A first-principles theoretical study of the electronic and optical properties of twisted bilayer GaN structures

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Abstract
Gallium nitride (GaN) is a well-investigated material that is applied in many advanced power electronic and optoelectronic devices due to its wide bandgap. However, derivatives of its monolayer form, such as bilayer structures, have rarely been reported. We study herein the electronic and optical properties of GaN bilayer structures that are rotated in the plane at several optimized angles by using the density functional theory method. To maintain the structural stability and use a small cell size, the twisting angles of the GaN bilayer structures are optimized to be 27.8°, 38.2°, and 46.8° using the crystal matching theory. The band-structure analysis reveals that the bandgap is wider for the twisted structures compared with the nontwisted case. The simulation results provide the absorption coefficient, extinction coefficient, reflectivity, and refractive index at these angles. The spectra of all these optical properties match with the bandgap values. The simulated refractive index of the bilayer structures at all the twisting angles including 0° is smaller than that of bulk GaN, indicating a reduced scattering loss for optoelectronics applications. Considering the results of this analysis, the possible applications may include low-loss integrated electronic and optical devices and systems.

Keywords GaN bilayer · Twisting effect · Optical properties

1 Introduction
GaN is a typical group III–V compound semiconductor material and has been widely applied in power semiconductor and optoelectronic devices. For optical applications, GaN can be used as a substrate to make violet (405 nm) laser diodes. Light-emitting diodes (LEDs) made using GaN can achieve high brightness and cover the range of primary colors, being applicable in daylight-visible full-color LED displays and white LEDs. Because of their low sensitivity to ionizing radiation, GaN-based solar cell arrays are suitable for use on satellites. Meanwhile, the working temperature and voltage limit of GaN transistors are higher than those of gallium arsenide (GaAs) transistors, leading to many applications in microwave power amplifier devices [1–4]. GaN has also become one of the important materials for use in next-generation smartphones and Internet technologies, as it can meet the requirements of high power, high temperature, low refractive index (2.3 for GaN compared with 3.5 for GaAs), and superior temperature stability of the refractive index (one order of magnitude higher than that of InP).

It was reported in past studies that devices with a suspended bilayer structure offer advantages in terms of enhanced electronic performance. Indeed, such two-dimensional (2D) material forms are deemed to have much thinner thickness than the bulk form, resulting in different electronic, mechanical, and optical performance characteristics [5–15]. This is due to the change of the crystallization and the quantum effects in 2D materials. Some previous work applying density functional theory to investigate the electronic properties of 2D materials has been conducted [16]. First-principles simulations of GaN as a single-layer III–V binary compound semiconductor have also been reported [17]. The band structures of single-layer GaN can be controlled via chemical functionalization [18, 19], adsorption...
of nonmetal materials [20–23], vacancies [24, 25], and doping [26, 27] methods. These results have indicated that 2D GaN has potential as a future developmental trend for high-performance nano electronic devices. Experimentally, it has been reported [28, 29] that 2D GaN can be synthesized by the migration-enhanced encapsulated growth (MEEG) technique utilizing epitaxial graphene. It can also be grown by using the surface-confined nitridation reaction method on liquid metals [30], demonstrating the potential to implement 2D GaN in real nano electronic devices [23]. Structures formed by stacking two single-layer materials via van der Waals (vdW) forces and rotating one of them against the other may display interesting characteristics. The first studies in this regard [31, 32] reported that the use of a particular rotating/twisting angle in bilayer graphene resulted in superior electrical conductivity. In that work, the twisting angle could be controlled by using graphene produced by the exfoliation method. These previous papers on the twisting effect in stacked 2D materials have stimulated researchers to conduct studies on other multilayer structures based on various monolayer materials, including heterostructures. Specifically, for 2D GaN, such investigations have been performed using the first-principles method to investigate the effect of twisting in bilayer GaN. The twisted GaN bilayer model is built and subsequently analyzed herein to reveal its electronic and optical properties. These results will lead to wider and deeper explorations of single-layer GaN in the hope its application in practical electronic and optoelectronic devices.

