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AlN Film Thickness Effect on Photoluminescence Properties of AlN/Carbon Nanotubes Shell/Core Nanostructures for Deep Ultra-Violet Optoelectronic Devices

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ABSTRACT

Aluminum nitride (AlN) nanostructures are very attractive in various optoelectronic applications such as deep ultraviolet light emitting devices. The fabrication of these AlN nanostructures with good crystalline quality and compatibility in line with other microfabrication processes has significant importance for practical applications. AlN films of different thickness values were deposited via DC reactive magnetron sputtering over vertically aligned multiwalled carbon nanotube (CNTs) arrays to obtain AlN/CNTs vertically-aligned shell/core nanostructure assembly. Such hybrid nanostructures were characterized using scanning electron microscope, transmission electron microscope, X-ray diffraction, Raman spectroscopy and time-resolved photoluminescence spectroscope (TR-PL) techniques. The results indicated that AlN/CNTs have a nanorods structure morphology with good AlN crystalline quality. The PL measurements revealed a maximum increase in the luminescent intensity of the exciton band in case of AlN/CNTs with 600 nm thick AlN layer, which is

many orders of magnitude higher than that of AlN film produced over silicon substrate. It is anticipated that synergistic effects of CNTs and AlN through an increase in the specific surface area and oxygen-induced defects cause enhancement in the photoluminescence properties, making these hybrid nanostructures a promising candidate for optoelectronic applications.

Keywords: Aluminum nitride, carbon nanotubes, reactive magnetron sputtering, hybrid nanostructures, Photoluminescence

1. Introduction

Aluminium nitride (AIN) is a material with remarkable properties including large optical band gap value (E_g 5.9-6.2eV), high hardness and good thermal conductivity [1]. It can be used in several electronic and optoelectronic devices, ultraviolet detector, light emitting diodes and thermal interface materials, to name a few [2, 3]. The large E_g value for AlN permits its incorporation into deep UV LEDs [4-6] and extreme UV detectors [7], the devices that can be employed for probing intrinsic florescence in protein, equipment/personnel decontamination, photocatalysis and astronomy [1, 8]. In order to obtain AlN with efficient UV emission, a fabrication process that will produce AlN with high structural quality, is desired. The UV emission of AlN can be enhanced by increasing its surface area, that in turn, may be possible in case of AlN nanostructures with morphologies as diverse as nanowires, nanotubes, nanorods, nanotips, hierarchical comb-like structures, pine like structures or nanobelts [9]. In most cases, however, the synthesis techniques carry disadvantages of high processing

temperature (800-1200 °C) [10, 11] and poor structural quality of the resulting AlN films with deterioration in band gap emission besides emission in the visible spectral region [12-14].

In this work, vertically aligned carbon nanotubes (CNTs) were grown over silicon substrates followed by sputter deposition of AlN films for any enhancement in the surface area and photoluminescence properties. Similar strategy was adopted by our group in case of zinc oxide (ZnO) [15] and titanium nitride (TiN) films produced over CNTs arrays [16]. Herein, a fabrication route compatible with micro-fabrication techniques was developed via dc reactive magnetron sputtering (dcMS) of AlN films over vertically-aligned CNTs arrays to obtain AlN/CNT shell/core nanostructures at low temperature. In addition, the ability to grow vertically aligned CNTs at relatively low temperature of 420 °C [17] offers the advantage of low cost processing.

In contrast with AlN films over polished silicon substrate, the AlN/CNT hybrid nanostructures exhibited band gap emission intensity of much larger magnitude in the deep UV spectral region with relatively little contribution from defect emission. Such shell/core hybrid nanostructures offer strong potential for used in deep UV light emitting devices and other optoelectronic applications.

