Vertically stacked GaN/WX$_2$ (X = S, Se, Te) heterostructures for photocatalysts and photoelectronic devices

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Tremendous attention has been paid to vertically stacked heterostructures owing to their tunable electronic structures and outstanding optical properties. In this work, we explore the structural, electronic and optical properties of vertically stacked GaN/WX$_2$ (X = S, Se, Te) heterostructures using density functional theory. We find that these stacking heterostructures are all semiconductors with direct band gaps of 1.473 eV (GaN/WTe$_2$), 2.102 eV (GaN/WSe$_2$) and 1.993 eV (GaN/WS$_2$). Interestingly, the GaN/WS$_2$ heterostructure exhibits a type-II band alignment, while the other two stackings of GaN/WSe$_2$ and GaN/WTe$_2$ heterostructures have type-I band alignment. The optical absorption of GaN/WX$_2$ heterostructures is very efficient in the visible light spectrum. Our results suggest that GaN/WX$_2$ heterostructures are promising candidates for photocatalytic water splitting and photoelectronic devices in visible light.

Introduction

Van der Waals (vdW) heterostructures (HS) are used to design unusual electronic devices because of their peculiar physical properties and immense performances.1 Additionally, vertically stacked heterostructures have innovative applications in electronic devices, such as ultrathin photodetectors,2 solar cells,3 memory devices,4 flexible sensors and transistors5,6 etc. Specifically, two-dimensional (2D) vdW heterostructures with type-II band alignments have promising potential in photovoltaics devices and photocatalysts7–11 due to their fascinating electronic and optical properties. In type-II heterostructures, the photo-generated holes and electrons become spatially located in different layers. As a result, the recombination of carriers can be effectively prevented and the light energy utilization is significantly enhanced.12 Moreover, 2D WX$_2$ (X = S, Se, Te) is a typical layered transition metal dichalcogenide family, which has a direct band gap from 1.0 eV to 2.4 eV.13–22 Among the transition metal dichalcogenides (TMD) materials, monolayer WS$_2$, WSe$_2$, and 1H WTe$_2$ are typically semiconductors with direct band gaps of 2.38 eV,18,19 2.11 eV,20,21 1.46 eV.22

Motivated by 2D graphene-like planar honeycomb structures, 2D group III–V compound semiconductors with planar hexagonal structures have been predicted to be stable.23–26 In particular, the monolayer hexagonal structure of gallium nitride (GaN) has recently received growing attractions, which has been directly experimentally synthesized via graphene encapsulation.27 Onen et al. reported that the indirect band gap of 2D GaN is 3.42 eV with HSE06 calculation,28 but the indirect bandgap hindered the practical applications in optoelectronics. In addition, heterostructures based on 2D GaN, such as GaN/blueP, GaN/MoS$_2$, GaN/MoSe$_2$, GaN/WS$_2$, MoSe$_2$/blueP, GaN/ZnO and GaN/BP heterostructures, are able to be formed to type-II band structures, which show good electronic and optical behaviors in photocatalyst and optoelectronic applications.29–33

It is also worth to note that 2D WX$_2$, together with 2D GaN, can form vdW heterostructures for their similar structures and same lattice constants. Generally, the superior structures and excellent properties of GaN/WX$_2$ (X = S, Se, Te) heterostructures are significantly valuable to explore, which could potentially provide a platform for applications in photocatalyst and optoelectronic devices.

Computational methods

Ab initio calculations were performed within the framework of density functional theory (DFT).35–38 as implemented in the Vienna Ab Initio Simulations Package (VASP).35 Electronic exchange and correlation effects were described with the Perdew–Burke–Ernzerhof (PBE)36 functional of generalized gradient approximation (GGA)37 and HSE06 hybrid function.38 Electron–ion interaction were treated with the projector augmented wave (PAW) method,39,40 with an energy cutoff of 550 eV. And Brillouin zone sampling of $K$-mesh was set to 9 x 9 x 1 for relaxation. Subsequently, $K$-mesh points for density of states (DOS) and energy band structures were set to 13 x 13 x 1 and the energy criteria were set as 10$^{-5}$ eV until the residual force was smaller than 10$^{-3}$ eV Å$^{-1}$. Especially, the vacuum layer thickness was set more than 25 Å to avoid spurious interactions in the neighboring images. As the long-range vdW interaction is

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important to hold the 2D heterostructure together, the vdW-D3 approach was used to describe long-range electron correlation effects.

