High-pressure Raman scattering in wurtzite indium nitride

J. Ibáñez,1 F. J. Manjón,2 A. Segura,3 R. Oliva,1 R. Cuscó,1 R. Vilaplana,4 T. Yamaguchi,5 Y. Nanishi,5 and L. Artús1

1) Institut Jaume Almera, Consell Superior d’Investigacions Científiques, 08028 Barcelona, Catalonia, Spain
2) Instituto de Diseño para la Fabricación y Producción Automatizada, MALTA Consolider Team-Universitat Politècnica de València, 46022 València, Spain
3) Departamento de Física Aplicada-ICMUV-MALTA Consolider Team, Universitat de València, 46100 Burjassot, València, Spain
4) Centro de Tecnologías Físicas, MALTA Consolider Team-Universitat Politècnica de València, 46022 València, Spain
5) Faculty of Science and Engineering, Ritsumeikan University, Shiga 525-8577, Japan

We perform Raman-scattering measurements at high hydrostatic pressures on $c$-face and $a$-face InN layers to investigate the high-pressure behavior of the zone-center optical phonons of wurtzite InN. Linear pressure coefficients and mode Grüneisen parameters are obtained, and the experimental results are compared with theoretical values obtained from ab initio lattice-dynamical calculations. Good agreement is found between the experimental and calculated results.
Over the last few years, indium nitride (InN) has received intensive research interest because of its great potential to develop a wide variety of applications such as high-frequency electronic devices, high-efficiency tandem solar cells or infrared light-emitting devices.\(^1\) Although the growth of high quality InN is still challenging, bulk material with residual electron densities lower than \(5 \times 10^{17} \text{ cm}^{-3}\) and electron mobilities higher than \(2000 \text{ cm}^2\text{V}^{-1}\text{s}^{-1}\) has been achieved by molecular beam epitaxy (MBE).\(^2\) In spite of the great interest of InN from both fundamental and applied points of view, many material properties of this compound remain to be investigated. The production of high-quality InN layers and the discovery of the 0.7 eV fundamental band gap of wurtzite InN (w-InN) made necessary to revise many material parameters of InN and InN-based alloys, including their high-pressure behavior.\(^3,4\)

While the available experimental data on the vibrational properties of InN stem mainly from Raman spectroscopy (see for instance Ref. 5 and references therein), a deeper understanding of the lattice dynamics in InN has only been accomplished very recently by means of grazing incidence inelastic x-ray scattering (IXS) measurements on high-quality InN epilayers.\(^6\) Regarding the high-pressure vibrational properties of InN, Pinquier et al.\(^7,8\) used Raman scattering to study the pressure dependence of the \(E_{2h}, A_1(\text{TO})\) and \(A_1(\text{LO})\) phonon modes of w-InN in a layer with a high background electron concentration \((2.3 \times 10^{19} \text{ cm}^{-3})\). These authors observed the wurtzite-to-rocksalt transition and studied the pressure dependence, up to 50 GPa, of broad bands arising from the rocksalt phase.\(^8\) However, the pressure dependence of the long-lived \(E_{2l}\) mode and of the \(E_1\) modes of w-InN was not reported in those works. Recently, Yao and co-workers employed Raman scattering to investigate the structural stability of poorly crystalline w-InN nanowires under high pressure.\(^9\) The pressure coefficients and mode Grüneisen parameters obtained by these authors for the \(A_1(\text{TO}), E_{2h}\) and \(A_1(\text{LO})\) modes were sizably lower than those reported in Refs. 7 and 8.

In this letter we present Raman-scattering measurements under high hydrostatic pressure on high-quality \(c\)-face and \(a\)-face InN epilayers grown by MBE. Experiments on both types of samples have allowed us to study the pressure behavior of the non-polar \(E_{2l}\) and \(E_{2h}\) optical phonons and of the polar \(A_1(\text{TO}), E_1(\text{TO})\) and LO phonons of w-InN. The linear pressure coefficients and mode Grüneisen parameters of these modes have been determined and found to be in good agreement with the results of \textit{ab initio} lattice-dynamical calculations based on density functional theory (DFT).

