Interface excitons at lateral heterojunctions in monolayer semiconductors

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We study the interface exciton at lateral type II heterojunctions of monolayer transition metal dichalcogenides (TMDs), where the electron and hole prefer to stay at complementary sides of the junction. We find that the 1D interface exciton has giant binding energy in the same order as 2D excitons in pristine monolayer TMDs although the effective radius (electron-hole separation) of interface exciton is much larger than that of 2D excitons. The binding energy, exciton radius and optical dipole strongly depends on the band offset at the junction. The inter-valley coupling induced by the electron-hole Coulomb exchange interaction and the quantum confinement effect at interface of a closed triangular shape are also investigated. Small triangles realize 0D quantum dot confinement of excitons, and we find a transition from non-degenerate ground state to degenerate ones when the size of the triangle varies. Our findings may facilitate the implementation of the optoelectronic devices based on the lateral heterojunction structures in monolayer semiconductors.

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I. INTRODUCTION

Heterostructures between conventional three-dimensional (3D) semiconductors has inspired the inventions of the modern electronic devices such as high speed transistor, 1 diode laser, 2 light-emitting diode, 3 and solar cell. 4 Thanks to the development of nanotechnology, we are able to engineer heterostructures on the nanoscale for high-speed opto-electronic devices. In III-V and II-VI semiconductors, various nanoscale heterostructures such as quantum wells, superlattices, and core-shell nanodots and nanowire have been widely studied. 5–7 Emerged as a new class of semiconductors in the two-dimensional (2D) limit, monolayers of group-VIB transition metal dichalcogenides (TMDs) possess visible range direct gap, exotic properties associated with valley degeneracy, and new geometries for realizing various heterostructures, which provide new platforms to study the physics and applications at semiconductor heterostructures. 8–10 By stacking different TMDs monolayers which are then bound together by the weak Van der Waals forces, vertical heterostructures have been realized recently, e.g. MoX2/WX2 (X=Se, S) heterobilayers 11,12 which can be analogs of the III-V semiconductor double heterojunctions.

Besides the vertical heterostructures, two-dimensional materials also make possible heterostructures of a unique planar geometry. Two different TMDs seamlessly connected in a single monolayer has been realized experimentally already. 13–15 A more recently development is on the growth of various lateral heterostructures, multi-heterostructures and superlattices for TMDs. 16 The possibility to form atomically sharp and straight lateral interface of different compounds 17–24, 35 point to exciting opportunities towards device applications based on the lateral heterojunctions, as well as a new geometry to realize quantum wires and even quantum dots in the monolayer semiconductors. The lateral heterojunctions can also be realized in an alternative way, by electrostatic gating to define lateral p-n junctions 16,17. The recent development shows that the width of the electric gate in the monolayer MoS2 can be narrowed down to 1nm by using a single-walled carbon nanotube as the gate electrode. 25

In most of the vertical and lateral heterostructures formed between different TMDs monolayers, they feature a type-II band alignment, where the conduction and valence band edge locate in different TMDs. The strong Coulomb interaction binds electron and hole to form exciton at the interface. In contrast to 2D exciton formed in pristine monolayer TMD, the electron and hole at the interface will be spatially separated because of the type-II band alignment, and such an interface exciton can have lower energy, being an excitonic ground state in the heterostructures. The properties of such interface excitons can be essential to determine the optical response of the lateral heterostructures of TMDs. In vertical heterojunctions MoX2/WX2 heterobilayers, such interface exciton has already been investigated theoretically and experimentally. 26–28 Due to the spatial separation of electron and hole, interlayer excitons in MoX2/WX2 heterobilayers have shown long lifetime exceeding nanoseconds 29–32 and electro-statically tunable resonance 33 which are highly desirable for the realization of excitonic circuits and condensation 34–36. And interestingly, the inevitable twisting and lattice mismatch in the heterobilayers can give rise to novel light coupling properties. 17,18

Albeit the novel and appealing properties discovered, the interface excitons in the heterobilayers of 2D semiconductors are analogues of those in the conventional heterostructures bulk semiconductors, for example the spatially indirect excitons in III-V double quantum well. The realization of lateral heterostructures in monolayer TMDs opens up new opportunity to extend the study of interface exciton from two-dimensional interface to the one-dimensional (1D) interface. The 1D interface exciton mode may shed light on novel optoelectronic devices.
based on these atomically thin 2D lateral heterostructures. Moreover, such 1D interface excitons may also become relevant in lateral p-n junctions in monolayers TMDs.\textsuperscript{38–40}

Here, we theoretically study the interface exciton states at lateral heterojunctions of the monolayer TMDs. The physical properties of one-dimensional type-II interface exciton such as the binding energy, exciton radius (i.e. electron-hole separation), longitudinal-transverse splitting by the electron-hole exchange and optical dipole are investigated as a function of band offset at the interface. We adopted two different approaches to calculate the interface exciton states. One approach bases on a real-space tight binding (TB) model, and the other approach uses the perturbation expansion in a hydrogen-like basis ineffective mass approximation. The numerical study shows with the increase of the band offset at the interface, the exciton radius grows and can become several times larger than that of the 2D excitons in homogeneous monolayer TMDs. In the meantime, the decrease in the exciton binding energy is not as significant, remaining in the same order as the 2D exciton, because of the weaker screening of Coulomb interaction as electron-hole separation increases. Due to spatial indirect nature of interface exciton, the optical transition dipole decreases fast with the increase of band offset, which, at a typical band offset of ~300 meV, is about one order of magnitude smaller than that of 2D exciton. We also investigated lateral heterostructures with a closed triangular shaped interface which effectively realize a 0D quantum dot confinement of exciton. Such quantum dot uniquely features the quantum confinement of one carrier by the band offset of the interface, and binding of the other carrier in the proximity exterior by the strong Coulomb. We find two distinct scenarios of energy level schemes and valley optical selection rules of the interface exciton at small and large quantum dot size respectively, which can be exploited for optical quantum controls.

The paper is organized as follows. In section II, we introduce the Hamiltonian of the exciton of lateral structures in the effective mass approximation. We study the interface exciton at the 1D p-n and p-n-p heterojunctions of monolayer semiconductors in Sec. III. The numerical results of the physical observables of interface exciton are also shown in section III. In Sec. IV, we show the numerical calculation of the interface exciton at the 0D quantum dot type triangular lateral heterostructure. We conclude in section V.

II. HAMILTONIAN IN THE EFFECTIVE MASS APPROXIMATION

The three-band model involving all d orbitals of the transition metal atom is usually applied to describe the single electron in the monolayer TMDs associated with valley index.\textsuperscript{38} In the low energy excitation limit where only the electron in the vicinity of the valance band edge is excited by light field to the vicinity of the conduction band edge, both the electron in the conduction band and the hole left in the valcence band can be approximately described by the effective mass model. In this sense, the periodic parts of the electron and hole Bloch wavefunctions are omitted and only the profiles of the electron and hole Bloch wavefunctions are taken into consideration in the following discussion. Together with the attractive Coulomb interaction and the lattice potentials, the type II interface exciton at the interface can be described by the following Hamiltonian

$$H = -\frac{\hbar^2}{2m_e} \nabla^2_r e - \frac{\hbar^2}{2m_h} \nabla^2_r h + V_C(|r_e - r_h|) + V_e(r_e) + V_h(r_h),$$

(1)

where $m_e$ ($m_h$) is the electron (hole) effective mass, and $r_e$ ($r_h$) denotes the position coordinates of the electron (hole). The lattice potentials of electron and hole $V_e(r_e)$ and $V_h(r_h)$ depends on the different geometries of the lateral heterostructures.