2. Experimental details

2.1. Plasma enhanced chemical vapour deposition growth of vertically aligned CNT arrays

Vertically aligned carbon nanotubes (CNTs) arrays were grown over silicon substrates using distributed electron-cyclotron-resonance (ECR) plasma enhanced chemical vapor deposition (PECVD) process. The details of the experimental setup are described elsewhere [18]. A 10

nm thick nickel (Ni) film was obtained via DC-plasma sputter deposition and then transferred to the PECVD reactor. Upon heating to 600 °C, the film transformed into Ni nanoparticles, \sim 50 nm in diameter, that subsequently catalyzed CNT growth in a PECVD reactor using ECR plasma and acetylene to ammonia C_2H_2 :NH₃ ratio of 1:4, other parameters being chamber pressure of 0.2 Pa, 125 W microwave power and synthesis time of 1 h.

2.2. DC plasma reactive sputtering of AlN films

Over both Si(100) substrates and CNTs arrays grown over Si(100), AlN films were deposited through DC reactive magnetron sputtering technique using aluminum target (50 mm in diameter, 99.999% purity), Argon/N₂ gas mixture atmosphere and a target-substrate distance of 30 mm. An unbalanced magnetron sputtering system powered by a 150 W DC power supply was used. Before deposition, the target was cleaned using argon gas discharge plasma for 15 min followed by a pre-sputtering step with a shutter shielding the sample under the same conditions as those during subsequent film deposition. Once residual pressure of <6×10⁻⁶ Pa was achieved, the chamber pressure was increased to 0.4 Pa and was kept at that level during AlN deposition. The total gas flow rate during deposition was maintained constant at 40 sccm with N₂ flow rate of 14 sccm. The substrates remained at a floating potential and were neither additionally heated nor cooled during the deposition. In order to check the effect of the thickness on the photoluminescence (PL) signal, different samples were prepared by varying the deposited AlN film thickness from 50 to 1500 nm.

2.3. Characterizations

The scanning electron microscope (SEM) studies were performed for morphology and thickness of the deposited hybrid nanostructures using a JEOL JSM 7600 apparatus. The images were recorded at an accelerating voltage of 5 kV. The morphology was also

investigated using field-emission gun scanning electron microscope (FEG-SEM) (JEOL; JSM7600F1) and field-emission gun transmission electron microscope (FEG-TEM; Hitachi HF2000) operating at 200 kV. For TEM observation, the samples were micro-delaminated with a diamond tip and directly transferred onto the TEM copper grids. The AlN/CNTs hybrid structures were also characterized using X-ray diffraction (XRD) apparatus (D500 MOXTEK) with monochromatic CuK α radiation (λ = 0.154 nm) in the θ –2 θ Bragg Brentano configuration. The room temperature Raman spectra were recorded with a Jobin Yvon micro Raman spectrometer (T64000) at 240 nm wavelength and 100 mW power. The time-resolved PL measurements were performed on a system consisting of a monochromator (1.3 m) and a frequency quadrupled 100 fs Ti:Saphire laser with excitation photon energy set at about 6.28 eV. A single photon counting detection system together with a micro-channel-plate photomultiplier tube (with 20 ps time resolution) or a streak camera (2 ps time resolution) was used to record time-resolved spectra.

3. Results and discussion

The AlN films with thickness values in the range of 50 to 1500 nm thick were produced over the CNT arrays, as shown in surface and cross-section SEM view of the AlN/CNT hybrid structures in Fig. 1. The cross-section view revealed vertically-aligned shell/core configuration of the AlN/CNT nanorod assembly with varying degrees of porosity and AlN film thickness values from 50 to 1500 nm (Fig1. b, d, f, h). Due to the line-of-sight deposition, the aligned CNT arrays undergo encapsulation of the individual nanotubes with AlN film at and near their ends, thus resulting in a convex shape whose size increases with an increase in the thickness of the deposited layer. While the surface microstructures, presented in Fig. 1(a, c, e, g) indicate the AlN/CNT grain size as well as the film density at the surface or near-surface area, the cross-section view reveals the degree of CNT encapsulation and