The optical properties of 2D GaN/WX2 heterostructures are described by the complex dielectric function \( \varepsilon(\omega) = \varepsilon_1(\omega) + i\varepsilon_2(\omega) \), where \( \varepsilon_1(\omega) \) and \( \varepsilon_2(\omega) \) are real and imaginary parts, respectively. At the same time, the imaginary part is calculated by summing up all possible transitions from the occupied to unoccupied states, which is closely related to the band structure in the absorption behaviors.\(^{43,44}\) The imaginary part is given by:

\[
\varepsilon_2(\hbar\omega) = \left( \frac{4\pi^2 e^2}{3m^*} \right) \sum_j \left| \langle j | M | i \rangle \right|^2 (1 - f_i^0) \times \delta(E_{jk} - E_{ij} - \omega) d^3k
\]

where \( e, m \) and \( \omega \) are the charge, mass of free electrons, and the frequency of incident photons, respectively. Meanwhile, \( M, i, j \) and \( f_i \) are the dipole momentum, indices of initial and final states, and the Fermi–Dirac distribution function for \( i \)th state with wave vector \( \mathbf{k} \), respectively. The real part \( \varepsilon_1(\omega) \) is obtained from the imaginary \( \varepsilon_2(\omega) \) by Kramers–Kronig relation:\(^{43,44}\)

\[
\varepsilon_1(\omega) = 1 + \frac{2}{\pi} \frac{\omega}{P} \int_0^\infty \frac{\omega' \varepsilon_2(\omega') d\omega'}{\omega'^2 - \omega^2}
\]

where \( P \) is the principal value. Absorption coefficient is then obtained as:\(^{43,44}\)

\[
\alpha(\omega) = \frac{\sqrt{2\omega}}{c} \left[ \sqrt{\varepsilon_1^2(\omega) + \varepsilon_2^2(\omega)} - \omega_1(\omega) \right]^{1/2}
\]

### Results and discussion

#### Stability and structures

Stacking patterns can modulate the electronic properties of the vdW heterostructures. Therefore, four typical stacking configurations (AA, AB, AC, AD) have been constructed. In the AA stacking, the Ga atom sits on the top of S/Se/Te atom, while the N atom is on top of W atom. In the AB stacking, the Ga atom is placed on the top of W atom, while the N atom is on the center of the hexagonal site. In the AC stacking, the N atom sits on the top of W atom, while the Ga atom is located on the center of the hexagonal site. In the AD stacking, the Ga atom sits on the top of W atom, while the N atom is on top of S/Se/Te atom. These relaxed structures are shown in Fig. 1. For GaN/WS2 heterostructure, the interface binding energies of AA, AB, AC and AD stacking patterns are \(-27.72 \text{ meV } \AA^{-2}, -27.01 \text{ meV } \AA^{-2}, -26.69 \text{ meV } \AA^{-2}, \text{ and } -22.58 \text{ meV } \AA^{-2} \) respectively. In consistency with other reports,\(^{27-28}\) after relaxation, AA stacking of GaN/WS2 heterostructures are the most stable due to the lowest interface binding energies. AA stacking patterns are only taken into consideration in the next calculation. The equilibrium lattice constants \( a(b) \), bond lengths, bond angles of GaN monolayer, WX2 monolayer and AA stacking GaN/WX2 heterostructures, calculated with GGA–PBE, are listed in Table 1. From Table 1, the calculated lattice constants, bond lengths/angles accord well with previous results.\(^{13,14,16,21,24,26,28}\) And the relaxed crystal structures of the GaN, WS2, WSe2, and WTe2 monolayers and GaN/WX2 bilayers are hexagonal. It is clear that the hexagonal unit parameters of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are 3.194 Å, 3.285 Å, 3.388 Å, respectively, which agree well with previous results.\(^{28,29}\) The lattice mismatches are expressed as:

\[
\omega = \frac{|a - c|}{a} \times 100\%
\]

where \( a \) and \( c \) are lattice constants of GaN/WX2 heterostructure and individual WX2 layer or GaN layer. The lattice mismatches of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are 1.7%, 3.0% and 3.4%, meaning that the differences are negligible.