Here, the Coulomb interaction $V_C(|r_e - r_h|)$ between the electron and hole in the 2D limit reads\textsuperscript{51}

$$V_C(r) = -\frac{e^2\pi}{2\alpha_0} \left( H_0\left(\frac{r}{\alpha_0}\right) - Y_0\left(\frac{r}{\alpha_0}\right) \right), \tag{2}$$

where $H_n$ and $Y_n$ denote Struve Function and Bessel Function of the Second Kind respectively. The former researches demonstrated that in monolayer TMDs the quasi-2D geometry leads to a distance-dependent effective dielectric screening.\textsuperscript{52–54} For monolayer TMDs, the parameter $\alpha_0$ is in the order of a few nm, which is comparable to the Bohr radius of a free 2D exciton.\textsuperscript{52–54}

III. INTERFACE EXCITON AT 1D P-N AND P-N-P HETEROJUNCTIONS

A. Type II interface in monolayer TMDs

A lateral type-II interface in monolayer TMDs can be implemented in two setups. The first is a lateral hetero-junction seamlessly formed between different TMDs,\textsuperscript{20,22} as shown in Fig. 1(a) and Fig. 1(d). In such cases, the type-II interface is atomically sharp. The conduction and valence band edges as functions of position are regarded as the step functions. The other setup shown in Fig. 1(b) and Fig. 1(e) is a lateral p-n or p-n-p junctions electrostatically created in a monolayer TMD by separate back gates, which has been studied experimentally.\textsuperscript{55–59} Such setup realizes a gentle type-II interface with a finite width of the interfaces $w$.

We are interested in the binding energy and wavefunction of the interface exciton ground state, which determines the stability and optoelectronics properties of the interface exciton. By the interface potentials, electron and hole prefer to stay at complementary sides of the interface, while the Coulomb interaction $V_C(|r_e - r_h|)$ attempts to bind the electron and hole. The properties
of the interface exciton therefore depends on the competition of the band offset and the Coulomb interaction, which are then tunable by the width $w$ of the interface and the magnitude of the conduction and valence band edge offsets $V_0$. The effective dielectric screening of Coulomb interaction varies with distance, and for large distance between electron and hole the screening effect is substantially reduced. As we will show, this is important for the interface exciton to have strong binding energy, even though the spatial separation between the electron and hole is much larger than the Bohr radius of the 2D exciton.

B. Solving the eigen problem using Bohr-Oppeheimer Approximation

Since $V_i(r_x,r_h) = V_e(r_x) + V_h(r_h)$ possesses translational symmetry while Coulomb interaction possesses rotational symmetry, incompatible symmetries make it impossible to obtain analytical solutions for the Schrödinger equation governed by the Hamiltonian in Eq. (1). We rewrite the above Hamiltonian with the center-of-mass motion and relative motion of the electron-hole pair as

$$H = -\frac{\hbar^2}{2M} \nabla^2_R - \frac{\hbar^2}{2\mu} \nabla^2_r + V_C(r) + V_i(R,r),$$  \hspace{1cm} (3)

where the center-of-mass and relative space coordinates are

$$\begin{aligned}
R &= \frac{1}{M} (m_e r_x + m_h r_h) \\
\mathbf{r} &= r_x - r_h
\end{aligned}$$  \hspace{1cm} (4)

with total mass $M = m_e + m_h$ and reduced mass $\mu = m_e m_h / (m_e + m_h)$. Due to the 2D nature of TMDs, these coordinates only have two components which means $R = (X,Y)$ and $\mathbf{r} = (x,y)$. Obviously, the total mass is at least four times greater than the reduced mass ($M \geq 4\mu$), which implies that the center-of-mass motion is a relatively slow one in comparison with the relative motion. Under this circumstance, we can apply the Bohr-Oppeheimer Approximation (BOA) here and in zeroth order BOA the eigen-wavefunction is a product state as

$$\Phi (\mathbf{R},\mathbf{r}) = \Psi (\mathbf{R}) \Theta (\mathbf{R},\mathbf{r}).$$  \hspace{1cm} (5)

For lateral heterojunctions $V_i(r_x,r_h)$ possesses trans-lational symmetry along $y$-direction as shown in Fig. 1(a) and Fig. 1(d). The interface potential is numerically modelled with $V_e(x_e) = \frac{V_0 + \delta}{2}(1 - \tanh (\frac{x_e}{w}))$ and $V_h(x_h) = -\frac{V_0 - \delta}{2}(1 - \tanh (\frac{x_h}{w}))$, where $w$ is the width of the interface which characterizing the sharpness of the band offset. As $V_i(r_x,r_h)$ is independent of $Y$, the envelope function remains to be a plane wave in $Y$-direction, so we rewrite the center-of-mass motion part as $\Psi (\mathbf{R}) = \Psi (X) e^{iP_Y Y}$. Since $P_Y$ stands for the $y$-component wave vector corresponding to a kinetic energy $\hbar^2 P_Y^2 / 2M$, obviously $P_Y = 0$ for the ground state of type-II interface exciton. Then the corresponding Schrödinger’s equations for the relative motion and center-of-mass motion read

$$\left[-\frac{\hbar^2}{2\mu} \nabla^2_r + V_C(r) + V_i(X,r)\right] \Theta (X,r) = E(X) \Theta (X,r),$$  \hspace{1cm} (6)

$$\left[-\frac{\hbar^2}{2M} \nabla^2_X + E(X)\right] \Psi (X) = E_g \Psi (X).$$  \hspace{1cm} (7)

The energy $E(X)$ plays the role of an effective potential in Eq. (7) which leads to the ground state $\Psi (X) \Theta (X,r)$ of type-II interface exciton with corresponding ground state energy $E_g$. To numerically solve Eq. (6), we adopted two different approaches: one is the solution based on a real-space tight binding model for the relative part Hamiltonian $H_r = -\frac{\hbar^2}{2\mu} \nabla^2_r + V_C(r) + V_i(X,r)$, and the other is a perturbative expansion of $H_r$ with a hydrogen-like basis of the effective mass model. Details of both approaches can be found in the numerical results section.

FIG. 1. (Color online) (a) (d) Schematics of the single- and double- heterojunctions formed between MoSe$_2$ and WS$_2$. The blue, red and yellow spheres respectively denote W, Mo and Se atoms. The purple shadow regions denotes the type II interfaces. (b) (e) Schematics of the monolayer TMDs p-n junction defined by the gate voltage. The red and blue regions denote the n and p regions, respectively. (c) (f) Illustration of the band edge profile across the single lateral junction in (a) or (b), and the double lateral junctions in (d) or (e). Here, $w$ is the width of the interface, which is zero for the case of (a) and (d), and has a finite value for the gate defined junctions in (b) and (e). $V_0$ is the average lattice potential of electron and hole in (a) or (d) and gate voltage in (b) or (e). Here, $\delta$ characterizes the difference of the band offsets for electron and hole.
C. Physical observables and the electron-hole overlap

Before we present the numerical results we would like to introduce several important physical observables first. When applying several BOA and obtaining the eigen-wavefunction of type-II interface excitons, we can straightforwardly calculate the binding energy, effective radius, optical dipole and the intervalley coupling of the interface excitons.

The binding energy of type-II interface exciton is defined as $E_b = E_f - E_q$, where $E_f$ is the energy of a non-interacting electron-hole pair at the interface. The effective radius is straightforwardly calculated as

$$ R_0 = \sqrt{\frac{\hbar}{2t}} \int dR dX r^2 |\Psi(X, r)|^2, \quad (8) $$

which measures the spatial separation between the electron and hole.