filling of the inter-nanotubes spacing by the incoming flux of reactively-sputtered AlN. Upon increasing the deposition time, the grain size, as estimated from the surface microstructure, increases from ~100 nm to ~1 µm with an associated decrease in the degree of surface porosity. Examination of the cross-section morphology indicated an increase in the degree of CNT encapsulation, both axially as well as radially, for longer deposition times as manifested by AlN deposition over greater length segments of the CNT forest and larger size of the AlN/CNT hybrid structure at and near the top of CNT array. Since the average length of the core/shell vertically-aligned nanostructure array (3.4 µm) is slightly higher than that for pure CNT array (2.2 µm) in case of 1500 nm thick AlN film, it is deduced that (i) AlN deposition does not completely fill the spacing between nanotubes in the aligned CNT array and (ii) beyond a certain deposition time, the top surface of the CNT array is completely buried underneath AlN thus resulting in a AlN-rich top layer. From SEM studies, it seems that AlN films with thickness values of 50, 400 and 600 nm maintain high surface area of the AlN/CNT hybrid structure. For these thickness values, the AlN deposition apparently induced an inverse pyramid- or conical shaped columnar morphology with a decrease in the degree of CNT encapsulation from its upper end to the base. The AlN attachment primarily to the upper end of the aligned nanotubes forest may result in higher luminescence intensity of band gap emission in such AlN/CNT shell/core nanostructures. In the case of 1500 nm thick AlN film, a continuous, compact layer albeit with less surface area is, likely to have formed over the top of CNT array. All these aspects will probably affect the band gap emission of AlN deposit.

The XRD patterns of the AlN films produced over vertically-aligned CNT arrays are presented in Fig. 2. The hybrid structures exhibit two different crystallographic orientations indexed as c-axis (002) and (101) planes, both of which may be associated with hexagonal Wurtzite structure of AlN (JCPDS: No. 08-0262) [19]. It was noticed that upon increasing the

AlN film thickness from 400 to 1500 nm, while the intensity of the (101) diffraction peak remains almost unchanged, there is a progressive increase in the peak intensity corresponding to the (002) plane.

One of the AlN@CNT core/shell hybrid nanostructure sample with AlN film thickness of 600 nm was examined under TEM, as shown in Fig. 3. From low magnification image of the region near the tip of AlN/CNT array (Fig. 3a), complete encapsulation of the nanotubes is noticed. Although, a cursory look at the HRTEM micorstructure of the tip area suggests layered petal-like morphology, a closer look reveals rodlike-morphology in the form of bundles (Fig. 3b). The dense, columnar morphology with c-axis (002)-preferred crystallographic orientation, produced at low deposition pressure (0.3 Pa) and ~35% N₂ gas concentration, is promoted by an increased in the mobility of adatoms and is in agreement with earlier studies on reactively sputtered AlN films [20, 21]. The selected area electron diffraction pattern (SAED) of a relatively large area of an individual AlN@CNT shell/core structure (Fig. 3c) exhibits diffraction rings characteristic of the (101) and (002) crystallographic orientations of the AlN lattice. The presence of the sharp, diffraction spots in a circular arrangement points towards polycrystalline nature of AlN with relatively large grains [22]. The SAED of a smaller region confirms (Fig. 3d) hexagonal structure of the AlN coating with a (002) preferential orientation.

Fig.4 shows the UV Raman spectrum in the range 200–1200 cm⁻¹ for AlN/CNT shell/core hybrid nanostructures with different AlN film thickness values. For 50 nm thick film, no spectroscopy mode was observed. All the three Raman active modes namely A₁, E₁ and E₂, are noticed to be present in the Raman spectra for AlN/CNT samples with AlN film thickness of 400, 600 and 1500 nm. Both A₁ and E₁ are polar modes with each of them having longitudinal (LO) and transverse (TO) modes arising from the interaction of the vibrations with the long-range Coulomb field [23]. From the Raman spectra, five (5) different peaks

