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Effects of Nitrogen Incorporation on Structural and Optical Properties of GaNAsP Nanowires

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Abstract

In this work we carry out a comprehensive investigation of structural and optical effects in GaNAsP nanowires (NWs), which are novel materials promising for advanced photovoltaic applications. Despite a significant mismatch in electronegativity between N and As/P atoms, we show that incorporation of nitrogen does not degrade structural quality of the nanowires and the fabricated NW arrays have excellent compositional uniformity among individual wires. From temperature-dependent photoluminescence (PL) measurements, statistical fluctuations of the alloy composition are shown to lead to localization of photo-excited carriers at low temperatures but do not affect material properties at room temperature. According to time-resolved PL measurements, the room-temperature carrier lifetime increases in the GaNAsP NWs as compared with the GaAsP NWs, which indicates reduced non-radiative recombination. Moreover, in spite of the very low N content in the studied NWs (up to 0.16 %), their bandgap energy can be tuned by more than 100 meV. This is accompanied by about 30% reduction in the temperature dependence of the bandgap energy. The presented results demonstrate that alloying of GaAsP with nitrogen provides an additional means of design optimization, beneficial for, e.g., NW-based intermediate band solar cells that are highly dependent on the optimum bandgap structure.

Keywords:
Dilute nitrides, GaNAsP, nanowire, photoluminescence, Raman scattering
Introduction

III-V semiconductor nanowires (NWs) have in recent years attracted an increasing research interest owing both to their interesting fundamental properties\(^1\) and to the prospects of numerous potential applications in electronics, optoelectronics and photovoltaics.\(^2\,3\) III-V materials are highly relevant to these applications due to their high electron mobility and often a direct bandgap. The NW geometry itself also has numerous advantages over the traditional bulk and epilayer geometries. For example, it relaxes lattice-matching constrains, which enables the integration of the III-V NWs with lattice-mismatched materials like Si. Additionally, it lowers fabrication costs due to reduced material consumption. Despite a small material volume in NW arrays as compared with that in an epilayer geometry, it has been shown that optical absorption efficiency is not reduced - even a sparse NW array can absorb nearly 100% of the incident light, a property that is very promising for photovoltaic applications.\(^4\,5\)

Recently there has been a rising interest in ternary III-III-V or III-V-V alloys, where the incorporation of an additional element provides a degree of freedom for, e.g., bandgap tuning. These types of alloys have been successfully implemented in NWs based on InGaAs\(^6\,7\), AlGaAs\(^8\) and GaAsP\(^9\). The latter is of particular interest for photovoltaic applications, since its bandgap can be varied between 1.4 eV and 2.3 eV, which is useful for development of multi-junction solar cells involving a bottom Si junction. Traditionally, GaAsP NWs have been grown using a gold droplet catalyst.\(^10\,11\) However, gold is known to introduce defects in III-V materials that degrade their optical quality.\(^12\,13\) Gold is also not compatible with the silicon technology, as it forms centers of efficient non-radiative recombination in Si detrimental for device performance.\(^14\) A combination of these effects makes gold-catalyzed NWs a less viable option for photovoltaic applications, prompting investigations into other growth methods. Recently it has become
possible to grow GaAsP NWs using self-catalyzed growth, where gold droplets are replaced by
gallium droplets, thereby eliminating the detrimental effects caused by gold incorporation. This
has led to the development of single-junction GaAsP NW solar cells with the efficiency
exceeding 10%, which were fabricated on a Si substrate.\textsuperscript{15,16}