Another important observable is the optical dipole defined as

$$ D = \left| D_0 \int dX \Psi(X, r) \Theta(X, 0) \right|, \quad (9) $$

which relates to the lifetime of type-II interface exciton in TMDs. Here, $D_0$ is the interband transition dipole element between the conduction band and valence band for 2D excitons. In contrast with the optical dipole of 2D excitons, there is an additional integral over $X$ direction whose value depends on the wavefunction profile along $X$ direction. Additionally, $r = 0$ in the relative part of the wavefunction $\Theta(X, 0)$ indicates that the recombination of the electron and the hole in the exciton occurs only when they exactly locate at the same position. In this sense, the electron and hole will become harder to recombine with each other and thus results in a longer lifetime due to the decreased optical dipole. As we will show that below the amount of overlap between the electron and hole can be controlled by $V_i(R, r)$.

As we can see, the electron-hole overlap plays an important role in Eq. (6). For a ground state of the 2D exciton, it closely resembles s-orbitals, so a very large optical dipole $D_0$ is expected. However for large $V_0$ the wavefunction overlap between the electron and hole is greatly reduced. Besides the separation of the electron-hole pair and optical dipole $D$, the electron-hole overlap also affects other properties of the type-II interface exciton such as intervalley coupling induced by Coulomb exchange interaction.

It was proved that under broken 3-fold rotation symmetry in a monolayer TMDs, the excitonic spectrum could have a finite valley exchange interaction even in the ground state which is induced by exchange Coulomb interaction between electrons and holes. In the presence of $V_i(R, r)$, translational symmetry is only preserved in the $y$-direction, breaking the 3-fold rotation symmetry. Thus lateral heterojunctions not only decrease the electron-hole overlap, but also results in a non-vanishing valley-exchange term $J$. Such term opens a coupling channel between $+K$ and $-K$ valleys which is normally suppressed in monolayer TMDs due to large momentum difference. In a quasi-1D system, the intervalley coupling strength is written as

$$ J = \left( \frac{a t}{E_g} \right)^2 \sum_{P_X} V_C(P_X, P_Y = 0) P_X^2 |\psi(P_X)|^2, \quad (10) $$

where $a$ is the lattice constant, $t$ is the hopping constant, and $V_C(P_X, P_Y)$ is Coulomb interaction in the momentum space. Here, $\psi(P_X) = \sum_X \Theta(X, r = 0) \Psi(X, P_X)$ is the electron-hole overlap in the $X$-component momentum space. Since the electron-hole overlap is controlled by the strength of the band offset, a tunable intervalley coupling is expected in the lateral heterojunction in TMDs.

It is important to note that the above described zeroth-order Bohr-Oppenheimer Approximation is valid only in the adiabatic limit where the gradient of the band offset caused transition probability is much smaller than the energy level spacing between the ground state and any excited state in Eq. (6). Detailed justification shall be referred to the appendix or literature about generalized Bohr-Oppenheimer Approximation (BOA). For the eigen-problem of type-II interface exciton in TMDs, we will numerically justify that the zeroth order BOA is sufficient.

We will take MoSe$_2$/WSe$_2$ heterojunctions as our example in subsequent sections of type-II interface excitons. It is trivial to generalize our method to other sharp TMDs lateral interfaces.

D. Numerical Results Based on TB Model

In order to obtain the TB model, we discretize Eq. (6) in the real space. We take unit in $x$-direction as $a$ and $y$-direction as $\sqrt{3}a$, where $a = 3.325\text{Å}$ is the lattice constant. The lattice constant of WSe$_2$ and MoSe$_2$ closely matches so it is legitimate to assume the same lattice constant across the heterojunctions. A $72 \times 84$ supercell and the open boundary conditions for both directions are taken into consideration. We consider an armchair interface in the following calculation, while it will give almost the identical results when changing the armchair edge to a zigzag one. Previous studies show that the conduction and valence bands are accurately described by $d$-orbitals of the metal atoms, while the orbits of the chalcogenides play a minor role. Hence we only consider the metal atoms in our TB model, and the nearest-neighbour hopping $t = -\hbar^2/3a^2\mu$ between metal atoms. The one of the advantages of applying TB model is that $V_C(r)$ and $V_i(R, r)$ are exactly diagonalized. It is noticed that the on-site electrostatic energies $U = V_C(r = 0)$ is divergent. Since for a type II alignment the electron and hole barely can occupy the same
site, a large value of the on-site electrostatic energies $U$ is assumed in our calculation in order to make the calculation convergent.

The effective masses of the electron and hole are chosen as $m_e = m_h = 0.32m_0$ and thus the reduced mass is $\mu = 0.16m_0$ with free electron mass $m_0$. The width of the interface is chosen as $w = 0.1a$ to model a very sharp band offset in order to simulate the single- or double-heterojunctions. Here, $r_0$ in the effective Coulomb interaction is chosen as $r_0 = 75 \text{ Å}$. A symmetric heterojunction ($\delta = 0$) is considered unless otherwise specified.

By solving Eq. (7), the binding energy $E_b$, effective radius $a_b$ and optical dipole $D$ versus different strength of the band offset $V_0$ are shown in Fig. 2. We also depict those physical observables when different on-site electrostatic energies $U$ are chosen. The red sphere, blue triangle and magenta diamond symbols respectively represent $U = -0.79, -1.19$ and $-2.98 \text{ eV}$. The physical observables converge to the same value at high voltage $V_0$ regardless of $U$, which actually implies that at large $V_0$ the electron and hole are well separated and thus there is almost no on-site electrostatic energy contribution in $E_b$. Basically there are two characteristic behaviours, the regime of small band offset ($V_0 < 0.1 \text{ eV}$) and large band offset ($V_0 > 0.4 \text{ eV}$). This reflects the competition between Coulomb interaction and the band offset. For small band offset $V_0$, $V_C(r)$ dominates over $V_I(R, r)$ so the exciton ground state is almost equivalent to a 2D exciton while $V_I(R, r)$ is regarded as a perturbation term. Therefore, the physical observables of type-II interface exciton are almost the same as the the ones for 2D exciton in this regime. However, for sufficiently large $V_0$, $V_I(R, r)$ dominates over $V_C(r)$. The effective radius $a_b$ shows a rapid rise while $D$ drops dramatically as the band offset increases. In this sense, we can control physical properties of type-II interface exciton by adjusting the band offset.

We obtained an interface exciton binding energy of about $0.2 \text{ eV}$ which is of the same order as a 2D exciton in TMDs. Such a large binding energy at a type-II interface is not present in most conventional semiconductor nanostructures. In fact, a type-II interface exciton is often considered unstable in conventional semiconductor heterostructures unless in the presence of other physical structures like a E-field or within a quantum dot. In a TMD lateral heterojunction, however, with a relatively large binding energy a type-II interface exciton is predicted to be stable with our calculations. It is also of concern whether the interface exciton changes back to a 2D exciton easily. From Fig. 2(b) we see that at $V_0 = 0.2eV$ the binding energy of the interface exciton is about $0.22eV$, which is about $0.1eV$ smaller than the binding energy of 2D exciton. In this sense, we may assume that for realistic configurations the interface exciton is a stable ground state of a lateral heterojunctions. The relatively large binding energy exactly results from the weaker screening effect of Coulomb interaction when the electron-hole separation increases as shown in Eq. (2).

It is noted that for a large $V_0$ the on-site Coulomb interaction for an interface exciton is irrelevant. This implies that the electron and hole are well-separated into opposite regions for sufficiently large $V_0$, while for small and intermediate $V_0$, even though qualitative behaviours are similar, numerical values obtained with different $U$ are quite different. Without loss of generality, $U = -0.79 \text{ eV}$ is assumed in the remainder of the paper as it gives the closest free exciton $E_b$ with Ref. [50] for 2D exciton.

