Above-bandgap ordinary optical properties of GaSe single crystal

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We report above-bandgap ordinary optical properties of \( \varepsilon \)-phase GaSe single crystal. Reference-quality pseudodielectric function \( \langle \varepsilon(E) \rangle = (\varepsilon_1(E)) + i(\varepsilon_2(E)) \) and pseudorefractive index \( \langle N(E) \rangle = (n(E)) + i(k(E)) \) spectra were measured by spectroscopic ellipsometry within the CP parabolic-band model. 24–26

I. INTRODUCTION

Gallium selenide (GaSe) crystallizes in a laminar structure characterized by a strong covalent interaction within the layers and a weak van der Waals binding between them as occurs in many other III-VI compounds. The GaSe-type structure can be described as a stacking of hexagonal layers and there are four possible stacking arrangements leading to the four polytypes: \( \beta-, \varepsilon-, \gamma-, \) and \( \delta- \)GaSe. \(^{1,12–15}\) With such a unique structural anisotropy, GaSe has long been an interesting material for both experimental \(^{1,12–15}\) and theoretical \(^{1,13–15}\) physics studies, and GaSe and related III-VI compounds have received considerable attention for their potential applications in various device structures. \(^{16–20}\) Furthermore, advances in van der Waals epitaxy \(^{21}\) have made it possible to grow single-phase high-quality III-VI compound thin films that renewed interests in GaSe.

Knowledge of optical properties of materials over a wide photon energy range is of importance for designing photonic and photovoltaic devices as well as for verifying the predictions made by the calculations of the electronic energy band structure. Consequently, a number of optical studies of GaSe have been done in the past but with large discrepancies in the results. \(^{2,12–10}\) Early studies \(^{2–10}\) reported the dielectric function \( \varepsilon \) calculated mostly from the Kramers–Kronig (KK) transformation of the reflectivity data. Spectroscopic ellipsometry (SE) possesses several advantages over conventional reflectivity techniques in terms of the accuracy of data and has been recognized as a highly suitable method of determining \( \varepsilon \) of materials without the use of KK transformation. \(^{22}\) Therefore, SE has been employed widely to investigate optical properties of many semiconductors in the past few decades. \(^{23}\)

For SE study of GaSe, Meyer et al. \(^{11}\) measured both “ordinary” and “extraordinary” \( \langle \varepsilon \rangle \) spectra in which the light is polarized perpendicular \( (\hat{E} \perp \hat{c}) \) and parallel \( (\hat{E} || \hat{c}) \), respectively, to the optic axis from 2 to 5 eV under ultrahigh vacuum environment. They observed that the two sets of optical functions were significantly different and attributed it to the different selection rules for optical transitions. However, the spectral range was limited and no critical-point (CP) analysis was performed. Later, Adachi and Shindo \(^{12}\) also reported the ordinary optical response of \( \varepsilon \)-type GaSe in the similar spectral range (2–5 eV). Although they employed the model dielectric function scheme to analyze their experimental data and extract CP parameters, only two distinct optical structures at \( \sim 3.6 \) and \( \sim 4.9 \) eV were studied. More CP structures have been predicted and observed in many other studies. \(^{2–10,13,14}\)

Here, we applied SE to investigate above-bandgap optical properties of a GaSe single crystal up to 6.45 eV at room temperature. The measured spectrum exhibits several optical structures associated with interband-transition CPs and their energies have been accurately determined by analyzing the numerically calculated second-energy derivative of \( \langle \varepsilon \rangle \) within the CP parabolic-band model. \(^{24–26}\)

II. EXPERIMENTS

GaSe single-crystal \( (\varepsilon \)-type) used in this study has been grown by the vertical Bridgman method from a melt of stoichiometric GaSe, and then cleaved and cut into a square shape \( (8 \times 8 \times \sim 0.2 \text{ mm}^3) \) with the face normal to the [001] direction from the ingot. \(^{27}\)

The ordinary pseudodielectric function \( \langle \varepsilon \rangle = (\varepsilon_1) + i(\varepsilon_2) \) spectrum was measured from 0.73 to 6.45 eV at room temperature using a spectroscopic rotating compensator-type ellipsometer (M2000-DI model, J.A. Woollam Inc.). The angle of incidence was 70° and the data were recorded after averaging 1000 compensator cycles (1000 revolutions per measurement) to increase the signal-to-noise ratio.