To further increase the solar cell efficiency above that of a standard single-junction cell, various
so-called third generation solar cell designs have been suggested.\textsuperscript{17} One such example is the
intermediate-band solar cell (IBSC),\textsuperscript{18} where an intermediate energy band within the bandgap
facilitates not only direct but also two-step two-photon transitions enabling absorption of low-
energy photons. The intermediate band can originate from either confined states in quantum
dots\textsuperscript{19-22} or from impurity states in the bandgap.\textsuperscript{23-25} Theoretically, the efficiency of IBSC is
expected to reach as high as 63% even for a cell based on a single junction,\textsuperscript{26} which by far
exceeds the Shockley-Queisser limit.\textsuperscript{27} The highest efficiency is predicted for IBSC with a total
bandgap energy of 1.95 eV, which splits into two sub-gaps of 0.71 and 1.24 eV respectively.\textsuperscript{18}
One promising material which can be tuned to the optimum band structure is the GaNAsP alloy
belonging to the so-called dilute nitrides.\textsuperscript{28,29} These materials have a number of fascinating
properties, including splitting of the conduction band (CB) into the two separate $E^{-}$ and $E^{+}$ bands
that diverge with increasing nitrogen composition. This splitting is promising for IBSC, where
the $E^{-}$ and $E^{+}$ bands serve as the intermediate band and conduction band, respectively.\textsuperscript{23}

Growth of GaNAsP in the form of epilayers\textsuperscript{30} and stacked quantum well structures\textsuperscript{31} has been
demonstrated and two-photon absorption via the intermediate band has been shown to occur in
such materials.\textsuperscript{32} Most recently GaNAsP was also fabricated in the NW geometry,\textsuperscript{33} which opens
a door for integration of the IBSC concept with the promising NW architecture. To enable this
new technology, the effects caused by introducing nitrogen in GaAsP NWs must be thoroughly
understood, yet are hitherto largely unknown. In this work, we address these important issues based on comprehensive structural and optical characterization studies.

**Experimental Methods**

The studied GaAsP and GaNAsP NWs were grown on Si (111) substrates by plasma assisted molecular beam epitaxy (MBE) using Ga droplets as a catalyst. Thermally cracked PH$_3$ and AsH$_3$ were used as P and As sources, respectively, while Ga atoms were supplied from a solid Ga-source through an effusion cell. Nitrogen in GaNAsP was supplied using an RF nitrogen plasma, which was excited with powers in the range of 200-250 W. The plasma was ignited after 60 seconds of the GaAsP NW growth. Full details of the growth procedure can be found in Ref. 33.

Four structures were studied including reference GaAsP NWs and GaNAsP NWs grown with increasing nitrogen flux (denoted as N1, N2 and N3, respectively). Growth parameters of the investigated structures are summarized in Table 1.

**Table 1:** Growth parameters of the investigated GaAsP and GaNAsP NWs.

<table>
<thead>
<tr>
<th>Sample</th>
<th>GaAsP</th>
<th>GaNAsP (N1)</th>
<th>GaNAsP (N2)</th>
<th>GaNAsP (N3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ga flux (ml/sec)</td>
<td>0.6</td>
<td>0.6</td>
<td>0.6</td>
<td>0.6</td>
</tr>
<tr>
<td>AsH$_3$ flux (sccm)</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>PH$_3$ flux (sccm)</td>
<td>4</td>
<td>4</td>
<td>4</td>
<td>4</td>
</tr>
<tr>
<td>N flux (sccm)</td>
<td>0</td>
<td>0.3</td>
<td>0.6</td>
<td>0.9</td>
</tr>
</tbody>
</table>
The NW morphology was investigated using a LEO 1550 Gemini scanning electron microscope (SEM), operated with an extraction voltage in the range of 2-4 kV. The crystallographic structure and alloy composition were analyzed using transmission electron microscopy (TEM) and energy dispersive X-ray spectroscopy (EDXS) performed with a FEI Tecnai G2 transmission electron microscope. For these purposes, the NWs were mechanically transferred to a TEM grid. The TEM images were obtained using a CCD camera, while scanning TEM (STEM) images were acquired by a high-angle annular dark-field detector.

