thereby skewing the desired deposition parameters and potentially creating particulates. To test this, six additional samples were grown to approximately 1 \( \mu \)m in thickness with mechanical shielding added between PLD targets to prevent cross-contamination. Energy dispersive x-ray characterization was performed with a Hitachi 3000 tabletop SEM with energy-dispersive X-ray spectroscopy (EDX) attachment. As shown in Fig. 3, the results were virtually identical within the error in terms of composition. However, the first XPS sample shown with a C:B ratio of \( \sim 0.25 \) demonstrates the first sample was contaminated as the EDX sample shows a carbon content of zero as expected, Fig. 3.

<table>
<thead>
<tr>
<th>Element</th>
<th>( E_B ) (eV)</th>
<th>Measurement intensity (a.u.)</th>
<th>Sensitivity factor</th>
<th>Normalized concentration</th>
<th>Quantity (atomic %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B 1s</td>
<td>191.0</td>
<td>1146.51</td>
<td>1.184</td>
<td>968.33</td>
<td>26.6</td>
</tr>
<tr>
<td>C 1s</td>
<td>285.0</td>
<td>2932.08</td>
<td>2.73</td>
<td>1074.02</td>
<td>29.6</td>
</tr>
<tr>
<td>N 1s</td>
<td>398.8</td>
<td>2187.55</td>
<td>3.789</td>
<td>577.34</td>
<td>15.9</td>
</tr>
<tr>
<td>O 1s</td>
<td>532.9</td>
<td>2406.50</td>
<td>2.37</td>
<td>1015.40</td>
<td>27.9</td>
</tr>
</tbody>
</table>

Fig. 3 XPS and EDX summary of B:N ratio (atomic %) and B:C ratio for each setpoint of carbon target pulses (%).

Table 2 X-ray photoelectron spectroscopy surface characterization of 50% carbon–50% BN sample. Measurement error is ±0.1 atomic %.

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PLD results in a direct stoichiometric transfer of material from target to substrate, but in vacuum also in all directions, hence the need to have a shield between targets to prevent cross-contamination. Adding a mechanical shield prevented cross-contamination issues.

When depositing from the BN target, there is a loss of nitrogen (Fig. 3, N:B ratio below 1) due to the fact that the deposition is being performed in vacuum, which is typical for multielement targets containing nitrogen or oxygen. One of many techniques to raise the resulting thin film nitrogen content is to perform depositions at a pressure of a few Pa of nitrogen. Figure 3 demonstrates that the composition, C:B ratio, can be adjusted with precision using PLD, but optimization process with regard to BCₙNₓ would need to be performed to obtain the desired nitrogen content as well. For single element targets, it is clear that subnanometer mixing can be accomplished using sequential laser pulses on each target.

3.2 Raman Characterization

Figure 4 shows four Raman spectra corresponding to each of the four samples and a carbon sample for the reference. A Renishaw inVia Raman microscope having a 514 nm laser excitation and 50× microscope objective was used for the measurements in a confocal mode providing a measurement depth of approximately 1 μm. The peaks shown in each of the spectra below 1000 cm⁻¹ are from the Si substrate and corresponding native oxide layer of SiO₂ (~2.25-nm thick). The carbon sample was deposited with the same conditions as the other four, but for a much shorter period of time yielding a film thickness of ~30 nm, and as such the Si and SiO₂ peaks are much larger than those observed in the other samples.

Raman spectra of carbon are expected to have a D peak at 1360 cm⁻¹ and a G peak at 1580 cm⁻¹, and boron nitride is expected to have Raman peaks at 1055 cm⁻¹ for cubic BN (c-BN) and 1364 cm⁻¹ for hexagonal BN (h-BN). As is shown in Fig. 4, the spectra above 1000 cm⁻¹ are as could be expected, essentially a D peak component, a much larger G peak component, and second harmonic of the D peak. By peak fitting the D and G components using two Gaussian functions, for pure carbon, the algorithm returns that there is dominantly graphitic carbon having a G peak centered at 1557 cm⁻¹ and a much smaller D peak centered at 1401 cm⁻¹, and for the pure BN sample, there is only h-BN with a broad peak centered around 1300 cm⁻¹. The D peak and h-BN peak overlap and the peak position of their summation shifts lower in wavenumber with the increasing BN content. Since the depositions were performed