positioned at frequencies of 248, 610, 657, 669 and 910 cm⁻¹ are seen. The weak phonon peak at 248 cm⁻¹ corresponds to the E_2 (low) (also denoted as E_2^L) mode, whereas the peak at 657 cm⁻¹ represents the E_2 (high) (also called E_2^H) mode. Similarly, while the phonon mode A_1 (TO) can be clearly identified at 610 cm⁻¹, the A_1 (LO) mode maybe present at 890 cm⁻¹ as a shoulder peak for the 910 cm⁻¹ peak. The peaks located at 669 and 910 cm⁻¹ correspond to the E_1 (TO) and E_1 (LO) phonon modes, respectively [23, 24]. Predominance of the active Raman modes depends on the direction of phonon propagation, whether parallel or perpendicular to the c-axis of the crystal [25]. Such discussion is beyond the scope of this research and the Raman studies presented herein confirm stoichiometric AlN film formation from dc reactive sputtering.

AlN has a relatively small c/a ratio of 1.601 and a band gap energy value of 6.113 eV including free exciton binding energy of 80 meV [26]. In bulk AlN ceramics, the excitation wavelength of 195 nm causes appearance of different bands in the emission spectrum at spectral positions of 300–500 nm, 600 nm, 700 nm and 1000 nm with respective characteristic features of a broad UV-blue band (300-500 nm), a narrower band and minor bands at 700 and 1000 nm. The broad complex UV-blue band is composed of two subbands at 400 and 480 nm, each of which has its own excitation spectrum. While the 400 nm emission band causes appearance of excitation band centred at 240-250 nm, the 480 nm emission band gives rise to an excitation band at 280-290 nm [27]. Both types of emission bands are also excited at ~210 nm (5.8-5.9 eV). The exciton generation is attributed to recombination of two defect centers, such as a complex donor defect formed via substitution of nitrogen ion O_N by an oxygen ion in a regular site and a neighboring aluminum vacancy v_{Al} (O_N - v_{Al})) and a closely situated oxygen ion (an acceptor O_N). In some cases, implantation of oxygen ions may cause damage to the AlN surface layers thus increasing oxygen concentration, and in turn, generating defect

centers [28]. At room temperature, therefore, the presence of impurities and defects cause radiative recombination energies at 5.69, 5.85 and 6.09 eV to represent bound exciton recombinations in AlN [29]. The room temperature photoluminescence (PL) spectra of different AlN/CNT vertically-aligned shell/core nanostructures were obtained, as presented graphically in Fig. 5. These were compared with the PL spectrum of the 1500-nm thick AlN film produced over silicon substrate (called ref. AlN). The choice for the ref. AlN sample was made owing to its high thickness value and superior structural quality. In case of ref. AlN, a weak exciton peak at ~5.8 eV is noticed, the presence of which may be attributed to the band gap emission at 400 and 480 nm. The reasons for such low luminescent intensity maybe high structural quality of the film and low defect density at the film surface due to very small amounts of residual oxygen in the chamber during film deposition.

When AlN is reactively sputter deposited over aligned CNT array as in the case of different AlN/CNT shell/core nanostructures, an intense, broad and asymmetric peak is seen at 5.85 eV in the PL spectra. The peak intensity increases with an increase in the AlN film thickness reaching a maximum value for 600 nm thick AlN film over the CNT array. While the intensity for this sample is much greater than that for 1500 nm thick AlN film over silicon, The PL intensity, however, drops once the AlN layer thickness over CNT is 1500 nm. Nevertheless, it still remains more intense than the one obtained for the ref AlN film. Furthermore, the relatively sharp deep-UV emission peak is an indication of good structural quality of the AlN film deposited over vertically-aligned CNT array. The increase of the band gap emission intensity of CNT/AlN compared to the ref AlN film maybe attributed to the increase in the surface-to-volume ratio of AlN/CNT shell/core nanostructures that exhibit a nanorod-like structure with a convex end i.e. nanoscale conical shape at the top of the vertically-aligned CNT forest. In fact, the peak intensity increases with AlN thickness up to 600 nm and then decreases for higher thicknesses due to the decrease in the surface area (Fig.