Optical properties and lattice dynamics of NWs were extensively characterized using microphotoluminescence (µPL) and µ-Raman spectroscopies. For measurements on individual wires, the NWs were mechanically transferred to a Si substrate, where the individual NWs could be resolved using an optical microscope. Both µPL and µ-Raman measurements were performed in a backscattering geometry using solid state lasers emitting at 532 or 660 nm for excitation. The laser beam was focused to an estimated spot diameter of 0.5-1 µm on a sample, using 50x (0.5 NA) or 100x (0.9 NA) objective lenses. The emitted (or scattered) light was dispersed using a grating monochromator and detected using a CCD camera. Photoluminescence excitation (PLE) measurements were performed with a tunable Ti:Sapphire laser as an excitation source. Transient photoluminescence studies were done at room temperature for NW arrays, excited by a frequency-doubled pulsed Ti:Sapphire laser at 440 nm with a pulse width of 2 ps. The PL signal was detected using a streak camera in combination with a single grating monochromator.

Results and Discussion

Electron microscopy
Effects of nitrogen incorporation on the NW morphology can be seen from Figures 1a-c, where representative SEM images of the NW arrays grown on Si substrates are shown. The GaAsP NWs are found to be uniform in both length (l) and diameter (d), with average values of approximately 2 µm and 90 nm, respectively—see Figure 1a. The average NW diameter does not
significantly change for the N1 (N2) GaNAsP wires grown with a nitrogen flux up to 0.6 sccm (Fig 1b,c), whereas a further increase in the N flux causes severe NW tapering accompanied by shortening of their lengths (Fig. 1d). This is attributed to the formation of strong Ga-N bonds, which limits the mobility and diffusion length of the Ga adatoms and accelerates consumption of the Ga droplet. After the Ga droplet was consumed, the initial axial growth via the vapor-liquid-solid (VLS) mechanism was terminated leading to the dominance of a radial vapor-solid (VS) epitaxial growth and, thus, enlargement of the NW sidewalls. With the highest N flux, the fast depletion of the Ga droplet may also promote the tapering, as the NW base can be expanded due to the transition from the VLS to VS epitaxial growth whereas the tip is unaffected by the VS growth and stays constant in diameter. All wires, however, retain a hexagonal cross-section characteristic for the axial growth along the <111> direction. This can be clearly seen from Figures 1e-f, where the top-view SEM images of single GaAsP and GaNAsP (N3) NWs are shown. The changes of the NW morphology with increasing N flux can also be seen by comparing the TEM images of single GaAsP and GaNAsP (N3) NWs shown in Figures 1g and h, respectively.

In contrast to the N-induced changes in the NW morphology, the incorporation of N does not affect the crystalline structure of the wires. Both GaAsP and GaNAsP wires are found to crystallize in the zinc-blende (ZB) lattice structure. The lower part of the NWs contains only a small number of rotational twin planes (seen as horizontal lines in Figures 1g and h), whereas in the upper part a significant increase in the twin plane density and minor inclusions of wurtzite segments are observed.

To evaluate effects of alloying with nitrogen on the alloy composition of the wires, we have performed scanning EDXS measurements and the representative results are shown in Figure 2.
Figure 2: (a) EDXS spectra, acquired from a single GaAsP and GaNAsP NW. The Gaussian fits of the Kα- peaks, used to estimate alloy content, are shown by the filled areas for one of the spectra. (b) and (c) show the EDXS line scans of the Kα-peak intensities for the GaAsP and GaNAsP NW, respectively. The corresponding TEM images of these NWs are shown in the low part of the figures. The scale bar is 500 nm.
The EDX signal was acquired by scanning the electron beam along the axial direction of the nanowire. Figure 2a shows the detected Kα-peaks from a single GaNAsP (N3) and GaAsP NW, where the signal has been integrated over the whole scan length and normalized to the Ga Kα-peak intensity. It can be seen that the relative peak intensities of the P-, As- and Ga-peaks are similar between the two samples. A slight increase (decrease) of the As (P) peak is seen with incorporation of nitrogen, but the change is close to the experimental accuracy. We, therefore, conclude that for the investigated N concentrations, the As/P ratio is not appreciably changed upon the formation of the GaNAsP alloy. This in turn implies that the N content in the GaNAsP NWs is rather low, i.e. below the detection limit of around 2%. Consistently, no N Kα-peak can be detected in the EDXS spectra. Based on the measured intensity ratios between the Kα-peaks, the P composition can be roughly estimated to be about $24 \pm 3\%$, which is in agreement with $[P] = 24\%$ deduced from the Raman and PL measurements to be discussed below. Moreover, within the experimental accuracy, the alloy composition is found to be practically constant along the nanowires suggesting that the nitrogen induced tapering does not appreciably affect the As/P ratio. This can be seen from Figures 2b and c, where the Kα peak intensities as a function of the scan position along the NW are shown together with the corresponding STEM images. The results of the EDXS study, therefore, show that alloying GaAsP with nitrogen results in no measurable change in the composition of the host material.