2 Computational methods

The modeling of the bilayer GaN structure is based on the Quantum Atomix Toolkit simulation tools. The simulations are conducted using density functional theory (DFT). The method of generalized gradient approximation (GGA) with the Perdew–Burke–Ernzerhof (PBE) parameterization at a temperature of 300 K with a mesh cutoff energy of 50 Hartrees and an $11 \times 11 \times 1$ $k$-point grid are used to optimize all the structures. Considering the influence of the vdW force applied to the bilayers, semiempirical corrections using the Grimme DFT-D2 model are applied to include the long-range van der Waals interaction. A large (> 20 Å) vacuum spacing is added to the structure along the $c$-direction to avoid the application of periodic boundary conditions in this direction. The maximum force on each atom is set to 0.01 eV/Å for the structural relaxation calculations. For the simulation of the electronic and optical properties, the TB09 meta-GGA method is used for the bandgap, projected density of states, and optical properties with a $c$ parameter of 1.76. Figure 1 shows the calculated $c$ parameters against the calculated bandgap. The $c$ value is calculated to match the experimental bandgap of 4.98 eV reported previously [28] for (untwisted) two-monolayer structures. There are no reports to date on the bandgap for two vdW-bonded GaN monolayers. The vdW bonding has less effect on the bandgap of the monolayer structure. It is reported that the vdW interaction does not affect the properties close to the Fermi level [33] for vdW-bonded heterostructures, indicating that the bandgap of the vdW-bonded GaN bilayer should be similar to that of the monolayer. Side views of the GaN bilayer structure are shown in Fig. 2. From this viewpoint, the structure is built by stacking two layers of a 2D GaN cell. Figure 2 also shows a bottom view of the different rotated structures. The rotation angles are selected according to the criteria of optimized stress and atom number, with the aim of achieving the most stable structures. The method of selecting optimal rotation angles was described previously [34]. Generally speaking, for the twisted bilayer structure, the number of atoms in a single cell increases as the rotation angle is increased. Table 1 presents the number of atoms and bandgap in each structure. However, when the angle equals 38.21°, the atom number decreases to 56, much less than for the rotation angles of 27.8° and 46.8°. All the models are passivated with hydrogen atoms. Figure 3 shows the band structures of the atomic model and the projected density of states (PDOS) for the bilayer GaN structures with the four rotation angles. The bilayer structures have bandgaps of 4.97 eV, 5.06 eV, 5.25 eV, and 5.22 eV for the angles of 0°, 27.8°, 38.2°, and 46.8°, respectively, indicating that bilayer GaN displays a typical wide-bandgap semiconducting property. The bandgap values are significantly larger than the value for bulk GaN. For all the structures, both the valence-band maximum (VBM) and the conduction-band minimum (CBM) occur at the $\Gamma$ point, indicating that the bandgap is of direct type. According to the PDOS results, the VBM is primarily composed of N 2$p$ and H 1$s$ orbitals while the CBM is a hybridization of the Ga 3$s$, Ga 3$p$, N 2$s$, and H
1s orbitals, although Ga 3p contributes the most. The CBM occurs in Ga 3p orbitals while the VBM occurs in N 2p. Figure 4 displays the results for the bandgap, CBM, and VBM versus the rotation angle. As the rotation angle is increased, the bandgap first increases (from 0° to 38.2°) then decreases slightly (from 38.2° to 46.8°). The CBM and VBM remain at the G point with increasing rotation angle. However, when

| Rotation (°) | 0  | 27.8 | 38.21 | 46.83 |
|-------------|----|------|-------|-------|
| Atom numbers in single cell | 8  | 104  | 56    | 152   |
| Angle between vectors (°) | 60 | 60   | 60    | 60    |
| Bandgap (eV) | 4.97 | 5.06 | 5.25  | 5.22  |

Fig. 2 A side view of the rotated structure, and bottom views of the structure with rotation of a 0°, b 38.2°, c 27.8°, and d 46.8°

Fig. 3 The bandgap and PDOS of the structure with rotation of a 0°, b 27.8°, c 38.2°, and d 46.8°

Table 1 The atom numbers and bandgap value of the different rotated structure
the angle is 27.8° or 38.2°, the absolute value of the CBM is larger than that of VBM, indicating that the Fermi level moves towards the valence band.