1). The band gap emission intensities of 50 and 400 nm thick AlN@CNT are lower than that of 600 nm AlN, even though the surface area values for the 50 and 400 nm thick CNT/AlN nanostructures seem to be higher (Fig. 1). This may be caused by an improvement in the crystalline quality of the AlN film upon a thickness increase from 400 to 600 nm [30, 31]. The maximization of the PL intensity upon varying AlN film thickness value maybe attributed to the combined effect of an increase in the specific surface area of the hybrid nanostructures as well as enhancement in the oxygen concentration with subsequently higher defect density.

4. Conclusions

The dc plasma reactive magnetron sputtering technique can be employed to obtain AIN/CNT shell/ core nanostructures over vertically aligned multiwalled carbon nanotubes. The AIN films produced at low chamber pressure and 35% N₂ gas flow are near-stoichiometric with predominantly c-axis (002) orientation and have dense, columnar morphology comprising of bundled nanorods configuration. While an increase in the AIN film thickness from of 50 to 600 nm over aligned CNT array promotes photoluminescence in the exciton spectrum owing to greater specific surface area and enhanced surface defect density, the CNT array is completed embedded by the AIN film upon further increase in thickness to 1500 nm, with subsequent deterioration in the PL properties. As compared to pure AIN film, the AIN/CNT hybrid shell/core assembly with 600 nm thick AIN film exhibits maximum photoluminescence intensity. The enhancement in the PL intensity of the AIN band gap emission in the deep UV regime presumably arises from the trade-off between AIN structural quality and changes in the defect density owing to any change in the specific surface area. The significant enhancement in the PL intensity of the AIN films upon interfacing with CNT

highlights the importance and potential utility of such shell/core nanostructure assemblies in multifunctional optoelectronic devices.

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List of Figures

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- **Fig. 3:** (a) TEM micrograph of the AlN/CNT sample with 600-nm AlN film over vertically-aligned CNT array, (b) High-resolution microstructure of the hybrid assembly near tip area and (c, d) SAED patterns of large and small areas of the microstructure presented in (b).
- **Fig.4:** Raman spectra of the AlN/CNT shell/core hybrid nanostructures with different thickness values for the AlN films.

Fig.5: PL spectra of the AlN/CNT shell/core nanostructures with different AlN film thickness and pure AlN films produced over silicon substrate.



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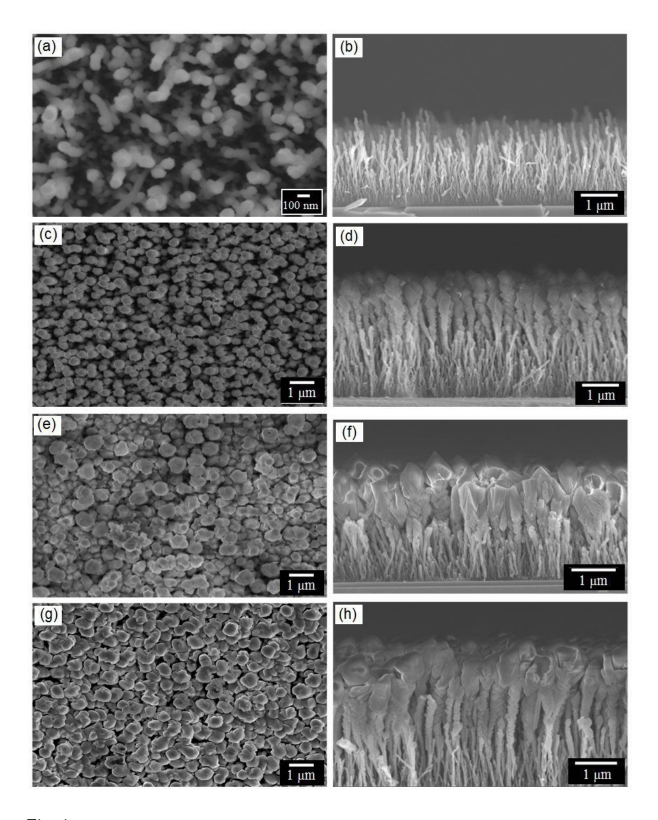