**Raman scattering**

To obtain evidence for N incorporation in the N1-N3 nanowires, and also to get further insight into the crystalline quality of the GaNAsP alloys, Raman spectroscopy was performed. Figure 3 shows unpolarized Raman spectra acquired from the studied NW arrays, which were performed
with the excitation and detection paths along the [111] crystallographic direction. All spectra show a bimodal behavior, where both GaAs- and GaP-like Raman modes are detected. In this geometry scattering involving zone-center longitudinal optical (LO) and transverse optical (TO) phonons are allowed from the Raman selection rules, and both types of the modes can indeed be seen for vibrations of the Ga-As and Ga-P bonds. The detected Raman spectra are characteristic for ZB GaAsP as the formation of the wurtzite phase should cause appearance of an additional
optical mode. In GaAs$_{1-x}$P$_x$ the spectral positions of the TO and LO modes are known to reflect the alloy composition, $x$. Based on the results reported by Im et al for GaAsP NWs, the alloy composition of our GaAs$_{1-x}$P$_x$ wires can be estimated as $x = 24 \pm 4 \%$. 

Previous Raman studies of dilute nitride GaNP and GaNAs alloys have shown that the incorporation of nitrogen into the host lattice may cause several effects on Raman spectra. They include: (i) shifts of the TO and LO modes caused by changes in lattice stiffness; (ii) appearance of an additional local mode representing vibrations of the Ga-N bonds; and also (iii) breaking of Raman selection rules and broadening of the Raman modes due to activation of forbidden scattering involving zone edge phonons in alloys with a high degree of lattice disorder. From Figure 3, the spectral positions of both TO and LO modes do not change in the GaNAsP NWs, which again suggests that the N content is rather low. It also indicates that the As/P-ratio does not change appreciably with nitrogen incorporation, in agreement with the EDXS results. The incorporation of nitrogen in the N1-N3 wires, however, can be confirmed based on the appearance on an additional mode centered at approximately 473 cm$^{-1}$ (labelled as LO$_2$ in Figure 3). Its observation only in the GaNAsP NWs is a clear indication that the mode appearance is an effect of nitrogen incorporation. The spectral position of the LO$_2$ mode is very close to that for the local mode due to the Ga-N bond vibrations in GaAs. On the other hand, the corresponding mode in GaNP has been reported at around 500 cm$^{-1}$, which is not observed in the present study. The fact that the LO$_2$ position corresponds almost exactly to that reported in GaNAs epilayers and NWs suggests that the local surroundings of N atoms is predominantly As-rich, which would result in the GaAs-like vibrations of the Ga-N bonds.
Finally it should be noted that the width of the Raman peaks, which is a measure of the structural disorder, is very similar in the GaAsP and GaNAsP NWs. This suggests that the structural quality does not deteriorate with nitrogen incorporation. This is further confirmed based on Raman measurement performed on single GaNAsP (a) and GaAsP (b) NWs transferred to a Si substrate.
The corresponding results are shown in Figures 4a and b, respectively. The measurements were performed with the excitation and detection beams directed along the [1\bar{1}0] crystallographic direction as shown in the inset in Figure 4. In this geometry, the LO mode is forbidden following the Raman selection rules in the ZB lattice. Accordingly, the LO mode is significantly suppressed as evident from a comparison of the TO/LO intensity ratios in the Raman spectra shown in Figures 4 and 3. However, the forbidden LO mode, although suppressed, is still visible in the spectra measured from the lying wires. This effect has been previously observed e.g. in polymorphic GaAs NWs and was attributed to the formation of (111) NW facets on the NW surface caused by the presence of rotational twins, which permits the LO scattering. The same mechanism is also applicable to the studied NWs. We, therefore, conclude that the Raman scattering generally obeys the selection rules in both GaAsP and GaNAsP NWs which implies that their crystalline quality is reasonably high.

