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Plasmonic modification of electron-longitudinal optical phonon coupling in Ag-nanoparticle embedded InGaN/GaN quantum wells

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ABSTRACT

Surface plasmon enhanced GaN and InGaN quantum wells (QWs) show promise for use as room-temperature light emitters. The effectiveness of the plasmon enhancement, however, is limited by the strong electron and hole longitudinal optical (LO) phonon coupling found in the III-V nitrides. The electron-phonon coupling within semiconductor QWs has been modified using Ag nanoparticles embedded with the QWs. Direct evidence is provided for this change via confocal Raman spectroscopy of the samples. This evidence is augmented by Angle-dependent photoluminescence (ADPL) experiments which show the alteration of the e-ph coupling strength through measurement of the emitted phonon replicas. Together these demonstrate a direct modification of carrier-phonon interactions within the system, opening up the possibility of controlling the coupling strength to produce high-efficiency room-temperature light emitters.

PACS Codes: 78.55.Cr; 78.67.De; 71.45.Gm
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The optical properties of confined excitons or carriers in III-nitrides alloys and quantum well based devices such as UV-Visible lasers, light emitting diodes (LEDs), solar blind UV detectors and high power optoelectronic devices is influenced by longitudinal phonon scattering. Due to the large LO phonon energy of InN (86 meV) and GaN (91 meV), the strength of the interaction of the carriers or excitons with the phonons effects the interband and intersubband optical transitions in these structures. There has been increased interest in realizing white light emitters and photovoltaic devices using nitride semiconductors that has a bandgap spanning from the UV to the near-infrared wavelength regime. GaN and its tertiary compounds has been utilized for realizing efficient UV [1] and visible LEDs [2]. Radiative recombination of excitons in semiconductors is accompanied by phonon replicas which influence the absorption and emission efficiency of the nitride based devices [3]. The large LO phonon energy should minimize non-radiative intersubband transitions due to LO phonon emission [3]. The strength of exciton-phonon coupling also provides useful information about an indirect Auger recombination process due to scattering, which plays a major role in efficiency droop in nitride based emitters [4].

Surface plasmon induced enhancement of light emission in nitrides is a potentially attractive way of increasing the efficiency of nitride LEDs due to the enhancement of the radiative recombination rate. The coupling between LO-phonon and exciton which has been studied extensively using photoluminescence (PL) measurements on III-nitrides has not yet been investigated in nitride-based plasmonic light emitters. Long range plasmons are normally dissipative due to the resonant coupling of the excitons or free carriers to the surface plasmon modes. A study of the exciton-phonon coupling mechanism in plasmonic light emitters and its role in the light emission efficiency can results in designing more efficient room-temperature nitride based light emitters. Long range surface plasmon mediated radiative recombination at room temperature is generally dominated by non-radiative recombination process yielding very low internal quantum efficiencies. One possible means of bypassing this issue would be to directly modify the electron-phonon coupling strength by using localized surface plasmon coupled to the emitter within the semiconductor. Previous work has demonstrated the possibility of producing coupled plasmon-exciton [5] and plasmon-phonon [6] modes. The exciton LO phonon coupling is
also influenced by the geometry of the structure due to the modification of the electromagnetic coupling in micro-cavities and can modify the radiative recombination process in nanostructures. There has, however, been no investigation to-date of whether these coupled modes can modify the electron-phonon and hole-phonon coupling strength interactions within the semiconductor.

We present here evidence of modification of this coupling strength in InGaN/GaN QWs by the introduction of resonant plasmons to the system via metallic nanoparticles (NPs). The ability to induce localized surface plasmons within a 2D quantum confined structure by infiltrating metal nanoparticles with InGaN QWs enables the modification of electron and hole-phonon interactions at the nanoscale limit. We provide direct experimental evidence of this change via observation of a collection-geometry forbidden Raman $A_1$[TO] mode. We also demonstrate that the introduction of metal NPs to the system also produces a significant change in the angle-dependence of the electron/hole-LO phonon (e/h-LOP) coupling strength. Finally, we briefly discuss the possible origins of this change in the e/h-LOP interactions, and the implications of the possibility of external modification of the coupling strength.