SE is a surface sensitive technique. \(^{22}\) Determination of the material’s intrinsic optical properties is therefore complicated by the presence of surface overlayers that generally appear as organic contaminants, a native oxide layer, or microscopic roughness, but it could be any other surface perturbation in practice. Two approaches are usually taken to deal with overlayer artifacts in the experimental spectrum: (1) Mathematical correction within the multilayer model \(^{23}\) or...
(2) actual removal of overlayers using a wet-chemical etching procedure developed by Aspnes.27,28 For GaSe, however, its interlayer binding is weak enough to peel the top few layers off easily along the (001) plane,10–12 which allows one to obtain a fresh surface immediately prior to SE measurements. The SE data were then acquired under a flowing N2 gas environment to minimize surface contamination during the measurement.

III. RESULTS AND DISCUSSION

A. Optical functions

Figure 1 shows our $\langle e_2 \rangle$ spectrum for GaSe along with digitized versions of those reported in the literature.6,10–12 A usual procedure for verifying successful reduction in overlayer artifacts is to maximize the $\langle e_2 \rangle$ peak in the high-energy region where the penetration depth of the probing light is small and therefore the surface sensitivity is enhanced,28 which occurs at $\sim 5$ eV for many semiconductors—the $E_g$ CP peak. Our $\langle e_2 \rangle$ value at 4.94 eV is 26.22 which is about 10% higher than the value reported in the previous SE studies11,12 for the same optical structure. Furthermore, a few additional weak optical structures are also seen in the higher energy region ($>4$ eV) of our data which have not been resolved in the previous SE studies.11,12 We thus believe that the spectrum presented here provides a better representation of ordinary $\langle e \rangle$ for GaSe than those available in the literature.2–12 Our $\langle e \rangle$ and $\langle N \rangle$ spectra for GaSe are presented in Figs. 2(a) and 2(b), respectively.

We note that our data show small oscillations in the low-energy region ($<2$ eV), which is probably a thickness interference due to the back side reflection from the transparent characteristic of the material below its fundamental absorption edge ($E_g$).29 The film thickness estimated from the analysis of the oscillations period30 was about 4 μm, which is much smaller than the actual sample thickness of $\sim 200$ μm. We suspect that a group of layers (with the total thickness of $\sim 4$ μm) has partially lost its binding to the remaining bulk due to its weak interlayer binding energy while we peeled off the top few layers. Both the amplitude of oscillations and the depolarization (not shown here) are damped to zero at $\sim 2$ eV, which is indeed a good indication29 of the $E_g$ of GaSe.$^1$

One may expect to see that $e_2$ and $k$ increase gradually with $E$ for the $E > E_g$. As also observed in the previous SE studies,11,12 however, those values for GaSe in Fig. 2 are nearly zero up to $\sim 3.2$ eV, that is, as much as 1 eV higher than its $E_g$. According to theoretical study15 of the electronic energy band structure for GaSe, the bottom of the conduction band is formed mainly by the Ga $p_z$ state. The top of the valence band is derived from the Se $4p$ states, and the cation/anion-anion interactions split this band into the uppermost valence band with $p_z$ character and the lower doubly degenerated pairs with $p_x, p_y$ character, and the spin-orbit interactions further break the degeneracy. The $E_g$ of GaSe can then be defined as the interband transition from the uppermost valence band (Ga $p_z$) to the bottom of the conduction band (Ga $p_z$), but the optical selection rules prohibit this transition for $\vec{E} \perp \vec{c}$. A schematic15 for the electronic energy band structure at the Brillouin zone (BZ) center in the vicinity of the $E_g$ is given in Fig. 3. It is known31 that SE measurement of the surface normal to the optic axis carries relatively little information on extraordinary optical property of a uniaxial crystal due to the attenuation of normal component of an applied field by the boundary condition on the displacement vector $\vec{D}$. This is perhaps why SE data do not show a clear optical absorption up to $\sim 3.2$ eV where the allowed transition from the deeper valence band ($p_x, p_y$ char-

FIG. 1. (Color online) Imaginary part of $\langle e \rangle$ spectra for GaSe. The solid line represents the present work. Digitized versions of previously reported data (Refs. 6 and 10–12) are also included for comparison.