![Fig. 4 Raman spectra of each of the five samples as deposited.](image-url)
without a heated substrate c-BN was not expected to be formed, which is confirmed by the absence of a 1055 cm\(^{-1}\) peak. The spectra in Fig. 4 for 80% BN-20% C deposited by PLD are equivalent to the Raman spectra presented in Ref. 13 from sputtering of B\(_4\)C in 10% N\(_2\) to obtain a film composition of 48% B, 12% C, and 40% N.\(^{13}\)

Beginning with the pure carbon sample Raman spectra, increasing the amount of BN causes the G peak to be reduced relative to the h-BN peak as the boron content is increased, and hence the h-BN peak of 1364 cm\(^{-1}\) is increased in intensity. The broad peak around 2700 cm\(^{-1}\) is the second harmonic of the \(\sim 1360\) cm\(^{-1}\) peak. The second harmonic of bulk h-BN (h-BN\(^{''}\)) is represented by a broad peak at 2765 cm\(^{-1}\), while the BN thin film second harmonic is \(\sim 2500\) cm\(^{-1}\), Fig. 4.\(^{14}\)

One noticeable detail is that with each of the four films having approximately the same thickness, the pure BN film has a Raman intensity at \(\sim 1300\) cm\(^{-1}\), which is much smaller than the substrate related to peak intensity at \(\sim 970\) cm\(^{-1}\). This lack of Raman response for pure BN would imply that either the film is not nearly as thick or that BN is perhaps much more transparent to 514 nm laser excitation than carbon. The film thickness was verified with ellipsometer modeling, AFM, and a KLA Tencor profilometer, so the lack of response must be from the actual material properties such as the absorption coefficient at 514 nm.

### 3.3 AFM Characterization

An MFP3D AFM (Asylum Research) was used to measure film thickness and surface root-mean-squared (RMS) roughness for each sample composition. The data shown in Fig. 5 were acquired in ac (tapping) mode by an Olympus ac160 silicon probe with a nominal tip radius of 9 ± 2 nm. Sample thickness was measured relative to the silicon substrate (disregarding an uneven edge formed by the removal of a mask adjacent to the deposited film). A measured thickness of 103.7 nm is indicated by the cross section in Fig. 5(a) for a BN-50:C-50 scan. An image analysis function in the MFP3D (Igor Pro-based) software was used to calculate the RMS roughness of unmodified topography data according to \(\sqrt{\frac{1}{n}} \sum \epsilon_i^2\), where \(n\) is the total number of \(z\) height values. Figure 5(b) shows a representative 10 \(\mu\)m scan of BN-80: C-20, which yielded a full image RMS value of 7.87 nm. RMS values were on the order of 7 to 11 nm for randomly sampled 10 \(\mu\)m areas. The horizontal lines in Fig. 5(b) were artifacts from the instrument of the tallest features in the scan line. The particulates captured in Fig. 5(b), displayed as white spots, were observed for each sample composition and could be attributed to the deposition of material onto the opposing PLD target with the ablation pulse. Each focused laser pulse was approximately 1.2 mm \(\times\) 0.4 mm, with each target material having

![Figure 5](http://nanophotonics.spiedigitallibrary.org/ on 03/27/2014 Terms of Use: http://spiedl.org/terms)
some 45-mm diameter active area of ablation, allowing buildup of particulates from many pulses onto each opposite target before an actual deposition pulse. In this manner, some degree of particulates are expected, but could be mitigated with the addition of a shield between the laser targets.\textsuperscript{15}

3.4 In Situ Ellipsometer Characterization

In situ ellipsometer data were collected for each of the four depositions, with the 50\% BN–50\% C and the 80\% BN–20\% C deposition dynamic data being shown in Fig. 6. Each deposition lasted approximately 15 min, with the deposition conditions being 300 mJ/laser pulse, and a 16 Hz pulse rate from the laser. During the deposition, the growth rate of the thin film thickness was linear as shown in Figs. 6(c) and 6(d). The value of $\Psi$ changed significantly during growth with each associated wavelength. The effect of different composition of materials and the difference in optical property of the material was significant, with the 80\% BN–20\% C having the largest change in the magnitude of $\Psi$. The change in $\Psi$ is most pronounced at 371.3-nm wavelength, Figs. 6(a) and 6(b).