Photoluminescence spectroscopy

To further understand effects of N incorporation on alloy properties and carrier recombination, we have performed detailed temperature-dependent PL and transient PL studies.

2a. Origin of the PL emission

Figure 5a shows PL spectra acquired at 4.5 K from the GaAsP and GaNAsP NW arrays. It is evident that alloying GaAsP with nitrogen leads to a significant red shift in the emission energy, that is characteristic for dilute nitrides due to the giant bandgap bowing effect. This red shift increases in the wires grown with a higher nitrogen flux, suggesting that the higher flux leads to a higher nitrogen content. The same conclusion can also be reached based on the performed PLE
measurements – see Figure 5b, where PLE-spectra acquired at 4.5 K from the three GaNAsP NW arrays are shown. (The bandgap energy of the GaAsP NWs was unfortunately outside the energy range of a tunable laser used for the PL excitation, which prevented us from performing the PLE studies of this N-free sample). Since absorption transitions in semiconductor alloys mainly involve extended band states with a higher density, the onset of the PLE spectrum corresponds to
the alloy bandgap. Taking into account that the studied alloys have a direct bandgap, their bandgap energies can be determined by linearly extrapolating the squared PL intensity ($I_{PL}^2$) as a function of excitation photon energy (shown by the dotted lines in Figure 5b). This yields the bandgap energies of 1.70, 1.70 and 1.68 eV for the structures N1, N2 and N3, respectively.

From Figure 5 it is apparent that incorporation of nitrogen also leads to a broader and more asymmetric lineshape of the monitored PL emission, with a tail extending to the low energy side of the PL spectra. This broadening does not arise chiefly from variations in N compositions between individual wires within the NW array as it is also observed in µPL spectra acquired from single wires – see the inset in Figure 5a, where µPL spectra of a single GaAsP and GaNAsP (N1) NWs are shown. The asymmetric PL lineshape and related broadening are characteristic for dilute nitrides. Previous studies of GaNAsP quantum well structures have shown that the emission at low temperatures is completely dominated by localized exciton (LE) recombination, where the excitons are localized within band tail states arising from compositional fluctuations. Similar results have also been reported in GaAs/GaNAs core/shell NWs. The observed large Stokes shift between the PL and PLE spectra provide further evidence that the low-temperature PL emission in the GaNAsP NWs entirely stems from the LE recombination. Results of µPL measurements show that, whereas the PL spectrum of the GaAsP wire consists of a single peak, the GaNAsP emission contains numerous sharp lines superimposed on a broad background. We have previously reported a similar behavior in GaAs/GaNAs core/shell NWs, where the sharp lines are attributed to quantum dot-like potential confinement of the excitons due to short-range alloy fluctuations in N composition.

To further investigate the localization potential within a single wire, temperature-dependent µPL studies were performed on individual GaAsP and GaNAsP (N2) NWs. Figure 6a shows the
resulting spectra from the GaNAsP NW. It is seen that the LE peak position redshifts with increasing temperature, $T$, due to exciton de-trapping from the localized states. At around 180 K, a new feature due to free exciton (FE) and/or free carrier recombination starts to appear at the high-energy side of the PL spectra, and becomes dominant at $T > 250$ K. The FE emission also shifts to lower energies with increasing $T$ due to thermal variation of the bandgap of the alloy. Similar behavior is found for the GaAsP NWs (not shown here), which means that at low