To describe the vdW interactions, the interface binding energies (\( \Delta E \)) of GaN/WX2 heterostructures are calculated as:

\[
\Delta E = (E_{\text{HS}} - E_{\text{GaN}} - E_{\text{WX2}})/S_0
\]

![Fig. 1](image_url) Relaxed structures of four typical stacking patterns of GaN/WX2 (X = S, Se, Te) heterostructures. Cyan, yellow, dark blue, and brown spheres indicate W, X, N and Ga atoms, respectively.
WTe2 and GaN/WTe2 heterostructures are direct band gap semiconductors. Both GaN/WX2 heterostructures have type-I band alignment semi-conductors for both the VBM and CBM of GaN/WSe2 and GaN/WTe2 heterostructures come from the W-dz2 orbitals of the WSe2 and WTe2 layer, which can be used in light emitting diodes (LED), indicating that the interlayer interactions between the GaN layer and the WX2 (X = Se, Te) layer reduce the band gap values. Therefore, the band gaps of the GaN/WX2 heterostructures could be significantly changed by vertically stacked heterostructures, which could provide a good opportunity to work on band engineering and electronic device.

Electronic properties

In the following section, projection-resolved band structures of GaN/WX2 heterostructures are depicted in Fig. 3. All the GaN/WX2 heterostructures are direct band gap semiconductors. Both the valence band maximum (VBM) and the conduction band minimum (CBM) of the GaN/WX2 (X = S, Se, Te) heterostructures locate at the K point of Brillouin zone (BZ). The band gaps of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are 3.479 eV, 3.821 eV, 3.386 eV, 3.479 eV, 3.821 eV, 3.386 eV, respectively. The interface binding energies of GaN/WS2, GaN/WSe2, and GaN/WTe2 heterostructures are shifted downwards by 1.80 eV, 5.40 eV, and 6.00 eV, respectively.

![Fig. 2. Plot of interface binding energies as function of interlayer distances for GaN/WS2, GaN/WSe2, and GaN/WTe2 heterostructures. 5th order polynomial has been employed for fitting. In order to clarify the equilibrium distances between the GaN layer and WX2 layer, the values of interface binding energies of both the GaN/WSe2 and GaN/WTe2 heterostructures are shifted downwards by 1.80 eV, 5.40 eV, respectively.](image-url)
designs. It is interesting that the band gap of GaN/WS₂ heterostructure is larger than the minimum required energy of the photocatalytic reaction (1.23 eV), which is potential for a photocatalyst in water splitting.

As reflected in the projection-resolved band structures, the band gap of WX₂ monolayer is decreased by the downshift of CBM as well as the upshift of VBM at K point. The contribution of VBM of GaN/WS₂ heterostructure is GaN layer and that of CBM is WS₂ layer. In Fig. 4(a), the blue and red lines present the DOS of GaN and WS₂ layer in GaN/WS₂ heterostructure. Both the contributions of VBM and CBM of GaN/WSe₂ and GaN/WTe₂ heterostructures are derived from WSe₂ and WTe₂ layer, as shown in Fig. 4(b and c). Obviously, for GaN/WS₂ heterostructure, the VBM and CBM are respectively confined in GaN and WS₂ layer, which also demonstrates that GaN/WS₂ heterostructure has a type-II band alignment.

Photocatalyst and optoelectronic performances

The absorption coefficients of GaN/WX₂ heterostructures are calculated, as illustrated in Fig. 5(a). It is clear that the first peaks in absorption spectrum of GaN/WS₂, GaN/WSe₂, GaN/WTe₂ heterostructures locate at 3.03, 2.69, 2.25 eV, which are mainly contributed by WX₂ layers. The absorption spectrum of GaN/WX₂ heterostructures covers the whole incident solar spectrum, which makes well suitable applications for optoelectronic devices in visible light.