The center-of-mass part $\Psi_g(X)$ and relative motion wavefunction with fixed electron position are respectively depicted in Fig. 3(a) and (b) for different $V_0$. From Fig. 3(a) we see how $\Psi_g(X)$ varies from $V_0 = 0$ to $0.4 \text{ eV}$. For small band offset $V_0 < 0.1 \text{ eV}$ $\Psi_g(X)$ is widespread across the supercell. This is expected since for small $V_0$ the electron-hole pair behaves as a 2D exciton. But for sufficiently high $V_0$, $\Psi_g(X)$ is localized around the interface at $X = 0$. The center-of-mass part of the wavefunction undergoes a transition from a plane wave to a localized state as $V_0$ increases, demonstrating the competition between the Coulomb interaction and the band offset. From Fig. 3(b), the biased relative motion wave-
function for large $V_0$ implies that the electron-hole pair tends to be separated apart well and thus electron-hole overlap is greatly reduced. To further demonstrate the separation nature of the type-II interface exciton, the reduced wavefunction of the electron and hole versus separation is shown in Fig. 4(a), where obviously the electron preferentially stays at left hand side of the interface while the the hole stays at the right hand side of the interface.

There is still a considerable optical dipole because of the tunnelling tail of the electron and hole reduced wavefunction. We find that a small but notable electron-hole overlap still survives. Such electron-hole overlap can be schematically demonstrated by Fig. 4(b). The finite magnitude of overlap for $V_0 > 0.1$ eV implies that the optoelectrical properties can be still detected for interface exciton. From Fig. 2(d) we see that at $V_0 = 0.2$ eV $D$ only drops by half that of 2D excitons, and by one order of magnitude at 0.3 eV. Thus at such $V_0$ interface exciton still can be excited by the pumping light. On the other hand, reduction of $D$ suggest a longer lifetime. For very large $V_0$, $D$ is a few orders smaller than 2D excitons, meaning that interface exciton may have a lifetime far exceeding 2D excitons.

Finally we calculated the inter-valley coupling strength $J$ for different $V_0$ at $U = -0.79$ eV for both a symmetric interface with $\delta = 0$ and an asymmetric interface with $\delta = 0.5$ eV in Fig. 5. For the symmetric case, it is expected that $J$ tends to zero for $V_0 = 0$ due to the emergence of 3-fold rotation symmetry. When $V_0$ increases, broken symmetry results in a dramatic increase of inter-valley coupling. However for an asymmetric interface with $\delta = 0.5eV$, the 3-fold rotation symmetry is broken at the beginning and thus there is a considerable $J$ at $V_0 = 0$. However as $V_0$ further increases, the broken symmetry plays a minor role and a very similar monotonic decreasing behavior in $J$ is observed for both interfaces. This manifests the reduced electron-hole overlap $\psi(P_X)$ as in the drop of $D$ when $V_0$ increases.

A non-zero inter-valley coupling between $\pm K$ implies that interface exciton ground state has a valley part of the form $\frac{1}{\sqrt{2}} (|K| \pm |-K|)$. This suggests that interface exciton couples with linearly polarized light instead of circularly polarized light as in 2D excitons which is shown in Fig. 7(b). Our calculations shows that $J$ has an order...
of a few meV in Fig. 5(c).

Under current parameters, we have numerically evaluated the first and second order terms of a more rigorous Generalized BOA\cite{58,59} and find that even within the intermediate regime of $V_0$, the corrections terms are in the order $10^{-6}$ eV, which is much smaller than the energy level spacing in the order of $10^{-1}$ eV. Thus the correction terms may safely be neglected and the zeroth-order BOA is sufficient for current circumstance.

E. Numerical Results Based on Continuous Model

When the size of the supercell is much larger than the lattice constant, we can also introduce a continuous model of type II interface exciton, where its Hamiltonian in Eq. (6) is diagonalized with a 2D hydrogenic basis. In the Hilbert space expanded by the 2D hydrogenic basis $\{ \phi_{nl} (r) \} (n = 0,1,\ldots,l = -n+1,\ldots,n-1)$ which satisfy the Schrödinger equation of the usual 2D hydrogen atom\cite{56}

$$\left(-\frac{\hbar^2}{2\mu}\nabla^2 + \frac{e^2}{\varepsilon r} \right) \phi_{nl} (r) = E_n \phi_{nl} (r) , \quad (11)$$

Eq. (6) can be rewritten as

$$a_{nl}^k (X) [E (X) - E_k (X)] + \sum_{n' = 1}^{\infty} \sum_{l' = -n+1}^{n-1} V_{n'n'}^{nl} (X) a_{nl}^{k*} (X) = 0 , \quad (12)$$

where $\Theta_k (X,r) = \sum_{n,l} a_{nl}^k (X) \phi_{nl} (r)$ has already been assumed as the linear combination of the basis with coefficients $\{ a_{nl}^k (X) \}$ and the elements of the electric potential are defined as $V_{n'n'}^{nl} (X) = \int d\mathbf{r} \phi_{nl}^* (\mathbf{r}) \left[ V_C (r) + V_I (X,r) \right] \phi_{n'l'} (\mathbf{r})$.

Since we need an infinite principal quantum number $n$ to complete the Hilbert space of Eq. (6) which is clearly impossible, we need to set a cutoff $n$ when both the binding energy and wavefunction of the ground state interface exciton are convergent. We plot the binding energy versus the principal number $n$ for different potential strength $V_0$ in Fig. 6. It is obvious that the ground state energies converge very quickly along with the principal quantum number even for relatively large potential strength. In the following discussion, the cutoff of $n$ is

![Figure 5](image1.png)

![Figure 6](image2.png)
set as \( n_{\text{cutoff}} = 7 \). It is also important to note that the dielectric constant \( \varepsilon \approx 1.10\varepsilon_0 \) for the 2D hydrogenic basis is fixed in the above numerical calculation in order to obtain the same binding energy \( E_b \approx 220\text{meV} \) as the one from TB model at \( V_0 = 0.2\ \text{eV} \). Here, \( \varepsilon_0 \) is the vacuum dielectric constant. In this sense, the binding energy of the 2D exciton is \( E_b = 396\text{meV} \). The other parameters are \( m_e = 0.434m_0, m_h = 0.533m_0 \) with free electron mass \( m_0 \). Based on the continuous model the numerical results of the binding energy \( E_b \), effective radius \( a_b \) and optical dipole \( D \) obtained with different strength of the band offset \( V_0 \) are shown in Fig. 7 as the blue triangle symbols and solid lines. Here, the numerical results based on the TB model is also shown in the same figure as the red sphere symbols and dashed lines. The numerical results especially the energies resembles each other reasonably, which implies the validity of both methods. For large band offset \( V_0 \), the difference between the numerical results of both methods becomes greater because the size of the supercell we chose is not sufficiently large.

F. Interface exciton at lateral double heterojunctions

The former discussion focuses on the properties of the 1-D interface exciton at single heterojunction as shown in Fig. 2(a). Another important case is the lateral double heterojunctions as shown in Fig. 2(d). When the interface exciton is generated in such structure, the electron is supposed to locate at the central region and the hole is supposed to locate at both hands side of the central region due to the lattice potential. However, because of the Coulomb interaction the electron and the hole have tendency to bind each other. Such competition will affect the properties of the interface exciton greatly. Since the lattice potential depends on the width of the double heterojunctions as well as the potential strength now, we calculate the binding energy of the interface exciton versus the width of the double heterojunctions \( L \) and the potential strength \( V_0 \) by applying the TB method.