Figure 6: (a) Temperature-dependent µPL spectra acquired from a single GaNAsP (N2) NW. (b) Peak positions of the LE (dots) and FE/carrier (triangles) emissions detected from the single GaAsP (filled orange symbols) and GaNAsP (blue open symbols) NWs. The star denotes the PLE onset in the GaNAsP NW, taken from Figure 4b. The dotted lines show the transition from the LE- to FE-dominated emission. The solid lines are the best fits to the experimental data using the Varshni equation for the GaAsP NW and the BAC model for the GaNAsP NW.
temperatures the emission spectra of both the ternary and quaternary alloys are dominated by the LE recombination, while thermal depopulation of the localized states at room temperature leads to the predominance of the FE/carrier emission. Figure 6b shows the peak positions of the two peaks as a function of temperature. The dark blue circles (triangles) represent the maximum positions of the LE (FE) emission from the GaNAsP NW, whereas the orange circles (triangles) show the peak positions of the corresponding PL bands from the GaAsP NW. The filled symbols indicate which recombination process is dominating. The star symbol shows the PLE onset. The transition of the dominant PL emission from the LE to FE emission with increasing T, combined with the temperature-dependent redshift of these PL bands, leads to the so-called ‘S-shape’ of the PL maximum position vs. temperature. It can be seen that the energy difference $E_{FE} - E_{LE}$ is larger for the GaNAsP wires and that the transition between the LE- and FE/carrier emission occurs at a higher temperature in GaNAsP as compared to that in GaAsP. These findings suggest that the localization potential becomes deeper due to alloying with nitrogen. Taking into account that the low-energy tail $I(E)$ of the LE emission reflects an exponential density distribution of the localized states, the localization potential ($E_0$) can be quantified from the slope of the PL tail as:

$$E_0 = \left( \frac{d}{dE} \ln I(E) \right)^{-1}$$

(1)

The $E_0$ values can be estimated to be about 20 and 70 meV for the GaAsP and GaNAsP NWs, respectively. These values are comparable to that reported previously in dilute nitride alloys. The observed increase in the localization potential in the N-containing wires is probably not surprising, since under the giant bowing effect of the bandgap energy in dilute nitrides a small variation in the nitrogen composition can lead to a large change of the bandgap energy.
2b. Alloy composition

The aforementioned results from the temperature-dependent PL measurements allow us to further verify the As/P ratio in the studied wires and also to accurately determine the N composition. Based on previous studies of dilute nitrides, temperature dependence of the bandgap energy ($E_{\text{g GaNASP}}$) in these materials can be accurately reproduced using the so-called band anticrossing (BAC) model\(^\text{50}\) as

$$E_{\text{g GaNASP}} = \frac{1}{2} \left( E_{\text{g GaAsP}}(T) + E^N \right) - \sqrt{E_{\text{g GaAsP}}(T) - E^N}^2 + 4V^2 y \right)$$

(2).

Here $E_{\text{g GaAsP}}(T)$ is the temperature-dependent bandgap of the GaAsP host matrix, $E^N$ is the energy position of the nitrogen level in GaAsP, $V$ is the parameter describing the interaction between the host and nitrogen states and $y$ is the nitrogen content. Both $E^N$ and $V$ are assumed to be temperature-independent. The parameters $E^N$ and $V$ are well known for both GaNAs and GaNP,\(^\text{51}\) and can be obtained for GaNAsP by a linear interpolation of the corresponding values as discussed in detail in Ref. 28 and 30. The $E_{\text{g GaAsP}}(T)$ and $E_{\text{g GaNASP}}(T)$ values are experimentally determined based on the measured peak positions of the FE/carrier emission and the PLE onset. This also allows us to verify the P composition $x$ as the bandgap of GaAs$_{1-x}$P$_x$ and its temperature dependence can be computed using the Vegard’s rule as

$$E_{\text{g GaAs}_{1-x}\text{P}_x}(T) = (1 - x)E_{\text{g GaAs}}(T) + x E_{\text{g GaP}}(T) - x(1 - x)\beta, \quad (3)$$