For the present study we used two different w-InN epilayers, with thicknesses of 5.7 \(\mu\text{m}\)
FIG. 1. Room-temperature Raman spectra of a $c$-face InN layer at different hydrostatic pressures in the up-stroke (a) in the low-frequency region and (b) in the high-frequency region.

($c$-face) and 500-nm ($a$-face). Both samples were grown by plasma assisted MBE on sapphire substrates. The background electron concentration of the samples was $\sim 1 \times 10^{18} \text{ cm}^{-3}$ ($c$-face) and $\sim 4 \times 10^{18} \text{ cm}^{-3}$ ($a$-face). Flakes of w-InN containing some residual sapphire were detached from the substrate and loaded in a gasketed membrane-type diamond anvil cell (DAC) with 400 $\mu$m culet-size diamonds. Methanol-ethanol-water (16:3:1) was employed as pressure transmitting medium, and the ruby fluorescence method was used to determine the applied pressure. Confocal micro-Raman measurements at room temperature were acquired with a HORIBA Jobin-Yvon LabRam-HR spectrometer. To reduce the Rayleigh radiation and detect the low-frequency $E_{2l}$ mode of w-InN by means of a single-grating spectrometer such as the LabRam-HR system, we used the 632.8-nm line of a He-Ne laser as excitation radiation. A 50× objective was employed to focus the laser beam and collect the scattered radiation. In the case of the $c$-face sample, the measurements were carried out up to the transition pressure, which was observed to begin at $\sim 12 \text{ GPa}$ in this sample. For the $a$-face sample, the experiments were performed up to a pressure of $\sim 10 \text{ GPa}$.

Figures 1 and 2 show selected Raman spectra at different pressures for the $c$-face and the $a$-face epilayers, respectively. As expected from the selection rules for Raman experiments in backscattering geometry on a $c$ face of wurtzite crystals, the spectra of Fig. 1 are dominated by the non-polar $E_{2l}$ and $E_{2h}$ modes of w-InN [Figs. 1(a) and (b), respectively]. The weak
FIG. 2. Room-temperature Raman spectra of a a-face InN layer at different hydrostatic pressures in the up-stroke (a) in the middle-frequency region and (b) in the high-frequency region.

band that appears in the high-frequency region of the spectra in Fig. 1(b) contains contributions of both $A_1(LO)$ and $E_1(LO)$ modes. Note that, at ambient pressure, the Raman spectra of as-grown c-face epilayers display the expected $A_1(LO)$ mode and also some weaker signal arising from forbidden $E_1(LO)$ mode. The observation of $E_1(LO)$ signal through impurity-induced Fröhlich interaction in the spectra of c-face InN layers is attributed to the relaxation of the selection rules induced by defects. When the InN material is loaded into the DAC, the LO band is found to broaden and shift to higher frequencies, while the dominant $E_{2h}$ peak remains basically unchanged. This suggests that sample misorientation and/or disorder generated during the sample loading gives rise to a further enhancement of the $E_1(LO)$ signal. Similarly, in the case of the a-face epilayer both the $A_1(LO)$ and $E_1(LO)$ modes are symmetry forbidden and they become visible through impurity scattering [Fig. 2(b)]. The spectra of the a-face sample also exhibit the $A_1(\text{TO})$ and $E_1(\text{TO})$ modes, which show up as weak features below the $E_{2h}$ peak [Fig. 2(a)]. Among all the modes observed in this work, the low-intensity TO peaks yield the highest error in the frequency measurement, which we estimate is lower than 2 cm$^{-1}$. The $E_{2l}$ peaks detected in the case of the a-face epilayer (not shown) were fairly weak, superimposed to a relatively strong background signal.

Both Figs. 1 and 2 show the expected frequency increase for all phonon modes with
FIG. 3. Pressure dependence of the optical phonon frequencies of w-InN measured in the upstroke in c-face InN (solid triangles) and in the a-face InN (open circles). The lines are the results of linear fits to the experimental data.