The binding energy versus the width of the double heterojunctions \( L \) is depicted in Fig. 8(a). The \( V_0 \) is chosen as \( 0.5\text{eV} \) in Fig. 8(a). The other parameters are chosen the same as those in Fig. 2. As shown in Fig. 8(a), the binding energy increases as the width of the double heterojunctions increases and eventually saturates to a constant value which is the binding energy of the 1D exciton shown in Fig. 2(a). It can be interpreted by the overlap of the 1D excitons locating at both interfaces. The reduced wavefunction of electron and hole for different widths of the double heterojunctions \( L \) are respectively depicted as blue solid lines and red dashed lines in Fig. 8(b). The typical effective radius of the 1D exciton for potential strength \( V_0 = 0.5\text{eV} \) is around \( 5\text{nm} \) as shown in Fig. 2(c). For a small double heterojunction...
tions with \(L < 5nm\), the lattice potential dominates the binding energy of the interface exciton. The consisting electron in 1D excitons locating at both interfaces has great overlap which results in that the electron can only locate at the very center of the double heterojunctions. When the double heterojunctions width increases to be larger than the typical effective radius of the 1D exciton such as \(L > 5nm\), the Coulomb interaction becomes dominating and the electron prefers to locate in the vicinity of the each interfaces which actually reduces the overlap of the electrons wavefunction. When the width \(L\) is much larger than the effective radius, the overlap tends to zero which results in the saturated value equaling to the binding energy of the 1D exciton.

The binding energy versus the potential strength \(V_0\) is depicted in Fig. 8(c), where \(L\) is chosen as 1.3nm. The binding energy decreases when the potential strength \(V_0\) increases. The reason is that the effective radius of the 1D interface exciton become larger as the potential strength \(V_0\) increases as shown in Fig. 8(b). Therefore the overlap of the electron wavefunction becomes smaller and eventually reduce the binding energy.

IV. INTERFACE EXCITON AT CLOSED TRIANGULAR SHARP INTERFACE AND EFFECTIVE QUANTUM DOT CONFINEMENT

A. Numerical results of closed triangular sharp interface without valley index

For all the discussion above, we have assumed the quasi 1D heterojunction as shown in Fig. 1. However, the realistic lateral heterostructures for TMDs present the triangular shape, whose characteristic length scale is about 5\(\mu\)m. Usually, the electron-hole separation of interface exciton is up to 10\(\mu\)m for large \(V_0\) from the above calculation of 1D interface. It is much smaller than the characteristic length scale of triangular heterostructures, which means the calculation of 1D interface is also valid for the closed triangular sharp interface in current experiments.

If the characteristic length scale of closed triangular sharp interface decreases to the same order of the electron-hole separation of interface exciton, the electron (hole) wavefunction will strongly affected by the boundaries of the triangular shape and thus such quantum confinement effect should be taken into consideration. Actually, such closed triangular interface effectively realizes 0D quantum dot confinement of the interface excitons. From the similar Hamiltonian in Eq. 6 but with triangular band offset as shown in Fig. 9, which reads

\[
V_e(r) = \begin{cases} 
0, & r \in \text{triangular quantum dot}, \\
V_0, & r \notin \text{triangular quantum dot}, 
\end{cases} 
\]  

\[
V_h(r) = \begin{cases} 
0, & r \in \text{triangular quantum dot}, \\
0, & r \notin \text{triangular quantum dot}. 
\end{cases} 
\]  

Here, \(L_{SC}\) and \(W_{SC}\) are the length and the width of the supercell adopted in the calculations in units of lattice constant \(a\), and \(R\) is the edge length of the regular triangular quantum dot.

Since the translational symmetry is no longer preserved in such closed triangular sharp interface, we need to develop another numerical method to calculate the physical properties of interface exciton. The complete orthonormal basis \(\{\phi_{e}^{(n)}(r_e) \otimes \phi_{h}^{(n,m)}(r_h)\}\) are introduced to expand the original Hamiltonian, where \(\phi_{e}^{(n)}(r_e)\) is the \(n\)-th eigen-state of the electron confined in the triangular region without hole part such as

\[
\left[ \frac{-\hbar^2}{2m_e} \nabla^2 + V_e(r_e) \right] \phi_{e}^{(n)}(r_e) = E_{e}^{(n)} \phi_{e}^{(n)}(r_e),
\]  

and the \(\phi_{h}^{(n,m)}(r_h)\) is the \(m\)-th eigen-state of the hole effective Hamiltonian \(H_{eff}(r_h) \phi_{h}^{(n,m)}(r_h) = E_{h}^{(n,m)} \phi_{h}^{(n,m)}(r_h)\), where the effective Hamiltonian of hole is obtained by averaging the original Hamiltonian on \(\phi_{e}^{(n)}(r_e)\) as

\[
H_{eff}(r_h) = \int d r_e \phi_{e}^{(n),*}(r_e) H \phi_{e}^{(n)}(r_e). 
\]  

The Hamiltonian matrix elements are straightforward calculated as

\[
H_{n,m,n',m'} = \int d r_e \int d r_h \phi_{e}^{(n'),*}(r_e) \phi_{h}^{(n,m)}(r_h) H \phi_{e}^{(n,m)}(r_h),
\]  

which can be simplified according to the orthogonality of the basis as

\[
H_{n,m,n',m'} = \begin{cases} 
E_{e}^{(n)} + E_{h}^{(n,m)}, & \text{if } n = n' \text{ and } m = m', \\
0, & \text{if } n = n' \text{ and } m \neq m', \\
V_C(n, n', m, m'), & \text{if } n \neq n',
\end{cases}
\]
with
\[ V_C(n, n', m, m') = \int dr_e \int dr_h \phi'^*_e(r_e) \phi_h(n', m') \times \int dr_e \phi'^*_e(r_e) \phi_h(n, m) \phi_h(r_h). \] (19)

We solve the eigen problem by diagonalizing the Hamiltonian matrix. We still need to set a cutoff for \( n \) and \( m \) when the binding energy of the ground state interface exciton are convergent. As shown in Fig. 10, where the parameters are chosen as \( R = 30a, L_{SC} = 60a, W_{SC} = 36\sqrt{3}a \), clearly the binding energy converges quickly along the quantum numbers, especially along \( n \). In the following calculation, we set the cutoff \( n_{\text{cutoff}} = m_{\text{cutoff}} = 15 \).

The numerical results of the binding energy versus the band offset \( V_0 \) and the size of the quantum dot \( R \) is depicted in Fig. 11. The binding energy monotonically decreases as the band offset increases as shown in Fig. 11(a), which results from the stronger quantum confinement. However, as shown in Fig. 11(b), the behavior of the binding energy versus \( R \) has a maximum value due to the competition between the quantum confinement and the Coulomb interaction. When \( R < 30a \), basically the ground state of the electron and hole dominates the wavefunction, and thus when \( R \) increases to decrease the quantum confinement, the binding energy of exciton increases. While when \( R > 30a \), the excited states of electron and hole start to appear in the wavefunction, which results in the decrement of binding energy. So if such decrement is greater than the increment of binding energy resulting from quantum confinement, the binding energy of exciton becomes to decrease as the size of quantum dot \( R \) increases. Therefore, there is a maximum binding energy for an optimal \( R \).