Previously it has been demonstrated that it is possible to incorporate metallic nanoparticles in the inverted hexagonal pits (IHPs) present in InGaN/GaN multi-quantum well (MQW) structures [7]. These IHPs arise from threading dislocations (TDs) propagating from the GaN/Sapphire interface [8-11], and provide a simple and elegant location for placing metallic NPs within the QWs without affecting their structure [7]. In this work, we have used a MQW structure which was grown via metal-organic chemical vapor deposition on sapphire substrate, and consists of 14-periods of 2.5 nm InGaN QWs separated by 7.5 nm GaN buffers. Two samples with this structure were grown, and one was infiltrated with Ag NPs with an average radius of 17 nm, while the other was left unaltered for use as a reference. The top down approach of infiltration of metal nanoparticles within quantum wells eliminates the possibility of any strain induced due to structural composition of the InGaN/GaN QWs. Characterization of the sample structure was performed via scanning electron microscopy (SEM) and atomic force microscopy (AFM). Figure 1(a) shows SEM of the sample structure with NPs embedded within the IHPs, along with the PL of the reference and Ag NP samples.
Raman excitation was provided by the 514 nm line of an Ar+ continuous-wave laser, and excitation and collection were along the c-axis of the sample with incident and scattered polarizations matched \([c \quad (a-a) \quad \bar{c}]\). A Horiba-T64000 UV-Visible Raman spectrometer was used for this experiment at room temperature. Figure 2 shows two important features resulting from the addition of the NPs. First, the \(E_2\)(high) phonon peak (569.9 cm\(^{-1}\)) is shifted ~1.1 cm\(^{-1}\) to the left of the reference peak. We may exclude strain modification as a source of this observed Raman shift due to the top down approach for infiltration of the nanoparticles into the MQWs. Secondly, a second peak, corresponding to the \(A_1\)(TO) phonon, appears at 527.9 cm\(^{-1}\). The second feature is particularly interesting, as the \(A_1\)(TO) phonon should not be observable using a \([c \quad (a-a) \quad \bar{c}]\) excitation/collection geometry, but instead under \([a \quad (c-c) \quad \bar{a}]\). [12] This suggests that the normal symmetries of e-ph coupling in GaN have been broken in the presence of the Ag NPs.

Taken together these constitute significant experimental evidence of a change in the e/h-ph interactions within the system due to the presence of the Ag NPs.

The samples were also studied using angle-dependent photoluminescence. The ADPL was performed using excitation from the 325 nm line of a HeCd laser, incident at -55 degrees from the surface normal. Photoluminescence was then captured in five degree increments from 0 degrees to 75 degrees using a fiber-optic light guide and iris. The guide and iris were mounted on an arm attached to a goniometer centered beneath the sample. The normalized PL spectra at normal incidence are shown in Figure 1(b). The measured spectra were then fitted using multiple Gaussians to extract the intensities of the phonon replicas using the model described in Ref [13]. Due to specular reflection of the excitation source, the fitting was not possible at 55 and 60 degrees and hence the fitting results for those angles were discarded.

Figure 3 shows the angle-dependent intensity of the QW emission for the zero-phonon line, as well as the first and second phonon replicas. The intensities are normalized such that the sum of the intensities of the three components is unity at all angles. It is immediately obvious that there is a significant deviation in the intensities of the three components between the reference and Ag NP sample. This implies a modification of the e/h-LO phonon coupling within the InGaN/GaN quantum wells induced by the metal nanoparticles. The primary changes in the relative intensities occur for emission near normal and for emission around 45 degrees. In the former case, the intensity of the first phonon replica (1-PR) is decreased for the Ag NP sample relative
to the reference. At 45 degrees, on the other hand, the 1-PR emission is strongly increased relative to the reference sample.

Figure 4 summarizes these results in terms of the Huang-Rhys parameters \((S_0, S_1)\), which are direct measures of the strength of e/h-LOP coupling in the system. The value of \(S_n\) has been determined empirically using the definition \(S_n = (n+1)^{1/2} I_{n+1}/I_n\) [14]. We have also measured the energy of the primary emission and the phonon replicas. As discussed previously [3,4], in InGaN the phonon replicas are separated from the main peak by the energy of one or more LO phonons \((\hbar \omega_{\text{LO}} = 92 \text{ meV})\). Since the distribution of phonons is not Gaussian, the peak of the first and second phonon replicas are offset from their expected positions by 1/2 and 3/2 kT respectively, where kT \(~26 \text{ meV at room temperature}\). In Figure 5 we show the energies of the three peaks measured as meV deviations from the reference emission energy of 2.984 eV. It is interesting to note that the first phonon replica energies line up well with the corrected phonon replica energy whereas the second phonon replica lines up more closely with the uncorrected values. The surface plasmon energy of the Ag nanoparticles overlaps with the bandgap energy of the quantum well. However, there is no change in the absorption edge of the Ag-NP-infiltrated InGaN QWs.

It is clear from the results above that there has been a modification in the relative intensities and emission energy of the zero-phonon line and the two phonon sidebands due to the presence of the Ag nanoparticles localized in the pits. This, in conjunction with the direct observation of a change in the symmetry of the Raman modes, presents compelling evidence of modification of the carrier-phonon interactions within the QWs. We discuss now two possible causes for the observed experimental results, and possible experiments and modelling that may distinguish between the two effects.