where $\beta=0.19$ is the bowing parameter of GaAsP\(^\text{52}\) and $E_{\text{g GaAs}}(T)$ and $E_{\text{g GaP}}(T)$ are the temperature dependent bandgaps of GaAs and GaP, respectively, which were taken from Ref. 52. The best fit to the experimental data using Eq. 2 and 3 (shown by the solid lines in Figure 6b)
yields $x = 24\%$ for the GaAsP NW, in accordance with both EDXS and Raman results. Taking into account that the As/P ratio is not appreciably affected by the presence of nitrogen, the nitrogen composition for the GaNAsP ($\text{N}_2$) NWs can then be determined as $y=0.14\%$, whereas it is found to be 0.12 and 0.16\% in the N1 and N3 NWs, respectively. We note that, in spite of the very low N compositions, the bandgap energy of the GaNAsP nanowires can be varied by more than 100 meV. It also becomes less sensitive to variations in the measurement temperature, as can be seen by comparing the $E_g(T)$ dependences for the GaAsP and GaNAsP wires shown in Figure 6b. Both of these properties are highly beneficial for future optoelectronic and photonic applications of such structures.

2c. Compositional uniformity between different nanowires

As discussed above, the band gap energy and emission characteristics of the studied wires critically depend on their alloy composition. Therefore, for the material system to be viable for applications, the composition uniformity among the NWs is of significant importance. To investigate deviations in the alloy composition among nanowires belonging to the same NW array, we have performed room-temperature µPL spectroscopy on 50 GaAsP and 50 GaNAsP (N2) NWs. Figure 7 shows the obtained histograms of the PL peak position, where the striped blue and solid orange columns represent the results from the GaNAsP (N2) and GaAsP NWs, respectively. By fitting the GaAsP data with a normal distribution function (shown by the solid
line in Figure 7), we find the mean value $\mu = 1.71 \text{ eV}$, the bandgap energy which corresponds to the P-composition of $x = 24 \%$. We also obtain $\sigma = 3.4 \text{ meV}$, meaning that within one standard deviation $\sigma$, the compositional variation of As and P among the GaAsP NWs within the fabricated NW array is $\pm 0.3 \%$. The fitting parameters for the GaNAsP (N2) NWs are $\mu = 1.63 \text{ eV}$ and $\sigma = 9.2 \text{ meV}$. Here $\sigma$ is slightly larger than that for the GaAsP sample, which is to be expected as the variation here is a combined effect of the compositional variation of three elements: As, P and N. If we assume that the fluctuation in the As/P ratio is not affected by the

Figure 7: Histograms showing statistical distributions of the peak positions in the $\mu$PL spectra measured at room temperature from 50 GaAsP (solid orange) and GaNAsP (striped blue) NWs. The data were fitted using a normal Gaussian distribution function as shown by the solid curves. The pink dotted curve shows the calculated energy fluctuations expected in the GaNAsP NW assuming the complete uniformity in the N content.
presence of nitrogen, it is possible to estimate the compositional variation of N. The nitrogen content among the GaNAsP (N2) wires is found to be $0.14 \pm 0.02 \%$. To illustrate the effect of the nitrogen related compositional non-uniformity, the variance of the GaNAsP emission energy was also computed assuming complete uniformity in nitrogen composition between the NWs. This is illustrated by the pink curve in Figure 7. The results of the statistical analysis show that the GaAsP NW array has excellent compositional uniformity, which also remains reasonably high within the GaNAsP NW array.

2d. Carrier lifetime

Carrier lifetime is a very important parameter of semiconductors as it reflects contributions of different radiative and non-radiative processes in the material and, therefore, its optical efficiency, and also determines transport properties. The carrier lifetime in III-V materials at room temperature is usually limited by the fast non-radiative recombination, which mainly stems from surface recombination in the NW geometry\textsuperscript{53-55} due to a large surface-to-volume ratio. To gain insights into effects of N incorporation on the lifetime of photo-excited carriers, we performed time-dependent PL measurements on the NW arrays. The results are summarized in Figure 8a, where time decays of the integrated PL intensity are shown. It can clearly be seen that the PL emission from the GaAsP NWs decays faster as compared with the three GaNAsP samples and the latter exhibit very similar transient characteristics. The transient PL decays can be fitted by a single exponential function,