Such competition can be also demonstrated in the reduced wavefunction of the electron and hole as shown in Fig. 12. The left and right panels respectively show the reduced wavefunction of electron and hole for increasing size of the quantum dot from the top to bottom. All reduced wavefunctions have the three-fold rotation symmetry inheriting from the symmetry of the regular triangular shape of the closed interface. For a small quantum dot such as \( R = 21a \), the electron is strongly confined in the quantum dot and the hole wavefunction spreads over the entire quantum dot. While for a large quantum dot such as \( R = 45a \), the wavefunctions of electron and hole only spread over the vicinity of the edges of the closed interface. Without the interplay between the wavefunctions at different edges, the closed triangular interface will degrade to 1D interface case. For a large quantum dot, the binding energy is about 140meV, which is consistent with the former 1D interface calculation as shown in Fig. 7(b).

We can imagine that when the size of the quantum dot is much larger than the effective radius of the 1D interface exciton, which is about 5nm according to the previous calculation, the interface exciton actually is split into three identical parts locating at the edges of triangular quantum dot and each part is analogy to the quasi-1D exciton. When the size of the triangular quantum dot is decreased, the three parts have considerable overlap at the corners of the triangular quantum dot when the size of the triangular quantum dot decreases. In this sense, an effective Hamiltonian is introduced to describe such three-fold rotational symmetric system as

\[ H_{\text{eff}} = \begin{bmatrix} E_0 & t e^{i\theta} & t e^{-i\theta} \\ t e^{-i\theta} & E_0 & t e^{i\theta} \\ t e^{i\theta} & t e^{-i\theta} & E_0 \end{bmatrix} \] (20)

with bases \( \{ |\Phi \rangle, C_3 |\Phi \rangle, C_3^2 |\Phi \rangle \} \). Here, \( |\Phi \rangle \) is the wavefunction of 1D interface exciton at one edge, \( C_3 \) and \( C_3^2 \) are rotation operators of three-fold rotational group, \( E_0 \) is the binding energy and \( t e^{i\theta} \) represents the transition between wavefunctions of 1D interface exciton at different edges. In order to satisfy the three-fold rotation symmetry, the phase factor can only be \( \theta = 0, \frac{2\pi}{3} \) or
In addition, the phase factors for the opposite valley should be opposite according to the time-reversal symmetry. Both binding energy $E_0$ and transition coefficient $t$ are determined by the numerical calculation based on the excitonic lattice model.

By diagonalizing the effective Hamiltonian, the lowest three excitonic states can be found as

\[
|\phi_1\rangle = \frac{1}{\sqrt{3}} \left( e^{i\pi/3} |\Phi\rangle + e^{i2\pi} C_3 |\Phi\rangle + e^{i4\pi} C_3^2 |\Phi\rangle \right),
\]

\[
|\phi_2\rangle = \frac{1}{\sqrt{3}} \left( e^{i7\pi/6} C_3 |\Phi\rangle + e^{i\pi} C_3^2 |\Phi\rangle \right),
\]

\[
|\phi_3\rangle = \frac{1}{\sqrt{3}} \left( e^{i7\pi/6} |\Phi\rangle + C_3 |\Phi\rangle + C_3^2 |\Phi\rangle \right),
\]

with corresponding eigen-energies $E_i = E_0 + 2t \cos \left( \frac{2\pi}{3} - \theta \right)$, for $\theta = \pm 1$ as an example. The energy level scheme is depicted in Fig. 13(a), where obviously $|\phi_2\rangle$ and $|\phi_3\rangle$ are degenerate states. More interest fact is that there is a transition when the absolute value of the transition coefficient $t$ varies from negative value to a positive one. When $t < 0$, the $|\phi_1\rangle$ is ground state. In contrast, when $t > 0$, the degenerate states $|\phi_2\rangle$ and $|\phi_3\rangle$ become ground states.

Such transition is depicted in the Fig. 13(b). The numerical calculation based on the lattice model show the transition occurs when the size of the triangular quantum dot is about 12.5nm (Fig 13(b)). The transition coefficient $t$ can also be parameterized by the numerical calculation, which is shown in Fig 13(c). The absolute value of transition coefficient $t$ strongly depends on the overlap of the quasi-1D excitonic wavefunctions at the corners of the triangular quantum dot. For a small quantum dot, the electron confined in the quantum dot and thus the overlap of the electron part supplies a relatively large attractive Coulomb interaction to overcome the kinetic energy. Therefore $t$ has negative value. In contrast for a large quantum dot, both the electron and hole spread over the vicinity of the edges of the quantum dot, and thus the overlap of the electron and hole are greatly decreased. In this sense, the Coulomb interaction part becomes smaller than the kinetic part resulting in positive $t$.

Among the above three excitons, only one state is bright exciton and the other two states are dark excitons when pumping them with right ($\sigma_+$) or left ($\sigma_-$) circularly polarized light. The optical transition matrix elements for the $i$th states are proportional to $\langle \phi_i | \mathbf{P}_\pm | \text{vac} \rangle$, where $\mathbf{P}_\pm$ are the dipole moments corresponding to the $\sigma_+$ or $\sigma_-$ circularly polarized light and $| \text{vac} \rangle$ denotes the initial states with full valence bands and empty conduction bands. Since under the three-fold rotation the transformations of the dipole moments are

\[
C_3^{-1} \mathbf{P}_\pm C_3 = e^{\mp \frac{2\pi i}{3}} \mathbf{P}_\pm, (C_3^3)^{-1} P_\pm C_3^2 = e^{\pm \frac{2\pi i}{3}} P_\pm, \tag{24}
\]

only the exciton states with appropriate phase factors of coefficients have nonzero optical transition matrix elements and thus are bright excitons. In this sense, the $\sigma_-$ circularly polarized light can pump the $|\phi_1\rangle$ in the $\tau = -1$ valley, and $\sigma_+$ circularly polarized light can pump the $|\phi_2\rangle$ in the $\tau = +1$ valley. The corresponding optical selection rule is shown in the Fig. 13(a).

B. Numerical results of closed triangular sharp interface with valley index

In the above discussion, only the interface excitons in the same valley are discussed. However, due to the geometry of the closed triangular sharp interface, the inter-valley couplings are inevitable which eventually couple interface excitons in the opposite valleys. We start from the effective Hamiltonian both including the three-fold rotational symmetric excitons and the inter-valley couplings as

\[
H_{\text{eff}}^{\text{inter}} = \begin{bmatrix}
E_0 & t e^{i\theta_+} & t e^{-i\theta_+} & p & q & q \\
t e^{-i\theta_+} & E_0 & t e^{i\theta_+} & q & p & q \\
t e^{i\theta_-} & t e^{-i\theta_-} & E_0 & q & q & p \\
p & q & q & E_0 & t e^{i\theta_-} & t e^{-i\theta_-} \\
q & p & q & t e^{-i\theta_-} & E_0 & t e^{i\theta_-} \\
q & q & p & t e^{i\theta_-} & t e^{-i\theta_-} & E_0
\end{bmatrix}, \tag{25}
\]

where the bases in the real space $(i = 1, 2, 3)$ are

\[
\langle \textbf{r}_\text{e}, \textbf{r}_\text{h}|\phi_i, \tau \rangle \approx \exp \left(i\tau \mathbf{K} \cdot (\textbf{r}_\text{e} - \textbf{r}_\text{h}) \right) \langle \textbf{r}_\text{e}, \textbf{r}_\text{h}|\phi_i \rangle \times u_\mathbf{K} (\textbf{r}_\text{e}, \textbf{r}_\text{h}) \tag{26}
\]
and $\tau = \pm 1$ is the valley index. The $|\phi_i\rangle$ are the envelopes of the excitonic states without considering the valley index which are defined in Eq. 123, and $u_K(r_x,r_y)$ is the periodic parts of the Bloch wavefunctions. We adopted the assumption that $u_{K+q}(r_x,r_y) \approx u_K(r_x,r_y)$ for the sake of simplicity. Here, $E_0 = \langle \phi_i, \tau | H | \phi_i, \tau \rangle$ is the binding energy of the 1D interface exciton, $t = \langle \phi_i, \tau | H | C_3 \phi_i, \tau \rangle$ is the intra-valley inter-edge hoppings, $p = \langle \phi_i, \tau | H | \phi_i, \bar{\tau} \rangle$ is the inter-valley intra-edge hoppings, and $q = \langle \phi_i, \tau | H C_3 | \phi_i, \bar{\tau} \rangle$ is the inter-valley inter-edge hoppings. Here, the original Hamiltonian $H$ is introduced in Eq. 1. In order to make sure that the ground state of the interface exciton still inherit the same optical selection rule, which means that the $\sigma_+ (\sigma_-)$ circularly polarized light only pump the ground states of the excitons in the $\tau = +1$ ($\tau = -1$) valley, the phase factors for both valleys are fixed as $\theta_+ = -\theta_- = -2\pi/3$.