The ellipsometer measures psi and del across the wavelength range of 370 to 1690 nm. CompleteEase software from J.A. Woollam provides for definition of optical layers, and in this case, a modeled layer for the Si wafer substrate, the SiO$_2$ native oxide layer, and a Cauchy model for each thin film of interest. Each optical layer is parameterized by the thickness, $n$ and $k$, and an appropriate model such as Tauc-Lorentz, Sellmeier, or in the simplest form Cauchy for transparent materials. The CompleEase software iteratively fits all measurements to all of the defined model layers to determine optimal model parameters. The Cauchy model, Eqs. (1) and (2), is an approximate function of the Sellmeier model in which a series expansion is used to compute $n(\lambda)$ and $k(\lambda)$:

\begin{align*}
\text{Wavelength (nm)} & \quad 371.3 \\
\text{606.7} & \quad 841.6 \\
\text{1171.2} & \quad 1678.2 \\
\end{align*}

\begin{align*}
\text{Psi} & \quad 30 \\
\text{15} & \quad 10 \\
\text{5} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Total thickness (Å)} & \quad 1000 \\
\text{500} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Time (min)} & \quad 15 \\
\text{10} & \quad 5 \\
\text{0} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Time (min)} & \quad 50 \\
\text{40} & \quad 30 \\
\text{25} & \quad 15 \\
\text{10} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Total thickness (Å)} & \quad 200 \\
\text{100} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Total thickness (Å)} & \quad 1000 \\
\text{500} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Time (min)} & \quad 25 \\
\text{20} & \quad 15 \\
\text{10} & \quad 5 \\
\text{0} & \quad 0 \\
\end{align*}

\begin{align*}
\text{Time (min)} & \quad 50 \\
\text{40} & \quad 30 \\
\text{25} & \quad 15 \\
\text{10} & \quad 0 \\
\end{align*}

\textbf{Fig. 6} Dynamic ellipsometer spectra collected during deposition: (a) BN-50:C-50 and (b) BN-80:C-20. In situ thickness modeled using ellipsometry (c) for BN-50:C-50 and (d) for BN-80:C-20.
In Eq. (1), the constant $A$ represents the approximate index of refraction and together with $B$ and $C$ define the index dispersion. Band edge is a parameter in nanometers that is set manually, whereas the $k_1$ and $k_2$ parameters determine the shape of the extinction coefficient dispersion. Once the best fit of the parameters $A$, $B$, $C$, $k_1$, and $k_2$ are obtained by CompleteEASE, where $n$ and $k$ and thickness are calculated.

From a Cauchy model fit of the psi and del in situ data of two deposition conditions, $n$ and $k$ were calculated at five representative wavelengths for the 50% BN–50% C and 80% BN–20% C cases, Fig. 7. The 50% BN case shows that particular film is very absorptive, especially at 514 nm where strong Raman spectra were collected, and the 80% BN case was not nearly as absorptive, Fig. 7. The extinction coefficient for pure BN, Fig. 7, was closest to zero at 514 nm as compared to the others, and was correspondingly the weakest Raman scattering thin film at 1364 cm$^{-1}$, Fig. 4.