A schematic plot of migrating carriers to depict the dynamic processes of photogenerated carriers at the GaN/WS₂ interface is shown in Fig. 5(b). In the GaN/WS₂ heterostructure, the photogenerated electrons start to transfer from the GaN layer the WS₂ layer in the conduction band, which is motivated by the conduction band offset (CBO) (0.71 eV). Meanwhile, the
photogenerated holes in the valence band move from the WS2 layer to the GaN layer, which are driven by a large valence band offset (VBO) (1.60 eV). Hence, the GaN/WS2 heterostructure, owing a type-II band alignment, could be utilized as a photocatalyst. As for the GaN/WSe2 and GaN/WTe2 heterostructures, one can see that all the photogenerated carriers of the GaN layer are likely to move to the WSe2 (WTe2) layer. The VBO and CBO between GaN and WSe2 (WTe2) layers are 1.98 eV (2.97 eV) and 0.86 eV (1.72 eV), respectively. As a result, the photogenerated carriers can be located in the WSe2 (WTe2) layers to build light-emitting diodes (LEDs). The band edge positions of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are illustrated in Fig. 5(c). The CBMs of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are −4.09, −3.99, −4.26 eV, respectively, whereas the VBM of GaN/WS2, GaN/WSe2, GaN/WTe2 heterostructures are −6.08, −6.09, −5.73 eV, respectively. Those are within the oxidation potential ($E_{O_2/H_2O}^0 = −5.67$ eV) of water and the reduction potential ($E_{H_2O2/H_2}^{0r} = −4.44$ eV). Especially, the redox reaction in the GaN/WS2 heterostructure occurs in different layers: water oxidation occurs in the WS2 layer, meanwhile water reduction takes place in the GaN layer.

It is well known that the lifetime of photogenerated carriers is significantly influenced by the build-in electric field. Hence, build-in electric field induced by ground state charge transfer were investigated. Based on Bader charge analysis, the charge transfer from the GaN layer to the WS2 layer is 0.029e, leading to an electric field pointing to GaN from WS2. This electric field will promote the migration of electrons from the GaN layer to WS2 layer and holes from the WS2 layer to GaN layer. The build-in electric field $E = \frac{P}{\epsilon_0 \epsilon_s S_{\text{eq}}}$ of $3.69 \times 10^3$ V m$^{-1}$ is acquired in the junction region of GaN/WS2 heterostructure. Where $P = \Delta q \cdot d_{\text{eq}} = 0.089e \cdot \text{Å}$ is the size of dipole moment, $\epsilon_0 = 8.850 \times 10^{-12}$ F m$^{-1}$ refers to the permittivity of free space, $\epsilon_s = 1$ represents the relative dielectric constant, $S_y = 8.835 \times 10^{-20}$ m$^2$ is the interfaces area of the WS2 layer, and $d_{\text{eq}} = 3.081 \times 10^{-10}$ m denotes the interlayer equivalent distance.

**Conclusions**

In summary, the electronic and optical properties of GaN/WX2 heterostructures have been performed within density functional theory. It is found that all the hexagonal GaN/WX2 heterostructures are stable semiconductors with direct band gaps. Interestingly, among them, the GaN/WS2 heterostructure has a type-II band alignment, which is beneficial in photocatalysts for water splitting. However, both the GaN/WSe2 and GaN/WTe2 heterostructures exhibit type-I band alignment, which can be utilized in LED. In the GaN/WS2 heterostructure, the charge transport from the GaN layer to the WS2 layer is 0.029e, based on Bader charge analysis, forming a build-in electric field favorable for charge separation. The build-in electric field $E = \frac{P}{\epsilon_0 \epsilon_s S_{\text{eq}}}$ of $3.69 \times 10^3$ V m$^{-1}$ is acquired in the junction region of GaN/WS2 heterostructure. The absorption behaviors of GaN/WX2 heterostructures are also performed, and results show that GaN/WX2 heterostructures are very efficient at absorbing the visible light to expand the application to photocatalyst and photoelectronic devices.

**Conflicts of interest**

There are no conflicts to declare.

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