Since the inter-valley terms $p(q)$ are at least one order smaller than the corresponding intra-valley terms $E_0(t)$ due to the large momentum difference, and the inter-valley terms results from the wavefunction overlap at the corners which obviously become smaller when the size of the quantum dot increases, the magnitudes of the parameters have the following relations $E_0 \gg t \sim p \gg q$. In the following calculation, we ignore the inter-valley inter-edge hoppings $q$. The numerical results of the inter-valley intra-edge hopping $p$ versus the size of quantum dot $R$ is shown in Fig. 14. The magnitude of $p$ almost decrease exponentially as the size of the quantum dot increases. For the large quantum dot, the value matches the previous inter-valley coupling results shown in Fig. 5 because the the interface exciton degrades to 1D interface exciton without the the wavefunction overlap at the corners. For small quantum dot such as $R < 5 \text{nm}$ it can reach to several meV.
Although $q$ is small in comparison with $t$, it can still couples the interface excitons in opposite valleys. By diagonalizing the effective Hamiltonian $H_{\text{eff}}$ in Eq. (25), we can obtain the lowest six interface excitonic states as

\begin{align}
|\Phi_1\rangle &= \cos \psi |\phi_2, \tau\rangle + \sin \psi |\phi_3, \tau\rangle, \\
|\Phi_2\rangle &= \cos \psi |\phi_1, \tau\rangle + \sin \psi |\phi_3, \tau\rangle, \\
|\Phi_3\rangle &= - \sin \psi |\phi_2, \tau\rangle + \cos \psi |\phi_3, \tau\rangle, \\
|\Phi_4\rangle &= - \sin \psi |\phi_1, \tau\rangle + \cos \psi |\phi_3, \tau\rangle, \\
|\Phi_5\rangle &= \frac{1}{\sqrt{2}} (|\phi_1, \tau\rangle - |\phi_2, \tau\rangle), \\
|\Phi_6\rangle &= \frac{1}{\sqrt{2}} (|\phi_1, \tau\rangle + |\phi_2, \tau\rangle),
\end{align}

with $\tan \psi = 2p/(3t)$ and the corresponding energies

\begin{align}
E_1 &= E_2 = E_0 + \frac{1}{2} \left( t + \sqrt{t^2 + 4p^2} \right), \\
E_3 &= E_4 = E_0 + \frac{1}{2} \left( t - \sqrt{t^2 + 4p^2} \right), \\
E_5 &= E_0 - t - p \\
E_6 &= E_0 - t + p.
\end{align}

Since the interval coupling $p$ actually couples one bright exciton and one dark exciton such as $|\phi_2, \tau\rangle$ and $|\phi_3, \tau\rangle$, both the $|\Phi_1\rangle$ and $|\Phi_3\rangle$ become bright but with different coupling strength with the same right circularly polarized light field. The energy level and the complete optical selection rules are shown in the Fig. 14(b). According to the orthogonality of the periodic parts of the Bloch wavefunctions, the envelope wavefunction of the six excitonic states with the valley index are analogous to the wavefunction without the valley index shown in Fig. 12.

V. CONCLUSION

In this paper, we theoretically study the interface exciton states at various lateral heterojunctions of monolayer semiconductors including single, double and closed triangular interfaces. When taking the distance dependent screening of Coulomb interaction into consideration, we numerically study the physical observables of type II interface exciton including the binding energy, effective radius between the electron and hole and optical dipole. Usually, such problem is quite difficult to be numerically solved by ab initio calculations. We adopted two different approaches to calculate excitons. One approach bases on a real-space tight binding model, and the other approach considers the perturbation expansion in a hydrogen-like basis in an effective mass model. The numerical study shows that even when the electron-hole separation is much larger compare to the 2D excitons in TMDs, type II interface exciton still has strong binding energy. When the effective radius between the electron and the hole is up to four times of the Bohr radius of 2D excitons, the binding energy remains 1/2 that of 2D excitons. This can be interpreted by the weaker screening of Coulomb interaction as the electron-hole spatial separation increases. Large energy separation between interface exciton and 2D excitons for band offset above 0.2 eV ensures that such 1D interface excitons are stable ones. Due to the spatial indirect nature of type II interface exciton, exciton radius increases while optical transition dipole decreases as band offset increases. Still, the optical dipole is comparable to that of 2D excitons at moderate band offset of 100meV or below. Inter-valley coupling that arises from electron-hole exchange is also studied, which may leads to the longitudinal-transverse splitting with the interface breaking the rotational symmetry. The lateral heterojunctions with closed triangular interface is also studied, which realize the 0D quantum dot confinement of exciton. The numerical study shows that the energy level schemes and valley optical selection rules of the exciton in quantum dot depends on the size of the quantum dot. Together with valley index, there are more exciton states in a single quantum dot which can be used to carry information. With its unique nature of having one carrier confined within the triangle by the band offset and the other carrier bounded to the proximity exterior of the triangle by the strong Coulomb, it is possible to realize the strong excitonic coupling between the neighbouring quantum dots for mediating controlled interplayer between spins at different dots. In this sense, our investigation may facilitate the quantum information procession based on the 2D monolayer semiconductors.

Appendix A: Generalized Born-Oppenheimer approximation

The regular Born-Oppenheimer approximation only consider the lowest order of the ratio between the reduced mass and the total mass $\kappa = \mu/M$ which corresponds to the relative motion and the center-of-mass motion, respectively. The total wavefunction is expressed as a product of the relative and center-of-mass parts when the adiabatic condition is satisfied. However, in the problem of our interface exciton there exist nonadiabatic processes and higher order terms in the ratio $\kappa$ should be taken into consideration. In this sense, we present the generalized Born-Oppenheimer approximation and the corresponding second-order perturbation theory here.