Increasing pressure, with the exception of the $E_{2l}$ mode, which displays a pressure-induced softening. This behavior is typical of shear phonon modes, i.e., zone-edge transverse acoustic (TA) modes in zincblende and diamond semiconductors. In general, it is found that the mode Grüneisen parameter of the soft mode linearly correlates to the transition pressure of the material. In the case of wurtzite compounds, the $E_{2l}$ mode corresponds to the TA(L) mode of the zincblende structure due to the folding of the zone-edge TA modes along the $\Gamma$-L direction of the Brillouin zone. The softening of the $E_{2l}$ mode has been observed in many different wurtzite compounds, including ZnO (Ref. 14) and GaN (Ref. 15). Only AlN (Refs. 13 and 16) and BeO (Ref. 17) seem not to exhibit this softening behavior. While the positive pressure coefficient for the $E_{2l}$ mode of BeO is probably linked with its high pressure stability, the case of AlN is somewhat puzzling, since the wurtzite-to-rocksalt transition occurs at a relatively low pressure ($\sim$ 20 GPa). The observation of positive pressure coefficients for the $E_{2l}$ mode of AlN might be related to the special balance between ionic and covalent restoring forces in this compound. In the case of InN, following the discussion of Ref. 12, the observed softening of the $E_{2l}$ mode is consistent with the observed transition pressures, around 12-14 GPa.

In Fig. 3 we have plotted the pressure dependence of the measured optical phonon fre-
quencies. Data obtained for the two samples studied in this work have been included in the figure. Note the similar pressure behavior observed for the $E_{2h}$ and LO modes in both samples. Table I displays the linear pressure coefficients ($a_i$) extracted with a linear fit to all the experimental data, i.e., employing the expression $\omega_i(P) = \omega_i(0) + a_i P$, where $\omega_i(P)$ is the frequency of the $i$th mode at a pressure $P$. Mode Grüneisen parameters for the $i$th mode, $\gamma_i$, are subsequently deduced by using the expression $\gamma_i = B_0 a_i / \omega_i(0)$, where $B_0$ is the bulk modulus of the material. Here we use a value of $B_0 = 143$ GPa, which we obtain from DFT $ab$ initio calculations (see details in Ref. 5). This value is in close agreement to that obtained recently with grazing incidence IXS measurements ($B_0 = 152$ GPa). We plot in Fig. 3 the results of linear fits to the experimental data (solid lines). The resulting mode Grüneisen parameters for the different phonons are displayed in the last column of Table I.

The $a_i$ values obtained with our high-pressure Raman experiments for the TO, $E_{2h}$ and LO modes of InN are lower than those reported in InN films with much higher residual electron densities. Taking into account that InN epilayers with higher densities of threading dislocations seem to exhibit higher background electron densities, we speculate that the poorer crystalline quality of the InN epilayers used in previous works might be affecting the pressure behavior of the Raman-active modes. Note that, in contrast, the $a_i$ values obtained by Yao et al. on poorly-crystalline nanowires are even lower than those measured here.

From the experimental data obtained with the present measurements (Table I), it can be seen that the pressure behavior of the $A_1$(TO) phonons of w-InN (ionic vibrations along the $c$-axis) is similar to that of the $E_{2h}$ and $E_1$(TO) modes (vibrations perpendicular to the $c$-axis). GaN exhibits a similar behavior. This is in contrast, for instance, to the case of w-AlN, where the measured pressure coefficients for the $E_1$ and $E_{2h}$ modes are much larger than for the $A_1$ modes.