3 Optical properties

The bilayer GaN models are simulated to determine several key optical properties such as the absorption coefficient and refractive index using the general mathematical procedure described in our previous work [35]. The susceptibility tensor at a frequency $\omega$, which characterizes the polarization of a material under an electrical field, can be written using the Kubo–Greenwood formula as

$$\chi_{ij}(\omega) = -\frac{e^2 h^4}{m^2 \epsilon_0 V \omega^2} \sum_{nm} f(E_n) - f(E_m) \frac{\epsilon_{nm}^* \epsilon_{mn}^*}{E_{nm} - \hbar \omega - i \Gamma_{nm}} $$

where $E_n$, $E_m$, and $E_{nm}$ represent the energy levels at the energy states $n$ and $m$ and the energy difference between $n$ and $m$. $f \Gamma$ and $V$ stand for the Fermi function broadening and volume, respectively. $\epsilon_{nm}^*$ is the $i$th component of the dipole matrix element between the states $n$ and $m$. $\epsilon$, $\hbar$, and $V$ represent the permittivity in vacuum, the reduced Planck’s constant, and the single electron charge, respectively. The susceptibility tensor is used to derive the relative permittivity ($\epsilon_r$), which determines the amount of polarization of a dielectric material under an external electric field. It is

$$\epsilon_r(\omega) = (1 + \chi(\omega)).$$

The polarization response of a material such as GaN under an electric field should be expressed in complex form, as there is always a phase change of the electric field. A simple expression for the complex permittivity at a frequency $\omega$ is

$$\epsilon_r(\omega) = \epsilon_r e^{i \epsilon'(\omega)} = \epsilon'(\omega) + i \epsilon''(\omega),$$

where $\epsilon_r$, $\epsilon'$, and $\epsilon''$ are the vacuum permittivity and the real and imaginary parts of the complex permittivity, respectively. Further expressions for the complex terms of the permittivity can be obtained from the Kramers–Kronig relations (bidirectional mathematical relations that connect the real and imaginary parts of any complex function) as follows:

$$\epsilon'(\omega) = 1 + \frac{2}{\pi} A \int_{0}^{\infty} \frac{\omega' \epsilon''(\omega')}{\omega'^2 - \omega^2} d\omega',$$

$$\epsilon''(\omega) = -\frac{2}{\pi} A \int_{0}^{\infty} \frac{\epsilon'(\omega')}{\omega'^2 - \omega^2} d\omega' + \frac{\sigma_0}{\epsilon_0 \omega},$$

where $\sigma_0$ and $A$ denote the direct-current (DC) conductivity and the Cauchy principal value. The refractive index ($n$) and the extinction coefficient ($\kappa$) can be written as

$$n = \sqrt{\frac{\epsilon'(\omega^2) + \epsilon''(\omega^2) + \epsilon'}{2}},$$

$$\kappa = \sqrt{\frac{\epsilon'(\omega^2) + \epsilon''(\omega^2) - \epsilon'}{2}}.$$

The optical absorption coefficient ($\alpha_\omega$) can then be written as

$$\alpha_\omega = \frac{2 \omega \kappa}{c},$$

where $c$ is the speed of light and $\omega$ is the angular frequency of the light. The reflectivity ($r$) is expressed as