The Schrödinger equation satisfied by one center-of-mass motion and multiple relative motions is $H \Phi (\{r\}) = E \Phi (\{r\})$ with Hamiltonian

\[ H = -\frac{\hbar^2}{2M} \nabla_\mathbf{R}^2 + H (\{r\}) + V (\{r\}), \]

where the first term is the kinetic energy of the center-of-mass motion, the second term describes the energy of the multiple relative motions $\{r\} = r_1, r_2, \ldots$ and the third term is coupling between the center-of-mass motion and the relative motions. For arbitrary center-of-mass space coordinate $\mathbf{R}$, the eigenvalue equation

\[ [H (\{r\}) + V (\{r\})] \Theta_k (\mathbf{R}, \{r\}) = E_k (\mathbf{R}) \Theta_k (\mathbf{R}, \{r\}) \]

can be solved to obtain the corresponding eigenvalues $E_k (\mathbf{R})$ and eigenfunctions $\Theta_k (\mathbf{R}, \{r\})$. Since these bases
\{\Theta_k (R, \{r\})\} are orthogonal and complete, one expands $\Phi (R, \{r\})$ in bases $\{\Theta_k (R, \{r\})\}$ as
\[
\Phi (R, \{r\}) = \sum_{k=1}^{\infty} \Psi_k (R) \Theta_k (R, \{r\}). \tag{A3}
\]

Obviously, this expanded wavefunction satisfies the original Schrödinger equation as well. The straightforward derivation gives the set of the effective motion equations of the coefficients $\Psi_k (R)$ as
\[
H_k (R) \Psi_k (R) + \sum_{k'} H_{k,k'} (R) \Psi_{k'} (R) = E \Psi_k (R), \tag{A4}
\]
where
\[
H_k (R) = H_0^k (R) + H^1_{k,k} (R), \tag{A5}
\]
\[
H_0^k (R) = -\frac{\hbar^2}{2M} (\nabla_R - i A_{k,k} (R))^2 + E_k (R), \tag{A6}
\]
\[
H^1_{k,k} (R) = \sum_{k' \neq k} \frac{\hbar^2}{2M} A_{k,k'} (R) \cdot A_{k',k} (R), \tag{A7}
\]
\[
H^1_{k,k'} (R) = i \frac{\hbar^2}{M} \sum_{k' \neq k} A_{k,k'} (R) \cdot \nabla_R \Psi_k (R) + i \frac{\hbar^2}{2M} \sum_{k' \neq k} \int dr [\nabla_R \cdot A_{k,k'} (R)] \Psi_{k'} (R), \tag{A8}
\]
and the Berry connections are defined as $A_{k,q} (R) \equiv i \int dr \Theta^*_k (R, \{r\}) \nabla_R \Theta_q (R, \{r\}).$ So far the effective motion equations are rigorous without any approximation. Here, the $H_0^k (R)$ and $H^1_{k,k} (R)$ are adiabatic terms because they only involve the $k$-th energy-level. However, $H^1_{k,k'} (R)$ involves the transitions between different energy-levels introducing the non-adiabatic processes.

To obtain the explicit expression for $A_{k,q} (R)$, differentiating the eigenvalue equation as Eq. (A2) leads to
\[
[H (\{r\}) + V (R, \{r\}) - E_p (R)] \nabla_R \Theta_p (R, \{r\}) = [\nabla_R E_p (R) - \nabla_R V (R, \{r\})] \Theta_p (R, \{r\}). \tag{A9}
\]

Multiplying $\Theta^*_k (R, \{r\})$ to both sides of the above equation and integrating over all relative space coordinates $\{r\}$ gives
\[
[E_k (R) - E_p (R)] \int dr \Theta^*_k (R, \{r\}) \nabla_R \Theta_p (R, \{r\}) = -\int dr \Theta^*_k (R, \{r\}) \nabla_R V (R, \{r\}) \Theta_p (R, \{r\}), \tag{A10}
\]

According to the definition of the Berry connection, the explicit expression of the absolute value of the Berry connections is
\[
|A_{k,p} (R)| = \left| \int dr \Theta^*_k (R, \{r\}) [\nabla_R V (R, \{r\})] \Theta_p (R, \{r\}) \right| \frac{E_k (R) - E_p (R)}{E_k (R) - E_p (R)}. \tag{A11}
\]

It is clear that the $H^1_{k} (R)$ and $H^1_{k,k} (R)$ are regarded as the perturbations when the partial derivation of the coupling $\nabla_R V (R, \{r\})$ is much smaller than the energy level spacing $|E_k (R) - E_p (R)|$. The order of the perturbations can be characterized by the number of the Berry connections. In this sense $H^1_{k} (R)$ is the second order perturbation and $H^1_{k,k} (R)$ contains the first order and the second order perturbations.

The Berry connections $A_{k,k} (R)$ in the $H^0_{k} (R)$ actually plays the role of a gauge field. It is important to indicate that for Eq. (A2) the phase of the bases $\{\Theta_k (R, \{r\})\}$ are not fixed because the the eigenvalue equation is unchanged under the transformation $\Theta_k (R, \{r\}) = \Theta_k (R, \{r\}) \exp [-i\theta (R)].$ However, the Berry connections of the transformed bases $A_{k,q} (R) \equiv i \int dr \Theta^*_k (R, \{r\}) \nabla_R \Theta_q (R, \{r\})$ accordingly become
\[
A_{k,q} (R) = \begin{cases} A_{k,q} (R), & k \neq q, \\ A_{k,k} (R) + \nabla_R \theta (R), & k = q. \end{cases} \tag{A12}
\]

Therefore $A_{k,k} (R)$ depends on the choice of the phase factor $\theta (R)$ and thus we can not decide its perturbation order. This is actually the $U(1)$ gauge transformation and the physical observations are not influenced by the specific choice of the phase factor. In our problem of the interface exciton, this induced gauge field can be cancelled out by choosing the proper bases as $A_{k,k} (R) = 0$ for any $k$.

To apply the standard perturbation theory, we rewrite Eq. (A3) in a matrix form as
\[
\Phi (R, \{r\}) = \Psi (R)^T \cdot \Theta (R, \{r\}), \tag{A13}
\]
where the coefficient vector $\Psi (R)$ and the base vector $\Theta (R, r)$ are
\[
\Psi (R) = \begin{bmatrix} \Psi_1 (R) \\ \Psi_2 (R) \\ \vdots \end{bmatrix}, \quad \Theta (R, r) = \begin{bmatrix} \Theta_1 (R, \{r\}) \\ \Theta_2 (R, \{r\}) \\ \vdots \end{bmatrix}, \tag{A14}
\]

And the Eqs. (A4) are rewritten as
\[
H^0 (R) + H^1 (R) \Psi (R) = E \Psi (R) \quad \text{with corresponding Hamiltonians in the matrix form as}
\]
\[
H^0 (R) = \begin{bmatrix} H_1^0 (R) & 0 & \cdots \\ 0 & H_2^0 (R) & \cdots \\ \vdots & \vdots & \ddots \end{bmatrix}, \tag{A15}
\]
and
\[
H^1 (R) = \begin{bmatrix} H_1^1 (R) & H_{12}^1 (R) & \cdots \\ H_{21}^1 (R) & H_2^1 (R) & \cdots \\ \vdots & \vdots & \ddots \end{bmatrix}. \tag{A16}
\]

Here all the first order and the second order perturbations are included into $H^1 (R)$. By applying the standard perturbation theory, the second order eigen-energy
and wavefunction respectively as $E_p = E_p^0 + E_p^1$ and
$\Psi_p (R) = \Psi_p^0 (R) + \Psi_p^1 (R)$, where

$$E_p^1 = \sum_{k \neq p} \frac{\hbar^2}{2M} \int dR \Psi_p^0 (R) \mathbf{A}_{p,k} (R) \cdot \mathbf{A}_{k,p} (R) \Psi_p^0 (R),$$

(A17)

$$\Phi_k^1 (R) = \sum_{k \neq p} \frac{\int dR' \Psi_k^{(0)*} (R') H_{kp}^1 (R') \Psi_p^0 (R')}{E_p^0 - E_k^0} \Psi_k^0 (R),$$

(A18)

the zero-th order eigen-energy and wavefunction are determined by $H^0 (R)$ as $H^0 (R) \Psi_p^0 (R) = E_p^0 \Psi_p^0 (R)$ and
$H_{kp}^1 (R')$ is the element of perturbation Hamiltonian $H^1 (R)$.

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