$$I_{PL}(t) = I_0 \exp\left(-t/\tau_{PL}\right),$$

where $\tau_{PL}$ is the PL lifetime and is given by $1/\tau_{PL} = 1/\tau_r + 1/\tau_{nr}$, with $\tau_r (\tau_{nr})$ being the radiative (non-
radiative) lifetime. By fitting the data in Figure 8a by Eq. (4), we find the PL lifetimes in the GaAsP and GaNAsP NWs to be 5 and 10 ps, respectively. The results of this fitting are shown by the solid lines in Figure 8a. The radiative lifetime in GaAs-based alloys is known to be of the order of ns even at low temperatures\textsuperscript{56,57} and is expected to further increase at elevated temperatures\textsuperscript{57} as the investigated NWs have too large diameters to experience quantum confinement effects. This means that the non-radiative lifetime in our structures controls the total
PL lifetime and that the increase in the PL lifetime upon nitrogen incorporation is an effect of reduced non-radiative recombination. Such behavior is opposite to that detected previously in dilute nitride epilayers and quantum structures, where alloying with nitrogen facilitated formation of various point defects acting as centers of efficient non-radiative recombination and controlling carrier lifetime.\textsuperscript{58-60} A similar decrease in non-radiative recombination has previously been seen in GaAs/GaNAs core/shell NW structures, however, where the incorporation of nitrogen also led to an increase in lifetime of the GaNAs-related emission. This was attributed to unintentional nitridation of the NW surface leading to the passivation of non-radiative surface states.\textsuperscript{56} The same effect may contribute to the PL lifetime increase in the GaNAsP wires seen in Figure 8a, which also suggests that it may be common for dilute nitride nanowires. Consistently, the GaNAsP NWs are found to be on average brighter than their GaAsP counterpart. This can be seen from Figures 8b and c, where the statistical distributions of integrated PL intensity of 50 NWs from each type of the structures are shown.

**Conclusions**

In summary, we have performed a comprehensive analysis of structural and optical properties of novel GaNAsP NWs using electron microscopy and a number of optical characterization techniques. Despite a significant mismatch in electronegativity between N and As/P atoms, the performed TEM and Raman studies show that alloying with nitrogen does not degrade the structural quality of the GaAsP NWs, though the high N flow during the growth promotes tapering. The statistical study of 50 GaAsP and GaNAsP NWs have revealed excellent compositional uniformity among the NWs constituting the NW arrays, which only slightly
deteriorates after nitrogen incorporation. From temperature-dependent PL measurements, statistical fluctuations of the alloy compositions are shown to lead to localization of photo-excited carriers at low temperatures, evident from the predominance of the LE emission in both GaAsP and GaNAsP NWs. The localization potential significantly increases upon nitrogen incorporation, promoted by the giant bandgap bowing in dilute nitrides. The room-temperature PL emission from both types of NWs, however, is dominated by free exciton/carrier recombination due to thermal activation of the photo-excited carriers from the localized tail states to extended band states. According to the time-resolved PL measurements, the carrier lifetime increases in the GaNAsP NWs, which indicates a reduced non-radiative recombination likely due to in-situ surface passivation with nitrogen. Moreover, in spite of the very low N content in the studied NWs (i.e. up to 0.16%), the bandgap energy of the GaNAsP nanowires can be tuned from 1.81 down to 1.68 eV (at 4 K). This is accompanied by a reduction in the temperature dependence of the bandgap energy, which is about 30% within the temperature range of 4.5 - 300K. Our study, therefore, demonstrates that alloying GaAsP with nitrogen represents a viable and attractive approach of bandgap engineering, which allows efficient tuning of the electronic properties of the GaNAsP NWs as required for future optoelectronic and photonic applications.

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References


33. La, R.; Pan, J. L.; Bastiman, F.; Tu, C. W. Self-Catalyzed Ga(N)AsP Nanowires and GaAsP/Ga(N)AsP Core-Shell Nanowires Grown on Si (111) by Gas-Source Molecular Beam Epitaxy. J. Vacuum Sci. & Tech. 2016, 34, 02L108.


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