Additionally, all samples $n$ and $k$ values are shown in Fig. 7: pure BN, 80% BN–20% C, 67% BN–33% C, 50% BN–50% C, and pure carbon. BN has the lowest value of index of refraction of the films grown at 2.0, and carbon the highest around 2.4.\textsuperscript{15} As demonstrated in Fig. 7, changing the stoichiometry by adding amounts of carbon will cause a shift in the actual optical properties of index of refraction and absorption coefficient, which can be desired for filters, optical lenses,

\begin{align}
n(\lambda) &= A + \frac{B}{\lambda^2} + \frac{C}{\lambda^4}, \quad (1) \\
k(\lambda) &= k_1 \cdot e^{k_2(E-E_B)} = k_1 \cdot e^{k_2[12400(\frac{1}{\text{Band edge}})]}. \quad (2)
\end{align}
coatings, metamaterials, or other applications. Increasing carbon concentrations, Table 3, show
that when compared at a specific wavelength such as 632 nm, the $n$ and $k$ values increase in an
increasing fashion. As the carbon concentration is linearly increased from 0 to some 41 atomic
%, the index of refraction increases from 2.03 to 2.36 at 632 nm and the value of $k$ increases from
$\sim 0.006$ to $\sim 0.304$. Typical values of index of refraction for h-BN are 1.8 and 3.3 for $B_4C$.

### Table 3 Changes of $n$ and $k$ according to carbon concentration at 632 nm.

<table>
<thead>
<tr>
<th>PLD A–carbon (%)</th>
<th>PLD B–BN (%)</th>
<th>Carbon (atomic %)</th>
<th>$n_{632}$</th>
<th>$k_{632}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>100</td>
<td>0</td>
<td>2.02</td>
<td>0.006</td>
</tr>
<tr>
<td>20</td>
<td>80</td>
<td>15</td>
<td>2.04</td>
<td>0.056</td>
</tr>
<tr>
<td>33</td>
<td>67</td>
<td>27</td>
<td>2.14</td>
<td>0.145</td>
</tr>
<tr>
<td>50</td>
<td>50</td>
<td>41</td>
<td>2.36</td>
<td>0.304</td>
</tr>
</tbody>
</table>

4 Conclusions

Nanotechnology research requires the ability to fabricate and test many different material samples
having defined, functionally graded, or static material properties or features at the 1 to 100 nm
scale. PLD is a highly energetic and efficient technique to create thin film materials, but typically
allows for only stoichiometric transfer of material from a single target material to substrate. As was
demonstrated, using more than one physical target and programming a desired number of laser
pulses onto each laser ablation target using a galvanometer resulted in a thin film material which
had tailorable subnanometer mixing throughout the thin film from two exemplary pure target
materials: BN and C. Typically, some 150 pulses at 300 mJ/pulse from the excimer laser resulted in
1 nm of material being deposited, demonstrating that island growth must occur on a per pulse basis.
By calibration of the number of laser pulses per target to XPS or EDX composition ratios, any
desired linear combination of B:C or other pure target materials can be achieved. By modeling the
carbon content and the resulting index of refraction, a film containing a desired index of refraction
(at 632 nm) between 2.02 and 2.36 could be deposited using this deposition technique. The amor-
phous nature and change of response with composition of each of the materials were verified
using Raman spectroscopy. AFM can be used to measure the thickness, grain size, and RMS
roughness of a sample, with typical roughness being shown to be on the order of 11 nm for samples
which were not biased nor heated during deposition. Optical ellipsometry in situ was shown to be
able to model the thickness of multiple layers, any combination of laser pulses, and all in real
time while the laser is operating growing a film. Optical ellipsometry allows for all wavelengths
from 370 to 1690 nm to be characterized simultaneously while depositing a thin film, and allows
for the implementation of process control to be able to deposit thin films having specific thick-
nesses or optical properties, as is required for optical metamaterials, thermal materials, and
nanotechnology applications.

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and implementation and optics train for excimer. P.T.M. aided in chamber design. N.R.M. was
involved in experimental apparatus implementation. L.S. conducted ellipsometric analysis of
samples. T.C.B. performed XPS. J.L.R. performed AFM. M.A.L. researched BCN details.
J.G.J. developed the overall apparatus design, authored deposition software, developed synchro-
nization of the laser pulses to the optical mirror system, performed thin film growth with in situ
ellipsometry measurements, and Raman and EDX characterization.
References


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