To gain further insight into the pressure behavior of the phonons of w-InN, we have performed $ab$ initio lattice-dynamical calculations of the linear pressure coefficients ($a_i$) and mode Grüneisen parameters ($\gamma_i$) for the zone-center optical phonons of this compound. The calculations, based on DFT within the local density approximation (LDA), were carried out with the ABINIT package. Calculations were performed at different pressures, and $a_i$ and $\gamma_i$ values were obtained with a linear fit to the data. Details of the calculations at ambient pressure can be found elsewhere. Since the ABINIT code fails to correctly predict the dispersion of the $E_1$(LO) mode in wurtzite materials, the calculated values for the LO
Phonon mode | $\omega_i(0)$ | $a_i$ | $\gamma_i$  
---|---|---|---
$E_{2l}$ | 88 (83.9) | -0.4 (-0.63) | -0.65 (-1.07) 
$A_1$(TO) | 449 (450.4) | 5.3 (4.69) | 1.69 (1.49) 
$E_1$(TO) | 476 (474.5) | 5.3 (4.88) | 1.59 (1.47) 
$E_{2h}$ | 492 (488.4) | 5.2 (5.18) | 1.51 (1.52) 
LO | 595 (591.8) | 5.1 (4.87) | 1.22 (1.17) 

TABLE I. Results of the linear fits to the pressure behavior of the optical phonons of w-InN. $\omega_i(0)$ and $a_i$ are the zero-pressure frequency and the linear pressure coefficient for the $i$th mode, respectively. A bulk modulus of 143 GPa was used to obtain the mode Gr"uneisen parameters ($\gamma_i$). Values in parentheses are the results of ab initio lattice-dynamical calculations. Theoretical values for the LO mode correspond to the $A_1$(LO) phonon.

phonon given in Table I correspond to the $A_1$(LO) phonon. The resulting $a_i$ and $\gamma_i$ values are displayed in parentheses in Table I. Good agreement is found between the experimental data obtained in the present work and the calculated values for the TO and the non-polar modes. In contrast, the theoretical $a_i$ values are systematically lower than those previously measured in bulk w-InN.$^{7,8}$ Remarkably, the DFT-LDA calculations predict the softening of the $E_{2l}$ mode, as observed with the present measurements. Note also that the DFT calculations predict a very small increase of the $A_1$(LO)-$A_1$(TO) splitting ($\sim 0.18$ cm$^{-1}$ GPa$^{-1}$). Instead, we observe a small decrease of $\sim -0.2$ cm$^{-1}$ GPa$^{-1}$ for the LO band in relation to both the $A_1$(TO) and the $E_1$(TO) modes. Within the experimental error of the present measurements, this result confirms that only a very small pressure dependence of the LO-TO splitting may be expected in w-InN. Finally, we would like to remark that, leaving aside the soft $E_{2h}$ mode, the ab initio calculations predict a very similar pressure behavior for all the Raman-active modes of w-InN regardless of the direction of the ionic vibrations, which is in agreement with the experimental results.

In conclusion, we have performed high-pressure Raman measurements on $c$-face and $a$-face layers to investigate the pressure behavior of the zone-center optical phonons of wurtzite InN. The linear pressure coefficient and mode Gr"uneisen parameters of the $E_{2l}$, $A_1$(TO), $E_1$(TO), $E_{2h}$ and LO modes of w-InN have been obtained. Good agreement between the experimental
data and theoretical values obtained with ab initio lattice-dynamical calculations has been found.

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REFERENCES

21The ABINIT code is a common project of the Université Catholique de Louvain, Corning Incorporated, and other contributors (http://www.abinit.org).
(a) $E_{2l}$

(b) $E_{2h}$

Raman shift (cm$^{-1}$)

Intensity (arb. units)

$c$-face InN

9.1 GPa
4.4 GPa
2.3 GPa
0.3 GPa

LO
a-face InN

(a) $A_1$(TO) $E_1$(TO) $E_2h$

(b) $E_2h$

Intensity (arb. units)

Raman shift (cm$^{-1}$)

6.8 GPa
4.5 GPa
2.2 GPa
0.3 GPa
Phonon frequency (cm$^{-1}$) vs Pressure (GPa)

- LO
- $E_{2h}$
- $E_1$(TO)
- $A_1$(TO)
- $E_2l$

Symbols:
- $\text{c-face}$
- $\text{a-face}$

Graph shows the variation of phonon frequencies with pressure.