$$r = \frac{(1 - n)^2 + \kappa^2}{(1 + n)^2 + \kappa^2}.$$

Figure 5 depicts the real and imaginary parts of the complex permittivity calculated for the rotated bilayer GaN structures for twisting angles of 0°, 27.8°, 38.2°, and 46.8°. In terms of the amplitude, the values are similar in all cases, except that the permittivity shows maximum peak values when the rotation angle is 38.2°. The peak wavelengths in all cases coincide with the bandgap values; i.e., there is a slight blue-shift for nonzero twisting angles. In Fig. 5a, the real part of the permittivity at a long wavelength of 400 nm is 1.7322, 1.7323, 1.7322, and 1.7322 for rotation angles of 0°, 27.8°, 38.2°, and 46.8°, respectively. The peak values of the imaginary permittivity appear between wavelengths of 200 nm and 250 nm in all cases according to Fig. 5b. Figure 6 shows the reflectivity, refractive index, absorption coefficient, and extinction coefficient spectra for the rotation angles of 0°, 27.8°, 38.2°, and 46.8°. Figure 6a shows the reflectivity curves against wavelength, being very similar in
all cases. The peak amplitude values lie in the range from $10^{-8}$ to $10^{-7}$. In Fig. 6b, the refractive index values for all the cases are much smaller than the values for bulk GaN material. In Fig. 6c, the maximum values of the absorption coefficient in the spectrum for all the cases are around 400 cm$^{-1}$. In Fig. 6d, the extinction coefficients in all cases follow the same trends as the absorption coefficient curves in terms of both the amplitude and wavelength. For the optical properties of the structure, a slight blue-shift is seen as a result of twisting. This occurs because the optical bandgap increases slightly as the rotation angle is increased. To explain these simulation results in terms of solid-state physics, a theoretical framework similar to that described in our previous paper [35] is utilized. Briefly, it is understood that the Lorentz model can be used to interpret the relations among all the above optical properties [36]. It is assumed that the bilayer structure and the incident optical light are sets of oscillating systems with harmonic resonant frequencies of $\omega_0$ and $\omega$, respectively. A high-absorption band appears when $\omega \approx \omega_0$. In the mismatched regions, this pseudomechanical theory applies in the interacting electron–photon system.

In optical applications, attenuation of the optical signal in a material is caused by absorption, scattering, and radiation of the optical power. In particular, the scattering loss occurs due to the refractive index caused by microscopic heterogeneities such as fluctuations in the material density or composition. The optical power loss due to scattering $I_s$ is proportional to the eighth power of the refractive index ($I_s \sim n^8$) [37], so a small reduction of the refractive index can significantly reduce the scattering loss. Interestingly, the refractive index of bilayer GaN is much smaller (~ 1.73) compared to that of bulk GaN (2.3), resulting in greatly reduced scattering loss of the optical power, which could enable promising applications in optoelectronic devices.
The optical power loss may be further reduced by the very low absorption of the bilayer structures simulated in this work, which is approximately three orders of magnitude lower compared with that of bulk GaN materials (reported to have an absorption coefficient of \( \sim 10^5 \text{ cm}^{-1} \) [38]). Moreover, the bandgap of two-layer GaN is much wider than that of bulk materials, introducing prospective applications in power electronics. The results of this theoretical investigation using the first-principles method reveal that the bandgap of bilayer GaN can be further increased by the twisting effect. Such twisted bilayer structures bonded by the vdW force can already be achieved experimentally [39]. Although experiments on other semiconductor materials predict that it should also be feasible for GaN.

4 Conclusions

The electronic and optical properties of bilayer GaN structures with rotation angles of 0°, 27.8°, 38.2°, and 46.8° are simulated using density functional theory. According to the results of these first-principles simulations, the bandgap, absorption coefficient, extinction coefficient, reflectivity, and refractive index vary with the rotation angle. The structure formed of two vdW-bonded GaN monolayers is found to exhibit a wide bandgap, which can be further increased by twisting the two layers at certain optimized angles. It is also found that the refractive index of the bilayer structure is much smaller compared with that of its bulk form, greatly reducing the Rayleigh scattering of the optical power. The absorption coefficient is significantly reduced at all twisting angles including 0°. The optical absorption coefficient (\( \sim 10^2 \text{ cm}^{-1} \)) calculated for bilayer GaN is much lower than that of bulk GaN (\( \sim 10^5 \text{ cm}^{-1} \)), reducing the power loss due to the absorption. Such bilayer structures thus offer the advantage of an increased optical bandgap and a reduced optical power loss due to the significant reduction of both the absorption coefficient and refractive index, making these bilayer GaN structures more suitable for monolithic integration of power electronic devices and optical devices. These results will be useful for future integrated low-loss optoelectronics applications.

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