FIG. 2. (Color online) The pseudodielectric function $\langle e \rangle$ and the pseudorefractive index $\langle N \rangle$ for $\varepsilon$-phase GaSe.
energies that we obtained are listed in Table I. The CP features are present in a narrow range of the spectrum.34,35 The CP features are controversial, in particular, when several CP structures have recently been observed experimentally by Segura et al.15 suggested two deeper valence bands with Se $p_x$, $p_y$, and $p_z$ symmetry at 1.27 and 1.59 eV below the uppermost valence band with Se $p_z$ symmetry for GaSe. The intraband transition of 1.5 eV from the deeper valence band to the uppermost one has recently been observed experimentally by Segura et al.39 from their nonlinear optical studies as well as the interband transitions from those two deeper valence bands to the bottom of the conduction band at $\sim 3.3$ and $\sim 3.6$ eV. Following the results from recent theoretical15 and nonlinear optical35 studies, we attribute tentatively the origin of the CP structure at 3.23 eV that corresponds to the transition from the second highest valence band with the Se $p_x$,$p_y$ character to the bottom of the conduction band as discussed in Sec. III A. This assignment implies that the energy separation between this deeper valence band and the uppermost valence band is 1.23 eV assuming the $E_g$ is 2 eV, which is in an excellent agreement with the theoretical prediction15 of 1.27 eV (see Fig. 3).

However, multiple interpretations have been made for the next CP structure at 3.75 eV. It had been once assumed9 to be a resonant exciton associated with a two-dimensional CP involving a transition from the uppermost valence band to the lowest group of the conduction bands along T and S valleys of the BZ, but this assignment appears not to be consistent with the cases for other similar compounds such as InSe.38 Piacentini et al.10 assigned this CP to the transition from the uppermost valence band to the third group of the conduction band at the BZ center (Γ-point). Later, Kuroda et al.15 suggested two deeper valence bands with Se $p_x$, $p_y$ symmetry at 1.27 and 1.59 eV below the uppermost valence band with Se $p_z$ symmetry for GaSe. The intraband transition of 1.5 eV from the deeper valence band to the uppermost one has recently been observed experimentally by Segura et al.39 from their nonlinear optical studies as well as the interband transitions from those two deeper valence bands to the bottom of the conduction band at $\sim 3.3$ and $\sim 3.6$ eV. Following the results from recent theoretical15 and nonlinear optical35 studies, we attribute tentatively the origin of the CP structure at 3.23 eV that corresponds to the transition from the second highest valence band with the Se $p_x$,$p_y$ character to the bottom of the conduction band as discussed in Sec. III A. This assignment implies that the energy separation between this deeper valence band and the uppermost valence band is 1.23 eV assuming the $E_g$ is 2 eV, which is in an excellent agreement with the theoretical prediction15 of 1.27 eV (see Fig. 3).

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tures at 3.23 and 3.75 eV to the transitions from the two deeper valence bands to the bottom of the conduction band at the BZ center.

The CP structures appearing in the higher energy region have been understood as the excitations of the chalcogen $p_x, p_y$ electrons to the second and the third conduction bands at the various locations of the BZ without specific assignments. Positive identification of the origins for those CP structures is beyond the scope of this work. Rather, we intend that this work will help theoreticians perform fine adjustments of the electronic energy band structure of III-VI compounds including GaSe.

**IV. CONCLUSION**

Ordinary ($\vec{E} \perp \vec{c}$) optical properties $\langle e \rangle$ and $\langle N \rangle$ for $e$-phase GaSe single crystal were measured by SE at room temperature. Good approximations to the intrinsic bulk optical properties were established. Measured spectrum exhibits seven optical structures associated with above-bandgap interband-transition CPs and their energies were obtained by analyzing the lineshapes of the second-energy-derivative spectrum assuming the CP parabolic band model.

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