Fig. 1

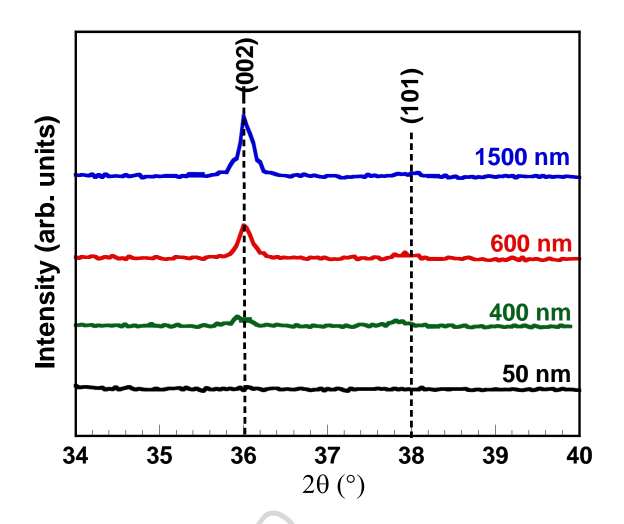


Fig. 2

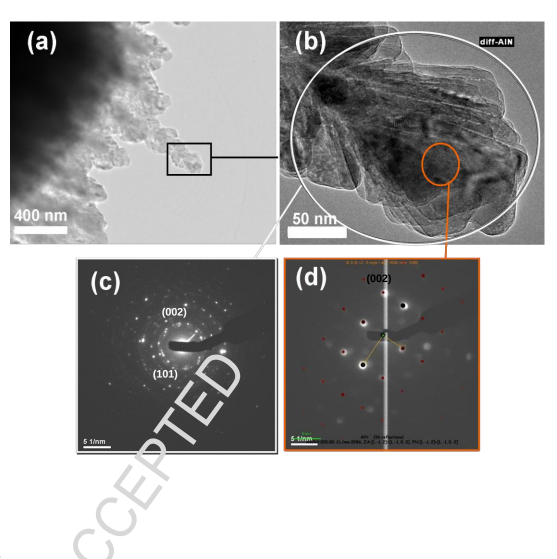


Fig. 3

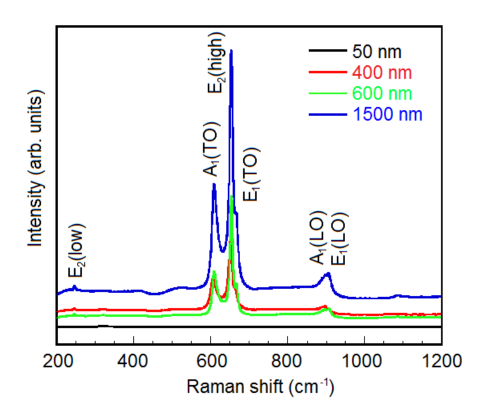
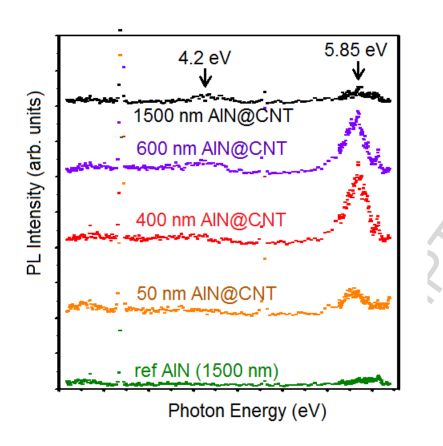


Fig. 4







Highlights

- AlN films with 50 to 1500 nm thickness were produced over vertically aligned CNTs.
- AlN/CNT shell/core assembly exhibits enhancement in PL properties.
- The 5.85 eV exciton band intensity maximizes for 600 nm think AlN film over CNTs.
- The AlN/CNT shell/core nanoassembly can be used for deep UV optoelectronic devices.