The first possibility is that the modification is due to out-of-sample scattering of wave-guided modes by the Ag NPs. This is not an unreasonable assumption as the LSP energy is near-resonant to the emission energy of the QW, and so the Ag NPs should be able to scatter the light within the QW. Essentially, for scattering to explain the observed effects, there would need to be a significant anisotropy with respect to \(S\) and the emission
direction. Such anisotropy clearly exists, as is evidenced in the angle dependence of $S_0$ and $S_1$ in the reference sample. This is due to the strong dependence of $S$ on the phonon wave-vectors [15]. Due to this anisotropy, photons emitted normal to the QW have a substantially different e/h-LOP coupling than those emitted parallel to the QW. Mie scattering calculations, however, show that this is unlikely to be the reason for the observed results. Figure 6 shows the relative scattering intensity for s- and p-polarized light interacting with an Ag NP within an IHP. These calculations show that only 5% of the light scattered by the NPs will be scattered out-of-sample. Furthermore, the scattered emission is essentially entirely composed of s-polarized light, and is isotropic, meaning that it cannot directly explain the changes in angle-dependence observed in the Ag NP sample. Additionally, neither the excitation source for the confocal Raman measurements nor the Raman scattered light are resonant to the Ag-NPs. This means that scattering by the Ag NPs cannot explain the appearance of the $A_1$(TO) mode in a forbidden geometry. Still, the Mie simulations only account for the situation where a single particle exists in a pit. It is possible that multiple particles per pit scatter wave-guided modes more strongly and maybe responsible for some of the observed angle-dependence.

The second, more interesting possibility is that the direct interaction between the plasmons and the carriers or the lattice is altering the e-ph coupling strength. Previous work has demonstrated the existence of coupled plasmon-exciton [5] and plasmon-phonon modes [6]. While little work has been done investigating the e/h-ph coupling strength in the presence of these coupled modes, it is reasonable to expect that they should have a direct effect on the coupling strength. These coupled modes might also result in broken symmetry, allowing for the observation of the $A_1$(TO) mode in normally forbidden geometries.

Additional experiments may be able to better distinguish between these causes. Of primary interest would be confocal Raman measurements in [a (c-c) ā] and other geometries. Observation of other normally forbidden modes, Raman-inactive modes, or the mode being partially observable in multiple geometries may give clues as to how the symmetry of the system has been altered, and conclusively rule out scattering as a source. Furthermore, while the Raman measurements are non-resonant, the ADPL measurements are resonant to the system and
may contain additional components from this resonant interaction. Angle-dependent
electroluminescence would allow for separating out the effects due to resonant interaction
between the excitation source from those due solely to interaction with the Ag NPs.

We have presented here strong evidence of modification of e/h-LOP coupling in
InGaN/GaN MQWs by embedded Ag NPs. This effect is likely a result of plasmonic
coupling to electron or phonon modes within the QWs. Due to the production of phonon-
replicas in the spectrum, the e/h-LOP coupling strength has a direct impact on the quality
and efficiency of emission of the system; therefore understanding the source of this
modification may provide an additional mechanism for using plasmonics to improve the
internal quantum efficiency of light emitters. Furthermore, since phonon replicas exist in
a many semiconductors, such a technique would be applicable over a wide range of light
emitters.

REFERENCES


FIGURE LEGENDS:

**Figure 1.** (a) SEM images showing Ag NPs placed within the inverted hexagonal pits. (b) PL spectrum of the reference (black, dashed) vs. the Ag NP sample (blue) taken normal to the surface demonstrating the change in spectrum due to the NPs.

**Figure 2.** Raman spectrum of the Reference (black, dashed) and Ag NP (blue) samples. The position of the $E_2$(high) and $A_1$(TO) phonons are marked for reference. Note that the $A_1$(TO) peak is absent in the reference measurement, and that the $E_2$(high) is shifted 1.1 cm$^{-1}$ to the left for the Ag NP.

**Figure 3.** Angle dependent PL intensity for the zero phonon, first (dashed) and second (dotted) phonon replica of the InGaN/GaN MQWs with (blue) and without (black) Ag nanoparticles normalized with respect to the total emission (i.e. $I_{PR0} + I_{PR1} + I_{PR2} = 1$). The gray shaded region represents missing values replaced by interpolation with a spline.

**Figure 4.** The Huang-Rhys parameters (a) $S_0$ and (b) $S_1$ as a function of emission angle for the reference (black squares) and Ag (blue triangles) system.

**Figure 5.** (a) Emission energy of the main emission for reference (black squares), Ag (blue triangles) and measured as a deviation from the reference emission energy of 2.984 eV. (b). Emission energy of the 1st phonon-replica, (c). Emission energy of the 2nd phonon replica.

**Figure 6.** (a) Simulated relative scattering intensity from an Ag NP within a GaN IHP as a function of scattering angle for s-polarized (dashed) and p-polarized (dotted) light. Red cone represents the angles over which light is transmitted out. 3-D scattering pattern for (b) s-polarized and (c) p-polarized light from the same Ag NP. The dashed green circle represents the angles over which light can escape at the GaN/Air interface.

"Plasmonic Modification of electron-LO phonon...."
Figure 1.

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